Testing Methods for Fertilizers
(2018)

Incorporated Administrative Agency
Food and Agricultural Materials Inspection Center
Introduction

“The Official Methods of Analysis of Fertilizers” stipulated by the Ministry of Agriculture, Forestry and Fisheries is the only analysis method to assess main components and harmful components in fertilizers in Japan and contributes to maintaining quality and securing safety of fertilizers. However, no new revision has been issued since “The Official Methods of Analysis of Fertilizers 1992” was issued. Some quarters such as fertilizer manufacturers and inspection instruction agencies have requested a revised edition of “The Official Methods of Analysis of Fertilizers” since new kinds of fertilizers and its new components were added into the public standard, and analysis instruments and technologies have progressed during the period.

Incorporated Administrative Agency, Food and Agricultural Materials Inspection Center (FAMIC) has revised the Official Methods of Analysis of Fertilizers by introducing the analysis conditions and the analysis methods, etc. which meet the latest situation. Additionally, FAMIC tried to study how to introduce the analysis methods or the new analysis instruments to cope with new effective components or harmful components and new fertilizers which are not documented in the Official Methods of Analysis of Fertilizers, and established new testing methods. At the same time, FAMIC conducted a validity test according to the requirements of ISO/IEC 17025 and opened the results and new testing methods which were discussed and approved by “the Technical Committee for Fertilizers etc.” including outside experts on FAMIC’s web site in 2008 as “The Testing Methods for Fertilizers 2008”. Since then, the contents have been annually added and updated. In this year, High Performance Liquid Chromatograph, etc. for uric acid within fluid fertilizers which have been studied anew in FY2017 were merged, and the “Testing Methods for Fertilizes 2018” was opened on FAMIC’s web site.

The “Testing Methods for Fertilizes” uses reagents and instruments which are stipulated in JIS standards, etc. and its validity of the testing method is checked by referencing IUPAC protocols and it is listed as an analysis method whose validity is checked in “The Management Handbook of Heavy Metal in Sludge Fertilizers” published by the Ministry of Agriculture, Forestry and Fisheries in August 2010. Therefore, FAMIC would like people engaged in the quality control and analysis of fertilizes to use this as a practical document.

June, 2018
Incorporated Administrative Agency
Food and Agricultural Materials Inspection Center
General Director
Makoto Kimura
The members of The Technical Committee
(Without honorifics, as of March, 2018)

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1. General Rule

1.1 Common Items

(1) Applicable range

The Testing Methods for Fertilizers stipulate the official method of analysis of fertilizers and fertilizer materials. The type of samples in the tests is shown in the summary of respective test items.

(2) General matters in common, procedures and terms

2.1 Terms related to laws and ordinances

a) Main components or major components: The main components or major components in fertilizers in Table 1 are stipulated as components to be calculated by a public notice of the Ministry of Agriculture, Forestry and Fisheries.

<table>
<thead>
<tr>
<th>Main component or major component</th>
<th>Component to be calculated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate</td>
<td>Phosphorus pentoxide (P$_2$O$_5$)</td>
</tr>
<tr>
<td>Potassium</td>
<td>Potassium oxide (K$_2$O)</td>
</tr>
<tr>
<td>Silicate</td>
<td>Silicon dioxide (SiO$_2$)</td>
</tr>
<tr>
<td>Magnesium</td>
<td>Magnesium oxide (MgO)</td>
</tr>
<tr>
<td>Manganese</td>
<td>Manganese oxide (MnO)</td>
</tr>
<tr>
<td>Boron</td>
<td>Diboron trioxide (B$_2$O$_3$)</td>
</tr>
<tr>
<td>Sulfur content</td>
<td>Sulfur trioxide (SO$_3$)</td>
</tr>
<tr>
<td>Lime</td>
<td>Calcium oxide (CaO)</td>
</tr>
</tbody>
</table>

(2.2) General matters and terms that cite Japanese Industrial Standards (JIS)

a) General notices: General matters common to analysis are according to JIS K 0050.

b) Definition: The definitions of major terms used in the Testing Methods for Fertilizers are according to JIS K 0211, JIS K 0214, JIS K 0215, JIS Z 8101-1, JIS Z 8101-2 or JIS Z 8101-3.

c) Laboratory sample: A sample transferred to a laboratory. Laboratory sample as specified in JIS K 0211.

d) Test sample: A sample obtained from a laboratory sample after pretreatment such as grinding. Test sample as specified in JIS K 0211.

e) Analytical sample: A sample measured from a laboratory sample or a test sample and to be used in one test. Test sample or analytical sample as specified in JIS K 0211.

f) Sample: A sample in the testing methods indicates e) a laboratory sample, f) a test sample or g) an analytical sample.

g) Rounding numbers: Methods of rounding the numbers are according to JIS Z 8401.

h) Absorptiometric analysis: General rules for absorptiometric analysis are according to JIS K 0115.

i) Atomic absorption spectrometry: Atomic absorption spectrometry includes flame atomic absorption spectrometry, electrically heated atomic absorption spectrometry (hereinafter

referred to as “electrically heated atomic absorption spectrometry”) and other atomic absorption spectrometry. General matters common to these are according to JIS K 0121.

j) **Gas chromatography**: General matters common to gas chromatography are according to JIS K 0114.

k) **Gas chromatography/Mass spectrometry**: General matters common to Gas chromatography/Mass spectrometry are according to JIS K 0123.

l) **Electrical conductivity measuring method**: General matters common to electrical conductivity measuring methods are according to JIS K 0130.

m) **Test sieving**: General matters common to test sieving are according to JIS Z 8815.

n) **High Performance Liquid Chromatography**: General matters common to High Performance Liquid Chromatography are according to JIS K 0124.

o) **High Performance Liquid Chromatography/Mass Spectrometry**: General matters common to High Performance Liquid Chromatography/mass spectrometry are according to JIS K 0136.

p) **ICP Optical Emission Spectrometry**: General matters common to ICP Optical Emission Spectrometry are according to JIS K 0116.

q) **ICP Mass Spectrometry**: General matters common to ICP Mass Spectrometry are according to JIS K 0133.

r) **Ion Chromatography**: General matters common to Ion Chromatography are according to JIS K 0127.

(2.3) **Description methods, procedures and terms in testing methods for fertilizers.**

a) **Reagent name**: Unless otherwise specified, conform to the names by the chemical nomenclature established by the Chemical Society of Japan [in accordance with the International Union of Pure and Applicable Chemistry (IUPAC) nomenclature of inorganic chemistry and nomenclature of organic chemistry] and the names of JIS reagents.

b) **Organic matters**: Fertilizers such as Organic fertilizers, sludge fertilizers and compost and fertilizer materials. However, organic compounds such as urea and urea compounds are excluded.

c) **Actual article**: A laboratory sample in original state.

d) **Drying matter**: The matter which remains after drying the actual article.

e) **Notes, comments, figures, tables and formulae**: Serial numbers for each test item should be recorded in notes, comments, figures, tables and formulae.

f) **Dilution of solution**: “Transfer accurately a predetermined amount (to a vessel)” means the procedure to measure any volume of solution with a measuring instrument specified in JIS K 0050 (into a vessel).

   Also, “dilute accurately a predetermined amount (with solvent or solution)” means the procedure to measure any volume of solution with a measuring instrument specified in JIS K 0050 into a volumetric flask of arbitrary volume and fill up to the marked line (with solvent or solution) (1).

g) **Description of mixture solution**: Mixture solutions are described as shown in 1) - 4).

1) **Reagent + reagent**: Describe as reagent name 1–reagent name 2 (V₁ + V₂). In this case, V₁ volume of reagent name 1 is mixed with V₂ volume of reagent name 2.

   Example: acetonitrile–water (1+1), hexane–ethyl acetate (2+1), methanol–buffer solution (3+1)

2) **Reagent + water**: Describe as reagent name 1 (V₁+V₂). In the case of reagents described in Table 1 in JIS K 0050, it means V₁ volume of reagent name 1 is diluted by mixing with V₂ volume of water.

   Example: hydrochloric acid (1+1), sulfuric acid (1+2), ammonia solution (1+3)
3) **Solution + reagent:** Describe as solution name a (concentration) - reagent name b \([V_1 + V_2]\). In this case, it means \(V_1\) volume of solution name a of a certain concentration is mixed with \(V_2\) volume of reagent name b.

Example: sodium hydroxide solution (4 g/L)–methanol [1+4]

4) **Diluted reagent + reagent:** Describe as reagents name a \((V_1 + V_2)\) - reagent name b \([V_3 + V_4]\). In this case, \(V_3\) volume of the solution in which \(V_1\) volume of reagent name a described in Table 1 in JIS K 0050 diluted by mixing with \(V_2\) volume of water, is mixed with \(V_4\) volume of reagent name b.

Example: hydrochloric acid (1+100)–methanol [2+3]

h) **Preparation of a calibration curve:** “Transfer A mL - B mL of the standard solution to volumetric flasks step-by-step.” means the procedure to transfer a volume of 4 - 6 steps (2) in the range from A mL to B mL of the standard solution to respective volumetric flasks step-by-step.

Prepare a calibration curve every time a test is conducted. Also, when the same test item is measured under the same conditions for multiple samples continuously, measure the standard solution at regular intervals to check the indicated value.

i) **Washing of apparatus:** Wash containers with a detergent and tap water before usage and wash sufficiently with water of A2 specified in JIS K 0577 or water that is confirmed not to affect a quantification value. In case of sampling a sample to test a metallic element and organic materials, after previous washing, dip with nitric acid (1+9) or hydrochloric acid (1+9) as appropriate and further wash sufficiently with water of A2, A3 or A4.

j) **Handling of reagents and liquid waste, etc.:** Handle with care and in compliance with relevant laws and regulations. When treatment methods are specified in respective test items, comply with the methods.

k) **Referential matters related to the validity of testing methods:** Information related to the validity of respective testing methods such as quantification range (minimum limits of quantification, etc.), mean recovery, repeatability, intermediate precision and reproducibility is described in a Comment, etc. Note that the numerical values such as Minimum Limit of Quantification, etc. are not standards to be targeted but examples.

**Note** (1) When the dilution factor is large, accuracy should be secured by procedures such as repeating the dilution procedure.

(2) Set according to the specification and operation method of the measurement instrument used. There is no need to include the minimum and the maximum values of the calibration curb range described in the Testing Methods for Fertilizers.

3) **Water**

a) **Water:** Water used in the Testing Methods for Fertilizers herein is water of A2 specified in JIS K 0557 or water that is confirmed not to affect a quantitation value. However, when otherwise specified in respective test items, use the specified water.

4) **Reagents**

a) **Reagents:** When the reagent is JIS-specified, use one of highest quality among those marked with the JIS symbol; when none of the reagent is marked with the JIS symbol, use one of quality that will not cause a problem in the test. Use reference materials for volumetric analysis specified in JIS K 8005 for the standardization of titration solutions.

b) **Reference materials:** The preparation of standard solutions or standardization of titration solutions using reference materials 1) - 2) below other than materials specified in respective testing items is possible.
1) **Reference materials provided by National Metrology Institute:** Reference materials traceable to International System of Unit (SI) provided by National Metrology Institute (NMI: National Institute of Advanced Industrial Science and Technology NMII, NIST, BAM, etc.) which signed CIPM MRA (Global Mutual Recognition Arrangement based on the Meter Convention)

2) **Reference materials for volumetric analysis:** Reference materials for volumetric analysis specified in JIS K 8005.

c) **Standard Solutions:** In the cases of specifying in the comment in respective testing items, the preparation of standard solutions for a calibration curve using the solution which is traceable to the National Metrology of 1) - 3) below other than specified in respective testing items is possible. However, use standard solutions which do not cause a problem in the test with the kinds and concentration of chemical compounds or added acid used. In addition, in (2.1) a) Main components or major components, calculate main components or major components using conversion factors specified in the comment of respective items.

1) **Standard Solutions provided by National Metrology Institute:** Standard solutions traceable to International System of Unit provided by National Metrology Institute (NMI: National Institute of Advanced Industrial Science and Technology NMII, NIST, BAM, etc.) which signed CIPM MRA.

2) **JCSS (Japan Calibration Service System) Standard Solutions:** Standard solutions prepared by JCSS (Japan Calibration Service System) the registered provider for Chemical Analysis, Atomic Absorption Spectrometry, ICP or Ion Chromatography traceable to specific reference materials based on Measurement Act Article 134. In addition, it is recommended to use standard solutions which indicate the uncertainty of concentration and factors.

3) **Standard Solutions traceable to National Metrology:** Standard Solutions traceable to National Metrology (National Institute of Advanced Industrial Science and Technology NMII, NIST, BAM reference materials, etc. traceable to International System of Unit) provide by National Metrology Institutes which signed CIPM MRA, but at the same time they are standard solutions for Chemical Analysis, Atomic Absorption Spectrometry, ICP or Ion Chromatography prepared by the providers who obtained the certification of ISO Guide 34(JIS Q 0034: General requirements for the competence of reference material producers).

In addition, it is recommended to use standard solutions which indicate the uncertainty of concentration and factors.

d) **Titrant:** A Titrant described in 1) is usable if it is specified in the comment of a testing item. In addition, the titrant may be diluted to a predetermined concentration as necessary. In this case, however, dilution treatment should be conducted when it is used and factors of the titrant before dilution should be applied.

1) **Titrant conforming to ISO/IEC 17025:** A Titrant which is prepared and standardized and whose factors are calculated by a laboratory which obtained an accreditation (accredited range: JIS K 8001 JA.5 solutions for titration) based on ISO/IEC 17205. In addition, it is recommended to use titrant which indicates the uncertainty of concentration and factors.

e) **Concentration of reagent solution:** Unless otherwise specified, the mass concentration is expressed as g/L or mg/L, while the molar concentration is expressed as mol/L or mmol/L. The concentration of the standard solution is expressed as the mass in 1 mL (mg/mL, µg/mL or ng/mL) except for the ion-selective electrode method.

f) **Concentration in parenthesis shown after the name of reagent solution:** It indicates that the solution is about that concentration except the standard solution. For example, sodium hydroxide solution (0.1 mol/L) means that it is about 0.1 mol/L sodium hydroxide solution. Also, the concentration shown in front of the name of solution means that it is the accurate concentration. However, the concentration is generally expressed as a round figure; calculate the factor separately.
(5) Apparatus

a) Glass apparatus: Unless otherwise specified, use glass apparatus specified in JIS R 3503 and JIS R 3505. Also, when a heating procedure is involved, use borosilicate glass-1 specified in JIS R 3503.

b) Non-glass apparatus: Unless otherwise specified, use plastic apparatus.

c) Desiccants for desiccators: Unless otherwise specified, use silica gel.

d) Porcelain crucibles and porcelain evaporating dishes: Use ones specified in JIS R 1301 and JIS R 1302.

e) Platinum crucibles and platinum evaporating dishes: Use ones specified in JIS H 6201 and JIS H 6202.

f) Filter paper: Use that specified in JIS P 3801. However, the type of filter paper is specified in respective test items.

g) Absorbance measurement (absorptiometric analysis) absorbance cells: Unless otherwise specified, use ones of 10 mm in optical path length.
1.2 Validity check of testing methods

The Testing Methods for Fertilizers are methods which have been discussed and approved by the Technical Committee for Fertilizers etc. or methods in the Official Methods of Analysis of Fertilizers (1992) whose style was rewritten according to the Testing Methods for Fertilizers. The Testing Methods for Fertilizers will be revised by adding, modifying, or deleting testing methods with the approval of the Technical Committee for Fertilizers etc. due to the needs such as progress in analytical techniques and changing social situation.

The procedure for the testing methods validity check is shown in the Appendix of this Testing Methods for Fertilizers. This procedure was made based on 5.4.5 Validation of methods in JIS Q 17025 “General requirements for the competence of testing and calibration laboratories” or 2.4 Tests for validation requirements in “Guidelines for the design and implementation of surveillance and monitoring and for the evaluation and publication of the results” which was issued by the Ministry of Agriculture, Forestry and Fisheries and with reference to the guideline of Codex Alimentarius Commission (CAC), IUPAC protocol and the guideline of AOAC INTERNATIONAL, etc. Validated testing methods are methods which are conducted according to this procedure and confirmed to conform to the standards such as required accuracy (trueness and precision), quantification range (maximum and minimum limits of quantitation) and so on.

In addition, according to the validation levels in Table 1, an individual testing method is classified from Type A to Type E in Table 2.

<table>
<thead>
<tr>
<th>Symbol of Validation, etc.</th>
<th>Validation method, etc.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Def-M (Defining method)</td>
<td>Procedures of a testing method define measurement items. No relation to a validation level.</td>
</tr>
<tr>
<td>Def-C (Defining calculation)</td>
<td>Only calculation methods in a testing method define measurement items. The part of the definition has no relation to a validation level.</td>
</tr>
<tr>
<td>Def-E (Defining extraction)</td>
<td>Only extraction procedures in a testing method define measurement items. The part of the definition has no relation to a validation level.</td>
</tr>
<tr>
<td>HCV (Harmonized collaborative validation)</td>
<td>Evaluation by a collaboration test using samples of no less than 5 concentrations in no less than 8 laboratories with a testing method validation method regarded as an international standard (Guideline of AOAC-International, IUPAC protocol, etc.).</td>
</tr>
<tr>
<td>MLV (Multi laboratory validation)</td>
<td>Evaluation of validation using multiple laboratories, though it does not satisfy HCV criteria.</td>
</tr>
<tr>
<td>SLV (SingleLaboratory validation)</td>
<td>Evaluation in a single laboratory using a testing method validation method regarded as an international standard (IUPAC/ISO/AOAC-International harmonized guideline, etc.).</td>
</tr>
<tr>
<td>RNV (Research non validated)</td>
<td>A testing method that does not carry out the validation by SLV or higher level.</td>
</tr>
</tbody>
</table>
### Table 2  Classification of an individual testing method

<table>
<thead>
<tr>
<th>Symbol of Classification</th>
<th>Validation level</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Type A</strong></td>
<td>A defined testing method</td>
</tr>
<tr>
<td><strong>Type B</strong></td>
<td>A testing method whose evaluation results by HCV and SLV satisfies the requirements of &quot;Appendix: The procedure to validate characteristics of testing methods&quot;.</td>
</tr>
<tr>
<td><strong>Type C</strong></td>
<td>A testing method whose evaluation results by MLV and SLV satisfies the requirements of &quot;Appendix: The procedure to validate characteristics of testing methods&quot;.</td>
</tr>
<tr>
<td><strong>Type D</strong></td>
<td>A testing method whose evaluation results by SLV satisfies the requirements of &quot;Appendix: The procedure to validate characteristics of testing methods&quot;.</td>
</tr>
<tr>
<td><strong>Type E</strong></td>
<td>A testing method that does not carry out the validation by SLV or higher level.</td>
</tr>
</tbody>
</table>
1.3 Procedure of testing methods
1.3.1 Competence evaluation of laboratory

When an individual testing method is used, it is recommended that the following competence evaluation of a laboratory is conducted.

As for the testing methods of Type A, Type B and Type C, conduct a replicate testing with five samples, whose measured component concentration is known and to which certified reference materials or standard solutions are added, to confirm trueness and precision in advance. As for the testing methods of Type D and Type E, conduct a validity check in a single laboratory anew.

In order to ensure the reliability of a series of testing, conduct internal quality control (internal quality assurance control, internal precision control) for each testing by a duplicate testing with samples, whose measured component concentration is known, to confirm trueness and precision.

If it is possible, participate in an external quality assessment (external precision control, competence test) in order to evaluate the consistency with testing results of other laboratories and confirm the evaluation by z score.

1.3.2 Evaluation of test result

The results of a testing method which substitutes for the Testing Methods for Fertilizers can be used if it conforms to criteria required in validation of a testing method. However, in case where the result of the testing method does not agree with the result of the Testing Methods for Fertilizers (1), the testing result of the latter is used to make a final judgement. In addition, if multiple methods are described for a testing component, it is recommended that the result of a testing method for final judgement is adopted in the following order: Type A, Type B, Type C, Type D and Type E.

Note (1) Refer to the separate sheet: “Target of trueness and criteria of precision in respective concentration levels” or the reproducibility of respective testing methods” in order to determine if there is mutual agreement.
2. **Handling of samples**

2.1 **Sampling**

Sampling is according to “2.1 Sampling in the Official Methods of Analysis of Fertilizers (1992)”.

**References**

1) National Institute for Agro-Environmental Sciences, the Ministry of Agriculture, Forestry and Fisheries: The Official Methods of Analysis of Fertilizers 1992, p.4 - 5, Japan Fertilizers Analysis Association, Tokyo (1992)

2.2 Storage of samples

(1) Summary
Put a sample in a container suitable for its characteristics and form and seal it tightly, and then store the sample at room or cold temperature. Care should be taken not to freeze it when it is stored at cold temperature.

(2) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Refrigerator: A refrigerator that can be adjusted to 1 ºC - 8 ºC.
   b) Storage container for a sample: A storage container for a sample should be clean, strong and completely sealed airtight. In particular, in case it contains sludge for raw materials, the container should be made of non-degradable, non-absorbable material. Additionally, it should be airtight, water-proof, vapor-proof and non-corrosive.

(3) Procedure: Conduct storage as shown below.
   a) Store a relatively stable sample in a tightly sealed container to avoid direct sunlight.
   b) Store a sample in a desiccator, etc. by tightly sealing it if test results are affected by moisture absorption.
   c) Store a sample in a tightly sealed container in a dark place at 1 ºC - 8 ºC if it is easily deteriorated by moisture.
2.3 Preparation of test samples

(1) Summary

a) Prepare a test sample by pre-drying, reducing, and grinding laboratory samples as necessary.

b) Conduct pre-drying if a laboratory sample is moist and hard to grind.

c) A laboratory sample made from such fertilizers as a fluid fertilizer or a particle-fertilizer, etc. that is sufficiently homogeneous can be used as a test sample.

d) If contamination by apparatus affects a test result, procedures such as pre-drying, reduction and grinding are prohibited.

e) Note that part of a test sample should not scatter, nor should surrounding fine particles or other alien substances be mixed with the test sample being prepared.

References


2) JIS K 0060: Sampling method of industrial wastes (1992)
2.3.1 Pre-drying

(1) Summary

This procedure is applicable to fertilizers whose laboratory sample is moist and hard to grind. The symbol of the procedure is 2.3.1-2017 or PD-1.

Conduct pre-drying using a drying apparatus, and measure the loss on drying in this procedure. In addition, calculate a conversion factor (actual article), if necessary, to convert the component content obtained in respective tests to the component content in a laboratory sample (actual article).

(2) Apparatus and instruments: Apparatus and instruments are shown below.

a) Drying apparatus: A drying apparatus that can be adjusted to the pre-drying temperature at ± 2 ºC.

b) Sample drying dish: Measure the mass to the order of 0.1g in advance. Additionally, use materials of a quality that do not affect the measurement of test components.

(3) Procedure: Conduct pre-drying as shown below.

a) Transfer 250 g - 1 kg of a laboratory sample to a sample drying dish, spread uniformly and measure the mass to the order of 0.1g.

b) Place a sample drying dish containing a laboratory sample in a drying apparatus and dry \(^{(1)}\).

c) Remove a sample drying dish from a drying apparatus and leave at rest at room temperature until it is balanced with atmospheric temperature \(^{(2)}\).

d) After leaving at rest, measure the mass of c) to the order of 0.1 g.

e) Calculate loss on drying in the pre-drying by the following formula (1). If necessary, calculate a conversion factor (actual article) by the following formula (2).

\[
\text{Loss on drying (\% (mass fraction))} = \left(\frac{W_1 - A}{W_1}\right) \times 100 \quad \ldots \ldots \quad (1)
\]

\[
\text{Conversion factor (actual article)} = \frac{A}{W_1} \quad \ldots \ldots \quad (2)
\]

\[W_1: \quad \text{Mass (g) of the sampled laboratory sample}\]

\[A: \quad \text{Mass (g) of the sampled laboratory sample}\]

Note

(1) Examples of drying temperature and drying time: About 70 hours at 40 ºC, no less than 5 hours at 65 ºC.

(2) An example of time to leave at rest: About 20 minutes

Comment 1

When calculating major components in a laboratory sample (actual article) such as compost and sludge fertilizers, etc. where the test sample is prepared conducting pre-drying, convert component contents in the analytical sample obtained in respective tests by the following formula.

\[
\text{Component content in a laboratory sample (actual article)} = B \times C
\]

\[B: \quad \text{Component content in an analytical sample obtained in each test}\]

\[C: \quad \text{Conversion factor (actual article)}\]

References

4) Flow sheet for pre-drying  The flow sheet for the pre-drying of a moist laboratory sample is shown below.

<table>
<thead>
<tr>
<th>250 g – 1 kg of Laboratory sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Transfer to a sample drying dish and spread uniformly. Measure the mass to the order of 0.1 g</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Drying</th>
</tr>
</thead>
<tbody>
<tr>
<td>Example: About 70 hours at 40 °C, no less than 5 hours at 65 °C</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Standing to cool</th>
</tr>
</thead>
<tbody>
<tr>
<td>Room temperature</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measure the mass to the order of 0.1 g</td>
</tr>
</tbody>
</table>

Figure  Flow sheet for pre-drying
2.3.2 Reduction (Separation)

(1) Summary
This procedure is applicable to fertilizers. The symbol of the procedure is 2.3.2-2017 or Red.-1.
In order to distinguish a test sample from a sample for granularity test and physical characteristics test, etc., reduce (separate) a laboratory sample with an increment reduction method, a riffle sampler method or a conical quartering method.

(2) Apparatus
a) Scoop for increment reduction: A scoop for increment reduction specified in the attached chart 1 of JIS M 8100.
b) Riffle sampler: A riffle sampler specified in the attached chart 3 of JIS M 8100.

(3) Procedure: Conduct reduction (separation) as shown below.
a) Increment reduction method: Conduct as indicated in 6.5.2 of JIS M 8100.
b) Riffle sampler method: Conduct as indicated in 6.5.3 of JIS M 8100.
c) Conical quartering method: Conduct as indicated in 6.5.4 of JIS M 8100.
2.3.3 Grinding

(1) Summary
This procedure is applicable to fertilizers. The symbol of the procedure is 2.3.3-2017 or GRD.1.
In order to prepare a homogeneous test sample, grind a laboratory sample with an adequate grinder until it completely passes through the designated granularity.

(2) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Grinder: Use a grinder (3) of a type and suitability for the granularity and the physical characteristics (1) of a laboratory sample. In addition, the grinder apparatus which come into contact with a laboratory sample should be made of materials (2) which do not affect the analytical value.
   b) Primary grinder: A grinder (3) that can primarily grind a large lump.
   c) Cutter machine: A cutter that can cut long stems, etc.
   d) Sieve: A sieve for the test specified in JIS Z 8801-1 or JIS Z 8801-2 or equivalents.

Note (1) The physical characteristics of a laboratory sample are defined by their solidity, toughness, specific gravity and adhesiveness.
   (2) (Ex.) Do not use stainless steel apparatus when preparing a test sample for chromium or nickel.
   (3) A centrifuging type grinder, a cutting mill, a vibration mill type grinder, etc.
   (4) A blender with an attachable blade, etc.

(3) Procedure: Conduct grinding as shown below.

(3.1) Fertilizers except ones specified in (3.2): Conduct as specified in 6.4 of JIS M 8100 and as shown below.
   a) Break or cut a laboratory sample with a primary grinder or a cutting machine as necessary.
   b) Grind with a grinding machine until it completely passes through a sieve of 500 µm - 1 mm aperture.
   c) Mix ground samples to make the test sample.

Comment 1 If the sampling amount of an analytical sample is less than 1 g, use a test sample which will completely pass through 500 µm aperture sieve. In addition, in case a test sample which suits the aforementioned condition cannot be obtained due to a deliquescent laboratory sample, etc., make one by crushing a test sample with a mortar and pestle, which completely passes through 1 mm aperture sieve.

(3.2) Fused phosphate fertilizer, slag silicate fertilizer, etc.: Conduct as specified in 6.4 of JIS M 8100 and as shown below.
   a) Grind a laboratory sample with a vibration mill type grinder.
   b) Transfer the ground laboratory sample to a sieve of 212 µm apertures.
   c) Incline the sieve about 20 degrees, supporting it with one hand or a bent arm, and tap the sieve frame with the other hand at the rate of about 120 times per minute. During the procedure, place the sieve in a horizontal position at the rate of 4 times per minute, rotate it 90 degrees and tap the sieve frame firmly one or two times.
   d) When fine powder attaches to the back side of a sieve screen, remove it gently from the back side to make minus sieve.
   e) Regarding the plus sieve of a sample, make them pass through by repeating the procedure in a) - d).
   f) Combine and mix the sample passed to make the test sample.
Comment 2 Conduct the procedures in (3.2) to obtain observed value of citric acid-soluble main components in a stable manner. For examples of applicable laboratory samples, fused matters such as fused phosphate and slag silicate fertilizer, fertilizers made from fused materials and calcined phosphate, etc. are listed.

Comment 3 The procedures in b) - d) are the procedures in 6.1.3 (1.4) of JIS Z 8815.
3. General tests
3.1 Moisture or moisture content
3.1.a Loss on drying method with drying apparatus

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type A (Def-M) and its symbol is 3.1.a-2017 or Mois.a-1.

Use drying apparatus under conditions suitable for the kind of fertilizers to be measured to heat analytical samples to measure loss on drying and obtain moisture in an analytical sample or the moisture content of a quality labeling standard of a special fertilizer (herein after referred to as “moisture”). Additionally, calculate a conversion factor (drying matter) to convert component content obtained by respective tests to component content in a drying matter as necessary.

This testing method corresponds to loss on heating in the Official Methods of Analysis of Fertilizers (1992).

(2) Apparatus and instruments: Apparatus and instruments are shown below.

a) Drying apparatus: Drying apparatus that can be adjusted to the test temperature ± 2 °C.

b) Ground-in stoppered weighing bottles (1): Low-form weighing bottles, 50 mm × 30 mm, specified in JIS R 3503. Dry by heating in advance in a drying apparatus at 75 °C - 130 °C, stand to cool in a desiccator, and measure the mass to the order of 1 mg.

Note (1) Aluminum weighing dishes described in the Handbook of the Feed Analysis Standards -2009- can also be used.

(3) Measurement: Conduct measurement as shown below.

a) Transfer 2 g-5 g of an analytical sample to a ground-in stoppered weighing bottle, spread so that the thickness is no more than 10 mm, and measure the mass to the order of 1 mg.

b) Place the ground-in stoppered weighing bottle containing the analytical sample in a drying apparatus at 100 °C ± 2 °C, and heat for 5 hours (2).

c) After heating, fit the stopper into the ground-in stoppered weighing bottle, and immediately transfer to a desiccator to let it stand to cool.

d) After standing to cool, remove the ground-in stoppered weighing bottle from the desiccator, and measure the mass to the order of 1 mg.

e) Calculate loss on drying in the analytical sample by the following formula (1) as moisture. If necessary, calculate a conversion factor (actual article) by the following formula (2).

\[
\text{Loss on drying (% (mass fraction))} = \left(\frac{W_1 - A}{W_1}\right) \times 100 \quad \ldots \quad (1)
\]

\[
\text{Conversion factor (drying matter)} = \frac{W_1}{A} \quad \ldots \quad \ldots \quad \ldots \quad (2)
\]

\[
W_1: \quad \text{Mass (g) of the sampled analytical sample}
\]

\[
A: \quad \text{Mass (g) of the analytical sample after drying}
\]

Note (2) Heat simultaneously the slightly moved or removed stopper of the ground-in stoppered weighing bottle.

Comment 1 When pre-drying a laboratory sample such as compost and sludge fertilizers to prepare a test sample, calculate the moisture of the laboratory sample (actual article) by the following formula:
Moisture (% (mass fraction)) in the laboratory sample (actual article) = \( B + C \times \left( \frac{100 - B}{100} \right) \)

- \( B \): Loss on drying (% (mass fraction)) of the laboratory sample (actual article) by the pre-drying procedure
- \( C \): Loss on drying (% (mass fraction)) in the analytical sample by moisture measurement

**Comment 2** When calculating harmful content in a drying matter of sludge fertilizers, etc., convert component content in a test sample obtained from respective tests by the following formula.

Component content in a drying matter = \( D \times E \)

- \( D \): Component content in an analytical sample obtained in each test
- \( E \): Conversion factor (drying matter)

**Comment 3** Use drying conditions in Table 1 for fertilizers of the types shown below:

<table>
<thead>
<tr>
<th>Type of fertilizers</th>
<th>Sampling amount of analytical samples</th>
<th>Drying temperature</th>
<th>Drying time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Superphosphate of lime, triple superphosphate of lime, or fertilizers containing these</td>
<td>About 5 g</td>
<td>100 °C ± 2 °C</td>
<td>3 hours</td>
</tr>
<tr>
<td>Ammonium sulfate, sodium nitrate, and potassium salts</td>
<td>2 g - 5 g</td>
<td>130 °C ± 2 °C</td>
<td>Until a constant weight is achieved.</td>
</tr>
<tr>
<td>Urea and urea-containing fertilizers</td>
<td>About 5 g</td>
<td>75 °C ± 2 °C</td>
<td>4 hours</td>
</tr>
<tr>
<td>Silica gel fertilizer and fertilizer containing silica gel, and silica hydrogel fertilizer</td>
<td>About 5 g</td>
<td>180 °C±5 °C</td>
<td>3 hours</td>
</tr>
</tbody>
</table>

**Comment 4** For samples containing volatile matters, subtract the volatile matter content by the following a) and b) from loss on drying to obtain moisture.

- a) Fertilizers containing guano or diammonium hydrogen phosphate, etc.: Determine total nitrogen in the test sample, and in the analytical sample after the drying procedure; convert the difference between the quantitation values into ammonia (NH\(_3\)) to make the volatile matter content.
- b) Potassium hydrogen carbonate: Determine carbon dioxide in the test sample, and in the analytical sample after the drying procedure; the difference between the quantitation values is the volatile matter content.
References


3) JIS Z 0701: Silica gel desiccants for packaging (1997)

4) Flow sheet for moisture: The flow sheet for moisture in fertilizers is shown below:

<table>
<thead>
<tr>
<th>2 g - 5 g</th>
</tr>
</thead>
<tbody>
<tr>
<td>analytical sample</td>
</tr>
</tbody>
</table>

Transfer to a ground-in stopper weighing bottle, spread so that the thickness is no more than 10 mm. Measure the mass to the order of 1 mg.

<table>
<thead>
<tr>
<th>Heating</th>
</tr>
</thead>
</table>

100 °C ± 2 °C, 5 hours

<table>
<thead>
<tr>
<th>Standing to cool</th>
</tr>
</thead>
<tbody>
<tr>
<td>Desiccator</td>
</tr>
</tbody>
</table>

Measure the mass to the order of 1 mg.

Figure Flow sheet for moisture in fertilizers by loss on drying with drying apparatus.
3.1.b  Loss on drying method by moisture analyzers

(1)  **Summary**

The testing method is applicable to sludge fertilizers, compost, and organic fertilizers, etc. This testing method is classified as Type B and its symbol is 3.1.b-2017 or Mois.b-1.

Use a moisture analyzer in the heat drying method to measure loss on drying and obtain moisture in an analytical sample or the moisture content of a quality labeling standard of a special fertilizer (herein after referred to as “moisture”). Additionally, calculate a conversion factor (drying matter) to convert component content obtained by respective tests to component content in a drying matter as necessary.

In addition, the performance of this testing method is shown in Comment 3.

(2)  **Instruments**: Instruments are as shown below:

   a)  **Moisture analyzer**: A moisture analyzer consisted of a heat source to heat an analytical sample (halogen lamp, infrared heater, ceramic heater, etc.) and a balance\(^{(1)}\) with calibration function.

   Note (1) There is a method to calibrate with calibration weights or a method to calibrate automatically with built-in weights.

(3)  **Measurement**: Conduct measurement as shown below. However, conduct in advance a comparative test with 3.1.a Loss on drying with drying apparatus using sludge fertilizers, compost, and organic fertilizers, etc., to confirm that there is no difference in the quantitation value of moisture.

   a)  Transfer 2 g - 5 g of an analytical sample to a ground-in stoppered weighing bottle, spread so that the thickness is no more than 10 mm, and measure the mass to the order of 1 mg.

   b)  Heat at 100 °C\(^{(2)}\), until a constant weight is achieved.

   c)  After the end of heating\(^{(2)}\), measure the mass to the order of 1 mg.

   d)  Calculate loss on drying in the analytical sample by the following formula (1) as moisture. If necessary, calculate a conversion factor (drying matter) by the following formula (2).

   \[
   \text{Loss on drying} \text{ (% (mass fraction))} = \left(\frac{W_1 - A}{W_1}\right) \times 100 \quad \ldots \; (1)
   \]

   \[
   \text{Conversion factor (drying matter)} = \frac{W_1}{A} \quad \ldots \; (2)
   \]

   \(W_1\):  Mass (g) of the sampled analytical sample

   \(A\):  Mass (g) of the analytical sample after drying

   Note (2) The setup of the drying program and the determination parameter for the end of heating (constant weight) is according to the specification and the operation method of the moisture analyzer used.

Comment 1  When pre-drying is conducted, calculate the moisture of the laboratory sample (actual article) by the following formula:

\[
\text{Moisture} \text{ (% (mass fraction)) in the laboratory sample (actual article)} = B + C \times \left(\frac{100 - B}{100}\right)
\]

\(B\):  Loss on drying (% (mass fraction)) of the laboratory sample (actual article) by the pre-drying procedure

\(C\):  Loss on drying (% (mass fraction)) in the analytical sample by moisture
Comment 2 When calculating harmful content in a drying matter of sludge fertilizers, etc., convert component content in a test sample obtained from respective tests by the following formula.

Component content in a drying matter $= D \times E$

$D$: Component content in an analytical sample obtained in each test  
$E$: Conversion factor (drying matter)

Comment 3 Table 1 shows the results of the comparison of the measurement values by loss on drying method with drying apparatus and the measurement values by loss on drying method with a moisture analyzer using organic fertilizers, compost and sludge fertilizers in order to evaluate trueness. Table 2 shows results and analysis results from a collaborative study for testing method validation.

<table>
<thead>
<tr>
<th>Symbol of measurement value</th>
<th>Sample</th>
<th>Range of yi~yj (%)</th>
<th>Regression coefficient $y = a + bx$</th>
<th>Correlation coefficient $r$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drying apparatus method 1)</td>
<td>Moisture method 2)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$x_i$</td>
<td>$y_i$</td>
<td>Sludge fertilizer 4)</td>
<td>26</td>
<td>5.50~90.61</td>
</tr>
<tr>
<td>$x_j$</td>
<td>$y_j$</td>
<td>Organic fertilizer 5)</td>
<td>25</td>
<td>2.96~12.33</td>
</tr>
</tbody>
</table>

1) 3.1.a Loss on drying method with drying apparatus  
2) 3.1.b Loss on drying method with a moisture analyzer  
3) Mass fraction  
4) Sewage sludge fertilizer, Human waste sludge fertilizer, Industrial sludge fertilizer, Composted sludge fertilizer  
5) Fish meal, Byproduct organic fertilizer of vegetable origin, Compost, Steamed leather meal, Rape seed meal and powdered rape seed meal
Table 2 Results and analysis results from a collaborative study for the validation of the moisture test method.

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (%)</th>
<th>$s_r$ (%)</th>
<th>$RSD_r$ (%)</th>
<th>$s_R$ (%)</th>
<th>$RSD_R$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sewage sludge fertilizer</td>
<td>9</td>
<td>21.93</td>
<td>0.32</td>
<td>1.4</td>
<td>0.47</td>
<td>2.1</td>
</tr>
<tr>
<td>Human waste sludge fertilizer</td>
<td>8</td>
<td>13.36</td>
<td>0.14</td>
<td>1.1</td>
<td>0.37</td>
<td>2.8</td>
</tr>
<tr>
<td>Industrial sludge fertilizer</td>
<td>9</td>
<td>34.28</td>
<td>0.21</td>
<td>0.6</td>
<td>0.50</td>
<td>1.5</td>
</tr>
<tr>
<td>Calcined sludge fertilizer</td>
<td>9</td>
<td>38.75</td>
<td>0.59</td>
<td>1.5</td>
<td>0.59</td>
<td>1.5</td>
</tr>
<tr>
<td>Composted sludge fertilizer</td>
<td>9</td>
<td>27.1</td>
<td>0.26</td>
<td>0.9</td>
<td>0.60</td>
<td>2.2</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n =$ number of laboratories $\times$ number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(4) Flow sheet for moisture: The flow sheet for moisture in sludge fertilizers, compost, and organic fertilizers, etc. is shown below:

```
<table>
<thead>
<tr>
<th>About 5 g analytical sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heating</td>
</tr>
<tr>
<td>End of drying</td>
</tr>
<tr>
<td>Measurement</td>
</tr>
</tbody>
</table>
```

Transfer to a weighing dish, spread so that the thickness is no more than 10 mm. Measure the mass to the order of 1 mg.

100 ºC

Constant weight

Measure the mass to the order of 1 mg.

Figure Flow sheet for moisture in sludge fertilizers, compost, and organic fertilizers, etc. by the loss on drying method with a moisture analyzer.
3.2 Ash content
3.2.1 Ignition residue method

(1) Summary
The method is applicable to organic fertilizers and fertilizers containing organic matters. This testing method is classified as Type A (Def-M) and its symbol is 3.2.a-2017 or Ash.a-1.
The method ignites an analytical sample with an electric furnace and measures residue on ignition to obtain ash content in an analytical sample.

(2) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Electric furnace: Electric furnace that can be adjusted to 550°C ± 5°C.
   b) Crucible: After heating a porcelain crucible for chemical analysis specified in JIS R 1301 with an electric furnace at 550 °C ± 5 °C, stand to cool in a desiccator in advance and measure the mass to the order of 1 mg.

(3) Measurement: Conduct measurement as shown below.
   a) Transfer about 2 g of an analytical sample into a crucible, and measure the mass to the order of 1 mg.
   b) Place it into an electric furnace, heat gently until carbonized (1).
   c) Heat at 550 °C ± 5 °C for no less than 4 hours (1).
   d) After heating, move the crucible into a desiccator and let it stand to cool.
   e) After standing to cool, remove the crucible from the desiccator and measure the mass to the order of 1 mg.
   f) Calculate the residue on ignition in the analytical sample by the following formula to make ash content.

\[
\text{Residue on ignition (\% (mass fraction))} = \left(\frac{A}{W}\right) \times 100
\]

\[W:\] Mass (g) of the sampled analytical sample
\[A:\] Mass (g) of the ignited analytical sample

Note (1) Example of carbonizing and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

(4) Flow sheet for ash content: The flow sheet for ash content in fertilizers is shown below:

<table>
<thead>
<tr>
<th>About 2 g of analytical sample</th>
<th>Transfer to a crucible.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charred</td>
<td>Heating gently with an electric furnace</td>
</tr>
<tr>
<td>Incineration</td>
<td>No less than 4 hours at 550 °C ± 5 °C</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Desiccator</td>
</tr>
<tr>
<td>Measurement</td>
<td>Measure the mass to the order of 1 mg</td>
</tr>
</tbody>
</table>

Figure Flow sheet for ash content in fertilizers.
3.3 pH
3.3.a Glass electrode method
(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type A (Def-M) and its symbol is 3.3.a-2017 or pH.a-1.
Measure the pH of fertilizers with a pH meter using a glass electrode.

(2) Reagent: Reagents are as shown below.
a) Oxalate pH standard solution: Oxalate pH standard solution class 2 traceable to National Metrology.
b) Phthalate pH standard solution: Phthalate pH standard solution class 2 traceable to National Metrology.
c) Neutral phosphate pH standard solution: Neutral phosphate pH standard solution class 2 traceable to National Metrology.
d) Borate pH standard solution: Borate pH standard solution class 2 traceable to National Metrology.
e) Carbonate pH standard solution: Carbonate pH standard solution class 2 traceable to National Metrology.

Comment 1 Respective pH standard solutions stored for long time should not be used since the pH value may change during storage period. In particular, note that borate pH standard solution and carbonate pH standard solution easily absorb carbon dioxide in the air, so that the pH values deteriorate. The pH standard solution that was used once or left exposed to the air should not be used.

(3) Instruments: Instruments are as shown below:
a) pH meter: Use type II specified in JIS Z 8802.

Comment 2 Conduct the calibration of a pH meter as indicated in JIS Z 8802. Actual calibration operation is according to the operation procedure of the pH meter used for measurement. When the pH of a sample solution is no more than 7, use neutral phosphate pH standard solution and oxalate pH standard solution, or phthalate pH standard solution. When it exceeds 7, use neutral phosphate pH standard solution and borate pH standard solution, or carbonate pH standard solution.

(4) Test procedures
(4.1) Preparation of sample solution: Conduct preparation of a sample solution as shown below.
(4.1.1) Fertilizers except inorganic fertilizers
a) Transfer a predetermined amount of an analytical sample (1) into a ground-in stopper flask and add water 5 - 10 times the volume.
b) Mix with a magnetic stirrer, filter with Type 3 filter paper to make a sample solution.

Note (1) In the case of a moist laboratory sample, it is recommended to use a sample that is not pre-dried.

(4.1.2) Inorganic fertilizers
a) Transfer a predetermined amount of an analytical sample (1) into a ground-in stopper flask and
add water 100 times the volume.
b) Mix with a magnetic stirrer, filter with Type 3 filter paper to make a sample solution.

**Comment 3** The procedure in (4.1.1) is the same as 3.4.a (4.1). Additionally, the sample solution prepared in 4.2.4.a (4.1) can be used instead of the sample solution prepared by (4.1.2).

(4.2) **Measurement**: Conduct measurement as shown below. Actual calibration operation is according to the operation procedure of the pH meter used for measurement.

a) Wash the read station of a calibrated pH meter repeatedly no less than 3 times with water and wipe out with clean and soft paper, etc.

b) Transfer a sample solution into a beaker (2), dip the read station in the solution and measure the pH value.

**Note** (2) It is necessary to transfer sufficient volume of sample solution to keep a measurement value stable.

**Comment 4** If a pH meter has a temperature correction dial or a digital switching, measure the pH value after adjusting the graduation of the pH meter with the temperature of a sample.

**References**

1) JIS Z 8802: Methods for determination of pH of aqueous solutions (2011)

(5) **Flow sheet for pH value**: The flow sheet for pH value in fertilizers is shown below.

```
<table>
<thead>
<tr>
<th>Predetermined volume of analytical sample</th>
<th>Ground-in stopper flask.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>← Water 5 -10 times or 100 times the volume</td>
</tr>
<tr>
<td>Stirring</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Filtration</td>
<td>pH meter</td>
</tr>
<tr>
<td>Measurement</td>
<td></td>
</tr>
</tbody>
</table>
```

Figure Flow sheet for pH in fertilizers.
3.4   Electrical conductivity

3.4.a  Measurement method with an electrical conductivity meter

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type A (Def-M) and its symbol is 3.4.a-2017 or EC.a-1.

Measure the electrical conductivity of organic fertilizers such as compost or sludge fertilizers with an electrical conductivity meter.

(2) Reagent: Reagents are as shown below.
   a) Potassium chloride: Grind potassium chloride used for measurement of electrical conductivity specified in JIS K 8121 with an agate mortar to powder and heat for 4 hours at 500 °C ± 5 °C, and then stand to cool in a desiccator.
   b) Potassium chloride standard solution (1): Measure predetermined volume (2) of potassium chloride of a) on a weighing dish, dissolve in a small amount of water, transfer it into a 1000-mL volumetric flask, and add up to the marked line with water.

Note   (1) Store potassium chloride standard solution in a polyethylene or borosilicate glass bottle and seal the bottle.
       (2) The volume that is recommended for an instrument or a cell used.

Comment 1 Potassium chloride standard solution used once or left in the air should not be used.

(3) Instruments: Instruments are as shown below:
   a) Electrical conductivity meter: An electrical conductivity meter specified in JIS K 0130

Comment 2 Check the indicated value as shown in 6.2 in JIS K 0130 as necessary. Actual procedure to check is according to the operation procedure of the electrical conductivity meter used for measurement.

(4) Test procedures
(4.1) Preparation of sample solution: Conduct preparation of a sample solution as shown below.
   a) Transfer the predetermined volume of an analytical sample (3) into a ground-in stopper flask and add water 10 times the equivalent volume of dry matter (4).
   b) Mix with a magnetic stirrer, filter with Type 3 filter paper to make a sample solution.

Note   (3) In the case of a moist laboratory sample, it is recommended to use a sample that is not pre-dried.
       (4) If the sample solution becomes hard to measure because it is gelled by the influence of flocculants in sludge fertilizer, etc., increase the volume of water added. However, this fact should be expressed in the test result.

Comment 3 The procedure in (4.1) is the same as 3.3.a (4.1.1).

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0130 and as shown below.
Actual measurement operation is according to the operation procedure of the electrical conductivity meter used for measurement.
   a) Wash the read station of an electrical conductivity meter repeatedly no less than 3 times with water.
   b) Transfer a sample solution into a beaker (5), dip the read station and measure electrical conductivity.
conductivity.

**Note (5)** It is necessary to transfer sufficient volume of sample solution to keep a measurement value stable.

**References**

(5) **Flow sheet for electrical conductivity**: The flow sheet for electrical conductivity is showed below.

![Flow sheet for electrical conductivity](image_url)

Figure Flow sheet for electrical conductivity in fertilizers.
3.5 Granularity

3.5.a Dry-type sieving testing method

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type A (Def-M) and its symbol is 3.5.a-2017 or P-size.a-1.

Measure the particle diameter distribution of solid fertilizers with a dry-type sieving analysis.

(2) Apparatus: Apparatus are shown below.
   a) Sieve: A sieve for testing specified in JIS Z 8815
   b) Clogging removal brush: A brush which is adequately hard for the apertures and does not damage the sieving screen.
   c) Weighing dish: A container that can contain about 250 g of a sample. Measure the mass to the order of 0.1g in advance.

(3) Dry-type sieving analysis procedure: Conduct sieving analysis corresponding to the aperture size of a sieve used as indicated in JIS Z 8815 and as shown below.

(3.1) More than 1mm and no more than 4mm
   a) Stack a large aperture sieve on an acceptor so that the large sieve is on top.
   b) Measure a laboratory sample to the order of 0.1g and transfer it to the sieve at the top section.
   c) After putting a stopper on it, hold the stacked sieves with both hands, and vibrate (1) them back and forth along a unidirectional and horizontal plane at about 60 times per minute with about 70 mm amplitude.
   d) Transfer respective plus and minus sieves to a weighing dish (2).

Note (1) Conduct more circular motion at the rate of about 3 revolutions per minute as necessary.
(2) Turning over the back side of a sieve, remove the clogged particles from the sieve screen with a clogging removal brush and combine them with the plus sieve.

(3.2) Less than or equal to 1mm
   a) Stack a large aperture sieve on an acceptor so that the large sieve is on top.
   b) Measure the laboratory sample or minus sieve of (3.1) c) to the order of 0.1g and transfer it to the sieve at the top section.
   c) After putting a stopper on it, incline the stacked sieves about 20 degrees, supporting with one hand or a bent arm, and tap the sieve frame with the other hand at the rate of about 120 times per a minute.
   d) During the procedure in c), place the sieve in a horizontal position at the rate of 4 times per minute, rotate it 90 degrees and tap the sieve frame firmly one or two times.
   e) Transfer respective plus and minus sieves (3) to a weighing dish (2).

Note (3) When fine powder attaches to the back side of a sieve screen, remove them gently from the back side with a clogging removal brush and combine them with the minus sieve.

(4) Measurement of granularity distribution: Calculate the granularity distribution in an analytical sample as shown below.
   a) Measure respective mass of plus and minus sieves to the order of 0.1g.
   b) Calculate “plus sieve percentage” and “integrated minus sieve percentage” with the following formula and round the results to the first decimal place.
   c) Confirm that the sum of the mass of the plus sieve and the mass of the minus sieve with the smallest aperture is in the range of ± 2 % of the mass of sample measured in (3.1) b) or (3.2)
b).

Mass percentage of plus sieve or minus sieve (%) \( (R) = \frac{A}{T} \times 100 \)

\( A \): Mass of plus sieve or minus sieve (g)  
\( T \): Sum of the mass of plus and minus sieve (g)

References
1) JIS Z 8815: Test sieving - General requirements (1994).

(5) Flow sheet for granularity: The flow sheet of granularity of solid fertilizers is shown below.

```
<table>
<thead>
<tr>
<th>Laboratory sample or Test sample</th>
<th>Measure the mass to the order of 0.1g</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry-type sieving</td>
<td></td>
</tr>
<tr>
<td>Mass measurement of respective plus sieve and minus sieve</td>
<td>Measure the mass to the order of 0.1g</td>
</tr>
</tbody>
</table>
```

Figure  Flow sheet for granularity of solid fertilizers
3.6 Oil content
3.6.a Diethyl ether extraction method

(1) Summary
This testing method is applicable to organic fertilizers. This testing method is classified as Type A (Def-M) and its symbol is 3.6.a-2017 or Oil.a-1.

Extract an analytical sample with diethyl ether using a Soxhlet extractor and measure the extract to obtain oil content in an analytical sample. The oil content contains not only fat but also fat-soluble pigments (carotenoid, chlorophyll, etc.), wax, and free fatty acids, etc.

(2) Reagent: Reagents are as shown below.
   a) Diethyl ether: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Drying apparatus: Drying apparatus that can be adjusted to the test temperature ± 2 °C.
   b) Soxhlet extractor: An inter-changeable Soxhlet extractor, cooling apparatus and weighing bottles (Example: JIS R 3503, attached figure 71)
   c) Water bath: A water bath that can be adjusted to about 60 °C
   d) Weighing bottle: A flat bottle flask connectable to a Soxhlet extractor. After heating with a drying apparatus at 100 °C - 105 °C in advance, stand to cool in a desiccator and measure the mass to the order of 1mg.
   e) Cylindrical filter paper: A cylindrical filter paper made of cellulose. Example: 22 mm external diameter, 20 mm internal diameter, 90 mm total length (1).

(4) Measurement: Conduct measurement as shown below.
   a) Weigh 2 g - 5 g of an analytical sample to the order of 1mg, and transfer it into a cylindrical filter paper.
   b) Place absorbent cotton on the upper end of an analytical sample (2), as if gently pushing it, and heat it at 100 °C - 105 °C for 2 hours.
   c) As soon as heating is complete, move the cylindrical filter paper to a desiccator and stand to cool.
   d) After standing to cool, transfer it into a Soxhlet extractor and connect it to a cooling apparatus.
   e) Transfer adequate volume of diethyl ether (3) into a weighing bottle, connect it to the Soxhlet extractor and heat (4) it for 8 hours to extract.
   f) Recover the diethyl ether (5).
   g) Disconnect the weighing bottle from the Soxhlet extractor and vaporize the diethyl ether (6).
   h) Heat the weighing bottle (7) at 100 °C - 105 °C for 3 hours.
   i) As soon as heating is complete, move the weighing bottle to the desiccator and stand to cool.
   j) After standing to cool, remove the weighing bottle from the desiccator and measure the mass to the order of 1mg.
   k) Calculate oil content with the following formula.

\[
\text{Oil content (\% (mass fraction)) = \left(\frac{B}{A}\right) \times 100}
\]

   \[A: \text{Mass (g) of the sampled analytical sample}\]
   \[B: \text{Mass of extract of diethyl ether (g)}\]

   Note (1) Select a scale according to the volume of a Soxhlet extractor.
(2) The purpose is to prevent overflow at the upper end of an analytical sample.
(3) The amount of diethyl ether depends on the volume of a weighing bottle.
(4) Adjust the temperature for diethyl ether to circulate 16 - 20 times per hour. (Target temperature is about 60 ºC)
(5) Remove the cylindrical filter paper from the Soxhlet extractor. In the case of a cock attached Soxhlet extractor, open the cock and recover it.
(6) It is dangerous if diethyl ether resides in a weighing bottle when the bottle is transferred to a drying apparatus.
(7) Wipe the outside of a weighing bottle since there is a risk of garbage or stain sticking to it.

References

(5) Flow sheet for oil content: The flow sheet for oil content in organic fertilizers is shown below:

<table>
<thead>
<tr>
<th>2 g - 5 g analytical sample</th>
<th>Weigh to the order of 1mg to a cylindrical filter paper</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-drying</td>
<td>100 ºC - 105 ºC, for 2 hours</td>
</tr>
<tr>
<td>Soxhlet extractor</td>
<td></td>
</tr>
<tr>
<td>Extraction</td>
<td>Diethyl ether, heating, for 8 hours</td>
</tr>
<tr>
<td>Weighing bottle</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>100 ºC - 105 ºC, for 3 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Desiccator</td>
</tr>
<tr>
<td>Measurement</td>
<td>Measure the mass to the order of 1mg</td>
</tr>
</tbody>
</table>

Figure Flow sheet for oil content in organic fertilizers.
4. **Main components, guaranteed components, etc**

4.1 **Nitrogen**

4.1.1 **Total nitrogen**

4.1.1.a **Kjeldahl method**

(1) **Summary**

This testing method is applicable to fertilizers containing no nitrate nitrogen. This testing method is classified as Type C and its symbol is 4.1.1.a-2017 or T-N.a-1.

Add sulfuric acid, potassium sulfate and copper (II) sulfate pentahydrate to an analytical sample, pretreat by Kjeldahl method to change total nitrogen (T-N) to ammonium ion, and add a sodium hydroxide solution to subject to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain the total nitrogen (T-N) in an analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain the total nitrogen (T-N) in an analytical sample. This testing method corresponds to the sulfuric acid method in the Official Methods of Analysis of Fertilizers (1992). In addition, the performance of this testing method is shown in Comment 8.

(2) **Reagent**: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

**Standardization:** Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL -300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

$$f_1 = \left( \frac{W_1 \times A}{0.01/97.095} \right) \times \left( \frac{V_1}{V_2} \right) \times \left( \frac{1000/V_3} \right) \times \left( \frac{1}{C_1} \right)$$

- $W_1$: Mass (g) of sulfamic acid sampled
- $A$: Purity (% (mass fraction)) of sulfamic acid
- $V_1$: Volume (mL) of sulfamic acid solution transferred
- $V_2$: Constant volume (250 mL) of sulfamic acid solution
- $V_3$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- $C_1$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) **Sulfuric acid**: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) **0.25 mol/L sulfuric acid**: (1) (2): Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

**Standardization:** Transfer a predetermined amount (3) of 0.25 mol/L sulfuric acid to a 200-mL -300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution,
and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\[ V_4/V_5 \]  

\[ \text{(1)} \]

Factor of 0.25 mol/L sulfuric acid \((f_2)\)

\[ (f_1 \times C_1 \times V_4/V_5)/(C_2 \times 2) \]  

\[ \text{(2)} \]

\(V_4\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\(V_5\): Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization

\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(C_2\): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

d) **Boric acid solution (40 g/L)**: Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.
e) **Catalyst** (5): Mix potassium sulfate specified in JIS K 8962 and copper (II) sulfate pentahydrate (6) specified in JIS K 8983 in the ratio of 9 to 1.
f) **Sodium hydroxide solution (200 g/L - 500 g/L)** (1): Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.
g) **Bromothymol blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.
h) **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.
i) **Methylene blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.
j) **Methyl red–methylene blue mixture solution**: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).
k) **Bromocresol green solution (0.5 g/100 mL)**: Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.
l) **Methyl red–bromocresol green mixture solution**: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

**Note**

(1) This is an example of preparation; prepare an amount as appropriate.
(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).
(3) 5 mL - 10 mL
(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.
(5) A tablet is commercially available.
(6) Crush into powder as appropriate.

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

**Comment 2** 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid
conforming to ISO/IEC 17025.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.

a) **Steam distillation apparatus**

b) **Digestion flask**: Kjeldahl flask
c) **Distillation flask**: A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.

(4) **Test procedures**

(4.1) **Kjeldahl method**: Conduct digestion as shown below.
a) Weigh 0.5 g - 5 g of an analytical sample to the order of 1 mg, and put it in a 300-mL - 500-mL digestion flask.
b) Add 5 g - 10 g of catalyst, and further add 20 mL - 40 mL of sulfuric acid, shake to mix \(^{7}\) and heat gently.
c) After bubbles cease to form, heat until white smoke of sulfuric acid evolves.
d) Ignite until organic matters are completely digested \(^{8}\).
e) After standing to cool, add a small amount of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water \(^{9}\), and further mix by shaking.
f) After cooling is complete, add water up to the marked line to make the digestion solution.

**Note**

- \(^{7}\) Leaving at rest overnight is preferable.
- \(^{8}\) When the solution has finished changing color, heat further for no less than 2 hours.
- \(^{9}\) When the entire sample solution volume is used in measurement, it is not necessary to transfer it to a volumetric flask.

**Comment 3** The procedure in (4.1) is the same as that in (4.1.1) a - f) in 4.2.1.a.

**Comment 4** In the case of fish meal containing amino acids that are not easily digested, use 0.5 g - 1 g analytical sample, 10 g catalyst and 30 mL - 40 mL sulfuric acid.

**Comment 5** In the case of nitrolime, moisten by adding a small amount of water before the procedure in (4.1) b). Care should be taken because bubbles are produced by the addition of sulfuric acid.

(4.2) **Distillation**: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.
a) Transfer a predetermined amount \(^{10}\) of 0.25 mol/L sulfuric acid to an acceptor \(^{11}\), add a few drops of methyl red—methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount \(^{10}\) of boric acid solution (40 g/L) to an acceptor \(^{11}\), add a few drops of methyl red—bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.
b) Transfer a predetermined amount of the digestion solution to a 300-mL distillation flask, add a proper amount of sodium hydroxide solution (200 g/L - 500 g/L) \(^{12}\), and immediately connect this distillation flask to the steam distillation apparatus.
c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.
d) Stop distilling when the distillate has reached 120 mL - 160 mL.
e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

**Note**

- \(^{10}\) 5 mL - 20 mL
- \(^{11}\) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL
beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).

(12) An amount sufficient to make the solution strong alkalinity. A blue color will appear.

(4.3) **Measurement:** Conduct measurement as shown below.

(4.3.1) When 0.25 mol/L sulfuric acid is used in (4.2):

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green.

b) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\begin{align*}
\text{Total nitrogen (T-N) (% (mass fraction)) in the analytical sample} & = (B \times V_6 - V_7) \times C_1 \times f_1 \times (V_9/V_8) \times (14.007/W_3) \times (100/1000) \\
& = (B \times V_6 - V_7) \times C_1 \times f_1 \times (V_9/V_8) \times (1.4007/W_3)
\end{align*}
\]

- **B:** Volume of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid
- **V_6:** Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)
- **V_7:** Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- **C_1:** Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- **f_1:** Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- **V_8:** Predetermined volume (mL) of the digestion solution in (4.1) f)
- **V_9:** Transferred amount (mL) of the digestion solution subjected to distillation in (4.2) b)
- **W_3:** Mass (g) of the analytical sample

(4.3.2) When a boric acid solution (40 g/L) is used in (4.2):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red

b) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\begin{align*}
\text{Total nitrogen (T-N) (% (mass fraction)) in the analytical sample} & = V_{10} \times C_2 \times 2 \times f_2 \times (V_{11}/V_{12}) \times (14.007/W_2) \times (100/1000) \\
& = V_{10} \times C_2 \times 2 \times f_2 \times (V_{11}/V_{12}) \times (2.8014/W_2) \times (100/1000)
\end{align*}
\]

- **V_{10}:** Volume (mL) of 0.25 mol/L sulfuric acid needed for titration
- **C_2:** Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid
- **f_2:** Factor of 0.25 mol/L sulfuric acid
- **V_{11}:** Predetermined volume (mL) of the digestion solution in (4.1) f)
- **V_{12}:** Transferred amount (mL) of the digestion solution subjected to distillation in (4.2) b)
- **W_2:** Mass (g) of the analytical sample

**Note** (13) The endpoint is reached when the color changes from green to light red.

**Comment 6** The titration procedures in (2) a) **Standardization**, (2) c) **Standardization** and (4.3) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

**Comment 7** The nitrogen content in the analytical sample can be measured by using an automatic

nitrogen analyzer (Kjeldahl method) instead of the test procedure in (4). The setup of the program and the parameter of the analyzer as well as vessels etc. are according to the specification and the operation method of the automatic nitrogen analyzer used. However, conduct in advance a comparative test with the test procedure in (4) using fertilizers containing no nitrate nitrogen, to confirm that there is no difference in the quantitation value of total nitrogen.

Comment 8 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 98.5 % - 100.6 % and 97.1%-99.2 % as total nitrogen (T-N) respectively.

The results of the collaborative study (limited to reported values with the Kjeldahl method) to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.2 % (mass fraction) for solid fertilizers, and 0.02 % (mass fraction) for fluid fertilizers.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Analysis results of the collaborative study to determine a certified reference material fertilizer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name of</td>
<td>Number of repetitions</td>
</tr>
<tr>
<td>certified reference material fertilizer</td>
<td>laboratory</td>
</tr>
<tr>
<td>FAMIC-A-10</td>
<td>11</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Kjeldahl method
2) Average (the number of laboratory ($p$) $\times$ test days (2) $\times$ the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

3) Takashi KUBOTA, Tomoko OSHIDA, Kozue YANAI, Yuzuru INOUE, Seiji MATSUI, Takaharu MATSUMOTO, Eiichi ISHIKURO and Akemi YASUI: Improvement of the Conditions for the Determination of Total Nitrogen in Fish Meal in Kjeldahl Method and Its Comparison with Dumas Method, Bunseki kagaku, 60, p. 67 - 74 (2011)
(5) Flow sheet for total nitrogen: The flow sheet for total nitrogen in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5 g - 5 g analytical sample</td>
<td>Weigh to the order of 1 mg into a 300-mL digestion flask.</td>
</tr>
<tr>
<td>5 g - 10 g catalyst</td>
<td>Gently after foam no longer evolves, ignite until organic matters are completely digested.</td>
</tr>
<tr>
<td>20 mL - 40 mL sulfuric acid</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td></td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Weigh to the order of 1 mg into a 300-mL digestion flask.</td>
</tr>
<tr>
<td>small amount of water</td>
<td></td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask, water</td>
</tr>
<tr>
<td>Cooling</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Digestion solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1  The flow sheet for total nitrogen in fertilizers (Kjeldahl method procedure)

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>300-mL distillation flask</td>
<td></td>
</tr>
<tr>
<td>Sodium hydroxide solution (200 g/L - 500 g/L)</td>
<td>Receiver: 200-mL - 300 - mL Erlenmeyer flask or beaker. A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution</td>
</tr>
<tr>
<td>Steam distillation apparatus</td>
<td>Distillation rate: 5 mL/min -7 mL/min</td>
</tr>
<tr>
<td>Steam distillation</td>
<td>120 mL – 160 mL distillate</td>
</tr>
<tr>
<td>Stop distilling</td>
<td>Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)</td>
</tr>
<tr>
<td>Titration</td>
<td>0.1 mol/L-0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or 0.25 mol/L sulfuric acid (until the solution becomes light red)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for total nitrogen in fertilizers (Distillation and measurement procedure)
4.1.1.b Combustion method

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 4.1.1.b-2017 or T-N.b-1.
Thermally decompose nitrogen compounds in an analytical sample using a total nitrogen analyzer by the combustion method to produce nitrogen gas and nitroxide gas. Reduce the nitroxide gas to nitrogen, and measure the nitrogen gas content with a thermal conductivity detector to obtain the total nitrogen (T-N) in an analytical sample. This testing method is also referred to as a modified Dumas’ method. In addition, the performance of this testing method is shown in Comment 4.

(2) Instruments: Instruments are as shown below:
   a) Total nitrogen analyzer by the combustion method: A total nitrogen analyzer configured on the basis of the principle of the combustion method (modified Dumas’ method).
      1) Turn on the total nitrogen analyzer by the combustion method (1), and adjust so that stable indicated values can be obtained.
         (i) Combustion gas: Oxygen having purity no less than 99.99 % (volume percentage)
         (ii) Carrier gas: Helium having purity no less than 99.99 % (volume percentage)

(3) Measurement: Conduct measurement as shown below. However, confirm in advance using an analytical sample that there is no difference from the measured value of total nitrogen obtained according to 4.1.1.a, 4.1.1.c, 4.1.1.d or 4.1.1.e.
   a) Measurement conditions for the total nitrogen analyzer by the combustion method: Set up the measurement conditions for the total nitrogen analyzer considering the following:
      Combustion temperature: No less than 870 ºC
   b) Calibration curve preparation
      1) Turn on the total nitrogen analyzer by the combustion method (1), and adjust so that stable indicated values can be obtained.
      2) Weigh a predetermined amount of the standard for calibration curves (2) to the order of 0.1 mg into a combustion vessel.
      3) Insert the combustion vessel into the total nitrogen analyzer by the combustion method, and read the indicated value.
      4) Conduct the procedure in 3) for another combustion vessel for a blank test, and read the indicated value.
      5) Prepare a curve for the relationship between the nitrogen content and the indicated value of the standard for calibration curves and the blank test for calibration curves.
   c) Sample measurement
      1) Weigh a predetermined amount of an analytical sample to the order of 0.1 mg into a combustion vessel.
      2) Insert the combustion vessel containing the analytical sample to the total nitrogen analyzer by the combustion method, and read the indicated value.
      3) Obtain the nitrogen content from the calibration curve, and calculate total nitrogen in the analytical sample.

Note (1) The setup of the program and the parameter of the analyzer are according to the specification and the operation method of the total nitrogen analyzer by the combustion method used.
   (2) Standard for calibration curves: DL-Aspartic acid (purity no less than 99 % (mass fraction)), EDTA (purity no less than 99 % (mass fraction)), hippuric acid (purity no less than 98 % (mass fraction)) or other reagents having equivalent purity recommended by the total nitrogen analyzer by the combustion method used.
Comment 1 Sample an analytical sample from a test sample prepared in 2.3.3 Grinding (3.1) by grinding with a mill until it completely passes through a sieve of 500 µm aperture or from a test sample prepared in 2.3.3 Grinding Comment 1. Additionally, the sampling amount of an analytical sample is as shown in Table 1. In addition, set the sampling amounts of analytical samples considering the estimated content of total nitrogen in the test sample and the measurement range of the total nitrogen analyzer by the combustion method.

<table>
<thead>
<tr>
<th>Type of fertilizers</th>
<th>Sampling amount (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed fertilizers and designated</td>
<td>0.02 - 0.5</td>
</tr>
<tr>
<td>blended fertilizers</td>
<td></td>
</tr>
<tr>
<td>Organic fertilizers and compost</td>
<td>0.05 - 0.5</td>
</tr>
<tr>
<td>Sludge fertilizers</td>
<td>0.05 - 0.5</td>
</tr>
</tbody>
</table>

Comment 2 Compound fertilizers, designated blended fertilizers and nitrolime may have high contents of phosphoric acid (P₂O₅), alkali metals (Na, K), alkaline earth metals (Ca, Mg), etc., causing contamination of packing or damage in quartz parts, etc. To avoid their influences, it is recommended to add tungsten oxide (elemental analysis reagent or heat-treated reagent) to completely cover the analytical sample.

Comment 3 When a sample with a low content of organic compounds, such as mixed fertilizers and designated blended fertilizers etc., and thus with low combustion efficiency is measured, it is recommended to add sucrose to the analytical sample so that the carbon content will be comparable to the standard for calibration curves. Additionally, confirm in advance that sucrose to be used has a nitrogen content that does not affect the measured value of total nitrogen of the analytical sample.

Comment 4 Table 2, in order to evaluate trueness, shows the results of the comparison of measurement values by the Combustion method and the Kjeldahl method with sludge fertilizers, organic fertilizers, and inorganic fertilizers, etc. Table 3 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction) for fluid fertilizers for home gardening, and 0.05 % (mass fraction) for the other fertilizers.

Table 2: Analysis results of comparison test results between methods

<table>
<thead>
<tr>
<th>Symbol of measurement value</th>
<th>Sample Kind</th>
<th>Number of samples</th>
<th>Range of $y_i - y_k$ (%)</th>
<th>Regression coefficient $y = a + bx$</th>
<th>Correlation coefficient $r$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_i$</td>
<td>Sludge fertilizers$^4$</td>
<td>81</td>
<td>0.31~8.35</td>
<td>-0.006</td>
<td>1.018</td>
</tr>
<tr>
<td>$x_j$</td>
<td>Organic fertilizers, etc.$^5$</td>
<td>31</td>
<td>1.10~12.90</td>
<td>0.009</td>
<td>1.012</td>
</tr>
<tr>
<td>$x_k$</td>
<td>Inorganic fertilizers, etc</td>
<td>36</td>
<td>0.60~46.35</td>
<td>0.000</td>
<td>1.004</td>
</tr>
</tbody>
</table>

1) 4.1.1.a Kjeldahl method
2) 4.1.1.b Combustion method
3) Mass fraction
4) Sewage sludge fertilizers, Human waste sludge fertilizers, Industrial sludge fertilizers, Calcined sludge fertilizers, Composted sludge fertilizers
5) Fish meal, Byproduct organic fertilizer of vegetable origin, Compost, Crustose fertilizer meal, Rape seed meal and powdered rape seed meal, etc.
6) Nitrogenous fertilizers, Compound fertilizers, Blended fertilizers, Fluid fertilizers

Table 3: Results and analysis results from a collaborative study for total nitrogen testing method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories$^1$</th>
<th>Mean$^2$ (%)</th>
<th>$s_r^4$ (%)</th>
<th>RSD$^5_r$ (%)</th>
<th>$s_R^6$ (%)</th>
<th>RSD$^7_R$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer (containing nitrate nitrogen)</td>
<td>11</td>
<td>9.32</td>
<td>0.07</td>
<td>0.8</td>
<td>0.25</td>
<td>2.7</td>
</tr>
<tr>
<td>Compound fertilizer (containing urea)</td>
<td>11</td>
<td>18.34</td>
<td>0.06</td>
<td>0.3</td>
<td>0.45</td>
<td>2.5</td>
</tr>
<tr>
<td>Designated blended fertilizer (containing organic fertilizer)</td>
<td>12</td>
<td>14.06</td>
<td>0.12</td>
<td>0.9</td>
<td>0.42</td>
<td>3.0</td>
</tr>
<tr>
<td>Nitrolime</td>
<td>8</td>
<td>19.96</td>
<td>0.07</td>
<td>0.4</td>
<td>0.17</td>
<td>0.8</td>
</tr>
<tr>
<td>Fish meal</td>
<td>10</td>
<td>8.34</td>
<td>0.04</td>
<td>0.4</td>
<td>0.10</td>
<td>1.3</td>
</tr>
<tr>
<td>Steamed wool waste</td>
<td>11</td>
<td>13.42</td>
<td>0.10</td>
<td>0.7</td>
<td>0.26</td>
<td>2.0</td>
</tr>
<tr>
<td>Rape seed meal and powdered rape seed meal</td>
<td>11</td>
<td>6.21</td>
<td>0.07</td>
<td>1.1</td>
<td>0.25</td>
<td>4.0</td>
</tr>
<tr>
<td>Composted sludge fertilizer A</td>
<td>13</td>
<td>6.20</td>
<td>0.02</td>
<td>0.3</td>
<td>0.09</td>
<td>1.4</td>
</tr>
<tr>
<td>Composted sludge fertilizer B</td>
<td>12</td>
<td>2.36</td>
<td>0.01</td>
<td>0.6</td>
<td>0.04</td>
<td>1.8</td>
</tr>
<tr>
<td>Human waste sludge fertilizer</td>
<td>11</td>
<td>4.44</td>
<td>0.02</td>
<td>0.4</td>
<td>0.06</td>
<td>1.3</td>
</tr>
<tr>
<td>Industrial sludge fertilizer</td>
<td>11</td>
<td>8.06</td>
<td>0.03</td>
<td>0.4</td>
<td>0.07</td>
<td>0.9</td>
</tr>
<tr>
<td>Calcined sludge fertilizer</td>
<td>13</td>
<td>8.80</td>
<td>0.02</td>
<td>2.8</td>
<td>0.03</td>
<td>4.3</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n = number of laboratories \times number of samples (2)$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

40
References

(4) Flow sheet for total nitrogen: The flow sheet for total nitrogen in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Analytical sample</th>
<th>Weigh to the order of 1 mg into a combustion vessel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measurement</td>
<td>Total nitrogen analyzer by the combustion method</td>
</tr>
</tbody>
</table>

Figure Flow sheet for total nitrogen in fertilizers by the combustion method.
Reference: Chromatograms of the standard for calibration curves and an analytical sample are shown below:

1) Standard for calibration curves (DL-aspartic acid)

2) Analytical sample (sludge fertilizer)

Reference figures Chromatograms of total nitrogen.

Measurement conditions for total nitrogen analyzer by the combustion method:
- Combustion gas: Highly pure oxygen, purity no less than 99.9999 % (volume fraction), flow rate 200 mL/min
- Carrier gas: Highly pure helium, purity no less than 99.9999 % (volume fraction), flow rate 80 mL/min
- Separation column: A silica gel stainless column (1m)
- Detector: Thermal conductivity detector (TCD)
- Measurement cycle: Purge time = 60 seconds, circulation combustion time = 200 seconds, measurement time = 100 seconds
- Current value of Detector: 160 mA
- Temperature conditions: Reaction furnace temperature: 870 °C
  - Reaction furnace temperature: 600 °C
  - Column oven temperature: 70 °C
  - Detector temperature: 100 °C
4.1.1.c Devarda’s alloy – Kjeldahl method

(1) Summary

The method is applicable to the fertilizers that contain nitrate nitrogen (N-N) and guarantee total nitrogen. This testing method is classified as Type E and its symbol is 4.1.1.c-2017 or T-N.c-1.

Add hydrochloric acid (1+1) and tin (II) chloride dihydrate to an analytical sample and further add devarda’s alloy to reduce nitrate nitrogen (N-N), and then add sulfuric acid (1+1), pretreat by Kjeldahl method to change total nitrogen (T-N) to ammonium ion and add sodium hydroxide to subject to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain the total nitrogen (T-N) in an analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain the total nitrogen (T-N) in an analytical sample. This testing method corresponds to the devarda’s alloy - sulfuric acid method in the Official Methods of Analysis of Fertilizers (1992).

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1). Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
W_1 \times A \times 0.01/97.095 \times (V_1/V_2) \times (1000/V_3) \times (1/C_1)
\]

\[W_1: \text{Mass (g) of sulfamic acid sampled} \]
\[A: \text{Purity (% (mass fraction)) of sulfamic acid} \]
\[V_1: \text{Volume (mL) of sulfamic acid solution transferred} \]
\[V_2: \text{Constant volume (250 mL) of sulfamic acid solution} \]
\[V_3: \text{Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration} \]
\[C_1: \text{Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution} \]

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid (1)(2). Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

Standardization: Transfer a predetermined amount (3) of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium
hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1).

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid
\[ V_4/V_5 \quad \quad \quad \quad (1) \]

Factor of 0.25 mol/L sulfuric acid \( f_2 \)
\[ (f_1 \times C_1 \times V_4/V_5)/(C_2 \times 2) \quad \quad \quad \quad (2) \]

\( V_4 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
\( V_5 \): Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization
\( C_1 \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
\( C_2 \): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

d) **Boric acid solution (40 g/L)**: Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

e) **Hydrochloric acid**: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

f) **Tin (II) chloride dihydrate**: A JIS Guaranteed Reagent specified in JIS K 8136 or a reagent of mercury analysis grade or equivalent quality.

g) **Devarda’s alloy**: A reagent of nitrogen analysis grade specified in JIS K 8653 or a reagent of equivalent quality.

h) **Sodium hydroxide solution (200 g/L - 500 g/L)**: Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

i) **Bromothymol blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.

j) **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

k) **Methylene blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

l) **Methyl red–methylene blue mixture solution**: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

m) **Bromocresol green solution (0.5 g/100 mL)**: Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

n) **Methyl red–bromocresol green mixture solution**: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

Note  
(1) This is an example of preparation; prepare an amount as appropriate.
(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).
(3) 5 mL - 10 mL
(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.

a) **Steam distillation apparatus**

b) **Digestion flask**: Kjeldahl flask
c) **Distillation flask**: A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

**Comment 2** 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.

(4) **Test procedures**

(4.1) **Reduction and Kjeldahl method**: Conduct reduction and digestion as shown below:

a) Weigh 0.5 g - 1 g (no more than the equivalents of N-N 50 mg) of an analytical sample to the order of 1 mg, and put it in a 300-mL- 500-mL digestion flask.

b) Add 60 mL of hydrochloric acid (1+1) and 2 g of tin (II) chloride dihydrate, and shake to mix and leave at rest for about 20 minutes.

c) Add 3.5 g of devarda’s alloy and leave at rest for about 40 minutes while sometimes shaking to mix.

d) Add 70 mL of sulfuric acid (1+1) and one boiling stone as necessary, and heat at low temperature.

 e) As soon as white smoke start evolving, strengthen heating gradually and further continue heating for about 90 minutes.

f) After standing to cool, add 100 mL - 200 mL of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water, and further mix by shaking.

g) After cooling is complete, add water up to the marked line to make the digestion solution.

**Note** (5) In the case of direct distillation, a 500-mL Kjeldahl flask connectable to a steam distillation apparatus is preferable.

(6) If the bubbles foam strongly and excessively, suspend heating for a little while.

(7) When the entire sample solution volume is used in measurement, it is not necessary to precisely adjust.

(4.2) **Distillation**: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.

a) Transfer a predetermined amount of 0.25 mol/L sulfuric acid to an acceptor, add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount of boric acid solution (40 g/L) to an acceptor, add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.

b) Transfer a predetermined amount of the digestion solution to a 300-mL distillation flask, add a proper amount of sodium hydroxide solution (200 g/L - 500 g/L), and connect this distillation flask to the steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

**Note** (8) 5 mL - 20 mL

(9) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be
immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).

(10) An amount sufficient to make the solution strong alkalinity. A blue color will appear.

(4.3) Measurement: Conduct measurement as shown below.
(4.3.1) When 0.25 mol/L sulfuric acid is used in (4.2):

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4).

b) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\text{Total nitrogen (T-N) (% (mass fraction)) in the analytical sample} = (B \times V_6 - V_7) \times C_1 \times f_1 \times (V_8/V_9) \times (14.007/W_2) \times (100/1000)
\]

- \( B \): Volume of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid
- \( V_6 \): Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)
- \( V_7 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- \( C_1 \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \( f_1 \): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \( V_8 \): Predetermined volume (mL) of the digestion solution in (4.1) g)
- \( V_9 \): Transferred amount (mL) of the digestion solution subjected to distillation in (4.2) b)
- \( W_2 \): Mass (g) of the analytical sample

(4.3.2) When a boric acid solution (40 g/L) is used in (4.2):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red (11).

b) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\text{Total nitrogen (T-N) (% (mass fraction)) in the analytical sample} = V_{10} \times C_2 \times 2 \times f_2 \times (V_{11}/V_{12}) \times (14.007/W_3) \times (100/1000)
\]

- \( V_{10} \): Volume (mL) of 0.25 mol/L sulfuric acid needed for titration
- \( C_2 \): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid
- \( f_2 \): Factor of 0.25 mol/L sulfuric acid
- \( V_{11} \): Predetermined volume (mL) of the digestion solution in (4.1) g)
- \( V_{12} \): Transferred amount (mL) of the digestion solution subjected to distillation in (4.2) b)
- \( W_3 \): Mass (g) of the analytical sample

Note (11) The endpoint is reached when the color changes from green to light red.

Comment 3 The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.3) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

References
1) Masayoshi KOSHINO: Second Revision of The Methods of Analysis of Fertilizers (Details), p.31-33, Yokendo, Tokyo (1988)
(5) **Flow sheet for total nitrogen**: The flow sheet for total nitrogen in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Operation</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weigh to the order of 1 mg into a 300-mL - 500-mL digestion flask.</td>
<td></td>
</tr>
<tr>
<td>60 mL of hydrochloric acid (1+1)</td>
<td>60 mL</td>
</tr>
<tr>
<td>2 g of tin (II) chloride dihydrate</td>
<td></td>
</tr>
<tr>
<td>3.5 g devardar's alloy</td>
<td>20 min</td>
</tr>
<tr>
<td>70 mL sulfuric acid (1+1)</td>
<td>40 min</td>
</tr>
<tr>
<td>1 boiling stone</td>
<td></td>
</tr>
<tr>
<td>Heat at low temperature, and as soon as white smoke evolves, strengthen heating gradually and further heat for about 90 minutes.</td>
<td></td>
</tr>
<tr>
<td>100 mL - 200 mL Water</td>
<td></td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask, water</td>
</tr>
<tr>
<td>Standing to cool</td>
<td></td>
</tr>
<tr>
<td>250 mL - 500 mL Water</td>
<td></td>
</tr>
<tr>
<td>Cooling</td>
<td></td>
</tr>
<tr>
<td>Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Digiton</td>
<td></td>
</tr>
<tr>
<td>Solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1: The flow sheet for total nitrogen in fertilizers (Reduction and Kjeldahl method procedure)

Figure 2 Flow sheet for total nitrogen in fertilizers (Distillation and measurement procedure).

- **Digestion Solution**
  - **Aliquot (predetermined volume)**
    - 300-mL distillation flask
    - Sodium hydroxide (200 g/L - 500 g/L)
      - Receiver: 200-mL - 300-mL Erlenmeyer flask or beaker
        - A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution
    - Steam distillation apparatus
      - Distillation rate: 5 mL/min - 7 mL/min
      - 120 mL – 160 mL distillate
        - Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)
  - Titration
    - 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green, or
    - 0.25 mol/L sulfuric acid (until the solution becomes light red)
4.1.1.d Reduced iron - Kjeldahl method

(1) Summary
The method is applicable to the fertilizers that contain nitrate nitrogen (N-N) and guarantee total nitrogen. This testing method is classified as Type E and its symbol is 4.1.1.d-2017 or TN.d.1.

Add water, reduced iron and sulfuric acid (1+1) to an analytical sample to reduce nitrate nitrogen (N-N) and heat at low temperature, and then add sulfuric acid and pretreat by Kjeldahl method to change total nitrogen (T-N) to ammonium ion, and add a sodium hydroxide solution to subject to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain the total nitrogen (T-N) in an analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain the total nitrogen (T-N) in an analytical sample. This testing method corresponds to the reduced iron - sulfuric acid method in the Official Methods of Analysis of Fertilizers (1992).

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (f)} = \left( W_1 \times A \times 0.01/97.095 \right) \times \left( V_1/V_2 \right) \times \left( 1000/V_3 \right) \times (1/C_1)
\]

\[ W_1: \text{Mass (g) of sulfamic acid sampled} \]
\[ A: \text{Purity (% (mass fraction)) of sulfamic acid} \]
\[ V_1: \text{Volume (mL) of sulfamic acid solution transferred} \]
\[ V_2: \text{Constant volume (250 mL) of sulfamic acid solution} \]
\[ V_3: \text{Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration} \]
\[ C_1: \text{Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution} \]

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid (1)(2): Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

Standardization: Transfer a predetermined amount (3) of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula.
(1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

Volume (B) of 0.1 mol/L -0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\[ = \frac{V_4}{V_5} \]  ............  (1)

Factor of 0.25 mol/L sulfuric acid \((f_2)\)

\[ = \frac{(f_1 \times C_1 \times V_4)/(C_2 \times 2)}{C_2} \]  ............  (2)

\(V_4\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\(V_5\): Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization

\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(C_2\): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

d) **Boric acid solution (40 g/L):** Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

e) **Reduced iron:** Nitrogen content is no more than 0.005 % (mass fraction)

f) **Catalyst**: Mix potassium sulfate specified in JIS K 8962 and copper (II) sulfate pentahydrate (6) specified in JIS K 8983 in the ratio of 9 to 1.

g) **Sodium hydroxide solution (200 g/L - 500 g/L)**: Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

h) **Bromothymol blue solution (0.1 g/100 mL):** Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8842, add water to make 100 mL.

i) **Methyl red solution (0.1 g/100 mL):** Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

j) **Methylene blue solution (0.1 g/100 mL):** Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

k) **Methyl red–methylene blue mixture solution:** To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

l) **Bromocresol green solution (0.5 g/100 mL):** Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

m) **Methyl red–bromocresol green mixture solution:** To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

**Note**  
(1) This is an example of preparation; prepare an amount as appropriate.  
(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).  
(3) 5 mL -10 mL  
(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.  
(5) A tablet is commercially available.  
(6) Crush into powder as appropriate.

(3) **Instruments:** Instruments are as shown below:  
a) **Steam distillation apparatus**  
b) **Digestion flask:** Kjeldahl flask  
c) **Distillation flask:** A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.
Comment 1 A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

Comment 2 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.

(4) Test procedures

(4.1) Reduction and Kjeldahl method: Conduct reduction and digestion as shown below:

a) Weigh 0.5 g - 1 g of an analytical sample to the order of 1 mg, and put it in a 300-mL - 500-mL digestion flask.

b) Add 30 mL of water and mix well.

c) As soon as 5 g of reduced iron and 30 mL of sulfuric acid (1+1) are added, insert a long stem funnel to a digestion flask and shake to mix gently while cooling the outside of the container under flowing water (7).

d) Leave at rest for about 5 minutes (8), boil in a low flame for about 15 minutes.

e) After standing to cool, add 5 g - 10 g of catalyst, 30 mL of sulfuric acid and, if necessary, one boiling stone, heat gradually until water evaporates and white smoke of sulfuric acid evolves (9).

f) Ignite until it is completely digested (10).

g) After standing to cool, add a small amount of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water, and further mix by shaking.

h) After cooling is complete, add water up to the marked line to make the digestion solution.

Note  (7) A sudden reaction generates heat, and unreacted nitric acid vaporizes or digests to make nitrogen oxide etc. through which process losses occur easily. Careful and efficient operation should be taken

(8) Until a sudden reaction is settled.

(9) If the bubbles foam strongly and excessively, suspend heating for a little while.

(10) When the solution has finished changing color, heat further for no less than 2 hours.

(4.2) Distillation: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.

a) Transfer a predetermined amount (11) of 0.25 mol/L sulfuric acid to an acceptor (12), add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount (11) of boric acid solution (40 g/L) to an acceptor (12), add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.

b) Transfer a predetermined amount of the digestion solution to a 300-mL distillation flask, add a proper amount of sodium hydroxide solution (200 g/L - 500 g/L) (13), and connect this distillation flask to the steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

Note  (11) 5 mL - 20 mL

(12) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL
beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).

(13) An amount sufficient to make the solution strong alkalinity. A blue or reddish-brown color will appear.

(4.3) Measurement: Conduct measurement as shown below.

(4.3.1) When 0.25 mol/L sulfuric acid is used in (4.2):

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green.

b) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\text{Total nitrogen (T-N) (% (mass fraction)) in the analytical sample} = (B \times V_6 - V_7) \times C_1 \times f_1 \times (V_6/V_9) \times \left(14.007/W_2\right) \times (100/1000)
\]

\[
B: \quad \text{Volume of 0.1 mol/L -0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid}
\]

\[
V_6: \quad \text{Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)}
\]

\[
V_7: \quad \text{Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration}
\]

\[
C_1: \quad \text{Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution}
\]

\[
f_1: \quad \text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution}
\]

\[
V_8: \quad \text{Predetermined volume (mL) of the digestion solution in (4.1) h)}
\]

\[
V_9: \quad \text{Transferred amount (mL) of the digestion solution subjected to distillation in (4.2) b)}
\]

\[
W_2: \quad \text{Mass (g) of the analytical sample}
\]

(4.3.2) When a boric acid solution (40 g/L) is used in (4.2):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red.

b) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\text{Total nitrogen (T-N) (% (mass fraction)) in the analytical sample} = V_{10} \times C_2 \times f_2 \times (V_{11}/V_{12}) \times \left(14.007/W_3\right) \times (100/1000)
\]

\[
V_{10}: \quad \text{Volume (mL) of 0.25 mol/L sulfuric acid needed for titration}
\]

\[
C_2: \quad \text{Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid}
\]

\[
f_2: \quad \text{Factor of 0.25 mol/L sulfuric acid}
\]

\[
V_{11}: \quad \text{Predetermined volume (mL) of the digestion solution in (4.1) h)}
\]

\[
V_{12}: \quad \text{Transferred amount (mL) of the digestion solution subjected to distillation in (4.2) b)}
\]

\[
W_3: \quad \text{Mass (g) of the analytical sample}
\]

Note (14) The endpoint is reached when the color changes from green to light red.

Comment 3 The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.3) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.
(5) **Flow sheet for total nitrogen**: The flow sheet for total nitrogen in fertilizers is shown below:

![Flow sheet for total nitrogen](image)

**Figure 1** The flow sheet for total nitrogen in fertilizers (Reduction and Kjeldahl method procedure)
A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution.

Distillation rate: 5 mL/min - 7 mL/min

120 mL – 160 mL distillate

0.1mol/L - 0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or
0.25 mol/L sulfuric acid (until the solution becomes light red)

Figure 2 Flow sheet for total nitrogen in fertilizers (Distillation and measurement procedure).
4.1.1.e Calculation with ammoniac nitrogen and nitrate nitrogen

(1) Summary
This testing method is applicable to the fertilizers that contain ammoniac nitrogen (A-N) and nitrate nitrogen (N-N), and that does not contain the fertilizers guaranteeing total nitrogen (T-N). This testing method is classified as Type A (Def-C) and its symbol is 4.1.1.e-2017 or T-N.e-1. Calculate total nitrogen (T-N) by adding the ammoniac nitrogen (A-N) obtained in 4.1.2 to the nitrate nitrogen (N-N) obtained in 4.1.3.

(2) The calculation of total nitrogen
a) Calculate the total nitrogen (T-N) in the analytical sample by the following formula:

\[
\text{Total nitrogen (T-N) } \left( \% \text{ (mass fraction)} \right) \text{ in the analytical sample} = (\text{A-N}) + (\text{N-N})
\]

A-N: Ammoniac nitrogen (\% (mass fraction)) in the analytical sample \(^{(1)}\) obtained by 4.1.2
N-N: Nitrate nitrogen (\% (mass fraction)) in the analytical sample \(^{(1)}\) obtained by 4.1.3

Note (1) A-N and N-N use raw data without rounding numerical value
4.1.2 Ammoniac nitrogen

4.1.2.a Distillation method

(1) Summary

The testing method is applicable to the fertilizers that contain ammonium salt. However, in some cases, it is not applicable to the fertilizers that contain such compounds as nitrolime that digests by heating. This testing method is classified as Type C and its symbol is 4.1.2.a-2017 or A-N.a-1.

Add water to an analytical sample, further add magnesium oxide or a sodium hydroxide solution to make the solution alkalinity and subject it to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain ammoniacal nitrogen (A-N) in an analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain ammoniacal nitrogen (A-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 7.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution\(^{(1)}\): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

**Standardization:** Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line\(^{(1)}\). Transfer a predetermined amount of the solution to a 200-mL -300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution } (f_1) = \frac{(W_1 \times A \times 0.01/97.095 \times (V_1/V_2) \times (1000/V_3) \times (1/C_1)}{V_1} \times \frac{1}{V_3}
\]

\(W_1: \) Mass (g) of sulfamic acid sampled

\(A: \) Purity (% (mass fraction)) of sulfamic acid

\(V_1: \) Volume (mL) of sulfamic acid solution transferred

\(V_2: \) Constant volume (250 mL) of sulfamic acid solution

\(V_3: \) Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\(C_1: \) Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Magnesium oxide: A JIS Guaranteed Reagent specified in JIS K 8432 or a reagent of equivalent quality.

c) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

d) 0.25 mol/L sulfuric acid \(^{(1)}(2)\): Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

**Standardization:** Transfer a predetermined amount \(^{(3)}\) of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the
solution becomes gray-green \(^{(4)}\). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid  
\[ V_4/V_5 \]  ........... (1)  

Factor of 0.25 mol/L sulfuric acid \((f_2)\)  
\[ f_2 = (f_1 \times C_1 \times V_4/V_5)/(C_2 \times 2) \]  ........... (2)  

\(V_4\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration  
\(V_5\): Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization  
\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution  
\(C_2\): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid  

- **Boric acid solution (40 g/L)**: Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.  
- **Sodium hydroxide solution (200 g/L - 500 g/L)** \(^{(1)}\): Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.  
- **Bromothymol blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.  
- **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.  
- **Methylene blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.  
- **Methyl red–methylene blue mixture solution**: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).  
- **Bromocresol green solution (0.5 g/100 mL)**: Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.  
- **Methyl red–bromocresol green mixture solution**: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).  

**Note**  
(1) This is an example of preparation; prepare an amount as appropriate.  
(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).  
(3) 5 mL -10 mL  
(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.  

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.  
**Comment 2** 0.25 mol/L sulfuric acid in (2) d) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.  

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.  
  a) **Steam distillation apparatus**  
  b) **Distillation flask**: A Kjeldahl flask or round bottom flask that can be connected to a steam
(4) Test procedures

(4.1) Sample solution preparation: Prepare a sample solution as shown below:

a) Weigh 0.25 g - 2 g (the equivalents of 20 mg - 100 mg as N) of an analytical sample to the order of 1 mg, and put it in a 300-mL - 500-mL distillation flask.

b) Add about 25 mL of water to make a sample solution.

Note (5) The sampling amount of the analytical sample is 5 g when there is less nitrogen content in the fertilizers such as a home garden-use fertilizer.

Comment 3 When it is a fertilizer which contains uric ammonium acid, humus acid ammonium or nitrate nitrogen, etc., or when it is not a fertilizer in which phosphate, ammonium and magnesium coexist, conduct the procedure in (4.1.1) a) - c) in 4.2.4.a or the procedure in (4.1.2) a) - c) in 3.2.4.a and transfer a predetermined amount of suspension (the equivalents of 20 mg - 100 mg as N) to a 300-mL - 500-mL distillation flask to make a sample solution.

(4.2) Distillation: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.

a) Transfer a predetermined amount of 0.25 mol/L sulfuric acid to an acceptor, add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount of boric acid solution (40 g/L) to an acceptor, add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.

b) Add no less than 2 g of magnesium oxide to the distillation flask, and connect this distillation flask to a steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

Note (6) 5 mL - 20 mL

(7) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).

(8) An amount sufficient to make the solution strong alkalinity.

(9) Add a small amount of silicone oil as necessary.

Comment 4 When the sample does not contain organic matters or urea, a proper amount of sodium hydroxide solution (200 g/L - 500 g/L) (8) can be added instead of magnesium oxide.

(4.3) Measurement: Conduct measurement as shown below.

(4.3.1) When 0.25 mol/L sulfuric acid is used in (4.2):

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green.

b) Calculate the ammoniacal nitrogen (A-N) in the analytical sample by the following formula:
Ammoniacal nitrogen (A-N) (% (mass fraction)) in the analytical sample

\[
\text{Ammoniacal nitrogen (A-N) (%) in the analytical sample} = (B \times V_6 - V_7) \times C_1 \times f_1 \times (14.007/W_2) \times (100/1000)
\]

\[
= (B \times V_6 - V_7) \times C_1 \times f_1 \times (1.4007/W_2)
\]

**B**: Volume of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

**V_6**: Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)

**V_7**: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

**C_1**: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

**f_1**: Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

**W_2**: Mass (g) of the analytical sample

(4.3.2) When a boric acid solution (40 g/L) is used in (4.2):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red\(^{(10)}\).

b) Calculate the ammoniacal nitrogen (A-N) in the analytical sample by the following formula:

\[
\text{Ammoniacal nitrogen (A-N) (%) in the analytical sample} = V_8 \times C_2 \times 2 \times f_2 \times (14.007/W_3) \times (100/1000)
\]

\[
= V_8 \times C_2 \times f_2 \times (2.8014/W_3)
\]

**V_8**: Volume (mL) of 0.25 mol/L sulfuric acid needed for titration

**C_2**: Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

**f_2**: Factor of 0.25 mol/L sulfuric acid

**W_3**: Mass (g) of the analytical sample

**Note** \(^{(10)}\) The endpoint is reached when the color changes from green to light red.

**Comment 5** If it is hard to confirm the endpoint due to the carbon dioxide resulting from carbonate in the extract when magnesium oxide is used, it is recommended to boil the extract for 1-2 minute(s) after distilling and cool, and then titrate.

**Comment 6** The titration procedures in (2) a) **Standardization**, (2) d) **Standardization** and (4.3) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

**Comment 7** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 21 % (mass fraction) and 1 % (mass fraction) are 102.2 % - 100.8 % and 102.5 % as ammoniacal nitrogen (A-N) respectively. The results of the collaborative study (limited to reported values with Distillation method) to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.1 % (mass fraction) for solid fertilizers, and 0.01 % (mass fraction) for fluid fertilizers.
Table 1: Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratories</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
<th>Reproducibility</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Repeatability</td>
<td>Intermediate precision</td>
<td>Reproducibility</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Average $^2)$</td>
<td>$s_r$ $^4)$</td>
<td>RSD$_r$ $^5)$</td>
</tr>
<tr>
<td>FAMIC-B-10</td>
<td>11</td>
<td>8.38</td>
<td>0.09</td>
<td>1.0</td>
</tr>
<tr>
<td>FAMIC-B-14</td>
<td>11</td>
<td>8.06</td>
<td>0.03</td>
<td>0.4</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Distillation method
2) Average (the number of laboratory ($p$) × test days ($2$) × the number of replicate testing ($3$))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

1) Masayoshi KOSHINO: Second Revision of The Methods of Analysis of Fertilizers (Details), p.36-37, Yokendo, Tokyo (1988)
(5) **Flow sheet for ammoniac nitrogen:** The flow sheet for ammoniac nitrogen in fertilizers is shown below.

0.25 g - 2 g analytical sample

- Weigh the equivalents of 20 mg – 100 mg as N to the order of 1 mg, and put it in a 300-mL distillation flask
- About 25 mL water
- 2 g Magnesium oxide to a distillation flask
- A small amount of defoaming agent as necessary

Steam distillation apparatus

- Receiver: 200-mL - 300-mL Erlenmeyer flask or beaker
- A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution
- Distillation rate: 5 mL/min - 7 mL/min

Stop distilling

- 120 mL – 160 mL distillate
- Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)

Titration

- 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or
- 0.25 mol/L sulfuric acid (until the solution becomes light red)

Figure  Flow sheet for ammoniac nitrogen in fertilizers.
4.1.2.b Formaldehyde method

(1) Summary

The testing method is applicable to the fertilizers which do not contain a large amount of flora and fauna sample. This testing method is classified as Type C and its symbol is 4.1.2.b-2017 or A-N.b-1. After adding water or hydrochloric acid (1+20) to an analytical sample to extract ammonium ion, add an aluminum chloride solution, drop a potassium hydroxide solution and precipitate phosphate and excessive aluminum to make a sample solution. Adjust the sample solution to slight acidity, add a formaldehyde solution and measure ammonium ion by complexometric titration with 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain the ammoniacal nitrogen (A-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (f)} = \left(\frac{W_1 \times A \times 0.01/97.095}{(V_1/V_2) \times (1000/V_3)} \times \frac{1}{C}\right)
\]

\[W_1\]: Mass (g) of sulfamic acid sampled

\[A\]: Purity (% (mass fraction)) of sulfamic acid

\[V_1\]: Volume (mL) of sulfamic acid solution transferred

\[V_2\]: Constant volume (250 mL) of sulfamic acid solution

\[V_3\]: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\[C\]: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Potassium chloride solution (1 mol/L) (1): Dissolve 75 g of potassium chloride specified in JIS K 8121 in water to make 1000 mL.

c) Aluminum chloride solution (1 mol/L) (1): Dissolve 240 g of aluminum chloride (III) hexahydrate specified in JIS K 8114 in water to make 1000 mL.

d) Potassium hydroxide solution (170 g/L) (1): Dissolve 170 g of potassium hydroxide in water to make 1000 mL.

e) Formaldehyde solution: Add one volume of water to one volume of the 36 % (mass fraction) - 38 % (mass fraction) formaldehyde specified in JIS K 8872.

f) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

g) Bromothymol blue solution (0.1 g/100 mL): Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, and add water to make 100
mL.

h) Methyl red solution (0.1 g/100 mL): Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

i) Bromothymol blue solution (1 g/100 mL): Dissolve 1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, and add water to make 100 mL.

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

Comment 2 Thymol blue can be dissolved if it is sodium salt. The thymol blue specified in JIS K 8643 is slightly hard to dissolve in ethanol and hard to dissolve in water. Therefore, add about 2.15 mL of sodium hydroxide solution (0.1 mol/L) per 0.1 g of thymol blue to neutralize, and then prepare the thymol blue solution (1 g/100 mL) through the same procedure as (2) i).

(3) Instruments: Instruments are as shown below:
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Ammonium salts
   a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

Comment 3 The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.4.a.

(4.1.2) Mixed fertilizers
   a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.
   b) Add about 300 mL of hydrochloric acid (1 + 20) and shake to mix at the rate of 30 - 40 revolutions/min for about 30 minutes.
   c) Add an aluminum chloride solution (1 mol/L) (2) to the solution and add a few drops of methyl red solution (1mol/L) as an indicator. After that, add immediately potassium hydroxide (170 g/L) while shaking the flask until the color of the solution changes to light yellow (3).
   d) Add water up to the marked line.
   e) Filter with Type 3 filter paper to make a sample solution.

Note (2) Add 3 mL of aluminum chloride per 0.04 g of P or 0.1 g of P₂O₅ in the sample solution.
(3) Form precipitate of aluminum hydroxide and aluminum phosphate to separate the phosphate.

Comment 4 In the procedure of (4.1.1) a) and (4.1.2) a), it is also allowed to weigh 2.5g of an analytical sample to the order of 1 mg, and put it in a 250-mL volumetric flask
Comment 5 When it is not a fertilizer in which phosphate, ammonium and magnesium coexist, about 400 mL of potassium chloride solution (1 mol/L) can be used instead of about 300 mL of hydrochloric acid (1+20) in the procedure of (4.1.2) b).

Comment 6 In the case of the mixed fertilizers containing bentonite, after shaking to mix using about 300 mL of hydrochloric acid (1+20) or about 400 mL of potassium chloride solution (1 mol/L) according to Comment 5 in (4.1.2) b), add water to the marked line, filter with Type 3 filter paper and transfer 50 ml - 100 mL to a 250-mL volumetric flask, and then conduct the procedure in (4.1.2) c) - e).

(4.2) Measurement: Conduct measurement as shown below.

a) Transfer a predetermined amount of sample solution (the equivalents of 50 mg as A-N) to a 300-mL volumetric flask (4).

b) Add water to the solution to make about 100 mL.

c) Add one or two drop(s) of methyl red solution (0.1 g/100 mL) and add hydrochloric acid (1+200) until the color of the solution changes to light pink.

d) Add 10 mL of formaldehyde solution.

e) Add one or two drop(s) of thymol blue solution (1 g/100 mL) and titrate with 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution changes to blue (5).

f) As a blank test, transfer 100 mL of water to another 300-mL Erlenmeyer flask and conduct the procedure in c) - e).

g) Calculate the ammoniacal nitrogen (A-N) in the analytical sample by the following formula:

Ammoniacal nitrogen (A-N) (% (mass fraction)) in the analytical sample

\[ \frac{(V_S - V_B) \times C \times f \times (V_1/V_2) \times (14.007/W_2) \times (100/1000)}{W_2} \]

Where:

- \( V_S \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration in (4.2) e)
- \( V_B \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration of the blank test in (4.2) f)
- \( C \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \( f \): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \( V_1 \): Predetermined volume (mL) of the sample solution in (4.1.1) c) or (4.1.2) d)
- \( V_2 \): Transferred amount (mL) of the sample solution in (4.2) a)
- \( W_2 \): Mass (g) of the analytical sample

Note (4) The volume to be transferred should be up to 100 mL.

(5) The endpoint is reached when the color changes from green to light red. It is easy to observe the change of color under fluorescent light.

Comment 7 The titration procedures in (2) a) Standardization and (4.2) e) - f) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

Comment 8 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 21 % (mass fraction) and 1 % (mass fraction) are 100.4 % - 101.0 % and 100.1 % as ammoniacal nitrogen (A-N) respectively.

The results of the collaborative study (limited to reported values by Formaldehyde...
method) to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.03 % (mass fraction) for solid fertilizers, and 0.02 % (mass fraction) for fluid fertilizers.

<table>
<thead>
<tr>
<th>Name of certified reference material fertilizer</th>
<th>Number of laboratory</th>
<th>Average $^1$</th>
<th>Repeatability $^2$</th>
<th>Intermediate precision $^5$</th>
<th>Reproducibility $^9$</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-A-10</td>
<td>10</td>
<td>10.66</td>
<td>0.07</td>
<td>0.09</td>
<td>0.16</td>
</tr>
<tr>
<td>FAMIC-A-13</td>
<td>9</td>
<td>10.36</td>
<td>0.06</td>
<td>0.08</td>
<td>0.21</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Formaldehyde method
2) Average (the number of laboratory ($p$) × test days (2) × the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References


(5) **Flow sheet for ammoniacal nitrogen**: The flow sheet for ammoniacal nitrogen in fertilizers is shown below.

![Flow sheet for ammoniacal nitrogen](image.png)
5 g analytical sample (compound fertilizer) → Weigh to the order of 1 mg into a 500-mL volumetric flask

<table>
<thead>
<tr>
<th>Shaking to mix</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rotary shaker (30 - 40 revolutions/minute), 30 minutes</td>
</tr>
<tr>
<td>→ About 300 mL of hydrochloric acid (1 + 20)</td>
</tr>
<tr>
<td>→ Aluminum chloride solution (1 mol/L)</td>
</tr>
<tr>
<td>→ One or two drop(s) of methyl red</td>
</tr>
<tr>
<td>→ Potassium hydroxide solution (170 g/L) (Until the solution changes to light yellow)</td>
</tr>
<tr>
<td>→ Water (up to the marked line)</td>
</tr>
</tbody>
</table>

Filtration → Type 3 filter paper

Sample solution

Figure 1-2 The flow sheet for ammoniac nitrogen in fertilizers (Extraction procedure (4.1.2))

Sample solution → Up to the equivalents of 50 mg as A-N

<table>
<thead>
<tr>
<th>Aliquot</th>
</tr>
</thead>
<tbody>
<tr>
<td>300-mL Erlenmeyer flask</td>
</tr>
<tr>
<td>→ Add water to make about 100 mL of solution</td>
</tr>
<tr>
<td>→ One or two drop(s) of methyl red</td>
</tr>
<tr>
<td>→ Hydrochloric acid (1+200) [light pink]</td>
</tr>
<tr>
<td>→ 10 mL of formaldehyde solution</td>
</tr>
<tr>
<td>→ One or two drop(s) of thymol blue solution (1 g/100 mL)</td>
</tr>
</tbody>
</table>

Titration → 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (until the solution becomes blue)

Figure 2 The flow sheet for ammoniac nitrogen in fertilizers (Measurement solution)
4.1.3 Nitrate nitrogen
4.1.3.a Devarda’s alloy - distillation method

(1) Summary
The testing method is applicable to the fertilizer containing nitrate. However, it is not applicable to fertilizers containing urea, nitrolime and organic matters that digest by heating and isolate ammonia. This testing method is classified as Type E and its symbol is 4.1.3.a-2017 or N-N.a-1.

Add water to an analytical sample to dissolve ammoniacal nitrogen (A-N) and nitrate nitrogen (N-N), and further add devarda’s alloy and sodium hydroxide to subject it to steam distillation. In this process, nitrate nitrogen (N-N) is reduced to ammonia. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain nitrogen content (N-N+A-N) in an analytical sample. Subtract separately obtained the ammoniacal nitrogen (A-N) by 4.1.2 to calculate nitrate nitrogen (N-N). This testing method corresponds to the devarda’s alloy method in the Official Methods of Analysis of Fertilizers (1992).

(2) Reagent: Reagents are as shown below.
   a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution: Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.
      Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line. Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (f_1)} = \left( W_1 \times A \times 0.01/97.095 \right) \times \left( V_1/V_2 \right) \times \left( 1000/V_3 \right) \times \left( 1/C_1 \right)
\]

\[\begin{align*}
W_1 & : \text{Mass (g) of sulfamic acid sampled} \\
A & : \text{Purity (% (mass fraction)) of sulfamic acid} \\
V_1 & : \text{Volume (mL) of sulfamic acid solution transferred} \\
V_2 & : \text{Constant volume (250 mL) of sulfamic acid solution} \\
V_3 & : \text{Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration} \\
C_1 & : \text{Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution}
\end{align*}\]

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid: Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

Standardization: Transfer a predetermined amount of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution,
and titrate with a 0.1 mol/L -0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green \(^{(4)}\). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

\[
\text{Volume (B) of 0.1 mol/L -0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid} = \frac{V_4}{V_5} \quad \text{......... (1)}
\]

\[
\text{Factor of 0.25 mol/L sulfuric acid (f_2)} = \frac{(f_1 \times C_1 \times V_4)}{(C_2 \times V_5)} \quad \text{......... (2)}
\]

\(V_4\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\(V_5\): Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization

\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(C_2\): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

d) **Boric acid solution (40 g/L):** Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

e) **Sodium hydroxide solution (200 g/L - 500 g/L) \(^{(1)}\):** Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

f) Devarda’s alloy: A reagent of nitrogen analysis grade specified in JIS K 8653 or a reagent of equivalent quality.

g) **Bromothymol blue solution (0.1 g/100 mL):** Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, and add water to make 100 mL.

h) **Methyl red solution (0.1 g/100 mL):** Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

i) **Methylene blue solution (0.1 g/100 mL):** Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

j) **Methyl red–methylene blue mixture solution:** To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

k) **Bromocresol green solution (0.5 g/100 mL):** Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

l) **Methyl red–bromocresol green mixture solution:** To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

**Note**

(1) This is an example of preparation; prepare an amount as appropriate.

(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).

(3) 5 mL -10 mL

(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

**Comment 2** 0.25 mol/L sulfuric acid in (2) e) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.
(3) **Apparatus and instruments**: Apparatus and instruments are shown below.
   a) **Steam distillation apparatus**
   b) **Distillation flask**: A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.

(4) **Test procedures**

(4.1) **Sample solution preparation**: Prepare a sample solution as shown below:
   a) Weigh 0.25 g - 1 g (the equivalents of 20 mg - 100 mg as N) of an analytical sample to the order of 1 mg, and put it in a 300-mL - 500-mL distillation flask
   b) Add about 25 mL of water to make a sample solution.

**Note** (5) Conduct the procedure in **Comment 3** when there is much nitrogen content in the fertilizers such as simple salt fertilizers.

**Comment 3** In the case of nitrate fertilizer, etc. containing much nitrogen content, weigh 2 g - 5g of an analytical sample to the order of 1 mg, put it into a 250- mL volumetric flask, dissolve it in water, and further add water up to the marked line. Put predetermined volume of suspension (the equivalents of 20 mg -100 mg as N) into a 300- mL - 500-mL distillation flask.

(4.2) **Distillation**: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.
   a) Transfer a predetermined amount (6) of 0.25 mol/L sulfuric acid to an acceptor (7), add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount (6) of boric acid solution (40 g/L) to an acceptor (7), add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.
   b) Add (10) no less than 3 mg of devarda’s alloy and adequate volume of sodium hydroxide (200 g/L - 500 g/L) (8)-(9) and connect this distillation flask to the steam distillation apparatus.
   c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.
   d) Stop distilling when the distillate has reached 120 mL - 160 mL.
   e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

**Note** (6) 5 mL - 20 mL
   (7) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).
   (8) Sudden reaction makes bubbles foam drastically and the bubbles overflow from a distillation flask. Therefore, it is required to add an alkali solution gradually and mix quietly.
   (9) An amount sufficient to make the solution strong alkalinity.
   (10) Add a small amount of silicone oil as necessary.

(4.3) **Measurement**: Conduct measurement as shown below.

(4.3.1) When 0.25 mol/L sulfuric acid is used in (4.2):
   a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4).
b) Calculate nitrogen content (N-N+A-N) in the analytical sample by the following formula:
c) Subtract the ammoniac nitrogen (A-N) separately obtained in 4.1.2 from the obtained nitrogen content (N-N+A-N) to calculate nitrate nitrogen (N-N) \(^{(11)}(12)\).

Nitrogen content (N-N+A-N) (% (mass fraction)) in the analytical sample
\[
= (B \times V_6 - V_7) \times C_1 \times f_1 \times \left(\frac{14.007}{W_2}\right) \times \left(\frac{100}{1000}\right)
\]

\(B\): Volume of 0.1 mol/L -0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid
\(V_6\): Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)
\(V_7\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration in (4.3) a)
\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
\(f_1\): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
\(W_2\): Mass (g) of the analytical sample

Note (11) The nitrogen content (N-N+A-N) and ammoniacal nitrogen (A-N) use raw data without rounding the numerical value.
(12) When no ammoniacal nitrogen (A-N) is contained, the nitrogen content (N-N+A-N) calculated in (4.3) b) is regarded as nitrate nitrogen (N-N).

(4.3.2) When a boric acid solution (40 g/L) is used in (4.2):
a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red (13).
b) Calculate nitrogen content (N-N+A-N) in the analytical sample by the following formula:
c) Subtract the ammoniac nitrogen (A-N) separately obtained in 4.1.2 from the obtained nitrogen content (N-N+A-N) to calculate nitrate nitrogen (N-N) \(^{(11)}(12)\).

Nitrogen content (N-N+A-N) (% (mass fraction)) in the analytical sample
\[
= V_{10} \times C_2 \times 2 \times f_2 \times \left(\frac{V_{11}}{V_{12}}\right) \times \left(\frac{14.007}{W_3}\right) \times \left(\frac{100}{1000}\right)
\]

\(V_{10}\): Volume (mL) of 0.25 mol/L sulfuric acid needed for titration
\(C_2\): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid
\(f_2\): Factor of 0.25 mol/L sulfuric acid
\(W_3\): Mass (g) of the analytical sample

Note (13) The endpoint is reached when the color changes from green to light red.

Comment 4 The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.3) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

References
(5) **Flow sheet for nitrate nitrogen**: The flow sheet for nitrate nitrogen in fertilizers is shown below.

<table>
<thead>
<tr>
<th>0.25 g - 1 g analytical sample</th>
<th>Weigh the equivalents of 20 mg - 100 mg as N to the order of 1 mg into a 300-mL distillation flask.</th>
</tr>
</thead>
<tbody>
<tr>
<td>➔ 25 mL of water</td>
<td>No less than 3 g of devarda's alloy</td>
</tr>
<tr>
<td>➔ Sodium hydroxide solution (200 g/L - 500 g/L)</td>
<td>A small amount of silicone oil as necessary</td>
</tr>
<tr>
<td>Receiver: 200-mL - 300-mL Erlenmeyer flask or beaker</td>
<td>A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution</td>
</tr>
</tbody>
</table>

**Steam distillation apparatus**

Distillation rate: 5 mL/min - 7 mL/min

Stop distilling

120 mL - 160 mL distillate

Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)

| Titration                      | 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or 0.25 mol/L sulfuric acid (until the solution becomes light red) |

Figure: The flow sheet for nitrate nitrogen in fertilizers
4.1.3.b Reduced iron-distillation method

(1) Summary

The testing method is applicable to the fertilizer containing nitrate. However, it is not applicable to fertilizers containing urea, nitrolime and organic matters that digest by heating and isolate ammonia. This testing method is classified as Type E and its symbol is 4.1.3.b-2017 or N-N.b-

Add water to an analytical sample to dissolve ammoniacal nitrogen (A-N) and nitrate nitrogen (N-N), and add reduced iron and a sulfuric acid solution to boil lightly. In this process, nitrate nitrogen (N-N) is reduced to ammonia. And further add a sodium hydroxide solution to distillate. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain nitrogen content (N-N+A-N) in an analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain nitrogen content (N-N+A-N) in an analytical sample. Subtract separately obtained the ammoniacal nitrogen (A-N) by 4.1.2 to calculate nitrate nitrogen (N-N). This testing method corresponds to the reduced iron method in the Official Methods of Analysis of Fertilizers (1992).

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution: Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL - 11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line. Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[ f_1 = \left( \frac{W_1 \times A \times 0.01/97.095}{V_1/V_2} \right) \times \left( \frac{1000}{V_3} \right) \times \left( \frac{1}{C_1} \right) \]

- \( W_1 \): Mass (g) of sulfamic acid sampled
- \( A \): Purity (% (mass fraction)) of sulfamic acid
- \( V_1 \): Volume (mL) of sulfamic acid solution transferred
- \( V_2 \): Constant volume (250 mL) of sulfamic acid solution
- \( V_3 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- \( C_1 \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid: Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

Standardization: Transfer a predetermined amount of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the
solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

$$= \frac{V_4}{V_5} \quad \text{........... (1)}$$

Factor of 0.25 mol/L sulfuric acid ($f_2$)

$$= \frac{f_1 \times C_1 \times V_4}{V_5} \times \frac{C_2 \times 2} {\text{........... (2)}}$$

$V_4$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

$V_5$: Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization

$C_1$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

$C_2$: Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

d) Boric acid solution (40 g/L): Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

e) Sodium hydroxide solution (200 g/L - 500 g/L) (1): Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

f) Reduced iron: Nitrogen content is no more than 0.005 % (mass fraction)

Bromothymol blue solution (0.1 g/100 mL): Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, and add water to make 100 mL.

h) Methyl red solution (0.1 g/100 mL): Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

i) Methylene blue solution (0.1 g/100 mL): Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

j) Methyl red–methylene blue mixture solution: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

k) Bromocresol green solution (0.5 g/100 mL): Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

l) Methyl red–bromocresol green mixture solution: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

Note (1) This is an example of preparation; prepare an amount as appropriate.

(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).

(3) 5 mL -10 mL

(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.

Comment 1 A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

Comment 2 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.

(3) Instruments: Instruments are as shown below:
a) **Steam distillation apparatus**
b) **Distillation flask:** A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.

(4) **Test procedures**

(4.1) **Preparation of sample solution:** Conduct preparation of a sample solution as shown below.

a) Weigh 0.5 g - 1 g \(^{(5)}\) (the equivalents of 20 mg - 100 mg as N) of an analytical sample to the order of 1 mg, and put it in a 300-mL - 500-mL distillation flask

b) Add about 30 mL of water and mix well.

c) As soon as adding 5 g of reduced iron and 10 mL of sulfuric acid \((1+1)\), insert a long stem funnel to a distillation flask and shake to mix gently while cooling the outside of the container under flowing water \(^{(6)}\).

d) After leaving at rest for about 5 minutes \(^{(7)}\), heat gradually by low temperature and boil in a low flame for about 15 minutes, and then stand to cool to make a sample solution.

Note \(^{(5)}\) Conduct the procedure in Comment 3 when there is much nitrogen content in the fertilizers such as simple salt fertilizers.

(6) Sudden reaction generates heat, and unreacted nitric acid vaporizes or digests to make nitrogen oxide etc. through which process losses occur easily. Careful and efficient operation should be taken

(7) Until a sudden reaction is settled.

Comment 3 In the case of nitrate fertilizer, etc. containing much nitrogen content, weigh 2 g - 5g of an analytical sample to the order of 1 mg, put it into a 250- mL volumetric flask, dissolve it in water, and further add water up to the marked line. Put predetermined volume of suspension (the equivalents of 20 mg - 100 mg as N) into a 300-mL - 500-mL distillation flask.

(4.2) **Distillation:** Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.

a) Transfer a predetermined amount \(^{(8)}\) of 0.25 mol/L sulfuric acid to an acceptor \(^{(9)}\), add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount \(^{(8)}\) of boric acid solution \((40 g/L)\) to an acceptor \(^{(9)}\), add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.

b) Add adequate volume of sodium hydroxide \((200 g/L - 500 g/L)\) \(^{(10)}\) and connect this distillation flask to the steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

Note \(^{(8)}\) 5 mL - 20 mL

(9) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution \((40 g/L)\).

(10) An amount sufficient to make the solution strong alkalinity.

(4.3) **Measurement:** Conduct measurement as shown below.
(4.3.1) When 0.25 mol/L sulfuric acid is used in (4.2) a),

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green.

b) Calculate nitrogen content (N-N+A-N) in the analytical sample by the following formula:

c) Subtract the ammoniac nitrogen (A-N) separately obtained in 4.1.2 from the obtained nitrogen content (N-N+A-N) to calculate nitrate nitrogen (N-N) (11)(12).

\[
\text{Nitrogen content (N-N+A-N) (% (mass fraction)) in the analytical sample} = \frac{B \times V_6 - V_7 \times C_1 \times f_1 \times (14.007/W_2) \times (100/1000)}{C_1 \times f_1 \times (1.4007/W_2)}
\]

\(B\): Volume of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\(V_6\): Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)

\(V_7\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(f_1\): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(W_2\): Mass (g) of the analytical sample

Note (11) The nitrogen content (N-N+A-N) and ammoniacal nitrogen (A-N) use raw data without rounding the numerical value.

(12) When no ammoniacal nitrogen (A-N) is contained, the nitrogen content (N-N+A-N) calculated in (4.3) b) is regarded as nitrate nitrogen (N-N).

(4.3.2) When a boric acid solution (40 g/L) is used in (4.2) a):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red.

b) Calculate nitrogen content (N-N+A-N) in the analytical sample by the following formula:

c) Subtract the ammoniac nitrogen (A-N) separately obtained in 4.1.2 from the obtained nitrogen content (N-N+A-N) to calculate nitrate nitrogen (N-N) (11)(12).

\[
\text{Nitrogen content (N-N+A-N) (% (mass fraction)) in the analytical sample} = \frac{V_{10} \times C_2 \times 2 \times f_2 \times (14.007/W_3) \times (100/1000)}{C_2 \times f_2 \times (2.8014/W_3)}
\]

\(V_{10}\): Volume (mL) of 0.25 mol/L sulfuric acid needed for titration

\(C_2\): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

\(f_2\): Factor of 0.25 mol/L sulfuric acid

\(W_3\): Mass (g) of the analytical sample

Note (13) The endpoint is reached when the color changes from green to light red.

Comment 4 The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.3) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

References

1) Masayoshi KOSHINO: Second Revision of The Methods of Analysis of Fertilizers
(5) **Flow sheet for nitrate nitrogen**: The flow sheet for nitrate nitrogen in fertilizers is shown below.

![Flow sheet for nitrate nitrogen in fertilizers](image)

Figure The flow sheet for nitrate nitrogen in fertilizers
4.1.3.c  Phenol sulfuric acid method

(1)  **Summary**

The testing method is applicable to the fertilizer containing nitrate. It is also applicable to the fertilizers containing chemical compounds such as urea, nitrolime and organic matters that digest by heating and isolate ammonia. This testing method is classified as Type D and its symbol is 4.1.3.c-2017 or N-N.c-1.

Add a copper sulfate - silver sulfate solution, calcium hydroxide and basic magnesium carbonate to an analytical sample, extract nitrate nitrogen (N-N) as well as removing chloride and organic matters, and measure the absorbance of nitro phenol ammonium sulfate formed by the reaction with phenol sulfuric acid and an ammonia solution to calculate nitrate nitrogen (N-N) in an analytical sample. In addition, the performance of this testing method is shown in **Comment 3**.

(2)  **Reagent**: Reagents are as shown below.

a)  **Nitrate standard solution (N-N 5 mg/mL)** (1): Heat potassium nitrate (no less than 99.9 (%) (mass fraction) in purity) at 110 ºC for no less than 1 hour, and after standing to cool in a desiccator, transfer 36.09 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.

b)  **Nitrate standard solution (N-N 0.01 mg/mL)**: Dilute predetermined volume of nitrate standard solution (N-N 5 mg/mL) with water to prepare a nitrate standard solution (N-N 0.01 mg/mL).

c)  **Copper sulfuric - silver sulfuric solution** (1): Dissolve 5 g of copper (II) sulfate pentahydrate specified JIS K 8983 in 900 mL of water, and dissolve while adding 4 g of silver sulfate specified in JIS K 8965 to make 1000 mL (2).

d)  **Phenol sulfuric acid**: Dissolve 15 g of phenol specified in JIS K 8798 in 100 mL of sulfuric acid specified in JIS K 8965, heat in a water bath at 80 ºC - 100 ºC for two hours and then let it stand to cool (2).

e)  **Ammonia solution**: A JIS Guaranteed (NH₃ 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.

f)  **Calcium hydroxide**: A JIS Guaranteed Reagent specified in JIS K 8575 or a reagent of equivalent quality.

g)  **Basic magnesium carbonate**: Basic magnesium carbonate that contains no nitrate nitrogen.

**Note** (1) This is an example of preparation; prepare an amount as appropriate.

(2)  Store in an amber bottle.

**Comment 1**  Instead of the nitrate standard solution in (2), a nitrate standard solution for a calibration curve can be prepared using nitrate nitrogen (NO₃-N 0.1 mg/mL or 1 mg/mL) traceable to National Metrology.

(3)  **Instruments**: Instruments are as shown below:

a)  **Rotary shaker**: A rotary shaker that can rotate a 250-mL volumetric flask upside down at 30 - 40 revolutions/min.

b)  **Spectrophotometer**: A spectrophotometer specified in JIS K 0115

c)  **Water bath**: Water bath that can be adjusted to no less than 80 ºC

(4)  **Test procedures**

(4.1)  **Extraction**: Conduct extraction as shown below.

a)  Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

b)  Add about 200 mL of copper sulfuric - silver sulfuric solution and shake to mix at 30 - 40
revolutions /min for about 20 minutes.

c) Add about 1 g of calcium hydroxide and about 1 mg of basic magnesium carbonate and shake to mix at 30 - 40 revolutions /min for about 10 minutes.

d) Add water up to the marked line.

e) Filter with Type 3 filter paper to make a sample solution (3).

**Note** (3) As soon as the sample solution is prepared, conduct the procedure in (4.2) a).

**Comment 2** If the filtrate of (4.1) e) is colored, add no more than 0.5 g of active carbon and filter with Type 3 filter paper to make a sample solution.

(4.2) **Coloring**: Conduct coloring as shown below.

a) Transfer a predetermined amount of sample solution (the equivalents of 0.01 mg - 0.1 mg as N-N) into a small evaporating dish (4).

b) Evaporate water until dry on a water bath at no less than 80 °C.

c) After standing to cool, swiftly add 2 mL of phenol sulfuric acid (5) and then rotate the evaporating dish so that the whole residue comes in contact with the acid.

d) After leaving at rest for about 10 minutes, add 20 mL of water (6).

e) After standing to cool, transfer it with water to a 100- mL volumetric flask.

f) Add an ammonia solution (1+2), until the color of a solution changes to light yellow, to allow it to be weak alkalinity, and further add 3 mL of ammonia solution (1+3) (7).

g) After cooling is complete, add water up to the marked line and leave at rest for 30 minutes.

**Note** (4) A round-bottom glass or porcelain evaporating dish is preferable.

(5) Add at the center of a small evaporating dish with Komagome pipet.

(6) If residue does not dissolve easily, grind it with a glass rod.

(7) As no color appears from a blank test solution for a calibration curve preparation, add almost the same volume of ammonia solution (1+2) as the nitrate standard solution.

(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) **Measurement conditions of spectrophotometer**: Set up the measurement conditions of spectrophotometer considering the following.

Detection wavelength: 410 nm

b) **Calibration curve preparation**

1) Transfer 1 mL - 10 mL of nitrate standard solution (N-N 0.01 mg/mL) to small evaporating dishes (4) step-by-step.

2) Conduct the same procedure as (4.2) b) - g) to make the nitrate standard solution for the calibration curve preparation.

3) Transfer 40 mL of water to a 100-mL volumetric flask, and shake to mix while gently adding 2 mL of phenol sulfuric acid. Let it stand to cool and conduct the same procedure as (4.2) f) - g) to make the blank test solution for calibration curve.

4) Measure absorbance at wavelength 410 nm of the nitrate standard solution for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.

5) Prepare the calibration curve of concentration of the nitrate nitrogen and absorbance of the nitrate standard solutions for the calibration curve preparation.

c) **Sample measurement**

1) Regarding the solution in (4.2) g), measure absorbance by the same procedure as b) 4).
2) Obtain the nitrate nitrogen (N-N) content from the calibration curve and calculate nitrate nitrogen (N-N) in the analytical sample.

Comment 3 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 16 % (mass fraction) and 1 % (mass fraction) - 3% (mass fraction) are 103.4 % and 101.1 % - 100.9 % as nitrate nitrogen (N-N) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction) for solid fertilizers, and 0.002 % (mass fraction) for fluid fertilizers.

References

(5) Flow sheet for nitrate nitrogen: The flow sheet for nitrate nitrogen in fertilizers is shown below.

```
<table>
<thead>
<tr>
<th>1 mg analytical sample</th>
<th>Weigh to the order of 1 mg to a 250-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shaking to mix</td>
<td>Rotary shaker (30 - 40 revolutions/min) for 20 minutes</td>
</tr>
<tr>
<td></td>
<td>← About 1 g of calcium hydroxide</td>
</tr>
<tr>
<td></td>
<td>← About 1 g of basic magnesium carbonate</td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>Rotary shaker (30 - 40 revolutions/min) for 10 minutes</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>
```

Figure 1 The flow sheet for nitrate nitrogen in fertilizers (Extraction procedure)
Sample solution

Aliquot (predetermined volume)

Small evaporating dish

Evaporation to dryness

No less than 80 °C Water bath

— 2 mL of phenol sulfuric acid

Leaving at rest

About 10 minutes after making phenol sulfuric acid contact with residue

— 20 mL of water

Standing to cool

Transfer

100-mL volumetric flask

— Ammonia solution (1+2) (until the color of a solution changes to light yellow)

— Ammonia solution (1+3) 3 mL

— Water (up to the marked line)

Leaving at rest

For about 30 minutes

Measurement

Spectrophotometer (410 mm)

Figure Flow sheet for nitrate nitrogen in fertilizers (Coloring and measurement procedure)
4.2 Phosphoric acid
4.2.1 Total phosphoric acid
4.2.1.a Ammonium vanadomolybdate absorptiometric analysis

(1) Summary
This testing method is applicable to fertilizers containing organic matters. This testing method is classified as Type C and its symbol is 4.2.1.a-2017 or T-P.a-1.

Add sulfuric acid, potassium sulfate and copper (II) sulfate pentahydrate to an analytical sample. Pretreat by Kjeldahl method or incineration-hydrochloric acid boiling to convert total phosphorus to phosphate ion, and measure the absorbance of phosphovanadomolybdate salt formed by the reaction with ammonium vanadate (V), hexaammonium heptamolybdate and nitric acid, to obtain total phosphoric acid (T-P_2O_5) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.

a) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.
b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
c) Nitric acid: A JIS Guaranteed (HNO_3 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
d) Ammonia solution: A JIS Guaranteed (NH_3 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.
e) Catalyst (1): Mix potassium sulfate specified in JIS K 8962 and copper (II) sulfate pentahydrate (2) specified in JIS K 8983 in the ratio of 9 to 1.
f) Coloring reagent solution (3)(4): Dissolve 1.12 g of ammonium vanadate (V) (5) specified in JIS K 8747 in water, add 250 mL of nitric acid, then add 27 g of hexaammonium heptamolybdate tetrahydrate (6) specified in JIS K 8905 dissolved in water, and further add water to make 1000 mL (7).
g) Phenolphthalein solution (1 g/100 mL): Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.
h) Phosphoric acid standard solution (P_2O_5 10 mg/mL) (3): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL-3 mL of nitric acid, and add water up to the marked line.
i) Phosphoric acid standard solution (P_2O_5 0.5 mg/mL) (3): Transfer 50 mL of phosphoric acid standard solution (P_2O_5 10 mg/mL) to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.

Note
(1) A tablet is commercially available.
(2) Crush into powder as appropriate.
(3) This is an example of preparation; prepare an amount as appropriate.
(4) This corresponds to reagent “a” reagent solution in the Official Methods of Analysis of Fertilizers (1992).
(5) This corresponds to ammonium metavanadate in the Official Methods of Analysis of Fertilizers (1992).
(6) This corresponds to ammonium molybdate in the Official Methods of Analysis of Fertilizers (1992).
(7) Store in an amber bottle.
Comment 1 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate the total phosphoric acid (T-P₂O₅) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.3) by a conversion factor (2.2914).

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) Spectrophotometer: A spectrophotometer specified in JIS K 0115
b) Electric furnace: An electric furnace that can be adjusted to 550 °C ± 5 °C.
c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.
d) Digestion flask: Kjeldahl flask

(4) Test procedures
(4.1) Sample solution preparation: Prepare a sample solution as shown below:

(4.1.1) Kjeldahl method

a) Weigh 0.5 g - 5 g of an analytical sample to the order of 1 mg, and put it in a 300-mL digestion flask.
b) Add 5 g - 10 g of catalyst, and further add 20 mL - 40 mL of sulfuric acid, shake to mix and heat gently.
c) After bubbles cease to form, heat until white smoke of sulfuric acid evolves.
d) Ignite until organic matters are completely digested (8).
e) After standing to cool, add a small amount of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water, and further mix by shaking.
f) After cooling is complete, add water up to the marked line
g) Filter with Type 3 filter paper to make a sample solution.

Note (8) When the solution has finished changing color, heat further for no less than 2 hours.

Comment 2 The procedure in (4.1.1) is the same as the procedure in (4.1) in 4.2.1.b. Also, the procedures in (4.1.1) a) - f) are the same as in (4.1) in 4.1.1.a.

(4.1.2) Incineration-hydrochloric acid boiling

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.
b) Put the tall beaker in an electric furnace, and heat gently to char (9).
c) Ignite at 550 °C ± 5 °C for no less than 4 hours to incinerate (9).
d) After standing to cool, moisten the residue with a small amount of water, gradually add about 10 mL of hydrochloric acid, and further add water to make 20 mL.
e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to boil for about 5 minutes.
f) After cooling is complete, transfer to a 250-mL - 500-mL volumetric flask with water.
g) Add water up to the marked line.
h) Filter with Type 3 filter paper to make a sample solution.

Note (9) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about
1 hour and further raise to 550 °C in 1 to 2 hours.

**Comment 3** The procedures in (4.1.2) are the same as the procedures in (4.1.1) in 4.3.1.a, (4.1.1) in 4.5.1.a, (4.1.1) in 4.6.1.a and (4.1) in 8.4.a.

(4.1.3) Incineration-aqua regia digestion

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char (10).

c) Ignite at 450 ºC ± 5 ºC for 8 - 16 hours to incinerate (10).

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass (11), and continue heating on the hot plate or sand bath to concentrate until nearly dried up (12).

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (13) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

**Comment 4** The procedures in (4.1.3) are the same as in (4.1.2) in 4.3.1.a, (4.1.2) in 4.5.1.a, (4.1.2) in 4.6.1.a, (4.1) in 4.9.1.a, (4.1) in 4.9.1.b, (4.1) in 4.10.1.a and (4.1) in 4.10.1.b. And also, the same as (4.1) a - h) in 5.3.a, (4.1) a - h) in 5.3.b, (4.1) a - h) in 5.4.a, (4.1) a - h) in 5.4.b, (4.1) a - h) in 5.5.a, (4.1) a - h) in 5.5.d, (4.1) a - h) in 5.6.a and (4.1) a - h) in 5.6.b.

(4.2) **Coloring:** Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 6 mg as P\textsubscript{2}O\textsubscript{5}) to a 100-mL volumetric flask.

b) Add 1 - 2 drop(s) of phenolphthalein solution (1 g/100 mL), and neutralize by adding ammonia solution (1+1) until the color of the solution becomes light red-purple (14).

c) Add nitric acid (1+10) until the light red-purple color of the solution disappears to make it slightly acidic, and add a proper amount of water (15).

d) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes.

**Note** (14) It is not necessary to add a phenolphthalein solution (1 g/100 mL) when determination can be done by the color change of copper ion (light blue → blue-purple).

(15) Without the addition of water, precipitate may be produced when a coloring reagent
solution is added.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.
   Detection wavelength: 420 nm

b) Calibration curve preparation
   1) Transfer 1 mL - 12 mL of phosphoric acid standard solution (P₂O₅ 0.5 mg/mL) to 100-mL volumetric flasks step-by-step.
   2) Add a proper amount of water (15), and conduct the same procedure as (4.2) d) to make the P₂O₅ 0.5 mg/100 mL - 6 mg/100 mL phosphoric acid standard solutions for the calibration curve preparation.
   3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.
   4) Measure absorbance at a wavelength of 420 nm of the phosphoric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control (16).
   5) Prepare the calibration curve of the phosphoric acid concentration and absorbance of the phosphoric acid standard solutions for the calibration curve preparation.

c) Sample measurement
   1) Regarding the solution in (4.2) d), measure absorbance by the same procedure as b) 4) (16).
   2) Obtain the phosphoric acid (P₂O₅) content from the calibration curve, and calculate total phosphoric acid (T-P₂O₅) in the analytical sample.

Note (16) Measure within 6 hours after adding the coloring reagent solution in the procedure in (4.2) d).

Comment 5 After the procedure in (4.2) a), it is also possible to measure soluble phosphoric acid at the same time by adding 4 mL of nitrate acid (1+1) and 2 mL of Petermans citrate solution and by conducting the procedures from (4.2) d) to (4.3) in 4.2.2.a (using b reagent solution in the Official Methods of Analysis of Fertilizers (1992)).

Further after the procedure in (4.2) a), it is also possible to measure citrate soluble phosphoric acid at the same time by adding 4 mL of nitrate acid (1+1) and 17 mL of citrate solution and by conducting the procedures from (4.2) d) to (4.3) in 4.2.3.a (using b reagent solution in the Official Methods of Analysis of Fertilizers (1992)).

Comment 6 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 99.4 % - 100.2 % and 101.0 % - 105.7 % as total phosphoric acid (T-P₂O₅) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.04 % (mass fraction) for solid fertilizers, and 0.01 % (mass fraction) for fluid fertilizers.

Table 1 Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratories used for analysis</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
<th>Reproducibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-C-12</td>
<td>9</td>
<td>8.62</td>
<td>0.03</td>
<td>0.04</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis
2) Average (the number of laboratory \( p \) \( \times \) test days \( 2 \) \( \times \) the number of replicate testing \( 3 \) )
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

Flow sheet for total phosphoric acid: The flow sheet for total phosphoric acid in fertilizers is shown below:

- Weigh to the order of 1 mg into a 300-mL Kjeldahl flask.
- About 10 g catalyst
- 20 mL - 40 mL sulfuric acid

- Gently
- After bubbles cease to form, ignite until organic matters are completely digested.
- Room temperature
- A small amount of water

- 250-mL - 500-mL volumetric flask, water

- Room temperature
- Water (up to the marked line)

- Type 3 filter paper

Figure 1-1 Flow sheet for total phosphate in fertilizers
(Kjeldahl method procedure (4.1.1))
Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker. Heat gently
Ignite at 550 °C ± 5 °C for no less than 4 hours
A small amount of water, moisten the residue
About 10 mL of hydrochloric acid
Water (up to about 20 mL)
Cover with a watch glass, and boil for 5 minutes.

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 g analytical sample</td>
<td></td>
</tr>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>Ignite at 550 °C ± 5 °C for no less than 4 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td></td>
<td>A small amount of water, moisten the residue</td>
</tr>
<tr>
<td></td>
<td>About 10 mL of hydrochloric acid</td>
</tr>
<tr>
<td></td>
<td>Water (up to about 20 mL)</td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and boil for 5 minutes.</td>
</tr>
<tr>
<td>Cooling</td>
<td></td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask, water</td>
</tr>
<tr>
<td></td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1-2  Flow sheet for total phosphate in fertilizers
(Incineration-hydrochloric acid boiling procedure (4.1.2))
Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker.

Heat gently
Ignite at 450 °C ± 5 °C for 8 - 16 hours

Room temperature
- A small amount of water
- About 10 mL of nitric acid
- About 30 mL of hydrochloric acid

Cover with a watch glass to digest

Slightly move the watch glass and remove acid

Room temperature
- 25 mL - 50 mL of hydrochloric acid (1:5)

Cover with a watch glass to dissolve

Standing to cool

Filtration
Type 3 filter paper

Sample solution

Figure 1-3  Flow sheet for total phosphate in fertilizers
(Incineration-aqua regia digestion orocedure (4.1.3))

Sample solution

Aliquot (predetermined amount)

100-mL volumetric flask

- 1-2 drop(s) of phenolphthalein solution (1 g/100 mL)
- Ammonia solution (1+1) [neutralization]
- Nitric acid (1+10) [slightly acidic]
- A proper amount of water
- 20 mL of coloring reagent solution
- Water (up to the marked line)

Leaving at rest

About 30 minutes

Measurement

Spectrophotometer (420 nm)

Figure 2 Flow sheet for total phosphate in fertilizers (Coloring and measurement procedure)
4.2.1.b  Quinoline gravimetric analysis

(1)  Summary
This testing method is applicable to fertilizers containing organic matters. It is suitable for the fertilizers with relatively a high content of phosphoric acid. This testing method is classified as Type E and its symbol is 4.2.1.b-2017 or T-P.b-1.

Add sulfuric acid, potassium sulfate and copper (II) sulfate pentahydrate to an analytical sample. Pretreat by the Kjeldahl method to convert the total phosphoric acid (T-P$_2$O$_5$) to phosphate ion, and measure the mass of quinolinium phosphomolybdate formed by the reaction with quinoline, molybdic acid and nitric acid to obtain the total phosphoric acid (T-P$_2$O$_5$) in an analytical sample.

(2)  Reagent: Reagents are as shown below.
   a)  **Sulfuric acid**: A JIS Guanteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.
   b)  **Nitric acid**: A JIS Guaranteed (HNO$_3$ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   c)  **Sodium molybdate solution**: Dissolve 70 g of sodium molybdate dihydrate in 150 mL of water.
   d)  **Quinoline solution**: Add 5 mL of quinolone specified in JIS K 8279 to the mixture solution of 35 mL of nitric acid and 100 mL of water.
   e)  **Quimosiac solution**: Add 60 g of citric acid monohydrate specified in JIS K 8283 to the mixture solution of 85 mL nitric acid and 150 mL of water to dissolve. Add gradually total volume of the sodium molybdate solution to mix. Add gradually the total volume of the quinoline solution while mixing the solution. After leaving at rest overnight, filter the total volume with Type 3 filter paper. Add 280 mL of acetone specified in JIS K 8034, and further add water to make 1000 mL.
   f)  **Catalyst**: Mix potassium sulfate specified in JIS K 8962 and copper (II) sulfate pentahydrate specified in JIS K 8983 in the ratio of 9 to 1.

Note (1)  This is an example of preparation; prepare an amount as appropriate.
(2)  A tablet is commercially available.
(3)  Crush into powder as appropriate.

(3)  **Apparatus and instruments**: Apparatus and instruments are shown below.
   a)  **Water bath**: Water bath that can be adjusted to 60 °C - 65 °C.
   b)  **Drying apparatus**: A drying apparatus that can be adjusted to 220 °C ± 5 °C.
   c)  **Crucible type glass filter**: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 220 °C ± 5 °C in advance and measure the mass to the order of 1 mg.
   d)  **Digestion flask**: Kjeldahl flask

(4)  Test procedures
(4.1)  **Kjeldahl method**: Conduct digestion as shown below.
   a)  Weigh 0.5 g - 5 g of an analytical sample to the order of 1 mg, and put it in a 300-mL digestion flask.
   b)  Add 5 g - 10 g of catalyst, and further add 20 mL - 40 mL of sulfuric acid, shake to mix and heat gently.
   c)  After bubbles cease to form, heat until white smoke of sulfuric acid evolves.
   d)  Ignite until organic matters are completely digested.
   e)  After standing to cool, add a small amount of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water.
f) After cooling is complete, add water up to the marked line

g) Filter with Type 3 filter paper to make a sample solution.

Note (4) When the solution has finished changing color, heat further for no less than 2 hours.

Comment 1 The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.1.a. In addition, the sample solution prepared in (4.1.2) in 4.2.1.a and (4.1.3) in 4.2.1.a can also be used.

(4.2) Measurement: Conduct measurement as shown below.

a) Transfer a predetermined volume (the equivalents of 10 mg - 30 mg as P₂O₅ and no more than 5 mL as sulfuric acid) of sample solution to a 300-mL tall beaker.

b) Add 5 mL of nitric acid and add water to make about 80 mL.

c) Cover with a watch glass. After boiling for about 3 minutes, wash the watch glass and the inside the tall beaker with water and add water to make about 100 mL.

d) Immediately, add 50mL of quimosiac solution, heat for about 15 minutes while sometimes mixing in a water bath at 60 °C - 65 °C to produce the precipitate of quinonium phosphomolybdate.

e) After standing to cool down to room temperature while sometimes mixing, filter under reduced pressure with a crucible type glass filter, wash the tall beaker 3 times with water and transfer the whole precipitate into a crucible type glass filter, and further wash 7 - 8 times with water.

f) Transfer the precipitate together with the crucible type glass filter into a drying apparatus and heat at 220 °C ± 5 °C for about 30 minutes.

g) As soon as heating is complete, move it into a desiccator and let it stand to cool.

h) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

i) Calculate total phosphoric acid (T-P₂O₅) in the analytical sample by the following formula.

\[
\text{Total phosphoric acid (T-P}_2\text{O}_5 \text{) (% (mass fraction)) in an analytical sample} = A \times 0.03207 \times (V_1/V_2) \times (1/W) \times 100
\]

\[A:\] Mass (g) of the precipitate in h)

\[W:\] Mass (g) of the analytical sample

\[V_1:\] Constant volume (mL) of sample solution

\[V_2:\] Volume (mL) of the sample solution transferred in a)

References

(5) **Flow sheet for total phosphoric acid**: The flow sheet for total phosphoric acid in fertilizers is shown below:

| 0.5 g - 5 g analytical sample | Weigh to the order of 1 mg into a 300-mL digestion flask. |
| 5 g - 10 g catalyst | ← |
| 20 mL - 40 mL sulfuric acid | ← |
| Heating | Gently |
| Heating | After bubbles cease to form, ignite until organic matters are completely digested |
| Standing to cool | Room temperature |
| Small mount of water | ← |
| Transfer | 250-mL - 500-mL volumetric flask, water |
| Cooling | ← Water (up to the marked line) |
| Filtration | Type 3 filter paper |
| Sample solution | |

Figure 1  Flow sheet for total phosphate in fertilizers  
(Kjeldahl method procedure)

Figure 2  The flow sheet for total phosphate in fertilizers (Measurement procedure)
4.2.2 Soluble phosphoric acid
4.2.2.a Ammonium vanadomolybdate absorptiometric analysis

(1) Summary
This testing method is applicable to fertilizers that do not contain matter not colored by hydrolysis with nitrate acids such as phosphonic acid. This testing method is classified as Type C and its symbol is 4.2.2.a-2017 or S-P.a-1.

Extract by adding water to an analytical sample, then extract by adding an ammonium citric acid solution, and combine respective pre-determined amounts of extract (equivalent volume). Heat after adding nitric acid (1+1), hydrolyze nonorthophosphoric acid to orthophosphate ion and measure the absorbance of phosphovanadomolybdate salt formed by the reaction with ammonium vanadate (V), hexaammonium heptamolybdate and nitric acid to obtain ammonia alkaline ammonium citrate-soluble phosphoric acid (soluble phosphoric acid (S-P₂O₅)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.
  a) Nitric acid: A JIS Guaranteed (HNO₃ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
  b) Ammonia solution: A JIS Guaranteed (NH₃ 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.
  c) Petermans citrate solution: Add 173 g of citric acid monohydrate specified in JIS K 8283 in water to dissolve and add gradually an ammonia solution equivalent to 42 g of nitrogen while cooling. After cooling is complete, add water to make 1000 mL. Additionally, check that the specific gravity of the solution is 1.082 - 1.083 (15 ºC) and the nitrogen content per 1 mL is 42 mg.
  d) Coloring reagent solution (1)(2): Dissolve 1.12 g of ammonium vanadate (V)(3) specified in JIS K 8747 in water, add 150 mL of nitric acid, then add 50 g of hexaammonium heptamolybdate tetrahydrate (4) specified in JIS K 8905 dissolved in water, and further add water to make 1000 mL (5).
  e) Phosphoric acid standard solution (P₂O₅ 10 mg/mL) (1): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL-3 mL of nitric acid, and add water up to the marked line.
  f) Phosphoric acid standard solution (P₂O₅ 0.5 mg/mL) (1): Transfer 50 mL of phosphoric acid standard solution (P₂O₅ 10 mg/mL) to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
  (2) This corresponds to reagent “b” reagent solution in the Official Methods of Analysis of Fertilizers (1992).
  (3) This corresponds to ammonium metavanadate in the Official Methods of Analysis of Fertilizers (1992).
  (4) This corresponds to ammonium molybdate in the Official Methods of Analysis of Fertilizers (1992).
  (5) Store in an amber bottle. However, the reagent solution cannot tolerate long term preservation.

Comment 1 The coloring reagent solution in d) can also be prepared by the following method. Dissolve 2.24 g of ammonium vanadate (V) (3) specified in JIS K 8747 in water, add 300 mL of nitric acid, and add water to make 1000 mL. Separately, add 100 g of
hexaammonium heptamolybdate tetrahydrate (4) specified in JIS K 8905 while dissolving in water, and further add water to make 1000 mL. In the case of usage, mix equal volumes of the two solutions.

**Comment 2** Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate soluble phosphoric acid (S-P$_2$O$_5$) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.3) by a conversion factor (2.2914).

(3) **Instruments**: Instruments are as shown below:
   a) **Water bath**: Water bath that can be adjusted to 65 °C ± 2 °C.
   b) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.
   c) **Spectrophotometer**: A spectrophotometer specified in JIS K 0115

(4) **Test procedures**

(4.1) **Extraction**: Conduct extraction as shown below.
   a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and put it in a small mortar.
   b) Add about 20 mL - 25 mL of water, grind well and filter the supernatant with Type 6 filter paper into a 250-mL volumetric flask.
   c) Further, after repeating procedure in b) 3 times, transfer non-dissolved matter in the small mortar onto a filter paper and wash with water until the filtrate becomes about 200 mL.
   d) Add a small amount of nitric acid to the filtrate, and further add water up to the marked line to make a sample solution (1).
   e) Transfer the non-dissolved matter on the filter paper together with the filter paper to another 250-mL volumetric flask, and add 100 mL of Petermans citrate solution and stopple. Then shake to mix until the filter paper breaks down.
   f) Heat the volumetric flask in e) in water bath at 65 °C ± 2 °C for 1 hour while shaking to mix every 15 minutes.
   g) After immediate cooling is complete, add water up to the marked line
   h) Filter with Type 6 filter paper to make a sample solution (2).

**Note** (6) It is recommended to use a 250-mL short-neck volumetric flask.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1) of 4.2.2.b.

**Comment 4** When the determination is affected by the coloring of the sample solution of d) and h), transfer the predetermined volume (equivalent volume) (7) of the sample solution (1) and the sample solution (2) to a 100-mL volumetric flask, add a few drops of hydrochloric acid (1+1) to acidify, then add no more than 0.1 g of active carbon. After leaving at rest for a little while, add water up to the marked line and filter. The filtrate is to be the mixture solution for the sample solution of (4.2) a). Additionally, as phosphorus contained in active carbon has the possibility to elute and affects the determination value, a blank test is required.

(4.2) **Coloring**: Conduct coloring as shown below.
   a) Transfer a predetermined amount (the equivalents of 0.5 mg - 6 mg as P$_2$O$_5$ and no more than the equivalents of 2 mL of Petermans citrate solution) (7) of the sample solution (1) and the sample solution (2) to a 100-mL volumetric flask.
   b) Add the solution to make Petermans citrate solution equivalent to 2 mL.
c) Add 4 mL of nitric acid (1+1)\(^{(8)}\), and heat to boil \(^{(9)}\).

d) After cooling is complete, add a proper amount of water \(^{(10)}\).

e) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes \(^{(8)}\).

**Comment 5** The volumetric flask used in the procedure in a) should be distinguished as a flask to be used for phosphate coloring operation and should not be used for other purposes.

**Note**

7) The transferred volume of the sample solution (1) and the sample solution (2) should be equivalent.

8) When the solution is muddled by adding nitric acid (1+1), conduct centrifugation after the procedure in e).

9) When it does not contain non-orthophosphate, the boiling operation is not necessary.

10) Without the addition of water, precipitate may be produced when a coloring reagent solution is added.

(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) **Measurement conditions of spectrophotometer**: Set up the measurement conditions of spectrophotometer considering the following.

Detection wavelength: 420 nm

b) **Calibration curve preparation**

1) Transfer 1 mL - 12 mL of phosphoric acid standard solution (P\(_2\)O\(_5\) 0.5 mg/mL) to 100-mL volumetric flasks step-by-step.

2) Add 2 mL of Petermans citrate solution, 4 mL of nitric acid (1+1) and a proper amount of water \(^{(10)}\), and conduct the same procedure as (4.2) e) to make the P\(_2\)O\(_5\) 0.5 mg/100 mL - 6 mg/100 mL phosphoric acid standard solutions for the calibration curve preparation.

3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.

4) Measure absorbance at a wavelength of 420 nm of the phosphoric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control \(^{(11)}\).

5) Prepare the calibration curve of the phosphoric acid concentration and absorbance of the phosphoric acid standard solutions for the calibration curve preparation.

c) **Sample measurement**

1) Regarding the solution in (4.2) e), measure absorbance by the same procedure as b) 4) \(^{(11)}\).

2) Obtain the phosphoric acid (P\(_2\)O\(_5\)) content from the calibration curve, and calculate soluble phosphoric acid (S-P\(_2\)O\(_5\)) in the analytical sample.

**Note**

11) Measure within 2 hours after adding the coloring reagent solution.

**Comment 6** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 99.4 % - 100.6 % and 98.6 % - 103.0 % as soluble phosphoric acid (S-P\(_2\)O\(_5\)) respectively. The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.
Additionally, the minimum limit of quantification of this testing method is about 0.08% (mass fraction).

<table>
<thead>
<tr>
<th>Name of certified reference material fertilizer</th>
<th>Number of laboratory</th>
<th>Average 2) (%) 3)</th>
<th>Repeatability 4)</th>
<th>Intermediate precision 5)</th>
<th>Reproducibility 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-B-10</td>
<td>10</td>
<td>8.62</td>
<td>0.04</td>
<td>0.05</td>
<td>0.06</td>
</tr>
<tr>
<td>FAMIC-B-14</td>
<td>10</td>
<td>9.18</td>
<td>0.03</td>
<td>0.04</td>
<td>0.09</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Ammonium vanadomolybdate absorptiometric analysis
2) Average (the number of laboratory ($p$) × test days (2) × the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References
(5) Flow sheet for soluble phosphoric acid: The flow sheet for soluble phosphoric acid in fertilizers is shown below:

2.5 g analytical sample

Repeat 3 times
Grinding

Supernatant filtration
Type 6 filter paper, 250-mL volumetric flask

<Residue>

Transfer to a filter paper
Washing with water, until filtrate reaches about 200 mL

<Residue>
<Filtrate>
A small amount of nitric acid
Water (up to the marked line)

Sample solution (1)

Transfer
Together with filter paper, 250-mL volumetric flask

Shaking to mix
Stopple and shake to mix until a filter paler breaks down

Heating
At 65 ºC ± 2 ºC for 1 hour while shaking to mix every 15 minutes.

Cooling
Immediately

Water (up to the marked line)
Filtration
Type 6 filter paper

Sample solution (2)

Figure 1  The flow sheet for soluble phosphoric acid in fertilizers (Extraction procedure)
Figure 2  The flow sheet for soluble phosphoric acid in fertilizers
(Coloring and measurement procedure)
4.2.2.b Quinoline gravimetric analysis

(1) Summary
This testing method is applicable to fertilizers containing no Phosphorous acid, etc. It is suitable for the fertilizers with relatively a high content of phosphoric acid. This testing method is classified as Type E and its symbol is 4.2.2.b-2017 or S-Pb-1.

Extract by adding water to an analytical sample, then extract by adding an ammonium citric acid solution, and combine respective pre-determined amounts of extract (equivalent volume). Heat after adding nitric acid and water, hydrolyze nonorthophosphoric acid to orthophosphate ion and measure the mass of quinolinium phosphomolybdate formed by the reaction with quinoline, molybdcic acid and nitric acid to obtain ammonia alkaline ammonium citrate soluble phosphoric acid (soluble phosphoric acid (S-P2O5)) in an analytical sample.

(2) Reagents: Reagents are as shown below.

a) Nitric acid: A JIS Guaranteed (HNO3 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.

b) Ammonia solution: A JIS Guaranteed (NH3 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.

c) Petermans citrate solution: Add 173 g of citric acid monohydrate specified in JIS K 8283 in water to dissolve and add gradually an ammonia solution equivalent to 42 g of nitrogen while cooling. After cooling is complete, add water to make 1000 mL. Additionally, check that the specific gravity of the solution is 1.082 - 1.083 (15 ºC) and the nitrogen content per 1 mL is 42 mg.

d) Sodium molybdate solution: Dissolve 70 g of sodium molybdate dihydrate in 150 mL of water.

e) Quinoline solution: Add 5 mL of quinolone specified in JIS K 8279 to the mixture solution of 35 mL of nitric acid and 100 mL of water.

f) Quimosiac solution: Add 60 g of citric acid monohydrate specified in JIS K 8283 to the mixture solution of 85 mL nitric acid and 150 mL of water to dissolve. Add gradually total volume of the sodium molybdate solution to mix. Add gradually the total volume of the quinoline solution while mixing the solution. After leaving at rest overnight, filter the total volume with Type 3 filter paper. Add 280 mL of acetone specified in JIS K 8034, and further add water to make 1000 mL.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) Water bath: Water bath that can be adjusted to 65 ºC ± 2 ºC and 60 ºC - 65 ºC.

b) Drying apparatus: A drying apparatus that can be adjusted to 220 ºC ± 5 ºC.

c) Crucible type glass filter: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 220 ºC ± 5 ºC in advance and measure the mass to the order of 1 mg.

(4) Test procedures

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and put it in a small mortar.

b) Add about 20 mL - 25 mL of water, grind well and filter the supernatant with Type 6 filter paper into a 250-mL volumetric flask.

c) Further, after repeating procedure in b) 3 times, transfer non-dissolved matter in the small mortar onto a filter paper and wash with water until the filtrate becomes about 200 mL.

d) Add a small amount of nitric acid to the filtrate, and further add water up to the marked line to make a sample solution (1).

e) Transfer the non-dissolved matter on the filter paper together with the filter paper to another
250- mL (1) volumetric flask, and add 100 mL of Petermans citrate solution and stopple. Then shake to mix until the filter paper breaks down completely.

f) Heat the volumetric flask in e) in water bath at 65 °C ± 2 °C for 1 hour while shaking to mix every 15 minutes.

g) After immediate cooling is complete, add water up to the marked line

h) Filter with Type 6 filter paper to make a sample solution (2).

Note (1) It is recommended to use a 250-mL short-neck volumetric flask.

Comment 1 The procedure in (4.1) is the same as the procedure in (4.1) of 4.2.2.a.

(4.2) Measurement: Conduct measurement as shown below.

a) Transfer a predetermined amount (the equivalents of 10 mg - 30 mg as P₂O₅ and no more than the equivalents of 2 mL of Petermans citrate solution) (2) of the sample solution (1) and the sample solution (2) to a 300-mL tall beaker.

b) Add 5 mL of nitric acid and add water to make about 80 mL.

c) Cover with a watch glass. After boiling for about 3 minutes, wash the watch glass and the inside the tall beaker with water and add water to make about 100 mL.

d) Immediately, add 50mL of quimosiac solution, heat for about 15 minutes while sometimes mixing in a water bath at 60 °C - 65 °C to produce the precipitate of quinonium phosphomolybdate.

e) After standing to cool down to room temperature while sometimes mixing, filter under reduced pressure with a crucible type glass filter, wash the tall beaker 3 times with water and transfer the whole precipitate into a crucible type glass filter, and further wash 7 - 8 times with water.

f) Transfer the precipitate together with the crucible type glass filter into a drying apparatus and heat at 220 °C ± 5 °C for about 30 minutes.

g) As soon as heating is complete, move it into a desiccator and let it stand to cool.

h) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

i) Calculate soluble phosphoric acid (S-P₂O₅) in the analytical sample by the following formula.

\[
\text{Soluble phosphoric acid (% (mass fraction)) in an analytical sample} = A \times 0.03207 \times \left( \frac{V_1}{V_2} \right) \times \left( \frac{1}{W} \right) \times 100
\]

\[
A: \quad \text{Mass (g) of the precipitate in h)}
\]

\[
W: \quad \text{Mass of an analytical sample (2.5 g)}
\]

\[
V_1: \quad \text{Predetermined volume (250 mL) of the sample solution}
\]

\[
V_2: \quad \text{Volume (mL) of the sample solution transferred in a)
\]

Note (2) The transferred volume of the sample solution (1) and the sample solution (2) should be equivalent.

References

(5) **Flow sheet for soluble phosphoric acid**: The flow sheet for soluble phosphoric acid in fertilizers is shown below:

2.5 g analytical sample  
Weigh to the order of 1 mg to a small mortar

Repeat 3 times

About 20 mL - 25 mL water

Grinding

Supernatant filtration

Type 6 filter paper, 250-mL volumetric flask

Transfer residue to a filter paper

Wash with water, until filtrate reaches about 200 mL

Residue

Filtrate

A small amount of nitric acid

Water (up to the marked line)

Sample solution (1)

Transfer

Together with filter paper, 250-mL volumetric flask

100 ml Petermans citrate solution

Shaking to mix

Stopple and shake to mix until a filter paler breaks down

Heating

At 65 ºC ± 2 ºC for 1 hour while shaking to mix every 15 minutes.

Cooling

Immediately

Water (up to the marked line)

Filtration

Type 6 filter paper

Sample solution (2)

Figure 1  Flow sheet for soluble phosphoric acid in fertilizers
(Extraction procedure)
Sample solutions (1) and (2) Aliquot (predetermined volume) Transfer the same volume of sample solutions (1) and (2) to 300-mL volumetric flasks

- 5 mL nitrate acid
- Water (to reach about 80 mL)

Boiling Cover with a watch glass for 3 minutes
Wash a watch glass and inside a tall beaker with water

- Water (to make about 100 mL)
- 50 mL of quimosiac solution

Forming precipitate 60 °C - 65 °C, for 15 minutes, sometimes mixing

Standing to cool Room temperature

Filtration under reduced pressure Crucible type glass filtering apparatus 1G4, 3 times with water

Washing Wash 7 - 8 times with water

Drying 220 °C ± 5 °C for about 30 minutes

Standing to cool Desiccator

Measurement Measure the mass to the order of 1 mg

Figure 2  Flow sheet for soluble phosphoric acid in fertilizers
(Measurement procedure)
4.2.3 Citrate-soluble phosphoric acid
4.2.3.a Ammonium vanadomolybdate absorptiometric analysis

(1) Summary
This testing method is applicable to fertilizers that do not contain matter not colored by hydrolysis with nitrate acids such as phosphonic acid. This testing method is classified as Type C and its symbol is 4.2.3.a-2018 or C-P.a-2.

Extract by adding a citric acid solution to an analytical sample. Heat after adding nitric acid (1+1), hydrolyze nonorthophosphoric acid to orthophosphate ion and measure the absorbance of phosphovanadomolybdate salt formed by the reaction with ammonium vanadate (V), hexaammonium heptamolybdate and nitric acid to obtain citrate soluble phosphoric acid (C-P$_2$O$_5$) in an analytical sample. In addition, the performance of this testing method is shown in Comment 9.

(2) Reagent: Reagents are as shown below.

a) Nitric acid: A JIS Guaranteed (HNO$_3$ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.

b) Citric acid solution: Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.

c) Coloring reagent solution: Dissolve 1.12 g of ammonium vanadate (V) specified in JIS K 8747 in water, add 150 mL of nitric acid, then add 50 g of hexaammonium heptamolybdate tetrahydrate specified in JIS K 8905 dissolved in water, and further add water to make 1000 mL.

d) Phosphoric acid standard solution (P$_2$O$_5$ 10 mg/mL): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL-3 mL of nitric acid, and add water up to the marked line.

e) Phosphoric acid standard solution (P$_2$O$_5$ 0.5 mg/mL): Transfer 50 mL of phosphoric acid standard solution (P$_2$O$_5$ 10 mg/mL) to a 1000-mL volumetric flask, add 2 mL-3 mL of nitric acid, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(2) This corresponds to reagent “b” reagent solution in the Official Methods of Analysis of Fertilizers (1992).

(3) This corresponds to ammonium metavanadate in the Official Methods of Analysis of Fertilizers (1992).

(4) This corresponds to ammonium molybdate in the Official Methods of Analysis of Fertilizers (1992).

(5) Store in an amber bottle. However, the reagent solution cannot tolerate long term preservation.

Comment 1 The coloring reagent solution in e) can also be prepared by the following method. Dissolve 2.24 g of ammonium vanadate (V) specified in JIS K 8747 in water, add 300 mL of nitric acid, and add water to make 1000 mL. Separately, add 100 g of hexaammonium heptamolybdate tetrahydrate specified in JIS K 8905 while dissolving in water, and further add water to make 1000 mL. In the case of usage, mix equal volumes of the two solutions.

Comment 2 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In
this case, calculate citrate-soluble phosphoric acid (C-P\(_2\)O\(_5\)) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.3) by a conversion factor (2.2914).

(3) **Instruments:** Instruments are as shown below:
   a) **Extractor:** Constant-temperature rotary shaker or reciprocating water bath shaker as described below.

   aa) **Constant-temperature rotary shaker:** A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

   ab) **Reciprocating water bath shaker:** A reciprocating water bath shaker that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/ minute with amplitude of 25 mm - 40 mm.

   b) **Hot plate:** A hot plate whose surface temperature can be adjusted up to 250 °C.

   c) **Spectrophotometer:** A spectrophotometer specified in JIS K 0115

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker:**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

   b) Add 150 mL of citric acid solution heated up to about 30 °C \((6)\), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.

   c) After immediate cooling is complete, add water up to the marked line

   d) Filter with Type 3 filter paper to make a sample solution.

**Note** (6) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 3** The procedure in (4.1.1) is the same as the procedure in (4.1.1) of 4.2.3.b, (4.1) of 4.2.3.c, (4.1) of 4.2.3.d, (4.1.1) of 4.3.2.a, (4.1) of 4.3.2.b, (4.1) of 4.3.2.c, (4.1.1) of 4.3.2.d, (4.1.1) of 4.6.3.a, (4.1.1) of 4.6.3.b, (4.1.1) of 4.7.2.a, (4.1.1) of 4.7.2.b, (4.1.1) of 4.8.1.a and (4.1.1) of 4.8.1.b.

(4.1.2) **Reciprocating water bath shaker:**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask \((6)\).

   b) Add 150 mL of citric acid solution heated up to about 30 °C \((6)\), and shake to mix by reciprocating horizontally at 160 times /min with amplitude of 25 mm - 40 mm (30 °C ± 1 °C) for 1 hour.

   c) After immediate cooling is complete, add water up to the marked line

   d) Filter with Type 3 filter paper to make a sample solution.

**Note** (7) Use a 250-mL flat-bottom volumetric flask to stabilize the shaking.

**Comment 4** The procedure in (4.1.2) is the same as the procedure in (4.1.2) of 4.2.3.b, (4.1.2) of 4.2.3.d, (4.1.2) of 4.3.2.a, (4.1.2) of 4.3.2.d, (4.1.2) of 4.6.3.a, (4.1.2) of 4.6.3.b, (4.1.2) of 4.7.2.a, (4.1.2) of 4.7.2.b, (4.1.2) of 4.8.1.a and (4.1.2) of 4.8.1.b.
Comment 5 For a by-product phosphate fertilizer, if the pH of the sample solution in (4.1.1) d) and (4.1.2) d) is neutral or basic, prepare a sample solution anew by replacing “1 g of an analytical sample” in the procedures in (4.1.1) a) and (4.1.2) a) with “0.5 g of an analytical sample”.

Comment 6 The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

Comment 7 When the determination is affected by the coloring of the sample solution of (4.1.1) d) and (4.1.2) d), transfer the predetermined volume of the sample solution to a 100-mL volumetric flask, add a few drops of hydrochloric acid (1+1) to acidify, then add no more than 0.1 g of active carbon. After leaving at rest for a little while, add water up to the marked line and filter. The filtrate is prepared as the sample solution of (4.2) a). Additionally, as phosphorus contained in active carbon has the possibility to elute and affects the determination value, a blank test is required.

(4.2) Coloring: Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 6 mg as P₂O₅ and no more than the equivalents of 17 mL of citric acid solution) to a 100-mL volumetric flask.
b) Add the solution to make citric acid solution equivalent to 17 mL.
c) Add 4 mL of nitric acid (1+1) (8), and heat to boil (9).
d) After cooling is complete, add a proper amount of water (10).
e) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes.

Comment 8 The volumetric flask used in the procedure in a) should be distinguished as a flask to be used for phosphate coloring operation and should not be used for other purposes.

Note (8) When the solution is muddled by adding nitric acid (1+1), conduct centrifugation after the procedure in e).
(9) When it does not contain non-orthophosphate, the boiling operation is not necessary.
(10) Without the addition of water, precipitate may be produced when a coloring reagent solution is added.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.
Detection wavelength: 420 nm
b) Calibration curve preparation
1) Transfer 1 mL - 12 mL of phosphoric acid standard solution (P₂O₅ 0.5 mg/mL) to 100-mL volumetric flasks step-by-step.
2) Add 17 mL of a citric acid solution, then add 4 mL of nitric acid (1+1), further add a proper amount of water (10) and conduct the same procedure as (4.2) e) to make the P₂O₅ 0.5 mg/100 mL - 6 mg/100 mL phosphoric acid standard solution for the calibration curve preparation.
3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.
4) Measure absorbance at a wavelength of 420 nm of the phosphoric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve
preparation as the control (11).

5) Prepare the calibration curve of the phosphoric acid concentration and absorbance of the phosphoric acid standard solutions for the calibration curve preparation.

c) Sample measurement

1) Regarding the solution in (4.2) e), measure absorbance by the same procedure as b) 4) (11).
2) Obtain the phosphoric acid (P₂O₅) content from the calibration curve, and calculate citrate soluble phosphoric acid (C-P₂O₅) in the analytical sample.

Note (11) Measure within 2 hours after adding the coloring reagent solution.

Comment 9 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 96.6 % - 103.4 % and 102.0 % - 103.8 % as citrate soluble phosphoric acid (C-P₂O₅) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.03 % (mass fraction) for solid fertilizers, and 0.01 % (mass fraction) for fluid fertilizers.

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of certified reference laboratory</th>
<th>Reproducibility</th>
<th>Intermediate precision</th>
<th>Repeatability</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-A-10</td>
<td>11</td>
<td>0.13</td>
<td>0.8</td>
<td>0.06</td>
</tr>
<tr>
<td>FAMIC-A-13</td>
<td>10</td>
<td>0.09</td>
<td>0.8</td>
<td>0.08</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Ammonium vanadomolybdate absorptiometric analysis
2) Average (the number of laboratory (p) × test days (2) × the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

(5) **Flow sheet for citrate soluble phosphoric acid:** The flow sheet for citrate soluble phosphoric acid in fertilizers is shown below:

![Flow sheet for citrate soluble phosphoric acid](image1)

Figure 1-1 Flow sheet for citrate soluble phosphoric acid in fertilizers (Extraction procedure (4.1.1))

![Flow sheet for citrate soluble phosphoric acid](image2)

Figure 1-2 Flow sheet for citrate soluble phosphoric acid in fertilizers (Extraction procedure (4.1.2))
<table>
<thead>
<tr>
<th>Sample solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>100-mL volumetric flask</td>
</tr>
<tr>
<td>← Citric acid solution, until it reaches the equivalents of 17 mL</td>
</tr>
<tr>
<td>← 4 mL of nitric acid (1+1)</td>
</tr>
<tr>
<td>Heating</td>
</tr>
<tr>
<td>Boiling</td>
</tr>
<tr>
<td>Cooling</td>
</tr>
<tr>
<td>← A proper amount of water</td>
</tr>
<tr>
<td>← 20 mL of coloring reagent solution</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Leaving at rest</td>
</tr>
<tr>
<td>For about 30 minutes</td>
</tr>
<tr>
<td>Measurement</td>
</tr>
<tr>
<td>Spectrophotometer (420nm)</td>
</tr>
</tbody>
</table>

**Figure 2** Flow sheet for citrate soluble phosphoric acid in fertilizers
(Coloring and measurement procedure)
4.2.3.b Ammonium vanadomolybdate absorptiometric analysis (Fertilizers containing phosphorous acid or phosphite)

(1) Summary
This testing method is applicable to the fertilizers containing phosphorous acid or phosphite. This testing method is classified as Type B and its symbol is 4.2.3.b-2018 or C-Pb-2.

Extract by adding a citric acid solution to an analytical sample, add hydrochloric acid - sulfuric acid to heat, and oxygenate phosphorous acid ion to orthophosphate ion, and then measure the absorbance of phosphovanadomolybdate salt formed by the reaction with ammonium vanadate (V), hexaammonium heptamolybdate and nitric acid to obtain citrate-soluble phosphoric acid (C-P2O5) in an analytical sample. In addition, the performance of this testing method is shown in Comment 7.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Nitric acid: A JIS Guaranteed (HNO₃ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.

c) Citric acid solution (1): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.

d) Coloring reagent solution (1)(2): Dissolve 1.12 g of ammonium vanadate (V) (3) specified in JIS K 8747 in water, add 150 mL of nitric acid, then add 50 g of hexaammonium heptamolybdate tetrahydrate (4) specified in JIS K 8905 dissolved in water, and further add water to make 1000 mL (5).

e) Phosphoric acid standard solution (P₂O₅ 10 mg/mL) (1): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.

f) Phosphoric acid standard solution (P₂O₅ 0.5 mg/mL) (1): Transfer 50 mL of phosphoric acid standard solution (P₂O₅ 10 mg/mL) to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(2) This corresponds to reagent “b” reagent solution in the Official Methods of Analysis of Fertilizers (1992).

(3) This corresponds to ammonium metavanadate in the Official Methods of Analysis of Fertilizers (1992).

(4) This corresponds to ammonium molybdate in the Official Methods of Analysis of Fertilizers (1992).

(5) Store in an amber bottle. However, the reagent solution cannot tolerate long term preservation.

Comment 1 The coloring reagent solution in d) can also be prepared by the following method. Dissolve 2.24 g of ammonium vanadate (V) (3) specified in JIS K 8747 in water, add 300 mL of nitric acid, and add water to make 1000 mL. Separately, add 100 g of hexaammonium heptamolybdate tetrahydrate (4) specified in JIS K 8905 while dissolving in water, and further add water to make 1000 mL. In the case of usage, mix equal volumes of the two solutions.

Comment 2 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard...
solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble phosphoric acid (C-P$_2$O$_5$) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.3) by a conversion factor (2.2914).

(3) **Instruments:** Instruments are as shown below:
   a) **Extractor:** Constant-temperature rotary shaker or reciprocating water bath shaker as described below.
      aa) **Constant-temperature rotary shaker:** A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.
      ab) **Reciprocating water bath shaker:** A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocating horizontally at 160 times/minute with amplitude of 25 mm - 40 mm.
   b) **Hot plate or sand bath:** A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.
   c) **Spectrophotometer:** A spectrophotometer specified in JIS K 0115

(4) **Test procedures**
(4.1) **Extraction:** Conduct extraction as shown below.
(4.1.1) **Constant-temperature rotary shaker:**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.
   b) Add 150 mL of citric acid solution heated up to about 30 °C (6), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.
   c) After immediate cooling is complete, add water up to the marked line
   d) Filter with Type 3 filter paper to make a sample solution.

**Note** (6) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 3** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker:**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (7).
   b) Add 150 mL of citric acid solution heated up to about 30 °C (6), and shake to mix by reciprocating horizontally at 160 times/min with amplitude of 25 mm - 40 mm (30 °C ± 1 °C) for 1 hour.
   c) After immediate cooling is complete, add water up to the marked line
   d) Filter with Type 3 filter paper to make a sample solution.

**Note** (7) Use a 250-mL flat-bottom volumetric flask to stabilize the shaking.

**Comment 4** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

**Comment 5** For a by-product phosphate fertilizer, if the pH of the sample solution in (4.1.1) d) and (4.1.2) d) is neutral or basic, prepare a sample solution anew by replacing “1 g of
an analytical sample” in the procedures in (4.1.1) a) and (4.1.2) a) with “0.5 g of an analytical sample”.

Comment 6 The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) Coloring: Conduct coloring as shown below.

a) Transfer a predetermined amount (up to 25 mL, the equivalents of 0.5 mg - 6 mg as P₂O₅) of the sample solution to a 100-mL - 200-mL tall beaker.
b) Add 3 mL of hydrochloric acid and 1 mL of nitric acid.
c) Cover the tall beaker with a watch glass, heat on a hot plate or sand bath at 200 ºC - 250 ºC and condense until the solution volume becomes about 2 mL.
d) After standing to cool, transfer it with water to a 100-mL volumetric flask.
e) Add the citric acid solution to make the equivalents of 17 mL of the citric acid and further add 2 mL of nitric acid (1+1).
f) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes.

Note (8) A watch glass should not be uncovered when bubbles form while heating because they may splash.

(9) It is recommended to transfer 2 mL of water to a 100-mL - 200-mL tall beaker in advance and confirm the volume.

(10) The volume of the solution after transferring should be up to about 50 mL.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.
Detection wavelength: 420 nm

b) Calibration curve preparation
1) Transfer 1 mL - 12 mL of phosphoric acid standard solution (P₂O₅ 0.5 mg/mL) to 100-mL volumetric flasks step-by-step.
2) Add 17 mL of a citric acid solution, then add 4 mL of nitric acid (1+1), further add a proper amount of water. Conduct the same procedure as (4.2) f) to make the P₂O₅ 0.5 mg/100 mL - 6 mg/100 mL phosphoric acid standard solutions for the calibration curve preparation.
3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.
4) Measure absorbance at a wavelength of 420 nm of the phosphoric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.
5) Prepare the calibration curve of the phosphoric acid concentration and absorbance of the phosphoric acid standard solutions for the calibration curve preparation.

c) Sample measurement
1) Regarding the solution in (4.2) f), measure absorbance by the same procedure as b) 4) (12).
2) Obtain the phosphoric acid (P₂O₅) content from the calibration curve, and calculate citrate soluble phosphoric acid (C-P₂O₅) in the analytical sample.

Note (11) If water is not added, precipitate may form when adding the coloring reagent solution.

(12) Measure within 2 hours after adding the coloring reagent solution in the procedure in
Comment 7 Additive recovery testing was conducted to evaluate trueness using solid fertilizers (10 samples) containing the equivalents of 1.03 % (mass fraction) - 51.04 % (mass fraction) as citrate soluble phosphoric acid. As a result, the average recovery is 99 % - 100 %.

The results of the repeatability tests on different days using a solid preparation sample to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Table 2 shows results and analysis results from a collaborative study for testing method validation.

Additionally, the minimum limit of quantification of this testing method is about 0.05 % (mass fraction).

Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Average ^2)</td>
<td>s_r ^4) RSD_r ^5)</td>
</tr>
<tr>
<td>Preparation sample (solid) 1</td>
<td>5</td>
<td>51.01</td>
<td>0.12 0.2</td>
</tr>
<tr>
<td>Preparation sample (solid) 2</td>
<td>5</td>
<td>2.57</td>
<td>0.01 0.6</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) x the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
Table 2  Results and analysis results from a collaborative study for the test method validation of citrate-soluble phosphoric acid

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^2)$</th>
<th>$s_r^{4)$</th>
<th>$RSD_r^{5)$</th>
<th>$s_R^{6)$</th>
<th>$RSD_R^{7)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Processed phosphorus fertilizers 1</td>
<td>12</td>
<td>47.21</td>
<td>0.13</td>
<td>0.3</td>
<td>0.69</td>
<td>1.5</td>
</tr>
<tr>
<td>Compound fertilizers 1</td>
<td>11</td>
<td>17.71</td>
<td>0.07</td>
<td>0.4</td>
<td>0.19</td>
<td>1.1</td>
</tr>
<tr>
<td>Compound fertilizers 2</td>
<td>12</td>
<td>5.08</td>
<td>0.08</td>
<td>1.6</td>
<td>0.17</td>
<td>3.3</td>
</tr>
<tr>
<td>Absorptive mixed fertilizer</td>
<td>11</td>
<td>14.32</td>
<td>0.06</td>
<td>0.4</td>
<td>0.18</td>
<td>1.2</td>
</tr>
<tr>
<td>Regent</td>
<td>11</td>
<td>50.89</td>
<td>0.14</td>
<td>0.3</td>
<td>0.57</td>
<td>1.1</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n = number of laboratories x number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(5) **Flow sheet for citrate-soluble phosphoric acid containing phosphorus acid, etc.:** The flow sheet for citrate-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. is shown below:

![Flow sheet diagram](image)

Figure 1-1 Flow sheet for citrate-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Extraction procedure (4.1.1))

![Flow sheet diagram](image)

Figure 1-2 Flow sheet for citrate-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Extraction procedure (4.1.2))
Figure 2 Flow sheet for citrate-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Coloring and measurement procedure)
4.2.3.c Quinoline gravimetric analysis

(1) **Summary**
This testing method is applicable to fertilizers containing no Phosphorous acid, etc. It is suitable for the fertilizers with relatively a high content of phosphoric acid. This testing method is classified as Type E and its symbol is 4.2.3.c-2017 or C-P.c-1.

Extract by adding a citric acid solution to an analytical sample. Heat after adding nitric acid and water, hydrolyze nonorthophosphoric acid to orthophosphate ion and measure the mass of quinolinium phosphomolybdate formed by the reaction with quinoline, molybdic acid and nitric acid to obtain citrate soluble phosphoric acid (C-P$_2$O$_5$) in an analytical sample.

(2) **Reagent**: Reagents are as shown below.
   a) **Nitric acid**: A JIS Guaranteed (HNO$_3$ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   b) **Citric acid solution** (1). Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   c) **Sodium molybdate solution**: Dissolve 70 g of sodium molybdate dihydrate in 150 mL of water.
   d) **Quinoline solution**: Add 5 mL of quinoline specified in JIS K 8279 to the mixture solution of 35 mL of nitric acid and 100 mL of water.
   e) **Quimosiac solution**: Add 60 g of citric acid monohydrate specified in JIS K 8283 to the mixture solution of 85 mL nitric acid and 150 mL of water to dissolve. Add gradually total volume of the sodium molybdate solution to mix. Add gradually the total volume of the quinoline solution while mixing the solution. After leaving at rest overnight, filter the total volume with Type 3 filter paper. Add 280 mL of acetone specified in JIS K 8034, and further add water to make 1000 mL.

**Note**  (1) This is an example of preparation; prepare an amount as appropriate.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.
   a) **Constant-temperature rotary shaker**: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.
   b) **Water bath**: Water bath that can be adjusted to 60 °C - 65 °C.
   c) **Drying apparatus**: A drying apparatus that can be adjusted to 220 °C ± 5 °C.
   d) **Crucible type glass filter**: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 220 °C ± 5 °C in advance and measure the mass to the order of 1 mg.

(4) **Test procedures**
(4.1) **Extraction**: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.
   b) Add 150 mL of citric acid solution heated up to about 30 °C (2), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.
   c) After immediate cooling is complete, add water up to the marked line
   d) Filter with Type 3 filter paper to make a sample solution.

**Note**  (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.
Comment 1  The procedures in (4.1) are the same as in (4.1.1) in 4.2.3.a.

Comment 2  For a by-product phosphate fertilizer or a fertilizer containing a by-product phosphate, if the pH of the sample solution of d) is neutral or basic, prepare a sample solution anew by replacing “1 g of an analytical sample” in the procedure in a) with “0.5 g of an analytical sample”.

Comment 3  The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedure of (4.1) b).

(4.2) Measurement: Conduct measurement as shown below.

a) Transfer a predetermined volume (the equivalents of 10 mg - 30 mg as P$_2$O$_5$) of sample solution to a 300-mL tall beaker.

b) Add 5 mL of nitric acid and add water to make about 80 mL.

c) Cover with a watch glass. After boiling for about 3 minutes, wash the watch glass and the inside the tall beaker with water and add water to make about 100 mL.

d) Immediately, add 50mL of quimosiac solution, heat for about 15 minutes while sometimes mixing in a water bath at 60 °C - 65 °C to produce the precipitate of quinolinium phosphomolybdate.

e) After standing to cool down to room temperature while sometimes mixing, filter under reduced pressure with a crucible type glass filter, wash the tall beaker 3 times with water and transfer the whole precipitate into a crucible type glass filter, and further wash 7 - 8 times with water.

f) Transfer the precipitate together with the crucible type glass filter into a drying apparatus and heat at 220 °C ± 5 °C for about 30 minutes.

 g) As soon as heating is complete, move it into a desiccator and let it stand to cool.

h) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

i) Calculate citrate soluble phosphoric acid (C-P$_2$O$_5$) in the analytical sample by the following formula.

\[
\text{Citrate soluble phosphoric acid (C-P$_2$O$_5$) (% (mass fraction)) in an analytical sample} = A \times 0.03207 \times \left( \frac{V_1}{V_2} \right) \times \left( \frac{1}{W} \right) \times 100
\]

$A$: Mass (g) of the precipitate in h)

$W$: Mass of an analytical sample (1 g)

$V_1$: Predetermined volume (250 mL) of the sample solution

$V_2$: Volume (mL) of the sample solution transferred in a)

References

(5) **Flow sheet for citrate soluble phosphoric acid**: The flow sheet for citrate soluble phosphoric acid in fertilizers is shown below:

<table>
<thead>
<tr>
<th>1 g analytical sample</th>
<th>Weigh to the order of 1 mg to a 250-mL volumetric flask ← 150 mL of citric acid solution [about 30 ºC]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shaking to mix</td>
<td>Constant-temperature rotary shaker (30 - 40 revolutions/min) 30 ºC ± 1 ºC, for 1 hour</td>
</tr>
<tr>
<td>Cooling</td>
<td>Immediately</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Filtration</td>
<td></td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1  Flow sheet for citrate soluble phosphoric acid in fertilizers (Extraction procedure)
Figure 2  Flow sheet for citrate soluble phosphoric acid in fertilizers (Measurement procedure)
4.2.3.d ICP Optical Emission Spectrometry

(1) Summary

This testing method is applicable to fertilizers. It is also suitable for the fertilizers containing phosphite. This testing method is classified as Type D and its symbol is 4.2.3.d-2018 or C-P.d-1. Extract by adding a citric acid solution to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the phosphorus at a wavelength of 178.287 nm to obtain citric acid-soluble phosphoric acid (citrate-soluble phosphoric acid (C-P₂O₅)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

(2) Reagent: Reagents are as shown below.


b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Citric acid solution: Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.

d) Phosphoric acid standard solution (P₂O₅ 10 mg/mL): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL-3 mL of nitric acid, and add water up to the marked line.

e) Phosphoric acid standard solution (P₂O₅ 1 mg/mL): Transfer 10 mL of phosphoric acid standard solution (P₂O₅ 10 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.

f) Phosphoric acid standard solution (P₂O₅ 20 µg/mL - 0.4 mg/mL) for the calibration curve preparation: Transfer 2 mL - 40 mL of phosphoric acid standard solution (P₂O₅ 1 mg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line.

g) Phosphoric acid standard solution ((P₂O₅ 5 µg/mL - 20 µg/mL) for the calibration curve preparation: Transfer 5 mL - 20 mL of phosphoric acid standard solution (P₂O₅ 0.1 mg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line.

h) Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in e) - g).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble phosphoric acid (C-P₂O₅) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.2) by a conversion factor (2.2914).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in f) and g) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a phosphoric acid
standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) **Instruments**: Instruments are as shown below:

a) **ICP Optical Emission Spectrometry**: A spectrophotometer specified in JIS K 0115

   1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

b) **Extractor**: Constant-temperature rotary shaker or reciprocating water bath shaker as described below.

   ba) **Constant-temperature rotary shaker**: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

   bb) **Reciprocating water bath shaker**: A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/minute with amplitude of 25 mm - 40 mm.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker**:

   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

   b) Add 150 mL of citric acid solution heated up to about 30 °C \(^{(2)}\) and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.

   c) After immediate cooling is complete, add water up to the marked line

   d) Filter with Type 3 filter paper to make a sample solution.

   **Note** (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

   **Comment 3** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker**:

   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask \(^{(3)}\).

   b) Add 150 mL of citric acid solution heated up to about 30 °C \(^{(2)}\), and shake to mix by reciprocating horizontally at 160 times/min with amplitude of 25 mm - 40 mm (30 °C ± 1 °C) for 1 hour.

   c) After immediate cooling is complete, add water up to the marked line

   d) Filter with Type 3 filter paper to make a sample solution.

   **Note** (3) Use a 250-mL flat-bottom volumetric flask to stabilize the shaking.

   **Comment 4** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

   **Comment 5** For a by-product phosphate fertilizer, if the pH of the sample solution in (4.1.1) d) and (4.1.2) d) is neutral or basic, prepare a sample solution anew by replacing “1 g of an analytical sample” in the procedures in (4.1.1) a) and (4.1.2) a) with “0.5 g of an analytical sample”.

   **Comment 6** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters.
after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
   Analytical line wavelength: 178.287 nm

b) Calibration curve preparation
   1) Spray the phosphoric acid standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 178.287 nm.
   2) Prepare a curve for the relationship between the phosphoric acid concentration and the indicated value of the phosphoric acid standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 40 mg as \( \text{P}_2\text{O}_5 \)) to a 100-mL volumetric flask.
   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the phosphoric acid (\( \text{P}_2\text{O}_5 \)) content from the calibration curve, and calculate citrate soluble phosphoric acid (C-\( \text{P}_2\text{O}_5 \)) in the analytical sample.

Comment 7 The simultaneous measurement of multiple elements by an ICP-OES is available. In this case, transfer a pre-determined amount of phosphoric acid standard solution (P 1 mg/mL or 10 mg/mL), potassium standard solution (K 1 mg/mL or 10 mg/mL), magnesium standard solution (Mg 1 mg/mL or 10 mg/mL), manganese standard solution (Mn 1 mg/mL or 10 mg/mL) and boron standard solution (B 1 mg/mL or 10 mg/mL) traceable to National Metrology to a volumetric flask to mix, add hydrochloric acid (1+5) to make an acid concentration of 0.5 mol/L and further add water up to the marked line to prepare a primary mixed standard solution. Transfer a pre-determined volume of primary mixed standard solution to volumetric flasks step-by-step, add hydrochloric acid (1+23) up to the marked line to prepare mixed standard solutions for calibration curve preparation within the concentration range in Table 1. In this case, multiply the respective concentrations of mixed standard solutions for calibration curve preparation or measurement values obtained in (4.2) by the conversion factors in Table 1 to calculate respective main components in an analytical sample. The measurement wavelengths of respective elements are according to Table 1. In addition, when preserving mixed standard solutions for calibration curve preparation, use a container, which can be sealed tightly, made of material such as PTFE that boron hardly elutes.
Comment 8 The comparison of the measurement value ($y_1$: 1.74 % (mass fraction) - 49.04 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of Ammonium vanadomolybdate absorptiometric analysis was conducted to evaluate trueness using processed phosphorus fertilizers (2 samples), compound fertilizers (12 samples), home garden-use mixed fertilizers (1 sample), mixed compost fertilizers (2 samples), mixed phosphate fertilizers (2 samples), designated blended fertilizers (4 samples), blended fertilizers (5 samples), byproduct mixed fertilizers (1 sample), byproduct phosphorus fertilizers (2 samples), organic compound fertilizers (1 sample) and fused phosphate fertilizers (1 sample). As a result, a regression equation was $y = -0.0027 + 1.001x$, and its correlation coefficient ($r$) was 1.000. In addition, recovery testing was conducted using a preparation sample. As a result, the average rate of recovery at the content level of 0.260 % (mass fraction) - 49.99 % (mass fraction) was 96.3 % - 100.8 %.

The results of the repeatability tests on different days using compound fertilizers and blended fertilizer to evaluate precision were analyzed by one-way analysis of variance. Table 2 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction).
<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average ( T^{(1)} ) ( (%)^{(3)} )</th>
<th>Repeatability ( s^{(4)} ) ( (%)^{(3)} )</th>
<th>( RSD_T^{(5)} ) ( (%) )</th>
<th>Intermediate precision ( s_{I(T)}^{(6)} ) ( (%) )</th>
<th>( RSD_{I(T)}^{(7)} ) ( (%) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>20.90</td>
<td>0.13</td>
<td>0.6</td>
<td>0.18</td>
<td>0.9</td>
</tr>
<tr>
<td>Blended fertilizer</td>
<td>7</td>
<td>6.44</td>
<td>0.06</td>
<td>0.9</td>
<td>0.06</td>
<td>1.0</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average \( (\)the number of test days \( T \) \( (\times\) the number of duplicate testing \( (2) \) \( )\)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

(5) Flow sheet for citrate soluble phosphoric acid: The flow sheet for citrate soluble phosphoric acid in fertilizers is shown below:

1 g analytical sample
→ Weigh to the order of 1 mg to a 250-mL volumetric flask

150 mL of citric acid solution [about 30 °C]

Shaking to mix
→ Constant-temperature rotary shaker (30 - 40 revolutions/min)

30 °C ± 1 °C, for 1 hour

Cooling
→ Immediately

Water (up to the marked line)

Filtration
→ Type 3 filter paper

Sample solution

Figure 1-1  Flow sheet for citrate soluble phosphoric acid in fertilizers
(Extraction procedure (4.1.1))
Weigh to the order of 1 mg to a 250-mL volumetric flask

→ 150 mL of citric acid solution [about 30 ºC]

Reciprocating water bath shaker (reciprocation horizontally at 160 times /min, with amplitude of 25 mm - 40 mm), at 30 ºC ± 1 ºC, for 1 hour

Immediately

← Water (up to the marked line)

Type 3 filter paper

Figure 1-2 Flow sheet for citrate soluble phosphoric acid in fertilizers
(Extraction procedure (4.1.2))

Sample solution

→ 100-mL volumetric flask

→ 25 mL of hydrochloric acid (1+5)

→ Water (up to the marked line)

Measurement

ICP Optical Emission Spectrometry (178.287 nm)

Figure 2 Flow sheet for citrate soluble phosphoric acid in fertilizers
(Measurement procedure)
4.2.4 Water-soluble phosphoric acid
4.2.4.a Ammonium vanadomolybdate absorptiometric analysis

(1) Summary
This testing method is applicable to fertilizers that do not contain matter not colored by hydrolysis with nitrate acids such as phosphonic acid. This testing method is classified as Type C and its symbol is 4.2.4.a-2017 or W-P.a-1.

Extract by adding water to an analytical sample, add nitric acid (1+1) to heat, and hydrolyze nonorthophosphoric acid to orthophosphate ion, then measure the absorbance of phosphovanadomolybdate salt formed by the reaction with ammonium vanadate (V), hexaammonium heptamolybdate and nitric acid to obtain water-soluble phosphoric acid (W-P$_2$O$_5$) in an analytical sample. In addition, the performance of this testing method is shown in Comment 9.

(2) Reagent: Reagents are as shown below.
   a) Nitric acid: A JIS Guaranteed (HNO$_3$ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   b) Ammonia solution: A JIS Guaranteed (NH$_3$ 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.
   c) Coloring reagent solution (1) (2). Dissolve 1.12 g of ammonium vanadate (V) (3) specified in JIS K 8747 in water, add 250 mL of nitric acid, then add 27 g of hexaammonium heptamolybdate tetrahydrate (4) specified in JIS K 8905 dissolved in water, and further add water to make 1000 mL (5).
   d) Phenolphthalein solution (1 g/100 mL): Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.
   e) Phosphoric acid standard solution (P$_2$O$_5$ 10 mg/mL) (1): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.
   f) Phosphoric acid standard solution (P$_2$O$_5$ 0.5 mg/mL) (1): Transfer 50 mL of phosphoric acid standard solution (P$_2$O$_5$ 10 mg/mL) to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) This corresponds to reagent “a” reagent solution in the Official Methods of Analysis of Fertilizers (1992).
   (3) This corresponds to ammonium metavanadate in the Official Methods of Analysis of Fertilizers (1992).
   (4) This corresponds to ammonium molybdate in the Official Methods of Analysis of Fertilizers (1992).
   (5) Store in an amber bottle.

Comment 1 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble phosphoric acid (W-P$_2$O$_5$) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.3) by a conversion factor (2.2914).
(3) **Instruments:** Instruments are as shown below:

a) **Extractor:** A rotary shaker or a vertical reciprocating shaker as described below.

aa) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.

ab) vertical reciprocating shaker: A vertical reciprocating shaker that can shake a 250-mL volumetric flask using an adapter to reciprocate vertically at 300 times/minute (amplitude of 40 mm).

b) **Hot plate:** A hot plate whose surface temperature can be adjusted up to 250 ºC.

c) **Spectrophotometer:** A spectrophotometer specified in JIS K 0115

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

**4.1.1) Powdery test sample**

(4.1.1.1) **Rotary shaker:**

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.

b) Add about 400 mL of water, and shake to mix at 30 - 40 revolutions/min for about 30 minutes.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

Comment 2 In the procedure of (4.1.1.1) a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it in a 250-mL volumetric flask

Comment 3 The procedure in (4.1.1.1) is the same as the procedure in (4.1.1) of 4.1.2.b, (4.1.1.1) of 4.2.4.b, (4.1) of 4.2.4.c, (4.1) of 4.2.4.d, (4.1.2.1) of 4.3.3.a, (4.1.2) of 4.3.3.b, (4.1.2) of 4.3.3.c, (4.1.1.1) of 4.7.3.a, (4.1.1) of 4.9.2.a, (4.1) of 4.9.2.c, (4.1.1) of 4.9.2.d, (4.1.2.1) of 4.3.3.a and (4.1.1) of 4.14.1.a.

Comment 4 When the determination is affected by the coloring of the sample solution of (4.1.1.1) d), (4.1.1.2) d) and (4.1.2) d), transfer the predetermined volume of the sample solution to a 100-mL volumetric flask, add a few drops of hydrochloric acid (1+1) to acidify, then add no more than 0.1 g of active carbon. After leaving at rest for a little while, add water up to the marked line and filter. The filtrate is prepared as the sample solution of (4.2) a). Additionally, as phosphorus contained in active carbon has the possibility to elute and affects the determination value, a blank test is required.

(4.1.1.2) **Vertical reciprocating shaker:**

a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

b) Add about 200 mL of water, and shake to mix by reciprocating vertically at 300 times/min (amplitude of 40 mm) for about 30 minutes.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

Comment 5 The procedure in (4.1.1.2) is the same as the procedures in (4.1.1.2) of 4.2.4.b, (4.1.1.2) of 4.3.3.a and (4.1.1.2) of 4.7.3.a.

(4.1.2) **Fluid test sample**

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 100-mL volumetric flask.

b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

**Comment 6** The procedure in (4.1.2) is the same as the procedure in (4.1.2) of 4.2.4.b, (4.1.3) of 4.3.3.a, (4.1) of 4.3.3.d, (4.1) of 4.5.3.b, (4.1) of 4.6.3.b, (4.1.2) of 4.7.3.a, (4.1) of 4.7.3.b, (4.1) of 4.8.2.b, (4.1.2) of 4.9.2.a, (4.1.2) of 4.9.2.b, (4.1.2) of 4.10.2.a, (4.1) of 4.10.2.b, (4.1.2) of 4.13.1.a, (4.1) of 4.13.1.b, (4.1.2) of 4.14.1.a, (4.1) of 4.14.1.b, (4.1.2) of 4.14.1.a, (4.1) of 4.14.1.b, (4.1) of 4.15.1.a and (4.1) of 4.15.1.b.

(4.2) **Coloring:** Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 6 mg as \( \text{P}_2\text{O}_5 \)) to a 100-mL volumetric flask.

b) Add 4 mL of nitric acid (1+1)\(^{(6)}\), and heat to boil \(^{(7)}\).

c) After cooling is complete, add 1 - 2 drop(s) of phenolphthalein solution (1 g/100 mL), and neutralize by adding ammonia solution (1+1) until the color of the solution becomes light red-purple \(^{(14)}\).

d) Add nitric acid (1+10) until the light red-purple color of the solution disappears to make it slightly acidic, and add a proper amount of water \(^{(8)}\).

e) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes \(^{(6)}\).

**Comment 7** The volumetric flask used in the procedure in a) should be distinguished as a flask to be used for phosphate coloring operation and should not be used for other purposes.

**Note** (6) When the solution is muddled by adding nitric acid (1+1), conduct centrifugation after the procedure in e).

(7) When it does not contain non-orthophosphate, the procedure in b) is not necessary.

(8) Without the addition of water, precipitate may be produced when a coloring reagent solution is added.

(4.3) **Measurement:** Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) **Measurement conditions of spectrophotometer:** Set up the measurement conditions of spectrophotometer considering the following.
Detection wavelength: 420 nm

b) **Calibration curve preparation**

1) Transfer 1 mL - 12 mL of phosphoric acid standard solution (\( \text{P}_2\text{O}_5 \) 0.5 mg/mL) to 100-mL volumetric flasks step-by-step.

2) Add a proper amount of water \(^{(8)}\), and conduct the same procedure as (4.2) e) to make the \( \text{P}_2\text{O}_5 \) 0.5 mg/100 mL - 6 mg/100 mL phosphoric acid standard solutions for the calibration curve preparation.

3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.

4) Measure absorbance at a wavelength of 420 nm of the phosphoric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control \(^{(9)}\).

5) Prepare the calibration curve of the phosphoric acid concentration and absorbance of the phosphoric acid standard solutions for the calibration curve preparation.

c) **Sample measurement**
1) Regarding the solution in (4.2) e), measure absorbance by the same procedure as b) 4) (9).

2) Obtain the phosphoric acid ($P_2O_5$) content from the calibration curve, and calculate water-soluble phosphoric acid (W-$P_2O_5$) in the analytical sample.

**Note** (9) Measure within 6 hours after adding the coloring reagent solution in the procedure in (4.2) e).

**Comment 8** After the procedure in (4.2) a), it is also possible to measure soluble phosphoric acid at the same time by adding 2 mL of Petermans citrate solution and by conducting the procedures from (4.2) d) to (4.3) in 4.2.2.a (using b reagent solution in the Official Methods of Analysis of Fertilizers (1992)).

Further after the procedure in (4.2) a), it is also possible to measure citrate soluble phosphoric acid at the same time by adding 17 mL of citrate solution and by conducting the procedures from (4.2) c) to (4.3) in 4.2.3.a (using b reagent solution in the Official Methods of Analysis of Fertilizers (1992)).

**Comment 9** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 100.5 % - 101.2 % and 99.0 % - 101.7 % as water-soluble phosphoric acid (W-$P_2O_5$) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

The comparison of the measurement value of extraction ($y_i$: 0.292 % (mass fraction) - 40.40 % (mass fraction)) with a vertical reciprocating shaker and the measurement value of extract with a rotary shaker ($x_i$) was conducted to evaluate trueness of the extraction of solid fertilizers using fertilizers (12 samples). As a result, a regression equation was $y = -0.041 + 0.999x$, and its correlation coefficient ($r$) was 1.000. Also, the results of the repeatability tests on different days using compound fertilizers and designated blended fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 2 shows the calculation results of intermediate precision and repeatability.

The comparison of the measurement value of simple extraction ($y_i$: 1.92 % (mass fraction) - 12.21 % (mass fraction )) and the measurement value of extract ($x_i$) with a rotary shaker was conducted to evaluate trueness of the extraction of fluid fertilizers using fluid fertilizers (12 samples). As a result, a regression equation was $y = 0.005+1.005x$, and its correlation coefficient ($r$) was 0.999. Also, the results of the repeatability tests on different days using fluid mixed fertilizers to evaluate the precision of the extraction of fluid fertilizers were analyzed by the one-way analysis of variance. Table 3 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.03 % (mass fraction) for solid fertilizers, and 0.004 % (mass fraction) for fluid fertilizers.
Table 1  Analysis results of the collaborative study to determine certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratories</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
<th>Reproducibility</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$p^{1)}$</td>
<td>Average $^{2)}$ ($%)^{3)}$</td>
<td>$s_r^{4)}$ ($%)^{3)}$</td>
<td>$RSD_r^{5)}$ ($%)^{3)}$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>($%)^{3)}$</td>
<td>($%)^{3)}$</td>
<td>($%)^{3)}$</td>
</tr>
<tr>
<td>FAMIC-B-10</td>
<td>9</td>
<td>7.00</td>
<td>0.02</td>
<td>0.3</td>
</tr>
<tr>
<td>FAMIC-B-14</td>
<td>15</td>
<td>6.70</td>
<td>0.02</td>
<td>0.3</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis; 2) Average (the number of laboratory $p \times$ test days $T$) $\times$ the number of replicate testing; 3) Mass fraction; 4) Repeatability standard deviation; 5) Repeatability relative standard deviation; 6) Intermediate standard deviation; 7) Intermediate relative standard deviation; 8) Reproducibility standard deviation; 9) Reproducibility relative standard deviation.

1) Masayoshi KOSHINO: Second Revision of The Methods of Analysis of Fertilizers.


Flow sheet for water-soluble phosphoric acid: The flow sheet for water-soluble phosphoric acid in fertilizers is shown below:

![Flow sheet for water-soluble phosphoric acid](image)

Figure 1-1 Flow sheet for water-soluble phosphoric acid in fertilizers (Extraction procedure (4.1.1.1))

![Flow sheet for water-soluble phosphoric acid](image)

Figure 1-2 Flow sheet for water-soluble phosphoric acid in fertilizers (Extraction procedure (4.1.1.2))

![Flow sheet for water-soluble phosphoric acid](image)

Figure 1-3 Flow sheet for water-soluble phosphoric acid in fertilizers (Extraction procedure (4.1.2))
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot (predetermined volume)</td>
<td>← 4 mL of nitric acid (1:1)</td>
</tr>
<tr>
<td>Heating</td>
<td>Boiling</td>
</tr>
<tr>
<td>Cooling</td>
<td>← 1-2 drop(s) of phenolphthalein solution (1 g/100 mL)</td>
</tr>
<tr>
<td></td>
<td>← Ammonia solution (1:1) [neutralization]</td>
</tr>
<tr>
<td></td>
<td>← Nitric acid (1:10) [slightly acidic]</td>
</tr>
<tr>
<td></td>
<td>← A proper amount of water</td>
</tr>
<tr>
<td></td>
<td>← 20 mL of coloring reagent solution</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Leaving at rest</td>
<td>For about 30 minutes</td>
</tr>
<tr>
<td>Measurement</td>
<td>Spectrophotometer (420 nm)</td>
</tr>
</tbody>
</table>

Figure 2 Flow sheet for water-soluble phosphoric acid in fertilizers
(Coloring and measurement procedure)
4.2.4.b Ammonium vanadomolybdate absorptiometric analysis (Fertilizers containing phosphorous acid or phosphate)

(1) Summary
This testing method is applicable to the fertilizers containing phosphorous acid or phosphate. This testing method is classified as Type B and its symbol is 4.2.4.b-2017 or W-P.b-1.

Extract by adding water to an analytical sample, add hydrochloric acid - sulfuric acid to heat, and oxygenate phosphonic acid ion to orthophosphate ion, and then measure the absorbance of phosphovanadomolybdate salt formed by the reaction with ammonium vanadate (V), hexaammonium heptamolybdate and nitric acid to obtain water-soluble phosphoric acid (W-P$_2$O$_5$) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Nitric acid: A JIS Guaranteed (HNO$_3$ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   c) Ammonia solution: A JIS Guaranteed (NH$_3$ 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.
   d) Coloring reagent solution (1) (2): Dissolve 1.12 g of ammonium vanadate (V) (3) specified in JIS K 8747 in water, add 250 mL of nitric acid, then add 27 g of hexaammonium heptamolybdate tetrahydrate (4) specified in JIS K 8905 dissolved in water, and further add water to make 1000 mL (5).
   e) Phenolphthalein solution (1 g/100 mL): Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.
   f) Phosphoric acid standard solution (P$_2$O$_5$ 10 mg/mL) (1): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.
   g) Phosphoric acid standard solution (P$_2$O$_5$ 0.5 mg/mL) (1): Transfer 50 mL of phosphoric acid standard solution (P$_2$O$_5$ 10 mg/mL) to a 1000-mL volumetric flask, add 2 mL - 3 mL of nitric acid, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) This corresponds to reagent “a” reagent solution in the Official Methods of Analysis of Fertilizers (1992).
   (3) This corresponds to ammonium metavanadate in the Official Methods of Analysis of Fertilizers (1992).
   (4) This corresponds to ammonium molybdate in the Official Methods of Analysis of Fertilizers (1992).
   (5) Store in an amber bottle.

Comment 1 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble phosphoric acid (W-P$_2$O$_5$) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.3) by a conversion factor (2.2914).
(3) **Instruments**: Instruments are as shown below:

a) **Extractor**: A rotary shaker or a vertical reciprocating shaker as described below.

aa) **Rotary shaker**: A rotary shaker that can rotate a 250-mL - 500-mL volumetric flask upside down at 30 - 40 revolutions/min.

ab) **Vertical reciprocating shaker**: A vertical reciprocating shaker that can shake a 250-mL volumetric flask using an adapter to reciprocate vertically at 300 times/minute (amplitude of 40 mm).

b) **Hot plate or sand bath**: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

c) **Spectrophotometer**: A spectrophotometer specified in JIS K 0115

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Powdery test sample**

(4.1.1.1) **Rotary shaker**:

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.

b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

**Comment 2** In the procedure of (4.1.1.1) a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it in a 250-mL volumetric flask.

**Comment 3** The procedure in (4.1.1.1) is the same as the procedure in (4.1.2.1.1) in 4.2.4.a.

(4.1.1.2) **Vertical reciprocating shaker**:

a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

b) Add about 200 mL of water, and shake to mix by reciprocating vertically at 300 times/minute (amplitude of 40 mm) for about 30 minutes.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

**Comment 4** The procedure in (4.1.1.2) is the same as the procedure in (4.1.1.2) in 4.2.4.a.

(4.1.2) **Fluid test sample**

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 100-mL volumetric flask.

b) Add about 50 mL of water, and shake to mix.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

**Comment 5** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.4.a.

(4.2) **Coloring**: Conduct coloring as shown below.

a) Transfer a predetermined amount (the equivalents of 0.5 mg - 6 mg as P₂O₅) of the sample solution to a 100-mL - 200-mL tall beaker.

b) Add 3 mL of hydrochloric acid and 1 mL of nitric acid.
c) Cover the tall beaker with a watch glass, heat on a hot plate or sand bath at 200 °C - 250 °C and condense until the solution volume becomes about 2 mL.

d) After standing to cool, transfer it with water to a 100- mL volumetric flask.

e) Add 1 - 2 drop(s) of phenolphthalein solution (1 g/100 mL), and neutralize by adding ammonia solution (1+1) until the color of the solution becomes light red-purple.

f) Add nitric acid (1+10) until the light red-purple color of the solution disappears to make it slightly acidic.

g) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes.

Note (6) It is recommended to transfer about 2 mL of water to a 100-mL - 200-mL tall beaker in advance and confirm the volume.

(7) Care should be taken not to dry and harden it. When it dries completely, the determined value becomes lower than usual in some cases.

(8) The volume of the solution after transferring should be up to about 50 mL.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.

Detection wavelength: 420 nm

b) Calibration curve preparation

1) Transfer 1 mL - 12 mL of phosphoric acid standard solution (P₂O₅ 0.5 mg/mL) to 100-mL volumetric flasks step-by-step.

2) Add a proper amount of water, and conduct the same procedure as (4.2) g) to make the P2O5 0.5 mg/100 mL - 6 mg/100 mL phosphoric acid standard solutions for the calibration curve preparation.

3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.

4) Measure absorbance at a wavelength of 420 nm of the phosphoric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.

5) Prepare the calibration curve of the phosphoric acid concentration and absorbance of the phosphoric acid standard solutions for the calibration curve preparation.

c) Sample measurement

1) Regarding the solution in (4.2) g), measure absorbance by the same procedure as b) 4)⁴¹⁰.

2) Obtain the phosphoric acid (P₂O₅) content from the calibration curve, and calculate water-soluble phosphoric acid (W-P₂O₅) in the analytical sample.

Note (9) If water is not added, precipitate may form when adding the coloring reagent solution.

(10) Measure within 6 hours after adding the coloring reagent solution in the procedure in (4.2) g).

Comment 6 Recovery testing was conducted to evaluate trueness using a fluid preparation sample. As a result, the average rate of recovery at the content level of 30 % (mass fraction) - 50 % (mass fraction), 10 % (mass fraction) - 20 % (mass fraction), 4 % (mass fraction) and 0.2 % (mass fraction) are 101.1 % - 101.8 %, 101.1 % - 101.5 %, 100.8 % and 102.5 % as water-soluble phosphoric acid (W-P₂O₅) respectively. In the case of using a solid preparation sample, the average rate of recovery at the content
level of 30 \% (mass fraction) - 59 \% (mass fraction), 12 \% (mass fraction) - 21 \% (mass fraction) and 1 \% (mass fraction) - 9 \% (mass fraction) are 99.5 \% - 100.4 \%, 99.3 \% - 100.3 \%, and 96.9 \% - 100.4\% respectively.

The results of the repeatability tests on different days using a solid preparation sample to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Also, table 2 and table 3 show results and analysis results from a collaborative study for testing method validation using fluid fertilizers and solid fertilizers.

Additionally, the minimum limit of quantification of this testing method is about 0.04 \% (mass fraction) for solid fertilizers, and 0.01 \% (mass fraction) for fluid fertilizers.

### Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$</th>
<th>Repeatability $^3$</th>
<th>Intermediate precision $^4$</th>
<th>Intermediate precision $^5$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preparation sample (solid) 1</td>
<td>7</td>
<td>59.36</td>
<td>0.09</td>
<td>0.2</td>
<td>0.13</td>
</tr>
<tr>
<td>Preparation sample (solid) 2</td>
<td>7</td>
<td>5.90</td>
<td>0.07</td>
<td>1.2</td>
<td>0.07</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) \times the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
### Table 2  Results and analysis results from a collaborative study for the test method validation of water-soluble phosphoric acid

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (%)</th>
<th>$s_r$ (%)</th>
<th>$RSD_r$ (%)</th>
<th>$s_R$ (%)</th>
<th>$RSD_R$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizers 1</td>
<td>12</td>
<td>33.56</td>
<td>0.25</td>
<td>0.7</td>
<td>0.59</td>
<td>1.8</td>
</tr>
<tr>
<td>Fluid mixed fertilizers 2</td>
<td>12</td>
<td>17.93</td>
<td>0.08</td>
<td>0.5</td>
<td>0.30</td>
<td>1.7</td>
</tr>
<tr>
<td>Fluid mixed fertilizers 3</td>
<td>12</td>
<td>7.99</td>
<td>0.12</td>
<td>1.5</td>
<td>0.31</td>
<td>3.8</td>
</tr>
<tr>
<td>Fluid mixed fertilizers 4</td>
<td>11</td>
<td>11.93</td>
<td>0.13</td>
<td>1.1</td>
<td>0.33</td>
<td>2.8</td>
</tr>
<tr>
<td>Fluid mixed fertilizers 5</td>
<td>11</td>
<td>24.1</td>
<td>0.08</td>
<td>0.3</td>
<td>0.47</td>
<td>2.0</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis  
2) Mean ($n =$ number of laboratories x number of samples (2))  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Reproducibility standard deviation  
7) Reproducibility relative standard deviation

### Table 3  Results and analysis results from a collaborative study for the test method validation of water-soluble phosphoric acid (Solid fertilizer)

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (%)</th>
<th>$s_r$ (%)</th>
<th>$RSD_r$ (%)</th>
<th>$s_R$ (%)</th>
<th>$RSD_R$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizers 1</td>
<td>12</td>
<td>58.47</td>
<td>0.13</td>
<td>0.2</td>
<td>0.42</td>
<td>0.7</td>
</tr>
<tr>
<td>Compound fertilizers 2</td>
<td>12</td>
<td>3.92</td>
<td>0.04</td>
<td>1.0</td>
<td>0.08</td>
<td>2.1</td>
</tr>
<tr>
<td>Compound fertilizers 3</td>
<td>12</td>
<td>13.37</td>
<td>0.10</td>
<td>0.7</td>
<td>0.20</td>
<td>1.5</td>
</tr>
<tr>
<td>Absorptive mixed fertilizers</td>
<td>12</td>
<td>7.16</td>
<td>0.03</td>
<td>0.4</td>
<td>0.16</td>
<td>2.3</td>
</tr>
<tr>
<td>Blended fertilizers</td>
<td>12</td>
<td>21.80</td>
<td>0.12</td>
<td>0.5</td>
<td>0.18</td>
<td>0.8</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis  
2) Mean ($n =$ number of laboratories x number of samples (2))  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Reproducibility standard deviation  
7) Reproducibility relative standard deviation
References
4) Masayuki YAMANISHI, Toshiaki HIROI and Fumika TAKATSU Determination Method for Citric Acid-Soluble and Water-Soluble Phosphorus in Solid Fertilizer Containing Phosphonic Acid or Phosphonate (Phosphite) using Spectrophotometer: A Collaborative Study, Vol.9, p. 59 - 68 (2016)

(5) Flow sheet for water-soluble phosphoric acid containing phosphorus acid, etc.: The flow sheet for water-soluble phosphoric acid in fertilizers containing phosphorus acid, etc.is shown below:

<table>
<thead>
<tr>
<th>5 g analytical sample (powdery)</th>
<th>Weigh to the order of 1 mg into a 500-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>← Water, about 400 mL.</td>
<td></td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>Rotary shaker (30 - 40 revolutions/min) for 30 minutes</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1-1 Flow sheet for water-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Extraction procedure (4.1.1.1))

<table>
<thead>
<tr>
<th>2.5 g analytical sample (powdery)</th>
<th>Weigh to the order of 1 mg into a 250-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>← Water, about 200 mL.</td>
<td></td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>Vertical reciprocating shaker(300 times/min, with amplitude of 40 mm), for 30 minutes</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1-2 Flow sheet for water-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Extraction procedure (4.1.1.2))
1g analytical sample (fluid) Weigh to the order of 1 mg into a 1000-mL volumetric flask
→ Water, about 200 mL
Shaking to mix
→ Water (up to the marked line)
Filtration Type 3 filter paper
Sample solution

Figure 1-3 Flow sheet for water-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Extraction procedure (4.12))

Sample solution

Aliquot (predetermined volume) 100-mL - 200-mL tall beaker
→ 3 mL of hydrochloric acid
→ 1 mL of nitric acid
Heating Cover the tall beaker with a watch glass, heat on a hot plate or sand bath at 200 ºC - 250 ºC, condense until the solution volume reaches about 2 mL
Standing to cool
Transfer 100-mL volumetric flask, water
→ 1-2 drop(s) of Phenolphthalein solution (1 g/100 mL)
→ Ammonia solution (1+1) [neutralization]
→ Nitric acid (1+10) [slightly acidic]
→ 20 mL of coloring reagent solution
→ Water (up to the marked line)
Leaving at rest For about 30 minutes
Measurement Spectrophotometer (420 nm)

Figure 2 Flow sheet for water-soluble phosphoric acid in fertilizers containing phosphorus acid, etc. (Coloring and measurement procedure)
4.2.4.c Quinoline gravimetric analysis

(1) Summary
This testing method is applicable to fertilizers containing no phosphorous acid, etc. It is suitable for the fertilizers with relatively a high content of phosphoric acid. This testing method is classified as Type E and its symbol is 4.2.4.c-2017 or W-P.c-1.

Extract by adding water to an analytical sample. Heat after adding nitric acid and water, hydrolyze nonorthophosphoric acid to orthophosphate ion and measure the mass of quinonium phosphomolybdate formed by the reaction with quinoline, molybdic acid and nitric acid to obtain water-soluble phosphoric acid (W-P₂O₅) in an analytical sample.

(2) Reagent: Reagents are as shown below.
   a) Nitric acid: A JIS Guaranteed (HNO₃ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   b) Sodium molybdate solution: Dissolve 70 g of sodium molybdate dihydrate in 150 mL of water.
   c) Quinoline solution: Add 5 mL of quinoline specified in JIS K 8279 to the mixture solution of 35 mL of nitric acid and 100 mL of water.
   d) Quimosiac solution: Add 60 g of citric acid monohydrate specified in JIS K 8283 to the mixture solution of 85 mL nitric acid and 150 mL of water to dissolve. Add gradually total volume of the sodium molybdate solution to mix. Add gradually the total volume of the quinoline solution while mixing the solution. After leaving at rest overnight, filter the total volume with Type 3 filter paper. Add 280 mL of acetone specified in JIS K 8034, and further add water to make 1000 mL.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Water bath: Water bath that can be adjusted to 60 ºC - 65 ºC.
   c) Drying apparatus: A drying apparatus that can be adjusted to 220 ºC ± 5 ºC.
   d) Crucible type glass filter: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 220 ºC ± 5 ºC in advance and measure the mass to the order of 1 mg.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30 - 40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

   Comment 1 In the procedure in a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it into a 250-mL volumetric flask.
   Comment 2 The procedure in (4.1) is the same as the procedure in (4.1) of 4.2.4.a.

(4.2) Measurement: Conduct measurement as shown below.
   a) Transfer a predetermined volume (the equivalents of 10 mg - 30 mg as P₂O₅ and no more than 20 mL as the total solution volume) of sample solution to a 300-mL tall beaker.
   b) Add 5 mL of nitric acid and add water to make 80 mL.
   c) Cover with a watch glass. After boiling for about 3 minutes, wash the watch glass and the
inside the tall beaker with water and add water to make 100 mL.

d) Immediately, add 50mL of quimiosiac solution, heat for about 15 minutes while sometimes mixing in a water bath at 60 °C - 65 °C to produce the precipitate of quinolinium phosphomolybdate.

e) After standing to cool down to room temperature while sometimes mixing, filter under reduced pressure with a crucible type glass filter, wash the tall beaker 3 times with water and transfer the whole precipitate into a crucible type glass filter, and further wash 7 - 8 times with water.

f) Transfer the precipitate together with the crucible type glass filter into a drying apparatus and heat at 220 °C ± 5 °C for about 30 minutes.

g) As soon as heating is complete, move it into a desiccator and let it stand to cool.

h) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

i) Calculate water-soluble phosphoric acid (W-P₂O₅) by the following formula.

\[
\text{Soluble phosphoric acid (% (mass fraction)) in an analytical sample} = A \times 0.03207 \times \left(\frac{V_1}{V_2}\right) \times \left(\frac{1}{W}\right) \times 100
\]

\(A\): Mass (g) of the precipitate in h)  
\(W\): Mass of an analytical sample (5 g)  
\(V_1\): Predetermined volume (500 mL) of the sample solution  
\(V_2\): Volume (mL) of the sample solution transferred in a)

References


(5) Flow sheet for water-soluble phosphoric acid: The flow sheet for water-soluble phosphoric acid in fertilizers is shown below:

- Weigh to the order of 1 mg to a 500-mL volumetric flask
- About 400 mL of water
- Shaking to mix: Rotary shaker (30 - 40 revolutions/min) for 30 minutes
- Water (up to the marked line)
- Filtration: Type 3 filter paper
- Sample solution

Figure 1 Flow sheet for water-soluble phosphoric acid in fertilizers
(Extraction procedure)
300-mL tall beaker

 ← 5 mL nitrate acid
 ← Water (to make about 80 mL)

Cover with a watch glass for 3 minutes
Wash a watch glass and the inside the tall beaker with water

 ← Water (to make about 100 mL)
 ← 50 mL quimosiac solution

60 °C - 65 °C, for 15 minutes, some times mix

60 °C - 65 °C, for 15 minutes, some times mix

Room temperature

Crucible type glass filtering apparatus 1G4, 3 times with water

Wash 7 - 8 times with water

220 °C ± 5 °C for about 30 minutes

Desiccator

Measure the mass to the order of 1 mg

Figure 2  Flow sheet for water-soluble phosphoric acid in fertilizers
(Measurement procedure)
4.2.4.d ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. It is also suitable for the fertilizers containing phosphorus acid (phosphite). This testing method is classified as Type D and its symbol is 4.2.4.d-2017 or W-P.d-1.

Extract by adding water to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the phosphorus at a wavelength of 178.287 nm to obtain water-soluble phosphoric acid W-P₂O₅ in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Phosphoric acid standard solution (P₂O₅ 10 mg/mL) (1): Heat potassium dihydrogen phosphate specified in JIS K 9007 at 105 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 19.17 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, add 2 mL-3 mL of nitric acid, and add water up to the marked line.

d) Phosphoric acid standard solution (P₂O₅ 1 mg/mL) (1): Transfer 10 mL of phosphoric acid standard solution (P₂O₅ 10 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.

e) Phosphoric acid standard solution (P₂O₅ 20 µg/mL - 0.4 mg/mL) for the calibration curve preparation (1): Transfer 2 mL - 40 mL of phosphoric acid standard solution (P₂O₅ 1 mg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line.

f) Phosphoric acid standard solution ((P₂O₅ 5 µg/mL - 20 µg/mL) for the calibration curve preparation (1): Transfer 5 mL - 20 mL of phosphoric acid standard solution (P₂O₅ 0.1 mg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line.

g) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the phosphoric acid standard solution in (2), a phosphoric acid standard solution for a calibration curve can be prepared using a phosphoric acid standard solution (P 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble phosphoric acid (W-P₂O₅) in an analytical sample by multiplying the concentration (P) of a phosphoric acid standard solution for a calibration curve or a measured value obtained in (4.2) by a conversion factor (2.2914).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in d) and e) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a phosphoric acid standard solution for the calibration curve in the concentration...
range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample (2) to the order of 1 mg, and put it in a 100-mL volumetric flask.
   b) Add about 50 mL of water, shake to mix and add water up to the marked line.
   c) Filter with Type 3 filter paper to make a sample solution.

Note  (2) The sampling amount of the analytical sample is 10 g when there is less phosphoric acid content in the fertilizers such as a home garden-use fertilizer.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.
   a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
      Analytical line wavelength: 178.287 nm
   b) Calibration curve preparation
      1) Spray the phosphoric acid standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 178.287 nm.
      2) Prepare a curve for the relationship between the phosphoric acid concentration and the indicated value of the phosphoric acid standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation.
   c) Sample measurement
      1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 40 mg as \( P_2O_5 \)) to a 100-mL volumetric flask.
      2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
      3) Subject to the same procedure as in b) 1) to read the indicated value.
      4) Obtain the phosphoric acid (\( P_2O_5 \)) content from the calibration curve, and calculate water-soluble phosphoric acid (W-P\( _2O_5 \)) in the analytical sample.

Comment 4 The simultaneous measurement of multiple elements by an ICP-OES is available. In this case, transfer a pre-determined amount of phosphoric acid standard solution (P 1 mg/mL or 10 mg/mL), potassium standard solution (K 1 mg/mL or 10 mg/mL), magnesium standard solution (Mg 1 mg/mL or 10 mg/mL), manganese standard solution (Mn 1 mg/mL or 10 mg/mL), boron standard solution (B 1 mg/mL or 10 mg/mL), calcium standard solution (Ca 1 mg/mL), iron standard solution (Fe 1 mg/mL), cobalt standard solution (Co 1 mg/mL), copper standard solution (Cu 1 mg/mL), zinc standard solution (Zn 1 mg/mL) and molybdenum standard solution (Mo 1 mg/mL) traceable to National Metrology to a volumetric flask to mix, add hydrochloric acid (1+5) to make an acid concentration of 0.5 mol/L and further add water up to the marked line to prepare a primary mixed standard solution. Transfer a
pre-determined volume of primary mixed standard solution to volumetric flasks step-by-step, add hydrochloric acid (1+23) up to the marked line to prepare mixed standard solutions for calibration curve preparation within the concentration range in Table 1. In this case, multiply the respective concentrations of mixed standard solutions for calibration curve preparation or measurement values obtained in (4.2) by the conversion factors in Table 1 to calculate respective main components in an analytical sample. The measurement wavelengths of respective elements are according to Table 1. In addition, when preserving mixed standard solutions for calibration curve preparation, use a container, which can be sealed tightly, made of material such as PTFE that boron hardly elutes.

Table 1  Preparation concentration and measurement wavelength of mixed standard solution for calibration curve preparation

<table>
<thead>
<tr>
<th>Name of test item</th>
<th>Concentration of element (µg/mL)</th>
<th>Concentration of the equivalents to oxide(µg/mL)</th>
<th>Conversion factor</th>
<th>Measurement wavelength (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water-soluble phosphoric</td>
<td>P 1~200</td>
<td>P₂O₅</td>
<td>2.921~458.2</td>
<td>2.2914</td>
</tr>
<tr>
<td>Water-soluble potassium</td>
<td>K 1~200</td>
<td>K₂O</td>
<td>1.205~241.0</td>
<td>1.2046</td>
</tr>
<tr>
<td>Water-soluble magnesium</td>
<td>Mg 0.1~20</td>
<td>MgO</td>
<td>0.1658~33.16</td>
<td>1.6583</td>
</tr>
<tr>
<td>Water-soluble manganese</td>
<td>Mn 0.05~10</td>
<td>MnO</td>
<td>0.06455~12.91</td>
<td>1.2912</td>
</tr>
<tr>
<td>Water-soluble boron</td>
<td>B 0.05~10</td>
<td>B₂O₃</td>
<td>0.1610~32.20</td>
<td>3.2199</td>
</tr>
<tr>
<td>Water-soluble calcium</td>
<td>Ca 0.1 - 20</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water-soluble zinc</td>
<td>Zn 0.1 - 20</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water-soluble copper</td>
<td>Cu 0.1 - 20</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water-soluble iron</td>
<td>Fe 0.1 - 20</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water-soluble molybdenum</td>
<td>Mo 0.1 - 20</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Water-soluble cobalt</td>
<td>Co 0.1 - 20</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

1) Factor to convert an element to oxide

Comment 5 The comparison of the measurement value (yi: 0.179 % (mass fraction) - 10.88 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value (xi) of Ammonium vanadomolybdate absorptiometric analysis was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was \( y = -0.022 + 1.008x \), and its correlation coefficient (\( r \)) was 0.999. Additionally, additive recovery testing was conducted using a fluid mixed fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery rates at additive level of 10 % (mass fraction) and 1 % (mass fraction) were 98.1 % and 101.9 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 2 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).
Table 2 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $T_{1}^{(1)}$</th>
<th>Repeatability $s_{r}^{(2)}$</th>
<th>RSD $RSD_{r}^{(3)}$</th>
<th>Intermediate precision $s_{I(T)}^{(4)}$</th>
<th>RSD $RSD_{I(T)}^{(5)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>10.83</td>
<td>0.10</td>
<td>0.9</td>
<td>0.14</td>
<td>1.3</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (fluid)</td>
<td>7</td>
<td>0.829</td>
<td>0.008</td>
<td>0.9</td>
<td>0.015</td>
<td>1.8</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $T$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

(5) Flow sheet: The flow sheet for water-soluble phosphoric acid of the fluid mixed fertilizers is shown below:

![Flow sheet](image)

Figure 1 The flow sheet for water-soluble phosphoric acid in fluid fertilizers

(Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot (predetermined volume)</td>
<td>← 25 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
<td>ICP-OES (178.287nm)</td>
</tr>
</tbody>
</table>

Figure 2  The flow sheet for water-soluble phosphoric acid in fluid fertilizers (Measurement procedure)
4.3 Potassium
4.3.1 Total potassium
4.3.1.a Flame atomic absorption spectrometry or flame photometry

(1) Summary
This testing method is applicable to fertilizers containing organic matters. This testing method is classified as Type C and its symbol is 4.3.1.a-2017 or T-K.a-1.

Pretreat an analytical sample with incineration and hydrochloric acid to convert the total potassium into potassium ion, add an interference suppressor solution, and then spray in an acetylene-air flame, and measure the atomic absorption with potassium at a wavelength of 766.5 nm or 769.9 nm to quantify the total potassium. Or, determine the intensity of the emission line at a wavelength of 766.5 nm or 769.9 nm produced in flame to obtain the total potassium (T-K₂O) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Nitric acid: A JIS Guaranteed (HNO₃ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   c) Interference suppressor solution: Weigh 12.5 g of calcium carbonate specified in JIS K 8617 into a 2000-mL beaker, add a small amount of water, gradually add 105 mL of hydrochloric acid, and heat for a little while. After cooling is complete, add water to make 1000 mL.
   d) Potassium standard solution (K₂O 1 mg/mL) (1): Heat potassium chloride specified in JIS K 8121 at 110 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 1.583 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   e) Potassium standard solution (K₂O 5 µg/mL - 50 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of potassium standard solution (K₂O 1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (2), and add water up to the marked line.
   f) Blank test solution for the calibration curve preparation (1): Transfer about 50 mL of interference suppressor solution to a 500-mL volumetric flask (2), and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the potassium standard solution in (2), a potassium standard solution for the calibration curve preparation can be prepared by using a potassium standard solution (K 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate total potassium (T-K₂O) in the analytical sample by multiplying the concentration (K) of a potassium standard solution for calibration curve preparation or a measurement value (K) obtained in (4.2) by a conversion factor (1.2046).

(3) Instruments: Instruments are as shown below:
   a) Analytical instrument: An atomic absorption spectrometer or a flame photometer as shown below:
      aa) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in
JIS K 0121.

1) **Light source**: A potassium hollow cathode lamp
2) **Gas**: Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

**Flame photometer**:
1) **Gas**: Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

**Electric furnace**: An electric furnace that can be adjusted to 550 ºC ± 5 ºC.

**Hot plate or sand bath**: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 ºC.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Incineration-hydrochloric acid boiling**

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.
b) Put the tall beaker in an electric furnace, and heat gently to char\(^{(3)}\).
c) Ignite at 550 ºC ± 5 ºC for no less than 4 hours to incinerate\(^{(3)}\).
d) After standing to cool, moisten the residue with a small amount of water, gradually add about 10 mL of hydrochloric acid, and further add water to make about 20 mL.
e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to boil for about 5 minutes.
f) After cooling is complete, transfer to a 250-mL - 500-mL volumetric flask with water.
g) Add water up to the marked line.
h) Filter with Type 3 filter paper to make a sample solution.

**Note** (3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

**Comment 2** The procedure in (4.1.1) is the same as the procedure in (4.1.2) in 4.2.1.a.

(4.1.2) **Incineration-aqua regia digestion**

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.
b) Put the tall beaker in an electric furnace, and heat gently to char\(^{(4)}\).
c) Ignite at 450 ºC ± 5 ºC for 8 - 16 hours to incinerate\(^{(4)}\).
d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
f) Slightly move the watch glass\(^{(5)}\), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

**Comment** The procedure in (4.1.1) is the same as the procedure in (4.1.2) in 4.2.1.a.
Note  (4) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.
(5) The watch glass can be removed.
(6) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 3 The procedures in (4.1.2) are the same as in (4.1.3) in 4.2.1.a. and (4.1) a - h) in 5.3.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer or flame photometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer or flame photometer: Set up the measurement conditions for the atomic absorption spectrometer or flame photometer considering the following:
Analytical line wavelength: 766.5 nm or 769.9 nm

b) Calibration curve preparation
1) Spray the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 766.5 nm or 769.9 nm.
2) Prepare a curve for the relationship between the potassium concentration and the indicated value of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as K₂O) to a 100-mL volumetric flask.
2) Add about 10 mL of interference suppressor solution (2), and add water up to the marked line.
3) Subject to the same procedure as in b) 1) to read the indicated value.
4) Obtain the potassium content from the calibration curve, and calculate total potassium (T-K₂O) in the analytical sample.

Comment 4 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 97.8 % - 100.1 % and 100.9 % - 103.1 % as total potassium acid (T-K₂O) respectively. The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.
Additionally, the minimum limit of quantification of this testing method is about 0.08 % (mass fraction) for solid fertilizers, and 0.03 % (mass fraction) for fluid fertilizers.
Table 1  Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of certified reference laboratory</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
<th>Reproducibility</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Repeatability</td>
<td>Intermediate precision</td>
<td>Reproducibility</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Average</td>
<td>s_R</td>
<td>RSD_R</td>
</tr>
<tr>
<td>FAMIC-C-12</td>
<td>11</td>
<td>0.584</td>
<td>0.005</td>
<td>0.9</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis
2) Average (the number of laboratory(p) × test days (2) × the number of replicate testing(3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

(5) **Flow sheet for total potassium**: The flow sheet for total potassium in fertilizers is shown below:

<table>
<thead>
<tr>
<th>5 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>550 °C ± 5 °C, no less than 4 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td></td>
<td>← A small amount of water, moisten the residue</td>
</tr>
<tr>
<td></td>
<td>← About 10 mL of hydrochloric acid</td>
</tr>
<tr>
<td></td>
<td>← Water (up to about 20 mL)</td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and boil for 5 minutes.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask, water</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

![Flow sheet for total potassium in fertilizers](Incineration-hydrochloric acid boiling procedure (4.1.1))
Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker.

Charring
Incineration

Heat gently
Ignite at 450 °C ± 5 °C for 8 - 16 hours

Standing to cool

Room temperature
← A small amount of water
← About 10 mL of nitric acid
← About 30 mL of hydrochloric acid

Heat

Cover with a watch glass to digest

Heating

Slightly move the watch glass and remove acid

Standing to cool

Room temperature
← 25 mL - 50 mL of hydrochloric acid (1:5)

Heat

Cover with a watch glass to dissolve

Standing to cool

Room temperature

Transfer

100-mL - 200-mL volumetric flask, water
← Water (up to the marked line)

Filtration

Type 3 filter paper

Sample solution

Figure 1-2  Flow sheet for total potassium in fertilizers
( Incineration-aqua regia digestion procedure (4.1.2))

Sample solution

Aliquot

100-mL volumetric flask, water
← About 10 mL interference suppressor solution
← Water (up to the marked line)

Measurement

Atomic absorption spectrometer or flame photometer

Figure 2  Flow sheet for total potassium in fertilizers (Measurement procedure)
4.3.1.b Sodium tetraphenylborate gravimetric analysis

(1) Summary
This testing method is applicable to fertilizers containing organic matters. It is suitable for fertilizers containing a relatively high content of potassium. This testing method is classified as Type D and its symbol is 4.3.1.b-2017 or T-K.b-1.

Pretreat an analytical sample by incineration and hydrochloric acid to convert the total potassium into potassium ion, mask co-existing ammonium and other salts with formaldehyde and ethylenediamine tetraacetate, and measure the mass of potassium tetraphenylborate formed by the reaction with tetraphenylborate to obtain the total potassium (T-K2O) in an analytical sample. In addition, the performance of this testing method is shown in Comment 2.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
b) Formaldehyde solution: A JIS Guaranteed Reagent specified in JIS K 8872 or a reagent of equivalent quality.
c) Sodium hydroxide solution (200 g/L) (1): Dissolve 200 g of sodium hydroxide specified in JIS K 8576 in water to make 1000 mL.
d) Aluminum chloride solution (1 mol/L) (1): Dissolve 12 g of aluminum chloride (III) hexahydrate specified in JIS K 8114 in water to make 100 mL.
e) Tetraphenylborate solution (1): Transfer 6.1 g of sodium tetraphenylborate specified in JIS K 9521 to a 250-mL volumetric flask, dissolve by adding about 200 mL of water and add 10 mL of aluminum chloride solution. Add a methyl red solution (0.1 g/100 mL) as an indicator, and neutralize with a sodium hydroxide solution (200 g/L) until the color of the solution changes to yellow, and then add water up to the marked line. Filter with Type 3 filter paper and add 0.5 mL of sodium hydroxide solution (200 g/L) to the total filtrate. Filter with Type 3 filter paper in the case of usage.
f) Tetraphenylborate washing solution (1): Dilute 40 mL of tetraphenylborate solution with water to make 1000 mL.
g) Ethylenediaminetetraacetate - Sodium hydroxide solution (1): Dissolve 10 g of ethylenediaminetetraacetic acid disodium dihydrogen dihydrate specified in JIS K 8107 and 8 g of sodium hydroxide specified in JIS K 8576 in a proper amount of water. Add 6 mL - 10 mL of tetraphenylborate solution while mixing according to the potassium content coexisting as impurity after standing to cool, and then add water to make 100 mL. After leaving at rest for about 30 minutes while sometimes mixing, filter with Type 3 filter paper.
h) Methyl red solution (0.1 g/100 mL): Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) Electric furnace: An electric furnace that can be adjusted to 550 °C ± 5 °C.
b) Drying apparatus: A drying apparatus that can be adjusted to 120 °C ± 2 °C.
c) Crucible type glass filter: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 120 °C ± 2 °C in advance and measure the mass to the order of 1 mg.
d) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.
(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh about 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char (2).

c) Ignite at 550 °C ± 5 °C for no less than 4 hours to incinerate (2).

d) After standing to cool, moisten the residue with a small amount of water, gradually add about 10 mL of hydrochloric acid, and further add water to make 20 mL.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to boil for about 5 minutes.

f) After cooling, transfer to a 250-mL - 500-mL volumetric flask with water.

g) Add water up to the marked line.

h) Filter with Type 3 filter paper to make a sample solution.

Note (2) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

Comment 1 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.1.a. In addition, the sample solution prepared in (4.1) a) - h) in 4.9.1.a can also be used.

(4.2) Measurement: Conduct measurement as shown below.

a) Transfer a predetermined amount (the equivalents of 15 mg - 30 mg as K₂O) of the sample solution to a 100-mL tall beaker.

b) Add water to the solution to reach 50 mL when the procedure in e) is complete.

c) Add hydrochloric acid (1+9), so that the hydrochloric acid becomes equivalent to 0.2 mL.

d) Add 5 mL of formaldehyde solution, and then add 5 mL of ethylenediamine tetraacetate–sodium hydroxide solution.

e) Add necessary volume (3) of tetraphenylborate solution at the rate of one or two drop(s) per second while mixing, and further add 4 mL of the same solution in the same manner.

f) Leave at rest for about 30 minutes while sometimes mixing to form the precipitate of potassium tetraphenylborate.

g) Filter supernatant under reduced pressure with a crucible type glass filter, wash the tall beaker 5 times with 5 mL of tetraphenylborate washing solution and transfer the whole precipitate to the crucible type glass filter and further wash 2 times with 2 mL of water.

h) Transfer the precipitate together with the crucible type glass filter into a drying apparatus and heat at 120 °C ± 2 °C for about 1 hour.

i) As soon as heating is complete, move it into a desiccator and let it stand to cool.

j) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

k) Calculate total potassium (T-K₂O) in the analytical sample by the following formula.

\[
\text{Total potassium (T-K}_2\text{O) (% (mass fraction)) in an analytical sample} = A \times 0.1314 \times (V_1/V_2) / W \times 100
\]

\[A: \text{ Mass (g) of the precipitate}\]

\[V_1: \text{ Predetermined volume (mL) of the sample solution in (4.1) g}\]

\[V_2: \text{ Transferred amount (mL) of the sample solution in (4.2) a}\]

\[W: \text{ Mass (g) of the analytical sample}\]
Note  (3)  About 3 ml of tetraphenylborate solution per 10 mg of K₂O is required to form the precipitate of potassium tetraphenylborate

Comment 2  Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 25 % (mass fraction) - 30 % (mass fraction) and 10 % (mass fraction) - 20 % (mass fraction) are 99.5 % - 100.8 % and 99.5 % - 100.6 % as total potassium acid (T-K₂O) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.3 % (mass fraction).

References
(5) **Flow sheet for total potassium:** The flow sheet for total potassium in fertilizers is shown below:

<table>
<thead>
<tr>
<th>5 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>Ignite at 550 °C ± 5 °C, no less than 4 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← A small amount of water, moisten the residue</td>
<td></td>
</tr>
<tr>
<td>← About 10 mL of hydrochloric acid</td>
<td></td>
</tr>
<tr>
<td>← Water (up to about 20 mL)</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and boil for 5 minutes.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1   Flow sheet for total potassium in fertilizers (Extraction procedure)

Sample solution

100-mL tall beaker

← Water (add tetraphenylborate solution to reach 50 mL)
← Hydrochloric acid (1+9) (the equivalents of 0.2 mL of hydrochloric acid)
← 5 mL of formaldehyde solution
← 5 mL of ethylenediamine tetraacetate- sodium hydroxide solution
← Tetraphenylborate solution (necessary volume + 4 mL)

For 30 minutes, sometimes mixing

Crucible type glass filtering apparatus 1G4,
5 times with 5 mL of tetraphenylborate washing solution

Wash 2 times with 2 mL of water

120 °C ± 5 °C for 1 hour.

Desiccator

Measure the mass to the order of 1 mg

Figure 2  Flow sheet for total potassium in fertilizers (Measurement procedure)
4.3.2 Citrate soluble potassium

4.3.2.a Flame atomic absorption spectrometry or flame photometry

(1) Summary
This testing method is applicable to fertilizers containing potassium silicate fertilizers, etc. This testing method is classified as Type D and its symbol is 4.3.2.a-2018 or C-K.a-2.

Extract by adding a citric acid solution to an analytical sample, add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with potassium at a wavelength of 766.5 nm or 769.9 nm to quantify citrate soluble potassium (C-K$_2$O). Or, measure the intensity of the emission line at a wavelength of 766.5 nm or 769.9 nm produced in flame to quantify citrate soluble potassium (C-K$_2$O) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Citric acid solution: Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   c) Interference suppressor solution: Weigh 12.5 g of calcium carbonate specified in JIS K 8617 into a 2000-mL beaker, add a small amount of water, gradually add 105 mL of hydrochloric acid, and heat for a little while. After standing to cool, add water to make 1000 mL.
   d) Potassium standard solution (K$_2$O 1 mg/mL): Heat potassium chloride specified in JIS K 8121 at 110 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 1.583 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   e) Potassium standard solution (K$_2$O 5 µg/mL - 50 µg/mL) for the calibration curve preparation: Transfer 2.5 mL - 25 mL of potassium standard solution (K$_2$O 1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution, and add water up to the marked line.
   f) Blank test solution for the calibration curve preparation: Transfer about 50 mL of interference suppressor solution to a 500-mL volumetric flask, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the potassium standard solution in (2), a potassium standard solution for the calibration curve preparation can be prepared by using a potassium standard solution (K 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble potassium (C-K$_2$O) in the analytical sample by multiplying the concentration (K) of a potassium standard solution for calibration curve preparation or a measurement value (K) obtained in (4.2) by a conversion factor (1.2046).

(3) Instruments: Instruments are as shown below:
   a) Extractor: A constant-temperature rotary shaker or a reciprocating water bath shaker as described below.
   aa) Constant-temperature rotary shaker: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 ºC ± 1 ºC, upside down at
30 - 40 revolutions/min.

**ab) Reciprocating water bath shaker:** A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/minute with amplitude of 25 mm - 40 mm

**b) Analytical instrument:** An atomic absorption spectrometer or a flame photometer as shown below:

**ba) Flame atomic absorption spectrometer:** An atomic absorption spectrometer specified in JIS K 0121.
1) **Light source:** A potassium hollow cathode lamp
2) **Gas:** Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

**bb) Flame photometer:**
1) **Gas:** Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

b) Add 150 mL of citric acid solution heated up to about 30 °C (3), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (3) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 2** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (4).

b) Add 150 mL of citric acid solution heated up to about 30 °C (3), and shake to mix by reciprocating horizontally at 160 times/min with amplitude of 25 mm - 40 mm (30 °C ± 1 °C) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (4) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

**Comment 3** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

**Comment 4** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0121 and as shown below.
Specific measurement procedures are according to the operation method of the atomic absorption spectrometer or flame photometer used in measurement.

a) **Measurement conditions for the atomic absorption spectrometer or flame photometer:**
   Set up the measurement conditions for the atomic absorption spectrometer or flame photometer considering the following:
   - Analytical line wavelength: 766.5 nm or 769.9 nm

b) **Calibration curve preparation**
   1) Spray the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 766.5 nm or 769.9 nm.
   2) Prepare a curve for the relationship between the potassium concentration and the indicated value of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as K₂O) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (2), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the potassium content from the calibration curve, and calculate citrate soluble potassium (C-K₂O) in the analytical sample.

**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 100.2 % - 101.7 % and 100.4 % - 101.8 % as citrate soluble potassium (C-K₂O) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.05 % (mass fraction) for solid fertilizers, and 0.06 % (mass fraction) for fluid fertilizers.

**References**
(5) **Flow sheet for citrate soluble potassium:** The flow sheet for citrate soluble potassium in fertilizers is shown below:

![Flow sheet for citrate soluble potassium](image)

Figure 1-1 Flow sheet for citrate-soluble potassium in fertilizers (Extract procedure 4.1.1)

![Flow sheet for citrate soluble potassium](image)

Figure 1-2 Flow sheet for citrate-soluble potassium in fertilizers (Extract procedure 4.1.2)

![Flow sheet for citrate soluble potassium](image)

Figure 2 Flow sheet for citrate-soluble potassium in fertilizers (Measurement procedure)
4.3.2.b Sodium tetraphenylborate gravimetric analysis

(1) Summary
This testing method is applicable to fertilizers containing potassium silicate fertilizers, etc. This testing method is classified as Type D and its symbol is 4.3.2.b-2017 or C-K.b.1.

Extract by adding a citric acid solution to an analytical sample, mask co-existing ammonium and other salts with formaldehyde and ethylenediamine tetraacetate and measure the mass of citric acid soluble potassium (citrate-soluble potassium (C-K_{2}O)) and the mass of potassium tetraphenylborate formed by the reaction with tetraphenylborate to obtain citrate soluble potassium (C-K_{2}O) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent: Reagents are as shown below.
   a) Citric acid solution (1): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   b) Formaldehyde solution: A JIS Guaranteed Reagent specified in JIS K 8872 or a reagent of equivalent quality.
   c) Sodium hydroxide solution (200 g/L) (1): Dissolve 200 g of sodium hydroxide specified in JIS K 8576 in water to make 1000 mL.
   d) Aluminum chloride solution (1 mol/L) (1): Dissolve 12 g of aluminum chloride (III) hexahydrate specified in JIS K 8114 in water to make 100 mL.
   e) Tetraphenylborate solution (1): Transfer 6.1 g of sodium tetraphenylborate specified in JIS K 9521 to a 250-mL volumetric flask, dissolve by adding about 200 mL of water and add 10 mL of aluminum chloride solution. Add a methyl red solution (0.1 g/100 mL) as an indicator, and neutralize with a sodium hydroxide solution (200 g/L) until the color of the solution changes to yellow, and then add water up to the marked line. Filter with Type 3 filter paper and add 0.5 mL of sodium hydroxide solution (200 g/L) to the total filtrate. Filter with Type 3 filter paper in the case of usage.
   f) Tetraphenylborate washing solution (1): Dilute 40 mL of tetraphenylborate solution with water to make 1000 mL.
   g) Ethylenediaminetetraacetate - Sodium hydroxide solution (1): Dissolve 10 g of ethylenediaminetetraacetic acid disodium dihydrogen dihydrate specified in JIS K 8107 and 8 g of sodium hydroxide specified in JIS K 8576 in a proper amount of water. Add 6 mL - 10 mL of tetraphenylborate solution while mixing according to the potassium content coexisting as impurity after standing to cool, and then add water to make 100 mL. After leaving at rest for about 30 minutes while sometimes mixing, filter with Type 3 filter paper.
   h) Methyl red solution (0.1 g/100 mL): Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Constant-temperature rotary shaker: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.
   b) Drying apparatus: A drying apparatus that can be adjusted to 120 °C ± 2 °C.
   c) Crucible type glass filter: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 120 °C ± 2 °C in advance and measure the mass to the order of 1 mg.

(4) Test procedure
   (4.1) Extraction: Conduct extraction as shown below.
Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

Add 150 mL of citric acid solution heated up to about 30 ºC (2), and shake to mix at 30 - 40 revolutions/min (30 ºC ± 1 ºC) for 1 hour.

After immediate cooling is complete, add water up to the marked line

Filter with Type 3 filter paper to make a sample solution.

**Note** (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 1** The procedures in (4.1) are the same as in (4.1.1) in 4.2.3.a.

**Comment 2** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedure of (4.1) b).

**Measurement:** Conduct measurement as shown below.

a) Transfer 20 mL of sample solution to a 100-mL tall beaker.

b) Add water to the solution to reach 50 mL when the procedure in d) is complete.

c) Add 5 mL of formaldehyde solution, and then add 5 mL of ethylenediamine tetraacetate - sodium hydroxide solution.

d) Add necessary volume (2) of tetraphenylborate solution at the rate of one or two drop(s) per second while mixing, and further add 4 mL of the same solution in the same manner.

e) Leave at rest for about 30 minutes while sometimes mixing to form the precipitate of potassium tetraphenylborate.

f) Filter supernatant under reduced pressure with a crucible type glass filter, wash the vessel 5 times with 5 mL of tetraphenylborate washing solution and transfer the whole precipitate to the crucible type glass filter and further wash 2 times with 2 ml of water.

g) Transfer the precipitate together with the crucible type glass filter into a drying apparatus and heat at 120 ºC ± 2 ºC for about 1 hour.

h) As soon as heating is complete, move it into a desiccator and let it stand to cool.

i) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

j) Calculate citrate soluble potassium (C-K_2O) by the following formula in the analytical sample.

\[
\text{Citrate soluble potassium (C-K}_2\text{O)} \ (% \text{ (mass fraction)) in an analytical sample} = \frac{A \times 0.1314 \times (V_1/V_2) \times W}{100} \\
\]

\[ A: \quad \text{Mass (g) of the precipitate} \\
V_1: \quad \text{Predetermined volume (mL) of the sample solution in (4.1) c)} \\
V_2: \quad \text{Transferred amount (mL) of the sample solution in (4.2) a)} \\
W: \quad \text{Mass (g) of the analytical sample} \]

**Note** (2) About 3 ml of tetraphenylborate solution per 10 mg of K_2O is required to form the precipitate of potassium tetraphenylborate

**Comment 3** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 25 % (mass fraction) - 30 % (mass fraction) and 10 % (mass fraction) - 20 % (mass fraction) are 98.6 % - 100.6 % and 100.6 % - 100.7 % as citrate soluble potassium (C-K_2O) respectively.
Additionally, the minimum limit of quantification of this testing method is about 0.6 % (mass fraction).

References

(5) Flow sheet for the testing method: The flow sheet for citrate soluble potassium in fertilizers is shown below:

1 g analytical sample

Weigh to the order of 1 mg into a 250-mL volumetric flask.

\[ \rightarrow 150 \text{mL citrate solution [about 30 °C]} \]

Shaking to mix

Constant temperature rotary shaker (30 - 40 revolutions/min) 
30 °C ± 1 °C, 1 hour

Cooling

Immediately

\[ \rightarrow \text{Water (up to the marked line)} \]

Filtration

Type 3 filter paper

Sample solution

Figure 1 Flow sheet for citrate-soluble potassium in fertilizers (Extraction procedure)
<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample solution</td>
<td>100-mL tall beaker</td>
</tr>
<tr>
<td>Aliquot (20 mL)</td>
<td>Water (add tetraphenylborate solution to reach 50 mL)</td>
</tr>
<tr>
<td></td>
<td>← 5 mL of formaldehyde solution</td>
</tr>
<tr>
<td></td>
<td>← 5 mL of ethylenediamine tetraacetate - sodium hydroxide solution</td>
</tr>
<tr>
<td></td>
<td>← Tetraphenylborate solution (the equivalents of potassium + 4 mL)</td>
</tr>
<tr>
<td>Forming precipitate</td>
<td>For 30 minutes, sometimes mixing</td>
</tr>
<tr>
<td>Filtration under reduced pressure</td>
<td>Crucible type glass filtering apparatus 1G4, 5 times with 5 mL of tetraphenylborate washing solution</td>
</tr>
<tr>
<td>Washing</td>
<td>Wash 2 times with 2 mL of water</td>
</tr>
<tr>
<td>Drying</td>
<td>120 °C ± 5 °C for 1 hour.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Desiccator</td>
</tr>
<tr>
<td>Measurement</td>
<td>Measure the mass to the order of 1 mg</td>
</tr>
</tbody>
</table>

Figure 2 Flow sheet for citrate-soluble potassium in fertilizers (Measurement procedure)
4.3.2.c Sodium tetraphenylborate volumetric analysis

(1) Summary
This testing method is applicable to fertilizers containing potassium silicate fertilizer, etc. but not organic matters. This testing method is classified as Type E and its symbol is 4.3.2.c-2017 or C-K.c-1.

Extract by adding a citric acid solution to an analytical sample, mask co-existing ammonium and other salts with formaldehyde, and make potassium ion and tetraphenylborate react with each other. Measure unconsumed tetraphenylborate by conducting a precipitate titration to obtain citrate-soluble potassium (C-K$_2$O) in an analytical sample.

(2) Reagent: Reagents are as shown below.
   a) Citric acid solution \(^{(1)}\): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   b) Formaldehyde solution: A JIS Guaranteed Reagent specified in JIS K 8872 or a reagent of equivalent quality.
   c) Sodium hydroxide solution (120 g/L) \(^{(1)}\): Dissolve 30 g of sodium hydroxide specified in JIS K 8576 in water to make 250 mL.
   d) Tetraphenylborate solution \(^{(1)}\): Transfer 12.2 g of sodium tetraphenylborate to a 1000-mL volumetric flask, dissolve by adding about 800 mL of water and add 3 mL of sodium hydroxide (120 g/L) to the total filtrate, and further add water up to the marked line. Filter with Type 3 filter paper in the case of usage.
   e) Benzalkonium chloride solution (3.3 g/500 mL) \(^{(1)}\): Dissolve 3.3 g of benzalkonium chloride in 500 mL of water.
   f) Methyl red solution (0.1 g/100 mL): Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.
   g) Titan Yellow solution (0.04 g/100 mL): Dissolve 0.04 g of Titan Yellow in 100 mL of water in the case of usage.
   h) Potassium standard solution (K$_2$O 2 mg/mL) \(^{(1)}\): Heat potassium chloride specified in JIS K 8121 at 110 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 3.166 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Instruments: Instruments are as shown below:
   a) Constant-temperature rotary shaker: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.
   b) Add 150 mL of citric acid solution heated up to about 30 °C \(^{(2)}\), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.
   c) After immediate cooling is complete, add water up to the marked line
   d) Filter with Type 3 filter paper to make a sample solution.

Note (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.
Comment 1 The procedures in (4.1) are the same as in (4.1.1) in 4.2.3.a.
Comment 2 The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedure of (4.1) b).

(4.2) Precipitate formation: Form precipitate as shown below.
   a) Transfer 5 mL - 15 mL (no more than the equivalents of 30 mg as K$_2$O) of the extract to a 100-mL volumetric flask.
   b) Add water to the solution to make about 30 mL.
   c) Add about 5 mL of formaldehyde solution and add 5 mL of sodium hydroxide solution (120 g/L).
   d) Add 25 mL of tetraphenylborate solution at the rate of one or two drop(s) per second while shaking to mix.
   e) After adding water up to the marked line, leave at rest for 10 minutes.
   f) Filter with Type 3 filter paper to make a sample solution.

(4.3) Measurement: Conduct measurement as shown below.
   a) Calibration curve preparation
      1) Transfer 1 mL - 15 mL of potassium standard solution (K$_2$O 2 mg/mL) to 100-mL volumetric flasks step-by-step.
      2) Conduct the same procedures as (4.2) b) - f) to make K$_2$O 2 mg/100 mL - 30 mg/100 mL of the potassium standard solutions for the calibration curve preparation.
      3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.
      4) Transfer 40 mL of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation to an Erlenmeyer flask respectively.
      5) Add a few drops of Titan Yellow solution.
      6) Titrate with a benzalkonium chloride solution (3.3 g/500 mL) until the color of the solution changes to light red $^{(2)}$.
      7) Prepare a curve for the relationship between the potassium concentration and the volume of the benzalkonium chloride solution (3.3 g/500 mL) required for the titration of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
   b) Sample measurement
      1) Transfer 40 mL of the sample solution of (4.2) f) to a 100-mL Erlenmeyer flask.
      2) Conduct similarly as in a) 5) - 6) to obtain the volume of the benzalkonium chloride solution (3.3 g/500 mL) required for the titration.
      3) Obtain the potassium content from the calibration curve, and calculate citrate soluble potassium (C-K$_2$O) in the analytical sample.

Note (3) If the solution temperature is no more than 20 °C, the reaction does not advance in some cases. Therefore, it is recommended to heat the solution up to about 30 °C.

References
(5) **Flow sheet for citrate soluble potassium:** The flow sheet for citrate soluble potassium in fertilizers is shown below:

```
1 g analytical sample  Weigh to the order of 1 mg into a 250-mL volumetric flask.
< 150 mL citrate solution [about 30 ºC]
Shaking to mix  Constant temperature rotary shaker (30 - 40 revolutions/min)
30 ºC ± 1 ºC, 1 hour
Standing to cool
< Water (up to the marked line)
Filtration  Type 3 filter paper
Cooling  Immediately
< Water (up to the marked line)
Filtration  Type 3 filter paper
Sample solution
```

Figure 1  Flow sheet for citrate-soluble potassium in fertilizers (Extraction procedure)

```
Sample solution

Aliquot  (5 mL - 15 mL)  100-mL volumetric flask
< Water (the volume to reach about 30 mL)
< About 5 mL of formaldehyde solution
< 5 mL of sodium hydroxide solution (120 g/L)
< 25 mL tetraphenylborate solution
(one or two drop(s) per second while shaking to mix)
< Water (up to the marked line)
Leaving at rest  10 minutes
Filtration  Type 3 filter paper
< A few drops of Titan Yellow solution
Aliquot  (40 mL)  100-mL Erlenmeyer flask
< A few drops of titan yellow solution
Titration  Benzaikonium chloride solution (3.3 g/500 mL)
(unti the solution changes to light red)
```

Figure 2  Flow sheet for citrate-soluble potassium in fertilizers (Measurement procedure)
4.3.2.d  ICP Optical Emission Spectrometry

(1)  Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 4.3.2.d-2018 or C-K.b-1.

Extract by adding a citric acid solution to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the potassium at a wavelength of 766.491 nm to obtain citric acid-soluble potassium (citrate-soluble potassium (C-K$_2$O)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2)  Reagents: Reagents are as shown below.
  b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
  c) Citric acid solution: Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
  d) Potassium standard solution (K$_2$O 1 mg/mL): Heat potassium chloride specified in JIS K 8121 at 110 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 1.583 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
  e) Potassium standard solution (K$_2$O 20 µg/mL - 0.16 mg/mL) for the calibration curve preparation: Transfer 2 mL - 16 mL of potassium standard solution (K$_2$O 1 mg/mL) to 100-mL volumetric flasks step-by-step, add about 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
  f) Potassium standard solution (K$_2$O 2 µg/mL - 20 µg/mL) for the calibration curve preparation: Transfer 2 mL - 20 mL of potassium standard solution (K$_2$O 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, add about 25 mL of hydrochloric acid (1+23), and add water up to the marked line.
  g) Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in e) and f).

Note  (1)  This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the potassium standard solution in (2), a potassium standard solution for the calibration curve preparation can be prepared by using a potassium standard solution (K 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble potassium (C-K$_2$O) in the analytical sample by multiplying the concentration (K) of a potassium standard solution for calibration curve preparation or a measurement value (K) obtained in (4.2) by a conversion factor (1.2046).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The axial observation mode does not apply to potassium since interference is serious.

(3)  Instruments: Instruments are as shown below:
  a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
     1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity
  b) Extractor: A constant-temperature rotary shaker or a reciprocating water bath shaker as described below.
  ba) Constant-temperature rotary shaker: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30
- 40 revolutions/min.

**bb) Reciprocating water bath shaker:** A reciprocating water bath shaker that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/minute with amplitude of 25 mm - 40 mm.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker:**

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

b) Add 150 mL of citric acid solution heated up to about 30 ºC (2), and shake to mix at 30 - 40 revolutions/min (30 ºC ± 1 ºC) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 3** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker:**

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (3).

b) Add 150 mL of citric acid solution heated up to about 30 ºC (2), and shake to mix by reciprocating horizontally at 160 times/min with amplitude of 25 mm - 40 mm (30 ºC ± 1 ºC) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (3) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

**Comment 3** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

**Comment 4** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b and (4.1.2) b.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer:** Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

Analytical line wavelength: 766.491 nm

b) **Calibration curve preparation**

1) Spray the potassium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 766.491 nm.

2) Prepare a curve for the relationship between the potassium concentration and the indicated value of the potassium standard solutions for the calibration curve preparation and the
blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.2 mg - 16 mg as K\textsubscript{2}O) to a 100-mL volumetric flask.

2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Obtain the potassium content from the calibration curve, and calculate citrate soluble potassium (C-K\textsubscript{2}O) in the analytical sample.

**Comment 5** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.3.d **Comment 7**.

**Comment 6** The comparison of the measurement value (y: 3.57 % (mass fraction) - 34.24 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value (x) of Flame atomic absorption spectrometry analysis was conducted to evaluate trueness using compound fertilizers (9 samples), mixed compost fertilizers (1 sample), designated blended fertilizers (1 sample), blended fertilizers (4 samples), and byproduct mixed fertilizers (1 sample). As a result, a regression equation was $y = -0.0058 + 1.0027x$, and its correlation coefficient ($r$) was 0.999. In addition, recovery testing was conducted using a preparation sample. As a result, the average rate of recovery at the content level of 0.329 % (mass fraction) - 63.18 % (mass fraction) was 98.0 % - 100.3 %.

The results of the repeatability tests on different days using compound fertilizers and blended fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.09 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average\textsuperscript{2\footnotesize{j}}</td>
<td>$s_r$\textsuperscript{3\footnotesize{j}}</td>
<td>$RSD_r$\textsuperscript{3\footnotesize{j}}</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>7 \textsuperscript{1\footnotesize{i}}</td>
<td>16.17</td>
<td>0.13</td>
</tr>
<tr>
<td>Blended fertilizer</td>
<td>7 \textsuperscript{1\footnotesize{i}}</td>
<td>4.42</td>
<td>0.04</td>
</tr>
</tbody>
</table>

**Table 1 Analysis results of the repeatability tests on different days**

1) The number of test days conducting a duplicate test

2) Average (the number of test days ($T$) × the number of duplicate testing (2))

3) Mass fraction

4) Repeatability standard deviation

5) Repeatability relative standard deviation

6) Intermediate standard deviation

7) Intermediate relative standard deviation
(5) **Flow sheet for citrate soluble potassium**: The flow sheet for citrate soluble potassium in fertilizers is shown below:

![Flow sheet for citrate soluble potassium](image)

Figure 1-1  Flow sheet for citrate-soluble potassium in fertilizers (Extract procedure 4.1.1)

![Flow sheet for citrate soluble potassium](image)

Figure 1-2  Flow sheet for citrate-soluble potassium in fertilizers (Extract procedure 4.1.2)

![Flow sheet for citrate soluble potassium](image)

Figure 2  Flow sheet for citrate-soluble potassium in fertilizers (Measurement procedure)
4.3.3 Water-soluble potassium
4.3.3.a Flame atomic absorption spectrometry or flame photometry

(1) Summary
This testing method is applicable to fertilizers containing potassium salts. This testing method is classified as Type C and its symbol is 4.3.3.a-2017 or W-K.a-1.

Extract by adding water to an analytical sample, add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with potassium at a wavelength of 766.5 nm or 769.9 nm to quantify water-soluble potassium (W-K₂O). Or, determine the intensity of the emission line at a wavelength of 766.5 nm or 769.9 nm produced in flame to quantify water-soluble potassium (W-K₂O) in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Interference suppressor solution: Weigh 12.5 g of calcium carbonate specified in JIS K 8617 into a 2000-mL beaker, add a small amount of water, gradually add 105 mL of hydrochloric acid, and heat for a little while. After cooling is complete, add water to make 1000 mL.

c) Potassium standard solution (K₂O 1 mg/mL) (1): Heat potassium chloride specified in JIS K 8121 at 110 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 1.583 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.

d) Potassium standard solution (K₂O 5 µg/mL - 50 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of potassium standard solution (K₂O 1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (2), and add water up to the marked line.

e) Blank test solution for the calibration curve preparation (1): Transfer about 50 mL of interference suppressor solution to a 500-mL volumetric flask (2), and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the potassium standard solution in (2), a potassium standard solution for the calibration curve preparation can be prepared by using a potassium standard solution (K 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble potassium (W-K₂O) in the analytical sample by multiplying the concentration (K) of a potassium standard solution for calibration curve preparation or a measurement value (K) obtained in (4.2) by a conversion factor (1.2046).

(3) Instruments: Instruments are as shown below:

a) Extractor: A constant-temperature rotary shaker or a vertical reciprocating shaker as described below.

aa) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.

ab) Vertical reciprocating shaker: A vertical reciprocating shaker that can vibrate a 250-mL volumetric flask using an adapter to reciprocate vertically at 300 times/ minute (amplitude of
40 mm).  

b) **Analytical instrument**: An atomic absorption spectrometer or a flame photometer as shown below:

ba) **Flame atomic absorption spectrometer**: An atomic absorption spectrometer specified in JIS K 0121.

1) **Light source**: A potassium hollow cathode lamp  
2) **Gas**: Gas for heating by flame  
   (i) Fuel gas: acetylene  
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

bb) **Flame photometer**:  
1) **Gas**: Gas for heating by flame  
   (i) Fuel gas: acetylene  
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

c) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.

(4) Test procedure  
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) **Compound fertilizers containing potassium salts and magnesium potassium sulfate**  

a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 300-mL tall beaker.  
b) Add about 200 mL of water, and cover with a watch glass and heat on a hot plate to boil for about 15 minutes.  
c) Immediately transfer it with water to a 250-mL volumetric flask.  
d) After immediate cooling is complete, add water up to the marked line  
e) Filter with Type 3 filter paper to make a sample solution.

**Comment 2** In the procedure in a), a 250-mL volumetric flask can be used instead of a 300-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. Additionally, “cover with a watch glass” in b) is replaced by “place a funnel”. Skip “transfer to a 250-mL volumetric flask with water” in the procedure in c).

**Comment 3** The procedure in (4.1.1) is the same as the procedures in (4.1.1) of 4.3.3.b, (4.1.1) of 4.3.3.c and (4.1) of 4.8.2.a.

(4.1.2) **Compound fertilizers containing no magnesium potassium sulfate**  

(4.1.2.1) **Rotary shaker**:  
a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.  
b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.  
c) Add water up to the marked line.  
d) Filter with Type 3 filter paper to make a sample solution.

**Comment 4** In the procedure of (4.1.2.1) a), it is also allowed to weigh 2.5g of an analytical sample to the order of 1 mg, and put it in a 250-mL volumetric flask.

**Comment 5** The procedure in (4.1.2.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.1.2.2) **Vertical reciprocating shaker**:  
a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.  
b) Add about 200 mL of water, and shake to mix by reciprocating vertically at 300 times/min (amplitude of 40 mm) for about 30 minutes.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 6 The procedure in (4.1.2.2) is the same as the procedure in (4.1.1.2) in 4.2.4.a.

(4.1.3) Fluid test sample
a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 7 The procedure in (4.1.3) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer or flame photometer used in measurement.
a) Measurement conditions for the atomic absorption spectrometer or flame photometer:
   Set up the measurement conditions for the atomic absorption spectrometer or flame photometer considering the following:
   Analytical line wavelength: 766.5 nm or 769.9 nm
b) Calibration curve preparation
   1) Spray the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 766.5 nm or 769.9 nm.
   2) Prepare a curve for the relationship between the potassium concentration and the indicated value of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as K₂O) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (²), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the potassium content from the calibration curve, and calculate water soluble potassium (W-K₂O) in the analytical sample.

Comment 8 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) - 20 % (mass fraction) and 1 % (mass fraction) - 5 % (mass fraction) are 97.9 % - 100.2 % and 97.3 % - 100.6 % as water soluble potassium (W-K₂O) respectively. The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

The comparison of the measurement value of extraction (y₁: 2.69 % (mass fraction) - 26.64 % (mass fraction) ) with a vertical reciprocating shaker and the measurement value of extract with a rotary shaker (x₁) was conducted to evaluate trueness of the extraction of solid fertilizers using fertilizers (12 samples). As a result, a regression equation was \( y = 0.022 + 1.001x \), and its correlation coefficient (r) was 1.000. Also, the results of the repeatability tests on different days using compound fertilizers and
designated blended fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 2 shows the calculation results of intermediate precision and repeatability.

The comparison of the measurement value of extraction ($y_i$: 2.69 % (mass fraction) - 26.64 % (mass fraction)) with a vertical reciprocating shaker and the measurement value of extract with a rotary shaker ($x_i$) was conducted to evaluate trueness of the extraction of fluid fertilizers using fertilizers (12 samples). As a result, a regression equation was $y = 0.022 + 1.001x$, and its correlation coefficient ($r$) was 1.000. Also, the results of the repeatability tests on different days using fluid mixed fertilizers to evaluate the precision of the extraction of fluid fertilizers were analyzed by the one-way analysis of variance. Table 3 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.04 % (mass fraction) for solid fertilizers, and 0.007 % (mass fraction) for fluid fertilizers.

### Table 1 Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratories used for analysis</th>
<th>Average of the number of laboratory (p) × test days (T) × the number of replicate testing (3)</th>
<th>Mass fraction</th>
<th>Repeatability standard deviation</th>
<th>Intermediate relative standard deviation</th>
<th>Reproducibility standard deviation</th>
<th>Reproducibility relative standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-A-10</td>
<td>11</td>
<td>13.59</td>
<td>0.09</td>
<td>0.6</td>
<td>0.16</td>
<td>1.2</td>
<td></td>
</tr>
<tr>
<td>FAMIC-A-13</td>
<td>10</td>
<td>13.07</td>
<td>0.08</td>
<td>0.6</td>
<td>0.11</td>
<td>0.8</td>
<td>0.16</td>
</tr>
<tr>
<td>FAMIC-B-10</td>
<td>9</td>
<td>8.85</td>
<td>0.04</td>
<td>0.4</td>
<td>0.07</td>
<td>0.7</td>
<td>0.12</td>
</tr>
<tr>
<td>FAMIC-B-14</td>
<td>14</td>
<td>8.32</td>
<td>0.03</td>
<td>0.4</td>
<td>0.07</td>
<td>0.8</td>
<td>0.13</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting frame atomic absorption spectrometry
2) Average (the number of laboratory (p) × test days (T) × the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation

### Table 2 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average of the number of test days conducting a duplicate test (T) × the number of duplicate testing (2)</th>
<th>Mass fraction</th>
<th>Repeatability standard deviation</th>
<th>Intermediate relative standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>19.67</td>
<td>0.5</td>
<td>0.15</td>
<td>0.7</td>
</tr>
<tr>
<td>Designated blended fertilizer</td>
<td>7</td>
<td>6.50</td>
<td>1.1</td>
<td>0.07</td>
<td>1.1</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) × the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
Table 3  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $t$</th>
<th>Repeatability $s_t$</th>
<th>Intermediate precision $s_{I(T)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer 1</td>
<td>7</td>
<td>9.66</td>
<td>0.02</td>
<td>0.07</td>
</tr>
<tr>
<td>Fluid mixed fertilizer 2</td>
<td>7</td>
<td>2.44</td>
<td>0.01</td>
<td>0.02</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $T$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

(5) Flow sheet for water soluble potassium: The flow sheet for water soluble potassium in fertilizers is shown below:

```
2.5 g analytical sample (potassium salts, etc.)
   ← About 200 mL water
   Heating
   Cooling
   → Water, 250-mL volumetric flask
   → Water (up to the marked line)
   Type 3 filter paper
   Filtration
   Sample solution
```

Figure 1-1  Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.1))
About 400 mL water
Rotary shaker (30 - 40 revolutions/min)
For 30 minutes
Water (up to the marked line)
Type 3 filter paper
Sample solution

Figure 1-2 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.2.1))

Weigh to the order of 1 mg into a 250-mL volumetric flask
About 200 mL water
Vertical reciprocating shaker (300 times/min, with amplitude of 40 mm), for 30 minutes
Water (up to the marked line)
Type 3 filter paper
Sample solution

Figure 1-3 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.2.2))

Weigh to the order of 1 mg into a 100-mL volumetric flask
About 50 mL water
Water (up to the marked line)
Type 3 filter paper
Sample solution

Figure 1-4 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.3))
Figure 2  Flow sheet for water-soluble potassium in fertilizers (Measurement procedure)
4.3.3.b Sodium tetraphenylborate gravimetric analysis

(1) Summary
This testing method is applicable to fertilizers containing potassium salts. This testing method is classified as Type D and its symbol is 4.3.3.b-2017 or W-K.b-1. Extract by adding water to an analytical sample, mask co-existing ammonium and other salts with formaldehyde and ethylenediamine tetracacetate and measure the mass of potassium tetraphenylborate formed by the reaction with tetraphenylborate to obtain water-soluble potassium (W-K₂O) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Formaldehyde solution: A JIS Guaranteed Reagent specified in JIS K 8872 or a reagent of equivalent quality.
   c) Sodium hydroxide solution (200 g/L) (1): Dissolve 200 g of sodium hydroxide specified in JIS K 8576 in water to make 1000 mL.
   d) Aluminum chloride solution (1 mol/L) (1): Dissolve 12 g of aluminum chloride (III) hexahydrate specified in JIS K 8114 in water to make 100 mL.
   e) Tetraphenylborate solution (1): Transfer 6.1 g of sodium tetraphenylborate specified in JIS K 9521 to a 250-mL volumetric flask, dissolve by adding about 200 mL of water and add 10 mL of aluminum chloride solution. Add a methyl red solution (0.1 g/100 mL) as an indicator, and neutralize with a sodium hydroxide solution (200 g/L) until the color of the solution changes to yellow, and then add water up to the marked line. Filter with Type 3 filter paper and add 0.5 mL of sodium hydroxide solution (200 g/L) to the total filtrate. Filter with Type 3 filter paper in the case of usage.
   f) Tetraphenylborate washing solution (1): Dilute 40 mL of tetraphenylborate solution with water to make 1000 mL.
   g) Ethylenediaminetetraacetate - Sodium hydroxide solution (1): Dissolve 10 g of ethylenediaminetetraacetic acid disodium dihydrogen dihydrate specified in JIS K 8107 and 8 g of sodium hydroxide specified in JIS K 8576 in a proper amount of water. Add 6 mL - 10 mL of tetraphenylborate solution while mixing according to the potassium content coexisting as impurity after standing to cool, and then add water to make 100 mL. After leaving at rest for about 30 minutes while sometimes mixing, filter with Type 3 filter paper.
   h) Methyl red solution (0.1 g/100 mL): Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Drying apparatus: A drying apparatus that can be adjusted to 120 °C ± 2 °C.
   c) Crucible type glass filter: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 120 °C ± 2 °C in advance and measure the mass to the order of 1 mg.
   d) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 °C.

(4) Test procedure
   (4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Compound fertilizers containing potassium salts and magnesium potassium sulfate

a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 300-mL tall beaker.
b) Add about 200 mL of water, and cover with a watch glass and heat on a hot plate to boil for about 15 minutes.
c) Immediately transfer it with water to a 250-mL volumetric flask.
d) After immediate cooling is complete, add water up to the marked line.
e) Filter with Type 3 filter paper to make a sample solution.

Comment 1 In the procedure in a), a 250-mL volumetric flask can be used instead of a 300-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. Additionally, “cover with a watch glass” in b) is replaced by “place a funnel”. Skip “transfer to a 250-mL volumetric flask with water” in the procedure in c).

Comment 2 The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.3.3.a.

(4.1.2) Compound fertilizers containing no magnesium potassium sulfate

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.
b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 3 In the procedure in a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it into a 250-mL volumetric flask.

Comment 4 The procedure in (4.1.2) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.2) Measurement: Conduct measurement as shown below.
a) Transfer a predetermined amount (the equivalents of 15 mg - 30 mg as K₂O) of the sample solution to a 100-mL tall beaker.
b) Add water to the solution to reach 50 mL when the procedure in e) is complete.
c) Add 2 mL of hydrochloric acid (1+9).
d) Add 5 mL of formaldehyde solution, and then add 5 mL of ethylenediamine tetraacetate - sodium hydroxide solution.
e) Add necessary volume (2) of tetraphenylborate solution at the rate of one or two drop(s) per second while mixing, and further add 4 mL of the same solution in the same manner.
f) Leave at rest for about 30 minutes while sometimes mixing to form the precipitate of potassium tetraphenylborate.
g) Filter supernatant under reduced pressure with a crucible type glass filter, wash the vessel 5 times with 5 mL of tetraphenylborate washing solution and transfer the whole precipitate to the crucible type glass filter and further wash 2 times with 2 ml of water.
h) Transfer the precipitate together with the crucible type glass filter into a drying apparatus adjusted to 120 °C ± 2 °C and heat for 1 hour.
i) As soon as heating is complete, move it into a desiccator and let it stand to cool.
j) After standing to cool, remove the ground-in stoppered weighing bottle from the desiccator, and measure the mass to the order of 1 mg.
k) Calculate water soluble potassium (W-K₂O) in the analytical sample by the following formula.

\[
\text{Water-soluble potassium (W-K}_2\text{O)} \:\% (\text{mass fraction}) = A \times 0.1314 \times \left(\frac{V_1}{V_2}\right) W \times 100
\]
A: Mass (g) of the precipitate

$V_1$: Predetermined volume (mL) of the sample solution in (4.1.1) d) or (4.1.2) e)

$V_2$: Transferred amount (mL) of the sample solution in (4.2) a)

$W$: Mass (g) of the analytical sample

**Note** (2) About 3 ml of tetraphenylborate solution per 10 mg of K$_2$O is required to form the precipitate of potassium tetraphenylborate

**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 30 % (mass fraction) - 50 % (mass fraction) and 10 % (mass fraction) - 20 % (mass fraction) are 100.2 % - 100.8 % and 99.3 % - 102.2 % as water soluble potassium (W-K$_2$O) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.7 % (mass fraction).

**References**

(5) **Flow sheet for water soluble potassium**: The flow sheet for water soluble potassium in fertilizers is shown below:

- **2.5 g analytical sample** (potassium salts, etc.)
- Weigh to the order of 1 mg into a 300-mL tall beaker
- **About 200 mL water**
- **Heating**
- Cover with a watch glass, boil for 15 minutes
- **Cooling**
- Immediately
- **Transfer**
- Water, 250-mL volumetric flask
- **Water (up to the marked line)**
- **Filtration**
- Type 3 filter paper
- **Sample solution**

Figure 1-1 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.1))

About 400 mL of water
Rotary shaker (30 - 40 revolutions/min)
For 30 minutes
Water (up to the marked line)
Type 3 filter paper

Sample solution

5 g analytical sample (compound fertilizer)
Weigh to the order of 1 mg into a 500-mL volumetric flask

Shake to mix

Filtration

Sample solution

Figure 1-2 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.2))

Sample solution

Aliquot (predetermined amount)

100-mL tall beaker

Water (add tetraphenylborate solution to reach 50 mL)
2 mL of hydrochloric acid (1+9)
5 mL of formaldehyde solution
5 mL of ethylenediamine tetraacetate-sodium hydroxide solution
Tetraphenylborate solution (the equivalents of potassium + 4 mL)

For 30 minutes, sometimes mixing

Crucible type glass filtering apparatus 1G4,
5 times with 5 mL of tetraphenylborate washing solution

Washing

Drying

Standing to cool

Measurement

Figure 2 Flow sheet for water-soluble potassium in fertilizers
(Coloring and measurement procedure)
4.3.3.c Sodium tetraphenylborate volumetric analysis

(1) Summary
This testing method is applicable to fertilizers containing potassium salt but not organic matters. This testing method is classified as Type E and its symbol is 4.3.3.c-2017 or W-K.c-1. Extract by adding water to an analytical sample, mask co-existing ammonium and other salts with formaldehyde, and make potassium ion and tetraphenylborate react with each other. Measure unconsumed tetraphenylborate by conducting a precipitate titration to obtain water-soluble potassium (W-K_2O) in an analytical sample.

(2) Reagent: Reagents are as shown below.
   a) Formaldehyde solution: A JIS Guaranteed Reagent specified in JIS K 8872 or a reagent of equivalent quality.
   b) Sodium hydroxide solution (120 g/L) \(^{(1)}\): Dissolve 30 g of sodium hydroxide specified in JIS K 8576 in water to make 250 mL.
   c) Tetraphenylborate solution \(^{(1)}\): Transfer 12.2 g of sodium tetraphenylborate to a 1000-mL volumetric flask, dissolve by adding about 800 mL of water and add 3 mL of sodium hydroxide (120 g/L) to the total filtrate, and further add water up to the marked line. Filter with Type 3 filter paper in the case of usage.
   d) Benzalkonium chloride solution (3.3 g/500 mL) \(^{(1)}\): Dissolve 3.3 g of benzalkonium chloride in 500 mL of water.
   e) Methyl red solution (0.1 g/100 mL): Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.
   f) Titan Yellow solution (0.04 g/100 mL): Dissolve 0.04 g of Titan Yellow in 100 mL of water in the case of usage.
   g) Potassium standard solution (K_2O 2 mg/mL) \(^{(1)}\): Heat potassium chloride specified in JIS K 8121 at 110 °C ± 2 °C for about 2 hours, let it stand to cool in a desiccator, and weigh 3.166 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.

   Note  (1) This is an example of preparation; prepare an amount as appropriate.

(3) Instruments: Instruments are as shown below:
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 °C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Compound fertilizers containing potassium salts and magnesium potassium sulfate
   a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 300-mL tall beaker.
   b) Add about 200 mL of water, and cover with a watch glass and heat on a hot plate to boil for about 15 minutes.
   c) Immediately transfer it with water to a 250- mL volumetric flask.
   d) After immediate cooling is complete, add water up to the marked line
   e) Filter with Type 3 filter paper to make a sample solution.

   Comment 1 In the procedure in a), a 250-mL volumetric flask can be used instead of a 300-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. Additionally, “cover with a watch glass” in b) is replaced by “place a funnel”. Skip “transfer to a 250-mL
volumetric flask with water” in the procedure in c).

Comment 2 The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.3.3.a.

(4.1.2) Compound fertilizers containing no magnesium potassium sulfate

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.

b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

Comment 3 In the procedure in a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it into a 250-mL volumetric flask.

Comment 4 The procedure in (4.1.2) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.2) Precipitate formation: Form precipitate as shown below.

a) Transfer 5 mL - 15 mL (no more than the equivalents of 30 mg as K₂O) of the extract to a 100-mL volumetric flask.

b) Add water to the solution to make about 30 mL.

c) Add about 5 mL of formaldehyde solution and add 5 mL of sodium hydroxide solution (120 g/L).

d) Add 25 mL of tetr phenylborate solution at the rate of one or two drop (s) per second while shaking to mix.

e) After adding water up to the marked line, leave at rest for 10 minutes.

f) Filter with Type 3 filter paper to make a sample solution.

(4.3) Measurement: Conduct measurement as shown below.

a) Calibration curve preparation

1) Transfer 1 mL - 15 mL of potassium standard solution (K₂O 2 mg/mL) to 100-mL volumetric flasks step-by-step.

2) Conduct the same procedures as (4.2) b) - f) to make K₂O 2 mg/100 mL - 30 mg/100 mL of the potassium standard solutions for the calibration curve preparation.

3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.

4) Transfer 40 mL of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation to an Erlenmeyer flask respectively.

5) Add a few drops of Titan Yellow solution.

6) Titrate with a benzalkonium chloride solution (3.3 g/500 mL) until the color of the solution changes to light red \(^{(2)}\).

7) Prepare a curve for the relationship between the potassium concentration and the volume of the benzalkonium chloride solution (3.3 g/500 mL) required for the titration of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

b) Sample measurement

1) Transfer 40 mL of the sample solution of (4.2) f) to a 100-mL Erlenmeyer flask.

2) Conduct similarly as in a) 5) - 6) to obtain the volume of the benzalkonium chloride solution (3.3 g/500 mL) required for the titration.

3) Obtain the potassium content from the calibration curve, and calculate water soluble potassium (W-K₂O) in the analytical sample.
Note (2) If the solution temperature is no more than 20 °C, the reaction does not advance in some cases. Therefore, it is recommended to heat the solution up to about 30 °C.

References

(5) Flow sheet for water soluble potassium: The flow sheet for water soluble potassium in fertilizers is shown below:

Figure 1-1 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.1))

Figure 1-2 Flow sheet for water-soluble potassium in fertilizers
(Extraction procedure (4.1.2))
<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
<tr>
<td>Aliquot (5 mL to 15 mL)</td>
<td>100-mL volumetric flask</td>
</tr>
<tr>
<td>Water (the liquid volume to reach about 30 mL)</td>
<td></td>
</tr>
<tr>
<td>About 5 mL of formaldehyde solution</td>
<td></td>
</tr>
<tr>
<td>5 mL of sodium hydroxide solution (120 g/L)</td>
<td></td>
</tr>
<tr>
<td>25 mL tetraphenylborate solution</td>
<td>(one or two drop(s) per second while shaking to mix)</td>
</tr>
<tr>
<td>Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Leaving at rest</td>
<td>10 minutes</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Aliquot (40 mL)</td>
<td>100-mL Erlenmeyer flask</td>
</tr>
<tr>
<td>A few drops of Titan Yellow solution</td>
<td></td>
</tr>
<tr>
<td>Titration</td>
<td>Benzaikonium chloride solution (3.3 g/500 mL)</td>
</tr>
<tr>
<td></td>
<td>(until the solution changes to light red)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for water-soluble potassium in fertilizers
(Precipitate formation and measurement procedure)
4.3.3.d ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.3.3.d-2017 or W-K.d-1.

Add water to an analytical sample to extract, introduce it to an ICP Optical Emission Spectrometer ("ICP-OES") and measure the potassium at a wavelength of 766.491 nm to obtain water-soluble potassium acid (W-K₂O). In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Potassium standard solution (K₂O 1 mg/mL) (1): Heat potassium chloride specified in JIS K 8121 at 110 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 1.583 g into a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   d) Potassium standard solution (K₂O 20 µg/mL - 0.16 mg/mL) for the calibration curve preparation (1): Transfer 2 mL - 16 mL of potassium standard solution (K₂O 1 mg/mL) to 100-mL volumetric flasks step-by-step, add about 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   e) Potassium standard solution (K₂O 2 µg/mL - 20 µg/mL) for the calibration curve preparation (1): Transfer 2 mL - 20 mL of potassium standard solution (K₂O 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in e).

Note  (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the potassium standard solution in (2), a potassium standard solution for the calibration curve preparation can be prepared by using a potassium standard solution (K 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble potassium (W-K₂O) in the analytical sample by multiplying the concentration (K) of a potassium standard solution for calibration curve preparation or a measurement value (K) obtained in (4.2) by a conversion factor (1.2046).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The axial observation mode does not apply to potassium since interference is serious.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 mg of an analytical sample (2) to the order of 1 g, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, shake to mix and add water up to the marked line.

c) Filter with Type 3 filter paper to make a sample solution.

Note (2) The sampling amount of the analytical sample is 10 g when there is less potassium content in the fertilizers such as a home garden-use fertilizer.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct measurement according to JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Optical Emission Spectrometer: Set up the measurement conditions for the ICP Optical Emission Spectrometer considering the following:
Analytical line wavelength: 766.491 nm

b) Calibration curve preparation
1) Spray the potassium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 766.491 nm.
2) Prepare a curve for the relationship between the potassium concentration and the indicated value of the potassium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
1) Transfer a predetermined amount of the sample solution (the equivalents of 0.2 mg - 16 mg as K2O) to a 100-mL volumetric flask.
2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
3) Subject to the same procedure as in b) 1) to read the indicated value.
4) Obtain the potassium content from the calibration curve, and calculate water soluble potassium (W-K2O) in the analytical sample.

Comment 4 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

Comment 5 The comparison of the measurement value (yi; 0.641 % (mass fraction) - 7.23 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value (xi) of Flame atomic absorption spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was y = −0.021 + 0.969x, and its correlation coefficient (r) was 0.999. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery rates at additive level of 5 % (mass fraction) and 0.4 % (mass fraction) were 102.3 % and 104.0 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.05 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$ (%)</th>
<th>Repeatability $^4$</th>
<th>Intermediate precision $^6$</th>
<th>Intermediate relative standard deviation $^7$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>5.69 (3%)</td>
<td>0.02 (3%)</td>
<td>0.06 (3%)</td>
<td>1.1 (1%)</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (liquid)</td>
<td>7</td>
<td>2.29 (3%)</td>
<td>0.02 (3%)</td>
<td>0.04 (3%)</td>
<td>1.6 (1%)</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $T$) × the number of duplicate testing (2)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References


(5) Flow sheet: The flow sheet for water-soluble potassium of the fluid mixed fertilizers is shown below:

1 g analytical sample ← Weigh to the order of 1 mg to a 100-mL volumetric flask
Shaking to mix ← Water, about 50 mL
Water (up to the marked line)
Filtration
Sample solution

Figure 1 Flow sheet for water-soluble potassium in fluid fertilizers
(Extraction procedure)

Sample solution
Aliquot (predetermined volume) ← 100-mL volumetric flask
25 mL of hydrochloric acid (1+5)
Water (up to the marked line)
Measurement ICP-OES (766.491nm)

Figure 2 Flow sheet for water-soluble potassium in fluid fertilizers
(Measurement procedure)
4.4 Silicic acid

4.4.1 Soluble silicic acid

4.4.1.a Potassium fluoride method

(1) Summary

This testing method is applicable to fertilizers containing no silica gel fertilizers. This testing method is classified as Type D and its symbol is 4.4.1.a-2017 or S-Si.a-1.

Extract by adding hydrochloric acid (1+23) to an analytical sample, add hydrochloric acid, potassium fluoride solution and potassium chloride and cool in a refrigerator, and then filter after forming precipitate as potassium silicofluoride ($K_2SiF_6$). Put the precipitate in water and heat, and measure potassium silicofluoride ($K_2SiF_6$) dissolved by precipitation titration to obtain the hydrochloric acid (1+23) soluble silicic acid (soluble silicic acid (S-SiO$_2$)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution} = \left( \frac{W_1 \times A \times 0.01/97.095}{V_1} \times \frac{V_2}{V_3} \times \frac{1000}{V_3} \times \frac{1}{C} \right)
\]

\[W_1:\] Mass (g) of sulfamic acid sampled
\[A:\] Purity (% (mass fraction)) of sulfamic acid
\[V_1:\] Volume (mL) of sulfamic acid solution transferred
\[V_2:\] Constant volume (250 mL) of sulfamic acid solution
\[V_3:\] Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
\[C:\] Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

c) Potassium chloride: A JIS Guaranteed Reagent specified in JIS K 8121 or a reagent of equivalent quality.

d) Potassium chloride solution (1): Add 250 mL of ethanol specified in JIS K 8101 to 750 mL of water to mix, and add 150 mL of potassium chloride to dissolve. Add a few drops of methyl red solution (0.1 g/100 mL) as an indicator and drop hydrochloric acid until the color of the solution becomes red to make it acidic. After leaving at rest for 1 day, neutralize with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution.

e) Potassium fluoride solution (1): Dissolve 58 g of potassium fluoride specified in JIS K 8815
in 1000 mL of water (2).

f) **Methyl red solution (0.1 g/100 mL):** Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

g) **Phenolphthalein solution (1 g/100 mL):** Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.

**Note**

1. This is an example of preparation; prepare an amount as appropriate.
2. Store in a container made of polymer that contains no silicon.

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

(3) **Apparatus and instruments:** Apparatus and instruments are shown below.

a) **Constant-temperature rotary shaker:** A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 ºC ± 1 ºC, upside down at 30 - 40 revolutions/min.

b) **Hot plate:** A hot plate whose surface temperature can be adjusted up to 250 ºC.

c) **Beaker made of polymer:** A beaker made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in (4.1).

d) **Filter made of polymer:** A Gooch crucible made of polymer (compatible filter diameter: 25 mm) or a funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) A Gooch crucible made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in (4.1).

**Comment 2** A funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) is sold under the name Polyethylene Kiriyama Funnel SB-21.

(4) **Test procedure**

**4.1 Extraction:** Conduct extraction as shown below.

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.

b) Add 150 mL of hydrochloric acid (1+23) heated up to about 30 ºC (3), and shake to mix at 30 - 40 revolutions/min (30 ºC ± 1 ºC) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

**Note**

3. Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1) of 4.4.1.d.

**Comment 4** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedure of (4.1) b).

**4.2 Measurement:** Conduct measurement as shown below.

a) Transfer a predetermined volume (the equivalents of 20 mg - 50 mg as SiO₂ and no more than 25 mL of liquid volume) to a 200-mL beaker made of polymer.

b) Add about 10 mL of hydrochloric acid and about 15 mL of potassium fluoride solution, and further add about 2 g of potassium chloride to dissolve, and then cool in a refrigerator for
about 30 minutes or more (4) to form the precipitate of potassium fluoride.

c) Filter under reduced pressure with a filter made of polymer (5) topped with Type 6 filter paper, and wash the container 3 times with a potassium chloride solution, then move the whole precipitate into the filter, and further wash 6 - 7 times with a small amount of potassium chloride solution (6).

d) Move the precipitate on the filter together with the filter paper into a 300-mL tall beaker with water, and further add water to make about 200 mL and heat it up to 70 °C - 80 °C on a hot plate.

e) Add a few drops of phenolphthalein solution (1 g/100 mL) to the sample solution as an indicator and titrate with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes light red.

f) Calculate soluble silicic acid (S-SiO₂) by the following formula.

\[
\text{Soluble silicic acid (S-SiO}_2\text{) (% (mass fraction)) in an analytical sample} = V_4 \times C \times f \times (\frac{V_5}{V_6}) \times (15.021/W_2) \times (100/1000)
\]

- \( V_4 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- \( C \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \( f \): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \( V_5 \): Predetermined volume (mL) of the extract in (4.1) c)
- \( V_6 \): Volume (mL) of the extract transferred in (4.2) a)
- \( W_2 \): Mass (g) of the analytical sample

Note (4) To be no more than 10 °C
(5) Filter paper pulp can be stuffed to restrain precipitate from outflowing.
(6) Until the filtrate becomes neutral.

Comment 5 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 25 % (mass fraction) - 40 % (mass fraction) and 10 % (mass fraction) are 98.4 % - 100.5 % and 101.0 % as soluble silicic acid (S-SiO₂) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.3 % (mass fraction).

References
5) **Flow sheet for soluble silicic acid:** The flow sheet for soluble silicic acid in fertilizers is shown below:

![Flow sheet for soluble silicic acid in fertilizers (Extraction procedure)](image)

**Figure 1** Flow sheet for soluble silicic acid in fertilizers (Extraction procedure)

![Flow sheet for soluble silicic acid in fertilizers (Measurement procedure)](image)

**Figure 2** Flow sheet for soluble silicic acid in fertilizers (Measurement procedure)
4.4.1.b Potassium fluoride method (Silica gel fertilizers, etc.)

(1) Summary
This testing method is applicable to silica gel fertilizers and silica hydrogel fertilizers. This testing method is classified as Type B and its symbol is 4.4.1.b-2017 or S-Si.b-1.

Extract by adding hydrochloric acid (1+23) to an analytical sample, add hydrochloric acid, potassium fluoride solution and potassium chloride and cool in a refrigerator, and then filter after forming precipitate as potassium silicofluoride (K$_2$SiF$_6$). Put the precipitate in water and heat, and measure potassium silicofluoride (K$_2$SiF$_6$) dissolved by precipitation titration to obtain the sodium hydroxide (20 g/L) soluble silicic acid (soluble silicic acid (S-SiO$_2$)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent: Reagents are as shown below.

a) $0.1 \text{ mol/L} - 0.2 \text{ mol/L sodium hydroxide solution}^{(1)}$: Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

**Standardization:** Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line $^{(1)}$. Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

$$ f_1 = \left( \frac{W_1 \times A \times 0.01}{97.095} \right) \times \left( \frac{V_1}{V_2} \right) \times \left( \frac{1000}{V_3} \right) \times \left( \frac{1}{C} \right) $$

$W_1$: Mass (g) of sulfamic acid sampled

$A$: Purity (% (mass fraction)) of sulfamic acid

$V_1$: Volume (mL) of sulfamic acid solution transferred

$V_2$: Constant volume (250 mL) of sulfamic acid solution

$V_3$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

$C$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.

c) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

d) Potassium chloride: A JIS Guaranteed Reagent specified in JIS K 8121 or a reagent of equivalent quality.

e) Potassium chloride solution $^{(1)}$: Add 250 mL of ethanol specified in JIS K 8101 to 750 mL of water to mix, and add 150 mL of potassium chloride to dissolve. Add a few drops of methyl red solution (0.1 g/100 mL) as an indicator and drop hydrochloric acid until the color of the solution becomes red to make it acidic. After leaving at rest for 1 day, neutralize with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution.

f) Potassium fluoride solution $^{(1)}$: Dissolve 58 g of potassium fluoride specified in JIS K 8815...
in 1000 mL of water \(^{(2)}\).

**g) Methyl red solution (0.1 g/100 mL):** Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

**h) Phenolphthalein solution (1 g/100 mL):** Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.

**Note**  
(1) This is an example of preparation; prepare an amount as appropriate.  
(2) Store in a container made of polyethylene, etc. that contains no silicon.

**Comment 1**  
A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in \((2)\) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

**3) Apparatus and instruments:** Apparatus and instruments are shown below.

**a) Water bath:** A water bath that can be adjusted to 65 °C ± 2 °C.

**b) Hot plate:** A hot plate whose surface temperature can be adjusted up to 250 °C.

**c) Volumetric flask and beaker made of polymer:** A flask and a beaker that are made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in \((4.1)\).

**d) Gooch crucible made of polymer:** A Gooch crucible made of polymer (compatible filter diameter: 25 mm) or a funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) A Gooch crucible made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in \((4.1)\).

**Comment 2**  
A funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) is sold under the name Polyethylene Kiriyama Funnel SB-21.

**4) Test procedure**

**4.1) Extraction:** Conduct extraction as shown below.

**a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask made of polymer.**

**b) Add about 150 mL of sodium hydroxide solution heated up to about 65 °C, and heat for 1 hour while shaking to mix at every 10 minutes in a water bath at 65 °C ± 2 °C.**

**c) After immediate cooling is complete, add water up to the marked line**

**d) Filter with Type 3 filter paper to make a sample solution.**

**4.2) Measurement:** Conduct measurement as shown below.

**a) Transfer a predetermined volume (the equivalents of 20 mg - 50 mg as SiO2 and no more than 25 mL of liquid volume) to a 200-mL beaker made of polymer.**

**b) Add about 10 mL of hydrochloric acid and about 15 mL of potassium fluoride solution, and further add about 2 g of potassium chloride to dissolve, and then cool in a refrigerator for about 30 minutes \(^{(3)}\) to form the precipitate of potassium fluoride.**

**c) Filter under reduced pressure with a filter made of polymer \(^{(4)}\) topped with Type 6 filter paper, and wash the container 3 times with a potassium chloride solution, then move the whole precipitate into the filter, and further wash 6 - 7 times with a small amount of potassium chloride solution \(^{(5)}\).**

**d) Move the precipitate on the filter together with the filter paper into a 300-mL tall beaker with water, and further add water to make about 200 mL and heat it up to 70 °C - 80 °C on a hot plate.**

**e) Add a few drops of phenolphthalein solution (1 g/100 mL) as an indicator and titrate with the
0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes light red.

f) Calculate soluble silicic acid (S-SiO$_2$) by the following formula.

\[
\text{Soluble silicic acid (S-SiO}_2\text{) (% (mass fraction)) in an analytical sample} = V_4 \times C \times f \times (V_5/V_6) \times (15.021/W_2) \times (100/1000)
\]

- $V_4$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- $C$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- $f$: Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- $V_5$: Predetermined volume (mL) of the extract in (4.1) e)
- $V_6$: Volume (mL) of the extract transferred in (4.2) a)
- $W_2$: Mass (g) of the analytical sample

Note
1) To be no more than 10 ºC
2) Filter paper pulp can be stuffed to restrain precipitate from outflowing.
3) Until the filtrate becomes neutral.

**Comment 3** Table 1 shows results and analysis results from a collaborative study for testing method validation.

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^2$ (%)</th>
<th>$s_r^3$ (%)</th>
<th>$RSD_r^4$ (%)</th>
<th>$s_R^5$ (%)</th>
<th>$RSD_R^6$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica gel fertilizers 1</td>
<td>8</td>
<td>79.37</td>
<td>0.23</td>
<td>0.3</td>
<td>0.55</td>
<td>0.7</td>
</tr>
<tr>
<td>Silica gel fertilizers 2</td>
<td>8</td>
<td>84.68</td>
<td>0.42</td>
<td>0.5</td>
<td>0.85</td>
<td>1.0</td>
</tr>
<tr>
<td>Silica gel fertilizers 3</td>
<td>8</td>
<td>89.58</td>
<td>0.4</td>
<td>0.4</td>
<td>0.51</td>
<td>0.6</td>
</tr>
<tr>
<td>Silica gel fertilizers 4</td>
<td>8</td>
<td>84.44</td>
<td>0.37</td>
<td>0.4</td>
<td>0.77</td>
<td>0.9</td>
</tr>
<tr>
<td>Silica gel fertilizers 5</td>
<td>8</td>
<td>85.77</td>
<td>0.46</td>
<td>0.5</td>
<td>0.59</td>
<td>0.7</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n$ = number of laboratories × number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

**References**


2) Akira SHIMIZU, Shin ABE and Jun ITO: Determination of Solubility Silicic Acid in Silica gel Fertilizer and Silica gel-including Fertilizer by Potassium Fluoride Method: A
5) **Flow sheet for soluble silicic acid:** The flow sheet for soluble silicic acid in fertilizers is shown below:

![Flow sheet for soluble silicic acid in silica gel fertilizers (Extraction procedure)](image1)

- **Weigh to the order of 1 mg** into a 250-mL volumetric flask made of polymer.
- **About 150 mL sodium hydroxide (20 g/L) [about 65 ºC]**
- **65 ºC ± 2 ºC, 1 hour, shaking to mix every 10 minutes**
- **Immediately**
- **Water (up to the marked line)**
- **Type 3 filter paper**
- **Filtration**
- **Sample solution**

**Figure 1** Flow sheet for soluble silicic acid in silica gel fertilizers (Extraction procedure)

![Flow sheet for soluble silicic acid in silica gel fertilizers (Measurement procedure)](image2)

- **Aliquot (predetermined amount)**
- **200-mL beaker made of polymer**
- **About 10 mL hydrochloric acid**
- **About 15 mL potassium fluoride solution**
- **About 2 g of potassium chloride**
- **For 30 minutes or more in a refrigerator**
- **Filtration under reduced pressure**
- **Gooch filter made of polymer, Type 6 filter paper**
- **Washing**
- **Wash 6 - 7 times with potassium chloride solution**
- **Transfer**
- **300-mL tall beaker, water**
- **Water (until the liquid volume reaches about 200 mL)**
- **Heating**
- **70 ºC - 80 ºC**
- **A few drops of phenolphthalein solution (1 g/100 mL)**
- **Titration**
- **0.1 mol/L - 0.2 mol/L sodium hydroxide solution**
- **(Until the solution becomes light red)**

**Figure 2** Flow sheet for soluble silicic acid in silica gel fertilizers (Measurement procedure)
4.4.1.c Potassium fluoride method (Fertilizers containing silica gel fertilizers)

(1) Summary
This testing method is applicable to fertilizers containing silica gel fertilizers. This testing method is classified as Type B and its symbol is 4.4.1.c-2017 or S-Si.c-1.

Mix the equivalent volumes of the extract which is extracted by adding hydrochloric acid (1+23) to an analytical sample and the liquid by extracting non-dissolved matter on a filter paper with sodium hydroxide (20 g/L), and add hydrochloric acid, a potassium fluoride solution and potassium chloride. Cool it in a refrigerator, and then filter after forming precipitate as potassium silicofluoride (K$_2$SiF$_6$). Then transfer water in the precipitate, heat and measure potassium silicofluoride (K$_2$SiF$_6$) dissolved by precipitation titration to obtain soluble silicic acid (S-SiO$_2$) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution $^{(1)}$: Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL - 11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line $^{(1)}$. Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (f)} = \left( W_1 \times A \times 0.01/97.095 \right) \times \left( V_1/V_2 \right) \times \left( 1000/V_3 \right) \times \left( 1/C \right)
\]

$W_1$: Mass (g) of sulfamic acid sampled
$A$: Purity (%) of sulfamic acid
$V_1$: Volume (mL) of sulfamic acid solution transferred
$V_2$: Constant volume (250 mL) of sulfamic acid solution
$V_3$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
$C$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.

c) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

d) Potassium chloride: A JIS Guaranteed Reagent specified in JIS K 8121 or a reagent of equivalent quality.

e) Potassium chloride solution $^{(1)}$: Add 250 mL of ethanol specified in JIS K 8101 to 750 mL of water to mix, and add 150 mL of potassium chloride to dissolve. Add a few drops of methyl red solution (0.1 g/100 mL) as an indicator and drop hydrochloric acid until the color of the solution becomes red to make it acidic. After leaving at rest for 1 day, neutralize with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution.
f) **Potassium fluoride solution** \(^{(1)}\): Dissolve 58 g of potassium fluoride specified in JIS K 8815 in 1000 mL of water \(^{(2)}\).

g) **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

h) **Phenolphthalein solution (1 g/100 mL)**: Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.

**Note**  
(1) This is an example of preparation; prepare an amount as appropriate.

(2) Store in a container made of polymer that contains no silicon.

**Comment 1**  
A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in \((2)\ a)\) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

(3) **Instruments**: Instruments are as shown below:

   a) **Water bath**: A water bath that can be adjusted to 65 °C ± 2 °C.

   b) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.

   c) **Volumetric flask and beaker made of polymer**: A flask and a beaker that are made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in \((4.1)\).

   d) **Filter made of polymer**: A Gooch crucible made of polymer (compatible filter diameter: 25 mm) or a funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) A Gooch crucible made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in \((4.1)\).

**Comment 2**  
A funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) is sold under the name Polyethylene Kiriyama Funnel SB-21.

(4) **Test procedure**

\((4.1)\) **Extraction**: Conduct extraction as shown below.

   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 300-mL tall beaker.

   b) Add 150 mL of hydrochloric acid (1+23) warmed up to about 30 °C, and warm it in a water bath at 30 °C ± 2 °C while stirring every 10 minutes with a glass rod for 1 hour.

   c) After immediate cooling is complete, is complete, filter with Type 6 filter paper to a 250-mL volumetric flask as an acceptor. Wash the tall beaker with water, then move the whole residue on the filter paper and add water up to the marked line to make a sample solution (1).

   d) Transfer the non-dissolved matter on the filter paper together with the filter paper to a 250-mL volumetric flask made of polymer.

   e) Add 150 mL of sodium hydroxide solution heated up to about 65 °C, and heat for 1 hour while shaking to mix at every 10 minutes in a water bath at 65 °C ± 2 °C.

   f) After immediate cooling is complete, add water up to the marked line and filter with Type 3 filter paper to make a sample solution (2).

\((4.2)\) **Measurement**: Conduct measurement as shown below.

   a) Transfer a predetermined volume (the equivalents of 20 mg - 50 mg as SiO2) \(^{(3)}\) of the sample solution (1) and (2) to a 200-mL beaker made of polymer.

   b) Add about 10 mL of hydrochloric acid and about 15 mL of potassium fluoride solution, and further add about 2 g of potassium chloride to dissolve, and then cool in a refrigerator for about 30 minutes or more \(^{(4)}\) to form the precipitate of potassium fluoride.

   c) Filter under reduced pressure with a filter made of polymer \(^{(5)}\) topped with Type 6 filter paper,
and wash the container 3 times with a potassium chloride solution, then move the whole precipitate into the filter, and further wash 6 - 7 times with a small amount of potassium chloride solution (6).

d) Move the precipitate on the filter together with the filter paper into a 300-mL tall beaker with water, and further add water to make about 200 mL and heat it up to 70 °C - 80 °C on a hot plate.

e) Add a few drops of phenolphthalein solution (1 g/100 mL) as an indicator and titrate with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes light red.

f) Calculate soluble silicic acid (S-SiO₂) by the following formula.

\[
\text{Soluble silicic acid (S-SiO}_2\text{)} \text{ (% (mass fraction)) in an analytical sample} = V_4 \times C \times f \times (V_5/V_6) \times (15.021/W_2) \times (100/1000)
\]

- \(V_4\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- \(C\): Estimated concentration (mol/L) of sodium hydroxide solution (0.1 mol/L - 0.2 mol/L)
- \(f\): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- \(V_5\): Predetermined volume (mL) of the sample solution in (4.1) c)
- \(V_6\): Transferred amount (mL) of the sample solution in (4.2) a)
- \(W_2\): Mass (g) of the analytical sample

**Note** (3) The transferred volume of the sample solution (1) and the sample solution (2) should be equivalent.

(4) To be no more than 10 °C

(5) Filter paper pulp can be stuffed to restrain precipitate from outflowing.

(6) Until the filtrate becomes neutral.

**Comment 3** Table 1 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this testing method is about 0.6 % (mass fraction).
Table 1  Results and analysis results from a collaborative study for the test method validation of soluble silicic acid in fertilizers including a silica gel fertilizer

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (2)</th>
<th>$s_r$ (4)</th>
<th>$RSD_r$ (5)</th>
<th>$s_R$ (6)</th>
<th>$RSD_R$ (7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed phosphate fertilizer 1</td>
<td>8</td>
<td>24.99</td>
<td>0.16</td>
<td>0.6</td>
<td>0.33</td>
<td>1.3</td>
</tr>
<tr>
<td>Mixed phosphate fertilizer 2</td>
<td>8</td>
<td>34.50</td>
<td>0.26</td>
<td>0.7</td>
<td>0.48</td>
<td>1.4</td>
</tr>
<tr>
<td>Compound fertilizers 1</td>
<td>8</td>
<td>30.30</td>
<td>0.13</td>
<td>0.4</td>
<td>0.60</td>
<td>2.0</td>
</tr>
<tr>
<td>Compound fertilizers 2</td>
<td>8</td>
<td>33.34</td>
<td>0.13</td>
<td>0.4</td>
<td>0.47</td>
<td>1.4</td>
</tr>
<tr>
<td>Compound fertilizers 3</td>
<td>8</td>
<td>15.76</td>
<td>0.11</td>
<td>0.7</td>
<td>0.21</td>
<td>1.3</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Total mean ($n = \text{number of laboratories} \times \text{number of replication(2)}$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(5) **Flow sheet for soluble silicic acid:** The flow sheet for soluble silicic acid in fertilizers including silica gel fertilizers is shown below:

```
<table>
<thead>
<tr>
<th>1 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 300-mL tall beaker</th>
</tr>
</thead>
<tbody>
<tr>
<td>← About 150 mL hydrochloric acid (1+23) [30 °C]</td>
<td>1 hour in water bath at 30 °C ± 2 °C</td>
</tr>
<tr>
<td>Warming</td>
<td>(Stir at every 10 minutes)</td>
</tr>
</tbody>
</table>

| Cooling               | Immediately                                          |
| Filtration (supernatant) | Type 6 filter paper, a 250-mL volumetric flask        |
| Transfer              | Move the whole residue to a filter paper              |
| ← Wash with water     | <Residue>                                             |
| ← Wash (up to the marked line) | <Filtrate>                                           |
| Sample solution (1)   |                                                     |

| Put                   | 250-mL volumetric flask made of polymer together with a filter paper |
|← Add about 150 mL of sodium hydroxide solution (20 g/L) [65°C] to stopple. |
| Heating               | 65 °C ± 2 °C, while shaking to mix at every 10 minutes for 1 hour |
| Cooling               | Immediately                                          |
| ← Water (up to the marked line) |                                                        |
| Filtration            | Type 3 filter paper                                   |
| Sample solution (2)   |                                                     |
```

Figure 1  Flow sheet for soluble silicic acid in fertilizers
(Extraction procedure)
About 10 mL of hydrochloric acid
About 15 mL of potassium fluoride solution
About 2 g of potassium chloride

For 30 minutes in a refrigerator

Filter made of polymer
Type 6 filter paper

Wash 6 - 7 times with potassium chloride solution

300-mL tall beaker, water

Water (until the liquid volume reaches about 200 mL)

70 ºC - 80 ºC

A few drops of phenolphthalein solution (1 g/100 mL)

0.1 mol/L - 0.2 mol/L sodium hydroxide solution
(Until the color of solution becomes light red)

Figure 2  Flow sheet for soluble silicic acid in fertilizers
(Measurement procedure)
4.4.1.d Perchloric acid method

(1) Summary
This testing method is applicable to fertilizers containing no silica gel fertilizers. This testing method is classified as Type E and its symbol is 4.4.1.d-2017 or S-Si.d-1.

Extract by adding hydrochloric acid (1+23) to an analytical sample, add perchloric acid and heat, and then measure the formed silicic acid anhydride to obtain hydrochloric acid (1+23) soluble silicic acid (soluble silicic acid (S-SiO2)) in an analytical sample.

(2) Reagents
a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
b) Perchloric acid: A JIS Guaranteed Reagent specified in JIS K 8223 or a reagent of equivalent quality.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
a) Constant-temperature rotary shaker: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.
b) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 °C.
c) Electric furnace: An electric furnace that can be adjusted to 1000 °C - 1100 °C.
d) Crucible: After heating a chemical analysis porcelain crucible specified in JIS R 1301 in an electric funnel at 1000 °C - 1100 °C, let it stand to cool in a desiccator and measure the mass to the order of 1 mg.

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.
a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.
b) Add 150 mL of hydrochloric acid (1+23) heated up to about 30 °C, and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.
c) After immediate cooling is complete, add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 1 The procedure in (4.1) is the same as the procedure in (4.1) of 4.4.1.a.

(4.2) Measurement: Conduct measurement as shown below.
a) Transfer a predetermined volume to a 100-mL tall beaker.
b) Add about 10 mL of perchloric acid and heat.
c) When white smoke from the perchloric acid starts evolving, cover with a watch glass, then heat for 15 - 20 minutes to form precipitate of silicone dioxide.
d) After standing to cool, add about 50 mL of hydrochloric acid (1+4) and cover with a watch glass and heat at 70 °C - 80 °C on a hot plate for several minutes.
e) Immediately after heating, filter with Type 5-C filter paper, wash the container with heated hydrochloric acid (1+10) and move the whole precipitate to a filter paper.
f) Wash the precipitate and the filter paper 2 times with heated hydrochloric acid (1+10), and further wash several times with hot water (1).
g) Put the precipitate together with the filter paper into the crucible.
h) Put the crucible into a drying apparatus and dry at about 120 °C for 1 hour.
i) After standing to cool, put the crucible into an electric funnel and heat gently to char (2).
j) Ignite at 1000 °C - 1100 °C for 1 hour (2).
k) After ignition, move the crucible to a desiccator and let it stand to cool.
l) After standing to cool, remove the crucible from the desiccator and measure the mass to the order of 1 mg.
m) Calculate soluble silicic acid (S-SiO$_2$) by the following formula.

\[
\text{Soluble silicic acid (S-SiO}_2\text{) (% (mass fraction)) in an analytical sample} = A \times \frac{V_1}{W} \times 100
\]

\(A\): Mass (g) of the precipitate
\(W\): Mass (g) of the analytical sample
\(V_1\): Predetermined volume (mL) of the sample solution in (4.1) c)
\(V_2\): Transferred amount (mL) of the sample solution in (4.2) a)

**Note**  
(1) Wash until no reaction by chloride appears in the filtrate.  
(2) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 1000 °C - 1100 °C in 1 to 2 hours.

**References**  

5) **Flow sheet for soluble silicic acid**: The flow sheet for soluble silicic acid in fertilizers is shown below:

<table>
<thead>
<tr>
<th>1 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 250-mL volumetric flask.</th>
</tr>
</thead>
<tbody>
<tr>
<td>← About 150 mL of hydrochloric acid (1+23) [about 30 °C]</td>
<td>Rotary shaker (30 - 40 revolutions/min)</td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>30 °C ± 1 °C, 1 hour</td>
</tr>
<tr>
<td>Cooling</td>
<td>Immediately</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Filtration</td>
<td>Sample solution</td>
</tr>
</tbody>
</table>

Figure 1  Flow sheet for soluble silicic acid in fertilizers (Extraction procedure)

Figure 2  Flow sheet for soluble silicic acid in fertilizers (Measurement procedure)
4.4.2 Water-soluble silicic acid

4.4.2.a Potassium fluoride method

(1) Summary
This testing method is applicable to liquid potassium silicate fertilizers. This testing method is classified as Type D and its symbol is 4.4.2.a-2017 or W-Si.a.1.

Extract by adding water to an analytical sample, add hydrochloric acid, a potassium fluoride solution, and potassium chloride, and cool in a refrigerator, and then filter after forming precipitate as potassium silicofluoride (K$_2$SiF$_6$) precipitate. Put the precipitate in water and heat, and measure potassium silicofluoride (K$_2$SiF$_6$) dissolved by precipitation titration to obtain the water-soluble silicic acid (W-SiO$_2$) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.
  a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution: Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL - 11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.
  b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
  c) Potassium chloride: A JIS Guaranteed Reagent specified in JIS K 8121 or a reagent of equivalent quality.
  d) Potassium chloride solution: Add 250 mL of ethanol specified in JIS K 8101 to 750 mL of water to mix, and add 150 mL of potassium chloride to dissolve. Add a few drops of methyl red solution (0.1 g/100 mL) as an indicator and drop hydrochloric acid until the color of the solution becomes red to make it acidic. After leaving at rest for 1 day, neutralize with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution.
  e) Potassium fluoride solution: Dissolve 58 g of potassium fluoride specified in JIS K 8815 in 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line. Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution } (f) = \left( \frac{W_1 \times A \times 0.01/97.095}{V_1} \right) \times \left( \frac{V_2}{V_3} \right) \times \left( \frac{1000}{V_3} \right) \times \left( \frac{1}{C} \right)
\]

\( W_1 \): Mass (g) of sulfamic acid sampled
\( A \): Purity (% (mass fraction)) of sulfamic acid
\( V_1 \): Volume (mL) of sulfamic acid solution transferred
\( V_2 \): Constant volume (250 mL) of sulfamic acid solution
\( V_3 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
\( C \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

c) Potassium chloride: A JIS Guaranteed Reagent specified in JIS K 8121 or a reagent of equivalent quality.

d) Potassium chloride solution: Add 250 mL of ethanol specified in JIS K 8101 to 750 mL of water to mix, and add 150 mL of potassium chloride to dissolve. Add a few drops of methyl red solution (0.1 g/100 mL) as an indicator and drop hydrochloric acid until the color of the solution becomes red to make it acidic. After leaving at rest for 1 day, neutralize with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution.

e) Potassium fluoride solution: Dissolve 58 g of potassium fluoride specified in JIS K 8815 in 1000 mL of water.
f) Methyl red solution (0.1 g/100 mL): Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

g) Phenolphthalein solution (1 g/100 mL): Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) Store in a container made of polyethylene, etc. that contains no silicon.

Comment 1 A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

(3) Instruments: Instruments are as shown below:
  a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
  b) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 ºC.
  c) Beaker made of polymer: A beaker made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in (4.1).
  d) Filter made of polymer: A Gooch crucible made of polymer (compatible filter diameter: 25 mm) or a funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) A Gooch crucible made of polyethylene, etc. of a quality of material which prevents silicic acid from eluting in the extract procedure in (4.1).

Comment 2 A funnel for reduced pressure filtering made of polymer (compatible filter diameter: 21 mm) is sold under the name Polyethylene Kiriyama Funnel SB-21.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
  a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.
  b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
  c) Add water up to the marked line.
  d) Filter with Type 3 filter paper to make a sample solution.

Comment 3 In the procedure in a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it into a 250-mL volumetric flask.

Comment 4 The procedure in (4.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.2) Measurement: Conduct measurement as shown below.
  a) Transfer a predetermined volume (the equivalents of 20 mg - 50 mg as SiO₂ and no more than 25 mL of liquid volume) to a 200-mL beaker made of polymer.
  b) Add about 10 mL of hydrochloric acid and about 15 mL of potassium fluoride solution, and further add about 2 g of potassium chloride to dissolve, and then cool in a refrigerator for about 30 minutes or more (3) to form the precipitate of potassium fluoride.
  c) Filter under reduced pressure with a filter made of polymer (4) topped with Type 6 filter paper, and wash the container 3 times with a potassium chloride solution, then move the whole precipitate into the filter, and further wash 6 - 7 times with a small amount of potassium chloride solution (5).
  d) Move the precipitate on the filter together with the filter paper into a 300-mL tall beaker with water, and further add water to make about 200 mL and heat it up to 70 ºC - 80 ºC on a hot
plate.

e) Add a few drops of phenolphthalein solution (1 g/100 mL) to the sample solution as an indicator and titrate with the 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes light red.

f) Calculate water soluble silicic acid (W-SiO$_2$) by the following formula.

$$\text{Water soluble silicic acid (W-SiO}_2\text{) (\% (mass fraction)) in an analytical sample} = V_4 \times C \times f \times (V_5/V_6) \times (15.021/W_2) \times (100/1000)$$

$V_4$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration  
$C$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution  
$f$: Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution  
$V_5$: Predetermined volume (mL) of the extract in (4.1) c  
$V_6$: Volume (mL) of the extract transferred in (4.2) a  
$W_2$: Mass (g) of the analytical sample

**Note**  
(3) To be no more than 10 °C  
(4) Filter paper pulp can be stuffed to restrain precipitate from outflowing.  
(5) Until the filtrate becomes neutral.

**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 30 % (mass fraction) and 12 % (mass fraction) - 20% (mass fraction) are 100.7 % and 99.5 % - 100.5 as water-soluble silicic acid (W-SiO$_2$) respectively. The results of the repeatability tests on different days using a liquid potassium silicate fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.2 % (mass fraction).

**Table 1 Analysis results of the repeatability tests on different days**

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$ (%)</th>
<th>Repeatability $^4$ ($%$)</th>
<th>Intermediate precision $^6$ ($%$)</th>
<th>RSD $^7$ ($%$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liquid potassium silicate fertilizer 1</td>
<td>7</td>
<td>24.01</td>
<td>0.07</td>
<td>0.3</td>
<td>0.08</td>
</tr>
<tr>
<td>Liquid potassium silicate fertilizer 2</td>
<td>7</td>
<td>16.07</td>
<td>0.03</td>
<td>0.2</td>
<td>0.04</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test  
2) Average (the number of test days ($T$) \times the number of duplicate testing (2))  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Intermediate standard deviation  
7) Intermediate relative standard deviation

**References**


2) Shinji KAWAGUCHI: Verification of Performance Characteristics of Testing Method

5) **Flow sheet for water-soluble silicic acid:** The flow sheet for water-soluble silicic acid in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>5 g analytical sample Weigh to the order of 1 mg to a 500-mL volumetric flask.</td>
</tr>
<tr>
<td>2.</td>
<td>About 400 mL of water</td>
</tr>
<tr>
<td>3.</td>
<td>Rotary shaker (30 - 40 revolutions/min) 30 minutes</td>
</tr>
<tr>
<td>4.</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>5.</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>6.</td>
<td>Filtration</td>
</tr>
<tr>
<td>7.</td>
<td>Sample solution</td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for water-soluble silicic acid in fertilizers (Extraction procedure)

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Sample solution</td>
</tr>
<tr>
<td>2.</td>
<td>Alquot (predetermined amount) 200-mL beaker made of polymer</td>
</tr>
<tr>
<td>3.</td>
<td>About 10 mL hydrochloric acid</td>
</tr>
<tr>
<td>4.</td>
<td>About 15mL potassium fluoride solution</td>
</tr>
<tr>
<td>5.</td>
<td>About 2 g of potassium chloride For 30 minutes or more in a refrigerator</td>
</tr>
<tr>
<td>6.</td>
<td>Cooling</td>
</tr>
<tr>
<td>7.</td>
<td>Filtration under reduced pressure Filter made of polymer Type 6 filter paper</td>
</tr>
<tr>
<td>8.</td>
<td>Washing Wash 6 - 7 times with potassium chloride solution</td>
</tr>
<tr>
<td>9.</td>
<td>Transfer 300-mL tall beaker, water</td>
</tr>
<tr>
<td>10.</td>
<td>Water (until the liquid volume reaches about 200 mL)</td>
</tr>
<tr>
<td>11.</td>
<td>Heating 70 °C - 80 °C</td>
</tr>
<tr>
<td>12.</td>
<td>A few drops of phenolphthalein solution (1 g/100 mL)</td>
</tr>
<tr>
<td>13.</td>
<td>Titration 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (Until the solution becomes light red)</td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for water-soluble silicic acid in fertilizers (Measurement procedure)
4.5 Lime, calcium and alkalinity
4.5.1 Total lime
4.5.1.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers containing organic matters. This testing method is classified as Type C and its symbol is 4.5.1.a-2017 or T-Ca.a-1.

Pretreat an analytical sample with incineration and hydrochloric acid, add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with calcium at a wavelength of 422.7 nm to quantify total lime (T-CaO) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Interference suppressor solution (1): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate (2) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.

c) Calcium standard solution (CaO 1 mg/mL) (1): Heat calcium carbonate specified in JIS K 8617 at 110 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 1.785 g into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add 20 mL of hydrochloric acid (1+3) to dissolve, and add water up to the marked line.

d) Calcium standard solution (CaO 5 µg/mL - 50 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of a calcium standard solution (CaO 1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (3), and add water up to the marked line (4).

e) Blank test solution for the calibration curve preparation (1): Transfer about 50 mL of interference suppressor solution to a 500-mL volumetric flask (3), and add water up to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.

(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

(4) For storage, use a sealable container made of materials which are not likely to dissolve calcium, such as borosilicate glass-1 specified in JIS R 3503 or Teflon.

Comment 1 Instead of the calcium standard solution in (2), a calcium standard solution for the calibration curve preparation can be prepared by using a calcium standard solution (Ca 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate total lime (T-CaO) in the analytical sample by multiplying the concentration (Ca) of the calcium standard solution for calibration curve preparation or a measurement value (Ca) obtained in (4.2) by a conversion factor (1.3992).

(3) Instruments: Instruments are as shown below:

a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121.

1) Light source: A calcium hollow cathode lamp
2) Gas: Gas for heating by flame
(i) Fuel gas: acetylene
(ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Electric furnace**: An electric furnace that can be adjusted to 550 °C ± 5 °C.

c) **Hot plate or sand bath**: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Incineration-hydrochloric acid boiling**

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char.

c) Ignite at 550 °C ± 5 °C for no less than 4 hours to incinerate.

d) After standing to cool, moisten the residue with a small amount of water, gradually add about 10 mL of hydrochloric acid, and further add water to make 20 mL.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to boil for about 5 minutes.

f) After standing to cool, transfer the solution to a 250-mL - 500-mL volumetric flask with water.

g) Add water up to the marked line.

h) Filter with Type 3 filter paper to make a sample solution.

Note (5) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

**Comment 2** The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.1.a.

(4.1.2) **Incineration-aqua regia digestion**

a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char.

c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate.

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass, and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

Note (6) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

(7) The watch glass can be removed.

(8) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric
flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

**Comment 3** The procedures in (4.1.2) are the same as in (4.1.3) in 4.2.1.a. and (4.1) a - h) in 5.3.a.

**4.2** Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 422.7 nm

b) **Calibration curve preparation**
   1) Spray the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 422.7 nm.
   2) Prepare a curve for the relationship between the calcium concentration and the indicated value of the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as CaO) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the calcium content from the calibration curve, and calculate total lime (T-CaO) in the analytical sample.

**Comment 4** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 15 % (mass fraction) and 1 % (mass fraction) are 101.8 % and 97.9% as total lime (T-CaO) respectively. The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.05 % (mass fraction).
Table 1 Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratories used for analysis</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
<th>Reproducibility</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$s_r$</td>
<td>$RSD_r$</td>
<td>$s_{I(T)}$</td>
</tr>
<tr>
<td>FAMIC-C-12</td>
<td>11</td>
<td>5.82</td>
<td>0.07</td>
<td>1.2</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting flame atomic absorbance spectrometry
2) Average (the number of laboratory ($p$) × test days (2) × the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References
(5) **Flow sheet for total lime**: The flow sheet for total lime in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 g analytical sample</td>
<td>Weigh to the order of 1 mg into a 200-mL- 300-mL tall beaker.</td>
</tr>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>Ignite at 550 °C ±5 °C, no less than 4 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← Small amount of water, moisten the residue</td>
<td></td>
</tr>
<tr>
<td>← About 10 mL of hydrochloric acid</td>
<td></td>
</tr>
<tr>
<td>← Water (up to about 20 mL)</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and boil for 5 minutes.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 1-1** Flow sheet for total lime in fertilizers.

*(Incineration-hydrochloric acid boiling procedure (4.1.1))*
Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker.

**Testing Methods for Fertilizers (2018)**

<table>
<thead>
<tr>
<th>5 g analytical sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charring</td>
</tr>
<tr>
<td>Incineration</td>
</tr>
<tr>
<td>Standing to cool</td>
</tr>
<tr>
<td>← A small amount of water</td>
</tr>
<tr>
<td>← About 10 mL of nitric acid</td>
</tr>
<tr>
<td>← About 30 mL of hydrochloric acid</td>
</tr>
<tr>
<td>Heating</td>
</tr>
<tr>
<td>Heating</td>
</tr>
<tr>
<td>Standing to cool</td>
</tr>
<tr>
<td>← 25 mL - 50 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td>Heating</td>
</tr>
<tr>
<td>Standing to cool</td>
</tr>
<tr>
<td>Transfer</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
</tr>
<tr>
<td>Type 3 filter paper</td>
</tr>
</tbody>
</table>

**Figure 1-2** Flow sheet for total lime in fertilizers
(Incineration-aqua regia digestion procedure (4.1.2))

<table>
<thead>
<tr>
<th>Sample solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alquot (predetermined amount)</td>
</tr>
<tr>
<td>← About 10 mL interference suppressor solution</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
</tr>
<tr>
<td>Atomic absorption spectrometer or frame photometer</td>
</tr>
</tbody>
</table>

**Figure 2** Flow sheet for total lime in fertilizers (Measurement)
4.5.2 Soluble lime
4.5.2.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers that guarantee alkalinity. This testing method is classified as Type D and its symbol is 4.5.2.a-2017 or S-Ca.a-1.

Add hydrochloric acid (1+23) to an analytical sample, boil to extract, and add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with calcium at a wavelength of 422.7 nm to quantify hydrochloric acid (1+23) soluble lime (soluble lime (S-CaO)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Interference suppressor solution: Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.

c) Calcium standard solution (CaO 1 mg/mL): Put calcium carbonate specified in JIS K 8617 in a drying apparatus, heat at 110 ºC ± 2 ºC for about 2 hours, let it stand to cool in a desiccator, and weigh 1.785 g into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add 20 mL of hydrochloric acid (1+3) to dissolve, and add water up to the marked line.

d) Calcium standard solution (CaO 5 µg/mL - 50 µg/mL) for the calibration curve preparation: Transfer 2.5 mL - 25 mL of a calcium standard solution (CaO 1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution that is 1/10 volume of the volume to be prepared, and add water up to the marked line.

e) Blank test solution for the calibration curve preparation: Transfer about 50 mL of interference suppressor solution to a 500-mL volumetric flask, and add water up to the marked line.

Note (1) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.

(2) This is an example of preparation; prepare an amount as appropriate.

(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

(4) For storage, use a sealable container made of materials which are not likely to dissolve calcium, such as borosilicate glass-1 specified in JIS R 3503 or Teflon.

Comment 1 Instead of the calcium standard solution in (2), a calcium standard solution for the calibration curve preparation can be prepared by using a calcium standard solution (Ca 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate soluble lime (S-CaO) in the analytical sample by multiplying the concentration (Ca) of the calcium standard solution for calibration curve preparation or a measurement value (Ca) obtained in (4.2) by a conversion factor (1.3992).

(3) Instruments: Instruments are as shown below:

a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121.

1) Light source: A calcium hollow cathode lamp
2) **Gas**: Gas for heating by flame  
   (i) Fuel gas: acetylene  
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.  

**b)** **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.  

(4) **Test procedure**  

(4.1) **Extraction**: Conduct extraction as shown below.  

a) Weigh 2 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL tall beaker.  

b) Add about 20 mL of hydrochloric acid (1+23), cover with a watch glass, and boil on a hot plate for about 5 minutes (5).  

c) After immediate cooling is complete, transfer to a 250-mL - 500-mL volumetric flask with water.  

d) Immediately add water up to the marked line  

e) Filter with Type 3 filter paper to make a sample solution.  

**Note** (5) Be aware that an analytical sample should not solidify in the bottom of a beaker.  

**Comment 2** In the case of a by-product magnesia fertilizer or a fertilizer containing a by-product magnesia, if the pH of the sample solution of d) is neutral or basic, prepare a sample solution anew by replacing “2 g of an analytical sample” in the procedure in a) with “1 g - 1.5 g of an analytical sample”.  

**Comment 3** In the procedure in a), a 500-mL volumetric flask can be used instead of a 500-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. In addition, “cover with a watch glass” in b) is replaced by “place a funnel”, and “transfer to a 250-mL - 500 mL volumetric flask with water” in the procedure in c) is skipped.  

**Comment 4** The procedure in (4.1) is the same as in (4.1) in 4.6.1.a and 4.7.1.a  

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.  

a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:  
   Analytical line wavelength: 422.7 nm  

b) **Calibration curve preparation**  

1) Spray the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 422.7 nm.  

2) Prepare a curve for the relationship between the calcium concentration and the indicated value of the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.  

c) **Sample measurement**  

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as CaO) to a 100-mL volumetric flask.  

2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.  

3) Subject to the same procedure as in b) 1) to read the indicated value.  

4) Obtain the calcium content from the calibration curve, and calculate soluble lime (S-CaO) in the analytical sample.  

**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample. As a
result, the average rate of recovery at the content level of 20 % (mass fraction) and 1 % (mass fraction) are 100.9 % and 101.1% as soluble lime (S-CaO) respectively.

References

(5) Flow sheet for soluble lime: The flow sheet for soluble lime in fertilizers is shown below:

![Flow sheet for soluble lime in fertilizers](image)

Figure 1  Flow sheet for soluble lime in fertilizers (Extraction procedure)

![Flow sheet for soluble lime in fertilizers](image)

Figure 2  Flow sheet for soluble lime in fertilizers (Measurement procedure)
4.5.3 Water-soluble calcium

4.5.3.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers that indicate calcium content as a response modifier. This testing method is classified as Type D and its symbol is 4.5.3.a-2017 or W-Ca,a-1.

Extract by adding water to an analytical sample, and add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with calcium at a wavelength of 422.7 nm to quantify water-soluble calcium (W-Ca) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Interference suppressor solution (1): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate (2) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.
   c) Calcium standard solution (Ca 1 mg/mL): Calcium standard solution (Ca 1 mg/mL) traceable to National Metrology.
   d) Calcium standard solution (Ca 5 µg/mL - 50 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of a calcium standard solution (Ca 1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (3), and add water up to the marked line (4).
   e) Blank test solution for the calibration curve preparation (1): Transfer about 50 mL of interference suppressor solution to a 500-mL volumetric flask (3), and add water up to the marked line (4).

Note  (1) This is an example of preparation; prepare an amount as appropriate.
    (2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
    (3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.
    (4) For storage, use a sealable container made of materials which are not likely to dissolve calcium, such as borosilicate glass-1 specified in JIS R 3503 or Teflon.

Comment 1 Instead of the calcium standard solution in (2), a calcium standard solution for the calibration curve preparation can be prepared by using a calcium standard solution (Ca 10 mg/mL) traceable to National Metrology.

Comment 2 Instead of the calcium standard solution in (2), a calcium standard solution (CaO 5 µg/mL - 50 µg/mL) for the calibration curve preparation prepared in (2) in 4.5.1.a can be used. In this case, calculate water-soluble calcium (W-Ca) in the analytical sample by multiplying the concentration (CaO) of a calcium standard solution for calibration curve preparation or a measurement value (CaO) obtained in (4.2) by a conversion factor (0.7147).

(3) Instruments: Instruments are as shown below:
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121.
1) **Light source**: A calcium hollow cathode lamp
2) **Gas**: Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.
   a) Weigh 1.00 g of an analytical sample, and put it in a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below.
   Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.
   a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
      Analytical line wavelength: 422.7 nm
   b) **Calibration curve preparation**
      1) Spray the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 422.7 nm.
      2) Prepare a curve for the relationship between the calcium concentration and the indicated value of the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
   c) **Sample measurement**
      1) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as Ca) to a 100-mL volumetric flask.
      2) Add about 10 mL of interference suppressor solution \(^{(3)}\), and add water up to the marked line.
      3) Subject to the same procedure as in b) 1) to read the indicated value.
      4) Obtain the calcium content from the calibration curve, and calculate water-soluble calcium (W-Ca) in the analytical sample by the following formula.

**Comment 3** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 1 % (mass fraction) - 5 % (mass fraction) is 98.1 % - 101.1% as water-soluble calcium (W-Ca) respectively.
Additionally, the minimum limit of quantification of this testing method is about 0.07 % (mass fraction) for solid fertilizers, and 0.04 % (mass fraction) for fluid fertilizers.

**References**
(5) **Flow sheet for water soluble calcium:** The flow sheet for water soluble calcium in fertilizers is shown below:

![Flow sheet for water soluble calcium](image)

Figure 1  Flow sheet for water-soluble calcium in fertilizers (Extraction procedure)

![Flow sheet for water soluble calcium](image)

Figure 2  Flow sheet for water-soluble calcium in fertilizers (Measurement procedure)
4.5.3.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.5.3.b-2017 or W-Ca.b-1.

Add water to an analytical sample to extract, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the calcium at a wavelength of 393.366 nm to obtain water-soluble calcium (W-Ca). In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Calcium standard solution (Ca 1 mg/mL): Calcium standard solution (Ca 1 mg/mL) traceable to National Metrology.
   d) Calcium standard solution (Ca 0.1 mg/mL) \(^{(1)}\): Transfer 10 mL of calcium standard solution (Ca 1 mg/mL) to a 100-mL flask and add hydrochloric acid (1+23) up to the marked line.
   e) Calcium standard solutions (Ca 1 µg/mL - 20 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 1 mL - 20 mL of calcium standard solution (Ca 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
   f) Calcium standard solutions (Ca 0.1 µg/mL - 1 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 1 mL - 10 mL of calcium standard solution (Ca 10 µg/mL) for the calibration curve preparation to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
   g) Blank test solution for the calibration curve preparation \(^{(1)}\): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the calcium standard solution in (2), a calcium standard solution for the calibration curve preparation can be prepared by using a calcium standard solution (Ca 10 mg/mL) traceable to National Metrology.

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a calcium standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample \(^{(2)}\) to the order of 1 mg, and put it in a 100-mL volumetric
flask.
b) Add about 50 mL of water, shake to mix and add water up to the marked line.
c) Filter with Type 3 filter paper to make a sample solution.

Note (2) The sampling amount of the analytical sample is 10 g when the content in the analytical sample is less than 0.01 % (mass fraction) as water-soluble calcium.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.
a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
   Analytical line wavelength: 393.366 nm
b) Calibration curve preparation
   1) Spray the calcium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 393.366 nm.
   2) Prepare a curve for the relationship between the calcium concentration and the indicated value of the calcium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Ca) to a 100-mL volumetric flask.
   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the calcium content from the calibration curve, and calculate water-soluble calcium (W-Ca) in the analytical sample by the following formula.

Comment 4 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

Comment 5 The comparison of the measurement value ($y_i$: 0.095 % (mass fraction) - 10.93 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of flame atomic absorbance spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = 0.005 + 0.978x$, and its correlation coefficient ($r$) was 0.999. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery at additive level of 0.01 % (mass fraction) and 0.1 % (mass fraction) were 105.9 % and 106.4 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).
Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>4) s_r</td>
<td>5) RSD_r</td>
</tr>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.02</td>
<td>0.7</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (fluid)</td>
<td>7</td>
<td>0.001</td>
<td>0.9</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \( T \) \( \times \) the number of duplicate testing \( 2 \))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

5) Flow sheet: The flow sheet for water-soluble calcium of the fluid mixed fertilizers is shown below:

```
1 g analytical sample  Weigh to the order of 1 mg to a 100-mL volumetric flask
← Water, about 50 mL
Shaking to mix
← Water (up to the marked line)
Filtration
Sample solution
```

Figure 1  The flow sheet for water-soluble calcium in fluid fertilizers
(Extraction procedure)

```
Sample solution

Aliquot (predetermined volume)  100-mL volumetric flask
← 25 mL of hydrochloric acid (1+5)
← Water (up to the marked line)
Measurement  ICP-OES (393.366 nm)
```

Figure 2  The flow sheet for water-soluble calcium in fluid fertilizers
(Measurement procedure)
4.5.4 Alkalinity
4.5.4.a Ethylenediamine tetraacetate method

(1) Summary
This testing method is applicable to fertilizers that guarantee alkalinity. This testing method is classified as Type E and its symbol is 4.5.4.a-2017 or AL.a-1.

Add hydrochloric acid (1+23) to an analytical sample, boil to extract, and mask with 2,2',2''-nitrilotriethanol and a potassium cyanide solution, add a 0.01 mol/L ethylenediamine tetraacetate standard solution, and conduct a chelatometric titration with a 0.01 mol/L magnesium standard solution to obtain alkalinity (AL). Or after masking, conduct a chelatometric titration with an ethylenediamine tetraacetate standard solution to obtain alkalinity (AL) in an analytical sample.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
b) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.
c) Ascorbic acid: A JIS Guaranteed Reagent specified in JIS K 9502 or a reagent of equivalent quality.
d) 2,2',2"-nitrilotriethanol: A JIS Guaranteed Reagent specified in JIS K 8663 or a reagent of equivalent quality.
e) Acetone: A JIS Guaranteed Reagent specified in JIS K 8034 or a reagent of equivalent quality.
f) Ammonia solution: A JIS Guaranteed (NH₃ 28 % (mass fraction)) reagent specified in JIS K 8085 or a reagent of equivalent quality.
g) 0.01 mol/L ethylenediamine tetraacetate standard solution: Dissolve 3.72 g of ethylenediaminetetraacetic acid dihydrogen disodium dihydrate in water to make 1000 mL.

(a) Standardization: Wash zinc reference material for volumetric analysis specified in JIS K 8005 with hydrochloric acid (1+3), water, ethanol (99.5) specified in JIS K 8101, and diethyl ether specified in JIS K 8103 successively, and immediately leave at rest in a desiccator for about 12 hours under no more than 2 kPa to dry, and then weigh about 0.65 mg to the order of 0.1 mg, transfer it to a 1000-mL volumetric flask, add about 10 mL of hydrochloric acid to dissolve and then add water up to the marked line. Transfer 25 mL of the solution to a 200-mL - 300-mL Erlenmeyer flask, add about 15 mL of water and about 5 mL of ammonium chloride buffer solution, and titrate with a 0.01 mol/L ethylenediamine tetraacetate standard solution, while adding Eriochrome Black T solution as an indicator, until the color of the solution becomes blue. Calculate the factor of a 0.01 mol/L ethylenediamine tetraacetate standard solution by the following formula.

\[
\text{Factor of 0.01 mol/L ethylenediamine tetraacetate standard solution (} f_1 \text{)} = \frac{W_1 \times (A/100) \times (1/65.409) \times (V_1/V_2) \times (1000/V_3) \times (1/C_1)}{W_1 \times A \times (1/65.409) \times (0.25/V_3)}
\]

\[
W_1: \text{ Mass (g) of d zinc sampled}
A: \text{ Purity (% (mass fraction)) of zinc}
V_1: \text{ Volume (25 mL) of zinc solution transferred}
V_2: \text{ Constant volume (1000 mL) of zinc solution}
V_3: \text{ Volume (mL) of 0.01 mol/L ethylenediamine tetraacetate standard solution needed for titration}
C_1: \text{ Set concentration (0.01 mol/L) of 0.01 mol/L ethylenediamine tetraacetate standard solution}
\]
h) **0.01 mol/L magnesium standard solution**: Put 0.24 g of magnesium specified in JIS K 8875 into a 1000-mL beaker, add about 10 mL of hydrochloric acid to dissolve, add a proper amount of water, and while adding a methyl red solution (0.1 g/100 mL) as an indicator, neutralize with an ammonia solution (1+3) until the color of the solution becomes yellow, and then add water to make 1000 mL.

**Standardization**: Transfer 25 mL of 0.01 mol/L magnesium standard solution to a 200-mL - 300-mL Erlenmeyer flask, add about 15 mL of water and about 5 mL of ammonium chloride buffer solution, and while adding an Eriochrome Black T solution as an indicator, titrate with 0.01 mol/L ethylenediamine tetraacetate standard solution until the color of the solution becomes blue. Calculate the factor of a 0.01 mol/L magnesium standard solution by the following formula.

\[
\text{Factor of 0.01 mol/L magnesium standard solution } (f_2) = (C_1 \times f_1 \times V_4) \times (1/V_5) \times (1/C_2)
\]

- **C_1**: Set concentration (0.01 mol/L) of 0.01 mol/L ethylenediamine tetraacetate standard solution
- **C_2**: Set concentration (0.01 mol/L) of 0.01 mol/L magnesium standard solution
- **f_1**: Factor of 0.01 mol/L ethylenediamine tetraacetate standard solution
- **V_4**: Volume (mL) of 0.01 mol/L ethylenediamine tetraacetate standard solution needed for titration
- **V_5**: Volume (mL) of 0.01 mol/L magnesium standard solution transferred.

i) **Ammonium chloride solution**: Dissolve 70 g of ammonium chloride specified in JIS K 8116 and 570 mL of ammonia solution in water to make 1000 mL.

j) **2-aminoethanol solution**: Add 400 mL of water to 150 mL of 2-aminoethanol specified in JIS K 8109, add gradually hydrochloric acid to a pH 10.6.

k) **Potassium cyanide solution**: Dissolve 100 g of potassium cyanide specified in JIS K 8443 in water to make 1000 mL.

l) **Eriochrome Black T solution**: Dissolve 0.5 g of Eriochrome Black T specified in JIS K 8736 and 4.5 g of hydroxyl-ammonium chloride in methanol-water (95+5) to make 100 mL.

m) **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.10 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

n) **Methanol**: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.

o) **Citric acid solution (2)**: Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.

**Note** (1) The reagent corresponds to triethanolamine in the Official Methods of Analysis of Fertilizers (1992).

(2) This is an example of preparation; prepare an amount as appropriate.

**Comment 1** Instead of a 0.01 mol/L ethylenediamine tetraacetate standard solution in (2) g, a 0.1 mol/L ethylenediaminetetraacetic acid dihydrogen disodium solution conforming to ISO/IEC 17025 can be used.

(3) **Instruments**: Instruments are as shown below:

a) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.
(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 2 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL tall beaker.
b) Add about 200 mL of hydrochloric acid (1+23), cover with a watch glass, and boil on a hot
   plate for about 5 minutes (3).
c) Immediately transfer to a 250-mL - 500-mL volumetric flask with water.
d) Immediately add water up to the marked line (4).
e) Filter with Type 3 filter paper to make a sample solution.

Note (3) Be aware that an analytical sample should not solidify in the bottom of a beaker.
(4) Conduct the procedure in Comment 5 when there is much manganese content in the
fertilizers.

Comment 2 In the case of a by-product magnesia fertilizer, if the pH of the sample solution of d) is
neutral or basic, prepare a sample solution anew by replacing “2 g of an analytical
sample” in the procedure in a) with “1 g - 1.5 g of an analytical sample”.

Comment 3 In the procedure in a), a 500-mL volumetric flask can be used instead of a 500-mL
tall beaker. However, the volumetric flask used should be distinguished as an
extraction flask and should not be used for the other purposes. In addition, “cover
with a watch glass” in b) is replaced by “place a funnel”, and “transfer to a 250-mL -
500 mL volumetric flask with water” in the procedure in c) is skipped.

Comment 4 The procedure in (4.1) is the same as the procedure in (4.1) of 4.5.2.a.

Comment 5 Add a predetermined volume of filtrate in (4.1) e), add a drop of methyl red solution
as an indicator and drop ammonia water (28 % (mass fraction)) specified in JIS K
8085 until the color of the solution changes light yellowish red from purplish red.
Add 20 mL of ammonium peroxodisulfate to boil (5). Immediately transfer it to a
100-mL - 200-mL volumetric flask with water. After immediate cooling is complete,
add water up to the marked line. Filter with Type 3 filter paper to make a sample
solution.

Note (5) Precipitation of manganese oxide, etc. is formed.

(4.2) Measurement: Conduct measurement as shown below. Two examples of titration are shown
as follows.
(4.2.1) Measurement (A): Titration with a magnesium standard solution (0.01 mol/L)
a) Transfer a predetermined volume (the equivalents of 5 mg - 20 mg as CaO + MgO) of sample
solution to a 200-mL - 300-mL Erlenmeyer flask.
b) Add a proper amount of water, add a drop of methyl red solution as an indicator and drop sodium hydroxide (5 g/100 mL) to neutralize until the color of the solution becomes yellow.
c) Add 0.1 g of ascorbic acid, 1 ml - 10 mL of 2,2’,2’”-nitrilotriethanol–water (1+3) and 1 mL -
10 mL of potassium cyanide solution (6).
d) Add a predetermined volume of 0.01 mol/L ethylenediamine tetraacetate standard solution (7).
e) Add 20 mL of ammonium chloride solution or 2-aminoethanol solution.
f) Add several drops of Eriochrome Black T solution, and titrate with a 0.01 mol/L magnesium
standard solution until the color of the solution becomes red.
g) Calculate the alkalinity (AL) content in an analytical sample by the following formula.

Alkalinity (AL) (% (mass fraction)) in an analytical sample

\[ \frac{\left( C_1 \times f_1 \times V_6 / 1000 \right) - \left( C_2 \times f_2 \times V_7 / 1000 \right) \times (56.077/W_2) \times (V_8/V_9) \times 100}{W} \]
\[ ((f_1 \times V_6) - (f_2 \times V_7)) \times (56.077/W_2) \times (V_2/V_4) \times (1/1000) \]

\[ C_1: \text{Set concentration (0.01 mol/L) of 0.01 mol/L ethylenediamine tetraacetate standard solution} \]

\[ C_2: \text{Set concentration (0.01 mol/L) of 0.01 mol/L magnesium standard solution} \]

\[ f_1: \text{Factor of 0.01 mol/L ethylenediamine tetraacetate standard solution} \]

\[ f_2: \text{Factor of 0.01 mol/L magnesium standard solution} \]

\[ V_6: \text{Additive volume (mL) of 0.01 mol/L ethylenediamine tetraacetate standard solution} \]

\[ V_7: \text{Volume (mL) of 0.01 mol/L magnesium standard solution needed for titration} \]

\[ V_8: \text{Predetermined volume (mL) of the sample solution in (4.1) d)} \]

\[ V_9: \text{Transferred volume (mL) of the sample solution subjected to titration in (4.2.1) a).} \]

\[ W_2: \text{Mass (g) of the analytical sample} \]

**Note (6)** If manganese is present, replace “1 mL - 10 mL of potassium cyanide solution” with “1g - 5g of potassium cyanide”.

**Note (7)** Add excess volume since 1.8 mL of ethylenediamine tetraacetate standard solution (0.01 mol/L) is required for 1 mg of CaO.

(4.2.2) **Measurement (B):** Titration with an ethylenediamine tetraacetate standard solution (0.01 mol/L)

a) Transfer a predetermined volume (the equivalents of 5 mg - 20 mg as CaO+MgO) of sample solution to a 200-mL - 300-mL Erlenmeyer flask.

b) Add a proper amount of water and 5 mL of citric acid solution \(^{(8)}\), add a drop of methyl red solution as an indicator and drop sodium hydroxide (5 g/100 mL) to neutralize until the color of the solution becomes yellow.

c) Add 0.1 g of ascorbic acid, 1 ml - 10 mL of 2,2',2"-nitrilotriethanol−water (1+3) and 1 mL - 10 mL of potassium cyanide solution \(^{(6)}\).

d) Add 20 mL of ammonium chloride solution or 2-aminoethanol solution.

e) Add several drops of an Eriochrome Black T solution, and immediately titrate with a 0.01 mol/L ethylenediamine tetraacetate standard solution until the color of the solution becomes blue-green.

f) Calculate the alkalinity (AL) content in an analytical sample by the following formula.

Alkalinity (AL) (% (mass fraction)) in an analytical sample

\[ = (C_1 \times f_1 \times V_{10}/1000) \times (56.077/W_3) \times (V_{11}/V_{12}) \times 100 \]

\[ = (f_1 \times V_{10}) \times (56.077/W_3) \times (V_{11}/V_{12}) \times (1/1000) \]

\[ C_1: \text{Set concentration (0.01 mol/L) of 0.01 mol/L ethylenediamine tetraacetate standard solution} \]

\[ f_1: \text{Factor of 0.01 mol/L ethylenediamine tetraacetate standard solution} \]

\[ V_{10}: \text{Volume (mL) of 0.01 mol/L ethylenediamine tetraacetate standard solution needed for titration} \]

\[ V_{11}: \text{Predetermined volume (mL) of the sample solution in (4.1) d)} \]

\[ V_{12}: \text{Transferred volume (mL) of the sample solution subjected to titration in (4.2.2) a).} \]

\[ W_3: \text{Mass (g) of the analytical sample} \]

**Note (8)** When the sample solution does not contain phosphate, silicate, etc., it is not necessary
Care should be fully taken in the case of using potassium cyanide and its solution in accordance with the Safety Data Sheet (SDS). In addition, observe laws and ordinances concerned such as the Poisonous and Deleterious Substance Control Law. Criteria of the abolition in the Poisonous and Deleterious Substance Control Law (for reference): Add a sodium hydroxide solution to make it alkalinity more than pH 11, and further add an oxidizer (sodium hypochlorite, bleaching powder) solution to conduct oxidative degradation processes. After dissolving CN ingredient, neutralize with sulfuric acid, and discard it after diluting it with a large amount of water. Take enough time to dissolve the CN ingredient with alkalinity.

References

(5) Flow sheet for alkalinity: The flow sheet for alkalinity in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 g analytical sample</td>
<td>Weigh to the order of 1 mg to a 500-mL tall beaker</td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass and boil for 5 minutes</td>
</tr>
<tr>
<td>Cooling</td>
<td>Immediately</td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL - 500-mL volumetric flask</td>
</tr>
<tr>
<td>Filtration</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Sample solution</td>
<td>Type 3 filter paper</td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for alkalinity in fertilizers (Extraction procedure)

Figure 2-1  Flow sheet for alkalinity in fertilizers (Measurement procedure (4.2.1) (A))

Sample solution

Aliquot (predetermined amount)
200-mL - 300-mL Erlenmeyer Flask

← Water (proper amount)
← A drop of methyl-red solution

Neutralization
Sodium hydroxide (5 g/100 mL)
(until the solution becomes yellow )
← 0.1 g ascorbic acid
← 1 mL - 10 mL of 2,2’,2”-nitrilotriethanol - water (1+3)
← 1 mL - 10 mL potassium cyanide solution
← Predetermined volume of 0.01 mol/L ethylenediamine tetraacetate standard solution
← 20 mL of ammonium chloride solution or 2-aminoethanol solution
← Several drops of Erio-chrome Black T solution

Titration
0.01 mol/L magnesium standard solution
(until the solution becomes red)

Figure 2-2  Flow sheet for alkalinity in fertilizers (Measurement procedure (4.2.2) (B))

Sample solution

Aliquot (predetermined amount)
200-mL - 300-mL Erlenmeyer Flask

← Water (proper amount)
← 5 mL citric acid solution
← A drop of methyl-red solution

Neutralization
Sodium hydroxide (5 g/100 mL)
(until the solution becomes yellow )
← 0.1 g of ascorbic acid
← 1 mL - 10 mL of 2,2’,2”-nitrilotriethanol - water (1+3)
← 1 mL - 10 mL of potassium cyanide solution
← Predetermined volume of 0.01 mol/L ethylenediamine tetraacetate standard solution
← 20 mL of ammonium chloride solution or 2-aminoethanol solution
← Several drops of Erio-chrome Black T solution

Titration
0.01 mol/L ethylenediamine tetraacetate standard solution
(until the solution becomes blue-green)
4.5.4.b Calculation with soluble lime and soluble magnesia

(1) Summary
This testing method is applicable to fertilizers that guarantee alkalinity (AL). This testing method is
classified as Type A (Def-C) and its symbol is 4.5.4.b-2017 or AL.b-1.
Multiply the soluble magnesia (S-MgO) obtained in 4.6.2 by the factor (1.3934) and add to the
soluble lime (S-CaO) obtained in 4.5.2 to calculate alkalinity (AL) in an analytical sample.

(2) Calculation of alkalinity
a) Calculate the alkalinity (AL) in a teat sample by the following formula.

\[
\text{Alkalinity (AL) (% (mass fraction)) in an analytical sample} = (S\text{-CaO}) + 1.3914 \times (S\text{-MgO})
\]

S-CaO: Soluble lime (% (mass fraction)) \(^{(1)}\) obtained in 4.5.2 in an analytical sample
S-MgO: Soluble magnesia (% (mass fraction)) \(^{(1)}\) obtained in 4.6.2 in an analytical sample

Note  (1) S-CaO and S-MgO use raw data without rounding numerical value
4.6 Magnesia
4.6.1 Total magnesia
4.6.1.a Flame atomic absorption spectrometry

(1) Summary
The testing method is applicable to compost, sludge fertilizers (except for calcined sludge fertilizer) and poultry manure ash. This testing method is classified as Type D and its symbol is 4.6.1.a-2018 or T-Mg.a-1.

Pretreat an analytical sample with incineration and hydrochloric acid or nitric acid−hydrochloric acid (3+1), add an interference suppressor solution, and then spray in an acetylene−air flame, and measure the atomic absorption with magnesium at a wavelength of 285.2 nm to quantify total magnesia (T-MgO) in an analytical sample.

In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A JIS Guaranteed Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   c) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   d) Interference suppressor solution (1): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate (2) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.
   e) Magnesium standard solution (MgO 1 mg/mL) (1): Weigh 0.603 g of magnesium (powder) specified in JIS K 8876 into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.
   f) Magnesium standard solution (MgO 0.1 mg/mL) (1): Transfer 10 mL of magnesium standard solution (MgO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   g) Magnesium standard solution (MgO 1 µg/mL-10 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL-25 mL of magnesium standard solution (MgO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution (3), and add water up to the marked line.
   h) Blank test solution for the calibration curve preparation (1): Transfer about 25 mL of interference suppressor solution to a 250-mL volumetric flask (3), and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
   (3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the magnesium standard solution in (2), a magnesium standard solution for the calibration curve preparation can be prepared by using a magnesium standard solution (Mg 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate total magnesia (T-MgO) in the analytical sample by multiplying the concentration (Mg) of the magnesium standard solution for calibration curve preparation or a measurement value (Mg) obtained in (4.2) by a
conversion factor (1.6583).

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121.
   b) Light source: A magnesium hollow cathode lamp
   c) Gas: Gas for heating by flame
      i) Fuel gas: acetylene
      ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   d) Electric furnace: An electric furnace that can be adjusted to 450 °C ± 5 °C or 550 °C ± 5 °C.
   e) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   (4.1.1) Incineration-hydrochloric acid boiling
      a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.
      b) Put the tall beaker in an electric furnace, and heat gently to char (4).
      c) Ignite at 550 °C ± 5 °C for no less than 4 hours to incinerate (4).
      d) After standing to cool, moisten the residue with a small amount of water, gradually add about 10 mL of hydrochloric acid, and further add water to make 20 mL.
      e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to boil for about 5 minutes.
      f) After standing to cool, transfer the solution to a 250-mL - 500-mL volumetric flask with water.
      g) Add water up to the marked line.
      h) Filter with Type 3 filter paper to make a sample solution.

   Note (4) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

   Comment 2 The procedure in (4.1.1) is the same as the procedure in (4.1.2) in 4.2.1.a.

   (4.1.2) Incineration-aqua regia digestion
      a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 200-mL - 300-mL tall beaker.
      b) Put the tall beaker in an electric furnace, and heat gently to char (5).
      c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (5).
      d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
      e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
      f) Slightly move the watch glass (6), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
      g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (7) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
      h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
Note  (5) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

(6) The watch glass can be removed.

(7) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 3 The procedures in (4.1.2) are the same as in (4.1.3) in 4.2.1.a and (4.1) a - h) in 5.3.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 285.2 nm

b) Calibration curve preparation
   1) Spray the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 285.2 nm.
   2) Prepare a curve for the relationship between the magnesium concentration and the indicated value of the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MgO) (8) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the magnesium content from the calibration curve, and calculate the total magnesia (T-MgO) in the analytical sample.

Note  (8) If there is a possibility that the concentration of total magnesia in the sample solution exceeds the maximum limit of the calibration curve, transfer a predetermined amount of the sample solution and dilute by adding water up to the marked line to make 0.1 mg - 1 mg as MgO.

Comment 4 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 5 % (mass fraction), 1 % (mass fraction) and 0.2 % (mass fraction) are 102.4 %, 101.7 % and 103.0 % as total magnesia (T-MgO) respectively. The results of the repeatability tests on different days using swine manure compost, composted sludge fertilizer and poultry manure ash (1 sample for each) to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 0.2 % (mass fraction).
The flow sheet for total magnesia in fertilizers is shown below:

Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T_{(1)}$</td>
<td>$s_{(1)}^{(4)}$</td>
<td>$RSD_{(1)}^{(5)}$</td>
</tr>
<tr>
<td></td>
<td>$T_{(2)}$</td>
<td>$s_{(2)}^{(4)}$</td>
<td>$RSD_{(2)}^{(5)}$</td>
</tr>
<tr>
<td>Swine manure compost</td>
<td>5</td>
<td>3.14</td>
<td>0.03</td>
</tr>
<tr>
<td>Composted sludge fertilizer</td>
<td>5</td>
<td>0.84</td>
<td>0.01</td>
</tr>
<tr>
<td>Poultry manure ash</td>
<td>5</td>
<td>3.97</td>
<td>0.03</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test  
2) Average (the number of test days $T$)  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Intermediate standard deviation  
7) Intermediate relative standard deviation

(5) Flow sheet for total magnesia: The flow sheet for total magnesia in fertilizers is shown below:

5 g analytical sample  Weigh to the order of 1 mg into a 200-mL-300-mL tall beaker.

Charring  Heat gently

Incineration  Ignite at 550 ºC ±5 ºC, no less than 4 hours

Standing to cool  Room temperature  
← Small amount of water, moisten the residue  
← About 10 mL of hydrochloric acid  
← Water (up to about 20 mL)

Heating  Cover with a watch glass, and boil for 5 minutes.

Standing to cool  Room temperature

Transfer  250-mL - 500-mL volumetric flask, water  
← Water (up to the marked line)

Filtration  Type 3 filter paper

Sample solution

Figure 1  Flow sheet for total magnesia in fertilizers.
(Incineration-hydrochloric acid boiling procedure (4.1.1))
Weigh to the order of 1 mg into a 200-mL - 300-mL tall beaker.

Charring
Ignite at 450 °C ± 5 °C for 8 - 16 hours

Incineration

Standing to cool
Room temperature
A small amount of water
About 10 mL of nitric acid
About 30 mL of hydrochloric acid

Heating
Cover with a watch glass to digest

Heating
Slightly move the watch glass and remove acid

Standing to cool
Room temperature
25 mL - 50 mL of hydrochloric acid (1+5)

Heating
Cover with a watch glass to dissolve

Standing to cool
Room temperature

Transfer
100-mL - 200-mL volumetric flask, water

Filtration
Type 3 filter paper

Sample solution

Figure 2 Flow sheet for total magnesia in fertilizers
(Incineration-aqua regia digestion procedure (4.1.2))

Sample solution

Aliquot
(predetermined amount)
100-mL volumetric flask

About 10 mL interference suppressor solution
Water (up to the marked line)

Measurement
Atomic absorption spectrometer

Figure 3 Flow sheet for total magnesia in fertilizers (Measurement procedure)
4.6.2 Soluble magnesia
4.6.2.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers containing by-product magnesia fertilizers and fertilizers that guarantee alkalinity. This testing method is classified as Type D and its symbol is 4.6.2.a-2018 or S-Mg.a-1.

Add hydrochloric acid (1+23) to an analytical sample, boil to extract and add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with magnesium at a wavelength of 285.2 nm to obtain hydrochloric acid (1+23) soluble magnesia (soluble magnesia (S-MgO)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Interference suppressor solution: Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.

c) Magnesium standard solution (MgO 1 mg/mL): Weigh 0.603 g of magnesium (powder) specified in JIS K 8876 into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.

d) Magnesium standard solution (MgO 0.1 mg/mL): Transfer 10 mL of magnesium standard solution (MgO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.

e) Magnesium standard solution (MgO 0.1 µg/mL-10 µg/mL) for the calibration curve preparation: Transfer 2.5 mL-25 mL of magnesium standard solution (MgO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution, and add water up to the marked line.

f) Blank test solution for the calibration curve preparation: Transfer about 25 mL of interference suppressor solution used in the procedure e) to a 250-mL volumetric flask, and add water up to the marked line.

Note
(1) This is an example of preparation; prepare an amount as appropriate.
(2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the magnesium standard solution in (2), a magnesium standard solution for the calibration curve preparation can be prepared by using a magnesium standard solution (Mg 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate soluble magnesia (S-MgO) in the analytical sample by multiplying the concentration (Mg) of a magnesium standard solution for calibration curve preparation or a measurement value (Mg) obtained in (4.2) by a conversion factor (1.6583).

(3) Instruments: Instruments are as shown below:

a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in
JIS K 0121.

1) **Light source**: A magnesium hollow cathode lamp

2) **Gas**: Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 2 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL tall beaker.

b) Add about 200 mL of hydrochloric acid (1+23), cover with a watch glass, and boil on a hot plate for about 5 minutes (4).

c) Immediately transfer to a 250-mL - 500-mL volumetric flask with water.

d) Immediately add water up to the marked line

e) Filter with Type 3 filter paper to make a sample solution.

*Note* (4) Be aware that an analytical sample should not solidify in the bottom of a beaker.

**Comment 2** In the case of a by-product magnesia fertilizer or a fertilizer containing a by-product magnesia, if the pH of the sample solution of d) is neutral or basic, prepare a sample solution anew by replacing “2 g of an analytical sample” in the procedure in a) with “1 g - 1.5 g of an analytical sample”.

**Comment 3** In the procedure in a), a 500-mL volumetric flask can be used instead of a 500-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. In addition, “cover with a watch glass” in b) is replaced by “place a funnel”, and “transfer to a 250-mL - 500 mL volumetric flask with water” in the procedure in c) is skipped.

**Comment 4** The procedure in (4.1) is the same as the procedure in (4.1) of 4.5.2.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 285.2 nm

b) **Calibration curve preparation**

1) Spray the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 285.2 nm.

2) Prepare a curve for the relationship between the magnesium concentration and the indicated value of the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MgO) to a 100-mL volumetric flask.

2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Obtain the magnesium content from the calibration curve, and calculate the soluble magnesia (S-MgO) in the analytical sample.
**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 15 % (mass fraction) and 1 % (mass fraction) are 101.7 % and 99.5% as soluble magnesia (S-MgO) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.2 % (mass fraction) for solid fertilizers, and 0.05 % (mass fraction) for fluid fertilizers.

**References**

(5) **Flow sheet for soluble magnesia**: The flow sheet for soluble magnesia in fertilizers is shown below:

![Flow sheet for soluble magnesia](image_url)

Figure 1  Flow sheet for soluble magnesia in fertilizers (Extraction procedure)

![Flow sheet for soluble magnesia](image_url)

Figure 2  Flow sheet for soluble magnesia in fertilizers (Measurement procedure)
4.6.3 Citrate-soluble magnesia
4.6.3.a Flame atomic absorption spectrometry

(1) Summary
This test method is applicable to fertilizers containing magnesia hydroxide fertilizers, etc. This testing method is classified as Type C and its symbol is 4.6.3.a-2018 or C-Mg.a-2.

Extract by adding a citric acid solution to an analytical sample and add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with magnesium at a wavelength of 285.2 nm to obtain citrate-soluble magnesia (C-MgO) in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Citric acid solution (1): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.

c) Interference suppressor solution (1): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate (2) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.

d) Magnesium standard solution (MgO 1 mg/mL) (1): Weigh 0.603 g of magnesium (powder) specified in JIS K 8876 into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.

e) Magnesium standard solution (MgO 0.1 mg/mL) (1): Transfer 10 mL of magnesium standard solution (MgO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.

f) Magnesium standard solution (MgO 1 µg/mL-10 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL-25 mL of magnesium standard solution (MgO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution (3), and add water up to the marked line.

g) Blank test solution for the calibration curve preparation (1): Transfer about 25 mL of interference suppressor solution used in the procedure f) to a 250-mL volumetric flask (3), and add water up to the marked line.

Note
(1) This is an example of preparation; prepare an amount as appropriate.
(2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the magnesium standard solution in (2), a magnesium standard solution for the calibration curve preparation can be prepared by using a magnesium standard solution (Mg 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble magnesia (C-MgO) in the analytical sample by multiplying the concentration (Mg) of a magnesium standard solution for calibration curve preparation or a measurement value (Mg) obtained in (4.2) by a conversion factor (1.6583).

(3) Instruments: Instruments are as shown below:

a) Extractor: Constant-temperature rotary shaker or reciprocating water bath shaker as
described below.

aa) **Constant-temperature rotary shaker:** A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

ab) **Reciprocating water bath shaker:** A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/minute with amplitude of 25 mm - 40 mm.

b) **Flame atomic absorption spectrometer:** An atomic absorption spectrometer specified in JIS K 0121.

1) **Light source:** A magnesium hollow cathode lamp
2) **Gas:** Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker:**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL tall beaker.
   b) Add 150 mL of citric acid solution heated up to about 30 °C (4), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.
   c) After immediate cooling is complete, add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

   **Note** (4) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

   **Comment 2** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker:**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (5).
   b) Add 150 mL of citric acid solution heated up to about 30 °C (4), and shake to mix horizontally at amplitude of 25 mm - 40 mm (30 °C ± 1 °C) at 160 times/minute for 1 hour.
   c) After immediate cooling is complete, add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

   **Note** (5) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

   **Comment 3** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

   **Comment 4** For a by-product magnesia fertilizer, if the pH of the sample solution in (4.1.1) d) and (4.1.2) d) is neutral or basic, prepare a sample solution anew by replacing “1 g of an analytical sample” in the procedures in (4.1.1) a) and (4.1.2) a) with “0.5 g of an analytical sample”.

   **Comment 5** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

   **Comment 6** In some slag silicate fertilizers, the variation of measurement value of citrate magnesia (C-MgO) may be observed according to the time variation of heating state after adding a citric acid solution. Therefore, in the case of slag silicate fertilizers, it
is necessary to conduct the procedures of (4.1.1) c) - d) and (4.1.2) c) - d) as quickly as possible after confirming the time of shaking to mix in the procedure in (4.1.1) b) and (4.1.2) b).

**Comment 7** In the case of a fertilizer containing Kieserite (magnesium sulfate fertilizers), wash the non-dissolved matters with water obtained while preparing the sample solution of water-soluble magnesia in (4.1) in 4.6.4.a to transfer it to a 250 mL volumetric flask, and then prepare a sample solution by the procedures in (4.1.1) b) - d) and (4.1.2) b) - d). Mix the magnesia obtained in (4.2) regarding this sample solution with the water-soluble magnesia in 4.6.4.a regarding the said fertilizers to make citrate-soluble magnesia.

**4.2 Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 285.2 nm

b) **Calibration curve preparation**
   1) Spray the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 285.2 nm.
   2) Prepare a curve for the relationship between the magnesium concentration and the indicated value of the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MgO) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the magnesium content from the calibration curve, and calculate the citrate soluble magnesia (C-MgO) in the analytical sample.

**Comment 8** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 1 % (mass fraction) - 5 % (mass fraction) are 98.9 % - 100.3 % as citrate-soluble magnesia (C-MgO) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this test method is about 0.06 % (mass fraction) for solid fertilizers.

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Table 1  Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratories used for analysis</th>
<th>Average (± standard deviation)</th>
<th>Intermediate precision (± standard deviation)</th>
<th>Reproducibility (± standard deviation)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>%</td>
<td>%</td>
<td>%</td>
</tr>
<tr>
<td>FAMIC-A-10</td>
<td>11</td>
<td>3.28 (0.07)</td>
<td>2.0 (0.08)</td>
<td>0.11 (0.11)</td>
</tr>
<tr>
<td>FAMIC-A-13</td>
<td>9</td>
<td>3.18 (0.03)</td>
<td>1.0 (0.04)</td>
<td>0.12 (0.12)</td>
</tr>
</tbody>
</table>

1) The number of laboratoties used for analysis
2) Average (the number of laboratory (p) × test days (2) × the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

(5) Flow sheet for citrate-soluble magnesia: The flow sheet for citrate-soluble magnesia in fertilizers is shown below:

```
1 g analytical sample

← About 150 mL of citrate solution [about 30 ºC]

Shaking to mix

Rotary shaker (30 - 40 revolutions/min),
30 ºC ± 1 ºC, for 1 hour

Cooling

Immediately

← Water (up to the marked line)

Filtration

Type 3 filter paper

Sample solution
```

Figure 1-1   Flow sheet for citrate-soluble magnesia in fertilizers
(Extraction procedure (4.1.1))
1 g analytical sample

Weigh to the order of 1 mg into a 250-mL tall beaker.

About 150 mL of citrate solution [about 30 °C]

Reciprocating water bath shaker (reciprocation horizontally at 160 times /min, with amplitude of 25 mm - 40 mm), at 30 °C ± 1 °C, for 1 hour

Shaking to mix

Immediately

Water (up to the marked line)

Filtration

Type 3 filter paper

Sample solution

Figure 1-2  Flow sheet for citrate-soluble magnesia in fertilizers
(Extraction procedure (4.1.2))

100-mL volumetric flask

Aliquot (predetermined amount)

About 10 mL of interference suppressor solution

Water (up to the marked line)

Measurement

Atomic absorption spectrometer (285.2 nm)

Figure 2  Flow sheet for citrate-soluble magnesia in fertilizers
(Measurement procedure)
4.6.3.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 4.6.3.b-2018 or C-Mg.b-1.

Extract by adding a citric acid solution to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the magnesium at a wavelength of 279.553 nm to obtain citric acid-soluble magnesia (citrate-soluble magnesia (C-MgO)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 9.

(2) Reagent: Reagents are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Citric acid solution (1): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   d) Magnesium standard solution (MgO 1 mg/mL) (1): Weigh 0.603 g of magnesium (powder) specified in JIS K 8876 into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.
   e) Magnesium standard solution (MgO 0.1 mg/mL): Transfer 10 mL of magnesium standard solution (MgO 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   f) Magnesium standard solution (MgO 2 µg/mL - 10 µg/mL) for the calibration curve preparation (1): Transfer 2 mL-16 mL of magnesium standard solution (MgO 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
   g) Magnesium standard solution (MgO 0.2 µg/mL - 2 µg/mL) for the calibration curve preparation (1): Transfer 2 mL - 20 mL of magnesium standard solution (MgO 10 µg/mL) for the calibration curve preparation to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
   h) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in e) - g).

Note  (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the magnesium standard solution in (2), a magnesium standard solution for the calibration curve preparation can be prepared by using a magnesium standard solution (Mg 1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble magnesia (C-MgO) in the analytical sample by multiplying the concentration (Mg) of a magnesium standard solution for calibration curve preparation or a measurement value (Mg) obtained in (4.2) by a conversion factor (1.6583).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in f) and g) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a magnesium standard solution for the calibration curve in the concentration range which is suitable to a
(3) **Instruments**: Instruments are as shown below:

a) **ICP Optical Emission Spectrometry**: A spectrophotometer specified in JIS K 0115

b) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

c) **Extractor**: Constant-temperature rotary shaker or reciprocating water bath shaker as described below.

   ba) **Constant-temperature rotary shaker**: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

   bb) **Reciprocating water bath shaker**: A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/ minute with amplitude of 25 mm - 40 mm.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL tall beaker.

b) Add 150 mL of citric acid solution heated up to about 30 °C (2), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 2** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (3).

b) Add 150 mL of citric acid solution heated up to about 30 °C (2), and shake to mix by reciprocating at amplitude of 25 mm - 40 mm (30 °C ± 1 °C) at 160 times/min for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (3) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

**Comment 3** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

**Comment 4** For a by-product magnesia fertilizer, if the pH of the sample solution in (4.1.1) d) and (4.1.2) d) is neutral or basic, prepare a sample solution anew by replacing “1 g of an analytical sample” in the procedures in (4.1.1) a) and (4.1.2) a) with “0.5 g of an analytical sample”.

**Comment 5** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

**Comment 6** In some slag silicate fertilizers, the variation of measurement value of citrate magnesia (C-MgO) may be observed according to the time variation of heating state after adding a citric acid solution. Therefore, in the case of slag silicate fertilizers, it
is necessary to conduct the procedures of (4.1.1) c - d and (4.1.2) c - d as quickly as possible after confirming the time of shaking to mix in the procedure in (4.1.1) b) and (4.1.2) b).

Comment 7 In the case of a fertilizer containing Kieserite (magnesium sulfate fertilizers), wash the non-dissolved matters with water obtained while preparing the sample solution of water-soluble magnesia in (4.1) in 4.6.4.a to transfer it to a 250 mL volumetric flask, and then prepare a sample solution by the procedures in (4.1.1) b) - d) and (4.1.2) b) - d). Mix the magnesia obtained in (4.2) regarding this sample solution with the water-soluble magnesia in 4.6.4.a regarding the said fertilizers to make citrate-soluble magnesia.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
   Analytical line wavelength: 279.553 nm

b) Calibration curve preparation
   1) Spray the magnesium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 279.553 nm.
   2) Prepare a curve for the relationship between the magnesium concentration and the indicated value of the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.02 mg - 1.6 mg as MgO) to a 100-mL volumetric flask.
   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the magnesium content from the calibration curve, and calculate the citrate soluble magnesia (C-MgO) in the analytical sample.

Comment 8 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.3.d Comment 7.

Comment 9 The comparison of the measurement value (yi: 1.59 % (mass fraction) - 15.06 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value (xi) of Flame atomic absorption spectrometry was conducted to evaluate trueness using processed phosphorus fertilizers (2 samples), compound fertilizers (11 samples), slag silicate fertilizer (1 sample), mixed compost fertilizer (1 sample), mixed phosphate fertilizers (2 samples), designated blended fertilizer (1 sample), blended fertilizers (5 samples), byproduct mixed fertilizer (1 sample), organic compound fertilizer (1 sample), and fused phosphate fertilizer (1 sample). As a result, a regression equation was \( y = 0.0271 + 1.0124x \), and its correlation coefficient \( r \) was 0.999. In addition, recovery testing was conducted using a preparation sample. As a result, the average rate of recovery at the content level of 0.232 % (mass fraction) - 18.81 % (mass fraction) was 94.9 % - 102.7 %.

The results of the repeatability tests on different days using compound fertilizers and blended fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.
Additionally, the minimum limit of quantification of this testing method is about 0.03 % (mass fraction).

Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Test days of average (T)</td>
<td>s_r (4) (%)</td>
<td>RSD_r (5) (%)</td>
</tr>
<tr>
<td>Blended fertilizer</td>
<td>7</td>
<td>0.09</td>
<td>1.1</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>0.03</td>
<td>1.6</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) × the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
Flow sheet for citrate-soluble magnesia: The flow sheet for citrate-soluble magnesia in fertilizers is shown below:

1 g analytical sample

Weigh to the order of 1 mg into a 250-mL tall beaker.

About 150 mL of citrate solution [about 30 °C]

Shaking to mix

Rotary shaker (30 - 40 revolutions/min),
30 °C ± 1 °C, for 1 hour

Cooling

Immediately

Water (up to the marked line)

Type 3 filter paper

Sample solution

Figure 1-1  Flow sheet for citrate-soluble magnesia in fertilizers (Extraction procedure (4.1.1))

1 g analytical sample

Weigh to the order of 1 mg into a 250-mL tall beaker.

About 150 mL of citrate solution [about 30 °C]

Shaking to mix

Reciprocating water bath shaker (reciprocation horizontally at 160 times /min, with amplitude of 25 mm - 40 mm), at 30 °C ± 1 °C, for 1 hour

Cooling

Immediately

Water (up to the marked line)

Type 3 filter paper

Sample solution

Figure 1-2  Flow sheet for citrate-soluble magnesia in fertilizers (Extraction procedure (4.1.2))

Sample solution

100-mL volumetric flask

Aliquot

(predetermined amount)

About 10 mL of interference suppressor solution

Water (up to the marked line)

Measurement

ICP-OES (279.553 nm)

Figure 2  Flow sheet for citrate-soluble magnesia in fertilizers (Measurement procedure)
4.6.4 Water-soluble magnesia
4.6.4.a Flame atomic absorption spectrometry

(1) Summary
This test method is applicable to fertilizers containing magnesia sulfate fertilizers, etc. This testing method is classified as Type D and its symbol is 4.6.4.a-2018 or W-Mg.a-1.

Add water to an analytical sample and boil to extract. Add an interference suppressor solution, then spray in an acetylene–air flame and measure the atomic absorption with magnesium at a wavelength of 285.2 nm to obtain water-soluble magnesia (W-MgO) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Interference suppressor solution: Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate (2) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.
   c) Magnesium standard solution (MgO 1 mg/mL): Weigh 0.603 g of magnesium (powder) specified in JIS K 8876 into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.
   d) Magnesium standard solution (MgO 0.1 mg/mL): Transfer 10 mL of magnesium standard solution (MgO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   e) Magnesium standard solution (MgO 0.1 mg/mL - 10 mg/mL) for the calibration curve preparation: Transfer 2.5 mL-25 mL of magnesium standard solution (MgO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution (3), and add water up to the marked line.
   f) Blank test solution for the calibration curve preparation: Transfer about 25 mL of interference suppressor solution used in the procedure d) to a 250-mL volumetric flask (3), and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the magnesium standard solution in (2), a magnesium standard solution for the calibration curve preparation can be prepared by using a magnesium standard solution (MgO 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble magnesia (W-MgO) in the analytical sample by multiplying the concentration (Mg) of a magnesium standard solution for calibration curve preparation or a measurement value (Mg) obtained in (4.2) by a conversion factor (1.6583).

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121.
   1) Light source: A magnesium hollow cathode lamp
2) **Gas:** Gas for heating by flame  
   (i) Fuel gas: acetylene  
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Extraction flask** (4): A 500-mL volumetric flask made of borosilicate glass
c) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.

**Note** (4) The volumetric flask used for extraction should be distinguished as an extraction flask and should not be used for the other purposes.

(4) **Test procedure**
(4.1) **Extraction**: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 400-mL tall beaker.
   b) Add 400 mL of water, and cover with a watch glass to boil on a hot plate for about 30 minutes (5).
   c) Immediately transfer it to a 500-mL volumetric flask with water.
   d) After immediate cooling is complete, add water up to the marked line
   e) Filter with Type 3 filter paper to make a sample solution.

**Note** (5) Be aware that an analytical sample should not solidify in the bottom of a beaker.

Comment 2 In the procedure in a), a 500-mL volumetric flask can be used instead of a 500-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. Additionally, “cover with a watch glass” in b) is replaced by “place a funnel”. Skip “transfer to a 500-mL volumetric flask with water” in the procedure in c).

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:  
   Analytical line wavelength: 285.2 nm

b) **Calibration curve preparation**
   1) Spray the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 285.2 nm.
   2) Prepare a curve for the relationship between the magnesium concentration and the indicated value of the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MgO) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the magnesium content from the calibration curve, and calculate the water-soluble magnesia (W-MgO) in the analytical sample.

Comment 3 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 1 % (mass fraction) - 5 % (mass fraction) are 100.4 % - 100.9 % as water-soluble magnesia (W-MgO)
respectively. Additionally, the minimum limit of quantification of this testing method is about 0.07 % (mass fraction).

References

(5) Flow sheet for water-soluble magnesia: The flow sheet for water-soluble magnesia in fertilizers is shown below:

![Flow sheet for water-soluble magnesia in fertilizers (Extraction procedure)](image)

![Flow sheet for water-soluble magnesia in fertilizers (Measurement procedure)](image)
4.6.4.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.6.4.b-2018 or W-Mg.b-1.

Add water to an analytical sample to extract, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the magnesium at a wavelength of 279.553 nm to obtain water-soluble magnesia (W-MgO) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Magnesium standard solution (MgO 1 mg/mL): Weigh 0.603 g of magnesium (powder) specified in JIS K 8876 into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.
   d) Magnesium standard solution (MgO 0.1 mg/mL): Transfer 10 mL of magnesium standard solution (MgO 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   e) Magnesium standard solution (MgO 0.2 µg/mL - 2 µg/mL) for the calibration curve preparation: Transfer 2 mL - 20 mL of magnesium standard solution (MgO 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
   f) Magnesium standard solution (MgO 10 µg/mL) for the calibration curve preparation: Transfer 2 mL - 20 mL of magnesium standard solution (MgO 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
   g) Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the magnesium standard solution in (2), a magnesium standard solution for the calibration curve preparation can be prepared by using a magnesium standard solution (Mg 1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble magnesia (W-MgO) in the analytical sample by multiplying the concentration (Mg) of a magnesium standard solution for calibration curve preparation or a measurement value (Mg) obtained in (4.2) by a conversion factor (1.6583).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a magnesium standard solution for the calibration curve in the concentration range which is suitable to a device used.
(3) **Apparatus and instruments**: Apparatus and instruments are shown below.
   
a) **ICP Optical Emission Spectrometry**: An Optical Emission Spectrometer specified in JIS K 0116.
   
   1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

   a) Weigh 1 mg of an analytical sample (2) to the order of 1 g, and put it in a 100-mL volumetric flask.

   b) Add about 50 mL of water, shake to mix and add water up to the marked line.

   c) Filter with Type 3 filter paper to make a sample solution.

**Note**  (2) The sampling amount of the analytical sample is 10 g when there is less magnesia content in the fertilizers such as a home garden-use fertilizer.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

   a) Measurement conditions for the ICP Optical Emission Spectrometer: Set up the measurement conditions for the ICP Optical Emission Spectrometer considering the following:
      Analytical line wavelength: 279.553 nm

   b) **Calibration curve preparation**

      1) Spray the magnesium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 279.553 nm.

      2) Prepare a curve for the relationship between the magnesium concentration and the indicated value of the magnesium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

   c) **Sample measurement**

      1) Transfer a predetermined amount of the sample solution (the equivalents of 0.02 mg - 1.6 mg as MgO) to a 100-mL volumetric flask.

      2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.

      3) Subject to the same procedure as in b) 1) to read the indicated value.

      4) Obtain the magnesium content from the calibration curve, and calculate the water-soluble magnesia (W-MgO) in the analytical sample.

**Comment 4** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

**Comment 5** The comparison of the measurement value ($y_i$: 0.160 % (mass fraction) - 9.36 % (mass fraction )) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of Flame atomic absorption spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = -0.006 + 0.985x - 0.006$, and its correlation coefficient ($r$) was 0.999. Additionally, additive recovery testing was conducted using a fluid mixed fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand) and a liquid microelement mixed fertilizer (1 sample). As a result, the mean recovery rates at additive level of 1 % (mass fraction) and 0.15 % (mass fraction) were 98.7 % - 102.8 % and 102.3 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.002 % (mass fraction).

### Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$T^{1)}$</td>
<td>$s^{2)}_r$</td>
</tr>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>1.18</td>
<td>0.004</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer</td>
<td>7</td>
<td>0.392</td>
<td>0.002</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test  
2) Average (the number of test days ($T$) × the number of duplicate testing (2))  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Intermediate standard deviation  
7) Intermediate relative standard deviation

### References


(5) **Flow sheet**: The flow sheet for water-soluble magnesia in fluid fertilizers is shown below:

```
1 g analytical sample                Weigh to the order of 1 mg to a 100-mL volumetric flask
    ← Water, about 50 mL
        Shaking to mix
    ← Water (up to the marked line)
        Filtration
        Sample solution
```

Figure 1 Flow sheet for water-soluble magnesia in fluid fertilizers  
(Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot (predetermined volume)</td>
<td>← 25 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
<td>ICP-OES (279.553 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for water-soluble magmesia in fluid fertilizers
(Measurement procedure)
4.7 Manganese
4.7.1 Soluble manganese
4.7.1.a Flame atomic absorption spectrometry

(1) Summary
This test method is applicable to fertilizers containing manganese carbonate fertilizers. This testing method is classified as Type D and its symbol is 4.7.1.a-2017 or S-Mn.a-1.

Add hydrochloric acid (1+23) to an analytical sample, boil to extract and add an interference suppressor solution, and then spray into an acetylene–air flame, and measure the atomic absorption with manganese at a wavelength of 279.5 nm to obtain the hydrochloric acid (1+23) soluble manganese (soluble manganese (S-MnO)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Interference suppressor solution (1): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate (2) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.
   c) Manganese standard solution (MnO 1 mg/mL) (1): Weigh 0.775 g of manganese powder (purity no less than 99 % (mass fraction)) into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.
   d) Manganese standard solution (MnO 0.1 mg/mL) (1): Transfer 10 mL of manganese standard solution (MnO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   e) Manganese standard solution (MnO 1 µg/mL-10 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL-25 mL of manganese standard solution (MnO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution (3), and add water up to the marked line.
   f) Blank test solution for the calibration curve preparation (1): Transfer about 25 mL of interference suppressor solution used in the procedure e) to a 250-mL volumetric flask (3), and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
   (3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the manganese standard solution in (2), a manganese standard solution for the calibration curve preparation can be prepared by using a manganese standard solution (Mn 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate soluble manganese (S-MnO) in the analytical sample by multiplying the concentration (Mn) of a manganese standard solution for calibration curve preparation or a measurement value (Mn) obtained in (4.2) by a conversion factor (1.2912).

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in

JIS K 0121.

1) **Light source**: A manganese hollow cathode lamp
2) **Gas**: Gas for heating by flame
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 °C.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 2 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL tall beaker.

b) Add about 200 mL of hydrochloric acid (1+23), cover with a watch glass, and boil on a hot plate for about 5 minutes (4).

c) Immediately transfer to a 250-mL - 500-mL volumetric flask with water.

d) After immediate cooling is complete, add water up to the marked line.

e) Filter with Type 3 filter paper to make a sample solution.

**Note** (4) Be aware that an analytical sample should not solidify in the bottom of a beaker.

Comment 2 In the procedure in a), a 500-mL volumetric flask can be used instead of a 500-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. In addition, “cover with a watch glass” in b) is replaced by “place a funnel”, and “transfer to a 250-mL - 500 mL volumetric flask with water” in the procedure in c) is skipped.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1) of 4.5.2.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 279.5 nm

b) **Calibration curve preparation**
   1) Spray the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 279.5 nm.
   2) Prepare a curve for the relationship between the manganese concentration and the indicated value of the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MnO) to a 100-mL volumetric flask.
   2) Add about 10 mL of interference suppressor solution (3), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the manganese content from the calibration curve, and calculate the soluble manganese (S-MnO) in the analytical sample.

Comment 4 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 5 % (mass fraction) and
0.1 % (mass fraction) are 100.5 % and 101.3 % as soluble manganese (S-MnO) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.006 % (mass fraction).

References

(5) Flow sheet for soluble manganese: The flow sheet for soluble manganese in fertilizers is shown below:

<table>
<thead>
<tr>
<th>2 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 500-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>↓</td>
<td>About 200 mL of hydrochloric acid (1+23)</td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and boil for about 5 minutes.</td>
</tr>
<tr>
<td>↓</td>
<td>Immediately</td>
</tr>
<tr>
<td>Cooling</td>
<td></td>
</tr>
<tr>
<td>Transfer</td>
<td>Water, 250-mL - 500-mL volumetric flask</td>
</tr>
<tr>
<td>↓</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1  Flow sheet for soluble manganese in fertilizers (Extraction procedure)

| Sample solution |
|↓ | |
|Aliquot (predetermined amount) | 100-mL volumetric flask |
|↓ | About 10 mL of interference suppressor solution |
|↓ | Water (up to the marked line) |
|Measurement | Atomic absorption spectrometer (279.5 nm) |

Figure 2  Flow sheet for soluble manganese in fertilizers (Measurement procedure)
4.7.2 Citrate-soluble manganese
4.7.2.a Flame atomic absorption spectrometry

(1) Summary
This test method is applicable to fertilizers containing manganese carbonate fertilizers. This testing method is classified as Type C and its symbol is 4.7.2.a-2018 or C-Mn.a-2.

Extract by adding a citric acid solution to an analytical sample and add an interference suppressor solution, and then spray into an acetylene–air flame, and measure the atomic absorption with manganese at a wavelength of 279.5 nm to obtain citric acid soluble manganese (citrate-soluble manganese(C-MnO)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
b) Citric acid solution (1): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
c) Interference suppressor solution (1): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.
d) Manganese standard solution (MnO 1 mg/mL) (1): Weigh 0.775 g of manganese powder (purity no less than 99 % (mass fraction)) into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.
e) Manganese standard solution (MnO 0.1 mg/mL): Transfer 10 mL of manganese standard solution (MnO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
f) Manganese standard solution (MnO 1 µg/mL - 10 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL-25 mL of manganese standard solution (MnO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution, and add water up to the marked line.
g) Blank test solution for the calibration curve preparation (1): Transfer about 25 mL of interference suppressor solution used in the procedure f) to a 250-mL volumetric flask, and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.

(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the manganese standard solution in (2), a manganese standard solution for the calibration curve preparation can be prepared by using a manganese standard solution (Mn 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble manganese (C-MnO) in the analytical sample by a multiplying the concentration (Mn) of manganese standard solution for calibration curve preparation or a measurement value (Mn) obtained in (4.2) by a conversion factor (1.2912).

(3) Instruments: Instruments are as shown below:
a) Extractor: Constant-temperature rotary shaker or reciprocating water bath shaker as
described below.

**aa) Constant-temperature rotary shaker**: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

**ab) Reciprocating water bath shaker**: A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/ minute with amplitude of 25 mm - 40 mm.

b) **Flame atomic absorption spectrometer**: An atomic absorption spectrometer specified in JIS K 0121.

1) **Light source**: A manganese hollow cathode lamp
2) **Gas**: Gas for heating by flame
   (i) **Fuel gas**: acetylene
   (ii) **Auxiliary gas**: Air sufficiently free of dust and moisture.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL tall beaker.

b) Add 150 mL of citric acid solution heated up to about 30 °C (4), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (4) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

**Comment 2** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (5).

b) Add 150 mL of citric acid solution heated up to about 30 °C (4), and shake to mix by reciprocating horizontally at amplitude of 25 mm - 40 mm (30 °C ± 1 °C) at 160 times/min for 1 hour.

c) Add water up to the marked line.

d) Filter with Type 3 filter paper to make a sample solution.

**Note** (5) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

**Comment 3** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

**Comment 4** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
Analytical line wavelength: 279.5 nm

b) **Calibration curve preparation**

1) Spray the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 279.5 nm.

2) Prepare a curve for the relationship between the manganese concentration and the indicated value of the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MnO) to a 100-mL volumetric flask.

2) Add about 10 mL of interference suppressor solution \(^{(3)}\), and add water up to the marked line.

3) Subject to the same procedure as in b) 1 to read the indicated value.

4) Obtain the manganese content from the calibration curve, and citrate soluble manganese (C-MnO) in the analytical sample.

**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 5 % (mass fraction) and 0.1 % (mass fraction) are 101.9 % and 100.5 % as citrate-soluble manganese (C-MnO) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.006 % (mass fraction).

<table>
<thead>
<tr>
<th>Certified reference material fertilizer</th>
<th>Number of laboratories</th>
<th>Average (^{(2)})</th>
<th>Repeatability (^{(3)})</th>
<th>Intermediate precision (^{(4)})</th>
<th>Reproducibility (^{(5)})</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-A-10</td>
<td>9</td>
<td>0.403</td>
<td>0.004</td>
<td>0.011</td>
<td>0.010</td>
</tr>
<tr>
<td>FAMIC-A-13</td>
<td>10</td>
<td>0.356</td>
<td>0.010</td>
<td>0.012</td>
<td>0.018</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting flame atomic absorbance spectrometry

2) Average (the number of laboratory \((p) \times\) test days \((2)\) \times the number of replicate testing \((3)\))

3) Mass fraction

4) Repeatability standard deviation

5) Repeatability relative standard deviation

6) Intermediate standard deviation

7) Intermediate relative standard deviation

8) Reproducibility standard deviation

9) Reproducibility relative standard deviation

**References**


(5) **Flow sheet for citrate-soluble manganese:** The flow sheet for citrate-soluble manganese in fertilizers is shown below:

```
<table>
<thead>
<tr>
<th>1 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 250-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>← About 150 mL of citrate-solution [about 30 °C]</td>
<td>Constant-temperature rotary shaker (30 - 40 revolutions/min), 30 °C ± 1 °C, for 1 hour</td>
</tr>
<tr>
<td>Shaking to mix</td>
<td></td>
</tr>
<tr>
<td>Cooling</td>
<td>Immediately</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Filtration</td>
<td></td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>
```

Figure 1-1 Flow sheet for citrate-soluble manganese in fertilizers
(Extraction procedure (4.1.1))

```
<table>
<thead>
<tr>
<th>1 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 250-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>← About 150 mL of citrate solution [about 30 °C]</td>
<td>Reciprocating water bath shaker (reciprocation horizontally at 160 times /min, with amplitude of 25 mm - 40 mm), at 30 °C ± 1 °C, for 1 hour</td>
</tr>
<tr>
<td>Shaking to mix</td>
<td></td>
</tr>
<tr>
<td>Cooling</td>
<td>Immediately</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Filtration</td>
<td></td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>
```

Figure 1-2 Flow sheet for citrate-soluble manganese in fertilizers
(Extraction procedure (4.1.2))
Figure 2  Flow sheet for citrate-soluble manganese in fertilizers  
(Measurement procedure)
4.7.2.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 4.7.2.b-2018 or C-Mn.b-1.

Extract by adding a citric acid solution to an analytical sample, introduce it to an ICP Optical Emission Spectrometer ("ICP-OES") and measure the manganese at a wavelength of 257.610 nm to obtain citric acid-soluble manganese (citrate-soluble manganese (C-MnO)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.
- Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
- Citric acid solution: Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
- Manganese standard solution (MnO 1 mg/mL): Weigh 0.775 g of manganese powder (purity no less than 99 % (mass fraction)) into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and further add hydrochloric acid (1+23) up to the marked line.
- Manganese standard solution (MnO 0.1 mg/mL): Transfer 10 mL of manganese standard solution (MnO 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
- Manganese standard solution (MnO 0.1 µg/mL - 2 µg/mL) for the calibration curve preparation: Transfer 2 mL - 8 mL of manganese standard solution (MnO 0.1 mg/mL) to 100-mL volumetric flasks step-by-step and add hydrochloric acid (1+23) up to the marked line.
- Manganese standard solution (MnO 0.1 µg/mL - 2 µg/mL) for the calibration curve preparation: Transfer 1 mL - 20 mL of manganese standard solution (MnO 10 µg/mL) to 100-mL volumetric flasks step-by-step and add hydrochloric acid (1+23) up to the marked line.
- Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in e) and g).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the manganese standard solution in (2), a manganese standard solution for the calibration curve preparation can be prepared by using a manganese standard solution (Mn 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble manganese (C-MnO) in the analytical sample by multiplying the concentration (Mn) of manganese standard solution for calibration curve preparation or a measurement value (Mn) obtained in (4.2) by a conversion factor (1.2912).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in f) and g) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a manganese standard solution for the calibration curve in the concentration range which is suitable to a
device used.

(3) **Instruments:** Instruments are as shown below:

a) **ICP Optical Emission Spectrometry:** A spectrophotometer specified in JIS K 0115
   1) **Gas:** Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity
b) **Extractor:** Constant-temperature rotary shaker or reciprocating water bath shaker as described below.

   ba) **Constant-temperature rotary shaker:** A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.

   bb) **Reciprocating water bath shaker:** A reciprocating water bath shaker: that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/ minute with amplitude of 25 mm - 40 mm.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker:**

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL tall beaker.

b) Add 150 mL of citric acid solution heated up to about 30 °C (2), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note**  (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

Comment 2 The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker:**

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask(3).

b) Add 150 mL of citric acid solution heated up to about 30 °C (2), and shake to mix by reciprocating horizontally at amplitude of 25 mm - 40 mm (30 °C ± 1 °C) at 160 times/min for 1 hour.

c) After immediate cooling is complete, add water up to the marked line

d) Filter with Type 3 filter paper to make a sample solution.

**Note**  (3) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

Comment 3 The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

Comment 4 The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer:** Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
Analytical line wavelength: 257.610 nm

b) **Calibration curve preparation**
1) Spray the manganese standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 257.610 nm.
2) Prepare a curve for the relationship between the manganese concentration and the indicated value of the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 0.8 mg as MnO) to a 100-mL volumetric flask.
2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
3) Subject to the same procedure as in b) 1) to read the indicated value.
4) Obtain the manganese content from the calibration curve, and calculate the citrate soluble manganese (C-MnO) in the analytical sample.

**Comment 5** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.3.d Comment 7.

**Comment 6** The comparison of the measurement value ($y_i$: 0.089 % (mass fraction) - 1.88 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of Flame atomic absorption spectrometry analysis was conducted to evaluate trueness using compound fertilizers (7 samples), mixed phosphate fertilizers (2 samples), solid fertilizers (2 samples), blended fertilizers (4 samples) and organic blended fertilizer (1 sample). As a result, a regression equation was $y = 0.0015 + 0.9988x$, and its correlation coefficient ($r$) was 0.999. In addition, recovery testing was conducted using a preparation sample. As a result, the average rate of recovery at the content level of 0.595 % (mass fraction) - 28.94 % (mass fraction) was 98.5 % - 105.5 %.

The results of the repeatability tests on different days using compound fertilizers and blended fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction).

| Name of sample          | Test days of repeatability $T$ | Average Repeatability $s$ | $RSD_R$ | Intermediate precision $s_{I(T)}$ $RSD_{I(T)}$
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$^{1)}$</td>
<td>$^{2)}$</td>
<td>$^{3)}$</td>
<td>$^{4)}$</td>
<td>$^{5)}$</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>0.54</td>
<td>0.01</td>
<td>2.3</td>
<td>0.02</td>
</tr>
<tr>
<td>Blended fertilizer</td>
<td>7</td>
<td>0.089</td>
<td>0.002</td>
<td>1.9</td>
<td>0.002</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) × the number of duplicate testing ($p$))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

Table 1  Analysis results of the repeatability tests on different days
(5) **Flow sheet for citrate-soluble manganese**: The flow sheet for citrate-soluble manganese in fertilizers is shown below:

![Flow sheet for citrate-soluble manganese in fertilizers](image1)

- **Extraction procedure (4.1.1)**
  - Weigh to the order of 1 mg into a 250-mL tall beaker.
  - About 150 mL of citrate solution [about 30 ºC]
  - Constant-temperature rotary shaker (30 - 40 revolutions/min), 30 ºC ± 1 ºC, for 1 hour
  - Shaking to mix
  - Cooling
    - Immediately
    - Water (up to the marked line)
  - Filtration
    - Type 3 filter paper
  - Sample solution

![Flow sheet for citrate-soluble manganese in fertilizers](image2)

- **Extraction procedure (4.1.2)**
  - Weigh to the order of 1 mg into a 250-mL tall beaker.
  - About 150 mL of citrate solution [about 30 ºC]
  - Reciprocating water bath shaker (reciprocation horizontally at 160 times /min, with amplitude of 25 mm - 40 mm), at 30 ºC ± 1 ºC, for 1 hour
  - Shaking to mix
  - Cooling
    - Immediately
    - Water (up to the marked line)
  - Filtration
    - Type 3 filter paper
  - Sample solution

![Flow sheet for citrate-soluble manganese in fertilizers](image3)

- **Measurement procedure**
  - Sample solution
  - Aliquot (predetermined amount)
  - 100-mL volumetric flask
  - About 25 mL of hydrochloric acid (1+5)
  - Water (up to the marked line)
  - Measurement
    - ICP-OES (257.610 nm)

Figure 1-1 Flow sheet for citrate-soluble manganese in fertilizers
(Extraction procedure (4.1.1))

Figure 1-2 Flow sheet for citrate-soluble manganese in fertilizers
(Extraction procedure (4.1.2))

Figure 2 Flow sheet for citrate-soluble manganese in fertilizers
(Measurement procedure)
4.7.3 Water-soluble manganese
4.7.3.a Flame atomic absorption spectrometry

(1) Summary
This test method is applicable to fertilizers containing manganese sulfate fertilizers. This testing method is classified as Type D and its symbol is 4.7.3.a-2017 or W-Mn.a-1.
Extract by adding water to an analytical sample and add an interference suppressor solution, and then spray in an acetylene–air flame, and measure the atomic absorption with manganese at a wavelength of 279.5 nm to obtain water-soluble manganese (W-MnO) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.

a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

b) Interference suppressor solution \(^{(1)}\): Weigh 60.9 g - 152.1 g of strontium chloride hexahydrate \(^{(2)}\) specified in JIS K 8132 into a 2000-mL beaker, add a small amount of water, add gradually 420 mL of hydrochloric acid to dissolve, and further add water to make 1000 mL.

c) Manganese standard solution (MnO 1 mg/mL) \(^{(1)}\): Weigh 0.775 g of manganese powder (purity no less than 99 % (mass fraction)) into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and add water up to the marked line.

d) Manganese standard solution (MnO 0.1 mg/mL) \(^{(1)}\): Transfer 10 mL of manganese standard solution (MnO 1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.

e) Manganese standard solution (MnO 1 µg/mL - 10 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 2.5 mL-25 mL of manganese standard solution (MnO 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, add about 25 mL of interference suppressor solution \(^{(3)}\), and add water up to the marked line.

f) Blank test solution for the calibration curve preparation \(^{(1)}\): Transfer about 25 mL of interference suppressor solution used in the procedure d) to a 250-mL volumetric flask \(^{(3)}\), and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) 29 g of lanthanum oxide (atomic absorption analysis grade or equivalents) can also be used.
(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the manganese standard solution in (2), a manganese standard solution for the calibration curve preparation can be prepared by using a manganese standard solution (Mn 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble manganese (C-MnO) in the analytical sample by a multiplying the concentration (Mn) of manganese standard solution for calibration curve preparation or a measurement value (Mn) obtained in (4.2) by a conversion factor (1.2912).

(3) Instruments: Instruments are as shown below.

a) Extractor: Constant-temperature rotary shaker or vertical reciprocating shaker as described below.

aa) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
ab) **Vertical reciprocating shaker**: A vertical reciprocating shaker that can shake a 250-mL volumetric flask using an adapter for a flask to reciprocate vertically at 300 times/minute (amplitude of 40 mm).

b) **Flame atomic absorption spectrometer**: An atomic absorption spectrometer specified in JIS K 0121.
   1) **Light source**: A manganese hollow cathode lamp
   2) **Gas**: Gas for heating by flame
      (i) **Fuel gas**: acetylene
      (ii) **Auxiliary gas**: Air sufficiently free of dust and moisture.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Powdery test sample**

(4.1.1.1) **Rotary shaker**
   a) Weigh 5 g of an analytical sample to the order of 1 mg, and transfer to a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

**Comment 2** In the procedure of (4.1.1.1) a), it is also allowed to weigh 2.5 g of an analytical sample to the order of 1 mg, and put it in a 250-mL volumetric flask.

**Comment 3** The procedure in (4.1.1.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.1.1.2) **Vertical reciprocating shaker**:
   a) Weigh 2.5 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask.
   b) Add about 200 mL of water, and shake to mix by reciprocating vertically at 300 times/min (amplitude of 40 mm) for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

**Comment 4** The procedure in (4.1.1.2) is the same as the procedure in (4.1.1.2) in 4.2.4.a.

(4.1.2) **Fluid test sample**
   a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 100-mL volumetric flask.
   b) Add about 50 mL of water, and shake to mix.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

**Comment 5** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.4.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.
   a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
      Analytical line wavelength: 279.5 nm
   b) **Calibration curve preparation**
1) Spray the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 279.5 nm.

2) Prepare a curve for the relationship between the manganese concentration and the indicated value of the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as MnO) to a 100-mL volumetric flask.

2) Add about 10 mL of interference suppressor solution \(^{(3)}\), and add water up to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Obtain the manganese content from the calibration curve, and water-soluble manganese (W-MnO) in the analytical sample.

**Comment 6** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 6 % (mass fraction) and 0.1 % (mass fraction) are 101.2 % and 101.1 % as water-soluble manganese (W-MnO) respectively.

The comparison of the measurement value of extraction \(y_i: 0.0330\%\) (mass fraction) - 6.18 % (mass fraction) by a vertical reciprocating shaker and the measurement value of extract with a rotary shaker \(x_i\) was conducted to evaluate trueness of the extraction of solid fertilizers using fertilizers (12 samples). As a result, a regression equation was \(y = -0.009 + 1.011x\), and its correlation coefficient \((r)\) was 1.000. Also, the results of the repeatability tests on different days using compound fertilizers and mixed micro element fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

The comparison of the measurement value of simple extraction \(y_i: 0.0590\%\) (mass fraction) - 1.27 % (mass fraction) and the measurement value of extract \(x_i\) with a rotary shaker was conducted to evaluate trueness of the extraction of fluid fertilizers using fluid fertilizers (12 samples). As a result, a regression equation was \(y = -0.001 + 1.006x\), and its correlation coefficient \((r)\) was 1.000. Also, the results of the repeatability tests on different days using fluid compound fertilizers and fluid mixed micro element fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 2 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.004 % (mass fraction).
### Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability (T)</th>
<th>Average (A) (%)</th>
<th>Repeatability (RSD)%</th>
<th>Intermediate precision (s) (%)</th>
<th>Intermediate precision (RSD)%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed micro element fertilizers</td>
<td>7</td>
<td>3.57</td>
<td>0.03</td>
<td>0.7</td>
<td>0.05</td>
</tr>
<tr>
<td>Blended fertilizer</td>
<td>7</td>
<td>0.226</td>
<td>0.002</td>
<td>1.0</td>
<td>0.004</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \(T\) × the number of duplicate testing \(2\))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

### Table 2  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability (T)</th>
<th>Average (A) (%)</th>
<th>Repeatability (RSD)%</th>
<th>Intermediate precision (s) (%)</th>
<th>Intermediate precision (RSD)%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid compound fertilizers</td>
<td>7</td>
<td>1.28</td>
<td>0.01</td>
<td>0.4</td>
<td>0.02</td>
</tr>
<tr>
<td>Liquid micro element compound fertilizers</td>
<td>7</td>
<td>0.230</td>
<td>0.001</td>
<td>0.5</td>
<td>0.003</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \(T\) × the number of duplicate testing \(2\))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

### References

(5) **Flow sheet for water-soluble manganese**: The flow sheet for water-soluble manganese in fertilizers is shown below:

<table>
<thead>
<tr>
<th>2.5 g analytical sample (Powdery)</th>
<th>Weigh to the order of 1 mg into a 250-mL volumetric flask.</th>
</tr>
</thead>
<tbody>
<tr>
<td>≈ About 200 mL of water</td>
<td>Vertical reciprocating shaker (300 times/min, with amplitude of 40 mm), for 30 minutes</td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1-2 Flow sheet for water-soluble manganese in fertilizers (Extraction procedure (4.1.2))

<table>
<thead>
<tr>
<th>1 g analytical sample (Fluid)</th>
<th>Weigh to the order of 1 mg into a 100-mL volumetric flask.</th>
</tr>
</thead>
<tbody>
<tr>
<td>≈ About 50 mL of water</td>
<td></td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1-3 Flow sheet for water-soluble manganese in fertilizers (Extraction procedure (4.1.2))
Sample solution

100-mL volumetric flask

Aliquot (predetermined amount)

← About 10 mL of interference suppressor solution
← Water (up to the marked line)

Measurement

Atomic absorption spectrometer (279.5 nm)

Figure 2  Flow sheet for water-soluble manganese in fertilizers
(Measurement procedure)
4.7.3.b ICP Optical Emission Spectrometry

(1) Summary
The test method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers, liquid by-product manganese fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.7.3.b-2017 or W-Mn.b-1.

Add water to an analytical sample to extract, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the manganese at a wavelength of 257.610 nm to obtain water-soluble manganese (W-MnO) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Manganese standard solution (MnO 1 mg/mL) \(^{(1)}\): Weigh 0.775 g of manganese powder (purity no less than 99 % (mass fraction)) into a weighing dish. Transfer to a 1000-mL volumetric flask with a small amount of water, add about 10 mL of hydrochloric acid to dissolve, and further add hydrochloric acid (1+23) up to the marked line.
   d) Manganese standard solution (MnO 0.1 mg/mL) \(^{(1)}\): Transfer 10 mL of manganese standard solution (MnO 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   e) Manganese standard solution (MnO 2 µg/mL - 8 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 2 mL - 8 mL of manganese standard solution (MnO 0.1 mg/mL) to 100-mL volumetric flasks step-by-step and add hydrochloric acid (1+23) up to the marked line.
   f) Manganese standard solution (MnO 0.1 µg/mL - 2 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 1 mL - 20 mL of manganese standard solution (MnO 10 µg/mL) to 100-mL volumetric flasks step-by-step and add hydrochloric acid (1+23) up to the marked line.
   g) Blank test solution for the calibration curve preparation \(^{(1)}\): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the manganese standard solution in (2), a manganese standard solution for the calibration curve preparation can be prepared by using a manganese standard solution (Mn 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble manganese (W-MnO) in the analytical sample by a multiplying the concentration (Mn) of manganese standard solution for calibration curve preparation or a measurement value (Mn) obtained in (4.2) by a conversion factor (1.2912).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in d) and e) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a manganese standard solution for the calibration curve in the concentration range which is suitable to a device used.
Apparatus and instruments: Apparatus and instruments are shown below.


1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 1 mg of an analytical sample to the order of 1 mg, and put it in a 100-mL volumetric flask.

b) Add about 50 mL of water, shake to mix and add water up to the marked line.

c) Filter with Type 3 filter paper to make a sample solution.

Note (2) The sampling amount of the analytical sample is 10 g when there is less manganese content in the fertilizers such as a home garden-use fertilizer.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Optical Emission Spectrometer: Set up the measurement conditions for the ICP Optical Emission Spectrometer considering the following:

Analytical line wavelength: 257.610 nm

b) Calibration curve preparation

1) Spray the manganese standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 257.610 nm.

2) Prepare a curve for the relationship between the manganese concentration and the indicated value of the manganese standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 0.8 mg as MnO) to a 100-mL volumetric flask.

2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Obtain the manganese content from the calibration curve, and water-soluble manganese (W-MnO) in the analytical sample.

Comment 4 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

Comment 5 The comparison of the measurement value (yi: 0.027 % (mass fraction) - 1.49 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value (xi) of Flame atomic absorption spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = -0.0013 + 1.025x$, and its correlation coefficient ($r$) was 0.999. Additionally, additive recovery testing was conducted using a fluid mixed fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand) and a liquid micro element mixed fertilizer (1 sample). As a result, the mean recovery rates at additive level of 0.15 % (mass fraction) - 0.2 % and 0.005 % (mass fraction) were 96.3 % - 96.5 % and 107.0 %
respectively. The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.0002 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^1$ (%)</th>
<th>$s_r^3$ (%)</th>
<th>$RSD_r^5$ (%)</th>
<th>Intermediate precision $^1$ (%)</th>
<th>$s_{I(T)}^6$ (%)</th>
<th>$RSD_{I(T)}^7$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid compound fertilizer</td>
<td>7</td>
<td>5.69</td>
<td>0.02</td>
<td>0.4</td>
<td>0.06</td>
<td>1.1</td>
<td></td>
</tr>
<tr>
<td>Home garden-use mixed fertilizers (Liquid)</td>
<td>7</td>
<td>2.29</td>
<td>0.02</td>
<td>0.8</td>
<td>0.04</td>
<td>1.6</td>
<td></td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) x the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Intermediate standard deviation
6) Intermediate relative standard deviation

**References**


(5) **Flow sheet**: The flow sheet for water-soluble manganese in fluid fertilizers is shown below:

1 g analytical sample Weigh to the order of 1 mg to a 100-mL volumetric flask

← Water, about 50 mL

Shaking to mix

← Water (up to the marked line)

Filtration

Sample solution

Figure 1 Flow sheet for water-soluble manganese in liquid fertilizers
(Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot (predetermined volume)</td>
<td>← 25 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
<td>ICP-OES (257.610 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for water-soluble manganese in liquid fertilizers
(Measurement procedure)
4.8 Boron
4.8.1 Citrate-soluble boron
4.8.1.a Azomethine-H method

(1) Summary
This testing method is applicable to fertilizers not containing organic matters. This testing method is classified as Type C and its symbol is 4.8.1.a-2018 or C-B.a-2.
Extract by adding a citric acid solution to an analytical sample, mask co-existing copper, iron and other salts with ethylenediamine tetraacetate and measure the absorbance with azomethine-H borate formed by the reaction with azomethine-H to obtain citrate soluble boron (C-B\(_2\)O\(_3\)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.
   a) **Citric acid solution** \((1)\): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   b) **Ethylenediamine tetraacetate solution** \((1)\): Dissolve 37.2 g of ethylenediaminetetraacetic acid dihydrogen disodium dihydrate specified in JIS K 8107 in water to make 1000 mL.
   c) **Ammonium acetate solution** \((1)\): Dissolve 250 g of ammonium acetate specified in JIS K 8359 in water to make 500 mL and adjust pH with sulfuric acid (1+4) to pH 5.2 ± 0.1.
   d) **Azomethine-H solution**: Add water to 0.6 g of azomethine-H and 2 g of L (+) – ascorbic acid specified in JIS K 9502, and heat up to 35 °C - 40 °C to dissolve and add water after cooling to make 100 mL.
   e) **Boron standard solution** \((B_2O_3 \ 2.5 \ mg/mL)\) \((1)\): After leaving boric acid specified in JIS K 8863 at rest in a desiccator for about 24 hours to dry, weigh 4.441 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   f) **Boron standard solution** \((B_2O_3 \ 0.05 \ mg/mL)\): Dilute the predetermined volume of boron standard solution \((B_2O_3 \ 2.5 \ mg/mL)\) with water exactly by a factor of 50.

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the boron standard solution in (2), a boron standard solution for the calibration curve preparation can be prepared by using a boron standard solution \((B \ 1 \ mg/mL \ or \ 10 \ mg/mL)\) traceable to National Metrology. In this case, calculate citrate-soluble boron \((C-B_2O_3)\) in the analytical sample by multiplying the concentration \((B)\) of a boron standard solution for calibration curve preparation or a measurement value \((B)\) obtained in (4.3) by a conversion factor (3.2199).

(3) Instruments: Instruments are as shown below:
   a) **Extractor**: Constant-temperature rotary shaker or reciprocating water bath shaker as described below.
      aa) **Constant-temperature rotary shaker**: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 °C ± 1 °C, upside down at 30 - 40 revolutions/min.
      ab) **Reciprocating water bath shaker**: A reciprocating water bath shaker that can be adjusted to 30 °C ± 1 °C and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/ minute with amplitude of 25 mm - 40 mm.
   b) **Spectrophotometer**: A spectrophotometer specified in JIS K 0115

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Constant-temperature rotary shaker:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL tall beaker.
b) Add 150 mL of citric acid solution heated up to about 30 °C \(^{(2)}\), and shake to mix at 30 - 40 revolutions/min (30 °C ± 1 °C) for 1 hour.
c) After immediate cooling is complete, add water up to the marked line
d) Filter with Type 3 filter paper to make a sample solution.

Note (2) Shake to mix the volumetric flask gently and disperse an analytical sample into the citric acid solution.

Comment 2 The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) Reciprocating water bath shaker:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask \(^{(3)}\).
b) Add 150 mL of citric acid solution heated up to about 30 °C \(^{(2)}\), and shake to mix by reciprocating horizontally at amplitude of 25 mm - 40 mm (30 °C ± 1 °C) at 160 times/min for 1 hour.
c) After immediate cooling is complete, add water up to the marked line
d) Filter with Type 3 filter paper to make a sample solution.

Note (3) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

Comment 3 The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

Comment 4 The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) Coloring: Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 1 mg as \(P_2O_5\) and no more than the equivalents of 15 mL of citric acid solution) to a 100-mL volumetric flask.
b) Add the solution to make citric acid solution equivalent to 15 mL.
c) Add 25 mL of ethylenediamine tetraacetate solution, and then add 10 mL of ammonium acetate solution.
d) Add 10 mL of azomethine-H solution, and further add water up to the marked line, and then leave at rest for about 2 hours.

Comment 5 If formaldehyde processed urea, a large quantity of aluminum, copper, iron zinc, organic matters, etc. coexists to affect quantification, transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 1 mg as \(B_2O_3\), no more than 10 mL of the solution) to a 100-mL separatory funnel, add 10 mL of hydrochloric acid (1+3), add water to about 20 mL and add 20 mL of 2-ethyl-1,3-hexanediol – 4-methyl-2-pentanone (1+9) to shake to mix with a shaking apparatus for about 1 minute. After allowing to stand still, remove the lower layer (aqueous phase) and add 20 mL of sodium hydroxide (20 mg/L) to shake to mix with a shaking apparatus for about 1 minute. After allowing to stand still, transfer the lower layer (aqueous phase) to a 100-mL volumetric flask, add a few drops of phenolphthalein solution (1 g/100mL) to neutralize with hydrochloric acid (1+3) until the color of the solution becomes achromatic, and conduct the
procedure in (4.2) b).

(4.3) **Measurement:** Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

**a)** **Measurement conditions of spectrophotometer:** Set up the measurement conditions of spectrophotometer considering the following.
Detection wavelength: 415 nm

**b)** **Calibration curve preparation**
1) Transfer 1 mL - 20 mL of boron standard solution (B₂O₃ 0.05 mg/mL) to 100-mL volumetric flasks step-by-step.
2) Add 15 mL of citric acid solution and conduct the same procedure as (4.2) c) to make the B₂O₃ 0.05 mg/100 mL - 1 mg/100 mL boron standard solution for the calibration curve preparation.
3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.
4) Measure absorbance at a wavelength of 415 nm of the boron standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.
5) Prepare a curve for the relationship between the boron concentration and absorbance of the boron standard solutions for the calibration curve preparation.

**c)** **Sample measurement**
1) Regarding the solution in (4.2) c), measure absorbance by the same procedure as b) 4).
2) Obtain the boron (B₂O₃) content from the calibration curve, and calculate citrate soluble boron (C-B₂O₃) in the analytical sample.

**Comment 6** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) and 0.05 % (mass fraction) were 101.5 % and 95.7 % as citrate-soluble boron (C-B₂O₃) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).
<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of certified reference laboratory (p)</th>
<th>Average (\bar{x}) (%)</th>
<th>Repeatability standard deviation (s_r) (%)</th>
<th>Repeatability relative standard deviation (RSD_r) (%)</th>
<th>Intermediate precision standard deviation (s_{I(T)}) (%)</th>
<th>Intermediate precision relative standard deviation (RSD_{I(T)}) (%)</th>
<th>Reproducibility standard deviation (s_R) (%)</th>
<th>Reproducibility relative standard deviation (RSD_R) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-A-10</td>
<td>11</td>
<td>0.209</td>
<td>0.004</td>
<td>2.0</td>
<td>0.005</td>
<td>2.2</td>
<td>0.006</td>
<td>3.1</td>
</tr>
<tr>
<td>FAMIC-A-13</td>
<td>10</td>
<td>0.203</td>
<td>0.004</td>
<td>1.8</td>
<td>0.005</td>
<td>2.5</td>
<td>0.009</td>
<td>4.7</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis
2) Average (the number of laboratory \(p\) × test days (2) \times the number of replicate testing (3))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

References

(5) **Flow sheet for citrate-soluble boron**: The flow sheet for citrate-soluble boron in fertilizers is shown below:

![Flow sheet for citrate-soluble boron in fertilizers](image)

**Figure 1-1** Flow sheet for citrate-soluble boron in fertilizers  
(Extraction procedure (4.1.1))

![Flow sheet for citrate-soluble boron in fertilizers](image)

**Figure 1-2** Flow sheet for citrate-soluble boron in fertilizers  
(Extraction procedure (4.1.2))
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot</td>
<td></td>
</tr>
<tr>
<td>(predetermined</td>
<td>Citric acid solution,</td>
</tr>
<tr>
<td>amount)</td>
<td>until the volume</td>
</tr>
<tr>
<td></td>
<td>reaches to the</td>
</tr>
<tr>
<td></td>
<td>equivalents of 15 mL</td>
</tr>
<tr>
<td></td>
<td>25 mL of ethylenediamine tetraacetate solution</td>
</tr>
<tr>
<td></td>
<td>10 mL of ammonium acetate solution</td>
</tr>
<tr>
<td></td>
<td>10 mL of azomethine-H solution</td>
</tr>
<tr>
<td></td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Leaving at rest</td>
<td>For about 2 hours</td>
</tr>
<tr>
<td>Measurement</td>
<td>Spectrophotometer (415 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for citrate-soluble boron in fertilizers
(Measurement procedure)
4.8.1.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 4.8.1.b-2018 or C-B.b-1.

Extract by adding a citric acid solution to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the boron at a wavelength of 249.773 nm to obtain citric acid-soluble boron (citrate-soluble boron acid (C-B$_2$O$_3$)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

(2) Reagent: Reagents are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Citric acid solution (1): Dissolve 20 g of citric acid monohydrate specified in JIS K 8283 in water to make 1000 mL.
   d) Boron standard solution (B$_2$O$_3$ 2.5 mg/mL) (1): After leaving boric acid specified in JIS K 8863 at rest in a desiccator for about 24 hours to dry, weigh 4.441 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   e) Boron standard solution (B$_2$O$_3$ 0.1 mg/mL) (1): Transfer 4 mL of boron standard solution (B$_2$O$_3$ 2.5 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line (2).
   f) Boron standard solution (B$_2$O$_3$ 2 µg/mL - 16 µg/mL) for the calibration curve preparation (1): Transfer 2 mL - 16 mL of boron standard solution (B$_2$O$_3$ 0.1 mg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line (2).
   g) Boron standard solution (B$_2$O$_3$ 0.2 µg/mL - 2 µg/mL) for the calibration curve preparation (1): Transfer 2 mL - 20 mL of boron standard solution (B$_2$O$_3$ 10 µg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line (2).
   h) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in e) - g).

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) When preserving, use a container, which can be sealed tightly, made of material such as PTFE that boron hardly elutes

Comment 1 Instead of the boron standard solution in (2), a boron standard solution for the calibration curve preparation can be prepared by using a boron standard solution (B 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate citrate-soluble boron (C-B$_2$O$_3$) in the analytical sample by multiplying the concentration (B) of a boron standard solution for calibration curve preparation or a measurement value (B) obtained in (4.2) by a conversion factor (3.2199).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in f) and g) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a boron standard solution
for the calibration curve in the concentration range which is suitable to a device used.

(3) **Instruments**: Instruments are as shown below:

a) **ICP Optical Emission Spectrometry**: A spectrophotometer specified in JIS K 0115

1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

b) **Extractor**: Constant-temperature rotary shaker or reciprocating water bath shaker as described below.

ba) **Constant-temperature rotary shaker**: A constant-temperature rotary shaker that can rotate a 250-mL volumetric flask, set up in a thermostat adjustable to 30 ºC ± 1 ºC, upside down at 30 - 40 revolutions/min.

bb) **Reciprocating water bath shaker**: A reciprocating water bath shaker: that can be adjusted to 30 ºC ± 1 ºC and shake a 250-mL volumetric flask, set up vertically on the water surface using a shaking rack, reciprocate horizontally at 160 times/ minute with amplitude of 25 mm - 40 mm.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

(4.1.1) **Constant-temperature rotary shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL tall beaker.

b) Add 150 mL of citric acid solution heated up to about 30 ºC (3), and shake to mix at 30 - 40 revolutions/min (30 ºC ± 1 ºC) for 1 hour.

c) After immediate cooling is complete, add water up to the marked line
d) Filter with Type 3 filter paper to make a sample solution.

**Note** (3) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

**Comment 3** The procedure in (4.1.1) is the same as the procedure in (4.1.1) in 4.2.3.a.

(4.1.2) **Reciprocating water bath shaker**:

a) Weigh 1 g of an analytical sample to the order of 1 mg, and transfer to a 250-mL volumetric flask (4).

b) Add 150 mL of citric acid solution heated up to about 30 ºC (3), and shake to mix by reciprocating horizontally at amplitude of 25 mm - 40 mm (30 ºC ± 1 ºC) at 160 times/min for 1 hour.

c) After immediate cooling is complete, add water up to the marked line
d) Filter with Type 3 filter paper to make a sample solution.

**Note** (4) Use a 250-mL flat-bottom volumetric flask to stabilize the vibration.

**Comment 4** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.3.a.

**Comment 5** The determination may be affected if an analytical sample cakes on the bottom of a 250-mL volumetric flask. Therefore, confirm the status of the non-dissolved matters after the procedures of (4.1.1) b) and (4.1.2) b).

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
Analytical line wavelength: 249.773 nm

b) **Calibration curve preparation**

1) Spray the boron standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 249.773 nm.

2) Prepare a curve for the relationship between the boron concentration and the indicated value of the boron standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.02 mg - 1.6 mg as \( \text{B}_2 \text{O}_3 \)) to a 100-mL volumetric flask.

2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Obtain the boron (\( \text{B}_2 \text{O}_3 \)) content from the calibration curve, and calculate citrate soluble boron (C-B\( \text{B}_2 \text{O}_3 \)) in the analytical sample.

**Comment 6** Wash the sample injector of an ICP-OES sufficiently with water because boron easily causes the memory effect.

**Comment 7** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.3.d Comment 7.

**Comment 8** The comparison of the measurement value \( (y_i; 0.073\% \text{ (mass fraction)} - 0.51\% \text{ (mass fraction)}) \) of ICP Optical Emission Spectrometry and the measurement value \( (x_i) \) of Flame atomic absorption spectrometry analysis was conducted to evaluate trueness using compound fertilizers (7 samples), mixed phosphate fertilizers (1 sample), solid fertilizers (2 samples), blended fertilizers (3 samples) and organic compound fertilizer (1 sample). As a result, a regression equation was \( y = -0.0408 + 1.0456x \), and its correlation coefficient \( (r) \) was 0.992. In addition, recovery testing was conducted using a preparation sample. As a result, the average rate of recovery at the content level of 0.601 \% (mass fraction) - 35.51 \% (mass fraction) was 97.0 \% - 102.0 \%.

The results of the repeatability tests on different days using compound fertilizers and blended fertilizers to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.01 \% (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average (^2) ( (% ) (^3)</th>
<th>( s_r ) (^4) ( (% ) (^3)</th>
<th>( RSD_r ) (^5) ( (% ) (^3)</th>
<th>Intermediate precision ( \times \text{the number of duplicate testing (2)} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>0.38</td>
<td>0.01</td>
<td>1.9</td>
<td>0.01 \times 3.1</td>
</tr>
<tr>
<td>Blended fertilizer</td>
<td>7</td>
<td>0.076</td>
<td>0.003</td>
<td>4.2</td>
<td>0.006 \times 7.5</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test 6) Intermediate standard deviation

2) Average \( \text{the number of test days (}T\) \) 7) Intermediate relative standard deviation

3) Mass fraction

4) Repeatability standard deviation
(5) **Flow sheet for citrate-soluble boron**: The flow sheet for citrate-soluble boron in fertilizers is shown below:

![Flow sheet for citrate-soluble boron](image1)

Figure 1-1  Flow sheet for citrate-soluble boron in fertilizers  
(Extraction procedure (4.1.1))

![Flow sheet for citrate-soluble boron](image2)

Figure 1-2  Flow sheet for citrate-soluble boron in fertilizers  
(Extraction procedure (4.1.2))

![Flow sheet for citrate-soluble boron](image3)

Figure 2  Flow sheet for citrate-soluble boron in fertilizers  
(Measurement procedure)
4.8.2 Water-soluble boron
4.8.2.a Azomethine-H method

(1) Summary
This test method is applicable to fertilizers containing borate fertilizers, etc. This testing method is classified as Type D and its symbol is 4.8.2.a-2017 or W-B.a-1.

Extract by adding water to an analytical sample, boil to extract, and mask co-existing copper, iron and other salts with ethylenediamine tetraacetate and measure the absorbance with azomethine-H borate formed by the reaction with azomethine-H to obtain water-soluble boron (W-B2O3). In addition, the performance of this testing method is shown in Comment 7.

(2) Reagent: Reagents are as shown below.
   a) Ethylenediamine tetraacetate solution (1): Dissolve 37.2 g of ethylenediaminetetraacetic acid dihydrogen disodium dihydrate specified in JIS K 8107 in water to make 1000 mL.
   b) Ammonium acetate solution (1): Dissolve 250 g of ammonium acetate specified in JIS K 8359 in water to make 500 mL and adjust pH with sulfuric acid (1+4) to pH 5.2 ± 0.1.
   c) Azomethine-H solution (1): Add water to 0.6 g of azomethine-H and 2 g of L (+) – ascorbic acid specified in JIS K 9502, and heat up to 35 ºC - 40 ºC to dissolve and add water after cooling to make100 mL.
   d) Boron standard solution (B2O3 2.5 mg/mL) (1): After leaving boric acid specified in JIS K 8863 at rest in a desiccator for about 24 hours to dry, weigh 4.441 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   e) Boron standard solution (B2O3 0.05 mg/mL): Dilute the predetermined volume of boron standard solution (B2O3 2.5 mg/mL) with water exactly by a factor of 50.

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the boron standard solution in (2), a boron standard solution for the calibration curve preparation can be prepared by using a boron standard solution (B 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble boron (W-B2O3) in the analytical sample by multiplying the concentration (B) of a boron standard solution for calibration curve preparation or a measurement value (B) obtained in (4.3) by a conversion factor (3.2199).

(3) Instruments: Instruments are as shown below:
   a) Spectrophotometer: A spectrophotometer specified in JIS K 0115
   b) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 ºC.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 2.5 g (2) of an analytical sample to the order of 1 mg, and transfer to a 300-mL tall beaker.
   b) Add about 200 mL of water, and cover with a watch glass and heat on a hot plate to boil for about 15 minutes.
   c) Immediately transfer it with water to a 250- mL volumetric flask.
   d) After immediate cooling is complete, add water up to the marked line
   e) Filter with Type 3 filter paper to make a sample solution.

Note (2) The sampling amount of the analytical sample is 1 g when there is a high amount of boric acid content in the fertilizers such as a borate fertilizer and boric acid fertilizer.
Comment 2  In the procedure in \((4.1.2)\) a) and \((4.1.2)\) b), a 250-mL volumetric flask can be used instead of a 300-mL tall beaker. However, the volumetric flask used should be distinguished as an extraction flask and should not be used for the other purposes. Additionally, “cover with a watch glass” in b) is replaced by “place a funnel”. Skip “transfer to a 250-mL volumetric flask with water” in the procedure in c).

Comment 3  The procedure in \((4.1.2)\) is the same as the procedure in \((4.1.1)\) in \(4.3.3.a\).

(4.2) Coloring: Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 1 mg as \(\text{B}_2\text{O}_3\)) to a 100-mL volumetric flask.

b) Add 25 mL of ethylenediamine tetraacetate solution, and then add 10 mL of ammonium acetate solution.

c) Add 10 mL of azomethine-H solution, and further add water up to the marked line, and then leave at rest for about 2 hours.

Comment 4  If formaldehyde processed urea, a large quantity of aluminum, copper, iron zinc, organic matters, etc. coexists to affect quantification, transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 1 mg as \(\text{B}_2\text{O}_3\), no more than 10 mL of the solution) to a 100-mL separatory funnel, add 10 mL of hydrochloric acid (1+3), add water to about 20 mL and add 20 mL of 2-ethyl-1,3-hexanediol–4-methyl-2-pentanone (1+9) to shake to mix with a shaking apparatus for about 1 minute. After allowing to stand still, remove the lower layer (aqueous phase) and add 20 mL of sodium hydroxide (20 mg/L) to shake to mix with a shaking apparatus for about 1 minute. After allowing to stand still, transfer the lower layer (aqueous phase) to a 100-mL volumetric flask, add a few drops of phenolphthalein solution (1 g/100mL) to neutralize with hydrochloric acid (1+3) until the color of the solution becomes achromatic, and conduct the procedure in \((4.2)\) b).

Comment 5  Water-soluble boron can be measured simultaneously with citrate soluble boron by adding 15 mL of citric acid solution before the procedure in \((4.2)\) b).

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.

   Detection wavelength: 415 nm

b) Calibration curve preparation

1) Transfer 1 mL - 20 mL of boron standard solution (\(\text{B}_2\text{O}_3 0.05 \text{mg/mL}\)) to 100-mL volumetric flasks step-by-step.

2) Conduct the same procedures as \((4.2)\) b) to make \(\text{B}_2\text{O}_3 0.05 \text{mg/100 mL} - 1 \text{mg/100 mL}\) of the boron standard solutions for the calibration curve preparation.

3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank test solution for the calibration curve preparation.

4) Measure absorbance at a wavelength of 415 nm of the boron standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.

5) Prepare a curve for the relationship between the boron concentration and absorbance of the boron standard solutions for the calibration curve preparation.
c) **Sample measurement**
1) Regarding the solution in (4.2) b), measure absorbance by the same procedure as b) 4).
2) Obtain the boron (B₂O₃) content from the calibration curve, and calculate water-soluble boron (W-B₂O₃) in the analytical sample.

**Comment 6** Water-soluble boron can be measured simultaneously with citrate-soluble boron by adding 15 mL of citric acid solution before the procedure in (4.3) b) 2).

**Comment 7** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 10 % (mass fraction) and 0.05 % (mass fraction) were 101.8 % and 107.1 % as water-soluble boron (W-B₂O₃) respectively. Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).

**References**

(5) **Flow sheet for water-soluble boron**: The flow sheet for water-soluble boron in fertilizers is shown below:

<table>
<thead>
<tr>
<th>2.5 g analytical sample</th>
<th>Weigh to the order of 1 mg into a 300-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>← About 200 mL of Water</td>
<td>Heating</td>
</tr>
<tr>
<td>Cooling</td>
<td>Cover with a watch glass, boil for 15 minutes</td>
</tr>
<tr>
<td>Transfer</td>
<td>Immediately</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Water, 250-mL volumetric flask</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for water-soluble boron in fertilizers (Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>← 25 mL of ethylenediamine tetraacetate solution</td>
</tr>
<tr>
<td></td>
<td>← 10 mL of ammonium acetate solution</td>
</tr>
<tr>
<td></td>
<td>← 10 mL of azomethine-H solution</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>Leaving at rest</td>
<td>For about 2 hours</td>
</tr>
<tr>
<td>Measurement</td>
<td>Spectrophotometer (415 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for water-soluble boron in fertilizers (Measurement procedure)
4.8.2.b  ICP Optical Emission Spectrometry

(1)  Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.8.2.b-2017 or W-B.b-1.

Extract by adding water to an analytical sample, further dilute the filtered solution, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the boron at a wavelength of 249.773 nm to obtain water-soluble boron (W-B\(_2\)O\(_3\)) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2)  Reagent, etc.: Reagents and water are as shown below.

   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Boron standard solution (B\(_2\)O\(_3\) 2.5 mg/mL)\(^{(1)}\): After leaving boric acid specified in JIS K 8863 at rest in a desiccator for about 24 hours to dry, weigh 4.441 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   d) Boron standard solution (B\(_2\)O\(_3\) 0.1 mg/mL)\(^{(1)}\): Transfer 4 mL of boron standard solution (B\(_2\)O\(_3\) 2.5 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line\(^{(2)}\).
   e) Boron standard solution (B\(_2\)O\(_3\) 2 µg/mL - 16 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 2 mL - 16 mL of boron standard solution (B\(_2\)O\(_3\) 0.1 mg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line\(^{(2)}\).
   f) Boron standard solution (B\(_2\)O\(_3\) 0.2 µg/mL - 2 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 2 mL - 20 mL of boron standard solution (B\(_2\)O\(_3\) 10 µg/mL) to a 100-mL volumetric flask step by step and add hydrochloric acid (1+23) up to the marked line\(^{(2)}\).
   g) Blank test solution for the calibration curve preparation \(^{(1)}\): Hydrochloric acid (1+23) used in the procedures in d), e) and f)\(^{(2)}\).

Note  (1) This is an example of preparation; prepare an amount as appropriate.
   (2) When preserving, use a container, which can be sealed tightly, made of material such as PTFE that boron hardly elutes

Comment 1 Instead of the boron standard solution in (2), a boron standard solution for the calibration curve preparation can be prepared by using a boron standard solution (B 1 mg/mL or 10 mg/mL) traceable to National Metrology. In this case, calculate water-soluble boron (W-B\(_2\)O\(_3\)) in the analytical sample by multiplying the concentration (B) of a boron standard solution for calibration curve preparation or a measurement value (B) obtained in (4.2) by a conversion factor (3.2199).

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in d) and e) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a boron standard solution for the calibration curve in the concentration range which is suitable to a device used.
(3) **Apparatus and instruments:** Apparatus and instruments are shown below.
   a) **ICP Optical Emission Spectrometry:** An Optical Emission Spectrometer specified in JIS K 0116.
      1) **Gas:** Argon gas specified in JIS K 1105 of no less than 99.5% (volume fraction) in purity

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.
   a) Weigh 1 mg of an analytical sample (3) to the order of 1 mg, and put it in a 100-mL volumetric flask.
   b) Add about 50 mL of water, shake to mix and add water up to the marked line.
   c) Filter with Type 3 filter paper to make a sample solution.

**Note** (3) The sampling amount of the analytical sample is 10 g when there is less boron content in the fertilizers such as a home garden-use fertilizer.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.
   a) **Measurement conditions for the ICP Optical Emission Spectrometer:** Set up the measurement conditions for the ICP Optical Emission Spectrometer considering the following:
      Analytical line wavelength: 249.773 nm
   b) **Calibration curve preparation**
      1) Spray the boron standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 249.773 nm.
      2) Prepare a curve for the relationship between the boron concentration and the indicated value of the boron standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation.
   c) **Sample measurement**
      1) Transfer a predetermined amount of the sample solution (the equivalents of 0.02 mg - 1.6 mg as B$_2$O$_3$) to a 100-mL volumetric flask.
      2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
      3) Subject to the same procedure as in b) 1) to read the indicated value.
      4) Obtain the boron content from the calibration curve, and calculate water-soluble boron (W-B$_2$O$_3$) in the analytical sample.

**Comment 4** Wash the sample injector of an ICP-OES sufficiently with water because boron easily causes the memory effect.

**Comment 5** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

**Comment 6** The comparison of the measurement value ($y_i$: 0.013 % (mass fraction) - 0.530 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of Azomethine-H method was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = -0.0041 + 0.986x$, and its correlation coefficient ($r$) was 0.999. Additionally, additive recovery testing was conducted using a fluid mixed fertilizer (1 brand) and a home garden-use mixed
fertilizer (1 brand) and a liquid micro element mixed fertilizer (1 sample). As a result, the mean recovery rates at additive level of 0.15 % (mass fraction) - 0.2 % and 0.01 % (mass fraction) were 95.5 % - 99.4 % and 96.5 % respectively. The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average 2)</th>
<th>Repeatability 4)</th>
<th>Intermediate precision 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.166</td>
<td>0.001</td>
<td>0.7</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (liquid)</td>
<td>7</td>
<td>0.0134</td>
<td>0.0001</td>
<td>1.0</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation

References


(5) Flow sheet: The flow sheet for water-soluble boron in fluid fertilizers is shown below:

- 1 g analytical sample
- Weigh to the order of 1 mg to a 100-mL volumetric flask
- → Water, about 50 mL
- Shaking to mix
- → Water (up to the marked line)
- Filtration
- Sample solution

Figure 1 Flow sheet for water-soluble boron in liquid fertilizers (Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>100-mL volumetric flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot</td>
<td>25 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td>(predetermined volume)</td>
<td>Water (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
<td>ICP-OES (249.773 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for water-soluble boron in liquid fertilizers (Measurement procedure)
4.9 Zinc
4.9.1 Total zinc
4.9.1.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type C and its symbol is 4.9.1.a-2017 or T-Zn.a-1.

Pretreat an analytical sample with incineration and nitric acid-hydrochloric acid (1+3), and then spray in an acetylene-air flame, and measure the atomic absorption with zinc at a wavelength of 213.9 nm to quantify the total zinc (T-Zn). In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Zinc standard solution (Zn 0.1 mg/mL): A zinc standard solution (Zn 0.1 mg/mL) traceable to National Metrology.
   e) Zinc standard solutions (Zn 0.5 µg - 5 µg/mL) for the calibration curve preparation: Transfer 2.5 mL - 25 mL of zinc standard solution (Zn 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in e).

Note  (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the zinc standard solution in (2), a zinc standard solution for the calibration curve preparation can be prepared by using a zinc standard solution (Zn 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction function.
   1) Light source: A zinc hollow cathode lamp (when the continuous source method as the background correction method is used, the light source is a deuterium lamp.)
   2) Gas: Gas for heating by flame
      (i) Fuel gas: acetylene
      (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can be adjusted to 450 °C ± 5 °C.
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

Note  (2) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char.
c) Ignite at 450 °C ± 3 °C for 8 - 16 hours to incinerate \(^{(3)}\).

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass \(^{(5)}\), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) \(^{(6)}\) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

**Note**  (3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

(4) The watch glass can be removed.

(5) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

**Comment 2** Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.b, (4.1) of 4.10.1.a, (4.1) of 4.10.1.b, (4.1) of 5.3.a, (4.1) of 5.3.b, (4.1) of 5.4.a, (4.1) of 5.4.b, (4.1) of 5.5.a, (4.1) of 5.5.d, (4.1) of (5.6.a) and (4.1) of 5.6.b. In addition, the sample solution can be used in 4.2.1.a, 4.2.1.b, 4.3.1.a, 4.3.1.b and 4.5.1.a.

**Comment 4** The sample solution prepared in (4.1.3) in 4.2.1.a can also be used.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:

Analytical line wavelength: 213.9 nm

b) **Calibration curve preparation**

1) Spray the zinc standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 213.9 nm.

2) Prepare a curve for the relationship between the zinc concentration and the indicated value of the zinc standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Subject the sample solution \(^{(6)}\) to the same procedure as in b) 1) to read the indicated value.

2) Subject the blank test solution to the same procedure as in b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.

3) Obtain the zinc content from the calibration curve, and calculate the total zinc (T-Zn) in the analytical sample.

**Note**  (6) If there is a possibility that the zinc concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount with
hydrochloric acid (1+23).

**Comment 5** The zinc concentration in the analytical sample can also be corrected by subjecting the blank test solution to the same procedures as in 1) and 3) to obtain the zinc content in the blank test solution.

**Comment 6** Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 1.2 % (mass fraction) and 90 mg/kg are 99.5 % and 97.8 % as the total zinc (T-Zn) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this test method is about 5 mg/kg for solid fertilizers.

<table>
<thead>
<tr>
<th>Name of certified reference material fertilizer</th>
<th>Number of laboratories</th>
<th>Number of test days</th>
<th>Repeatability standard deviation ($s_r$) (mg/kg)</th>
<th>Intermediate precision relative standard deviation ($RSD_{I(T)}$) (%)</th>
<th>Reproducibility relative standard deviation ($RSD_R$) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-C-12</td>
<td>12</td>
<td>992</td>
<td>11</td>
<td>1.1</td>
<td>32</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Cold vapor atomic absorption spectrometry  
2) Average (the number of laboratory ($p$) × test days (2) × the number of replicate testing (3))  
3) Repeatability standard deviation  
4) Repeatability relative standard deviation  
5) Intermediate standard deviation  
6) Intermediate relative standard deviation  
7) Reproducibility standard deviation  
8) Reproducibility relative standard deviation

**References**


(5) **Flow sheet for total zinc**: The flow sheet for total zinc in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>5.00 g analytical sample</td>
</tr>
<tr>
<td>2.</td>
<td>Charring</td>
</tr>
<tr>
<td>3.</td>
<td>Incineration</td>
</tr>
<tr>
<td>4.</td>
<td>Standing to cool</td>
</tr>
<tr>
<td></td>
<td>← A small amount of water</td>
</tr>
<tr>
<td></td>
<td>← About 10 mL of nitric acid</td>
</tr>
<tr>
<td></td>
<td>← About 30 mL of hydrochloric acid</td>
</tr>
<tr>
<td>5.</td>
<td>Heating</td>
</tr>
<tr>
<td>6.</td>
<td>Heating</td>
</tr>
<tr>
<td>7.</td>
<td>Standing to cool</td>
</tr>
<tr>
<td></td>
<td>← 25 mL - 50 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td>8.</td>
<td>Heating</td>
</tr>
<tr>
<td>9.</td>
<td>Standing to cool</td>
</tr>
<tr>
<td>10.</td>
<td>Transfer</td>
</tr>
<tr>
<td></td>
<td>← Water (up to the marked line)</td>
</tr>
<tr>
<td>11.</td>
<td>Filtration</td>
</tr>
<tr>
<td>12.</td>
<td>Sample solution</td>
</tr>
<tr>
<td>13.</td>
<td>Measurement</td>
</tr>
</tbody>
</table>

Figure  Flow sheet for total zinc in fertilizers
ICP Optical Emission Spectrometry

1) Summary
The testing method is applicable to sludge fertilizers, etc. This testing method is classified as Type D and its symbol is 4.9.1.b-2017 or T-Zn.b-1.

Pretreat an analytical sample with incineration, nitric acid–hydrochloric acid (1+3), introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the emission with zinc at a wavelength of 206.191 nm to obtain the total zinc (T-Zn) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Zinc standard solution (Zn 0.1 mg/mL): A zinc standard solution (Zn 0.1 mg/mL) traceable to National Metrology.
   e) Zinc standard solution (Zn 25 µg/mL): Dilute a predetermined amount of zinc standard solution (Zn 0.1 mg/mL) with hydrochloric acid (1+23) to prepare a zinc standard solution (Zn 25 µg/mL).

   Note (1) This is an example of preparation; prepare an amount as appropriate.

   Comment 1 Instead of the zinc standard solution in (2), a zinc standard solution for the calibration curve preparation can be prepared by using a zinc standard solution (Zn 1 mg/mL or 10 mg/mL) traceable to National Metrology.

3) Instruments: Instruments are as shown below:
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity
      b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
      c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 ºC.

4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char (2).
   c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (2).
   d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
   e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
   f) Slightly move the watch glass (3), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
   g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (4) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
   h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
   i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the
Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

The watch glass can be removed.

Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in b), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 2 Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) Measurement: Conduct measurement (Standard Addition Method) according to JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

- **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
  - Analytical line wavelength: 206.191 nm

- **Calibration curve preparation and sample measurement**
  1) Put 5mL of sample solution to three 10-mL volumetric flasks respectively.
  2) Add 2mL and 4 mL of zinc standard solution (0.25 μg/mL) to volumetric flasks of 1) above, then add hydrochloric acid (1+23) up to the marked line to make a sample solution of Standard Addition Method.
  3) Add hydrochloric acid (1+23) to the marked line of the remaining volumetric flask of 1) above to make a sample solution without a standard solution.
  4) Spray the sample solution of Standard Addition Method and the sample solution without a standard solution into the induction plasma, and read the indicated value at a wavelength of 206.191 nm.
  5) Transfer 5 mL of blank test solution to a 10-mL volumetric flask, conduct the same procedures as in 3) - 4) to read the indicated value, and correct the indicated value obtained from the respective sample solutions.
  6) Prepare a curve for the relationship between the added zinc concentration and the corrected indicated value of the sample solution for Standard Addition Method and the sample solution without a standard solution.
  7) Obtain the zinc content from the intercept of the calibration curve to calculate the total zinc (T-Zn) in the analytical sample.

Comment 4 The total zinc (T-Zn) in the analytical sample can also be corrected by subjecting the blank test solution to the same procedures as in b) 1) - b) 4) and b) 6) - b) 7) to obtain the zinc content in the blank test solution.

Comment 5 The simultaneous measurement of multiple elements by an ICP-OES is available. In this case, transfer a pre-determined amount of copper standard solution (Cu 0.1 mg/mL, 1 mg/mL or 10 mg/mL), zinc standard solution (Zn 0.1 mg/mL, 1 mg/mL or 10 mg/mL), cadmium standard solution (Cd 0.1 mg/mL, 1 mg/mL or 10 mg/mL), nickel standard solution (Ni 0.1 mg/mL, 1 mg/mL or 10 mg/mL), chromium standard solution (Cr 0.1 mg/mL, 1 mg/mL or 10 mg/mL) and lead standard solution (Pb 0.1 mg/mL) into the volumetric flask respectively.
mg/mL, 1 mg/mL or 10 mg/mL) to a volumetric flask to mix, add hydrochloric acid (1+5) to make acid concentration 0.5 mol/L and further add water up to the marked line to prepare a primary mixed standard solution. Transfer a pre-determined amount of primary mixed standard solution to a volumetric flask, add hydrochloric acid (1+23) up to the marked line to prepare a mixed standard solution for addition within the concentration range in Table 1. The measurement wavelengths of respective elements are according to Table 1.

In addition, the additive amount of mixed standard solution for addition and the additive concentrations of respective elements in a sample solution are shown in the Table below.

<table>
<thead>
<tr>
<th>Test item</th>
<th>Concentration of mixed standard solution for addition (µg/mL)</th>
<th>Additive concentration of element in sample solution (µg/mL)</th>
<th>Measurement wavelength (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total zinc</td>
<td>Zn 25</td>
<td>0 2 4</td>
<td>206.191</td>
</tr>
<tr>
<td>Total copper</td>
<td>Cu 25</td>
<td>0 2 4</td>
<td>324.754</td>
</tr>
<tr>
<td>Cadmium</td>
<td>Cd 0.25</td>
<td>0 0.05 0.1</td>
<td>228.802</td>
</tr>
<tr>
<td>Nickel</td>
<td>Ni 2.5</td>
<td>0 0.5</td>
<td>231.604</td>
</tr>
<tr>
<td>Chromium</td>
<td>Cr 2.5</td>
<td>0 0.5</td>
<td>205.552</td>
</tr>
<tr>
<td>Lead</td>
<td>Pb 2.5</td>
<td>0 0.5</td>
<td>220.351</td>
</tr>
</tbody>
</table>

1) Additive amount of mixed standard solution for addition

Comment 6 The comparison of the measurement value ($x_i$: 65.0 mg/kg - 3310 mg/kg) of ICP Optical Emission Spectrometry and the measurement value ($y_i$) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (49 samples). As a result, a regression equation was $y = -47.6 + 1.080x$ and its correlation coefficient ($r$) was 0.995. Triplicates measurement for each one sample of sewage sludge fertilizer, human waste sludge fertilizer, industrial sludge fertilizer, mixed sludge fertilizer, calcined sludge fertilizers and composted sludge fertilizer was conducted. As a result, a repeatability obtained was 0.1 % - 2.3 % as a relative standard deviation. Additionally, the minimum limit of quantification of this testing method is about 8 mg/kg.

References
(5) **Flow sheet for total zinc**: The flow sheet for total zinc in fertilizers is shown below:

![Flow sheet for total zinc in fertilizers (Extraction procedure)](image1)

![Flow sheet for total zinc in fertilizers (Measurement procedure)](image2)
4.9.2 Water-soluble zinc
4.9.2.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers that indicate zinc content as a response modifier. This testing method is classified as Type D and its symbol is 4.9.2.a-2017 or W-Zn.a-1.

Extract by adding water to an analytical sample, spray in an acetylene–air flame and measure the atomic absorption with zinc at a wavelength of 213.9 nm to obtain water-soluble zinc (W-Zn) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   c) Zinc standard solution (Zn 0.1 mg/mL): A zinc standard solution (Zn 0.1 mg/mL) traceable to National Metrology.
   d) Zinc standard solutions (Zn 0.5 µg - 5 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of zinc standard solution (Zn 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   e) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in d).

Note  (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the zinc standard solution in (2), a zinc standard solution for the calibration curve preparation can be prepared by using a zinc standard solution (Zn 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (2) function.
      1) Light source: A zinc hollow cathode lamp (when the continuous source method as the background correction method is used, the light source is a deuterium lamp.)
      2) Gas: Gas for heating by flame
         (i) Fuel gas: acetylene
         (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

Note  (2) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Powdery test sample
   a) Weigh 5.00 g of an analytical sample, and put it in a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

Comment 2 In the procedure in (4.1.1) a), it is also allowed to weigh 2.50 g of the analytical
sample and transfer to a 250-mL volumetric flask.

**Comment 3** The procedure in (4.1.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.1.2) **Fluid test sample**

a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

**Comment 4** The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.4.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   - Analytical line wavelength: 213.9 nm

b) **Calibration curve preparation**
   1) Spray the zinc standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 213.9 nm.
   2) Prepare a curve for the relationship between the zinc concentration and the indicated value of the zinc standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 0.5 mg as Zn) to a 100-mL volumetric flask.
   2) Add about 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the zinc content from the calibration curve, and calculate the water-soluble zinc (W-Zn) in the analytical sample.

**Comment 5** Recovery testing was conducted to evaluate trueness using a preparation sample (solid). As a result, the average rate of recovery at the content level of 10 % (mass fraction), 2 % (mass fraction) and 0.01 % (mass fraction) are 101.6 %, 101.9 % and 98.9 % as water-soluble zinc (W-Zn) respectively. In addition, recovery testing was conducted using a preparation sample (fluid). As a result, the average rate of recovery at the content level of 1 % (mass fraction), 0.05 % (mass fraction) and 20 mg/kg are 99.6 %, 100.4 % and 100.6 % as water-soluble zinc respectively. The results of the repeatability tests on different days using a fluid mixed fertilizer and a liquid microelement mixed fertilizer to evaluate the extract precision of fluid fertilizers were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 10 mg/kg (solid fertilizers) and 0.9 mg/kg (fluid fertilizers).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average (^{2)})</th>
<th>Repeatability (^{3)})</th>
<th>Intermediate precision (^{4)})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>1.28</td>
<td>0.01</td>
<td>0.4</td>
</tr>
<tr>
<td>Liquid microelement mixed fertilizers</td>
<td>7</td>
<td>0.230</td>
<td>0.001</td>
<td>0.5</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \(T\) \(\times\) the number of duplicate testing \(2\))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

(5) **Flow sheet for water-soluble zinc:** The flow sheet for water-soluble zinc in fertilizers is shown below:

```
5.00 g analytical sample (powdery) 500-mL volumetric flask.
← Water, About 400 mL
Shaking to mix Rotary shaker (30 - 40 revolutions/min), for 30 minutes
← Water (up to the marked line)
Filtration Type 3 filter paper
Sample solution
```

Figure 1-1 Flow sheet for water-soluble zinc in fertilizers (Extraction procedure(4.1.1))
1.00 g analytical sample (fluid) → 100-mL volumetric flask.

→ Water, About 50 mL

Shaking to mix

→ Water (up to the marked line)

Filtration

Type 3 filter paper

Sample solution

Figure 1-2 Flow sheet for water-soluble zinc in fertilizers
(Extraction procedure(4.1.2))

Sample solution

Aliquot (predetermined amount) → 100-mL volumetric flask

→ 25 mL of hydrochloric acid (1+5)

→ Water (up to the marked line)

Measurement Atomic absorption spectrometer (213.9 nm)

Figure 2 Flow sheet for water-soluble zinc in fertilizers
(Measurement procedure)
4.9.2.b ICP Optical Emission Spectrometry

(1) Summary

This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.9.2.b-2017 or W-Zn.b-1.

Extract by adding water to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the zinc at a wavelength of 213.856 nm to obtain water-soluble zinc (W-Zn) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Zinc standard solution (Zn 1 mg/mL): A zinc standard solution (Zn 1 mg/mL) traceable to National Metrology.
   d) Zinc standard solution (Zn 0.1 mg/mL)\(^{(1)}\): Transfer 10 mL of zinc standard solution (Zn 1 mg/mL) to a 100-mL flask and add hydrochloric acid (1+23) up to the marked line.
   e) Zinc standard solutions (Zn 1 µg - 20 µg/mL) for the calibration curve preparation\(^{(1)}\): Transfer 1 mL - 20 mL of zinc standard solution (Zn 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Zinc standard solutions (Zn 0.1 µg - 1 µg/mL) for the calibration curve preparation\(^{(1)}\): Transfer 1 mL - 10 mL of zinc standard solution (Zn 10 µg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation\(^{(1)}\): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the zinc standard solution in (2), a zinc standard solution for the calibration curve preparation can be prepared by using a zinc standard solution (Zn 10 mg/mL) traceable to National Metrology.

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, in case of using the ICP-OES of an axial observation mode, it is recommended to prepare a zinc standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1.00 g\(^{(2)}\) of an analytical sample, and put it in a 100-mL volumetric flask.
   b) Add about 50 mL of water, shake to mix and add water up to the marked line.
c) Filter with Type 3 filter paper to make a sample solution.

Note (2) The sampling amount of the analytical sample is 10 g when the content in the sample is less than 0.01 % (mass fraction) as water-soluble zinc.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
   Analytical line wavelength: 213.856 nm

b) Calibration curve preparation
   1) Spray the zinc standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 213.856 nm.
   2) Prepare a curve for the relationship between the zinc concentration and the indicated value of the zinc standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Zn) to a 100-mL volumetric flask.
   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the zinc content from the calibration curve, and calculate the water-soluble zinc (W-Zn) in the analytical sample.

Comment 4 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

Comment 5 The comparison of the measurement value ($y; 0.0109 \% \text{ (mass fraction)} - 0.0827 \% \text{ (mass fraction)}$) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of flame atomic absorbance spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = -0.0007 + 0.984x$, and its correlation coefficient ($r$) was 0.998. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery at additive level of 0.01 % (mass fraction) and 0.1 % (mass fraction) were 91.6 % and 95.9 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Test days of intermediate precision</th>
<th>Average (T) (%)</th>
<th>Repeatability standard deviation (%)</th>
<th>Intermediate relative standard deviation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.0677</td>
<td>0.0004</td>
<td>0.6</td>
<td>0.0005</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (liquid)</td>
<td>7</td>
<td>0.0107</td>
<td>0.0003</td>
<td>2.3</td>
<td>0.0004</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) × the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation

References

(5) Flow sheet for water-soluble zinc: The flow sheet for water-soluble zinc in fluid fertilizers is shown below:

1.00 g analytical sample

Weigh to the order of 1 mg to a 100-mL volumetric flask

← Water, about 50 mL

Shaking to mix

← Water (up to the marked line)

Filtration

Sample solution

Figure 1 Flow sheet for water-soluble zinc in liquid fertilizers (Extraction procedure)

Sample solution

Aliquot (predetermined volume)

100-mL volumetric flask

← 25 mL of hydrochloric acid (1+5)

← Water (up to the marked line)

Measurement

ICP-OES (213.856 nm)

Figure 2 Flow sheet for water-soluble zinc in liquid fertilizers (Measurement procedure)
4.10 Copper
4.10.1 Total copper
4.10.1.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type C and its symbol is 4.10.1.a-2017 or T-Cu.a-1.

Pretreat an analytical sample with incineration and nitric acid–hydrochloric acid (1+3), and then spray in an acetylene–air flame, and measure the atomic absorption with copper at a wavelength of 324.8 nm to obtain the total copper (T-Cu) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Copper standard solution (Cu 0.1 mg/mL): A copper standard solution (Cu 0.1 mg/mL) traceable to National Metrology.
   e) Copper standard solutions (Cu 0.5 µg - 5 µg/mL) for the calibration curve preparation
      (1): Transfer 2.5 mL - 25 mL of copper standard solution (Cu 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Blank test solution for the calibration curve preparation
      (1): Hydrochloric acid (1+23) used in the procedures in e).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the copper standard solution in (2), a copper standard solution for the calibration curve preparation can be prepared by using a copper standard solution (Cu 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction function.
   1) Light source: A copper hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
   2) Gas: Gas for heating by flame
      (i) Fuel gas: acetylene
      (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

Note (2) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char.

c) Ignite at 450 °C ± 3 °C for 8 - 16 hours to incinerate (3).
d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
f) Slightly move the watch glass (5), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (6) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note (3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.
(4) The watch glass can be removed.
(5) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 2 Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.
Comment 3 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.
Comment 4 The sample solution prepared in (4.1.2) in 4.2.1.a can also be used.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.
a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 324.8 nm
b) Calibration curve preparation
   1) Spray the copper standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 324.8 nm.
   2) Prepare a curve for the relationship between the copper concentration and the indicated value of the copper standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
c) Sample measurement
   1) Subject the sample solution (6) to the same procedure as in b) 1) to read the indicated value.
   2) Subject the blank test solution to the same procedure as in b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.
   3) Obtain the copper content from the calibration curve, and calculate the total copper (T-Cu) in the analytical sample.

Note (6) If there is a possibility that the copper concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount with hydrochloric acid (1+23).
Comment 5 The copper concentration in the analytical sample can also be corrected by subjecting the blank test solution to the same procedures as in 1) and 3) to obtain the copper content in the blank test solution.

Comment 6 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 0.15 % (mass fraction) and 0.03 % (mass fraction) are 100.4 % and 99.6 % as total copper (T-Cu) respectively.

The results of the collaborative study to determine a certified reference material fertilizer were analyzed by using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this test method is about 4 mg/kg.

<table>
<thead>
<tr>
<th>Name of certified reference laboratory</th>
<th>Number of laboratory</th>
<th>Average (mg/kg)</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
<th>Reproducibility</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>s_r^1 (mg/kg)</td>
<td>RSD_r^2 (%)</td>
<td>s_{I(T)}^3 (mg/kg)</td>
<td>RSD_{I(T)}^4 (%)</td>
</tr>
<tr>
<td>FAMIC-C-12</td>
<td>11</td>
<td>583</td>
<td>7</td>
<td>11</td>
<td>1.9</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Cold vapor atomic absorption spectrometry
2) Average (the number of laboratory × test days × the number of replicate testing)
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Intermediate standard deviation
6) Intermediate relative standard deviation
7) Reproducibility standard deviation
8) Reproducibility relative standard deviation

References
Flow sheet for total copper: The flow sheet for total copper in fertilizers is shown below:

- **5.00 g analytical sample**
- 200-mL - 300-mL tall beaker.
- **Charring**
- Heat gently
- **Incineration**
- Ignite at 450 ºC ± 5 ºC, 8 - 16 hours
- **Standing to cool**
- Room temperature
- ← A small amount of water
- ← About 10 mL of nitric acid
- ← About 30 mL of hydrochloric acid
- **Heating**
- Cover with a watch glass, and digest
- **Heating**
- Slightly move a watch glass to remove acid
- **Standing to cool**
- Room temperature
- ← 25 mL - 50 mL of hydrochloric acid (1+5)
- **Heating**
- Cover with a watch glass, and dissolve
- **Standing to cool**
- Room temperature
- **Transfer**
- 100-mL - 200-mL volumetric flask, water
- ← Water (up to the marked line)
- **Filtration**
- Type 3 filter paper
- **Sample solution**
- **Measurement**
- Atomic absorption spectrometer (324.8nm)

Figure Flow sheet for total copper in fertilizers
4.10.1.b ICP Optical Emission Spectrometry

(1) Summary

The test method is applicable to sludge fertilizers, etc. This testing method is classified as Type D and its symbol is 4.10.1.b-2017 or T-Cu.b-1.

Pretreat an analytical sample with incineration, nitric acid–hydrochloric acid (1+3), introduce it to ICP Optical Emission Spectrometry (“ICP-OES”) and measure the emission with copper at a wavelength of 324.754 nm to quantify the total copper (T-Cu) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Hydrochloric acid:

d) Copper standard solution (Cu 0.1 mg/mL): A copper standard solution (Cu 0.1 mg/mL) traceable to National Metrology.

e) Copper standard solutions (Cu 25 µg/mL) (1): Dilute a predetermined amount of copper standard solution (Cu 0.1 mg/mL) with hydrochloric acid (1+23) to prepare a copper standard solution (Cu 25 µg/mL). 

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the copper standard solution in (2), a copper standard solution for the calibration curve preparation can be prepared by using a copper standard solution (Cu 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:


1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

b) Electric furnace: An electric furnace that can keep the test temperature at 450 ºC ± 5 ºC.

c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 ºC.

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char (2).

c) Ignite at 450 ºC ± 5 ºC for 8 - 16 hours to incinerate (2).

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass (3), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (4) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.
Note  (2) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.
(3) The watch glass can be removed.
(4) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 2 Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.
a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
Analytical line wavelength: 324.754 nm
b) Calibration curve preparation and sample measurement
1) Put 5mL of sample solution to three 10-mL volumetric flasks respectively.
2) Add 2mL and 4 mL of copper standard solution (2.5 μg/mL) to volumetric flasks of 1) above, then add hydrochloric acid (1+23) to the marked line to make a sample solution of Standard Addition Method.
3) Add hydrochloric acid (1+23) to the marked line of the remaining volumetric flask of 1) above to make a sample solution without a standard solution.
4) Spray the sample solution of Standard Addition Method and the sample solution without a standard solution into the induction plasma, and read the indicated value at a wavelength of 324.754 nm.
5) Transfer 5 mL of blank test solution to a 10-mL volumetric flask, conduct the same procedures as in 3) - 4) to read the indicated value, and correct the indicated value obtained from the respective sample solutions.
6) Prepare a curve for the relationship between the added copper concentration and the corrected indicated value of the sample solution for Standard Addition Method and the sample solution without a standard solution.
7) Obtain the copper content from the intercept of the calibration curve to calculate the total copper (T-Cu) in the analytical sample.

Comment 4 The copper concentration in the analytical sample can also be corrected by subjecting the blank test solution to the same procedures as in b) 1) - b) 4) and b) 6) - b) 7) to obtain the copper content in the blank test solution.

Comment 5 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.9.1.b Comment 5.

Comment 6 The comparison of the measurement value \(x_i:12.0 \text{ mg/kg} - 1400 \text{ mg/kg}\) of ICP Optical Emission Spectrometry and the measurement value \(y_i\) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (49 samples). As a result, a regression equation was \(y = -5.5 + 1.062x\) and its correlation coefficient \((r)\) was 0.997. Triplicates measurement for each one sample of
sewage sludge fertilizer, human waste sludge fertilizer, industrial sludge fertilizer, mixed sludge fertilizer, calcined sludge fertilizers and composted sludge fertilizer was conducted. As a result, a repeatability obtained was 0.6 % - 1.8 % as a relative standard deviation. Additionally, the minimum limit of quantification of this testing method is about 3 mg/kg.

References


(5) Flow sheet for total copper: The flow sheet for total copper in fertilizers is shown below:

<table>
<thead>
<tr>
<th>5.00 g analytical sample</th>
<th>200-mL - 300-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>Ignite at 450 °C ± 5 °C, 8 - 16 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← A small amount of water</td>
<td></td>
</tr>
<tr>
<td>← About 10 mL of nitric acid</td>
<td></td>
</tr>
<tr>
<td>← About 30 mL of hydrochloric acid</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and digest</td>
</tr>
<tr>
<td>Heating</td>
<td>Slightly move a watch glass to remove acid</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← 25 mL - 50 mL of hydrochloric acid (1+5)</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and dissolve</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Transfer</td>
<td>100-mL - 200-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for total copper in fertilizers (Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aliquot 5 mL</strong></td>
</tr>
<tr>
<td>10-mL volumetric flask, 3 flasks</td>
</tr>
<tr>
<td>← 0, 2 and 4 mL of copper standard solution (25 μg/mL)</td>
</tr>
<tr>
<td>← Hydrochloric acid (1+23) (up to the marked line)</td>
</tr>
<tr>
<td><strong>Measurement</strong></td>
</tr>
<tr>
<td>ICP Optical Emission Spectrometer (324.754 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for total copper in fertilizers (Measurement procedure)
4.10.2 Water-soluble copper
4.10.2.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers that indicate copper content as a response modifier. This testing method is classified as Type D and its symbol is 4.10.2.a-2017 or W-Cu.a-1.
Extract by adding water to an analytical sample, spray in an acetylene–air flame and measure the atomic absorption with copper at a wavelength of 324.8 nm to obtain water-soluble copper (W-Cu) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   c) Copper standard solution (Cu 0.1 mg/mL): A copper standard solution (Cu 0.1 mg/mL) traceable to National Metrology.
   d) Copper standard solutions (Cu 0.5 µg - 5 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of copper standard solution (Cu 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   e) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in d).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the copper standard solution in (2), a copper standard solution for the calibration curve preparation can be prepared by using a copper standard solution (Cu 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (2) function.
   1) Light source: A copper hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
   2) Gas: Gas for heating by flame
      (i) Fuel gas: acetylene
      (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

Note (2) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Powdery test sample
   a) Weigh 5.00 g of an analytical sample, and put it in a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.
Comment 2 In the procedure in (4.1.1) a), it is also allowed to weigh 2.50 g of the analytical sample and transfer to a 250-mL volumetric flask.

Comment 3 The procedure in (4.1.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 4 The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 324.8 nm

b) Calibration curve preparation
   1) Spray the copper standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 324.8 nm.
   2) Prepare a curve for the relationship between the copper concentration and the indicated value of the copper standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 0.5 mg as Cu) to a 100-mL volumetric flask.
   2) Add about 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the copper content from the calibration curve, and calculate the water-soluble copper (W-Cu).

Comment 5 Recovery testing was conducted to evaluate trueness using a preparation sample (solid). As a result, the average rate of recovery at the content level of 10 % (mass fraction), 1 % (mass fraction) and 0.03 % (mass fraction) are 100.7 %, 99.4 % and 102.6 % as water-soluble copper (W-Cu) respectively. In addition, recovery testing was conducted using a preparation sample (fluid). As a result, the average rate of recovery at the content level of 1 % (mass fraction), 0.05 % (mass fraction) and 20 mg/kg are 98.8 %, 99.3 % and 101.4 % as water-soluble copper respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a liquid microelement mixed fertilizer to evaluate the extract precision of fluid fertilizers were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 10 mg/kg (solid fertilizers) and 3 mg/kg (fluid fertilizers).
### Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>$T^{1)}$</td>
<td>$\text{Average}^{2)}$</td>
</tr>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.0540</td>
<td>0.0003</td>
</tr>
<tr>
<td>Liquid microelement mixed fertilizers</td>
<td>7</td>
<td>0.0172</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) \times the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

### References


(5) **Flow sheet for water-soluble copper**: The flow sheet for water-soluble copper in fertilizers is shown below:

```
5.00 g analytical sample (powdery) 500-mL volumetric flask.
                        ← Water, About 400 mL
Shaking to mix            Rotary shaker (30 - 40 revolutions/min), for 30 minutes
                        ← Water (up to the marked line)
Filtration               Type 3 filter paper
                        ← Sample solution
```

Figure 1-1 Flow sheet for water-soluble copper in fertilizers
(Extraction procedure (4.1.1))
Figure 1-2 Flow sheet for water-soluble copper in fertilizers
(Extraction procedure (4.1.2))

1.00 g analytical sample (fluid) 100-mL volumetric flask.
← Water, About 50 mL
Shaking to mix
← Water (up to the marked line)
Filtration Type 3 filter paper
Sample solution

Figure 2 Flow sheet for water-soluble copper in fertilizers
(Measurement procedure)

Sample solution

Aliquot (predetermined amount) 100-mL volumetric flask
← 25 mL of hydrochloric acid (1+5)
← Water (up to the marked line)
Measurement Atomic absorption spectrometer (324.8 nm)
4.10.2.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.10.2.b-2017 or W-Cu.b-1.

Extract by adding water to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the copper at a wavelength of 327.396 nm to obtain water-soluble copper (W-Cu) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Copper standard solution (Cu 1 mg/mL): A copper standard solution (Cu 1 mg/mL) traceable to National Metrology.
   d) Copper standard solutions (Cu 0.1 mg/mL) for the calibration curve preparation (1):
      Transfer 10 mL of copper standard solution (Cu 1 mg/mL) to 100-mL volumetric flask, and add hydrochloric acid (1+23) to the marked line.
   e) Copper standard solutions (Cu 0.1 µg - 20 µg/mL) for the calibration curve preparation (1):
      Transfer 1 mL - 20 mL of copper standard solution (Cu 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Copper standard solutions (Cu 0.1 µg - 1 µg/mL) for the calibration curve preparation (1):
      Transfer 1 mL - 10 mL of copper standard solution (Cu 10 µg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation (1):
      Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the copper standard solution in (2), a copper standard solution for the calibration curve preparation can be prepared by using a copper standard solution (Cu 10 mg/mL) traceable to National Metrology.

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, in case of using the ICP-OES of an axial observation mode, it is recommended to prepare a copper standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample (2) to the order of 1 mg, and put it in a 100-mL volumetric
flask.

b) Add about 50 mL of water, shake to mix and add water up to the marked line.

c) Filter with Type 3 filter paper to make a sample solution.

**Note (2)** The sampling amount of the analytical sample is 10 g when the content in the sample is less than 0.01 % (mass fraction) as water-soluble copper.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

   Analytical line wavelength: 327.396 nm

b) **Calibration curve preparation**

   1) Spray the copper standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 327.396 nm.

   2) Prepare a curve for the relationship between the copper concentration and the indicated value of the copper standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Cu) to a 100-mL volumetric flask.

   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.

   3) Subject to the same procedure as in b) 1) to read the indicated value.

   4) Obtain the copper content from the calibration curve, and calculate the water-soluble copper (W-Cu).

**Comment 4** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d **Comment 4**.

**Comment 5** The comparison of the measurement value \( y_i; 0.00982 \% \) (mass fraction) - 0.0819 % (mass fraction) of ICP Optical Emission Spectrometry and the measurement value \( x_i \) of flame atomic absorbance spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was \( y = -0.0006 + 0.966x \), and its correlation coefficient \( (r) \) was 0.999. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery at additive level of 0.01 % (mass fraction) and 0.1 % (mass fraction) were 93.5 % and 95.3 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T(1)</td>
<td>Average(2)</td>
<td>Repeatability standard deviation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(%) (3)</td>
<td>s(4) (%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>RSD(5) (%)</td>
</tr>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.0643</td>
<td>0.0006</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (liquid)</td>
<td>7</td>
<td>0.00976</td>
<td>0.00006</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) x the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

(5) Flow sheet for water-soluble copper: The flow sheet for water-soluble copper in fluid fertilizers is shown below:

![Flow sheet for water-soluble copper in fertilizers (Extraction procedure)](image1)

![Flow sheet for water-soluble copper in fertilizers (Measurement procedure)](image2)
4.11 Organic carbon and carbon-nitrogen ratio

4.11.1 Organic carbon

4.11.1.a Dichromate oxidation

(1) Summary

This test method is applicable to sludge fertilizers and compost, etc. This testing method is classified as Type C and its symbol is 4.11.1.a-2017 or O-C.a-1.

Add a potassium dichromate-sulfuric acid solution to an analytical sample and heat to oxidize organic carbon with potassium dichromate. Quantify unconsumed potassium dichromate by oxidation-reduction titration to obtain organic carbon (O-C). This test method is also referred to as the Method of Tyulin. In addition, the performance of this testing method is shown in Comment 2.

(2) Reagent: Reagents are as shown below.


b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.2 mol/L ammonium iron (II) sulfate solution (1): Weigh 80 g of ammonium iron (II) sulfate hexahydrate specified in JIS K 8979 into a 2000-mL beaker, and add 1000 mL of sulfuric acid (1+50) to dissolve.

Standardization: Grind potassium dichromate reference material for volumetric analysis specified in JIS K 8005 in an agate mortar to powder, heat at 150 ºC ± 2 ºC for 1 hour, let it stand to cool in a desiccator, and then transfer about 1 g to a weighing dish, and weigh the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 100-mL volumetric flask, and add water up to the marked line to make the potassium dichromate standard solution (1)(2). On each day to use a 0.2 mol/L ammonium iron (II) sulfate solution, transfer 10 mL of the potassium dichromate standard solution to a 100-mL Erlenmeyer flask, add about 5 mL of sulfuric acid (1+2), and then conduct the procedures in (4.2) b) - c), and calculate the factor of a 0.2 mol/L ammonium iron (II) sulfate solution by the following formula:

Factor \((f)\) of 0.2 mol/L ammonium iron (II) sulfate solution

\[
W_1 \times (A/100) \times (6/294.18) \times (V_1/V_2) \times (1000/V_3)/C = (W_1 \times A/V_3) \times (30/294.18)
\]

\(W_1\): Mass (g) of potassium dichromate weighed

\(A\): Purity (% (mass fraction)) of potassium dichromate

\(V_1\): Volume (10 mL) of potassium dichromate solution transferred

\(V_2\): Constant volume (100 mL) of potassium dichromate solution

\(V_3\): Volume (mL) of 0.2 mol/L ammonium iron (II) sulfate solution needed for titration

\(C\): Set concentration (0.2 mol/L) of 0.2 mol/L ammonium iron (II) sulfate solution

d) Potassium dichromate-sulfuric acid solution (1): Weigh 40 g of potassium dichromate specified in JIS K 8517 to a 3000-mL beaker. Add 1000 mL of water to dissolve, and further add gradually 1000 mL of sulfuric acid while cooling and mixing.

e) N-Phenylanthranilic acid solution: Dissolve 0.2 g of N-phenylanthranilic acid of no less than 98 % (mass fraction) in purity and 0.2 g of sodium carbonate specified in JIS K 8625 in a small amount of water, and add water to make 100 mL.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(2) This corresponds to the standard potassium dichromate solution (0.2 M (1/6 K₂Cr₂O₇) solution) in 7.1 B 1) in the Official Methods of Analysis of Fertilizers (1992).
(3) **Apparatus and instruments**: Apparatus and instruments are shown below.

a) **Hot plate**: A hot plate whose surface temperature can be adjusted up to 250 ºC.

b) **Sample digestion flask**: A 100-mL borosilicate glass volumetric flask 100 mL (180 mm total height, 13 mm mouth diameter)

**Note** (3) Distinguish the volumetric flask used in digestion as a sample digestion flask and do not use it for any other purposes.

(4) **Test procedure**

(4.1) **Dichromate oxidation**: Conduct oxidation as follows:

a) Weigh 0.05 g of an analytical sample to the order of 0.1 mg (4), and transfer to a sample digestion flask.

b) Add 25 mL of potassium dichromate-sulfuric acid solution.

c) Heat on a hot plate at 200 ºC until organic matters are completely digested (5).

d) After immediate cooling is complete, precisely adjust to 100 mL with water to make a sample solution.

e) As a blank test, conduct the procedures in b) and d) using another sample digestion flask to prepare the blank test solution.

**Note** (4) Up to about 28 mg as organic carbon (O-C).

(5) Heat for no less than 1 hour after boiling

**Comment 1** Sample an analytical sample from a test sample prepared in 2.3.3 Grinding (3.1) by grinding with a mill until it completely passes through a sieve of 500 µm aperture or from a test sample prepared in 2.3.3 Grinding Comment 1.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below.

a) Transfer 20 mL of the sample solution to a 100-mL Erlenmeyer flask.

b) Add a 0.2 mol/L ammonium iron (II) sulfate solution drop-by-drop until the brown color of dichromate ion almost disappears from the sample solution.

c) Add about a 0.25 mL of N-phenylanthranilic acid solution (6), and titrate with a 0.2 mol/L ammonium iron (II) sulfate solution until the color of the solution changes from dark red-purple to blue-green.

d) Transfer 20 mL of the blank test solution to a 100-mL Erlenmeyer flask, and conduct the procedures in b) - e) to titrate.

e) Calculate the organic carbon (O-C) in the analytical sample by the following formula:

\[
\text{Organic carbon (mass fraction)}) = \frac{(V_6 - V_7) \times f \times (12.011/40)}{W_2 \times (100/1000)} \times (V_4/ V_5)
\]

\[
V_4: \text{Volume (mL) of 0.2 mol/L ammonium iron (II) sulfate solution needed for the titration of the blank test solution}
\]

\[
V_5: \text{Volume (mL) of 0.2 mol/L ammonium iron (II) sulfate solution needed for the titration of the sample solution}
\]

\[
C: \text{Set concentration (0.2 mol/L) of 0.2 mol/L ammonium iron (II) sulfate solution}
\]

\[
f: \text{Factor of 0.2 mol/L ammonium iron (II) sulfate solution}
\]

\[
V_6: \text{Predetermined volume (100 mL) of the sample solution and the blank test solution in (4.1) d)
\]
$V_7$: Transferred volume (20 mL) of the sample solution and the blank test solution subjected to titration in (4.2) a) and (4.2) d)

$W_2$: Mass (g) of the analytical sample

**Note** (6) About 5 drops with a 1-mL - 2-mL Komagome pipette. Add the same volume to the sample solution and the blank test solution.

**Comment 2** The results of the collaborative study to determine a certified reference material fertilizer were analyzed using a three-level nesting analysis of variance. Table 1 shows the calculation results of reproducibility, intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 1.5 % (mass fraction).

**Table 1** Analysis results of the collaborative study to determine a certified reference material fertilizer

<table>
<thead>
<tr>
<th>Name of certified reference material fertilizer</th>
<th>Number of laboratory $p^{1)}$</th>
<th>Average $2)^{2)}$</th>
<th>Repeatability $s_r^{4)}$ (%) $RSD_r^{5)}$ (%)</th>
<th>Intermediate precision $s_{IR}^{6)}$ (%) $RSD_{IR}^{7)}$ (%)</th>
<th>Reproducibility $s_R^{8)}$ (%) $RSD_R^{9)}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAMIC-C-12</td>
<td>12</td>
<td>20.2</td>
<td>0.3</td>
<td>1.5</td>
<td>0.5</td>
</tr>
</tbody>
</table>

1) The number of laboratories used for analysis conducting Dichromate oxidation
2) Average (the number of laboratory $(p)$ × test days $(2)$ × the number of replicate testing $(3)$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
8) Reproducibility standard deviation
9) Reproducibility relative standard deviation

**References**


5) **Flow sheet for organic carbon**: The flow sheet for organic carbon in sludge fertilizers and compost, etc. is shown below:
Weigh to the order of 0.1 mg into a sample digestion flask. (Up to about 28 mg as organic carbon.)

Immediately

Weigh to the order of 0.1 mg into a sample digestion flask.

25 mL potassium dichromate-sulfuric acid solution

Boil for about 1 hour, 200 °C

Water (fill up to 100 mL)

100-mL Erlenmeyer flask

0.05 g analytical sample

Heating

Cooling

20 mL aliquot

Sample solution

Figure 1 Flow sheet for organic carbon in sludge fertilizers and compost, etc. (Dichromate oxidation procedure)

Adding drop by drop

0.2 mol/L Ammonium iron (II) sulfate solution

(0.2 mol/L ammonium iron (II) sulfate solution (until the solution becomes blue-green)

(untill the brown color of dichromate ion almost disappears from the solution)

About 0.25 mL N-phenylantranilic acid solution

Figure 2 Flow sheet for organic carbon in sludge fertilizers and compost, etc. (Measurement procedure)
4.11.1.b Combustion method

(1) Summary
This testing method is applicable to compost and sludge fertilizers. This testing method is classified as Type B and its symbol is 4.11.1.b-2017 or O-C.b-1.
Drop hydrochloric acid (1+3) to an analytical sample and evaporate inorganic carbon as carbon dioxide, then thermally decompose carbon compounds using a total nitrogen-total carbon analyzer by the combustion method to measure carbon dioxide gas with a thermal conductivity detector and obtain organic carbon (O-C) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent: Reagents are as shown below.
   a) Sea sand: Particle diameter 425 µm - 850 µm
   b) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

Comment 1 Sea sand (particle diameter 425 µm - 850 µm) is commercially sold by FUJIFILM Wako Pure Chemical Co., Ltd. and YONEYAMA YAKUHIN KOGYO Co., Ltd.

(3) Instruments: Instruments are as shown below:
   a) Total nitrogen-total carbon analyzer by the combustion method: A total nitrogen-total carbon analyzer configured on the basis of the principle of the combustion method (modified Dumas’ method).
      1) Turn on the total nitrogen-total carbon analyzer by the combustion method (1), and adjust so that stable indicated values can be obtained.
         (i) Combustion gas: Oxygen having purity no less than 99.99% (volume percentage)
         (ii) Carrier gas: Helium having purity no less than 99.99% (volume percentage)
   b) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 °C.
   c) Drying apparatus: A drying apparatus that can be adjusted to 105 °C ± 2 °C.

Note (1) The setup of the program and the parameters of the analyzer are according to the specification and the operation method of the total nitrogen-total carbon analyzer by the combustion method used.

(4) Test procedures: Conduct measurement as shown below. However, confirm that there is no difference from the measured value of organic carbon obtained in advance according to 4.11.1.a by using an analytical sample.

(4.1) Hydrochloric acid treatment
   a) Weigh 0.05 g of an analytical sample to the order of 0.1 mg, and transfer to a container for combustion.
   b) Cover the analytical sample with about 0.2 g of sea sand and moisten the analytical sample by dropping a few drops of water.
   c) After dropping 0.5 mL - 0.7 mL of hydrochloric acid (1+3) little by little (2), drop about 0.3 mL of water (3)(4).
   d) Heat a container for combustion on a hot plate at 100 °C for 90 minutes to dry and harden it.
   e) Put the container for combustion at 105 °C ± 2 °C into a drying apparatus and heat for 30 minutes (5).
   f) After heating, let it stand to cool to make a test sample.

Note (2) The additive amount of hydrochloric acid (1+3) is merely a target. It is enough to make
the whole analytical sample come into contact with hydrochloric acid. Let it stand for a short time in the case of producing bubbles.

(3) In some cases, it is not necessary to add water due to the capacity of a container.
(4) Shake calmly the container for combustion to make the analytical sample come into complete contact with hydrochloric acid.
(5) Remove hydrochloric acid completely

Comment 2 Sample an analytical sample from a test sample prepared in 2.3.3 Grinding (3.1) by grinding with a mill until it completely passes through a sieve of 500 µm aperture or from a test sample prepared in 2.3.3 Grinding Comment 1.
Comment 3 When it is confirmed that, for example, the volatilization of hydrogen chloride is not detected by using a test paper, etc. and hydrochloric is completely removed in the procedure d), the procedure e) can be skipped.

4.2 Measurement: Specific measurement procedures are according to the operation method of a total nitrogen-total carbon analyzer by the combustion method.

a) Measurement conditions for the total nitrogen-total carbon analyzer by the combustion method: Set up the measurement conditions for the total nitrogen-total carbon analyzer considering the following:
   Combustion temperature: No less than 870 ºC

b) Calibration curve preparation
   1) Turn on the total nitrogen-total carbon analyzer by the combustion method (1), and adjust so that stable indicated values can be obtained.
   2) Weigh a predetermined amount of the standard for calibration curves (6) to the order of 0.1 mg into a combustion vessel.
   3) Insert the combustion vessel into the total nitrogen-total carbon analyzer by the combustion method, and read the indicated value.
   4) Conduct the procedure in 3) for another combustion vessel for a blank test, and read the indicated value.
   5) Prepare a curve for the relationship between the carbon content and the indicated value of the standard for calibration curves and the blank test for calibration curves.

c) Sample measurement
   1) Insert the combustion vessel containing the test sample to the total nitrogen-total carbon analyzer by the combustion method, and read the indicated value.
   2) Obtain the carbon content from the calibration curve, and calculate organic carbon in the analytical sample.

Note (6) Standard for calibration curves: DL-Aspartic acid (purity no less than 99 % (mass fraction)), EDTA (purity no less than 99 % (mass fraction)), hippuric acid (purity no less than 98 % (mass fraction)) or other reagents having equivalent purity recommended by the total nitrogen-total carbon analyzer by the combustion method used.

Comment 4 The comparison of the measurement value ($y_i$: 0.21 % (mass fraction) - 45.40 % (mass fraction)) of Dichromate oxidation was conducted to evaluate trueness using sludge fertilizers and compost (total 25 samples). As a result, a regression equation was $y = 0.004 + 1.009x$ and its correlation coefficient ($r$) were 0.999.
Table 1 shows results and analysis results from a collaborative study for test method validation.
Additionally, the minimum limit of quantification of this testing method is about
0.05 % (mass fraction).

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^2$ (%) $^3$</th>
<th>$s_r$ $^4$ (%) $^3$</th>
<th>RSD $^5_r$ (%)</th>
<th>$s_R$ $^6$ (%) $^3$</th>
<th>RSD $^7_R$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human waste sludge</td>
<td>8</td>
<td>34.96</td>
<td>0.07</td>
<td>0.2</td>
<td>0.62</td>
<td>1.8</td>
</tr>
<tr>
<td>Industrial sludge</td>
<td>8</td>
<td>15.13</td>
<td>0.20</td>
<td>1.3</td>
<td>0.42</td>
<td>2.8</td>
</tr>
<tr>
<td>Calcined sludge</td>
<td>9</td>
<td>9.45</td>
<td>0.17</td>
<td>1.8</td>
<td>0.38</td>
<td>4.0</td>
</tr>
<tr>
<td>Composted sludge</td>
<td>9</td>
<td>38.20</td>
<td>0.27</td>
<td>0.7</td>
<td>0.73</td>
<td>1.9</td>
</tr>
<tr>
<td>Compost</td>
<td>9</td>
<td>20.50</td>
<td>0.76</td>
<td>3.7</td>
<td>0.94</td>
<td>4.6</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis  
2) Mean ($n = number of laboratories \times number of samples (2)$)  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Reproducibility standard deviation  
7) Reproducibility relative standard deviation

References


(5) **Flow sheet for organic carbon**: The flow sheet for organic carbon in sludge fertilizers and compost, etc. is shown below:

- **0.05 g analytical sample**: Weigh to the order of 0.1 mg into a combustion vessel.
- **Cover the analytical sample with 0.2 g of sea sand and drop a few drops of water.**
- **Drop 0.5mL - 0.7mL of hydrochloric acid (1+3) little by little**
- **Drop 0.3mL of water**
- **Heating**: Heat to make it dry for 90 minutes, at 100 ºC
- **Drying**: Heat to make it dry for 30 minutes, at 105 ºC
- **Test sample**
- **Measurement**: Total nitrogen-total carbon analyzer by the combustion method

Figure Flow sheet for organic carbon by the combustion method
Reference: Chromatograms of the standard for calibration curves and an analytical sample are shown below:

1) Total carbon in a standard calibration curve (DL-Aspartic acid)

2) Total carbon in an analytical sample (sludge fertilizer)

Reference figures Chromatograms of organic carbon.

Measurement conditions for total nitrogen-total carbon analyzer by the combustion method

Combustion gas: Highly pure oxygen, purity no less than 99.99995 % (volume fraction), flow rate 200 mL/min
Carrier gas: Highly pure helium, purity no less than 99.9999 % (volume fraction), flow rate 80 mL/min
Separation column: A silica gel stainless column (length 1m)
Detector: Thermal conductivity detector (TCD)
Measurement cycle: Purge time = 60 seconds, circulation combustion time = 300 seconds, measurement time = 270 seconds
Current value of Detector: 160 mA
Temperature conditions: Reaction furnace temperature: 870 °C
Reaction furnace temperature: 600 °C
Column oven temperature: 70 °C
Detector temperature: 100 °C
4.11.2 Carbon-nitrogen ratio
4.11.2.a Calculation with organic carbon and total nitrogen

(1) Summary
This test method is applicable to compost and sludge fertilizers. This testing method is classified as Type A (Def-C) and its symbol is 4.11.2.a-2017 or C/N.a-1. Calculate carbon-nitrogen ratio (CN ratio) by dividing the organic carbon obtained in 4.11.1 by the total nitrogen obtained in 4.1.1.

(2) The calculation of carbon-nitrogen ratio
a) Calculate the carbon-nitrogen ratio (CN ratio) in an analytical sample by the following formula:

\[
\text{Carbon-nitrogen ratio in an analytical sample} = \frac{O-C}{T-N}
\]

O-C: Organic carbon (% (mass fraction)) \(^{(1)}\) in the analytical sample obtained in 4.11.1
T-N: Total nitrogen (% (mass fraction)) \(^{(1)}\) in the analytical sample obtained in 4.1.1

Note (1) O-C and T-N use raw data without rounding numerical value
4.12 Sulfur
4.12.1 Total sulfur content
4.12.1.a Potassium permanganate analysis

(1) Summary
This testing method is applicable to fertilizers mainly containing ferrous sulfate (iron (II) sulfate) (FeSO₄) among sulfur and its compound. This testing method is classified as Type D and its symbol is 4.12.1.a-2017 or T-S.a-1.
Dissolve an analytical sample in water and diluted sulfuric acid, add phosphoric acid and then titrate ferrous sulfate (iron (II) sulfate) (FeSO₄) by oxidation-reduction with a potassium permanganate solution to obtain the total sulfate (T-SO₃). In addition, the performance of this testing method is shown in Comment 1.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.
   c) Phosphoric acid: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.
   d) 0.02 mol/L potassium permanganate solution: Dissolve 3.16 g of potassium permanganate specified in JIS K 8247 in about 800 mL of water to boil and add water to make 1000 mL and leave at rest 1 - 2 day(s). Further filter with a funnel type glass filter (G4) and store in a colored bottle. Or, a reagent of equivalent quality (volumetric analysis grade) that is commercially available.

Standardization: Dry sodium oxalate of reference materials for volumetric analysis specified in JIS K 8005 at 200 ºC for 1 hour and let it stand to cool in a desiccator, and then put about 0.3 g into a weighing dish to measure the mass to the order of 0.1 mg. Add about 250 mL of sulfuric acid (1+20) cooled down to 25 ºC - 30 ºC after boiling and dissolve. Add about 40 mL of 0.02 mol/L potassium permanganate solution while gently shaking for about 1 minute. Heat up to 55 ºC - 60 ºC after the red color of a potassium permanganate solution disappears. Titrate with a 0.02 mol/L potassium permanganate solution while keeping the temperature and continue titrating until the color of the solution becomes light red (1). Calculate the factor of a 0.02 potassium permanganate solution by the following formula.

Factor ($f$) of 0.02 mol/L potassium permanganate solution  
$$f = W₁ \times (A/100) \times ((2/5)/133.999) \times ((1000/V₁)/C)$$  
$$f = W₁ \times (A/V₁) \times 1.4925$$

$W₁$: Mass (g) of sodium oxalate sampled  
$A$: Purity (% (mass fraction)) of sodium oxalate sampled  
$V₁$: Volume (mL) of 0.02 mol/L potassium permanganate needed for titration  
$C$: 0.02 mol/L potassium permanganate solution

Note  (1) The endpoint is reached when the color of solution keeps as it is for 30 seconds after coloring

(3) Instruments: Instruments are as shown below:
   a) Magnetic stirrer:

(4) Test procedure
(4.1) Measurement: Conduct measurement as shown below.
Weigh 0.5 g - 1 g of an analytical sample to the order of 0.1 mg, and transfer to a 200-mL tall beaker.

Add about 50 mL of water and about 15 mL of sulfuric acid (1+5) and shake with a magnetic stirrer to dissolve.

Immediately add about 1 mL of phosphate, and then titrate with a 0.02 mol/L potassium permanganate solution until the color of the solution becomes light red.

As a blank test, conduct the procedures in b) - c) using another 200-mL tall beaker to titrate.

Calculate the total sulfur content (T-SO$_3$) in an analytical sample by the following formula.

Total sulfur content (% (mass fraction)) = $(5 \times 0.02 \times f \times (V_2-V_3)/1000 \times 80.064)/W_2 \times 100$

$W_2$: Mass (g) of the sampled analytical sample  
$V_2$: Volume (mL) of 0.02 mol/L potassium permanganate needed for titration  
$V_3$: Volume (mL) of 0.02 mol/L potassium permanganate needed for titration  
$f$: Factor of 0.02 mol/L potassium permanganate solution

Note (2) Titrate using a brown burette.

Comment 1 Recovery testing was conducted using a reagent (ferrous sulfate heptahydrate); as a result, the average rate of recovery was 29.1 % (mass fraction) as total sulfate content (T-SO$_3$). The recovery rate to a theoretical value was 101.0 %.

Additionally, the minimum limit of quantification of this testing method is about 0.04 % (mass fraction).

References


(5) Flow sheet for total sulfur content: The flow sheet for total sulfur content in fertilizers mainly containing ferrous sulfate is shown below.

- Weigh to the order of 0.1 mg into a 200-mL tall beaker
- About 50 mL of water
- About 15 mL of sulfuric acid (1+5)
- Shake to mix
- About 1 mL of phosphate
- 0.02 mol/L potassium permanganate solution (until the solution becomes light red)

Figure Flow sheet for total sulfur content (raw material: ferrous sulfate)
4.12.1.b Barium chloride gravimetric analysis

(1) Summary
This test method is applicable to fertilizers mainly containing sulfur or sulfuric acid among sulfur and its compound. This testing method is classified as Type B and its symbol is 4.12.1.b-2017 or T.S.b-1.

Dissolve an analytical sample in a potassium hydroxide-ethanol solution and add hydrogen peroxide to oxidize, and measure the mass of barium sulfate (BaSO₄) formed by reaction with barium chloride to obtain the total sulfate content (T-SO₃). In addition, the performance of this testing method is shown in Comment 1.

(2) Reagent: Reagents are as shown below.
   a) Potassium hydroxide/ethanol solution: Dissolve 10 g of potassium hydroxide specified in JIS K 8574 in 50 mL of ethanol (95) specified in JIS K 8102, further add 50 mL of water.
   b) Hydrogen peroxide: A JIS Guaranteed Reagent (30 % (mass fraction)) specified in JIS K 8230 or a reagent of equivalent quality.
   c) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   d) Nitric acid: A JIS Guaranteed (HNO₃ 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   e) Barium chloride solution: Dissolve 100 g of barium chloride dihydrate specified in JIS K 8155 in water to make 1000 mL.
   f) Silver nitrate solution (2 g/100mL): Dissolve 2 g of silver nitrate specified in JIS K 8550 in water to make 100 mL.
   g) Phenolphthalein solution (1 g/100 mL): Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 °C.
   b) Water bath: Water bath that can be adjusted to 80 °C - 90 °C.
   c) Crucible: After heating porcelain crucible or platinum crucible in an electric furnace at 800 °C in advance, let it stand to cool in a desiccator and measure the mass to the order of 0.1 mg.
   d) Drying apparatus: Drying apparatus that can be adjusted to 110 °C - 120 °C.
   e) Electric furnace: An electric furnace that can be kept at 800 °C ± 5 °C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as follows:
   a) Weigh 1 g - 5 g of an analytical sample to the order of 0.1 mg, and transfer to a 200-mL tall beaker.
   b) Add about 50 mL of potassium hydroxide/ethanol solution, cover with a watch glass and heat on a hot plate to boil.
   c) After standing to cool, transfer to a 250-mL volumetric flask and add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

Note (2) Omit extraction if fluid fertilizers are made from only sulfuric acid and all materials are dissolved
(3) Until sulfur content is dissolved. About 5 minutes when raw materials, etc. are not dissolved.
(4) Omit the procedures in d) when all materials are dissolved.

(4.2) **Measurement**: Conduct measurement as shown below.

a) Transfer a predetermined volume (the equivalents of 30 mg - 170 mg as SO$_3$) of sample solution to a 300-mL tall beaker.

b) Add about 50 mL of water and about 5 mL of hydrogen peroxide and heat in a water bath at 80 °C - 90 °C for about 1 hour while sometimes shaking.

c) After standing cool, add 1 - 2 drops of phenolphthalein solution (1 g/100 mL) and add hydrochloric acid (2+1) until the color of the solution disappears.

d) Further add hydrochloric acid (2+1), add water to make about 100 mL and boil on a hotplate for about 5 minutes.

e) Immediately add about 6 mL of thermal barium chloride solution while sometimes shaking in a water bath at 80 °C - 90 °C.

f) After leaving at rest for a few minutes, add a few drops of thermal barium chloride solution and check that new precipitate of barium sulfate is no longer formed.

g) Further add about 2 mL of thermal barium chloride solution (100 g/L) while shaking to mix.

h) After heating in a water bath at 80 °C - 90 °C for about 2 hours, stop the heat source of water bath and let it stand to cool for no less than 4 hours.

i) Filter with filter paper (Type 5-C), wash a container with water and move the whole precipitate to a filter paper.

j) Wash the precipitate and the filter paper (Type 5-C) with water several times.

k) Put the precipitate together with the filter paper into the crucible.

l) Put the crucible into a drying apparatus and dry at about 110 °C - 120 °C for 1 hour.

m) After standing to cool, put the crucible into an electric funnel and heat gently to char.

n) Ignite at about 800 °C ± 5 °C for 2 hours.

o) After ignition, move the crucible to a desiccator and let it stand to cool.

p) Measure the mass of crucible to the order of 0.1 mg.

q) Calculate the total sulfur content (T-SO$_3$) in an analytical sample by the following formula.

\[
\text{Total sulfur content (% (mass fraction)) = } \frac{A \times 0.343}{W \times V_2/V_1} \times 100 = \frac{34.3 \times A \times V_1}{W \times V_2}
\]

A: Mass (g) of the precipitate in p)

W: Mass (g) of the analytical sample

V$_1$: Constant volume (mL) of sample solution

V$_2$: Volume (mL) of sample solution transferred

**Note (5)** Weigh 1g - 5g of an analytical sample to the order of 0.1 mg if the fluid fertilizers of the analytical sample are made from only sulfuric acid and all materials are dissolved.

(6) It can be suspended after the procedures end.

(7) A pH meter can be used for neutralization.

(8) Omit the procedures in e) if fluid fertilizers are made from only sulfuric acid and all materials are dissolved.

(9) Heated up to 70 °C - 80 °C in a water bath in advance.

(10) Add drop-by-drop.

(11) Until precipitate settles.

(12) Add a barium chloride solution slightly excessively to reduce the solubility of barium sulfate.

(13) Wash the precipitate until a white turbidity is no longer formed when about 5 mL of
nitric acid (1+2) and about 1 mL of silver nitrate solution (2 g/100 mL) are added to about 20 mL of washing.

(14) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 800 °C in 1 to 2 hours.

(15) To prevent a crucible from chipping, it is recommended to let it stand to cool gently in an electric furnace until the temperature of an electric furnace falls to no more than 200 °C.

(16) Time to let it stand to cool in a desiccator should be constant. In the case of a porcelain crucible, it is about 45 - 60 minutes.

Comment 1 Testing was conducted using sulfur simple substance fertilizers containing no materials (2 samples); as a result, the quantitative value of the total sulfur content (T-SO₃) was 99.9 % - 100.1 % to a theoretical value. Table 1 shows results and analysis results from a collaborative study for testing method validation.

Additionally, the minimum limit of quantification of this testing method is about 0.4 % (mass fraction).

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories 1)</th>
<th>Mean 2) (%)</th>
<th>Mean 3) (%)</th>
<th>sʳ 4) (%)</th>
<th>RSDr 5) (%)</th>
<th>sR 6) (%)</th>
<th>RSDr 7) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfur material a</td>
<td>8</td>
<td>8.32</td>
<td>3.33</td>
<td>0.02</td>
<td>0.7</td>
<td>0.05</td>
<td>1.4</td>
</tr>
<tr>
<td>Sulfur material b</td>
<td>10</td>
<td>12.71</td>
<td>5.09</td>
<td>0.03</td>
<td>0.6</td>
<td>0.14</td>
<td>2.8</td>
</tr>
<tr>
<td>Sulfur material c</td>
<td>9</td>
<td>247.6</td>
<td>99.17</td>
<td>0.24</td>
<td>0.2</td>
<td>1.39</td>
<td>1.4</td>
</tr>
<tr>
<td>Sulfur material d</td>
<td>8</td>
<td>245.6</td>
<td>98.37</td>
<td>0.18</td>
<td>0.2</td>
<td>0.30</td>
<td>0.3</td>
</tr>
<tr>
<td>Sulfur material e</td>
<td>8</td>
<td>1.41</td>
<td>0.564</td>
<td>0.002</td>
<td>0.4</td>
<td>0.003</td>
<td>0.6</td>
</tr>
<tr>
<td>Sulfur material f</td>
<td>9</td>
<td>2.89</td>
<td>1.157</td>
<td>0.001</td>
<td>0.1</td>
<td>0.010</td>
<td>0.9</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Total mean as sulfur trioxide (SO₃) (n = number of laboratories × number of repetition (2))
3) Total mean as sulfur (S) obtained by dividing the total mean in 2) by 2.4969
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation
8) Mass fraction

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References

1) JIS K 8088: Sulfur (Regent) (2010)
3) Edited by KANTO CHEMICAL CO., INC.: Technique on Chemical Analysis of Regent - Practical Basic Technique and Knowledge, p. 112 - 120 (2009)

(5) Flow sheet for total sulfur content: The flow sheet for total sulfur content in fertilizers mainly containing sulfur and sulfuric acid is shown below.

<table>
<thead>
<tr>
<th>1 g - 5 g analytical sample</th>
<th>Weigh to the order of 0.1 mg into a 200-mL tall beaker</th>
</tr>
</thead>
<tbody>
<tr>
<td>← About 50 mL of potassium hydroxide/ethanol solution</td>
<td>Heating</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Cover with a watch glass and boil</td>
</tr>
<tr>
<td>Transfer</td>
<td>250-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Filtration</td>
</tr>
<tr>
<td>Type 3 filter paper</td>
<td>Sample solution</td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for total sulfur content in fertilizers (Extraction procedure)
## Sample solution

- **Aliquot (predetermined volume)**
  - 300-mL tall beaker
  - About 50 mL of water
  - About 5 mL of hydrogen peroxide

### Heating
- 80 °C - 90 °C, 1 hour

### Standing to cool
- 1 - 2 drop(s) of phenolphthalein solution (1 g/100 mL)

### Neutralization
- Hydrochloric acid (2+1) (until the solution becomes transparent)
  - About 1mL of hydrochloric acid (2+1)
  - Water (the liquid volume to reach about 100 mL)

### Heating
- Boil, for 5 minutes
  - About 6 mL of thermal barium chloride solution, while shaking

### Leaving at rest
- For several minutes
  - A few drops of thermal barium chloride solution
    - Check that new precipitate is no longer formed
  - About 2 mL of thermal barium chloride solution, while shaking

### Heating
- 80 °C - 90 °C, 2 hour

### Standing to cool
- For no less than 4 hours, in a water bath whose heat source is stopped

### Filtration
- Type 5-C filter paper

### Transfer
- Type 5-C filter paper, water
  - Wash with water (until reaction of chloric matters disappears in filtrate)

### Put
- Crucible

### Drying
- Drying apparatus, about 100 °C - 120 °C, 1 hour

### Standing to cool

### Charring
- Heat gently in an electric furnace

### Incineration
- Ignite at 800 °C ± 5°C, 2 hours

### Standing to cool

### Measurement
- Measure the mass to the order of 0.1 mg

---

**Figure 2** Flow sheet for total sulfur content in fertilizers

(Measurement procedure)
4.12.1.c Transmitted light analysis

(1) Summary
This test method is applicable to fertilizers mainly containing sulfur or sulfuric acid among sulfur and its compound. This testing method is classified as Type D and its symbol is 4.12.1.c-2017 or T-S.c-1.

Dissolve an analytical sample in a potassium hydroxide-ethanol solution and add hydrogen peroxide to oxidize, and then measure the intensity of transmitted light of suspension of barium sulfate (BaSO₄) formed by the reaction with barium chloride as the absorbance to obtain total sulfur content (T-SO₃) in an analytical sample. In addition, the performance of this testing method is shown in Comment 2.

(2) Reagent, etc.: Reagents are as shown below.
   b) Potassium hydroxide/ethanol solution: Dissolve 10 g of potassium hydroxide specified in JIS K 8574 in 50 mL of ethanol (95) specified in JIS K 8102, further add 50 mL of water.
   c) Hydrogen peroxide: A JIS Guaranteed Reagent (H₂O₂ 30 % (mass fraction)) specified in JIS K 8230 or a reagent of equivalent quality.
   d) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   e) Glycerin – ethanol solution (1+1): Add 250 mL of ethanol (95) specified in JIS K 8102 to 250 mL of glycerin specified in JIS K 8295.
   f) Sodium chloride solution (1): Dilute 240 g of sodium chloride specified on JIS K 8150 in water containing 20 mL of hydrochloric acid specified in JIS K 8180, further add water to make 1000 mL.
   g) Barium chloride solution: Sieve barium chloride dihydrate specified in JIS K 8155 to make barium chloride whose particle size is between 710 μm - 500 μm.
   h) Sulfate standard solution (SO₃ 2 mg/mL) (1): Heat potassium sulfate specified in JIS K 8962 until it becomes constant weight value at 800 °C in advance, stand to cool it in a desiccator, and then transfer 4.3531 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   i) Sulfate standard solution (SO₃ 0.02 mg/mL - 0.1 mg/mL): Transfer 2mL - 10 mL of sulfate standard solution (SO₃ 2 mg/mL) to 200-mL volumetric flasks step by step and add water up to the marked line.
   j) Phenolphthalein solution (1 g/100 mL): Dissolve 1 g of phenolphthalein specified in JIS K 8799 in 100 mL of ethanol (95) specified in JIS K 8102.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Hot plate: A hot plate whose surface temperature can be adjusted up to 250 °C.
   b) Water bath: Water bath that can be adjusted to 30 °C ± 2 °C and 80 °C - 90 °C.
   c) Magnetic stirrer:
   d) Spectrophotometer: A spectrophotometer specified in JIS K 0115

(4) Test procedure
(4.1) Extraction: Conduct extraction as follows (2):
   a) Weigh 1 g - 2 g of an analytical sample to the order of 0.1 mg, and transfer to a 200-mL tall beaker.
   b) Add about 50 mL of potassium hydroxide/ethanol solution, cover with a watch glass and heat on a hot plate to boil (3).
c) After standing to cool, transfer to a 250-mL volumetric flask and add water up to the marked line.

d) Filter with Type 3 filter paper \(^{(4)}\) to make the extract.

Note (2) Omit extraction if fluid fertilizers are made from only sulfuric acid and all materials are dissolved

(3) Until sulfur content is dissolved. About 5 minutes when raw materials, etc. are not dissolved.

(4) Omit the procedures in d) when all materials are dissolved.

(4.2) Oxidation: Conduct oxidation as follows:

a) Transfer a predetermined volume (the equivalents of 5 mg - 200 mg as SO\(_3\)) of sample solution to a 300-mL tall beaker \(^{(5)}\).

b) Add about 50 mL of water and about 5 mL of hydrogen peroxide and heat in a water bath at 80 °C - 90 °C for about 1 hour while sometimes shaking \(^{(6)}\).

c) After standing cool, add 1 - 2 drop(s) of phenolphthalein solution (1 g/100 mL) \(^{(7)}\), and add hydrochloric acid (2+1) until the color of the solution disappears \(^{(8)}\).

d) After standing to cool, transfer to a 200-mL volumetric flask and add water up to the marked line.

e) Filter with 0.3μm glass filter paper.

Note (5) Weigh 1g - 5g of an analytical sample to the order of 0.1 mg if the fluid fertilizers of the analytical sample are made from only sulfuric acid and all materials are dissolved.

(6) It can be suspended after the procedures end.

(7) A pH meter can be used for neutralization.

(8) Omit the procedures in c) if fluid fertilizers are made from only sulfuric acid and all materials are dissolved.

(4.3) Precipitate formation: Form precipitate as shown below.

a) Transfer 50 mL of filtrate to a 100-mL Erlenmeyer flask with screw cap.

b) Add about 10 mL of glycerin – ethanol solution (1+1) and about 5 mL of sodium chloride solution in the Erlenmeyer flask with screw cap.

c) Warm up on a water bath at 30ºC ± 2 ºC.

d) After warming, add 0.30 g of barium chloride and stir with a magnetic stirrer for about 2 minutes.

e) Warm up on a water bath at 30ºC ± 2 ºC for about 4 minutes.

f) After warming, stir with a magnetic stirrer for about 3 minutes to make a sample solution.

g) As a blank test, conduct the procedures in a) - c) and f) using another 100-mL Erlenmeyer flask with screw cap to prepare the blank test solution.

(4.4) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.

Detection wavelength: 450 nm

b) Calibration curve preparation

1) Transfer 50 mL of sulfate standard solution (SO\(_3\) 0.02 mg/mL - 0.1 mg/mL) to a 100-mL Erlenmeyer flask with screw cap and conduct the procedure (4.3) b) - f) to make the SO\(_3\) 1 mg/65 mL - 5 mg/65 mL sulfate standard solution for the calibration curve preparation.
2) Transfer 50 mL of water to another 100-mL Erlenmeyer flask with screw cap and conduct the same procedures as 1) to make the blank test solution for the calibration curve preparation.

3) Measure absorbance at a wavelength of 450 nm of the sulfuric acid standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control (8) (9).

4) Prepare the calibration curve of the sulfuric acid concentration and absorbance of the sulfuric acid standard solutions for the calibration curve preparation.

c) Sample measurement
1) For the sample solution, conduct procedures similarly as in b) 3) to measure absorbance.
2) For the blank test solution, conduct procedures similarly as in 1) to measure absorbance, and correct the absorbance obtained for the sample solution.
3) Obtain the sulfate (SO\textsubscript{3}) content from the calibration curve, and calculate the total sulfur content (T-SO\textsubscript{3}) in the analytical sample.

Note  (8) Measure right after stirring because barium sulfate easily precipitates.
(9) A spectrophotometer with automatic sample introducing device is preferable.

Comment 1 The range of calibration curve with linearity is SO\textsubscript{3} 1 mg/65 mL - 5 mg/65 mL and the curve does not pass through the origin.

Comment 2 Testing was conducted using sulfur simple substance fertilizers containing no materials (2 samples); as a result, the quantitative value of the total sulfur content (T-SO\textsubscript{3}) was 98.4% - 99.4% to a theoretical value. Additionally, the minimum limit of quantification of this testing method is about 1% (mass fraction).

References
1) JIS K 8001: General rule for test methods of reagents (2009)
2) JIS K 8088: Sulfur (Regent) (2010)
3) JAPAN Sewage Works Association: Sewage Sludge Analysis -2007-, p. 132 - 134 Tokyo (1964)
4) Edited by KANTO CHEMICAL CO., INC.: Technique on Chemical Analysis of Regent -Practical Basic Technique and Knowledge, p. 131 - 135 (2009)
(5) Flow sheet for total sulfur content: The flow sheet for total sulfur content in fertilizers mainly containing sulfur and sulfuric acid is shown below.

**Flow sheet for total sulfur content in fertilizers (Extraction procedure):**

- Weigh to the order of 0.1 mg into a 200-mL tall beaker
- About 50 mL of potassium hydroxide/ethanol solution
- Cover with a watch glass and boil
- 250-mL volumetric flask, water
  - Water (up to the marked line)
  - Type 3 filter paper

**Flow sheet for total sulfur content in fertilizers (Oxidation procedure):**

- 300-mL tall beaker
- About 50 mL of water
- About 5 mL of hydrogen peroxide
- 80 °C - 90 °C, 1 hour
- 1 - 2 drop(s) of phenolphthalein solution (1 g/100 mL)
- Hydrochloric acid (2+1) (until the color of solution becomes transparent)
- 200-mL volumetric flask, water
- 0.3μm glass filter paper
Figure 2 Flow sheet for total sulfur content in fertilizers  
(precipitate formation and measurement procedure)
4.13 Iron
4.13.1 Water-soluble iron
4.13.1.a Flame atomic absorption spectrometry

(1) Summary
This test method is applicable to fertilizers that indicate iron content as a response modifier. This testing method is classified as Type D and its symbol is 4.13.1.a-2017 or W-Fe.a-1.
Extract by adding water to an analytical sample, spray in an acetylene–air flame and measure the atomic absorption with iron at a wavelength of 248.3 nm to obtain water-soluble iron (W-Fe) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
 b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
 c) Iron standard solution (Fe 0.1 mg/mL): An iron standard solution (Fe 0.1 mg/mL) traceable to National Metrology.
 d) Iron standard solutions (Fe 0.5 µg/mL - 5 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of iron standard solution (Fe 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line.
 e) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in d).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the iron standard solution in (2), an iron standard solution for the calibration curve preparation can be prepared by using an iron standard solution (Fe 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
 a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
 b) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (2) function.
 1) Light source: An iron hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
 2) Gas: Gas for heating by flame
 (i) Fuel gas: acetylene
 (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

Note (2) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Powdery test sample
 a) Weigh 5.00 g of an analytical sample, and put it in a 500-mL volumetric flask.
 b) Add about 400 mL of water, and shake to mix at 30 - 40 revolutions/min for about 30 minutes.
 c) Add water up to the marked line.
 d) Filter with Type 3 filter paper to make a sample solution.
Comment 2 In the procedure in (4.1.1) a), it is also allowed to weigh 2.50 g of the analytical sample and transfer to a 250-mL volumetric flask.

Comment 3 The procedure in (4.1.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 4 The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 248.3 nm
b) Calibration curve preparation
   1) Spray the iron standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 248.3 nm.
   2) Prepare a curve for the relationship between the iron concentration and the indicated value of the iron standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 mg - 0.5 mg as Fe) to a 100-mL volumetric flask.
   2) Add about 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the iron content from the calibration curve, and calculate the water-soluble iron (W-Fe) in the analytical sample.

Comment 5 Recovery testing was conducted to evaluate trueness using a preparation sample (solid). As a result, the average rate of recovery at the content level of 10 % (mass fraction), 5 % (mass fraction) and 0.05 % (mass fraction) are 101.1 %, 102.8 % and 107.0 % as water-soluble iron (W-Fe) respectively. In addition, recovery testing was conducted using a preparation sample (fluid). As a result, the average rate of recovery at the content level of 1 % (mass fraction), 0.1 % (mass fraction) and 0.01 % (mass fraction) are 103.6 %, 105.7 % and 105.1 % as a water-soluble iron (W-Fe) respectively. The results of the repeatability tests on different days using a fluid mixed fertilizer and a liquid microelement mixed fertilizer to evaluate the extract precision of fluid fertilizers were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 40 mg/kg (solid fertilizers) and 4 mg/kg (fluid fertilizers).
Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^3$</th>
<th>Repeatability standard deviation $s_r$</th>
<th>Repeatability relative standard deviation $RSD_r$</th>
<th>Intermediate precision $s_{I(T)}$</th>
<th>Intermediate relative standard deviation $RSD_{I(T)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid compound fertilizer</td>
<td>7</td>
<td>0.244</td>
<td>0.002</td>
<td>0.6</td>
<td>0.003</td>
<td>1.4</td>
</tr>
<tr>
<td>Liquid microelement mixed fertilizer</td>
<td>7</td>
<td>0.099</td>
<td>0.001</td>
<td>0.5</td>
<td>0.003</td>
<td>2.9</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) \times the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

5) Flow sheet for water-soluble iron: The flow sheet for water-soluble iron in fertilizers is shown below:

```
5.00 g analytical sample (powdery)  500-mL volumetric flask.
← Water, About 400 mL

Shaking to mix Rotary shaker (30 - 40 revolutions/min), for 30 minutes
← Water (up to the marked line)

Filtration Type 3 filter paper

Sample solution
```

Figure 1-1  Flow sheet for water-soluble iron in fertilizers
(Extraction procedure (4.1.1))
1.00 g analytical sample (Fluid) → 100-mL volumetric flask.
  ← Water, About 50 mL
Shaking to mix
  ← Water (up to the marked line)
Filtration → Type 3 filter paper
Sample solution

Figure 1-2  Flow sheet for water-soluble iron in fertilizers
(Extraction procedure (4.1.2))

Sample solution

Aliquot (predetermined amount) → 100-mL volumetric flask
  ← 25 mL of hydrochloric acid (1+5)
  ← Water (up to the marked line)
Measurement → Atomic absorption spectrometer (248.3 nm)

Figure 2  Flow sheet for water-soluble iron in fertilizers
(Measurement procedure)
4.13.1.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.13.1.b-2017 or W-Fe.b-1.

Extract by adding water to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the iron at a wavelength of 259.940 nm to obtain water-soluble iron(W-Fe) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Iron standard solution (Fe 1 mg/mL): An iron standard solution (Fe 1 mg/mL) traceable to National Metrology.
   d) Iron standard solution (Fe 0.1 mg/mL) (1): Transfer 10 mL of iron standard solution (Fe 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   e) Iron standard solutions (Fe 1 µg - 20 µg/mL) for the calibration curve preparation (1): Transfer 1 mL - 20 mL of iron standard solution (Fe 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Iron standard solutions (Fe 0.1 µg - 1 µg/mL) for the calibration curve preparation (1): Transfer 1 mL - 10 mL of iron standard solution (Fe 10 µg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the iron standard solution in (2), an iron standard solution for the calibration curve preparation can be prepared by using an iron standard solution (Fe 10 mg/mL) traceable to National Metrology.

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, in case of using the ICP-OES of an axial observation mode, it is recommended to prepare an iron standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample (2) to the order of 1 mg, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, shake to mix and add water up to the marked line.
c) Filter with Type 3 filter paper to make a sample solution.

**Note (2)** The sampling amount of the analytical sample is 10 g when the content in the sample is less than 0.01 % (mass fraction) as water-soluble iron.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

**4.2 Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
   - Analytical line wavelength: 259.940 nm

b) **Calibration curve preparation**
   1) Spray the iron standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 259.940 nm.
   2) Prepare a curve for the relationship between the iron concentration and the indicated value of the iron standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Fe) to a 100-mL volumetric flask.
   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the iron content from the calibration curve, and calculate the water-soluble iron (W-Fe) in the analytical sample.

**Comment 4** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

**Comment 5** The comparison of the measurement value ($y_i$: 0.0191 % (mass fraction) - 0.517 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of flame atomic absorbance spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = 0.001 + 0.968x$, and its correlation coefficient ($r$) was 0.999. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery at additive level of 0.01 % (mass fraction) and 0.1 % (mass fraction) were 96.5 % and 93.9 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average (^2) ((% ))</th>
<th>Repeatability (s_t ) ((% ))</th>
<th>RSD ((% ))</th>
<th>Intermediate precision (s_{II(T)} ) ((% ))</th>
<th>RSD (I(T) ) ((% ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.145</td>
<td>0.001</td>
<td>0.6</td>
<td>0.002</td>
<td>1.1</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (fluid)</td>
<td>7</td>
<td>0.0485</td>
<td>0.003</td>
<td>0.5</td>
<td>0.0005</td>
<td>0.9</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test  
2) Average (the number of test days \(T\)) \times the number of duplicate testing \((2)\)  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation

References


(5) Flow sheet: The flow sheet for water-soluble iron in fluid fertilizers is shown below:

![Figure 1](image1.png)

Figure 1 The flow sheet for water-soluble iron in fertilizers (Extraction procedure)

![Figure 2](image2.png)

Figure 2 The flow sheet for water-soluble iron in fertilizers (Measurement procedure)
4.14 Molybdenum

4.14.1 Water-soluble molybdenum

4.14.1.a Sodium thiocyanate absorpiometric analysis

(1) Summary
This test method is applicable to fertilizers that indicate molybdenum content as a response modifier. This testing method is classified as Type D and its symbol is 4.14.1.a-2017 or W-Mo.a-1.

Extract by adding water to an analytical sample, add sulfuric acid (1+1) and perchloric acid, further add a sodium thiocyanate solution and a tin (II) chloride solution, and measure the absorbance with thiocyanate complex formed by the reaction of reduced molybdenum (V) with thiocyanate ion to obtain water-soluble molybdenum (W-Mo) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent: Reagents are as shown below.
   a) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.
   b) Perchloric acid: A JIS Guaranteed Reagent specified in JIS K 8223 or a reagent of equivalent quality.
   c) Iron (III) sulfate solution (1): Dissolve 5 g of iron (III) sulfate specified in JIS K 8981 in about 10 mL of sulfuric acid (1+1) and a proper amount of water, and further add water to make 100 mL.
   d) Sodium thiocyanate solution (1): Dissolve 50 g of sodium thiocyanate specified in JIS K 9002 in water to make 500 mL.
   e) Tin (II) chloride solution (1): Dissolve 20 g of tin (II) chloride dihydrate specified in JIS K 8136 in 80 mL of hydrochloric acid (1+1) while heating, then add water to make 200 mL.
   f) Molybdenum standard solution (Mo 1 mg/mL) (1): After leaving at rest molybdenum (VI) oxide (2) in a desiccator for about 24 hours to dry, put it to a 1,500 g weighing dish. Dissolve in a small amount of water, transfer to a 1000 mL volumetric flask and add about 5 g of sodium hydroxide specified in JIS K 8576 to dissolve and add water up to the marked line.
   g) Molybdenum standard solution (Mo 0.01 mg/mL): Dilute a predetermined amount of molybdenum standard solution (Mo 1 mg/mL) precisely by a factor of 100 with water.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) A reagent of no less than 99.5 % (mass fraction) in purity is commercially sold as molybdenum (VI) oxide.

Comment 1 Instead of the molybdenum standard solution in (2), a molybdenum standard solution for the calibration curve preparation can be prepared by using a molybdenum standard solution (Mo 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) Spectrophotometer: A spectrophotometer specified in JIS K 0115

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Powdery test sample
a) Weigh 5.00 g of an analytical sample, and put it in a 500-mL volumetric flask.
b) Add about 400 mL of water, and shake to mix at 30-40 revolutions/min for about 30 minutes.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 2 In the procedure in (4.1.1) a), it is also allowed to weigh 2.50 g of the analytical sample and transfer to a 250-mL volumetric flask.

Comment 3 The procedure in (4.1.1) is the same as the procedure in (4.1.1.1) in 4.2.4.a.

Comment 4 When the sample solution in d) contains organic matters that affect the determination, put a predetermined amount of the sample solution to a 100-mL tall beaker, add a small amount of sulfuric acid and nitric acid to heat and digest the organic matters until white smoke of sulfuric acid evolves. After standing to cool, transfer the solution to a 100-mL volumetric flask and add water up to the marked line to filter. The filtrate is prepared as the sample solution of (4.2) a).

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line.
d) Filter with Type 3 filter paper to make a sample solution.

Comment 5 The procedure in (4.1.2) is the same as the procedure in (4.1.2) in 4.2.4.a.

(4.2) Coloring: Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 0.3 mg as Mo) to a 100-mL volumetric flask.
b) Add about 5 mL of sulfuric acid (1+1), about 5 mL of perchloric acid and about 2 mL of iron (III) sulfate solution.
c) Add about 16 mL of sodium thiocyanate solution and about 10 mL of tin (II) chloride solution successively while shaking to mix and further add water up to the marked line (3).

Note (3) When the solution becomes muddy, centrifuge after the procedure in c). However, if it is presumed that copper (I) thiocyanate caused the muddying, centrifuge after leaving at rest for 1 hour.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.
   Detection wavelength: 460 nm

b) Calibration curve preparation
   1) Transfer 1 mL - 30 mL of molybdenum standard solution (Mo 0.01 mg/mL) to 100-mL volumetric flasks step-by-step.
   2) Conduct the same procedure as (4.2) b) - c) to make the 0.01 mg/100 mL - 0.3 mg/100 mL molybdenum standard solution for the calibration curve preparation.
   3) Conduct the same procedures as 2) for another 100-mL volumetric flask to make the blank
test solution for the calibration curve preparation.

4) Measure absorbance at wavelength 460 nm of the molybdenum standard solution for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.

5) Prepare a curve for the relationship between the molybdenum concentration and the absorbance of the molybdenum standard solutions for the calibration curve preparation.

c) Sample measurement
1) Regarding the solution in (4.2) c), measure absorbance by the same procedure as b) 4).

2) Obtain the molybdenum content from the calibration curve, and calculate the water-soluble molybdenum (W- Mo).

Comment 6 Recovery testing was conducted to evaluate trueness using a preparation sample. As a result, the average rate of recovery at the content level of 2.5 % (mass fraction) and 0.1 % (mass fraction) are 100.2 % and 100.8 % as water-soluble molybdenum (W-Mo) respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a liquid microelement mixed fertilizer to evaluate the extract precision of fluid fertilizers were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 60 mg/kg (solid fertilizers) and 6 mg/kg (fluid fertilizers).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average ( T^1 ) (( % )) ( ^2 )</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>( s_r^4 ) (( % )) ( ^3 )</td>
<td>( RSD_r^5 ) (( % ))</td>
<td>( s_{I(T)}^6 ) (( % )) ( ^3 )</td>
</tr>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.242</td>
<td>0.001</td>
<td>0.4</td>
</tr>
<tr>
<td>Liquid microelement mixed fertilizers</td>
<td>7</td>
<td>0.0228</td>
<td>0.0001</td>
<td>0.4</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \( T \) × the number of duplicate testing \( 2 \))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation

Table 1 Analysis results of the repeatability tests on different days

References
(5) **Flow sheet for water-soluble molybdenum**: The flow sheet for water-soluble molybdenum in fertilizers is shown below:

![Flow sheet for water-soluble molybdenum](image)

Figure 1-1 Flow sheet for water-soluble molybdenum in fertilizers (Extraction procedure (4.1.1))

![Flow sheet for water-soluble molybdenum](image)

Figure 1-2 Flow sheet for water-soluble molybdenum in fertilizers (Extraction procedure (4.1.2))

Figure 2 Flow sheet for water-soluble molybdenum in fertilizers (Measurement procedure)
4.14.1.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.14.1.b-2017 or W-Mo.b-1.
Extract by adding water to an analytical sample, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the molybdenum at a wavelength of 202.030 nm to obtain water-soluble molybdenum (W-Mo) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Molybdenum standard solution (Mo 1 mg/mL): A molybdenum standard solution (Mo 1 mg/mL) traceable to National Metrology.
   d) Molybdenum standard solution (Mo 0.1 mg/mL) (1): Transfer 10 mL of molybdenum standard solution (Mo 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   e) Molybdenum standard solutions (Mo 1 µg - 20 µg/mL) for the calibration curve preparation (1): Transfer 1 mL - 20 mL of molybdenum standard solution (Mo 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Molybdenum standard solutions (Mo 0.1 µg - 1 µg/mL) for the calibration curve preparation (1): Transfer 1 mL - 10 mL of molybdenum standard solution (Mo 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the molybdenum standard solution in (2), a molybdenum standard solution for the calibration curve preparation can be prepared by using a molybdenum standard solution (Mo 10 mg/mL) traceable to National Metrology.

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, when using the ICP-OES of an axial observation mode, it is recommended to prepare a molybdenum standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample (2) to the order of 1 mg, and put it in a 100-mL volumetric
flask.

b) Add about 50 mL of water, shake to mix and add water up to the marked line.

c) Filter with Type 3 filter paper to make a sample solution.

**Note** (2) The sampling amount of the analytical sample is 10 g when the content in the sample is less than 0.01 % (mass fraction) as water-soluble molybdenum.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

   - Analytical line wavelength: 202.030 nm

b) **Calibration curve preparation**

   1) Spray the molybdenum standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 202.030 nm.

   2) Prepare a curve for the relationship between the molybdenum concentration and the indicated value of the molybdenum standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Mo) to a 100-mL volumetric flask.

   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.

   3) Subject to the same procedure as in b) 1) to read the indicated value.

   4) Obtain the molybdenum content from the calibration curve, and calculate the water-soluble molybdenum (W-Mo).

**Comment 4** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.2.4.d Comment 4.

**Comment 5** The comparison of the measurement value ($y_i$: 0.00342 % (mass fraction) - 0.20374 % (mass fraction)) of ICP Optical Emission Spectrometry and the measurement value ($x_i$) of the Sodium thiocyanate absorptiometric analysis was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was $y = 0.0004 + 0.982x$ and its correlation coefficient ($r$) was 0.999. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery at additive level of 0.01 % (mass fraction) and 0.1 % (mass fraction) were 95.4 % and 97.6 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$ (%)</th>
<th>$s^2$ $^4$ (%)</th>
<th>RSD $^5$ (%)</th>
<th>Intermediate precision $^6$ (%)</th>
<th>RSD $^7$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.124</td>
<td>0.001</td>
<td>0.5</td>
<td>0.001</td>
<td>1.2</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (fluid)</td>
<td>7</td>
<td>0.00359</td>
<td>0.00001</td>
<td>0.3</td>
<td>0.00014</td>
<td>4.0</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

**References**


(5) **Flow sheet**: The flow sheet for water-soluble molybdenum in fluid fertilizers is shown below:

![Flow sheet image]

Figure 1 Flow sheet for water-soluble molybdenum in liquid fertilizers
(Extraction procedure)

![Flow sheet image]

Figure 2 Flow sheet for water-soluble molybdenum in liquid fertilizers
(Measurement procedure)
4.15 Cobalt
4.15.1 Water-soluble cobalt
4.15.1.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers
and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as
Type E and its symbol is 4.15.1.a-2017 or W-Co.a-1.
Extract by adding water to an analytical sample, spray in an acetylene–air flame and measure the
atomic absorption with cobalt at a wavelength of 240.7 nm to obtain water-soluble cobalt (W-Co) in
an analytical sample.

(2) Reagent, etc.: Reagents and water are as shown below.
  b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or
equivalents.
  c) Cobalt standard solution (Co 0.1 mg/mL): A cobalt standard solution (Co 0.1 mg/mL)
traceable to National Metrology.
  d) Cobalt standard solutions (Co 0.5 µg - 5 µg/mL) for the calibration curve preparation (1):
Transfer 1 mL - 25 mL of cobalt standard solution (Co 0.1 mg/mL) to 500-mL volumetric
flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
  e) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used
in the procedures in d).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the cobalt standard solution in (2), a cobalt standard solution for the
 calibration curve preparation can be prepared by using a cobalt standard solution (Co
1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
  a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in
JIS K 0121 with the background correction (2) function.
  1) Light source: An cobalt hollow cathode lamp (In case of background correction system
using continuous spectrum source, the light source is a deuterium lamp.)
  2) Gas: Gas for heating by flame
     (i) Fuel gas: acetylene
     (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

Note (2) There are the continuous source method, the Zeeman method, the non-resonance
 spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
  a) Weigh 1 g of an analytical sample (3) to the order of 1 mg, and put it in a 100-mL volumetric
flask.
  b) Add about 50 mL of water, shake to mix and add water up to the marked line.
  c) Filter with Type 3 filter paper to make a sample solution.

Note (3) The sampling amount of the analytical sample is 10 g when the content in the sample
is less than 0.01 % (mass fraction) as water-soluble cobalt.
Comment 2 The procedure in (4.1) is the same as the procedure in (4.1.2) of 4.2.4.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
   Analytical line wavelength: 240.7 nm

b) Calibration curve preparation
   1) Spray the cobalt standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 240.7 nm.
   2) Prepare a curve for the relationship between the cobalt concentration and the indicated value of the cobalt standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Co) to a 100-mL volumetric flask.
   2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
   3) Subject to the same procedure as in b) 1) to read the indicated value.
   4) Obtain the cobalt content from the calibration curve, and calculate the water-soluble cobalt (W-Co) in the analytical sample.

(5) Flow sheet: The flow sheet for water-soluble cobalt in fluid fertilizers is shown below:

![Flow sheet](attachment:flow_sheet.jpg)

Figure 1 The flow sheet for water-soluble cobalt in fertilizers
(Extraction procedure)

![Flow sheet](attachment:flow_sheet2.jpg)

Figure 2 The flow sheet for water-soluble cobalt in fertilizers
(Measurement procedure)
4.15.1.b ICP Optical Emission Spectrometry

(1) Summary
This testing method is applicable to fluid mixed fertilizers, liquid microelement mixed fertilizers and the fluid fertilizers of home garden-use mixed fertilizers. This testing method is classified as Type D and its symbol is 4.15.1.b-2017 or W-Co.b-1.

Extract by adding water to an analytical sample, introduce it to an ICP Optical Emission Spectrometer ("ICP-OES") and measure the cobalt at a wavelength of 228.616 nm to obtain water-soluble cobalt (W-Co) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Cobalt standard solution (Co 1 mg/mL): A cobalt standard solution (Co 1 mg/mL) traceable to National Metrology.
   d) Cobalt standard solution (Co 0.1 mg/mL) \(^{(1)}\): Transfer 10 mL of cobalt standard solution (Co 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   e) Cobalt standard solutions (Co 1 µg - 20 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 1 mL - 20 mL of cobalt standard solution (Co 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Copper standard solutions (Cu 0.1 µg - 1 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 1 mL - 10 mL of copper standard solution (Cu 10 µg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation \(^{(1)}\): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 Instead of the cobalt standard solution in (2), a cobalt standard solution for the calibration curve preparation can be prepared by using a cobalt standard solution (Co 10 mg/mL) traceable to National Metrology.

Comment 2 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in e) and f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, in case of using the ICP-OES of an axial observation mode, it is recommended to prepare a cobalt standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0115
   1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1 g of an analytical sample \(^{(2)}\) to the order of 1 mg, and put it in a 100-mL volumetric
flask.

b) Add about 50 mL of water, shake to mix and add water up to the marked line.

c) Filter with Type 3 filter paper to make a sample solution.

**Note** (2) The sampling amount of the analytical sample is 10 g when the content in the sample is less than 0.01 % (mass fraction) as water-soluble cobalt.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1.2) of **4.2.4.a**.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:
Analytical line wavelength: 228.616 nm

b) **Calibration curve preparation**
1) Spray the cobalt standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 228.616 nm.
2) Prepare a curve for the relationship between the cobalt concentration and the indicated value of the cobalt standard solution for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**
1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as Co) to a 100-mL volumetric flask.
2) Add 25 mL of hydrochloric acid (1+5), and add water up to the marked line.
3) Subject to the same procedure as in b) 1) to read the indicated value.
4) Obtain the cobalt content from the calibration curve, and calculate the water-soluble cobalt (W-Co) in the analytical sample.

**Comment 4** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see **4.2.4.d Comment 4**.

**Comment 5** The comparison of the measurement value \(y_i\): 0.00105 % (mass fraction) - 0.0213 % (mass fraction) of ICP Optical Emission Spectrometry and the measurement value \(x_i\) of flame atomic absorbance spectrometry was conducted to evaluate trueness using fluid fertilizers (12 samples). As a result, a regression equation was \(y = 0.0001 + 0.927x\), and its correlation coefficient \((r)\) was 0.996. Additionally, additive recovery testing was conducted using a fluid compound fertilizer (1 brand) and a home garden-use mixed fertilizer (1 brand). As a result, the mean recovery at additive level of 0.01 % (mass fraction) and 0.1 % (mass fraction) were 94.6 % and 98.4 % respectively.

The results of the repeatability tests on different days using a fluid mixed fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.0005 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability $T^{1)}$</th>
<th>Average $^{2)}$</th>
<th>Repeatability $^{3)}$</th>
<th>Intermediate precision $^{4)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluid mixed fertilizer</td>
<td>7</td>
<td>0.0554</td>
<td>0.0010</td>
<td>1.7</td>
</tr>
<tr>
<td>Home garder-use mixed fertilizer (fluid)</td>
<td>7</td>
<td>0.0105</td>
<td>0.0003</td>
<td>3.3</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $(T)$ × the number of duplicate testing $(2))$
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References


(5) Flow sheet: The flow sheet for water-soluble cobalt in fluid fertilizers is shown below:

Figure 1 The flow sheet for water-soluble cobalt in liquid fertilizers
(Extraction procedure)

1 g analytical sample
← Water, about 50 mL
Shaking to mix
← Water (up to the marked line)
Filtration
Sample solution

Figure 2 The flow sheet for water-soluble cobalt in liquid fertilizers
(Measurement procedure)

Sample solution

Aliquot (predetermined volume)
← 25 mL of hydrochloric acid (1+5)
← Water (up to the marked line)
Measurement ICP-OES (228.616 nm)
5. Harmful components

5.1 Mercury

5.1.a Cold vapor atomic absorption spectrometry

(1) Summary

The test method is applicable to fertilizers excluding fluid sludge fertilizers. This testing method is classified as Type B and its symbol is 5.1.a-2017 or Hg.a-1.

Pretreat an analytical sample with nitric acid-perchloric acid, and then reduce mercury (II) in the solution with tin (II) chloride. Aerate this solution, and measure the atomic absorption for generated mercury vapor at a wavelength of 253.7 nm to obtain mercury (Hg) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Perchloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

d) Sulfuric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

e) Tin (II) chloride solution: To 10 g of tin (II) chloride dihydrate specified in JIS K 8136, add 60 mL of sulfuric acid (1+20), and stir while heating to dissolve. After cooling is complete, add water to make 100 mL.

f) L-cysteine solution: To 10 mg of L-cysteine (HSCH₂CH(NH₂)COOH) of no less than 98.0 % in purity, add 100 mL of water and 2 mL of nitric acid to dissolve, and further add water to make 1000 mL. Store in a refrigerator, and do not use after 6 months after preparation.

g) Tri-n-butyl phosphate: A reagent of no less than 98 % in purity.

h) Mercury standard solution (Hg 0.1 mg/mL): A mercury standard solution (Hg 0.1 mg/mL) traceable to National Metrology.

i) Mercury standard solution (Hg 10 µg/mL): Transfer 10 mL of mercury standard solution (Hg 0.1 mg/mL) to a 100 mL of volumetric flask and add an L-cysteine solution up to the marked line.

j) Mercury standard solution (Hg 0.1 µg/mL): Dilute a predetermined amount of mercury standard solution (Hg 10 µg/mL) with an L-cysteine solution to prepare a mercury standard solution (Hg 0.1 µg/mL).

Note  (1) Use a reagent with low mercury content, such as for mercury analysis or for harmful metal analysis.

(2) Use as an anti-forming agent.

(3) This is an example of preparation; prepare an amount as appropriate.

(4) Store in a refrigerator, and do not use after 4 months after preparation.

(5) Store in a refrigerator, and do not use after 1 months after preparation.

Comment 1 Instead of the mercury standard solution in (2), a mercury standard solution for the calibration curve preparation can be prepared by using a mercury standard solution (Hg 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) Mercury atomic absorption spectrometer: Mercury atomic absorption spectrometer using a method for producing the atomic vapor by reduction specified in JIS K 0121.

1) Light source: Low-pressure mercury lamp

b) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can
be set to 180 °C - 200 °C.

c) **Sample digestion flask** (6): A 100-mL borosilicate glass volumetric flask 100 mL (180 mm total height, 13 mm mouth diameter)

**Note** (6) Distinguish the volumetric flask used in digestion as a sample digestion flask and do not use it for any other purposes.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and transfer to a sample digestion flask.

b) Add about 10 mL of nitric acid, and heat on a hot plate or sand bath for a short time (7).

c) After standing to cool, add about 10 mL of perchloric acid, and digest by heating on a hot plate or sand bath at 180 ºC - 200 ºC for about 30 minutes - 1 hour (8).

d) After standing to cool, fill up with water to 100 mL to make a sample solution.

e) As a blank test, conduct the procedures in b) - d) using another sample digestion flask to prepare the blank test solution.

**Note** (7) If it foams vigorously, leave at rest overnight.

(8) The sample solution and the blank test solution should be stored when they are cooled after the procedure in (4.1) c). After filling up the sample solution and the blank test solution with water, immediately conduct the procedure in (4.2).

(4.2) **Measurement**: Conduct measurement by cold vapor atomic absorption spectrometry specified in JIS K 0121. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used. An example of measurement using a mercury atomic absorption spectrometer is shown below:

a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:

   Analytical line wavelength: 253.7 nm

b) **Calibration curve preparation**

1) Transfer 1 mL - 20 mL of mercury standard solution (Hg 0.1 µg/mL) to 100-mL volumetric flasks step-by-step, and add water up to the marked line. Transfer 5 mL of these solutions to respective reduction vessels, add 1 drop of tri-n-butyl phosphate (9), to make mercury standard solutions for the calibration curve preparation.

2) Add 5 mL of water to another reduction vessel, and add 1 drop of tri-n-butyl phosphate (9), to make the blank test solution for the calibration curve preparation.

3) Connect the reduction vessel to the mercury atomic absorption spectrometer, and introduce sulfuric acid (1+1) and a tin (II) chloride solution, and circulate air.

4) Read the indicated value at a wavelength of 253.7 nm.

5) Prepare a curve for the relationship between the mercury content (µg) and the indicated value of the mercury standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer 5 mL of a sample solution to respective reduction vessels, add 1 drop of tri-n-butyl phosphate (9), and conduct similarly as in b) 3) - 4) to read the indicated value.

2) Transfer 5 mL of the blank test solution to a reduction vessel, add 1 drop of tri-n-butyl phosphate (9), and conduct similarly as in b) 2) - 4) to read the indicated value, and correct the indicated value obtained for the sample solution.

3) Obtain the mercury content (µg) from the calibration curve, and calculate mercury (Hg) in the sample.
Note (9) It is not required to add tri-n-butyl phosphate if not needed.

Comment 2 Instead of the correction method in c) 2), mercury (Hg) can be corrected by obtaining the mercury content in the blank test.

Comment 3 Recovery testing was conducted to evaluate trueness using an industrial sludge fertilizer (1 sample), composted sludge (3 samples) and a human waste sludge fertilizer (1 sample). As a result, the average rate of recovery at the content level of 2 mg/kg and 0.2 mg/kg are 98.7 % - 101.6 % and 100.7 % - 105.4 % as mercury (Hg) respectively. In addition, recovery testing was conducted using soybean meal, rape seed meal, compound fertilizers (2 samples) and blended fertilizers. As a result, the average rate of recovery at the content level of 40 mg/kg and 0.5 mg/kg are 98.5 % - 101.5 % and 100.4 % - 103.3 % as mercury (Hg) respectively.

Table 1 shows results and analysis results from a collaborative study for test method validation. Additionally, the minimum limit of quantification of this test method is about 0.01 mg/kg.

Table 1 Results and analysis results from a collaborative study for mercury test method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (mg/kg)</th>
<th>RSD (%)</th>
<th>RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human waste sludge fertilizer A</td>
<td>11</td>
<td>0.651</td>
<td>5.3</td>
<td>11.6</td>
</tr>
<tr>
<td>Human waste sludge fertilizer B</td>
<td>11</td>
<td>1.10</td>
<td>6.3</td>
<td>10.2</td>
</tr>
<tr>
<td>Composted sludge fertilizer A</td>
<td>11</td>
<td>0.489</td>
<td>6.8</td>
<td>10.2</td>
</tr>
<tr>
<td>Composted sludge fertilizer B</td>
<td>11</td>
<td>0.822</td>
<td>8.1</td>
<td>13.1</td>
</tr>
<tr>
<td>Composted sludge fertilizer C</td>
<td>9</td>
<td>0.182</td>
<td>10.6</td>
<td>10.6</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (n = number of laboratories x number of samples (2))
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

References


(5) **Flow sheet for mercury**: The flow sheet for mercury in fertilizers is shown below:

```
<table>
<thead>
<tr>
<th>1.00 g analytical sample</th>
<th>Sample digestion flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heating</td>
<td>About 10 mL nitric acid</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Decomposition</td>
<td>About 10 mL perchloric acid</td>
</tr>
<tr>
<td>Heating</td>
<td>Heating on a hot plate or sand bath at 180 °C - 200 °C for 30 minutes - 1 hour</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Water (fill up to 100 mL)</td>
</tr>
<tr>
<td>Sample solution</td>
<td>Atomic absorption spectrometer (253.7 nm) for mercury</td>
</tr>
<tr>
<td>Measurement</td>
<td></td>
</tr>
</tbody>
</table>
```

Figure Flow sheet for mercury in fertilizers
5.1.b Cold vapor atomic absorption spectrometry (Fluid sludge fertilizers)

(1) **Summary**

The test method is applicable to fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.1.b-2017 or Hg.b-1.

Pretreat an analytical sample with nitric acid-perchloric acid, and then reduce mercury (II) in the solution with tin (II) chloride. Aerate this solution, and measure the atomic absorption for generated mercury vapor at a wavelength of 253.7 nm to obtain mercury (Hg) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) **Reagent, etc.:** Reagents and water are as shown below.

   a) **Water:** Water of A3 specified in JIS K 0557.

   b) **Nitric acid:** A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

   c) **Hydrogen peroxide:** A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.

   d) **Sulfuric acid:** A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

   e) **Tin (II) chloride solution:** To 10 g of tin (II) chloride dihydrate specified in JIS K 8136, add 60 mL of sulfuric acid (1+20), and stir while heating to dissolve. After cooling is complete, add water to make 100 mL.

   f) **L-cystein solution:** To 10 mg of L-cystein (HSCH₂CH(NH₂)COOH) of no less than 98.0 % in purity, add 100 mL of water and 2 mL of nitric acid to dissolve, and further add water to make 1000 mL. Store in a refrigerator, and do not use after 6 months after preparation.

   g) **Mercury standard solution (Hg 0.1 mg/mL):** A mercury standard solution (Hg 0.1 mg/mL) traceable to National Metrology.

   h) **Mercury standard solution (Hg 10 µg/mL) (2) (3):** Transfer 10 mL of mercury standard solution (Hg 0.1 mg/mL) to a 100 mL of volumetric flask and add an L-cystein solution up to the marked line.

   i) **Mercury standard solution (Hg 0.1 µg/mL) (2) (4):** Dilute a predetermined amount of mercury standard solution (Hg 10 µg/mL) with an L-cystein solution to prepare a mercury standard solution (Hg 0.1 µg/mL).

**Note**

   (1) Use a reagent with low mercury content, such as for mercury analysis or for harmful metal analysis.

   (2) This is an example of preparation; prepare an amount as appropriate.

   (3) Store in a refrigerator, and do not use after 4 months after preparation.

   (4) Store in a refrigerator, and do not use after 1 month after preparation.

**Comment 1** Instead of the mercury standard solution in (2), a mercury standard solution for the calibration curve preparation can be prepared by using a mercury standard solution (Hg 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) **Apparatus and instruments:** Apparatus and instruments are shown below.

   a) **Mercury atomic absorption spectrometer:** Mercury atomic absorption spectrometer using a method for producing the atomic vapor by reduction specified in JIS K 0121.

   b) **Pressure vessel decomposing device:** A device which pressurizes the inside of a vessel by putting acid, etc. to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.

   1) **Light source:** Low-pressure mercury lamp

   2) **Pressure vessel decomposing device:** In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and
temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.

2) **Exhaust system**: A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.

3) **Airtight vessel**: A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.

c) **Centrifugal separator**: A centrifugal separator that can work at about $1700 \times g$.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 20.0 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.
c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave.
d) Ignite at 240 ºC ±5 ºC for no less than 10 minutes to decompose.
e) After standing to cool, transfer the solution to a 50-mL volumetric flask with water.
f) Add water up to the marked line and transfer it to a 50-mL ground-in stopper centrifugal precipitate tube.
g) Centrifuge it at $1700 \times g$ centrifugal force for about five minutes and use the supernatant as the sample solution.
h) As a blank test, conduct the procedures in b) - g) using another airtight vessel to prepare the blank test solution.

**Note**  
(5) The maximum limit of solid content, which is converted from moisture content, in the sampling volume 20.0 g of an analytical sample is about 0.5 g. If solid content is likely to exceed the limit, reduce sampling volume as necessary.

(6) Condition examples for a microwave decomposing device: 0 min (room temperature) → 10 min (240 ºC) → 20 min (240 ºC) → 40 min (room temperature), initial output 1400 W

(7) When organic matters still remain, for example the digestion solution is colored, repeat the procedures in (4.1) b) - c).

(8) The vessel should be made of polypropylene, etc. to not affect the measurement.

(9) 16.5-cm of radius and 3000 rpm of revolutions makes about $1700 \times g$ centrifugal force.

**Comment 2** The procedure in (4.1) is the same as the procedure in (4.1) in 5.2.c, 5.3.c, 5.4.c, 5.5.e and 5.6.c.

(4.2) **Measurement**: Conduct measurement by cold vapor atomic absorption spectrometry specified in JIS K 0121. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used. An example of measurement using a mercury atomic absorption spectrometer is shown below:

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following: Analytical line wavelength: 253.7 nm

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b) **Calibration curve preparation**

1) Transfer 0.4 mL - 10 mL of mercury standard solution (Hg 0.1 µg/mL) to 100-mL volumetric flasks step-by-step, and add water up to the marked line. Transfer 5 mL of these solutions to respective reduction vessels to make mercury standard solutions for the calibration curve preparation.

2) Add 5 mL of water to another reduction vessel to make the blank test solution for the calibration curve preparation.

3) Connect the reduction vessel to the mercury atomic absorption spectrometer, and introduce sulfuric acid (1+1) and a tin (II) chloride solution, and circulate air.

4) Read the indicated value at a wavelength of 253.7 nm.

5) Prepare a curve for the relationship between the mercury content (µg) and the indicated value of the mercury standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Transfer 5 mL of a sample solution to respective reduction vessels, and conduct similarly as in b) 3) - 4) to read the indicated value.

2) Transfer 5 mL of the blank test solution to a reduction vessel and conduct similarly as in b) 2) - 4) to read the indicated value, and correct the indicated value obtained for the sample solution.

3) Obtain the mercury content (µg) from the calibration curve, and calculate mercury (Hg) in the analytical sample.

**Comment 3** Instead of the correction method in c) 2), mercury (Hg) can be corrected by obtaining the mercury content in the blank test.

**Comment 4** Triplicates additive recovery testing was conducted to evaluate trueness using fluid industrial sludge fertilizers (2 samples), and composted sludge fertilizers (6 samples). As a result, the mean recovery at the concentration level of 0.2 mg/kg - 0.4 mg/kg, 0.01 mg/kg - 0.09 mg/kg and 0.7 µg /kg - 7 µg /kg are 100.0 % - 109.1 %, 99.0 % - 114.6 % and 100.4 % - 113.4 % as mercury (Hg) in an actual article respectively.

The results of the repeatability tests on different days using two kinds of fluid sludge fertilizers to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.2 µg/kg.

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average (^1) (mg/kg)</th>
<th>Repeatability (\overline{\sigma}^2) (mg/kg)</th>
<th>RSD (^1) (%)</th>
<th>Intermediate precision (\overline{\sigma}_{I(T)}^5) (mg/kg)</th>
<th>RSD (_{I(T)}^6) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composted sludge fertilizer 1</td>
<td>5</td>
<td>0.0577</td>
<td>0.0009</td>
<td>1.5</td>
<td>0.0014</td>
<td>2.5</td>
</tr>
<tr>
<td>Composted sludge fertilizer 2</td>
<td>5</td>
<td>0.0142</td>
<td>0.0002</td>
<td>1.7</td>
<td>0.0003</td>
<td>2.2</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \(T\) × the number of duplicate testing \(2\))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Intermediate standard deviation
6) Intermediate relative standard deviation
(5) **Flow sheet for mercury**: The flow sheet for mercury in fluid sludge fertilizers is shown below:

![Flow sheet for mercury in fluid sludge fertilizers](image-url)

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**References**

5.2 Arsenic

5.2.a Hydride generation atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.2.a-2017 or As.a-1.

Pretreat an analytical sample with nitric acid-sulfuric acid-perchloric acid, and then generate arsenic hydride by the addition of sodium tetrahydroborate in the acidic condition with hydrochloric acid, introduce it with argon gas to a heated absorption cell, and measure the atomic absorption with arsenic at a wavelength of 193.7 nm to obtain arsenic (As) in an analytical sample. In addition, the performance of this testing method is shown in Comment 7.

(2) Reagent, etc.: Reagents and water are as shown below.
  b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
  c) Sulfuric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
  d) Perchloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
  e) Hydrochloric acid: Hydrochloric acid for arsenic analysis specified in JIS K 8180, or of harmful metal analysis grade, microanalysis grade or equivalents.
  f) Potassium iodide solution (1): Dissolve 20 g of potassium iodide specified in JIS K 8913 in water to make 100 mL.
  g) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.
  h) Sodium tetrahydroborate solution (1): Dissolve 10 g of sodium tetrahydroborate (NaBH₄) for atomic absorption spectrometry in a sodium hydroxide solution (4 g/L) to make 1000 mL.
  i) Arsenic standard solution (As 0.1 mg/mL): An arsenic standard solution (As 0.1 mg/mL) traceable to National Metrology.
  j) Arsenic standard solution (As 1 µg/mL) (2)(3): Dilute a predetermined amount of arsenic standard stock solution (As 0.1 mg/mL) accurately with hydrochloric acid (1+100) to prepare an arsenic standard solution (As 1 µg/mL).
  k) Arsenic standard solution (As 0.1 µg/mL) (2)(4): Dilute a predetermined amount of arsenic standard solution (As 0.1 µg/mL) with hydrochloric acid (1+100) to prepare an arsenic standard solution (As 0.1 µg/mL).

Note (1) The concentrations of a potassium iodide solution and a sodium tetrahydroborate solution vary depending on the instrument used.
(2) This is an example of preparation; prepare an amount as appropriate.
(3) Store in a refrigerator, and do not use after 6 months after preparation.
(4) Store in a refrigerator, and do not use after 1 month after preparation.

Comment 1 Instead of the arsenic standard solution in (2), an arsenic standard solution for the calibration curve preparation can be prepared by using an arsenic standard solution (As 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
  a) Atomic absorption spectrometer: To an atomic absorption spectrometer specified in JIS K 0121, connect a hydride generator and parts shown below. Also, an atomic absorption spectrometer with a built-in hydride generator can be used.
  1) Light source: An arsenic hollow cathode lamp or an arsenic high-intensity discharge lamp.
  2) Atomizer: Heated absorption cell (5)
3) **Gas:** Gases for heating the heated absorption cell.
   (i) Fuel gas: acetylene
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Hydride generator:** A batch-type or continuous-type hydride generator specified in JIS K 0121. For continuous hydride generators, there is a method to introduce a potassium iodide solution on-line in addition to a sample solution, hydrochloric acid, and a sodium tetrahydroborate solution.

1) **Argon:** Argon of grade 2 specified in JIS K 1105 or equivalents.

c) **Hot plate or sand bath:** A hot plate whose surface temperature can be adjusted up to 350 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to no less than 300 °C.

Note (5) For cell heating, there are a method of electric heating and a method of flame heating.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 1.00 g - 2.00 g of an analytical sample, and transfer to a 200-mL - 300-mL tall beaker.

b) Add about 10 mL of nitric acid and about 5 mL of sulfuric acid, cover the tall beaker with a watch glass, and leave at rest overnight.

c) Heat mildly on a hot plate or sand bath at 170 °C - 220 °C for no less than 30 minutes. After bubbles cease to form, set the temperature of the hot plate or sand bath to no less than 300 °C, and heat until nitroxide (yellow-brown smoke) is no longer generated.

d) After standing to cool, add about 5 mL of perchloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or sand bath at no less than 300 °C for 2 - 3 hours to digest.

f) Slightly move the watch glass, and keep on heating on the hot plate or sand bath to concentrate until the liquid volume becomes no more than 2 mL.

g) After standing to cool, add about 5 mL of hydrochloric acid (1+10) and about 20 mL of water, cover the tall beaker with a watch glass and heat mildly to dissolve.

h) After standing to cool, transfer the solution to a 100-mL - 200-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note (6) Carbonization (degradation) of organic matters by sulfuric acid begins by heating when nitric acid no longer remains. In this state, As$^{5+}$ may be reduced to As$^{3+}$ and evaporate; therefore, stop heating immediately after the end of the generation of nitroxide (yellow-brown smoke).

(7) Oxidation of organic matters by perchloric acid progresses extremely rapidly and explosively. For that reason, add perchloric acid after fully degrading organic matters with nitric acid to avoid danger.

(8) When the white smoke of perchloric acid is generated, if the solution is colored such as black-brown or brown, stop heating immediately, and after standing to cool, add nitric acid, and heat again to degrade remaining organic matters.

(9) The watch glass can be removed.

(10) The generation of arsenic hydride is inhibited by the presence of nitric acid; therefore, remove nitric acid by sufficiently generating the white smoke of sulfuric acid.

**Comment 2** The procedure in (4.1) is the same as in (4.1) in 5.2.b and 5.5.c. However, the sampling amount of the analytical sample in (4.1) a) in 5.5.c is 1.00 g.
Comment 3 When the analytical sample solidifies in the procedure in (4.1) b), moisten the analytical sample with a small amount of water as necessary in advance.

Comment 4 It is not necessary to conduct the procedure in (4.1) b) “leave at rest overnight” in the case of fertilizers not containing organic matters.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used. Two examples of measurement procedures with a continuous hydride generator are shown below.

(4.2.1) Measurement (A): Leaving at rest after adding a potassium iodide solution.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
Analytical line wavelength: 193.7 nm

b) Calibration curve preparation
1) Transfer 2.5 mL - 10 mL of arsenic standard solution (As 0.1 µg/mL) to 50-mL volumetric flasks step-by-step.
2) Add 5 mL of hydrochloric acid and 5 mL of potassium iodide solution, leave at rest for about 15 minutes, and then add water up to the marked line, to make 5 ng/mL -20 ng/mL arsenic standard solutions for the calibration curve preparation.
3) Conduct the same procedures as 2) for another 50-mL volumetric flask to make the blank test solution for the calibration curve preparation.
4) While letting argon flow, introduce the arsenic standard solution for the calibration curve preparation for each step and the blank test solution for the calibration curve preparation respectively, further introduce hydrochloric acid (1+1) and a sodium tetrahydroborate solution to the hydride generator to generate arsenic hydride.
5) Separate arsenic hydride and liquid waste, and then introduce the gas containing arsenic hydride to a heated absorption cell, and read the indicated value at a wavelength of 193.7 nm.
6) Prepare a curve for the relationship between the arsenic concentration and the indicated value of the arsenic standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
1) Transfer a predetermined amount of the sample solution to a 50-mL volumetric flask, and conduct procedures similarly as in b) 2) and b) 4) - 5) to read the indicated value.
2) Transfer a predetermined amount of the blank test solution to a 50-mL volumetric flask, and conduct procedures similarly as in b) 2) and b) 4) - 5) to read the indicated value, and correct the indicated value obtained for the sample solution.
3) Obtain the arsenic content from the calibration curve, and calculate arsenic (As) in the analytical sample.

(4.2.2) Measurement (B): On-line introduction of a potassium iodide solution.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
Analytical line wavelength: 193.7 nm

b) Calibration curve preparation
1) Transfer 5 mL - 25 mL of arsenic standard solution (As 0.1 µg/mL) to 50-mL volumetric flasks step-by-step, add water up to the marked line, to make the 10 ng/mL - 50 ng/mL arsenic standard solution for the calibration curve preparation. Use water as the blank test solution for the calibration curve preparation.
2) While letting argon flow, introduce arsenic standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation, respectively, and
further introduce a potassium iodide solution, hydrochloric acid (1+1) and a sodium tetrahydroborate solution to the hydride generator to generate arsenic hydride.

3) Separate arsenic hydride and liquid waste, and then introduce the gas containing arsenic hydride to a heated absorption cell, and read the indicated value at a wavelength of 193.7 nm.

4) Prepare a curve for the relationship between the arsenic concentration and the indicated value of the arsenic standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement

1) Transfer a predetermined amount of the sample solution to a 50-mL volumetric flask, and conduct procedures similarly as in b) 2) - 3) to read the indicated value.

2) Transfer a predetermined amount of the blank test solution to a 50-mL volumetric flask, add water up to the marked line, and conduct similarly as in b) 2) - 3) to read the indicated value, and correct the indicated value obtained for the sample solution.

3) Obtain the arsenic content from the calibration curve, and calculate arsenic (As) in the analytical sample.

Comment 5 The coexistence of iron, nickel, and cobalt at over 5, 10, 80 folds amount of arsenic, respectively, inhibits the generation of arsenic hydride. However, the inhibition of arsenic hydride generation even in the coexistence of iron at 1000 folds amount can be removed by adding or introducing a potassium iodide solution.

Comment 6 Instead of the correction method in c) 2), the arsenic (As) in the analytical sample can also be corrected by obtaining the arsenic content in the blank test solution.

Comment 7 A recovery testing was conducted using industrial sludge fertilizer, composted sludge fertilizer (3 samples) and human waste sludge fertilizer; as a result, the recovery at the concentration level of 50 mg/kg and 5 mg/kg were 94.6 % - 100.6 % and 99.9 % - 103.3 % as arsenic (As), respectively. Also, a recovery testing was conducted using processed slug phosphate fertilizer, soybean meal, rape seed meal, compound fertilizer and potassium-magnesium-sulfate fertilizer; as a result, the recovery at the concentration level of 50 mg/kg and 5 mg/kg were 98.5 % - 109.8 % and 103.5 % - 108.6 %, respectively.

Table 1 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of this test method is about 0.1 mg/kg.
Table 1 Results and analysis results from a collaborative study for arsenic test method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (^2) (mg/kg)</th>
<th>(RSD_t) (^3) (%)</th>
<th>(RSD_R) (^4) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sewage sludge fertilizer</td>
<td>11</td>
<td>6.42</td>
<td>3.5</td>
<td>10.7</td>
</tr>
<tr>
<td>Human waste sludge fertilizer</td>
<td>10</td>
<td>4.62</td>
<td>4.9</td>
<td>7.0</td>
</tr>
<tr>
<td>Industrial sludge fertilizer</td>
<td>12</td>
<td>0.632</td>
<td>5.7</td>
<td>19.7</td>
</tr>
<tr>
<td>Calcined sludge fertilizer</td>
<td>12</td>
<td>5.08</td>
<td>4.1</td>
<td>9.5</td>
</tr>
<tr>
<td>Composted sludge fertilizer</td>
<td>10</td>
<td>1.23</td>
<td>6.1</td>
<td>11.4</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean \((n = \text{number of laboratories} \times \text{number of samples})\)
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

References


(5) **Flow sheet for arsenic**: The flow sheet for arsenic in fertilizers is shown below:

![Flow sheet for arsenic in fertilizers](image)

Figure Flow sheet for arsenic in fertilizers
5.2.b Silver diethyl dithiocarbamate absorptiometric analysis

(1) Summary
This test method is applicable to fertilizers other than sulfur and its compound. This testing method is classified as Type E and its symbol is 5.2.b-2017 or As.b-1.

Pretreat an analytical sample with nitric acid - sulfuric acid - perchloric acid, and then put the predetermined volume into an arsenic hydride generation bottle, and generate arsenic hydride by adding a potassium iodide solution, a tin chloride solution and zinc successively in the acidic condition with hydrochloric acid to react with silver diethyl dithiocarbamate in pyridine. Measure the absorbance with the silver diethyl dithiocarbamate solution, the coloring liquid, at a wavelength of 510 nm or 519 nm to obtain arsenic (As) in an analytical sample.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Sulfuric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Perchloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   e) Hydrochloric acid: Hydrochloric acid for arsenic analysis specified in JIS K 8180, or of harmful metal analysis grade, microanalysis grade or equivalents.
   f) Potassium iodide solution: Dissolve 20 g of potassium iodide specified in JIS K 8913 in water to make 100 mL.
   g) Tin (II) chloride solution: Dissolve 15 g of tin (II) chloride dihydrate specified in JIS K 8136 in 100 mL of hydrochloric acid (1+1), add a small amount of granular tin specified in JIS K 8580 then store in a colored bottle.
   h) Ascorbic acid: A JIS Guaranteed Reagent specified in JIS K 9502 or a reagent of equivalent quality.
   i) Zinc: A reagent of arsenic analysis grade specified in JIS K 8012 or equivalents. (1 mm - 1.5 mm particle diameter)
   j) Lead acetate glass wool: Glass wool air-dried after moisturizing with 100 mL of the solution, where 10 g of lead acetate (II) trihydrate specified in JIS K 8374 is dissolved in water.
   k) Silver diethyl dithiocarbamate solution: Dissolve 0.5 g of silver N, N-diethyl dithiocarbamate specified in JIS K 9512 in 100 ml of pyridine specified in JIS K 8777, then store in cool and dark place
   l) Arsenic standard solution (As 0.1 mg/mL): An arsenic standard solution (As 0.1 mg/mL) traceable to National Metrology.
   m) Arsenic standard solution (As 1 µg/mL) (1) (2): Dilute a predetermined amount of arsenic standard stock solution (As 0.1 mg/mL) accurately with hydrochloric acid (1+100) to prepare an arsenic standard solution (As 1 µg/mL).

Note  (1) This is an example of preparation; prepare an amount as appropriate.
(2) Store in a refrigerator, and do not use after 6 months after preparation.

Comment 1 Instead of the arsenic standard solution in (2), an arsenic standard solution for the calibration curve preparation can be prepared by using an arsenic standard solution (As 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Arsenic hydride generator: An arsenic hydride generator specified in 61.1 in JIS K 0102 or equivalents.
   b) Spectrophotometer: A spectrophotometer specified in JIS K 0115
c) **Hot plate or sand bath**: A hot plate whose surface temperature can be adjusted up to 350 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to no less than 300 °C.

(4) Test procedure

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 1.00 g - 2.00 g of an analytical sample, and transfer to a 200-mL - 300-mL tall beaker.

b) Add about 10 mL of nitric acid and about 5 mL of sulfuric acid, cover the tall beaker with a watch glass, and leave at rest overnight.

c) Heat mildly on a hot plate or sand bath at 170 °C - 220 °C for no less than 30 minutes. After bubbles cease to form, set the temperature of the hot plate or sand bath to no less than 300 °C, and heat until nitroxide (yellow-brown smoke) is no longer generated.\(^{(3)}\)^{(4)}

d) After standing to cool, add about 5 mL of perchloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or sand bath at no less than 300 °C for 2 - 3 hours to digest.\(^{(5)}\)

f) Slightly move the watch glass,\(^{(6)}\) and keep on heating on the hot plate or sand bath to concentrate until the liquid volume becomes no more than 2 mL.\(^{(7)}\)

g) After standing to cool, add about 5 mL of hydrochloric acid (1+10) and about 20 mL of water, cover the tall beaker with a watch glass and heat mildly to dissolve.

h) After standing to cool, transfer the solution to a 100-mL - 200-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

i) As a blank test, conduct the procedures in (b) - (h) using another tall beaker to prepare the blank test solution.

**Note**

(3) Carbonization (degradation) of organic matters by sulfuric acid begins by heating when nitric acid no longer remains. In this state, As\(^{5+}\) may be reduced to As\(^{3+}\) and evaporate; therefore stop heating immediately after the end of the generation of nitroxide (yellow-brown smoke).

(4) Oxidation of organic matters by perchloric acid progresses extremely rapidly and explosively. For that reason, add perchloric acid after fully degrading organic matters with nitric acid to avoid danger.

(5) When the white smoke of perchloric acid is generated, if the solution is colored such as black-brown or brown, stop heating immediately, and after standing to cool, add nitric acid, and heat again to degrade remaining organic matters.

(6) The watch glass can be removed.

(7) The generation of arsenic hydride is inhibited by the presence of nitric acid; therefore, remove nitric acid by sufficiently generating the white smoke of sulfuric acid.

**Comment 2** The procedure in (4.1) is the same as the procedure in (4.1) of 5.2.a.

**Comment 3** When the analytical sample solidifies in the procedure in (4.1) b), moisten the analytical sample with a small amount of water as necessary in advance.

**Comment 4** It is not necessary to conduct the procedure in (4.1) b) “leave at rest overnight” in the case of fertilizers not containing organic matters.

(4.2) **Reaction**: Conduct reaction as shown below.

a) Put a predetermined amount (the equivalents of 1 μg - 20 μg of As and liquid volume is no more than 40 mL) of the sample solution into an arsenic hydride generation bottle.

b) Add water to the solution to make about 40 mL.

c) Add hydrochloric acid to make it equivalent to 10 mL of hydrochloric acid.

d) Add about 2 mL of potassium iodide solution, shake and leave at rest for a few minutes.
Add about 1 mL of tin (II) chloride solution, shake and leave at rest for about 10 minutes (8).

Connect arsenic hydride generation bottle, glass tube lightly stuffed with lead acetate glass wool in advance and 5 mL of silver diethyl dithiocarbamate solution (9), and quickly put 2.5 g of zinc into an arsenic hydride generation bottle.

Leave at rest at room temperature (15ºC - 25ºC) for about 45 minutes, and allow generated arsenic hydride to absorb into the silver diethyl dithiocarbamate solution and color.

Transfer a predetermined amount of blank test solution into the arsenic hydride generation bottle, conduct procedures similarly as in (4.2) b) - g) to allow generated arsenic hydride to absorb into the silver diethyl dithiocarbamate solution and color.

Note (8) When a large amount of iron is contained, add 1 g of ascorbic acid and 2 mL of tin (II) chloride solution and shake to mix to leave at rest for about 10 minutes instead of the procedure in e).

(9) Apply a small amount of silicone grease, etc. to the connecting parts of an arsenic hydride generation bottle, glass tube and arsenic hydride absorption tube to maintain their airtightness.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used.

a) Measurement conditions of spectrophotometer: Set up the measurement conditions of spectrophotometer considering the following.
   Detection wavelength: 510 nm or 519 nm

b) Calibration curve preparation
   1) Transfer 2.5 mL - 20 mL of arsenic standard solution (1 µg/mL) to arsenic hydride generation bottles step-by-step.
   2) Conduct procedures similarly as in (4.2) b) - g) and allow them to react.
   3) For another arsenic hydride generation bottle, the silver diethyl dithiocarbamate solution prepared similarly as in the procedure in 2) is used as the blank test solution for the calibration curve preparation.
   4) Measure the absorbance at wavelength 510 nm of the silver diethyl dithiocarbamate solution of the arsenic standard solution for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.
   5) Prepare a curve for the relationship between the arsenic concentration and the indicated value of the arsenic standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
   1) For the silver diethyl dithiocarbamate solution of (4.2) g), conduct procedures similarly as in b) 4) to measure absorbance.
   2) For the silver diethyl dithiocarbamate solution of (4.2) h), conduct procedures similarly as in b) 4) to measure absorbance, and correct the absorbance obtained for the sample solution.
   3) Obtain the arsenic content from the calibration curve, and calculate arsenic (As) in the analytical sample.

Comment 5 Instead of the correction method in c) 2), the arsenic (As) in the analytical sample can also be corrected by obtaining the arsenic content in the blank test solution.

References
(5) **Flow sheet for arsenic**: The flow sheet for arsenic in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Analytical Sample</th>
<th>Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00 g - 2.00 g</td>
<td>200-mL - 300-mL tall beaker</td>
</tr>
<tr>
<td>Leaving at rest overnight</td>
<td>A small amount of water to moisten analytical sample (if needed).</td>
</tr>
<tr>
<td>Heating</td>
<td>About 10 mL nitric acid, about 5 mL sulfuric acid. Cover with a watch glass.</td>
</tr>
<tr>
<td>Heating</td>
<td>Heat mildly on a hot plate or sand bath at 170 °C - 220 °C for no less than 30 minutes.</td>
</tr>
<tr>
<td>Heating</td>
<td>Heat on a hot plate or sand bath at no less than 300 °C until yellow-brown smoke is no longer generated.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature.</td>
</tr>
<tr>
<td>Heating</td>
<td>About 5 mL of perchloric acid. Cover with a watch glass, and heat on a hot plate or sand bath at no less than 300 °C for 2 - 3 hours.</td>
</tr>
<tr>
<td>Heating</td>
<td>Slightly move the watch glass, and concentrate until it becomes no more than 2 mL.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature.</td>
</tr>
<tr>
<td>Heating</td>
<td>About 5 mL of hydrochloric acid (1+10), about 20 mL of water. Cover with a watch glass, and dissolve.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature.</td>
</tr>
<tr>
<td>Transfer</td>
<td>100-mL volumetric flask.</td>
</tr>
<tr>
<td>Filtration</td>
<td>Water (up to the marked line). Type 3 filter paper.</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

*Figure 1  Flow sheet for arsenic in fertilizers (Extraction procedure)*
Transfer into an arsenic hydride generation bottle

- Water (until liquid volume becomes about 40 mL)
- Hydrochloric acid (until hydrochloric acid becomes equivalent to 10 mL)
- About 2 mL of potassium iodide solution

**Aliquot (predetermined volume)**

- Shaking to mix
- For a few minutes
- About 1 mL of tin (II) chloride solution

**Shaking to mix**

- Leaving at rest
- For about 10 minutes
- 2.5 g of zinc

**Generation of arsenic hydride**

- Room temperature (15°C - 25°C), about 45 minutes

**Absorption**

- 5 mL of silver diethyldithiocarbamate solution

**Measurement**

- Spectrophotometer (wavelength 510 nm or 519 nm)

Figure 2  Flow sheet for arsenic in fertilizers
(Reaction and measurement procedure)
5.2.c ICP Mass Spectrometry (Fluid sludge fertilizers)

(1) Summary
The testing method is applicable to fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.2.c-2017 or As.c-1.

Add nitric acid – hydrogen peroxide to an analytical sample to heat and extract by microwave irradiation. After adding an internal standard element, introduce it to an ICP Mass Spectrometer (ICP-MS) and measure the respective indicated values of arsenic and an internal standard element with mass/charge number (m/z) and obtain arsenic (As) in the analytical sample from the ratio of the indicated value for arsenic and the indicated value for an internal standard element. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Nitric acid: Nitric acid used to dilute a standard solution and a sample solution is a highly pure reagent specified in JIS K 9901.

d) Hydrogen peroxide: A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.

e) Rhodium standard solution (Rh 1 mg/mL): A rhodium standard solution (Rh 1 mg/mL) traceable to National Metrology.

f) Rhodium standard solution (Rh 0.1 µg/mL) \(^{(1)(2)(3)}\): Dilute a predetermined amount of rhodium standard solution (Rh 1 mg/mL) with nitric acid (1+19) to prepare a rhodium standard solution (Rh 0.1 µg/mL).

g) Rhenium standard solution (Re 1 mg/mL) \(^{(4)}\): A rhenium standard solution (Re 1 mg/mL) traceable to National Metrology.

h) Rhenium standard solution (Re 0.1 µg/mL) \(^{(1)(2)(3)(4)}\): Dilute a predetermined amount of rhenium standard solution (Re 1 mg/mL) with nitric acid (1+19) to prepare a rhenium standard solution (Re 0.1 µg/mL).

i) Arsenic standard solution (As 0.1 mg/mL): An arsenic standard solution (As 0.1 mg/mL) traceable to National Metrology.

j) Arsenic standard solution (As 5 µg/mL) \(^{(1)(2)(3)}\): Transfer 5 mL of arsenic standard solution (As 0.1 mg/mL) to a 100-mL volumetric flask and add nitric acid (1+19) to the marked line.

k) Arsenic standard solutions (As 1 ng/mL - 100 ng/mL) for the calibration curve preparation \(^{(1)(2)(3)}\): Transfer 0.02 mL - 2 mL of arsenic standard solution (As 5 µg/mL) to 100-mL volumetric flasks step-by-step, and add 10 mL of rhodium standard solutions (Rh 0.1 µg/mL) as internal standard respectively \(^{(5)}\) and add nitric acid (1+19) up to the marked line.

l) Blank test solution for the calibration curve preparation \(^{(1)(2)(3)}\): Transfer 10 mL of rhodium standard solution (Rh 0.1 µg/mL) to 100-mL volumetric flasks as internal standard \(^{(5)}\) and add nitric acid (1+19) up to the marked line.

Note
(1) This is an example of preparation; prepare an amount as appropriate.
(2) Store in a refrigerator, and do not use after 1 months after preparation.
(3) For storage, use a sealable container made of materials such as polypropylene containing no arsenic.
(4) Use when measuring lead simultaneously.
(5) Add an internal solution that is 1/10 of the volume to be prepared.

Comment 1 Instead of the arsenic standard solution in (2), an arsenic standard solution for the calibration curve preparation can be prepared by using an arsenic standard solution (As 1 mg/mL or 10 mg/mL) traceable to National Metrology.
(3) **Instruments:** Instruments are as shown below:

a) **ICP Mass Spectrometer:** High-frequency plasma mass spectrometer specified in JIS K 0133.

b) **Pressure vessel decomposing device:** A device which pressurizes the inside of a vessel by putting acid, etc. to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.

1) **The main part of a decomposing device:** In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.

2) **Exhaust system:** A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.

3) **Airtight vessel:** A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.

4) **Centrifugal separator:** A centrifugal separator that can work at about $1700 \times g$.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 20.0 g (6) of an analytical sample, and put it in a 100-mL volumetric flask.

b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.

c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave (7).

d) Ignite at 240 °C ± 5 °C for no less than 10 minutes (7) to decompose (8).

Note (6) The maximum limit of solid content, which is converted from moisture content, in the sampling volume 20.0 g of an analytical sample is about 0.5g. If solid content is likely to exceed the limit, reduce sampling volume as necessary.

(7) Condition examples for a microwave decomposing device: 0 min (room temperature) → 10 min (240 °C) → 20 min (240 °C) → 40 min (room temperature), initial output 1400 W

(8) When organic matters still remain, for example the digestion solution is colored, repeat the procedures in (4.1 b) - d).

(9) The vessel should be made of polypropylene, etc. to not affect the measurement.

(10) 16.5-cm of radius and 3000 rpm of revolutions makes about $1700 \times g$ centrifugal force.

Comment 2  The procedure in (4.1) is the same as the procedure in (4.1) of 5.1.b.

(4.2) Measurement: Conduct measurement (Internal Standard Method) according to JIS K 0113 and as shown below. Specific measurement procedures are according to the operation method of the ICP Mass Spectrometer used in measurement.

a) Measurement conditions for the ICP Mass Spectrometer: Set up the measurement conditions for the ICP Mass Spectrometer considering the following:
   Arsenic: monitor ion (m/z): 75
   Rhodium: monitor ion (m/z): 103

b) Calibration curve preparation
   1) Spray arsenic standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into the inductively coupled plasma, and read the ratio of the respective ion count number for the monitor ion of an element subjected to measurement and an internal standard element.
   2) Prepare a curve for the relationship between the concentration and the ratio of the ion count number of an element subjected to measurement.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 µg - 5 µg as As) to a 50-mL volumetric flask (9).
   2) Add 5mL of internal standard solution (5) and add nitric acid (1+19) to the marked line.
   3) Conduct procedures similarly as in b) 1) to read the ratio of the ion count numbers.
   4) For blank test solution, conduct procedures similarly as in 1) - 3) to correct the ratio of the ion count numbers obtained for the sample solution.
   5) Obtain the arsenic content from the calibration curve, and calculate arsenic (As) in the analytical sample.

Comment 3 Instead of the correction method in c) 4), the arsenic (As) in the analytical sample can also be corrected by obtaining the arsenic content in the blank test solution.

Comment 4 Triplicates additive recovery testing was conducted to evaluate trueness using fluid industrial sludge fertilizers (2 samples) and composted sludge fertilizers (6 samples). As a result, the mean recovery at the concentration level of 1 mg/kg - 9 mg/kg, 0.1 mg/kg - 0.9 mg/kg and 0.02 mg /kg - 0.04 mg /kg are 85.0 % - 105.9 %, 90.6 % - 108.5 % and 95 % as arsenic (As) in an actual article respectively.

The results of the repeatability tests on different days using two kinds of fluid sludge fertilizers to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 3 µg /kg.
Simultaneous measurement of multiple elements by an ICP-MS is available. In this case, transfer a predetermined amount of cadmium standard solution (Cd 0.1 mg/mL, 1 mg/mL or 10 mg/mL), lead standard solution (Pb 0.1 mg/mL, 1 mg/mL or 10 mg/mL), nickel standard solution (Ni 0.1 mg/mL, 1 mg/mL or 10 mg/mL), chromium standard solution (Cr 0.1 mg/mL, 1 mg/mL or 10 mg/mL) and arsenic standard solution (As 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology to a volumetric flask to mix, dilute with nitric acid (1+19) to prepare a mixed standard solution (Cd 0.5 µg/mL, Pb 5 µg/mL, Ni 5 µg/mL, Cr 5 µg/mL, As 5 µg/mL) (1)(2)(3). Transfer 0.02 mL - 2 mL of mixed standard solutions to 100-mL volumetric flasks (9) step-by-step, add 10 mL of rhodium standard solution (Rh 0.1 µg/mL) and rhenium standard solution (Re 0.1 µg/mL) (5) respectively as internal standard and add nitric acid (1+19) up to the marked line to prepare mixed standard solutions for calibration curve preparation (1)(2)(3) within the concentration range in Table 2. Conduct procedures similarly as (4.2) b) - c) under the measurement conditions in Table 2 and calculate the respective element concentrations in the analytical samples.

**Comment 5** Simultaneous measurement of multiple elements by an ICP-MS is available. In this case, transfer a predetermined amount of cadmium standard solution (Cd 0.1 mg/mL, 1 mg/mL or 10 mg/mL), lead standard solution (Pb 0.1 mg/mL, 1 mg/mL or 10 mg/mL), nickel standard solution (Ni 0.1 mg/mL, 1 mg/mL or 10 mg/mL), chromium standard solution (Cr 0.1 mg/mL, 1 mg/mL or 10 mg/mL) and arsenic standard solution (As 0.1 mg/mL, 1 mg/mL or 10 mg/mL) traceable to National Metrology to a volumetric flask to mix, dilute with nitric acid (1+19) to prepare a mixed standard solution (Cd 0.5 µg/mL, Pb 5 µg/mL, Ni 5 µg/mL, Cr 5 µg/mL, As 5 µg/mL) (1)(2)(3). Transfer 0.02 mL - 2 mL of mixed standard solutions to 100-mL volumetric flasks (9) step-by-step, add 10 mL of rhodium standard solution (Rh 0.1 µg/mL) and rhenium standard solution (Re 0.1 µg/mL) (5) respectively as internal standard and add nitric acid (1+19) up to the marked line to prepare mixed standard solutions for calibration curve preparation (1)(2)(3) within the concentration range in Table 2. Conduct procedures similarly as (4.2) b) - c) under the measurement conditions in Table 2 and calculate the respective element concentrations in the analytical samples.

<table>
<thead>
<tr>
<th>Name of test item</th>
<th>Concentration of element (ng/mL)</th>
<th>Monitor ion (m/z)</th>
<th>Concentration of element (ng/mL)</th>
<th>Monitor ion (m/z)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arsenic</td>
<td>As 1 - 100</td>
<td>75</td>
<td>Rh 10</td>
<td>103</td>
</tr>
<tr>
<td>Cadmium</td>
<td>Cd 0.1 - 10</td>
<td>111, 114</td>
<td>Rh 10</td>
<td>103</td>
</tr>
<tr>
<td>Nickel</td>
<td>Ni 1 - 100</td>
<td>60, 58</td>
<td>Rh 10</td>
<td>103</td>
</tr>
<tr>
<td>Chromium</td>
<td>Cr 1 - 100</td>
<td>53, 52, 50</td>
<td>Rh 10</td>
<td>103</td>
</tr>
<tr>
<td>Lead</td>
<td>Pb 1 - 100</td>
<td>208, 206, 207</td>
<td>Re 10</td>
<td>187</td>
</tr>
</tbody>
</table>

1) Rh: Rhodium, Re: Rhenium
Comment 6  The presence and the extent of interference to the measurement mass number of an element subjected to measurement and an internal standard element can be estimated by qualitative analysis using an ICP-MS in advance to quantitation. Select a measurement mass number considering the extent of interference. However, a mass number cannot be changed in the measurement of arsenic. A magnetic sector double-focusing mass spectrometer or a collision reaction cell specified in JIS K0133 can be used as a method to reduce spectrum interference.

References

(5) Flow sheet for arsenic: The flow sheet for arsenic in fluid sludge fertilizers is shown below:

Figure 1  Flow sheet for arsenic in fluid sludge fertilizers
(Extraction procedure)

Figure 2  Flow sheet for arsenic in fluid sludge fertilizers
(Measurement procedure)
5.3 Cadmium
5.3.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.3.a-2017 or Cd.a-1.

Pretreat an analytical sample with incineration and nitric acid-hydrochloric acid (1+3), spray into an acetylene-air flame, and measure the atomic absorption with cadmium at a wavelength of 228.8 nm to obtain cadmium (Cd) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Cadmium standard solution (Cd 0.1 mg/mL): A cadmium standard solution (Cd 0.1 mg/mL) traceable to National Metrology.
   e) Cadmium standard solution (Cd 10 µg/mL): Transfer 10 mL of cadmium standard solution (Cd 0.1 mg/mL) to a 100-mL volumetric flask, and add hydrochloric acid (1+23) up to the marked line.
   f) Cadmium standard solutions (Cd 0.05 µg - 0.5 µg/mL) for the calibration curve preparation: Transfer 2.5 mL - 25 mL of cadmium standard solution (Cd 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in e) and f).

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the cadmium standard solution in (2), a cadmium standard solution for the calibration curve preparation can be prepared by using a cadmium standard solution (Cd 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction function.
      1) Light source: A cadmium hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
      2) Gas: Gas for heating by flame
         (i) Fuel gas: acetylene
         (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

Note (3) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char (4).

c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (4).

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass (5), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (6) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

**Note**

(4) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

(5) The watch glass can be removed.

(6) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

**Comment 2**

Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

**Comment 3**

The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:

   Analytical line wavelength: 228.8 nm

b) **Calibration curve preparation**

   1) Spray the cadmium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 228.8 nm.

   2) Prepare a curve for the relationship between the cadmium concentration and the indicated value of the cadmium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

   1) Subject the sample solution (7) to the same procedure as in b) 1) to read the indicated value.

   2) Subject the blank test solution to the same procedure as in b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.

   3) Obtain the cadmium content from the calibration curve, and calculate cadmium (Cd) in the analytical sample.

**Note**

(7) If there is a possibility that the cadmium concentration in the sample solution will
exceed the maximum limit of the calibration curve, dilute a predetermined amount with hydrochloric acid (1+23).

**Comment 4** Instead of the correction method in c) 2), the cadmium (Cd) in the analytical sample can also be corrected by obtaining the cadmium content in the blank test solution.

**Comment 5** Recovery testing was conducted using industrial sludge fertilizer and composted sludge fertilizer (5 samples); as a result, the recovery at the concentration level of 5 mg/kg and 0.5 mg/kg were 97.5 % - 99.2 % and 96.7 % - 99.7 %, respectively. Table 1 shows results and analysis results from a collaborative study for test method validation. Additionally, the minimum limit of quantification of this test method is about 0.1 mg/kg.

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories 1)</th>
<th>Mean 2) (mg/kg)</th>
<th>RSD 3) (%)</th>
<th>RSD 4) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sewage sludge fertilizer a</td>
<td>10</td>
<td>1.50</td>
<td>5.5</td>
<td>6.4</td>
</tr>
<tr>
<td>Sewage sludge fertilizer b</td>
<td>10</td>
<td>3.35</td>
<td>1.2</td>
<td>4.2</td>
</tr>
<tr>
<td>Composted sludge fertilizer a</td>
<td>10</td>
<td>1.96</td>
<td>1.0</td>
<td>4.4</td>
</tr>
<tr>
<td>Composted sludge fertilizer b</td>
<td>11</td>
<td>3.81</td>
<td>1.9</td>
<td>3.2</td>
</tr>
<tr>
<td>Composted sludge fertilizer c</td>
<td>10</td>
<td>1.80</td>
<td>3.5</td>
<td>4.9</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (n = number of laboratories x number of samples (2))
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

**References**


Flow sheet for cadmium: The flow sheet for cadmium in fertilizers is shown below:

- **5.00 g analytical sample**: 200-mL - 300-mL tall beaker.
- **Charring**: Heat gently
- **Inceration**: Ignite at 450 °C ± 5 °C, 8 - 16 hours
- **Standing to cool**: Room temperature
  - A small amount of water
  - About 10 mL of nitric acid
  - About 30 mL of hydrochloric acid
- **Heating**: Cover with a watch glass, and digest
- **Heating**: Slightly move a watch glass to remove acid
- **Standing to cool**: Room temperature
  - About 25 mL - 50 mL of hydrochloric acid (1+5)
- **Heating**: Cover with a watch glass, and dissolve
- **Standing to cool**: Room temperature
- **Transfer**: 100-mL - 200-mL volumetric flask, water
  - Water (up to the marked line)
- **Filtration**: Type 3 filter paper
- **Sample solution**
- **Measurement**: Atomic absorption spectrometer (228.8 nm)

Figure Flow sheet for cadmium in fertilizers
5.3.b ICP Optical Emission Spectrometry

(1) Summary
The test method is applicable to sludge fertilizers, etc. This testing method is classified as Type D and its symbol is 5.3.b-2017 or Cd.b-1.
Pretreat an analytical sample with incineration, nitric acid - hydrochloric acid (1+3), introduce it to ICP Optical Emission Spectrometry ("ICP-OES") and measure the emission with cadmium at a wavelength of 228.802 nm to quantify the cadmium (Cd) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.

   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Cadmium standard solution (Cd 0.1 mg/mL): A cadmium standard solution (Cd 0.1 mg/mL) traceable to National Metrology.
   e) Cadmium standard solution (Cd 0.25 µg/mL) (1) (2): Dilute a predetermined amount of cadmium standard solution (Cd 0.1 mg/mL) with hydrochloric acid (1+23) to prepare a cadmium standard solution (Cd 0.25 µg/mL)

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the cadmium standard solution in (2), a cadmium standard solution for the calibration curve preparation can be prepared by using a cadmium standard solution (Cd 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity.
      b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
      c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char (3).
   c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (3).
   d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
   e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
   f) Slightly move the watch glass (4), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
   g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (5) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
   h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

**Note**  
(3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

(4) The watch glass can be removed.

(5) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

**Comment 2** Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) **Measurement**: Conduct measurement (Standard Addition Method) according to JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

**a) Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

- Analytical line wavelength: 228.802 nm

**b) Calibration curve preparation and sample measurement**

1) Put 5mL of sample solution to three 10-mL volumetric flasks respectively.

2) Add 2mL and 4 mL of cadmium standard solution (0.25 μg/mL) to volumetric flasks of 1) above, and further add hydrochloric acid (1+23) to the marked line to make a sample solution of Standard Addition Method.

3) Add hydrochloric acid (1+23) to the marked line of the remaining volumetric flask of 1) above to make a sample solution without a standard solution.

4) Spray the sample solution of Standard Addition Method and the sample solution without a standard solution into the induction plasma, and read the indicated value at a wavelength of 228.802 nm.

5) Transfer 5 mL of blank test solution to a 10-mL volumetric flask, conduct the same procedures as in 3) - 4) to read the indicated value, and correct the indicated value obtained from the respective sample solutions.

6) Prepare a curve for the relationship between the added cadmium concentration and the corrected indicated value of the sample solution for Standard Addition Method and sample solution without a standard solution.

7) Obtain the cadmium content from the intercept of the calibration curve to calculate cadmium (Cd) in the analytical sample

**Comment 4** Instead of the correction method in c) 5), the cadmium (Cd) in the analytical sample can also be corrected by obtaining the cadmium content in the blank test solution.

**Comment 5** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.9.1.b Comment 5.

**Comment 6** The comparison of the measurement value (\(x_i\): 0.003 mg/kg - 3.32 mg/kg) of ICP Optical Emission Spectrometry and the measurement value (\(y_i\)) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (49 samples). As a result, a regression equation was \(y = -0.03 + 1.009x\) and its
correlation coefficient ($r$) was 0.996. Triplicates measurement for each one sample of sewage sludge fertilizer, human waste sludge fertilizer, industrial sludge fertilizer, mixed sludge fertilizer, calcined sludge fertilizers and composted sludge fertilizer was conducted. As a result, a repeatability obtained was 0.8 % - 4.1 % as a relative standard deviation. Additionally, the minimum limit of quantification of this testing method is 0.2 mg/kg.

References

(5) Flow sheet for cadmium: The flow sheet for cadmium in fertilizers is shown below:

![Flow sheet for cadmium](image)

Figure 1 Flow sheet for cadmium in sludge fertilizers (Extraction procedure)
Sample solution

Aliquot 5 mL
10-mL volumetric flask, 3 flasks
0, 2 and 4 mL of cadmium standard solution (0.25 μg/mL)

Measurement
Hydrochloric acid (1:23) (up to the marked line)
ICP Optical Emission Spectrometer (228.802 nm)

Figure 2  Flow sheet for cadmium in sludge fertilizers
(Measurement procedure)
5.3.c ICP Mass Spectrometry (Fluid sludge fertilizers)

(1) **Summary**
The test method is applicable to fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.3.c-2017 or Cd.c-1.

Add nitric acid – hydrogen peroxide to an analytical sample to heat and extract by microwave irradiation. After adding an internal standard element, introduce it to an ICP Mass Spectrometer (ICP-MS) and measure the respective indicated values of cadmium and an internal standard element with mass/charge number (m/z) and obtain cadmium (Cd) in the analytical sample from the ratio of the indicated value for cadmium and the indicated value for an internal standard element. In addition, the performance of this testing method is shown in **Comment 4**.

(2) **Reagent, etc.:** Reagents and water are as shown below.
   a) **Water:** Water of A4 specified in JIS K 0557.
   b) **Nitric acid:** A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) **Nitric acid:** Nitric acid used to dilute a standard solution and a sample solution is a highly pure reagent specified in JIS K 9901.
   d) **Hydrogen peroxide:** A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.
   e) **Rhodium standard solution (Rh 1 mg/mL):** A rhodium standard solution (Rh 1 mg/mL) traceable to National Metrology.
   f) **Rhodium standard solution (Rh 0.1 µg/mL) \(^{(1)(2)(3)}\):** Dilute a predetermined amount of rhodium standard solution (Rh 1 mg/mL) with nitric acid (1+19) to prepare a rhodium standard solution (Rh 0.1 µg/mL).
   g) **Rhenium standard solution (Re 1 mg/mL) \(^{(4)}\):** A rhenium standard solution (Re 1 mg/mL) traceable to National Metrology.
   h) **Rhenium standard solution (Re 0.1 µg/mL) \(^{(1)(2)(3)(4)}\):** Dilute a predetermined amount of rhenium standard solution (Re 1 mg/mL) with nitric acid (1+19) to prepare a rhenium standard solution (Re 0.1 µg/mL).
   i) **Cadmium standard solution (Cd 0.1 mg/mL):** A cadmium standard solution (Cd 0.1 mg/mL) traceable to National Metrology.
   j) **Cadmium standard solution (Cd 0.5 µg/mL) \(^{(1)(2)(3)}\):** Dilute a predetermined amount of cadmium standard solution (Cd 1 mg/mL) with nitric acid (1+19) to prepare a cadmium standard solution (Cd 0.5 µg/mL).
   k) **Cadmium standard solutions (Cd 0.1 ng/mL - 100 ng/mL) for the calibration curve preparation \(^{(1)(2)(3)}\):** Transfer 0.02 mL - 2 mL of cadmium standard solution (Cd 0.5 µg/mL) to 100-mL volumetric flasks step-by-step, and add 10 mL of rhodium standard solutions (Rh 0.1 µg/mL) as internal standard respectively \(^{(5)}\) and add nitric acid (1+19) up to the marked line.
   l) **Blank test solution for the calibration curve preparation \(^{(1)(2)(3)}\):** Transfer 10 mL of rhodium standard solution (Rh 0.1 µg/mL) to 100-mL volumetric flasks as internal standard and add nitric acid (1+19) up to the marked line.

**Note**  
(1) This is an example of preparation; prepare an amount as appropriate.
(2) Store in cool and dark place, and do not use after 1 month after preparation.
(3) For storage, use a sealable container made of materials such as polypropylene containing no cadmium.
(4) Use when measuring lead simultaneously.
(5) Add an internal solution that is 1/10 of the volume to be prepared.

**Comment 1** Instead of the cadmium standard solution in (2), a cadmium standard solution for the calibration curve preparation can be prepared by using a cadmium standard solution
(Cd 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) **Instruments**: Instruments are as shown below:
   a) **ICP Mass Spectrometer**: High-frequency plasma mass spectrometer specified in JIS K 0133.
   1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.995 % in purity.
   b) **Pressure vessel decomposing device**: A device which pressurizes the inside of a vessel by putting acid, etc. to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.
   1) **The main part of a decomposing device**: In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.
   2) **Exhaust system**: A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.
   3) **Airtight vessel**: A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.
   c) **Centrifugal separator**: A centrifugal separator that can work at about 1700 × g.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.
   a) Weigh 20.0 g (6) of an analytical sample, and put it in a 100-mL volumetric flask.
   b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.
   c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave (7).
   d) Ignite at 240 °C ± 5 °C for no less than 10 minutes (7) to decompose (8).
   e) After standing to cool, transfer the solution to a 50-mL volumetric flask (9) with water.
   f) Add water up to the marked line and transfer it to a 50-mL ground-in stopper centrifugal precipitate tube (9).
   g) Centrifuge it at 1700 × g centrifugal force for about five minutes (10) and use the supernatant as the sample solution.
   h) As a blank test, conduct the procedures in b) - g) using another airtight vessel to prepare the blank test solution.

**Note** (6) The maximum limit of solid content, which is converted from moisture content, in the sampling volume 20.0 g of an analytical sample is about 0.5 g. If solid content is likely to exceed the limit, reduce sampling volume as necessary.

(7) Condition examples for a microwave decomposing device: 0 min (room temperature) → 10min (240 °C) → 20 min (240 °C) → 40 min (room temperature), initial output 1400 W

(8) When organic matters still remain, for example the digestion solution is colored, repeat the procedures in (4.1) b) - d).

(9) The vessel should be made of polypropylene, etc. to not affect the measurement.

(10) 16.5-cm of radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.
Comment 2  The procedure in (4.1) is the same as the procedure in (4.1) of 5.1.b.

(4.2) Measurement: Conduct measurement (Internal Standard Method) according to JIS K 0113 and as shown below. Specific measurement procedures are according to the operation method of the ICP Mass Spectrometer used in measurement.

a) Measurement conditions for the ICP Mass Spectrometer: Set up the measurement conditions for the ICP Mass Spectrometer considering the following:
   Cadmium: monitor ion (m/z): 111, 114
   Rhodium: monitor ion (m/z): 103

b) Calibration curve preparation
   1) Spray cadmium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into the inductively coupled plasma, and read the ratio of the respective ion count numbers for the monitor ion of an element subjected to measurement and an internal standard element.
   2) Prepare a curve for the relationship between the concentration and the ratio of the ion count number of an element subjected to measurement.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.005 µg - 0.5 µg as Cd) to a 50-mL volumetric flask.
   2) Add 5mL of internal standard solution and add nitric acid (1+19) to the marked line.
   3) Conduct procedures similarly as in b) 1) to read the ratio of the ion count numbers.
   4) For blank test solution, conduct procedures similarly as in 1) - 3) to correct the ratio of the ion count number obtained for the sample solution.
   5) Obtain the cadmium content from the calibration curve, and calculate cadmium (Cd) in the analytical sample.

Comment 3 Instead of the correction method in c) 4), the cadmium (Cd) in the analytical sample can also be corrected by obtaining the cadmium content in the blank test solution.

Comment 4 Triplicates additive recovery testing was conducted to evaluate trueness using fluid industrial sludge fertilizers (2 samples), and composted sludge fertilizers (6 samples). As a result, the mean recovery at the concentration level of 0.1 mg/kg - 0.9 mg/kg, 0.01 mg/kg - 0.09 mg/kg and 2 µg /kg - 4 µg /kg are 89.4 % - 108.5 %, 91.0 % - 112.0 % and 96.3 % - 108.5 % as cadmium (Cd) in an actual article respectively. The results of the repeatability tests on different days using two kinds of fluid sludge fertilizers to evaluate repeatability were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 2 µg /kg.
Comment 5 Simultaneous measurement of multiple elements by an ICP-MS is available. In that case, see 5.2.c Comment 5.

Comment 6 The presence and the extent of interference to the measurement mass number of an element subjected to measurement and an internal standard element can be estimated by qualitative analysis using an ICP-MS in advance to quantitation. Select a measurement mass number considering the extent of interference. However, a mass number cannot be changed in the measurement of arsenic. A magnetic sector double-focusing mass spectrometer or a collision reaction cell specified in JIS K0133 can be used as a method to reduce spectrum interference.

References

(5) Flow sheet for cadmium: The flow sheet for cadmium in fluid sludge fertilizers is shown below:

![Flow sheet for cadmium](image)

Figure 1 Flow sheet for cadmium in liquid sludge fertilizers (Extraction procedure)
Figure 2  Flow sheet for cadmium in liquid sludge fertilizers
(Measurement procedure)
5.3.d ICP Mass Spectrometry

(1) Summary

The testing method is applicable to sludge fertilizers excluding fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.3.d-2017 or Cd.d-1.

Add nitric acid–hydrogen peroxide to an analytical sample to heat and extract by microwave irradiation. After adding an internal standard element, introduce it to an ICP Mass Spectrometer (ICP-MS) and measure the respective indicated values of cadmium and an internal standard element with mass/ charge number \((m/z)\) and obtain cadmium (Cd) in the analytical sample from the ratio of the indicated value for cadmium and the indicated value for an internal standard element. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Nitric acid: Nitric acid used to dilute a standard solution and a sample solution is a highly pure reagent specified in JIS K 9901.

d) Hydrogen peroxide: A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.

e) Rhodium standard solution (Rh 1 mg/mL): A rhodium standard solution (Rh 1 mg/mL) traceable to National Metrology.

f) Rhodium standard solution (Rh 0.1 µg/mL)\(^{(1)}\)(\(^{(2)}\)(\(^{(3)}\))): Dilute a predetermined amount of rhodium standard solution (Rh 1 mg/mL) with nitric acid (1+19) to prepare a rhodium standard solution (Rh 0.1 µg/mL).

g) Cadmium standard solution (Cd 0.1 mg/mL): A cadmium standard solution (Cd 0.1 mg/mL) traceable to National Metrology.

h) Cadmium standard solution (Cd 0.5 µg/mL)\(^{(1)}\)(\(^{(2)}\)(\(^{(3)}\))): Dilute a predetermined amount of cadmium standard solution (Cd 1 mg/mL) with nitric acid (1+19) to prepare a cadmium standard solution (Cd 0.5 µg/mL).

i) Cadmium standard solutions (Cd 0.1 ng/mL - 100 ng/mL) for the calibration curve preparation\(^{(1)}\)(\(^{(2)}\)(\(^{(3)}\))): Transfer 0.02 mL - 2 mL of cadmium standard solution (Cd 0.5 µg/mL) to 100-mL volumetric flasks step-by-step, and add 10 mL of rhodium standard solutions (Rh 0.1 µg/mL) as internal standard respectively \(^{(4)}\) and add nitric acid (1+19) up to the marked line.

j) Blank test solution for the calibration curve preparation\(^{(1)}\)(\(^{(2)}\)(\(^{(3)}\))): Transfer 10 mL of rhodium standard solution (Rh 0.1 µg/mL) to 100-mL volumetric flasks as internal standard \(^{(4)}\) and add nitric acid (1+19) up to the marked line.

Note  
(1) This is an example of preparation; prepare an amount as appropriate.

(2) Store in cool and dark place, and do not use after 1 month after preparation.

(3) For storage, use a sealable container made of materials such as polypropylene containing no cadmium.

(4) Add an internal solution that is 1/10 of the volume to be prepared.

Comment 1  Instead of the cadmium standard solution in (2), a cadmium standard solution for the calibration curve preparation can be prepared by using a cadmium standard solution (Cd 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:

a) ICP Mass Spectrometer: High-frequency plasma mass spectrometer specified in JIS K 0133.

b) Pressure vessel decomposing device: A device which pressurizes the inside of a vessel by
putting acid, etc. to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.

1) **The main part of a decomposing device**: In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.

2) **Exhaust system**: A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.

3) **Airtight vessel**: A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.

c) **Centrifugal separator**: A centrifugal separator that can work at about 1700 × g.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 0.20 g of an analytical sample, and put it into an airtight vessel.

b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.

c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave.

d) Ignite at 240 °C ± 5 °C for no less than 10 minutes to decompose.

e) After standing to cool, transfer the solution to a 50-mL volumetric flask with water.

f) Add water up to the marked line and transfer it to a 50-mL ground-in stopper centrifugal precipitate tube.

g) Centrifuge it at 1700 × g centrifugal force for about five minutes and use the supernatant as the sample solution.

h) As a blank test, conduct the procedures in b) - g) using another airtight vessel to prepare the blank test solution.

Note (5) Leaving at rest overnight is preferable.

(6) Condition examples for a microwave decomposing device: 0 min (room temperature) → 10 min (240 °C) → 20 min (240 °C) → 40 min (room temperature), initial output 1400 W

(7) When organic matters still remain, for example the digestion solution is colored, repeat the procedures in (4.1) b) - d).

(8) The vessel should be made of polypropylene, etc. to not affect the measurement.

(9) 16.5-cm of radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

(4.2) **Measurement**: Conduct measurement (Internal Standard Method) according to JIS K 0113 and as shown below. Specific measurement procedures are according to the operation method of the ICP Mass Spectrometer used in measurement.

a) **Measurement conditions for the ICP Mass Spectrometer**: Set up the measurement conditions for the ICP Mass Spectrometer considering the following:

   Cadmium: monitor ion (m/z): 111

   Rhodium: monitor ion (m/z): 103
b) **Calibration curve preparation**
1) Spray cadmium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into the inductively coupled plasma, and read the ratio of the respective ion count numbers for the monitor ion of an element subjected to measurement and an internal standard element.
2) Prepare a curve for the relationship between the concentration and the ratio of the ion count number of an element subjected to measurement.

c) **Sample measurement**
1) Transfer a predetermined amount of the sample solution (the equivalents of 0.005 µg - 0.5 µg as Cd) to a 50-mL volumetric flask (8).
2) Add 5mL of internal standard solution (4) and add nitric acid (1+19) to the marked line.
3) Conduct procedures similarly as in 1) to read the ratio of the ion count number.
4) For blank test solution, conduct procedures similarly as in 1) - 3) to correct the ratio of the ion count number obtained for the sample solution.
5) Obtain the cadmium content from the calibration curve, and calculate cadmium (Cd) in the analytical sample.

**Comment 2** Instead of the correction method in c) 4), the cadmium (Cd) in the analytical sample can also be corrected by obtaining the cadmium content in the blank test solution.

**Comment 3** The comparison of the measurement value (x_i: 0.18 mg/kg - 3.02 mg/kg) of ICP-MS and the measurement value (y_i) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (26 samples). As a result, a regression equation was y = 0.0402 + 1.01x and its correlation coefficient (r) was 0.997.

The results of the repeatability tests on different days using two kinds of sludge fertilizers to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 0.1 mg/kg.

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T^1)</td>
<td>Average^2)</td>
<td>s_x^3)</td>
</tr>
<tr>
<td>Composted sludge fertilizer 1</td>
<td>5</td>
<td>0.263</td>
<td>0.022</td>
</tr>
<tr>
<td>Composted sludge fertilizer 2</td>
<td>5</td>
<td>2.39</td>
<td>0.04</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) × the number of duplicate testing (2))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Intermediate standard deviation
6) Intermediate relative standard deviation

**Comment 4** The presence and the extent of interference to the measurement mass number of an element subjected to measurement and an internal standard element can be estimated by qualitative analysis using an ICP-MS in advance to quantitation. Select a measurement mass number considering the extent of interference. A magnetic sector double-focusing mass spectrometer or a collision reaction cell specified in JIS K0133.
can be used as a method to reduce spectrum interference.

**References**


(5) **Flow sheet for cadmium:** The flow sheet for cadmium in solid sludge fertilizers is shown below:

![Flow sheet for cadmium](#)

- 0.20 g analytical sample
- Airtight vessel
- → Nitric acid, 2.5 mL
- → Hydrogen peroxide, 2 mL
- Microwave decomposition
- Transfer
- 50-mL volumetric flask, water
- → Water (up to the marked line)
- Centrifugal separation
- Ground-in stopper centrifugal precipitate tube,

![Flow sheet for cadmium](#)

- Sample solution

Figure 1 Flow sheet for cadmium in solid sludge fertilizers
(Extraction procedure)

- Sample solution

![Flow sheet for cadmium](#)

- Aliquot (predetermined volume)
- 50-mL volumetric flask
- → Rhodium standard solution (Rh 0.1µg/mL) 5 mL (internal standard)
- → Nitric acid (1+19) (up to the marked line)
- Measurement
- ICP-MS (Cd: m/z 111 Rh: m/z 103)

Figure 2 Flow sheet for cadmium in solid sludge fertilizers
(Measurement procedure)
5.4 Nickel
5.4.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.4.a-2017 or Ni.a-1.
Pretreat an analytical sample with incineration and nitric acid–hydrochloric acid (1+3), spray into an acetylene–air flame, and measure the atomic absorption with nickel at a wavelength of 232.0 nm to obtain nickel (Ni) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Nickel standard solution (Ni 0.1 mg/mL): A nickel standard solution (Ni 0.1 mg/mL) traceable to National Metrology.
   e) Nickel standard solutions (Ni 0.5 µg/mL - 5 µg/mL) for the calibration curve preparation (1) (2): Transfer 2.5 mL - 25 mL of nickel standard solution (Ni 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Blank test solution for the calibration curve preparation (1) (2): Hydrochloric acid (1+23) used in the procedures in e).

Note (1) This is an example of preparation; prepare an amount as appropriate.
       (2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the nickel standard solution in (2), a nickel standard solution for the calibration curve preparation can be prepared by using a nickel standard solution (Ni 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (3) function.
      1) Light source: A nickel hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
      2) Gas: Gas for heating by flame
         (i) Fuel gas: acetylene
         (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C .
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

Note (3) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char (4).
c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (4).
d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
f) Slightly move the watch glass (5), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (6) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note  (4) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.  
(5) The watch glass can be removed.  
(6) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 2 Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:  
Analytical line wavelength: 232.0 nm

b) Calibration curve preparation
1) Spray the nickel standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 232.0 nm.
2) Prepare a curve for the relationship between the nickel concentration and the indicated value of the nickel standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
1) Subject the sample solution (7) to the same procedure as in b) 1) to read the indicated value.
2) Subject the blank test solution to the same procedure as in b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.
3) Obtain the nickel content from the calibration curve, and calculate nickel (Ni) in the analytical sample.

Note  (7) If there is a possibility that the nickel concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount with hydrochloric acid (1+23).
Comment 4 Instead of the correction method in c) 2), the nickel (Ni) in the analytical sample can also be corrected by obtaining the nickel content in the blank test solution.

Comment 5 Recovery testing was conducted using industrial sludge fertilizer and composted sludge fertilizer (5 samples); as a result, the recovery at the concentration level of 300 mg/kg and 30 mg/kg were 98.5 % - 100.3 % and 97.1 % - 99.9 %, respectively. Table 1 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of this test method is about 1 mg/kg.

Table 1 Results and analysis results from a collaborative study for nickel test method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (^2) (mg/kg)</th>
<th>(RSD_r) (^3) (%)</th>
<th>(RSD_R) (^4) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sewage sludge fertilizer a</td>
<td>11</td>
<td>56.9</td>
<td>1.1</td>
<td>4.6</td>
</tr>
<tr>
<td>Sewage sludge fertilizer b</td>
<td>11</td>
<td>21.8</td>
<td>2.2</td>
<td>3.9</td>
</tr>
<tr>
<td>Composted sludge fertilizer a</td>
<td>11</td>
<td>28.9</td>
<td>1.3</td>
<td>6.4</td>
</tr>
<tr>
<td>Composted sludge fertilizer b</td>
<td>11</td>
<td>28.5</td>
<td>1.8</td>
<td>4.4</td>
</tr>
<tr>
<td>Composted sludge fertilizer c</td>
<td>12</td>
<td>58.3</td>
<td>1.6</td>
<td>4.4</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean \( (n = \text{number of laboratories x number of samples (2)}) \)
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

References


Flow sheet for nickel: The flow sheet for nickel in fertilizers is shown below:

- 5.00 g analytical sample
  - 200-mL - 300-mL tall beaker.
  - Heat gently
  - Ignite at 450 °C ± 5 °C, 8 - 16 hours
- Charring
  - Room temperature
  - A small amount of water
  - About 10 mL of nitric acid
  - About 30 mL of hydrochloric acid
  - Cover with a watch glass, and digest
- Incineration
  - Slightly move a watch glass to remove acid
  - Room temperature
  - About 25 mL - 50 mL of hydrochloric acid (1+5)
  - Cover with a watch glass, and dissolve
- Standing to cool
- Heating
  - Room temperature
  - 100-mL - 200-mL volumetric flask, water
  - Water (up to the marked line)
  - Type 3 filter paper
- Standing to cool
  - Transfer
  - Filtration
  - Sample solution
  - Measurement
  - Atomic absorption spectrometer (232.0 nm)

Figure Flow sheet for nickel in fertilizers
5.4.b ICP Optical Emission Spectrometry

(1) Summary

The test method is applicable to sludge fertilizers, etc. This testing method is classified as Type D and its symbol is 5.4.b-2017 or Ni.b-1.

Pretreat an analytical sample with incineration and nitric acid–hydrochloric acid (1+3), and then introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the emission with nickel at a wavelength of 231.604 nm to obtain nickel (Ni) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

d) Nickel standard solution (Ni 0.1 mg/mL): A nickel standard solution (Ni 0.1 mg/mL) traceable to National Metrology.

e) Nickel standard solutions (Ni 2.5 µg/mL): Dilute a predetermined amount of nickel standard solution (Ni 0.1 mg/mL) with hydrochloric acid (1+23) to prepare a nickel standard solution (Ni 2.5 µg/mL)

Note (1) This is an example of preparation; prepare an amount as appropriate.

(2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the nickel standard solution in (2), a nickel standard solution for the calibration curve preparation can be prepared by using a nickel standard solution (Ni 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:


1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.

c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char

c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass, and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

**Note** (3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.
(4) The watch glass can be removed.
(5) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

**Comment 1** Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

**Comment 2** The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) **Measurement**: Conduct measurement (Standard Addition Method) according to JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) **Measurement conditions for the ICP Atomic Emission Spectrometer**: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

Analytical line wavelength: 231.604 nm

b) **Calibration curve preparation and sample measurement**

1) Put 5mL of sample solution to three 10-mL volumetric flasks respectively.
2) Add 2 mL and 4 mL (0.25 μg/mL) of nickel standard solution to volumetric flasks of 1) above, then add hydrochloric acid (1+23) to the marked line to make a sample solution of Standard Addition Method.
3) Add hydrochloric acid (1+23) to the marked line of the remaining volumetric flask of 1) above to make a sample solution without a standard solution.
4) Spray the sample solution of Standard Addition Method and the sample solution without a standard solution into the induction plasma, and read the indicated value at a wavelength of 231.604 nm.
5) Transfer 5 mL of blank test solution to a 10-mL volumetric flask, conduct the same procedures as in 3) - 4) to read the indicated value, and correct the indicated value obtained from the respective sample solutions.
6) Prepare a curve for the relationship between the added nickel concentration and the corrected indicated value of the sample solution for Standard Addition Method and the sample solution without a standard solution.
7) Obtain the nickel content from the intercept of the calibration curve to calculate nickel (Ni) concentration in the analytical sample.

**Comment 3** Instead of the correction method in c) 5), the nickel (Ni) in the analytical sample can also be corrected by obtaining the nickel content in the blank test solution.

**Comment 4** The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.9.1.b **Comment 5**.

**Comment 5** The comparison of the measurement value (\(x_i\): 8.4 mg/kg - 129 mg/kg) of ICP Optical Emission Spectrometry and the measurement value (\(y_i\)) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (49 samples). As a result, a regression equation was \(y = -0.96 + 1.010x\) and its
correlation coefficient ($r$) was 0.995. Triplicates measurement for each one sample of sewage sludge fertilizer, human waste sludge fertilizer, industrial sludge fertilizer, mixed sludge fertilizer, calcined sludge fertilizers and composted sludge fertilizer was conducted. As a result, a repeatability obtained was 1.0% - 2.6% as a relative standard deviation. Additionally, the minimum limit of quantification of this testing method is about 8 mg/kg.

References

(5) Flow sheet for nickel: The flow sheet for nickel in fertilizers is shown below:

![Flow sheet for nickel]

Figure 1  Flow sheet for nickel in sludge fertilizers (Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
<th>10-mL volumetric flask, 3 flasks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot 5 mL</td>
<td>← 0, 2 and 4 mL of nickel standard solution (0.25 μg / mL)</td>
</tr>
<tr>
<td></td>
<td>← Hydrochloric acid (1:23) (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
<td>ICP Optical Emission Spectrometer (231.604 nm)</td>
</tr>
</tbody>
</table>

Figure 2 Flow sheet for nickel in sludge fertilizers (Measurement procedure)
5.4.c ICP Mass Spectrometry (Fluid sludge fertilizers)

(1) Summary
The test method is applicable to fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.4.c-2017 or Ni.c-1.

Add nitric acid–hydrogen peroxide to an analytical sample to heat and extract by microwave irradiation. After adding an internal standard element, introduce it to an ICP Mass Spectrometer (ICP-MS) and measure the respective indicated values of nickel and an internal standard element with mass/charge number \((m/z)\) and obtain nickel \((Ni)\) in the analytical sample from the ratio of the indicated value for nickel and the indicated value for an internal standard element. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent, etc.: Reagents and water are as shown below.

\(a\) Water: Water of A4 specified in JIS K 0557.

\(b\) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

\(c\) Nitric acid: Nitric acid used to dilute a standard solution and a sample solution is a highly pure reagent specified in JIS K 9901.

\(d\) Hydrogen peroxide: A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.

\(e\) Rhodium standard solution \((Rh 1 \text{ mg/mL})\): A rhodium standard solution \((Rh 1 \text{ mg/mL})\) traceable to National Metrology.

\(f\) Rhodium standard solution \((Rh 0.1 \mu g/mL)\) \(^{(1)}\)\(^{(2)}\)\(^{(3)}\): Dilute a predetermined amount of rhodium standard solution \((Rh 1 \text{ mg/mL})\) with nitric acid \((1+19)\) to prepare a rhodium standard solution \((Rh 0.1 \mu g/mL)\).

\(g\) Rhenium standard solution \((Re 1 \text{ mg/mL})\) \(^{(4)}\): A rhenium standard solution \((Re 1 \text{ mg/mL})\) traceable to National Metrology.

\(h\) Rhenium standard solution \((Re 0.1 \mu g/mL)\) \(^{(1)}\)\(^{(2)}\)\(^{(3)}\)\(^{(4)}\): Dilute a predetermined amount of rhenium standard solution \((Re 1 \text{ mg/mL})\) with nitric acid \((1+19)\) to prepare a rhenium standard solution \((Re 0.1 \mu g/mL)\).

\(i\) Nickel standard solution \((Ni 0.1 \text{ mg/mL})\): A nickel standard solution \((Ni 0.1 \text{ mg/mL})\) traceable to National Metrology.

\(j\) Nickel standard solution \((Ni 5 \mu g/mL)\) \(^{(1)}\)\(^{(2)}\)\(^{(3)}\): Transfer 5 mL of nickel standard solution \((Ni 0.1 \text{ mg/mL})\) to a 100-mL volumetric flask and add nitric acid \((1+19)\) to the marked line.

\(k\) Nickel standard solutions \((Ni 1 \text{ ng/mL} - 100 \text{ ng/mL})\) for the calibration curve preparation \(^{(1)}\)\(^{(2)}\)\(^{(3)}\): Transfer 0.02 mL - 2 mL of nickel standard solution \((Ni 5 \mu g/mL)\) to 100-mL volumetric flasks step-by-step, and add 10 mL of rhodium standard solutions \((Rh 0.1 \mu g/mL)\) as internal standard respectively \(^{(5)}\) and add nitric acid \((1+19)\) up to the marked line.

\(l\) Blank test solution for the calibration curve preparation \(^{(1)}\)\(^{(2)}\)\(^{(3)}\): transfer 10 mL of rhodium standard solution \((Rh 0.1 \mu g/mL)\) to 100-mL volumetric flasks as internal standard \(^{(5)}\) and add nitric acid \((1+19)\) up to the marked line.

Note \(1\) This is an example of preparation; prepare an amount as appropriate.

\(2\) Store in cool and dark place, and do not use after 1 month after preparation.

\(3\) For storage, use a sealable container made of materials such as polypropylene containing no nickel.

\(4\) Use when measuring lead simultaneously.

\(5\) Add an internal solution that is 1/10 of the volume to be prepared.

Comment 1 Instead of the nickel standard solution in \((2)\), a nickel standard solution for the calibration curve preparation can be prepared by using a nickel standard solution \((Ni 1 \text{ mg/mL} \text{ or } 10 \text{ mg/mL})\) traceable to National Metrology.
(3) **Instruments**: Instruments are as shown below:

a) **ICP Mass Spectrometer**: High-frequency plasma mass spectrometer specified in JIS K 0133.

1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.995 % in purity

b) **Pressure vessel decomposing device**: A device which pressurizes the inside of a vessel by putting acid, etc.to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.

1) **The main part of a decomposing device**: In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.

2) **Exhaust system**: A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.

3) **Airtight vessel**: A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.

c) **Centrifugal separator**: A centrifugal separator that can work at about 1700 \( \times g \).

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 20.0 g \((6)\) of an analytical sample, and put it into an airtight vessel.

b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.

c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave \((7)\).

d) Ignite at 240 °C ± 5 °C for no less than 10 minutes \((7)\) to decompose \((8)\).

e) After standing to cool, transfer the solution to a 50-mL volumetric flask \((9)\) with water.

f) Add water up to the marked line and transfer it to a 50-mL ground-in stopper centrifugal precipitate tube \((9)\).

g) Centrifuge it at 1700 \( \times g \) centrifugal force for about five minutes \((10)\) and use the supernatant as the sample solution.

h) As a blank test, conduct the procedures in b) - g) using another airtight vessel to prepare the blank test solution.

**Note** \((6)\) The maximum limit of solid content, which is converted from moisture content, in the sampling volume 20.0 g of an analytical sample is about 0.5 g. If solid content is likely to exceed the limit, reduce sampling volume as necessary.

(7) **Condition examples for a microwave decomposing device**: 0 min (room temperature) → 10min (240 °C) → 20 min (240 °C) → 40 min (room temperature), initial output 1400 W

(8) **When organic matters still remain, for example the digestion solution is colored**, repeat the procedures in (4.1) b) - d).

(9) The vessel should be made of polypropylene, etc. to not affect the measurement.

(10) 16.5-cm of radius and 3000 rpm of revolutions makes about 1700 \( \times g \) centrifugal force.
Comment 2  The procedure in (4.1) is the same as the procedure in (4.1) of 5.1.b.

4.2 Measurement: Conduct measurement (Internal Standard Method) according to JIS K 0113 and as shown below. Specific measurement procedures are according to the operation method of the ICP Mass Spectrometer used in measurement.

a) Measurement conditions for the ICP Mass Spectrometer: Set up the measurement conditions for the ICP Mass Spectrometer considering the following:
   Nickel: monitor ion (m/z): 60, 58
   Rhodium: monitor ion (m/z): 103

b) Calibration curve preparation
   1) Spray nickel standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into the inductively coupled plasma, and read the ratio of the respective ion count numbers for the monitor ion of an element subjected to measurement and an internal standard element.
   2) Prepare a curve for the relationship between the concentration and the ratio of the ion count number of an element subjected to measurement.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 µg - 5 µg as Ni) to a 50-mL volumetric flask (9).
   2) Add 5mL of internal standard solution (5) and add nitric acid (1+19) to the marked line.
   3) Conduct procedures similarly as in b) 1) to read the ratio of the ion count number.
   4) For blank test solution, conduct procedures similarly as in 1) - 3) to correct the ratio of the ion count number obtained for the sample solution.
   5) Obtain the nickel content from the calibration curve, and calculate nickel (Ni) in the analytical sample.

Comment 3  Instead of the correction method in c) 4), the nickel (Ni) in the analytical sample can also be corrected by obtaining the nickel content in the blank test solution.

Comment 4  Triplicates additive recovery testing was conducted to evaluate trueness using fluid industrial sludge fertilizers (2 samples) and composted sludge fertilizers (6 samples). As a result, the mean recovery at the concentration level of 10 mg/kg - 60 mg/kg, 1 mg/kg - 9 mg/kg and 0.1 mg/kg - 0.9 mg/kg are 89.6 % - 99.2 %, 91.5 % - 114.7 % and 96.1 - 103.7 % as nickel (Ni) in an actual article respectively. The results of the repeatability tests on different days using two kinds of fluid sludge fertilizers to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 6 µg /kg.
Comment 5 Simultaneous measurement of multiple elements by an ICP-MS is available. In that case, see 5.2.c Comment 5.

Comment 6 The presence and the extent of interference to the measurement mass number of an element subjected to measurement and an internal standard element can be estimated by qualitative analysis using an ICP-MS in advance to quantitation. Select a measurement mass number considering the extent of interference. However, a mass number cannot be changed in the measurement of arsenic. A magnetic sector double-focusing mass spectrometer or a collision reaction cell specified in JIS K0133 can be used as a method to reduce spectrum interference.

References

(5) Flow sheet for nickel: The flow sheet for nickel in fluid sludge fertilizers is shown below:

```
<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T^1)</td>
<td>Average^2)</td>
<td>S_r^3)</td>
</tr>
<tr>
<td>Composted sludge fertilizer 1</td>
<td>5</td>
<td>8.60</td>
<td>0.44</td>
</tr>
<tr>
<td>Composted sludge fertilizer 2</td>
<td>5</td>
<td>2.04</td>
<td>0.13</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test 5) Intermediate standard deviation
2) Average (the number of test days (T) × the number of duplicate testing (2)) 6) Intermediate relative standard deviation
3) Repeatability standard deviation
4) Repeatability relative standard deviation
```

Flow sheet for nickel:
```
Airtight vessel ← 20.0 g analytical sample
← Nitric acid, 2.5 mL
← Hydrogen peroxide, 2 mL
Microwave decomposition
Transfer 50-mL volumetric flask, water
← Water (up to the marked line)
Centrifugal separation
Ground-in stopper centrifugal precipitate tube, Anout 1700 × g, 5 minutes
Sample solution
```

Figure 1 Flow sheet for nickel in fluid sludge fertilizers (Extraction procedure)
Figure 2  Flow sheet for nickel in fluidsludge fertilizers
(Measurement procedure)
5.5 Chromium

5.5.a Flame atomic absorption spectrometry (Fertilizers containing organic matters)

(1) Summary
This testing method is applicable to fertilizers containing organic matters. This testing method is classified as Type B and its symbol is 5.5.a-2017 or Cr.a-1.

Pretreat an analytical sample with incineration and nitric acid–hydrochloric acid (1+3), spray into an acetylene–air flame, and measure the atomic absorption with chromium at a wavelength of 357.9 nm or 359.3 nm to obtain chromium (Cr) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Interference suppressor solution (1): Dissolve 100 g of potassium disulfate specified in JIS K 8783 in water to make 1000 mL.
   e) Chromium standard solution (Cr 0.1 mg/mL): A chromium standard solution (Cr 0.1 mg/mL) traceable to National Metrology
   f) Chromium standard solutions (Cr 0.5 µg/mL - 5 µg/mL) for the calibration curve preparation (1)(2): Transfer 2.5 mL - 25 mL of chromium standard solution (Cr 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (3), and further add hydrochloric acid (1+23) to the marked line.
   g) Blank test solution for the calibration curve preparation (1)(2): Transfer about 50 mL of interference suppressor solution (3) to a 500-mL volumetric flask, and add hydrochloric acid (1+23) used in the procedure in f) to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) Store at room temperature, and do not use after 6 months after preparation.
   (3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the chromium standard solution in (2), a chromium standard solution for the calibration curve preparation can be prepared by using a chromium standard solution (Cr 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (4) function.
      1) Light source: A chromium hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
      2) Gas: Gas for heating by flame
         (i) Fuel gas: acetylene
         (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

Note (4) There are the continuous source method, the Zeeman method, the non-resonance
spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char \(^{(5)}\).
   c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate \(^{(5)}\).
   d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
   e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
   f) Slightly move the watch glass \(^{(6)}\), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
   g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid \((1+5)\) \(^{(7)}\) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
   h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
   i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note \(^{(5)}\) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.
   \(^{(6)}\) The watch glass can be removed.
   \(^{(7)}\) Add hydrochloric acid \((1+5)\) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid \((1+23)\). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid \((1+5)\) should be added.

Comment 2 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below.
   Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.
   a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
      Analytical line wavelength: 357.9 nm or 359.3 nm \(^{(8)}\)
   b) Calibration curve preparation
      1) Spray the chromium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame \(^{(9)}\), and read the indicated value at a wavelength of 357.9 nm or 359.3 nm \(^{(8)}\).
      2) Prepare a curve for the relationship between the chromium concentration and the indicated value of the chromium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
   c) Sample measurement
      1) Transfer 25 mL of the sample solution \(^{(10)}\) to a 100-mL volumetric flask.
      2) Add about 10 mL of interference suppressor solution \(^{(3)}\), and add hydrochloric acid \((1+23)\) to the marked line.
      3) Subject to the same procedure as in b) 1) to read the indicated value.
      4) Subject the blank test solution to the same procedure as in 1) - 2) and b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.
5) Obtain the chromium content from the calibration curve, and calculate chromium (Cr) in the analytical sample.

Note (8) When background correction is conducted by the Zeeman method, 359.3 nm is recommended as the analytical line wavelength.

(9) Use low-fuel acetylene–air flame. Acetylene–nitrous oxide flame can also be used.

(10) If there is a possibility that the chromium concentration in the sample solution will exceed the maximum limit of the calibration curve, decrease the amount to be transferred.

Comment 3 In an acetylene–air flame, sensitivity is enhanced in high-fuel flame, but interference by coexisting substances such as iron and nickel will also be enhanced. In an acetylene–nitrous oxide flame, such interference hardly affects the result.

Comment 4 Instead of the correction method in e) 4), the chromium (Cr) in the analytical sample can be corrected by obtaining the chromium content in the blank test solution.

Comment 5 Recovery testing was conducted using industrial sludge fertilizer and composted sludge fertilizer (5 samples); as a result, the recovery at the concentration level of 500 mg/kg and 50 mg/kg were 97.5 % - 100.0 % and 95.9 % - 101.9 %, respectively. Table 1 shows results and analysis results from a collaborative study for test method validation. Additionally, the minimum limit of quantification of this test method is about 1 mg/kg.

Table 1  Results and analysis results from a collaborative study for chromium test method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories 1)</th>
<th>Mean 2) (mg/kg)</th>
<th>RSD 3) (%)</th>
<th>RSD 4) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sewage sludge fertilizer a</td>
<td>12</td>
<td>33.6</td>
<td>5.3</td>
<td>15.6</td>
</tr>
<tr>
<td>Sewage sludge fertilizer b</td>
<td>12</td>
<td>26.3</td>
<td>4.9</td>
<td>18.7</td>
</tr>
<tr>
<td>Composted sludge fertilizer a</td>
<td>11</td>
<td>41.3</td>
<td>2.1</td>
<td>11.0</td>
</tr>
<tr>
<td>Composted sludge fertilizer b</td>
<td>12</td>
<td>30.2</td>
<td>5.5</td>
<td>13.8</td>
</tr>
<tr>
<td>Composted sludge fertilizer c</td>
<td>12</td>
<td>85.0</td>
<td>6.4</td>
<td>12.5</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n$ = number of laboratories x number of samples (3))
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

References


2) Yoshinari SAKAKIBARA and Manabu MATSUZAKI: Determination of Cadmium,
Flow sheet for chromium: The flow sheet for chromium in fertilizers containing organic matters is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>5.00 g analytical sample placed in 200-mL - 300-mL tall beaker.</td>
</tr>
<tr>
<td>2.</td>
<td>Charring, Incineration at 450 ºC ± 5 ºC, 8-16 hours.</td>
</tr>
<tr>
<td>3.</td>
<td>Heating up to Room temperature</td>
</tr>
<tr>
<td>4.</td>
<td>A small amount of water, moisten the residue</td>
</tr>
<tr>
<td>5.</td>
<td>About 10 mL of nitric acid</td>
</tr>
<tr>
<td>6.</td>
<td>About 30 mL of hydrochloric acid</td>
</tr>
<tr>
<td>7.</td>
<td>Standing to cool</td>
</tr>
<tr>
<td>8.</td>
<td>Slightly move a watch glass to remove acid</td>
</tr>
<tr>
<td>9.</td>
<td>Heating up to Room temperature</td>
</tr>
<tr>
<td>10.</td>
<td>25 mL - 50 mL of hydrochloric acid (1+5)</td>
</tr>
<tr>
<td>11.</td>
<td>Heating up to Room temperature</td>
</tr>
<tr>
<td>12.</td>
<td>Transfer 100-mL - 200-mL volumetric flask, water</td>
</tr>
<tr>
<td>13.</td>
<td>Filtration Type 3 filter paper</td>
</tr>
<tr>
<td>14.</td>
<td>Aliquot 25 mL 100-mL volumetric flasks</td>
</tr>
<tr>
<td>15.</td>
<td>About 10 mL of interference suppressor solution</td>
</tr>
<tr>
<td>16.</td>
<td>Hydrochloric acid (1+23) (up to the marked line)</td>
</tr>
<tr>
<td>17.</td>
<td>Standing to cool</td>
</tr>
<tr>
<td>18.</td>
<td>Measurement Atomic absorption spectrometer (357.9 nm or 359.3 nm)</td>
</tr>
</tbody>
</table>

Figure: Flow sheet for chromium in fertilizers containing organic matters.
5.5.b Flame atomic absorption spectrometry (Fertilizers mainly containing molten matters, slag, etc.)

(1) Summary
This testing method is applicable to fertilizers mainly containing molten matters, slag, etc. This testing method is classified as Type B and its symbol is 5.5.b-2017 or Cr.b-1.

Add ammonium sulfate to prevent the bumping of an analytical sample, pretreat the analytical sample with nitric acid–sulfuric acid–perchloric acid, and then spray into an acetylene–air flame. Measure the atomic absorption with chromium at a wavelength of 357.9 nm or 359.3 nm to obtain chromium (Cr) in an analytical sample. The performance of this testing method is shown in Comment 4.

In addition, fertilizers containing no organic matters can also be measured by the method of 5.5.c. However, this method should be used for fertilizers that bump when heating. (Fertilizers made of molten matters, slag, etc. may bump in many cases).

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Ammonium sulfate: A reagent of atomic absorption analysis grade or equivalents
   c) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Sulfuric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   e) Perchloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   f) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   g) Interference suppressor solution (1): Dissolve 100 g of potassium disulfate specified in JIS K 8783 in water to make 1000 mL.
   h) Chromium standard solution (Cr 0.1 mg/mL): A chromium standard solution (Cr 0.1 mg/mL) traceable to National Metrology
   i) Chromium standard solutions (Cr 0.01 mg/mL) for the calibration curve preparation (1): Transfer 10 mL of chromium standard solution (Cr 0.1 mg/mL) to 100-mL volumetric flask, and add hydrochloric acid (1+23) to the marked line.
   j) Chromium standard solutions (Cr 0.5 µg/mL - 5 µg/mL) for the calibration curve preparation (1)(2): Transfer 2.5 mL - 25 mL of chromium standard solution (Cr 0.1 mg/mL) or chromium standard solution (Cr 0.01 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (3), and further add hydrochloric acid (1+23) to the marked line.
   k) Blank test solution for the calibration curve preparation (1)(2): Transfer about 50 mL of interference suppressor solution (3) to a 500-mL volumetric flask, and add hydrochloric acid (1+23) used in the procedure in i) and j) to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.
(2) Store at room temperature, and do not use after 6 months after preparation.
(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the chromium standard solution in (2), a chromium standard solution for the calibration curve preparation can be prepared by using a chromium standard solution (Cr 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in
JIS K 0121 with the background correction\(^{(4)}\) function.

1) **Light source**: A chromium hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)

2) **Gas**: Gas for heating by flame  
   (i) Fuel gas: acetylene  
   (ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Hot plate or sand bath**: A hot plate whose surface temperature can be adjusted up to 350 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to no less than 300 °C.

**Note** (4) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) **Test procedure**
(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample and transfer to a 300-mL tall beaker.

b) Add 4 g of ammonium sulfate and moisten the analytical sample with a small amount of water.

c) Add about 10 mL of nitric acid, about 5 mL of sulfuric acid and about 5 mL of perchloric acid. Cover the tall beaker with a watch glass, heat on a hot plate or sand bath gently at 170 °C - 220 °C for no less than 1 hour, and then warm to raise temperature gradually to no less than 300 °C taking no less than 30 minutes\(^{(5)}\) and further heat at no less than 300 °C for 2-3 hours.

d) Slightly move the watch glass\(^{(6)}\), and keep on heating on the hot plate or sand bath to concentrate until the liquid volume becomes about 3 mL\(^{(7)}\).

e) After standing to cool, add about 5 mL of hydrochloric acid (1+10) and about 20 mL of water, cover the tall beaker with a watch glass and heat mildly for about 10 minutes to dissolve\(^{(8)}\).

f) After standing to cool, transfer the solution to a 100-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

g) As a blank test, conduct the procedures in b) - f) using another tall beaker to prepare the blank test solution. However, the heating procedure after adding acid in the procedures in e) should be conducted as shown below. Cover the tall beaker with a watch glass, heat on a hot plate or sand bath at about 170 °C for a while\(^{(8)}\), slightly move the watch glass\(^{(6)}\) and then heat gently for no less than 1 hour to vapor nitric acid. After concentrating to about 15 mL\(^{(9)}\), heat and raise the temperature gradually to no less than 300 °C in more than 30 minutes\(^{(5)}\). When white smoke starts evolving, cover the tall beaker with a watch glass and further heat at no less than 300 °C for 2-3 hours.

**Note** (5) Sudden temperature rise may cause bumping. Therefore, increase the temperature gradually.

(6) If there is no possibility of bumping, the watch glass can be removed.

(7) Drying and hardening it up may cause the chromium to become insoluble. Therefore, care should be taken not to concentrate too much.

(8) Heating at high temperature may cause bumping. Therefore, it is recommended to increase the temperature gradually from about 170 °C to the temperature at which it boils slightly.

(9) If much nitric acid remains, care should be taken because bumping can occur easily when heating temperature rises.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic
absorption spectrometer used in measurement.

a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:

Analytical line wavelength: 357.9 nm or 359.3 nm \(^{(10)}\)

b) Calibration curve preparation

1) Spray the chromium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a low-fuel acetylene-air flame \(^{(11)}\), and read the indicated value at a wavelength of 357.9 nm or 359.3 nm \(^{(10)}\).

2) Prepare a curve for the relationship between the chromium concentration and the indicated value of the chromium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement

1) Transfer 25 mL of the sample solution \(^{(12)}\) to a 100-mL volumetric flask.

2) Add about 10 mL of interference suppressor solution \(^{(3)}\), and add hydrochloric acid (1+17) to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Subject the blank test solution to the same procedure as in 1) - 2) and b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.

5) Obtain the chromium content from the calibration curve, and calculate chromium (Cr) in the analytical sample.

Note \(^{(10)}\) When background correction is conducted by the Zeeman method, 359.3 nm is recommended as the analytical line wavelength.

\(^{(11)}\) Acetylene–nitrous oxide flame can also be used.

\(^{(12)}\) If there is a possibility that the concentration of chromium in the sample solution exceeds the maximum limit of the calibration curve, transfer no more than 10 mL of the sample solution to a 100-mL volumetric flask and add about 10 mL of interference suppressor solution and about 67 mL of hydrochloric acid (1+17), and add water up to the marked line.

Comment 2 In an acetylene–air flame, sensitivity is enhanced in high-fuel flame, but interference by coexisting substances such as iron and nickel will also be enhanced. In an acetylene–nitrous oxide flame, such interference hardly affects the result.

Comment 3 Instead of the correction method in c) 4), the chromium (Cr) in the analytical sample can also be corrected by obtaining the chromium content in the blank test solution.

Comment 4 The comparison of the measurement value \((y_i): 54 \text{ mg/kg} - 4649 \text{ mg/kg}\) of this testing method and the measurement value \((x_i)\) of 5.8 Chromium 5.8.2 Atomic Absorption Spectrometry in Official Methods of Analysis of Fertilizers (1992) was conducted to evaluate trueness using fertilizers (29 samples). As a result, a regression equation was \(y = -6.842 + 0.998x\) and its correlation coefficient \((r)\) was 0.999. The results of the repeatability tests on different days using fused silicic phosphate fertilizer slag silicic fertilizer and compound fertilizer (1 sample for each) to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this test method is about 6 mg/kg.
Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Average&lt;sup&gt;2&lt;/sup&gt;</td>
<td>s&lt;sub&gt;r&lt;/sub&gt;&lt;sup&gt;3&lt;/sup&gt;</td>
</tr>
<tr>
<td>Fused silicic phosphate fertilizer</td>
<td>5</td>
<td>4628</td>
<td>37</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>5</td>
<td>545</td>
<td>5.9</td>
</tr>
<tr>
<td>Slag silicic fertilizer</td>
<td>5</td>
<td>319</td>
<td>3.8</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) × the number of duplicate testing (2))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Intermediate standard deviation
6) Intermediate relative standard deviation

Table 2  Results and analysis results from a collaborative study for the test method validation of chromium

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories&lt;sup&gt;1&lt;/sup&gt;</th>
<th>Mean&lt;sup&gt;2&lt;/sup&gt;</th>
<th>s&lt;sub&gt;r&lt;/sub&gt;&lt;sup&gt;3&lt;/sup&gt;</th>
<th>RSD&lt;sub&gt;r&lt;/sub&gt;&lt;sup&gt;4&lt;/sup&gt;</th>
<th>s&lt;sub&gt;R&lt;/sub&gt;&lt;sup&gt;5&lt;/sup&gt;</th>
<th>RSD&lt;sub&gt;R&lt;/sub&gt;&lt;sup&gt;6&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slag silicic fertilizer</td>
<td>12</td>
<td>63.75</td>
<td>2.02</td>
<td>3.2</td>
<td>3.87</td>
<td>6.1</td>
</tr>
<tr>
<td>Mixed phosphate fertilizer</td>
<td>12</td>
<td>912.9</td>
<td>13.0</td>
<td>1.4</td>
<td>37.3</td>
<td>4.1</td>
</tr>
<tr>
<td>Slag manganese fertilizer</td>
<td>12</td>
<td>2962</td>
<td>74</td>
<td>2.5</td>
<td>176</td>
<td>5.9</td>
</tr>
<tr>
<td>Fused silicic phosphate fertilizer</td>
<td>10</td>
<td>4662</td>
<td>135</td>
<td>2.9</td>
<td>166</td>
<td>3.6</td>
</tr>
<tr>
<td>Compound fertilizers</td>
<td>10</td>
<td>543.6</td>
<td>10.2</td>
<td>1.9</td>
<td>15.4</td>
<td>2.8</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (n = number of laboratories x number of samples (2))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Reproducibility standard deviation
6) Reproducibility relative standard deviation

References

(5) **Flow sheet for chromium**: The flow sheet for chromium in fertilizers mainly containing molten matters, slag is shown below:

- 1.00 g analytical sample
- 300-mL tall beaker.
- 4 g of ammonium sulfate
- A small amount of water, moisten the analytical sample
- About 10 mL of nitric acid
- About 5 mL of sulfuric acid
- About 5 mL of perchloric acid
- Cover with a watch glass, and heat mildly on a hot plate or sand bath at 170 °C - 220 °C for no less than 1 hour
- Raise temperature gradually to no less than 300 °C in more than 30 minutes
- Digest on a hot plate or sand bath for 2 -3 hours
- Slightly move a watch glass and concentrate until the liquid volume becomes about 3 mL
- Room temperature
- About 5 mL of hydrochloric acid (1+5)
- About 20 mL of water
- Cover with a watch glass, and dissolve (heat for about 10 minutes)
- Room temperature
- Transfer 100-mL volumetric flask
- Water (up to the marked line)
- Type 3 filter paper
- Sample solution

Figure 1  Flow sheet for chromium in fertilizers mainly containing molten matters and slag, etc. (Extraction procedure)
<table>
<thead>
<tr>
<th>Sample solution</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aliquot</strong></td>
</tr>
<tr>
<td>(no more than 25 mL)</td>
</tr>
<tr>
<td>100-mL volumetric flasks</td>
</tr>
<tr>
<td>← About 10 mL of interference suppressor solution</td>
</tr>
<tr>
<td>← Hydrochloric acid (1+17) (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
</tr>
<tr>
<td>Atomic absorption spectrometer (357.9 nm or 359.3 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for chromium in fertilizers

mainly containing molten matters and slag, etc. (Measurement procedure)
5.5.c Flame atomic absorption spectrometry (Fertilizers not containing organic matters)

(1) Summary
This testing method is applicable to fertilizers not containing organic matters (including calcined sludge fertilizer). This testing method is classified as Type B and its symbol is 5.5.c-2017 or Cr.c-1. In addition, for fertilizers mainly made of molten matters, slag, etc., care should be taken because they may bump while heating. 5.5.b applies to fertilizers that may bump.

Pretreat an analytical sample with nitric acid–sulfuric acid–perchloric acid, and then spray into an acetylene–air flame, and measure the atomic absorption with chromium at a wavelength of 357.9 nm or 359.3 nm to obtain chromium (Cr) in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

(2) Reagent, etc.: Reagents and water are as shown below.

b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
c) Sulfuric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
d) Perchloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
e) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
f) Interference suppressor solution (1): Dissolve 100 g of potassium disulfate specified in JIS K 8783 in water to make 1000 mL.
g) Chromium standard solution (Cr 0.1 mg/mL): A chromium standard solution (Cr 0.1 mg/mL) traceable to National Metrology.
h) Chromium standard solutions (Cr 0.01 mg/mL) for the calibration curve preparation (1): Transfer 10 mL of chromium standard solution (Cr 0.1 mg/mL) to 100-mL volumetric flask, and add hydrochloric acid (1+23) to the marked line.
i) Chromium standard solutions (Cr 0.5 µg/mL - 5 µg/mL) for the calibration curve preparation (1)(2): Transfer 2.5 mL - 25 mL of chromium standard solution (Cr 0.1 mg/mL) or chromium standard solution (Cr 0.01 mg/mL) to 500-mL volumetric flasks step-by-step, add about 50 mL of interference suppressor solution (3), and further add hydrochloric acid (1+23) to the marked line.
j) Blank test solution for the calibration curve preparation (1)(2): Transfer about 50 mL of interference suppressor solution (3) to a 500-mL volumetric flask, and add hydrochloric acid (1+23) used in the procedure in h) and i) to the marked line.

Note
(1) This is an example of preparation; prepare an amount as appropriate.
(2) Store at room temperature, and do not use after 6 months after preparation.
(3) Add an interference suppressor solution that is 1/10 volume of the volume to be prepared.

Comment 1 Instead of the chromium standard solution in (2), a chromium standard solution for the calibration curve preparation can be prepared by using a chromium standard solution (Cr 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (4) function.
1) Light source: A chromium hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
2) Gas: Gas for heating by flame
(i) Fuel gas: acetylene
(ii) Auxiliary gas: Air sufficiently free of dust and moisture.

b) **Hot plate or sand bath:** A hot plate whose surface temperature can be adjusted up to 350 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to no less than 300 ºC.

**Note** (4) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
b) Add about 10 mL of nitric acid and about 5 mL of sulfuric acid, cover the tall beaker with a watch glass, and leave at rest overnight.
c) Heat mildly on a hot plate or sand bath at 170 ºC - 220 ºC for no less than 30 minutes. After bubbles cease to form, set the temperature of the hot plate or sand bath to no less than 300 ºC (5), and heat until nitroxide (yellow-brown smoke) is no longer generated (6).
d) After standing to cool, add about 5 mL of perchloric acid.
e) Cover the tall beaker with a watch glass, and heat on a hot plate or sand bath at no less than 300 ºC for 2 - 3 hours to digest (7).
f) Slightly move the watch glass (8), and keep on heating on the hot plate or sand bath to concentrate until the liquid volume becomes no more than 2 mL (9).
g) After standing to cool, add about 5 mL of hydrochloric acid (1+10) and about 20 mL of water, cover the tall beaker with a watch glass and heat mildly to dissolve.
h) After standing to cool, transfer the solution to a 100-mL - 200-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

**Note** (5) If there is vigorous bumping, increase the temperature gradually.

(6) Oxidation of carbides by perchloric acid progresses extremely rapidly and explosively. For that reason, add perchloric acid after fully degrading carbides with nitric acid to avoid danger.

(7) When the white smoke of perchloric acid is generated, if the solution is colored such as black-brown or brown, stop heating immediately, and after standing to cool, add nitric acid, and heat again to degrade remaining carbides.

(8) If there is no possibility of bumping, the watch glass can be removed.

(9) When drying and hardening it, chromium may not dissolve completely in g) and the concentration may become a low value.

**Comment 2** The procedure in (4.1) is the same as the procedure in (4.1) of 5.2.a.

**Comment 3** When the analytical sample solidifies in the procedure in (4.1) b), moisten the analytical sample with a small amount of water as necessary in advance.

**Comment 4** It is not necessary to conduct the procedures in (4.1) b) “leave at rest overnight” because the range subjected to analysis does not contain organic matters.

**Comment 5** In some cases, about 10 minutes heating in procedure in (4.1) g) is required.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.
a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following: Analytical line wavelength: 357.9 nm or 359.3 nm

b) Calibration curve preparation
1) Spray the chromium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a low-fuel acetylene–air flame, and read the indicated value at a wavelength of 357.9 nm or 359.3 nm.
2) Prepare a curve for the relationship between the chromium concentration and the indicated value of the chromium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement
1) Transfer 25 mL of the sample solution to a 100-mL volumetric flask.
2) Add about 10 mL of interference suppressor solution, and add hydrochloric acid (1+17) to the marked line.
3) Subject to the same procedure as in b) 1) to read the indicated value.
4) Subject the blank test solution to the same procedure as in 1) - 2) and b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.
5) Obtain the chromium content from the calibration curve, and calculate chromium (Cr) in the analytical sample.

Note (10) When background correction is conducted by the Zeeman method, 359.3 nm is recommended as the analytical line wavelength.

(11) Acetylene–nitrous oxide flame can also be used.

(12) If there is a possibility that the concentration of chromium in the sample solution exceeds the maximum limit of the calibration curve, transfer no more than 10 mL of the sample solution to a 100-mL volumetric flask and add about 10 mL of interference suppressor solution and about 67 mL of hydrochloric acid (1+17), and add water up to the marked line.

Comment 6 In an acetylene-air flame, sensitivity is enhanced in high-fuel flame, but interference by coexisting substances such as iron and nickel will also be enhanced. In an acetylene–nitrous oxide flame, such interference hardly affects the result.

Comment 7 Instead of the correction method in c) 4), the chromium (Cr) in the analytical sample can also be corrected by obtaining the chromium content in the blank test solution.

Comment 8 The comparison of the measurement value \( y_i \): 52 mg/kg - 4052 mg/kg) of this testing method and the measurement value \( x_i \) of 5.8 Chromium 5.8.2 Atomic Absorption Spectrometry in Official Methods of Analysis of Fertilizers (1992) was conducted to evaluate trueness using fertilizers (27 samples). As a result, a regression equation was \( y = -0.405 + 0.994x \) and its correlation coefficient \( r \) was 0.999. The results of the repeatability tests on different days using mixed phosphate fertilizer, compound fertilizer and slag silicic fertilizer (1 sample for each) to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation using calcined sludge fertilizers. Additionally, the minimum limit of quantification of this test method is about 6 mg/kg.
Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$ (mg/kg)</th>
<th>Repeatability $^3$ s$_r$ (mg/kg)</th>
<th>$RSD_r$ $^4$ (%)</th>
<th>Intermediate precision $^5$ s$_{I(T)}$ (mg/kg)</th>
<th>$RSD_{I(T)}$ $^6$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixed phosphate fertilizer</td>
<td>5</td>
<td>3966</td>
<td>96</td>
<td>2.4</td>
<td>107</td>
<td>2.7</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>5</td>
<td>542</td>
<td>6.1</td>
<td>1.1</td>
<td>8.6</td>
<td>1.6</td>
</tr>
<tr>
<td>Slag silicic fertilizer</td>
<td>5</td>
<td>288</td>
<td>7.0</td>
<td>2.4</td>
<td>13</td>
<td>4.4</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) × the number of duplicate testing ($n$))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Intermediate standard deviation
6) Intermediate relative standard deviation

Table 2  Results and analysis results from a collaborative study for chromium test method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories $^1$</th>
<th>Mean $^2$ (mg/kg)</th>
<th>$RSD_r$ $^3$ (%)</th>
<th>$RSD_R$ $^4$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcined sludge fertilizer 1</td>
<td>10</td>
<td>107</td>
<td>5.0</td>
<td>9.7</td>
</tr>
<tr>
<td>Calcined sludge fertilizer 2</td>
<td>9</td>
<td>136</td>
<td>3.4</td>
<td>3.6</td>
</tr>
<tr>
<td>Calcined sludge fertilizer 3</td>
<td>9</td>
<td>182</td>
<td>1.1</td>
<td>2.6</td>
</tr>
<tr>
<td>Calcined sludge fertilizer 4</td>
<td>9</td>
<td>213</td>
<td>1.1</td>
<td>3.9</td>
</tr>
<tr>
<td>Calcined sludge fertilizer 5</td>
<td>9</td>
<td>117</td>
<td>1.8</td>
<td>4.0</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n$ = number of laboratories x number of samples (2))
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

References


(5) **Flow sheet for chromium:** The flow sheet for chromium in fertilizers not containing calcined sludge fertilizer and organic matters is shown below: However, fertilizers that may bump are excluded from the scope of application.

<table>
<thead>
<tr>
<th>Step</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00 g analytical sample</td>
<td>200-mL - 300-mL tall beaker.</td>
</tr>
<tr>
<td>← A small amount of water, moisten the analytical sample (as necessary)</td>
<td></td>
</tr>
<tr>
<td>← About 10 mL of nitric acid</td>
<td></td>
</tr>
<tr>
<td>← About 5 mL of sulfuric acid</td>
<td></td>
</tr>
<tr>
<td>Leaving at rest overnight</td>
<td>Cover with a watch glass</td>
</tr>
<tr>
<td>Heating</td>
<td>Heat gently on a hot plate or a sand bath at 170 °C - 220 °C for no less than 30 minutes</td>
</tr>
<tr>
<td>Heating</td>
<td>Heat on a hot plate or sand bath at no less than 300 °C, until yellow-brown smoke no longer evolves.</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← About 5 mL of perchloric acid</td>
<td>Cover with a watch glass, digest on a hot plate or a sand bath at 300 °C for 2-3 hours</td>
</tr>
<tr>
<td>Heating</td>
<td>Slightly move the watch glass, concentrate until it becomes no more than 2mL</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← About 5 mL of nitric acid</td>
<td>Cover with watch glass, dissolve</td>
</tr>
<tr>
<td>← About 20 mL of water</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td></td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Heating</td>
<td></td>
</tr>
<tr>
<td>Transfer</td>
<td>100-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1  Flow sheet for chromium in fertilizers not containing calcined sludge fertilizer and organic matters (Extraction procedure)
(Excluding fertilizers which may bump while heating)
<table>
<thead>
<tr>
<th>Sample solution</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Aliquot 25 mL</strong></td>
</tr>
<tr>
<td>100-mL volumetric flask</td>
</tr>
<tr>
<td>← About 10 mL of interference suppressor solution</td>
</tr>
<tr>
<td>← Hydrochloric acid (1+17) (up to the marked line)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Measurement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atomic absorption spectrometer (357.9 nm or 359.3 nm)</td>
</tr>
</tbody>
</table>

Figure 2 Flow sheet for chromium in fertilizers not containing calcined sludge fertilizer and organic matters (Measurement procedure)
(Excluding fertilizers which may bump while heating)
5.5.d ICP Optical Emission Spectrometry

(1) Summary
The testing method is applicable to sludge fertilizers, etc. (except for calcined sludge fertilizer) This testing method is classified as Type D and its symbol is 5.5.d-2017 or Cr.d-1.

Pretreat an analytical sample with incineration, nitric acid–hydrochloric acid (1+3), introduce it to ICP Optical Emission Spectrometry (“ICP-OES”) and measure the emission with chromium at a wavelength of 205.552 nm to quantify the chromium (Cr) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Chromium standard solution (Cr 0.1 mg/mL): A chromium standard solution (Cr 0.1 mg/mL) traceable to National Metrology
   d) Chromium standard solutions (Cr 2.5 µg/mL) (1)(2): Dilute a predetermined amount of chromium standard solution (Cr 0.1 mg/mL) with hydrochloric acid (1+23) to prepare a chromium standard solution (Cr 2.5 µg/mL)

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the chromium standard solution in (2), a chromium standard solution for the calibration curve preparation can be prepared by using a chromium standard solution (Cr 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   b) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity
   c) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
   d) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 ºC.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char (3).
   c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (3).
   d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
   e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
   f) Slightly move the watch glass (4), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
   g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (5) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
   h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note (3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

(4) The watch glass can be removed.

(5) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 2 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) Measurement: Conduct measurement (Standard Addition Method) according to JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Optical Emission Spectrometer: Set up the measurement conditions for the ICP Optical Emission Spectrometer considering the following:
   Analytical line wavelength: 205.552 nm

b) Calibration curve preparation and sample measurement
   1) Put 5mL of sample solution to three 10-mL volumetric flasks respectively.
   2) Add 2mL and 4 mL of chromium standard solution (2.5 μg/mL) to volumetric flasks of 1) above, then add hydrochloric acid (1+23) to the marked line to make a sample solution of Standard Addition Method.
   3) Add hydrochloric acid (1+23) to the marked line of the remaining volumetric flask of 1) above to make a sample solution without a standard solution.
   4) Spray the sample solution of Standard Addition Method and the sample solution without a standard solution into the induction plasma, and read the indicated value at a wavelength of 205.552 nm.
   5) Transfer 5 mL of blank test solution to a 10-mL volumetric flask, conduct the same procedures as in 3) - 4) to read the indicated value, and correct the indicated value obtained from the respective sample solutions.
   6) Prepare a curve for the relationship between the added chromium concentration and the corrected indicated value of the sample solution for Standard Addition Method and the sample solution without a standard solution.
   7) Obtain the chromium content from the intercept of the calibration curve to calculate chromium (Cr) in the analytical sample.

Comment 3 Instead of the correction method in e) 5), the chromium (Cr) in the analytical sample can also be corrected by obtaining the chromium content in the blank test solution.

Comment 4 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.9.1.b Comment 5.

Comment 5 The comparison of the measurement value ($x_i$: 12.9 mg/kg - 193 mg/kg) of ICP Optical Emission Spectrometry and the measurement value ($y_i$) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (49 samples). As a result, a regression equation was $y = 1.74+0.971x$ and its correlation coefficient ($r$) was 0.991. Triplicates measurement for each one sample of sewage sludge fertilizer, human waste sludge fertilizer, industrial sludge fertilizer,
mixed sludge fertilizer and composted sludge fertilizer was conducted. As a result, a repeatability obtained was 0.9 % - 2.5 % as a relative standard deviation. Additionally, the minimum limit of quantification of this testing method is 4 mg/kg.

References


(5) Flow sheet for cadmium: The flow sheet for cadmium in fertilizers is shown below:

<table>
<thead>
<tr>
<th>5.00 g analytical sample</th>
<th>200-mL - 300-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>Ignite at 450 ºC ± 5 ºC, 8 - 16 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← A small amount of water, moisten the residue</td>
<td>Cover with a watch glass, and digest for 30 minutes</td>
</tr>
<tr>
<td>← About 10 mL of nitric acid</td>
<td>Slightly move a watch glass to remove acid</td>
</tr>
<tr>
<td>← About 30 mL of hydrochloric acid</td>
<td>Heating</td>
</tr>
<tr>
<td>Heating</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← 25 mL - 50 mL of hydrochloric acid (1+5)</td>
<td>Cover with a watch glass, and dissolve</td>
</tr>
<tr>
<td>Heating</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Transfer</td>
<td>100-mL - 200-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Filtration</td>
<td>Sample solution</td>
</tr>
</tbody>
</table>

Figure 1  Flow sheet for chromium in sludge fertilizers (Extraction procedure)

<table>
<thead>
<tr>
<th>Sample solution</th>
<th>10-mL volumetric flask, 3 flasks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot 5 mL</td>
<td>← 0, 2 and 4 mL of chromium standard solution (2.5 μg / mL) respectively</td>
</tr>
<tr>
<td></td>
<td>← Hydrochloric acid (1+23) (up to the marked line)</td>
</tr>
<tr>
<td>Measurement</td>
<td>ICP Optical Emission Spectrometer (205.552 nm)</td>
</tr>
</tbody>
</table>

Figure 2  Flow sheet for chromium in sludge fertilizers (Measurement procedure)
5.5.e ICP Mass Spectrometry (Fluid sludge fertilizers)

(1) **Summary**
The test method is applicable to fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.5.e-2017 or Cr.e-1.

Add nitric acid–hydrogen peroxide to an analytical sample to heat and extract by microwave irradiation. After adding an internal standard element, introduce it to an ICP Mass Spectrometer (ICP-MS) and measure the respective indicated values of chromium and an internal standard element with mass/charge number \(m/z\) and obtain chromium (Cr) in the analytical sample from the ratio of the indicated value for chromium and the indicated value for an internal standard element. In addition, the performance of this testing method is shown in Comment 4.

(2) **Reagent, etc.** Reagents and water are as shown below.

a) **Water**: Water of A4 specified in JIS K 0557.

b) **Nitric acid**: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) **Nitric acid**: Nitric acid used to dilute a standard solution and a sample solution is a highly pure reagent specified in JIS K 9901.

d) **Hydrogen peroxide**: A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.

e) **Rhodium standard solution (Rh 1 mg/mL)**: A rhodium standard solution (Rh 1 mg/mL) traceable to National Metrology.

f) **Rhodium standard solution (Rh 0.1 µg/mL)** \((1)(2)(3)\): Dilute a predetermined amount of rhodium standard solution (Rh 1 mg/mL) with nitric acid (1+19) to prepare a rhodium standard solution (Rh 0.1 µg/mL).

h) **Rhenium standard solution (Re 0.1 µg/mL)** \((1)(2)(3)\): Dilute a predetermined amount of rhenium standard solution (Re 1 mg/mL) with nitric acid (1+19) to prepare a rhenium standard solution (Re 0.1 µg/mL).

i) **Chromium standard solution (Cr 0.1 mg/mL)**: A chromium standard solution (Cr 0.1 mg/mL) traceable to National Metrology.

j) **Chromium standard solution (Cr 5 µg/mL)** \((1)(2)(3)\): Transfer 5 mL of chromium standard solution (Cr 0.1 mg/mL) to a 100-mL volumetric flask and add nitric acid (1+19) to the marked line.

k) **Chromium standard solutions (Cr 1 ng/mL - 100 ng/mL) for the calibration curve preparation** \((1)(2)(3)\): Transfer 0.02 mL - 2 mL of chromium standard solution (Cr 5 µg/mL) to 100-mL volumetric flasks step-by-step, and add 10 mL of rhodium standard solutions (Rh 0.1 µg/mL) as an internal standard respectively \((5)\) and add nitric acid (1+19) up to the marked line.

l) **Blank test solution for the calibration curve preparation** \((1)(2)(3)\): transfer 10 mL of rhodium standard solution (Rh 0.1 µg/mL) to 100-mL volumetric flasks as internal standard \((5)\) and add nitric acid (1+19) up to the marked line.

**Note**

(1) This is an example of preparation; prepare an amount as appropriate.

(2) Store in cool and dark place, and do not use after 1 month after preparation.

(3) For storage, use a sealable container made of materials such as polypropylene containing no chromium.

(4) Use when measuring lead simultaneously.

(5) Add an internal solution that is 1/10 of the volume to be prepared.

**Comment 1** Instead of the chromium standard solution in (2), a chromium standard solution for
the calibration curve preparation can be prepared by using a chromium standard solution (Cr 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) **Instruments**: Instruments are as shown below:

a) **ICP Mass Spectrometer**: High-frequency plasma mass spectrometer specified in JIS K 0133.

1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.995 % in purity

b) **Pressure vessel decomposing device**: A device which pressurizes the inside of a vessel by putting acid, etc.to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.

1) **The main part of a decomposing device**: In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.

2) **Exhaust system**: A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.

3) **Airtight vessel**: A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.

c) **Centrifugal separator**: A centrifugal separator that can work at about 1700 × g.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 20.0 g \(^{(6)}\) of an analytical sample, and put it into an airtight vessel.

b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.

c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave \(^{(7)}\).

d) Ignite at 240 °C ± 5 °C for no less than 10 minutes \(^{(7)}\) to decompose \(^{(8)}\).

e) After standing to cool, transfer the solution to a 50-mL volumetric flask \(^{(9)}\) with water.

f) Add water up to the marked line and transfer it to a 50-mL ground-in stopper centrifugal precipitate tube \(^{(9)}\).

g) Centrifuge it at 1700 × g centrifugal force for about five minutes \(^{(10)}\) and use the supernatant as the sample solution.

h) As a blank test, conduct the procedures in b) - g) using another airtight vessel to prepare the blank test solution.

**Note** (6) The maximum limit of solid content, which is converted from moisture content, in the sampling volume 20.0 g of an analytical sample is about 0.5 g. If solid content is likely to exceed the limit, reduce sampling volume as necessary.

(7) Condition examples for a microwave decomposing device: 0 min (room temperature) → 10min (240 °C) → 20 min (240 °C) → 40 min (room temperature), initial output 1400 W

(8) When organic matters still remain, for example the digestion solution is colored, repeat the procedures in (4.1) b) - d).

(9) The vessel should be made of polypropylene, etc. to not affect the measurement.

(10) 16.5-cm of radius and 3000 rpm of revolutions makes about 1700 × g centrifugal
Comment 2  The procedure in (4.1) is the same as the procedure in (4.1) of 5.1.b.

(4.2) Measurement: Conduct measurement (Internal Standard Method) according to JIS K 0113 and as shown below. Specific measurement procedures are according to the operation method of the ICP Mass Spectrometer used in measurement.

a) Measurement conditions for the ICP Mass Spectrometer: Set up the measurement conditions for the ICP Mass Spectrometer considering the following:
Chromium: monitor ion (m/z): 52, 53, 50
Rhodium: monitor ion (m/z): 103

b) Calibration curve preparation
1) Spray chromium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into the inductively coupled plasma, and read the ratio of the respective ion count numbers for the monitor ion of an element subjected to measurement and an internal standard element.
2) Prepare a curve for the relationship between the concentration and the ratio of the ion count number of an element subjected to measurement.

c) Sample measurement
1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 µg - 5 µg as Cr) to a 50-mL volumetric flask.
2) Add 5mL of internal standard solution and add nitric acid (1+19) to the marked line.
3) Conduct procedures similarly as in b) 1) to read the ratio of the ion count number.
4) For blank test solution, conduct procedures similarly as in 1) - 3) to correct the ratio of the ion count number obtained for the sample solution.
5) Obtain the chromium content from the calibration curve, and calculate chromium (Cr) in the analytical sample.

Comment 3 Instead of the correction method in c) 4), the chromium (Cr) in the analytical sample can also be corrected by obtaining the chromium content in the blank test solution.

Comment 4 Triplicates additive recovery testing was conducted to evaluate trueness using fluid industrial sludge fertilizers (2 samples) and composted sludge fertilizers (6 samples). As a result, the mean recovery at the concentration level of 10 mg/kg - 90 mg/kg, 1 mg/kg - 9 mg/kg and 0.1 mg/kg - 0.9 mg/kg are 92.4 % - 108.8 %, 94.3 % - 115.4 % and 105.8 - 106.8 % as chromium (Cr) in an actual article respectively. The results of the repeatability tests on different days using two kinds of fluid sludge fertilizers to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 1 µg/kg.
Comment 5 Simultaneous measurement of multiple elements by an ICP-MS is available. In that case, see 5.2.c Comment 5.

Comment 6 The presence and the extent of interference to the measurement mass number of an element subjected to measurement and an internal standard element can be estimated by qualitative analysis using an ICP-MS in advance to quantitation. Select a measurement mass number considering the extent of interference. However, a mass number cannot be changed in the measurement of arsenic. A magnetic sector double-focusing mass spectrometer or a collision reaction cell specified in JIS K0133 can be used as a method to reduce spectrum interference.

References
(5) **Flow sheet for chromium:** The flow sheet for chromium in fluid sludge fertilizers is shown below:

![Flow sheet for chromium in fluid sludge fertilizers](image)

**Figure 1** Flow sheet for chromium in liquid sludge fertilizers  
(Extraction procedure)

![Flow sheet for chromium in liquid sludge fertilizers](image)

**Figure 2** Flow sheet for chromium in liquid sludge fertilizers  
(Measurement procedure)
5.6 Lead
5.6.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.6.a-2017 or Pb.a-1.

Pretreat an analytical sample with incineration and nitric acid–hydrochloric acid (1+3), spray into an acetylene–air flame, and measure the atomic absorption with lead at a wavelength of 217.0 nm or 283.3 nm to obtain the lead (Pb) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Lead standard solution (Pb 0.1 mg/mL): A lead standard solution (Pb 0.1 mg/mL) traceable to National Metrology.
   e) Lead standard solutions (Pb 0.5 µg - 5 µg/mL) for the calibration curve preparation (1): Transfer 2.5 mL - 25 mL of lead standard solution (Pb 0.1 mg/mL) to 500-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   f) Blank test solution for the calibration curve preparation (1)(2): Hydrochloric acid (1+23) used in the procedures in e).

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the lead standard solution in (2), a lead standard solution for the calibration curve preparation can be prepared by using a lead standard solution (Pb 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121 with the background correction (3) function.
      1) Light source: A lead hollow cathode lamp (In case of background correction system using continuous spectrum source, the light source is a deuterium lamp.)
      2) Gas: Gas for heating by flame
         (i) Fuel gas: acetylene
         (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can keep the test temperature at 450 °C ± 5 °C.
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.

Note (3) There are the continuous source method, the Zeeman method, the non-resonance spectrum method, and the self-reversal method, etc.

(4) Test procedure
(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.

b) Put the tall beaker in an electric furnace, and heat gently to char (4).

c) Ignite at 450 °C ± 5 °C for 8 - 16 hours to incinerate (4).

d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.

f) Slightly move the watch glass (5), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.

g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (6) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.

h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

**Note** (4) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250°C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

(5) The watch glass can be removed.

(6) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

**Comment 2** Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

**Comment 3** The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.

a) **Measurement conditions for the atomic absorption spectrometer**: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
- Analytical line wavelength: 217.0 nm or 283.3 nm

b) **Calibration curve preparation**

1) Spray the lead standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 217.0 nm or 283.3 nm.

2) Prepare a curve for the relationship between the lead concentration and the indicated value of the lead standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) **Sample measurement**

1) Subject the sample solution (7) to the same procedure as in b) 1) to read the indicated value.

2) Subject the blank test solution to the same procedure as in b) 1) to read the indicated value, and correct the indicated value obtained for the sample solution.

3) Obtain the lead content from the calibration curve, and calculate lead (Pb) in the analytical sample.
**Note** (7) If there is a possibility that the lead concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount with hydrochloric acid (1+23).

**Comment 4** Instead of the correction method in c) 2), the lead (Pb) in the analytical sample can also be corrected by obtaining the lead content in the blank test solution.

**Comment 5** Recovery testing was conducted using industrial sludge fertilizer and composted sludge fertilizer (5 samples); as a result, the recovery at the concentration level of 100 mg/kg and 10 mg/kg were 99.1 % - 100.6 % and 97.5 % - 99.6 %, respectively. Table 1 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of this test method is about 1 mg/kg.

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (mg/kg)</th>
<th>RSD$_r$ (%)</th>
<th>RSD$_R$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sewage sludge fertilizer a</td>
<td>10</td>
<td>25.2</td>
<td>4.6</td>
<td>3.9</td>
</tr>
<tr>
<td>Sewage sludge fertilizer b</td>
<td>11</td>
<td>29.4</td>
<td>3.7</td>
<td>4.3</td>
</tr>
<tr>
<td>Composted sludge fertilizer a</td>
<td>10</td>
<td>18.6</td>
<td>3.2</td>
<td>5.0</td>
</tr>
<tr>
<td>Composted sludge fertilizer b</td>
<td>10</td>
<td>22.2</td>
<td>1.8</td>
<td>7.0</td>
</tr>
<tr>
<td>Composted sludge fertilizer c</td>
<td>11</td>
<td>86.8</td>
<td>1.3</td>
<td>4.0</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n =$ number of laboratories x number of samples (2))
3) Repeatability relative standard deviation
4) Reproducibility relative standard deviation

**References**


(5) **Flow sheet for lead:** The flow sheet for lead in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Action</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>5.00 g analytical sample</td>
<td>200-mL - 300-mL tall beaker.</td>
</tr>
<tr>
<td>2.</td>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>3.</td>
<td>Incineration</td>
<td>Ignite at 450 °C ± 5 °C, 8 - 16 hours</td>
</tr>
<tr>
<td>4.</td>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>5.</td>
<td>A small amount of water, moisten the residue</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>About 10 mL of nitric acid</td>
<td></td>
</tr>
<tr>
<td>7.</td>
<td>About 30 mL of hydrochloric acid</td>
<td></td>
</tr>
<tr>
<td>8.</td>
<td>Cover with a watch glass, and digest</td>
<td></td>
</tr>
<tr>
<td>9.</td>
<td>Heating</td>
<td>Slightly move a watch glass to remove acid</td>
</tr>
<tr>
<td>10.</td>
<td>Heating</td>
<td></td>
</tr>
<tr>
<td>11.</td>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>12.</td>
<td>About 25 mL - 50 mL of hydrochloric acid (1+5)</td>
<td></td>
</tr>
<tr>
<td>13.</td>
<td>Cover with a watch glass, and dissolve</td>
<td></td>
</tr>
<tr>
<td>14.</td>
<td>Heating</td>
<td></td>
</tr>
<tr>
<td>15.</td>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>16.</td>
<td>Transfer</td>
<td>100-mL - 200-mL volumetric flask, water</td>
</tr>
<tr>
<td>17.</td>
<td>Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>18.</td>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>19.</td>
<td>Sample solution</td>
<td></td>
</tr>
<tr>
<td>20.</td>
<td>Measurement</td>
<td>Atomic absorption spectrometer (217.0 nm or 283.3 nm)</td>
</tr>
</tbody>
</table>

Figure Flow sheet for lead in fertilizers
5.6.b ICP Optical Emission Spectrometry

(1) Summary
The test method is applicable to sludge fertilizers, etc. This testing method is classified as Type D and its symbol is 5.6.b-2017 or Pb.b-1.

Pretreat an analytical sample with incineration, nitric acid–hydrochloric acid (1+3), introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the emission with lead at a wavelength of 220.351 nm to obtain the lead (Pb) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Hydrochloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Lead standard solution (Pb 0.1 mg/mL): A lead standard solution (Pb 0.1 mg/mL) traceable to National Metrology.
   e) Lead standard solutions (Pb 2.5 µg/mL): Dilute a predetermined amount of lead standard solution (Pb 0.1 mg/mL) with hydrochloric acid (1+23) to prepare a lead standard solution (Pb 2.5 µg/mL).

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) Store at room temperature, and do not use after 6 months after preparation.

Comment 1 Instead of the lead standard solution in (2), a lead standard solution for the calibration curve preparation can be prepared by using a lead standard solution (Pb 1 mg/mL or 10 mg/mL) traceable to National Metrology.

(3) Instruments: Instruments are as shown below:
       1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity
       b) Electric furnace: An electric furnace that can keep the test temperature at 450 ºC ± 5 ºC.
       c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 ºC. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 ºC.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Put the tall beaker in an electric furnace, and heat gently to char (3)
   c) Ignite at 450 ºC ± 5 ºC for 8 - 16 hours to incinerate (3).
   d) After standing to cool, moisten the residue with a small amount of water, and add about 10 mL of nitric acid and about 30 mL of hydrochloric acid.
   e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to digest.
   f) Slightly move the watch glass (4), and continue heating on the hot plate or sand bath to concentrate until nearly dried up.
   g) After standing to cool, add 25 mL - 50 mL of hydrochloric acid (1+5) (5) to the digest, cover the tall beaker with the watch glass, and heat quietly to dissolve.
   h) After standing to cool, transfer to a 100-mL - 200 mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.
As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note (3) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 450 °C in 1 to 2 hours.

(4) The watch glass can be removed.

(5) Add hydrochloric acid (1+5) so that the hydrochloric acid concentration of the sample solution will be hydrochloric acid (1+23). For example, when a 100-mL volumetric flask is used in the procedure in h), about 25 mL of hydrochloric acid (1+5) should be added.

Comment 2 Do not conduct the procedures in (4.1) b) - c) in the case of fertilizers not containing organic matters.

Comment 3 The procedure in (4.1) is the same as the procedure in (4.1) of 4.9.1.a.

(4.2) Measurement: Conduct measurement (Standard Addition Method) according to JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Optical Emission Spectrometer: Set up the measurement conditions for the ICP Optical Emission Spectrometer considering the following:

Analytical line wavelength: 220.351 nm

b) Calibration curve preparation and sample measurement

1) Put 5mL of sample solution to three 10-mL volumetric flasks respectively.
2) Add 2 mL and 4 mL of lead standard (0.25 μg/mL) solution to volumetric flasks of 1) above, then add hydrochloric acid (1+23) to the marked line to make a sample solution of Standard Addition Method.
3) Add hydrochloric acid (1+23) to the marked line of the remaining volumetric flask of 1) above to make a sample solution without a standard solution.
4) Spray the sample solution of Standard Addition Method and the sample solution without a standard solution into the induction plasma, and read the indicated value at a wavelength of 220.351 nm.
5) Transfer 5 mL of blank test solution to a 10-mL volumetric flask, conduct the same procedures as in 3) - 4) to read the indicated value, and correct the indicated value obtained from the respective sample solutions.
6) Prepare a curve for the relationship between the added lead concentration and the corrected indicated value of the sample solution for Standard Addition Method and the sample solution without a standard solution.
7) Obtain the lead content from the intercept of the calibration curve, and calculate lead (Pb) in the analytical sample.

Comment 4 Instead of the correction method in c) 5), the lead (Pb) in the analytical sample can also be corrected by obtaining the lead content in the blank test solution.

Comment 5 The simultaneous measurement of multiple elements by an ICP-OES is available. In that case, see 4.9.1.b Comment 5.

Comment 6 The comparison of the measurement value \((x_i: 1.1 \text{ mg/kg} - 69.0 \text{ mg/kg})\) of ICP Optical Emission Spectrometry and the measurement value \((y_i)\) of Flame Atomic Absorption Spectrometry was conducted to evaluate trueness using sludge fertilizers (49 samples). As a result, a regression equation was \(y = -0.31 + 1.045x\) and its
correlation coefficient \((r)\) was 0.993. Triplicates measurement for each one sample of sewage sludge fertilizer, human waste sludge fertilizer, industrial sludge fertilizer, mixed sludge fertilizer, calcined sludge fertilizers and composted sludge fertilizer was conducted. As a result, a repeatability obtained was 0.9 % - 3.3 % as a relative standard deviation. Additionally, the minimum limit of quantification of this testing method is about 5 mg/kg.

References


(5) **Flow sheet for lead**: The flow sheet for lead in fertilizers is shown below:

<table>
<thead>
<tr>
<th>5.00 g analytical sample</th>
<th>200-mL - 300-mL tall beaker.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Charring</td>
<td>Heat gently</td>
</tr>
<tr>
<td>Incineration</td>
<td>Ignite at 450 °C ± 5 °C, 8 - 16 hours</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← A small amount of water</td>
<td></td>
</tr>
<tr>
<td>← About 10 mL of nitric acid</td>
<td></td>
</tr>
<tr>
<td>← About 30 mL of hydrochloric acid</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and digest</td>
</tr>
<tr>
<td>Heating</td>
<td>Slightly move a watch glass to remove acid</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>← 25 mL - 50 mL of hydrochloric acid (1+5)</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Cover with a watch glass, and dissolve</td>
</tr>
<tr>
<td>Standing to cool</td>
<td>Room temperature</td>
</tr>
<tr>
<td>Transfer</td>
<td>100-mL - 200-mL volumetric flask, water</td>
</tr>
<tr>
<td>← Water (up to the marked line)</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1  Flow sheet for lead in sludge fertilizers (Extraction procedure)
Sample solution

Aliquot 5 mL

10-mL volumetric flask, 3 flasks

0 mL, 2 mL and 4 mL of lead standard solution (2.5 μg/mL) respectively

Hydrochloric acid (1+23) (up to the marked line)

Measurement

ICP Optical Emission Spectrometer (220.351 nm)

Figure 2  Flow sheet for lead in sludge fertilizers (Measurement procedure)
5.6.c ICP Mass Spectrometry (Fluid sludge fertilizers)

(1) **Summary**
The test method is applicable to fluid sludge fertilizers. This testing method is classified as Type D and its symbol is 5.6.c-2017 or Pb.c-1.

Add nitric acid–hydrogen peroxide to an analytical sample to heat and extract by microwave irradiation. After adding an internal standard element, introduce it to an ICP Mass Spectrometer (ICP-MS) and measure the respective indicated values of lead and an internal standard element with mass/charge number (m/z) and obtain lead (Pb) in the analytical sample from the ratio of the indicated value for lead and the indicated value for an internal standard element. In addition, the performance of this testing method is shown in Comment 4.

(2) **Reagent, etc.:** Reagents and water are as shown below.

a) **Water:** Water of A4 specified in JIS K 0557.

b) **Nitric acid:** A reagent of harmful metal analysis grade, microanalysis grade or equivalents.

c) **Nitric acid:** Nitric acid used to dilute a standard solution and a sample solution is a highly pure reagent specified in JIS K 9901.

d) **Hydrogen peroxide:** A JIS Guaranteed Reagent specified in JIS K 8230 or a reagent of equivalent quality.

e) **Rhenium standard solution (Re 1 mg/mL):** A rhenium standard solution (Re 1 mg/mL) traceable to National Metrology.

f) **Rhenium standard solution (Re 0.1 µg/mL)** (1)(2)(3): Dilute a predetermined amount of rhenium standard solution (Re 1 mg/mL) with nitric acid (1+19) to prepare a rhenium standard solution (Re 0.1 µg/mL).

g) **Rhodium standard solution (Rh 1 mg/mL)** (4): A rhodium standard solution (Rh 1 mg/mL) traceable to National Metrology.

h) **Rhodium standard solution (Rh 0.1 µg/mL)** (1)(2)(3): Dilute a predetermined amount of rhodium standard solution (Rh 1 mg/mL) with nitric acid (1+19) to prepare a rhodium standard solution (Rh 0.1 µg/mL).

i) **Lead standard solution (Pb 0.1 mg/mL):** A lead standard solution (Pb 0.1 mg/mL) traceable to National Metrology.

j) **Lead standard solution (Pb 5 µg/mL)** (1)(2)(3): Transfer 5 mL of lead standard solution (Pb 0.1 mg/mL) to a 100-mL volumetric flask and add nitric acid (1+19) to the marked line.

k) **Lead standard solutions (Pb 1 ng/ mL - 100 ng/ mL) for the calibration curve preparation** (1)(2)(3): Transfer 0.02 mL - 2 mL of lead standard solution (Pb 5 µg/mL) to 100-mL volumetric flasks step-by-step, and add 10 mL of rhenium standard solutions (Re 0.1 µg/mL) as an internal standard respectively (5) and add nitric acid (1+19) up to the marked line.

l) **Blank test solution for the calibration curve preparation** (1)(2)(3): transfer 10 mL of rhenium standard solution (Re 0.1 µg/mL) to 10-mL volumetric flasks as internal standard respectively (5) and add nitric acid (1+19) up to the marked line.

**Note**

(1) This is an example of preparation; prepare an amount as appropriate.

(2) Store in cool and dark place, and do not use after 1 month after preparation.

(3) For storage, use a sealable container made of materials such as polypropylene containing no lead.

(4) Use when measuring arsenic, cadmium, nickel or chromium simultaneously.

(5) Add an internal solution that is 1/10 of the volume to be prepared.

**Comment 1** Instead of the lead standard solution in (2), a lead standard solution for the calibration curve preparation can be prepared by using a lead standard solution (Pb 1 mg/mL or
10 mg/mL) traceable to National Metrology.

(3) **Instruments**: Instruments are as shown below:

a) **ICP Mass Spectrometer**: High-frequency plasma mass spectrometer specified in JIS K 0133.

1) **Gas**: Argon gas specified in JIS K 1105 of no less than 99.995 % in purity

b) **Pressure vessel decomposing device**: A device which pressurizes the inside of a vessel by putting acid, etc. to heat in the airtight vessel, and decomposes a sample by the interaction of heating, pressurizing and acid. The following requirements should be met.

1) **The main part of a decomposing device**: In the case of the microwave heating method, a device should be able to produce a high-frequency wave using a frequency which is permitted at an industrial frequency facility. It is desirable to be able to monitor pressure and temperature, etc. in the airtight vessel with a sensor inside the device. The interior of a device should be acid-resistant treated and should have a high temperature durability and a high level of safety.

2) **Exhaust system**: A system which has an exhaust fan of acid-resistant specification, and has an air-cool function inside the device to allow constant airflow, keeping operating temperature below a certain temperature level.

3) **Airtight vessel**: A vessel which is heat-resistance and pressure-tight and has the durability required to decompose particles, and has resistance to internal contamination. It should have safety functions such as causing overheat prevention valve to work and internal pressure to drop by emitting gas, and preventing gas bumping in the case of exceeding a pressure limitation.

c) **Centrifugal separator**: A centrifugal separator that can work at about $1700 \times g$.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 20.0 g of an analytical sample, and put it into an airtight vessel.

b) Gradually add 2.5 mL of nitric acid and 2 mL of hydrogen peroxide.

c) Put the airtight vessel into the main part of a decomposing device and heat using a microwave.

d) Ignite at $240 ^\circ C \pm 5 ^\circ C$ for no less than 10 minutes to decompose.

e) After standing to cool, transfer the solution to a 50-mL volumetric flask with water.

f) Add water up to the marked line and transfer it to a 50-mL ground-in stopper centrifugal precipitate tube.

g) Centrifuge it at $1700 \times g$ centrifugal force for about five minutes and use the supernatant as the sample solution.

h) As a blank test, conduct the procedures in b) - g) using another airtight vessel to prepare the blank test solution.

**Note** (6) The maximum limit of solid content, which is converted from moisture content, in the sampling volume 20.0 g of an analytical sample is about 0.5 g. If solid content is likely to exceed the limit, reduce sampling volume as necessary.

(7) Condition examples for a microwave decomposing device: 0 min (room temperature) → 10min ($240 ^\circ C$) → 20 min ($240 ^\circ C$) → 40 min (room temperature), initial output 1400 W

(8) When organic matters still remain, for example the digestion solution is colored, repeat the procedures in (4.1) b) - d).

(9) The vessel should be made of polypropylene, etc. to not affect the measurement.

(10) 16.5-cm of radius and 3000 rpm of revolutions makes about $1700 \times g$ centrifugal force.
Comment 2  The procedure in (4.1) is the same as the procedure in (4.1) of 5.1.b.

(4.2) Measurement: Conduct measurement (Internal Standard Method) according to JIS K 0113 and as shown below. Specific measurement procedures are according to the operation method of the ICP Mass Spectrometer used in measurement.

a) Measurement conditions for the ICP Mass Spectrometer: Set up the measurement conditions for the ICP Mass Spectrometer considering the following:
   Lead: monitor ion \((m/z)\): 208, 206, 207
   Rhenium: monitor ion \((m/z)\): 187

b) Calibration curve preparation
   1) Spray lead standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into the inductively coupled plasma, and read the ratio of the respective ion count numbers for the monitor ion of an element subjected to measurement and an internal standard element.
   2) Prepare a curve for the relationship between the concentration and the ratio of the ion count number of an element subjected to measurement.

c) Sample measurement
   1) Transfer a predetermined amount of the sample solution (the equivalents of 0.05 µg - 5 µg as Pb) to a 50-mL volumetric flask \(^{(9)}\).
   2) Add 5mL of internal standard solution \(^{(5)}\) and add nitric acid (1+19) to the marked line.
   3) Conduct procedures similarly as in b) 1) to read the ratio of the ion count number.
   4) For blank test solution, conduct procedures similarly as in 1) - 3) to correct the ratio of the ion count number obtained for the sample solution.
   5) Obtain the lead content from the calibration curve, and calculate lead (Pb) in the analytical sample.

Comment 3  Instead of the correction method in c) 4), the lead (Pb) in the analytical sample can also be corrected by obtaining the lead content in the blank test solution.

Comment 4  Triplicates additive recovery testing was conducted to evaluate trueness using fluid industrial sludge fertilizers (2 samples) and composted sludge fertilizers (6 samples). As a result, the mean recovery at the concentration level of 10 mg/kg - 20 mg/kg, 1 mg/kg - 5 mg/kg and 0.1 mg /kg - 0.9 mg /kg are 91.2 % - 103.1 %, 85.0 % - 113.9 % and 93.2 - 108.1 % as lead (Pb) in an actual article respectively.

The results of the repeatability tests on different days using two kinds of fluid sludge fertilizers to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this test method is about 4 µg /kg.
Comment 5 Simultaneous measurement of multiple elements by an ICP-MS is available. In that case, see 5.2.c Comment 5.

Comment 6 The presence and the extent of interference to the measurement mass number of an element subjected to measurement and an internal standard element can be estimated by qualitative analysis using an ICP-MS in advance to quantitation. Select a measurement mass number considering the extent of interference. However, a mass number cannot be changed in the measurement of arsenic. A magnetic sector double-focusing mass spectrometer or a collision reaction cell specified in JIS K0133 can be used as a method to reduce spectrum interference.

References

(5) Flow sheet for lead: The flow sheet for lead in fluid sludge fertilizers is shown below:

```
<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Average (s_T) (s_{\text{RSD}})</td>
<td>(s_{\text{RSD}})</td>
</tr>
<tr>
<td>Composted sludge fertilizer 1</td>
<td>5</td>
<td>2.80</td>
<td>0.09</td>
</tr>
<tr>
<td>Composted sludge fertilizer 2</td>
<td>5</td>
<td>0.740</td>
<td>0.014</td>
</tr>
</tbody>
</table>
```

1)  The number of test days conducting a duplicate test
2)  Average (the number of test days \(T\) \times\) the number of duplicate testing (2)
3)  Repeatability standard deviation
4)  Repeatability relative standard deviation
5)  Intermediate standard deviation
6)  Intermediate relative standard deviation

Table 1 Analysis results of the repeatability tests on different days

Figure 1  Flow sheet for lead in liquid sludge fertilizers (Extraction procedure)
Figure 2  Flow sheet for lead in liquid sludge fertilizers (Measurement procedure)
5.7 Sulfamic acid (amidosulfuric acid)

5.7.a Ion Chromatography

(1) Summary
The testing method is applicable to ammonium sulfate. This testing method is classified as Type D and its symbol is 5.7.a-2017 or AS-acid.a-1.

Add water to an analytical sample to extract sulfamic acid, introduce it to an Ion Chromatograph (IC) or a High-Performance Liquid Chromatograph (HPLC) to isolate it with an ion exchange column, then measure the sulfamic acid with an electric conductivity detector to obtain sulfamic acid (amidosulfuric acid) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

Sulfamic acid and sulfurized cyanide (ammonium thiocyanate) can be simultaneously quantified by using this method. (Refer to Comment 4).

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Phthalic acid: A reagent of no less than 98 % (mass fraction) in purity.
   c) p-hydroxybenzoic acid: A reagent of no less than 95 % (mass fraction) in purity.
   d) 1-sodium octane sulfonate: A reagent of no less than 98 % (mass fraction) in purity.
   e) 1-sodium hexane sulfonate: A reagent of no less than 98 % (mass fraction) in purity.
   f) Boric acid: A JIS Guaranteed Reagent specified in JIS K 8863 or a reagent of equivalent quality.
   g) Eluent: Weigh 0.083 g of phthalic acid, 0.552 g of p-hydroxybenzoic acid, 0.195 g of 1-sodium octane sulfonate, 0.376 g of 1-sodium hexane sulfonate and 6.183 g of boric acid to a 1000-mL volumetric flask, add about 500 mL of water to dissolve and add water up to the marked line. Filter with a membrane filter (aperture diameter: no more than 0.5 µm) made of hydrophilic PTFE.
   h) Sulfamic acid standard solution (1000 mg/L): Put 0.1 g of sulfamic acid, reference material for volumetric analysis (HOSO₂NH₂: dried for 48 hours in a silica gel desiccator), in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
   i) Sulfamic acid standard solution (10 mg/L): At the time of usage, put 2.5 mL of sulfamic acid standard solution (1000 mg/L) to a 250-mL volumetric flask and add water up to the marked line.
   j) Sulfamic acid standard solution for the calibration curve preparation (0.3 mg/L - 3 mg/L): At the time of usage, put 3 mL - 30 mL of sulfamic acid standard solution (10 mg/L) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) The concentration of prepared solutions is phthalic acid 0.5 mmol/L, p-hydroxybenzoic acid 4.0 mmol/L, 1-sodium octane sulfonate 0.9 mmol/L, 1-sodium hexane sulfonate 2.0 mmol/L and boric acid 100 mmol/L.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Ion Chromatograph (IC) or High-Performance Liquid Chromatograph (HPLC): IC specified in JIS K 0127 or HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm inner diameter 100-mm long stainless-steel column tube filled with hydrophilic methacrylate-gel, to which 5-µm particle diameter class 4 ammonium group chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 55 ºC - 60 ºC.
3) **Detection unit:** An electric conductivity detector

b) **Membrane filters:** Pore size is no more than 0.5 μm, made of hydrophilic PTFE

**Note** (3) A column is commercially sold under the name Shodex IC NI-424, etc.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.

b) Add about 50 mL of water and shake to dissolve, and then add water up to the marked line.

c) Transfer a predetermined amount of the solution, and dilute exactly by a factor of 12.5 with water.

d) Filter with a membrane filter (pore size: no more than 0.5 μm) to make a sample solution.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0127 and as shown below. Specific measurement procedures are according to the operation method of the Ion Chromatograph (IC) or the High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for the Ion Chromatograph (IC) or High-Performance Liquid Chromatograph (HPLC):** An example of measurement conditions is shown below. Set up the measurement conditions considering it:

1) **Column:** A hydrophilic methacrylate-gel column (4-mm inner diameter, 100-mm long, 5-μm particle diameter) to which quaternary ammonium group chemically bonds.

2) **Column bath temperature:** 58 °C

3) **Eluent:** Prepared by the procedures in (2) g

4) **Flow rate:** 1 mL/min

5) **Injection volume:** 20 μL

6) **Detection unit:** An electric conductivity detector

b) **Calibration curve preparation**

1) Inject 20 μL of respective standard solutions for the calibration curve preparation to an IC or an HPLC, and record the chromatogram of electric conductivity to obtain peak area.

2) Prepare a curve for the relationship between the concentration and the peak area of electric conductivity of respective standard solutions for the calibration curve preparation. Prepare a calibration curve when the sample is measured.

**Comment 1** In the measurement of a sample solution, there is a possibility that the recovery rate becomes lower than actual due to the influence of matrix if the concentration is calculated with peak height. Therefore, prepare a calibration curve using peak area.

c) **Sample measurement**

1) Subject 20 μL of sample solution to the same procedure as in b) 1)

2) Obtain the sulfamic acid content from calibration curve by peak area to calculate the sulfamic acid (amidosulfuric acid) in the analytical sample.

**Comment 2** Calculate the concentration by the peak area similarly as the calibration curve preparation to prevent the influence of matrix.

**Comment 3** Note that it takes time to stabilize the baseline due to the usage of the ion-pairing reagent in the elute. It is recommended to take about 120 minutes for stabilization time before starting measurement.

**Comment 4** It is possible for the simultaneous measurement of sulfamic acid and ammonium
thiocyanate in this testing method. In that case, mix a predetermined amount of sulfamic acid standard solution (1000 mg/L) and ammonium thiocyanate standard solution (1000 mg/L), dilute with water to prepare a mixture standard solution (10 mg/L) \(^{(1)}\) and use it instead of respective standard solutions (10 mg/L) in (2) \(i\). After that, conduct the same procedure in (4.2) \(b\) to calculate the respective concentrations of materials subjected to measurement in the analytical sample.

**Comment 5** A recovery testing of ammonium sulfate (3 brands) was conducted. As a result, the mean recovery at additive level of 0.25 % (mass fraction) and 0.075 % (mass fraction) was 94.4 % - 103.5 % and 94.4 % - 100.8 %. Additionally, the minimum limit of quantification of this testing method is about 0.04 % (mass fraction).

**References**

(5) **Flow sheet for testing method:** The flow sheet for sulfamic acid in ammonium sulfate is shown below:

```
1.00 g  Weigh into a 100-mL volumetric flask
 ← About 50 mL of water
Shaking to mix  Stopple the volumetric flask and dissolve
 ← Water (up to the marked line)
Dilution  Dilute 12.5 times, water
Filtration  Membrane filter (no more than 0.5-µm)
Sample solution
Measurement  Ion Chromatograph
```

Figure Flow sheet for sulfamic acid in ammonium sulfate
Reference: The IC chromatogram of sulfamic acid and thiocyanic acid of the standard solution for the calibration curve preparation and sample solution (ammonium sulfate) are shown below.

(A) Mixture standard solution (the equivalents of 60 ng as sulfamic acid and ammonium thiocyanate (3 mg/L, 20 μL), respectively)

(B) Sample solution (the equivalents of 0.25 % (mass fraction) as sulfamic acid and ammonium thiocyanate added in ammonium sulfate (2500 μg/g), respectively)

Reference diagram: IC chromatogram of sulfamic acid and thiocyanic acid.
(Peak: 1. Sulfamic acid, 2. Ammonium thiocyanate)

IC measurement conditions
Column: Shodex IC NI-424 (4.6-mm inner diameter, 100-mm long, 5μm-particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
5.7.b High-Performance Liquid Chromatograph Mass Spectrometry

(1) **Summary**
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.7.b-2017 or AS-acid.b-1.

Add water to an analytical sample to extract sulfamic acid, introduce it to a High-Performance Liquid Chromatograph Mass Spectrometer to isolate it with a silica gel column, to which crosslink-type diol chemically bonds, and measure with a Selected Ion Monitoring (SIM) method to obtain sulfamic acid (amidosulfuric acid) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2) **Reagent, etc.:** Reagents and water are as shown below.
   a) **Water:** Water of A3 specified in JIS K 0557. Note that water of A4 or equivalent quality should be used as the eluent which is introduced to LC-MS.
   b) **Acetonitrile:** A reagent of LC-MS analysis grade or equivalents.
   c) **Formic acid:** A reagent of LC-MS analysis grade or equivalents.
   d) **Ammonium formate buffer solution (pH 3.2):** Dilute 3.153 g of ammonium formate (no less than 95 % (mass fraction) in purity) with water to make 500 mL and adjust to pH 3.2 with formic acid.
   e) **Sulfamic acid standard solution (1 mg/L):** Put 0.1 g of sulfamic acid, reference material for volumetric analysis specified in JIS K 8005, in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
   f) **Sulfamic acid standard solution (10 μg/L) (1):** At the time of usage, put 2.5 mL of standard solution (1 mg/L) to a 250-mL volumetric flask and add water up to the marked line.
   g) **Sulfamic acid standard solution (200 ng/L) (1):** At the time of usage, put 5 mL of standard solution (10 μg/L) to a 250-mL volumetric flask and add water up to the marked line.
   h) **Sulfamic acid standard solution for the calibration curve preparation (10 ng/L - 600 ng/L):** At the time of usage, put 2.5 mL - 6 mL of sulfamic acid standard solution (10 μg/L) to 100-mL volumetric flasks step-by-step and add water up to the marked line. Similarly put 5 mL - 50 mL of sulfamic acid standard solution (200 ng/L) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

**Note** (1) This is an example of preparation; prepare an amount as appropriate.

(3) **Apparatus and instruments:** Apparatus and instruments are shown below.
   a) **High-Performance Liquid Chromatograph/Mass Spectrometer (LC-MS):** LC-MS specified in JIS K 0136 that satisfies the following requirements.
      1) **High-Performance Liquid Chromatograph**
         (i) **Column bath:** A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.
         (ii) **Column:** A 2-mm - 3-mm inner diameter 100-mm - 150 mm long stainless-steel column tube filled with silica gel to which 5-μm crosslink-type diol chemically bonds or polyhydroxymethacrylate. This should comply with the specification of a Mass Spectrometer.
      2) **Mass Spectrometer**
         (i) **Ionization method:** Electro-Spray Ionization (ESI) method
         (ii) **Ion detection method:** Selected Ion Monitoring (SIM) method
   b) **Magnetic stirrer:**
   c) **Centrifugal separator:** A centrifugal separator that can work at about 1700 × g.
   d) **High speed centrifugal separator:** A centrifugal separator that can work at 8000 × g - 10000 × g.
Comment 1 A high Performance Liquid Chromatograph/Mass Spectrometer (LC-MS/MS) can be used instead of a High-Performance Liquid Chromatograph/Mass Spectrometer (LC-MS). In this case, set up the measurement conditions considering (4.3) a) The measurement conditions of High-Performance Liquid Chromatograph/Mass spectrometer and confirm that a calibration curve can be prepared by the procedure in b) in advance.

Comment 2 A column is sold under the name LUNA HILIC or Shodex ODP2 HP-2D.

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Powdery test sample
a) Weigh 1.00 g of an analytical sample, and put it into a 200-mL ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at 1700 × g centrifugal force for about five minutes (2) and use the supernatant as the extract.

Note (2) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

Comment 3 Instead of the procedures in (4.1.1) c) and d), it is allowed to filter with Type 3 filter paper and the filtrate can be the extract.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water to the marked line to make the extract

(4.2) Extraction: Conduct extraction as shown below.
a) Transfer 2 mL of the sample solution to a 200-mL Erlenmeyer flask.
b) Add water up to the marked line and transfer it to a 1.5-mL ground-in stopper centrifugal precipitate tube (3).
c) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (4) and use the supernatant as the extract.

Note (3) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(4) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

Comment 4 Instead of the procedures in (4.2) b) and c), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5 µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0136 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph Mass Spectrometer used in measurement.
a) The measurement conditions of High-Performance Liquid Chromatograph/Mass spectrometer: An example of measurement conditions is shown below. Set up the measurement conditions considering it:

1) High-Performance Liquid Chromatograph
   (i) Column: A silica gel column to which crosslink-type diol chemically bonds or polyhydroxymethacrylate (2-mm - 3-mm inner diameter, 100-mm - 150-mm long, 5-µm particle diameter).
   (ii) Flow rate: 0.2 mL/min
   (iii) Eluent: Ammonium formate buffer solution - Acetonitrile (1+9)
   (iv) Temperature of column bath: 40 °C
   (v) Injection rate: 1 µL
   (vi) Measurement time: About 20 minutes

2) Mass Spectrometer
   (i) Ionization method: Electro-Spray Ionization (ESI) method
   (ii) Mode: Negative
   (iii) Capillary voltage: −3.5 kv
   (iv) Ion source temperature: 300 °C
   (v) Nebulizer gas rate: 1.5 L/min
   (vi) Desolvation temperature: 250 °C
   (vii) Monitor ion: m/z 95.9

b) Calibration curve preparation
   1) Inject 1 µL of respective standard solutions for the calibration curve preparation to an LC-MS, and record the chromatogram of monitor ion (m/z) to obtain peak area.
   2) Prepare a curve for the relationship between the sulfamic acid concentration and the peak area of monitor ion of respective standard solutions for the calibration curve preparation.

c) Sample measurement
   1) Subject 1 µL of sample solution to the same procedure as in b) 1)
   2) Obtain the sulfamic acid content from calibration curve to calculate the sulfamic acid (amidosulfuric acid) in the analytical sample.

Comment 5 Recovery testing was conducted using samples that sulfamic acid equivalent to 1/5 - 4 times of permissible content are added to ammonium sulfate fertilizer (1 brand), by-product nitrogen fertilizer (1 brand), by-product mixed fertilizer (1 brand), compound fertilizer (1 brand) and fluid mixed fertilizer (1 brand). As a result, the mean recovery at the additive level of 0.1 % (mass fraction), 0.025 % (mass fraction) and 0.005 % (mass fraction) are 97.6 % - 104.2 %, 95.2 % - 107.0 % and 96.4 % - 111.2 % respectively.

The results of the repeatability tests on different days using ammonium sulfate fertilizer, by-product nitrogen fertilizer and by-product mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation. In addition, sufficient reproducibility by sulfamic acid concentration 0.0116 % (mass fraction) was not obtainable. But sufficient reproducibility was obtained in the scope of sulfamic acid concentration 0.0386 % (mass fraction) - 0.401 % (mass fraction).
Table 1: Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average&lt;sup&gt;2)&lt;/sup&gt; (T&lt;sup&gt;1)&lt;/sup&gt; (%)&lt;sup&gt;3)&lt;/sup&gt;</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4) s&lt;sub&gt;r&lt;/sub&gt; (%&lt;sup&gt;4)&lt;/sup&gt;</td>
<td>5) RSD&lt;sub&gt;r&lt;/sub&gt; (%&lt;sup&gt;5)&lt;/sup&gt;</td>
<td>6) s&lt;sub&gt;r(I)&lt;/sub&gt; (%&lt;sup&gt;6)&lt;/sup&gt;</td>
<td>7) RSD&lt;sub&gt;r(I)&lt;/sub&gt; (%&lt;sup&gt;7)&lt;/sup&gt;</td>
</tr>
<tr>
<td>Ammonium sulfate</td>
<td>5</td>
<td>0.0974</td>
<td>0.0011</td>
<td>1.1</td>
</tr>
<tr>
<td>By-product nitrogen fertilizer</td>
<td>5</td>
<td>0.0656</td>
<td>0.0014</td>
<td>2.1</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>5</td>
<td>0.005</td>
<td>0.000</td>
<td>2.4</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) × the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

Table 2: Results and analysis results from a collaborative study for sulfamic acid testing method validation

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories&lt;sup&gt;1)&lt;/sup&gt;</th>
<th>Mean&lt;sup&gt;2)&lt;/sup&gt; (%)&lt;sup&gt;3)&lt;/sup&gt;</th>
<th>4) s&lt;sub&gt;r&lt;/sub&gt; (%)&lt;sup&gt;4)&lt;/sup&gt;</th>
<th>5) RSD&lt;sub&gt;r&lt;/sub&gt; (%)&lt;sup&gt;5)&lt;/sup&gt;</th>
<th>6) s&lt;sub&gt;R&lt;/sub&gt; (%)&lt;sup&gt;6)&lt;/sup&gt;</th>
<th>7) RSD&lt;sub&gt;R&lt;/sub&gt; (%)&lt;sup&gt;7)&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ammonium sulfate</td>
<td>9</td>
<td>0.203</td>
<td>0.021</td>
<td>10.4</td>
<td>0.024</td>
<td>11.9</td>
</tr>
<tr>
<td>By-product nitrogen fertilizer</td>
<td>9</td>
<td>0.401</td>
<td>0.030</td>
<td>7.5</td>
<td>0.035</td>
<td>8.8</td>
</tr>
<tr>
<td>By-product mixed fertilizer</td>
<td>7</td>
<td>0.0957</td>
<td>0.0043</td>
<td>4.5</td>
<td>0.0043</td>
<td>4.5</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>9</td>
<td>0.0166</td>
<td>0.0028</td>
<td>16.8</td>
<td>0.0048</td>
<td>29.1</td>
</tr>
<tr>
<td>Fluid mixed fertilizer 1</td>
<td>9</td>
<td>0.0381</td>
<td>0.0022</td>
<td>5.8</td>
<td>0.0049</td>
<td>12.8</td>
</tr>
<tr>
<td>Fluid mixed fertilizer 2</td>
<td>9</td>
<td>0.243</td>
<td>0.011</td>
<td>4.5</td>
<td>0.018</td>
<td>7.6</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (n = number of laboratories × number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References


(5) **Flow sheet for testing method:** The flow sheet for sulfamic acid in fertilizers is shown below:

![Flow sheet for sulfamic acid in fertilizers](Extraction procedure (4.1.1))

- **1.00 g analytical sample (powdery)**
- **200-mL ground-in stopper Erlenmeyer flask**
- **← 100 mL of water**
- **Extraction**
- **Stir to mix, 10 minutes**
- **Centrifugal separation**
- **Ground-in stopper centrifugal precipitate tube, 1700 ×g, 5 minutes**
- **Extract**

Figure 1-1 Flow sheet for sulfamic acid in fertilizers
(Extraction procedure (4.1.1))

![Flow sheet for sulfamic acid in fertilizers](Extraction procedure (4.1.2))

- **1.00 g analytical sample (fluid)**
- **100-mL volumetric flask**
- **← About 50 mL of water**
- **Extraction**
- **Shake to mix**
- **← Water (up to the marked line)**
- **Extract**

Figure 1-2 Flow sheet for sulfamic acid in fertilizers
(Extraction procedure (4.1.2))

![Flow sheet for sulfamic acid in fertilizers](Dilution and measurement procedure)

- **Extract**
- **Aliquot (5 mL)**
- **100-mL volumetric flask**
- **← Water (up to the marked line)**
- **Centrifugal separation or filtration**
- **Ground-in stopper centrifugal precipitate tube, 8000 ×g - 10000 ×g, 5 minutes**
- **Sample solution**
- **Measurement**
- **Liquid Chromatograph Mass Spectrometer**

Figure 2 Flow sheet for sulfamic acid in fertilizers
(Dilution and measurement procedure)
Reference: The chromatogram of sulfamic acid for the calibration curve preparation is shown below.

(A) Standard solution (The equivalent of 0.6 ng as sulfamic acid)

(B) Sample solution (The equivalent of 0.1 % of mass fraction as sulfamic acid is added to a compound fertilizer solution)

Reference diagram: Chromatogram of sulfamic acid

LC-MS measurement conditions
Column: LUNA HILIC (2.0-mm inner diameter, 100-mm long, 5-μm particle diameter) 
Other conditions are according to the example of LC-MS measurement conditions in (4.3) a)
5.8 Ammonium thiocyanate (Sulfurized cyanide)
5.8.a Ion Chromatography

(1) Summary
The testing method is applicable to ammonium sulfate. This testing method is classified as Type D and its symbol is 5.8.a-2017 or SCN.a-1.

Add water to an analytical sample to extract ammonium thiocyanate (sulfurized cyanide), introduce it to an Ion Chromatograph (IC) or a High-Performance Liquid Chromatograph (HPLC) to isolate it with an ion exchange column, then measure the thiocyanic acid with an electric conductivity detector to obtain ammonium thiocyanate (sulfurized cyanide) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

Sulfamic acid and ammonium thiocyanate (sulfurized cyanide) can be simultaneously quantified by using this method. (Refer to Comment 4).

(2) Reagent, etc.: Reagents and water are as shown below.


b) Phthalic acid: A reagent of no less than 98 % (mass fraction) in purity.

c) p-hydroxybenzoic acid: A reagent of no less than 95 % (mass fraction) in purity.

d) 1-sodium octane sulfonate: A reagent of no less than 98 % (mass fraction) in purity.

e) 1-sodium hexane sulfonate: A reagent of no less than 98 % (mass fraction) in purity.

f) Boric acid: A JIS Guaranteed Reagent specified in JIS K 8863 or a reagent of equivalent quality.

g) Eluent\(^{(1)(2)}\): Weigh 0.083 g of phthalic acid, 0.552 g of p-hydroxybenzoic acid, 0.195 g of 1-sodium octane sulfonate, 0.376 g of 1-sodium hexane sulfonate and 6.183 g of boric acid to a 1000-mL volumetric flask, add about 500 mL of water to dissolve and add water up to the marked line. Filter with a membrane filter (aperture diameter: no more than 0.5 µm) made of hydrophilic PTFE.

h) Ammonium thiocyanate standard solution (1000 mg/L) \(^{(1)}\): Put 0.1 g of ammonium thiocyanate \(^{(3)}\), specified in JIS K 9000 in weighing dish, and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.

i) Ammonium thiocyanate standard solution (10 mg/L) \(^{(1)}\): At the time of usage, put 10 mL of ammonium thiocyanate standard solution (1000 mg/L) to a 100-mL volumetric flask and add water up to the marked line.

j) Ammonium thiocyanate standard solution for the calibration curve preparation (0.3 mg/L - 3 mg/L): At the time of usage, put 3 mL - 30 mL of ammonium thiocyanate solutions (10 mg/L) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(2) The concentration of prepared solutions is phthalic acid 0.5 mmol/L, p-hydroxybenzoic acid 4.0 mmol/L, 1-sodium octane sulfonate 0.9 mmol/L, 1-sodium hexane sulfonate 2.0 mmol/L and boric acid 100 mmol/L.

(3) It is recommended to store in a desiccator because of deliquescence.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) Ion Chromatograph (IC) or High-Performance Liquid Chromatograph (HPLC): IC specified in JIS K 0127 or HPLC specified in JIS K 0124 that satisfies following requirements.

1) Column: A 4-mm inner diameter 100-mm long stainless steel column tube filled with hydrophilic methacrylate-gel, to which 5-µm particle diameter class 4 ammonium group chemically bonds \(^{(4)}\).
2) **Column bath**: A column bath whose temperature can be adjusted to 55 °C - 60 °C.
3) **Detection unit**: An electric conductivity detector
b) **Membrane filters**: Pore size is no more than 0.5 μm, made of hydrophilic PTFE

**Note** (4) A column is commercially sold under the name Shodex IC NI-424, etc.

(4) **Test procedure**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.

b) Add about 50 mL of water and shake to dissolve, and then add water up to the marked line.

c) Transfer a predetermined amount of the solution, and dilute exactly by a factor of 12.5 with water.

d) Filter with a membrane filter (pore size: no more than 0.5 μm) to make a sample solution.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0127 and as shown below. Specific measurement procedures are according to the operation method of the Ion Chromatograph (IC) or the High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for the Ion Chromatograph (IC) or High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions for an Ion Chromatograph (IC) is shown below. Set up the measurement conditions considering it:

1) **Column**: A hydrophilic methacrylate-gel column (4-mm inner diameter, 100-mm long, 5-μm particle diameter) to which quaternary ammonium group chemically bonds.

2) **Column bath temperature**: 58 °C

3) **Eluent**: Prepared by the procedures in (2) g)

4) **Flow rate**: 1 mL/min

5) **Injection volume**: 20 μL

6) **Detection unit**: An electric conductivity detector

b) **Calibration curve preparation**

1) Inject 20 μL of respective standard solutions for the calibration curve preparation to an IC or an HPLC, and record the chromatogram of electric conductivity to obtain peak area.

2) Prepare a curve for the relationship between the concentration and the peak area of electric conductivity of respective standard solutions for the calibration curve preparation. Prepare a calibration curve when the sample is measured.

**Comment 1** In the measurement of a sample solution, there is a possibility that the recovery rate becomes lower than actual due to the influence of matrix if the concentration is calculated with peak height. Therefore, prepare a calibration curve using peak area.

c) **Sample measurement**

1) Subject 20 μL of sample solution to the same procedure as in b) 1)

2) Obtain the ammonium thiocyanate content from the calibration curve by peak area to calculate ammonium thiocyanate (sulfurized cyanide) in the analytical sample.

**Comment 2** Calculate the concentration by the peak area similarly as the calibration curve preparation to prevent the influence of matrix.

**Comment 3** Note that it takes time to stabilize the baseline due to the usage of the ion-pairing reagent in the elute. It is recommended to take about 120 minutes for stabilization time before starting measurement.
Comment 4 It is possible for the simultaneous measurement of sulfamic acid and ammonium thiocyanate (sulfurized cyanide) in this testing method. In that case, mix a predetermined amount of sulfamic acid standard solution (1000 mg/L) and ammonium thiocyanate standard solution (1000 mg/L), dilute with water to prepare a mixture standard solution (10 mg/L) (1) and use it instead of respective standard solutions (10 mg/L in (2) i). After that, conduct the same procedure in (4.2 b) to calculate the respective concentrations of materials subjected to measurement in the analytical sample.

Comment 5 A recovery testing of ammonium sulfate (3 brands) was conducted. As a result, the mean recovery at additive level of 0.25 % (mass fraction) and 0.075 % (mass fraction) was 101.8 % - 103.7 % and 93.9 % - 97.4 %.
Additionally, the minimum limit of quantification of this testing method is about 0.04 % (mass fraction).

References

(5) Flow sheet for testing method: The flow sheet for ammonium thiocyanate in ammonium sulfate is shown below:

| 1.00 g | Weigh into a 100-mL volumetric flask |
| Shaking to mix | ← About 50 mL of water |
| Water (up to the marked line) | Stopple the volumetric flask and dissolve |
| Dilution | Dilute 12.5 times, water |
| Filtration | Membrane filter (no more than 0.5-µm) |
| Sample solution | |
| Measurement | Ion Chromatograph |

Figure Flow sheet for ammonium thiocyanate in ammonium sulfate
**Reference**: The IC chromatogram of sulfamic acid and thiocyanic acid of the standard solution for the calibration curve preparation and sample solution (ammonium sulfate) are shown below.

(A) Mixture standard solution (the equivalents of 60 ng as sulfamic acid and ammonium thiocyanate (3 mg/L, 20 μL), respectively)

(B) Sample solution (the equivalents of 0.25 % (mass fraction) as sulfamic acid and ammonium thiocyanate added in ammonium sulfate (2500 µg/g), respectively)

Reference diagram: IC chromatogram of sulfamic acid and thiocyanic acid.
(Peak: 1. Sulfamic acid, 2. Thiocyanic acid)

IC measurement conditions
Column: Shodex IC NI-424 (4.6-mm inner diameter, 100-mm long, 5μm-particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
5.8.b High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.8.b-2017 or SCN.b-1.

Add water to an analytical sample, extract ammonium thiocyanate (sulfurized cyanide) and adjust pH as necessary. Introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with a silica gel column to which amino group chemically bonds or a vinyl alcohol polymer column to which amino group chemically bonds, and measure at wavelength 210 nm to obtain ammonium thiocyanate (sulfurized cyanide) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

Nitrous acid and ammonium thiocyanate (sulfurized cyanide) can be simultaneously quantified by using this method. (Refer to Comment 4).

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.
   c) Disodium hydrogen phosphate dodecahydrate: A JIS Guaranteed Reagent specified in JIS K 9019 or a reagent of equivalent quality.
   d) Sodium dihydrogen phosphate dihydrate: A JIS Guaranteed Reagent specified in JIS K 9009 or a reagent of equivalent quality.
   e) Sodium perchlorate monohydrate: A JIS Guaranteed Reagent specified in JIS K 8227 or a reagent of equivalent quality.
   f) Ammonium thiocyanate standard solution (1 mg/L) (1): Put 0.1 g of ammonium thiocyanate specified in JIS K 9000 in weighing dish, and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
   g) Ammonium thiocyanate standard solution (10 mg/L) (1): At the time of usage, put 10 mL of ammonium thiocyanate standard solution (1 mg/L) to a 100-mL volumetric flask and add water up to the marked line.
   h) Ammonium thiocyanate standard solution for the calibration curve preparation (1 μg/mL - 20 μg/mL): At the time of usage, put 1 mL - 20 mL of ammonium thiocyanate standard solution (100 μg/mL) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm to 6-mm inner diameter 150-mm to 250-mm long stainless steel column tube filled with poly vinyl alcohol or silica gel (2), to which 5-μm particle diameter amino group chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 210 nm.
   b) Magnetic stirrer:
   c) Centrifugal separator: A centrifugal separator that can work at about 1700 × g.
   d) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.
   e) pH test paper: A pH test paper infilled with indicator and dried, which can measure the
value from pH 1 to pH 11 and a color change chart with the pH interval value 1 is attached.

**Note (2)** Remaining silanol group of silica gel affects the measurement of ion in some cases. Therefore, use a column which does not affect the measurement of sodium thiocyanate by treating the silanol group. As an example of the treatment, silica gel is to be entirely covered with the uniform membrane of silicone polymer.

**Comment 1** A column is sold under the names CAPCELL PAK NH2 UG80 or Asahipak NH2P-50 4E.

**Comment 2** pH test paper is sold under the name UNIV Test Paper, etc.

(4) **Test procedure**

(4.1) **Extraction:** Conduct extraction as shown below.

(4.1.1) **Powdery test sample**

a) Weigh 1.00 g of an analytical sample, and put it in a 200-mL volumetric flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at $1700 \times g$ centrifugal force for about five minutes (3) and use the supernatant as the extract.

**Note (3)** 16.5-cm of rotor radius and 3000 rpm of revolutions makes about $1700 \times g$ centrifugal force.

(4.1.2) **Fluid test sample**

a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water to the marked line to make the extract

(4.2) **pH adjustment:** Conduct pH adjustment as shown below.

a) Transfer a small amount of the extract to confirm pH value using a pH-test paper.
b) If the pH value in a) is pH 5 or more, transfer the extract to a 1.5-mL ground-in stopper centrifugal precipitate tube (4) and conduct the procedure in f) to prepare a sample solution.
c) If the pH value in a) is pH 4 or less, transfer 40 mL of the extract to a 100-mL beaker.
d) Add a sodium hydroxide solution (5 mg/mL), adjust it to pH 5 to pH 7 with a pH meter and transfer to a 50-mL volumetric flask with water.
e) Add water up to the marked line and transfer it to a 1.5-mL ground-in stopper centrifugal precipitate tube (4).
f) Centrifuge it at $8000 \times g - 10000 \times g$ centrifugal force for about five minutes (5) and use the supernatant as the extract.

**Note (4)** The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about $8100 \times g - 10000 \times g$ centrifugal force.

**Comment 3** Instead of procedures in (4.2) b) and e) - f), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.
(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions for an Ion Chromatograph (IC) is shown below. Set up the measurement conditions considering it:

1) **Column**: A vinyl alcohol polymer column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino group chemically bonds or a silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino group chemically bonds.

2) **Column bath temperature**: 30 °C - 40 °C

3) **Eluent**: Dissolve 1.79 g of disodium hydrogen phosphate dodecahydrate, 0.78 g of sodium dihydrogen phosphate dihydrate and 14.04 g of sodium perchlorate monohydrate in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.

4) **Flow rate**: 0.9 mL/min - 1.0 mL/min

5) **Injection volume**: 10 µL

6) **Detection unit**: An absorptiometric detector, measurement wavelength: 210 nm

b) **Calibration curve preparation**

1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 210 nm and obtain the peak area.

2) Prepare a curve for the relationship between the concentration and the peak area at wavelength 210 nm of the respective standard solutions for the calibration curve preparation.

c) **Sample measurement**

1) Subject 10 µL of sample solution to the same procedure as in b) 1)

2) Obtain the ammonium thiocyanate content from the calibration curve by peak area to calculate ammonium thiocyanate (sulfurized cyanide) in the analytical sample.

**Comment 4** This testing method enables the simultaneous measurement of nitrous acid and ammonium thiocyanate (sulfurized cyanide). In this case, mix a predetermined amount of nitrous acid standard solution (1 mg/mL) and ammonium thiocyanate standard solution (1 mg/mL), dilute with water to prepare a mixture standard solution (100 µg/mL) \(^{(1)}\) and use it instead of (2) h) ammonium thiocyanate standard solution (100 µg/mL). After that, conduct the same procedure in (4.3) b) to calculate the respective concentrations of materials subjected to measurement in the analytical sample.

**Comment 5** A recovery testing was conducted using samples that ammonium thiocyanate equivalent to 1/5 - 5 times of permissible content are added to an ammonium sulfate fertilizer (1 brand), a coating nitrogen fertilizer (1 brand), a blended fertilizer (2 brands), a compound fertilizer (1 brand) and a fluid mixed fertilizer (1 brand). As a result, the mean recovery at the additive level of 0.025 % (mass fraction), 0.01 % (mass fraction), 0.005 % (mass fraction) and 0.0025 % (mass fraction) are 95.4 % - 100.5 %, 94.7 % - 103.8 %, 83.8 % - 109.0 % and 87.2 % - 103.3 % respectively. Table 1 shows results and analysis results from a collaborative study for test method validation. Ammonium thiocyanate had sufficient reproducibility in the range from 0.00476 % (mass fraction) to 0.204 %. (mass fraction). Additionally, the minimum limit of quantification of this testing method is about 0.002 % (mass fraction).
<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^{2)}$ (%)$^{3)}$</th>
<th>$s_r^{4)}$ (%)$^{3)}$</th>
<th>$RSD_r^{5)}$ (%)</th>
<th>$s_R^{6)}$ (%)$^{3)}$</th>
<th>$RSD_R^{7)}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Home garden-use mixed fertilizer 1</td>
<td>10</td>
<td>0.00476</td>
<td>0.00019</td>
<td>4.1</td>
<td>0.00060</td>
<td>12.7</td>
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<tr>
<td>Home garden-use mixed fertilizer 2</td>
<td>9</td>
<td>0.00976</td>
<td>0.00029</td>
<td>2.9</td>
<td>0.00050</td>
<td>4.7</td>
</tr>
<tr>
<td>Home garden-use mixed fertilizer 3</td>
<td>9</td>
<td>0.0506</td>
<td>0.0019</td>
<td>3.7</td>
<td>0.0022</td>
<td>4.3</td>
</tr>
<tr>
<td>Compound fertilizer 1</td>
<td>10</td>
<td>0.10</td>
<td>0.002</td>
<td>2.3</td>
<td>0.003</td>
<td>2.6</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>11</td>
<td>0.204</td>
<td>0.006</td>
<td>2.7</td>
<td>0.008</td>
<td>3.7</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>10</td>
<td>0.00989</td>
<td>0.00037</td>
<td>3.8</td>
<td>0.00060</td>
<td>6.5</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis  
2) Mean ($n =$ number of laboratories $\times$ number of samples (2))  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Reproducibility standard deviation  
7) Reproducibility relative standard deviation

References


(5) **Flow sheet for testing method:** The flow sheet for ammonium thiocyanate in fertilizers is shown below:

**Figure 1-1**  Flow sheet for ammonium thiocyanate in fertilizers
(Extraction procedure (4.1.1))

```
1.00 g analytical sample  200-mL ground-in stopper Erlenmeyer flask
(powdery)← 100 mL of water

Extraction  Stir to mix, 10 minutes

Centrifugal separation  Ground-in stopper centrifugal precipitate tube,
1700 ×g , 5 minutes

Filtrate
```

**Figure 1-2**  Flow sheet for ammonium thiocyanate in fertilizers
(Extraction procedure (4.1.2))

```
1.00 g analytical sample  100-mL volumetric flask
(fluid)← 50 mL - 70 mL of water

Extraction  Shake to mix
← Water (up to the marked line)

Filtrate
```
Figure 2 Flow sheet for ammonium thiocyanate in fertilizers (pH adjustment and measurement procedure)
Reference: HPLC chromatogram of nitrous acid and ammonium thiocyanate are shown below.

(A) Mixture standard solution
(The equivalents of 100 ng (10 µg/mL, 10 µL) as nitrous acid and ammonium thiocyanate, respectively)

(B) Sample solution
(The equivalents of 0.1 % (mass fraction) as nitrous acid and ammonium thiocyanate added in blended fertilizer respectively)

Reference diagram: HPLC chromatogram of nitrous acid and ammonium thiocyanate
(Peak: 1. Nitrous acid, 2. Thiocyanate)

Measurement conditions for HPLC
Column: CAPCELL PAK NH2 UG80 (4.6-mm inner diameter, 250-mm long, 5-µm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.3) a)
5.9 Nitrous acid
5.9.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.9.a-2017 or NO2.a-1.

Add water to an analytical sample, extract nitrous acid and adjust pH as necessary. Introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with a silica gel column to which amino group chemically bonds or a vinyl alcohol polymer column to which amino group chemically bonds, and measure at wavelength 210 nm to obtain nitrous acid in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

Nitrous acid and ammonium thiocyanate (sulfurized cyanide) can be simultaneously quantified by using this method. (Refer to Comment 4).

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Sodium hydroxide: A JIS Guaranreed Reagent specified in JIS K 8576 or a reagent of equivalent quality.
   c) Disodium hydrogen phosphate dodecahydrate: A JIS Guaranreed Reagent specified in JIS K 9019 or a reagent of equivalent quality.
   d) Sodium dihydrogen phosphate dihydrate: A JIS Guaranreed Reagent specified in JIS K 9009 or a reagent of equivalent quality.
   e) Sodium perchlorate monohydrate: A JIS Guaranreed Reagent specified in JIS K 8227 or a reagent of equivalent quality.
   f) Nitrous Acid standard solution (1 mg/mL)\(^{(1)}\): Put 0.147 g of sodium nitrite specified in JIS K 8019 in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
   g) Nitrous acid standard solution (100 μg/mL)\(^{(1)}\): At the time of usage, put 10 mL of nitrous acid standard solution (1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   h) Nitrous acid standard solution for the calibration curve preparation (1 μg/mL - 20 μg/mL): At the time of usage, put 1 mL - 20 mL of nitrous acid standard solution (100 μg/mL) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube filled with poly vinyl alcohol or silica gel \(^{(2)}\), to which 5-μm particle diameter amino group chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 30 °C - 45 °C.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 210 nm.
   b) Magnetic stirrer:
   c) Centrifugal separator: A centrifugal separator that can work at about 1700 × g.
   d) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.
   e) pH test paper: A pH test paper infilled with indicator and dried, which can measure the value from pH 1 - pH 11 and a color change chart with the pH interval value 1 is attached.
Note (2) Remaining silanol group of silica gel affects the measurement of ion in some cases. Therefore, use a column which does not affect the measurement of nitrous acid by treating the silanol group. As an example of the treatment, silica gel is to be entirely covered with the uniform membrane of silicone polymer.

Comment 1 A column is sold under the names CAPCELL PAK NH2 UG80 or Asahipak NH2P-50 4E.

Comment 2 pH test paper is sold under the name UNIV Test Paper, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Powdery test sample
a) Weigh 1.00 g of an analytical sample, and put it into a 200-mL ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at 1700 \( \times g \) centrifugal force for about five minutes \(^{(3)}\) and use the supernatant as the extract.

Note (3) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 \( \times g \) centrifugal force.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water to the marked line to make the extract

(4.2) pH adjustment: Conduct pH adjustment as shown below.
a) Transfer a small amount of the extract to confirm pH value using a pH-test paper.
b) If the pH value in a) is pH 5 or more, transfer the extract to a 1.5-mL ground-in stopper centrifugal precipitate tube \(^{(4)}\) and conduct the procedure in f) to prepare a sample solution.
c) If the pH value in a) is pH 4 or less, transfer 40 mL of the extract to a 100-mL beaker.
d) Add a sodium hydroxide solution (5 mg/mL), adjust it to pH 5 to pH 7 with a pH meter and transfer to a 50-mL volumetric flask with water.
e) Add water up to the marked line and transfer it to a 1.5-mL ground-in stopper centrifugal precipitate tube \(^{(4)}\).
f) Centrifuge it at 8000 \( \times g \) - 10000 \( \times g \) centrifugal force for about five minutes \(^{(5)}\) and use the supernatant as the extract.

Note (4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 \( \times g \) - 10000 \( \times g \) centrifugal force.

Comment 3 Instead of procedures in (4.2) b) and e) - f), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-\( \mu \)m) made of hydrophilic PTFE and the filtrate can be the sample solution.
(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of High-Performance Liquid Chromatograph (HPLC) used in measurement.

**a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions for an Ion Chromatograph (IC) is shown below. Set up the measurement conditions considering it:

1) **Column**: A vinyl alcohol polymer column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino group chemically bonds or a silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino group chemically bonds.

2) **Column bath temperature**: 30 °C - 40 °C

3) **Eluent** (i): Dissolve 1.79 g of disodium hydrogen phosphate dodecahydrate, 0.78 g of sodium dihydrogen phosphate dihydrate and 14.04 g of sodium perchlorate monohydrate in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.

4) **Flow rate**: 0.9 mL/min - 1.0 mL/min

5) **Injection volume**: 10 µL

6) **Detection unit**: An absorptiometric detector, measurement wavelength: 210 nm

**b) Calibration curve preparation**

1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 210 nm and obtain the peak area.

2) Prepare a curve for the relationship between the concentration and the peak area at wavelength 210 nm of the respective standard solutions for the calibration curve preparation.

**c) Sample measurement**

1) Subject 10 µL of sample solution to the same procedure as in b) 1)

2) Obtain the nitrous acid content by the peak area using the calibration curve to calculate the concentration of nitrous acid in the analytical sample.

**Comment 4** This testing method enables the simultaneous measurement of nitrous acid and ammonium thiocyanate (sulfurized cyanide). In this case, mix a predetermined amount of nitrous acid standard solution (1 mg/mL) and ammonium thiocyanate standard solution (1 mg/mL), dilute with water to prepare a mixture standard solution (100 µg/mL) (i) and use it instead of (2) h) nitrous acid standard solution (100 µg/mL). After that, conduct the same procedure in (4.3) b) to calculate the respective concentrations of materials subjected to measurement in the analytical sample.

**Comment 5** Recovery testing was conducted using samples that nitrous acid equivalent to 1/5 - 5 times of permissible content are added to ammonium sulfate fertilizer (1 brand), coating nitrogen fertilizer (1 brand), blended fertilizer (2 brands), compound fertilizer (1 brand) and fluid mixed fertilizer (1 brand). As a result, the mean recovery at the additive level of 0.1 % (mass fraction), 0.04 % (mass fraction), 0.02 % (mass fraction) and 0.01 % (mass fraction) are 99.0 % - 100.8 %, 100.4 % - 102.0 %, 103.1 % - 106.6 % and 101.2 % - 105.9 % respectively.

Table 1 shows results and analysis results from a collaborative study for test method validation. Nitrous acid had sufficient reproducibility in the range from 0.0255 % (mass fraction) to 0.291 %. (mass fraction).

Additionally, the minimum limit of quantification of this testing method is about 0.0003 % (mass fraction).
Table 1 Results and statistical analysis results from a collaborative study for the nitrous acid method

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean 3) (%)</th>
<th>Repeatability standard deviation 4) (%)</th>
<th>Repeatability relative standard deviation 5) (%)</th>
<th>Reproducibility standard deviation 6) (%)</th>
<th>Reproducibility relative standard deviation 7) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Home garden-use mixed fertilizer 1</td>
<td>10</td>
<td>0.0502</td>
<td>0.0005</td>
<td>1.1</td>
<td>0.0009</td>
<td>1.7</td>
</tr>
<tr>
<td>Home garden-use mixed fertilizer 2</td>
<td>11</td>
<td>0.0255</td>
<td>0.00070</td>
<td>2.6</td>
<td>0.0009</td>
<td>3.5</td>
</tr>
<tr>
<td>Home garden-use mixed fertilizer 3</td>
<td>9</td>
<td>0.150</td>
<td>0.004</td>
<td>2.9</td>
<td>0.005</td>
<td>3.6</td>
</tr>
<tr>
<td>Compound fertilizer 1</td>
<td>10</td>
<td>0.202</td>
<td>0.004</td>
<td>1.9</td>
<td>0.004</td>
<td>2.2</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>10</td>
<td>0.291</td>
<td>0.004</td>
<td>1.3</td>
<td>0.005</td>
<td>1.7</td>
</tr>
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<td>Compound fertilizer 3</td>
<td>10</td>
<td>0.0498</td>
<td>0.0007</td>
<td>1.4</td>
<td>0.0010</td>
<td>2.0</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (n = number of laboratories \times number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(5) **Flow sheet for testing method:** The flow sheet for nitrous acid in fertilizers is shown below:

![Flow sheet for nitrous acid in fertilizers](image1)

![Flow sheet for nitrous acid in fertilizers](image2)

*Figure 1-1  Flow sheet for nitrous acid in fertilizers (Extraction procedure (4.1.1))

*Figure 1-2  Flow sheet for nitrous acid in fertilizers (Extraction procedure (4.1.2))

Figure 2  Flow sheet for nitrous acid in fertilizers

- Filtrate
- pH value confirmation
- pH 5 or more
  - pH test paper
- pH 4 or less
  - Aliquot (40 mL)
  - 100-mL beaker
  - pH adjustment (pH 5 - pH 7)
  - Sodium hydroxide solution (5 mg/mL)
  - Transfer
  - 50-mL volumetric flask, water
    - Water (up to the marked line)
  - Centrifugal separation
    - Ground-in stopper centrifugal precipitate tube, 8000 × g - 10000 × g, 5 minutes
  - Sample
  - Measurement
    - High Performance Liquid Chromatograph
**Reference:** HPLC chromatogram of nitrous acid and ammonium thiocyanate are shown below.

(A) Mixture standard solution
(The equivalents of 100 ng (10 μg/mL, 10 μL) as nitrous acid and ammonium thiocyanate, respectively)

(B) Sample solution
(The equivalents of 0.1 % (mass fraction) as nitrous acid and ammonium thiocyanate added in blended fertilizer respectively)

Reference diagram: HPLC chromatogram of nitrous acid and ammonium thiocyanate

Measurement conditions for HPLC
Column: CAPCELL PAK NH2 UG80 (4.6-mm inner diameter, 250-mm long, 5-μm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.3) a)
5.10 Biuret nitrogen
5.10.a High-Performance Liquid Chromatography
(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 5.10.a-2017 or B-N.a-1.
Add water to an analytical sample to extract biuret, introduce it to a High-Performance Chromatograph (HPLC) to isolate it with a weak acid ion-exchange column, and then measure at wavelength 190 nm to obtain biuret nitrogen (B-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.
Dicyandiamide nitrogen (Dd-N), urea nitrogen (U-N), guanidine nitrogen (Gd-N) and guanyleurea nitrogen (GU-N) and can be simultaneously quantified by using this method. (Refer to Comment 5).

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Potassium dihydrogen phosphate: A JIS Guaranteed Reagent specified in JIS K 9007 or a reagent of equivalent quality.
   c) Phosphoric acid: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.
   d) Biuret nitrogen standard solution (B-N 2 mg/mL): Put 0.491 g of biuret [C2H5N3O2] in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water, transfer to a 100-mL volumetric flask and warm up to 50ºC to dissolve. After standing to cool, add water up to the marked line.
   e) Biuret nitrogen standard solution (B-N 200 µg/mL): Put 10 mL of biuret nitrogen standard solution (B-N 2 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   f) Biuret nitrogen standard solution (B-N 50 µg/mL - 100 µg/mL): Put 25 mL - 50 mL of biuret nitrogen standard solution (B-N 200 µg/mL) to 100-mL volumetric flasks and add water up to the marked line.
   g) Biuret nitrogen standard solution for the calibration curve preparation (B-N 1 µg/mL - 50 µg/mL): At the time of usage, put 1 mL - 50 mL of biuret nitrogen standard solution (B-N 100 µg/mL) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) A reagent of no less than 97% (mass fraction) in purity is commercially sold as biuret.
   (3) If biuret nitrogen standard solution (B-N 2 mg/mL) is stored in a refrigerator, precipitates may appear. Therefore, it is recommended to store it at room temperature. In addition, sudden cooling should be avoided.

Comment 1 Biuret is sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Tokyo Chemical Industry Co., Ltd.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 7.5-mm inner diameter 100-mm long stainless steel column tube filled with weak acid ion-exchange resin.
      2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45ºC.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 190 nm.
   b) Magnetic stirrer:
   c) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000...
× g.

Comment 2 A column is sold under the production name Asahipak ES-502C 7C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Powdery test sample
a) Weigh 1.00 g of an analytical sample, and put it into a 200-mL ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution (4) to a 1.5-mL ground-in stopper centrifugal precipitate tube (5).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (6) and use the supernatant as the extract.

Note (4) If there is a possibility that the biuret nitrogen (B-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

(5) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(6) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line (7) and transfer to a 1.5-mL ground-in stopper centrifugal precipitate tube (5).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (6) and use the supernatant as the extract.

Note (7) If there is a possibility that the biuret nitrogen (B-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of precisely adjusted solution with water.

Comment 3 Instead of procedures in (4.1.1) c) - d) or (4.1.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions is shown below. Set up the measurement conditions considering it:
1) Column: A weak acid ion-exchange resin column (7.5-mm inner diameter, 100-mm long, 5-µm - 10-µm particle diameter)
2) Column bath temperature: 40 °C
3) Eluent (1): Dissolve 3.92 g of potassium dihydrogen phosphate and 0.12 g of phosphoric acid in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than
0.5-µm) made of hydrophilic PTFE.

4) **Flow rate:** 0.6 mL/min  
5) **Injection volume:** 10 µL  
6) **Detection unit:** An absorptiometric detector, measurement wavelength: 190 nm

**Comment 4** Eluent can be prepared as follows. Dissolve 19.6 g of potassium dihydrogen phosphate and 0.584 g of phosphoric acid with water to make 500 mL and store in a refrigerator. At the time of usage, dilute a predetermined volume of the solution by a factor of 10 and filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.

**b) Calibration curve preparation**  
1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 190 nm and obtain the peak height.  
2) Prepare a curve for the relationship between the biuret nitrogen (B-N) concentration and the peak height at wavelength 190 nm of the respective standard solutions for the calibration curve preparation.

**c) Sample measurement**  
1) Subject 10 µL of sample solution to the same procedure as in b) 1)  
2) Obtain the biuret nitrogen (B-N) content from the peak height using the calibration curve to calculate the biuret nitrogen (B-N) in the analytical sample.

**Comment 5** This testing method enables the simultaneous measurement of biuret nitrogen (B-N), dicyandiamide nitrogen (Dd-N), urea nitrogen (U-N), guanidine nitrogen (Gd-N) and a guanylurea nitrogen standard solution (GU-N). In this case, mix a predetermined amount of biuret nitrogen (B-N 2 mg/mL), an urea nitrogen standard solution (U-N 2 mg/mL), a dicyandiamide nitrogen standard solution (Dd-N 2 mg/mL), a guanidine nitrogen standard solution (Gd-N 2 mg/mL) and a guanylurea nitrogen standard solution (GU-N 2 mg/mL), dilute with water to prepare a mixture standard solution (200 µg/mL) (1) and use it instead of (2) e) a biuret nitrogen standard solution (B-N 200 µg/mL). After that, conduct the same procedure in (4.2) b) to calculate the respective concentrations of materials subjected to measurement in the analytical sample.

**Comment 6** Additive recovery testing was conducted using one brand of an acetaldehyde condensed urea fertilizer, a compound fertilizer, a blended fertilizer, a fluid compound fertilizer and a home garden-use mixed fertilizer respectively. As a result, the mean recovery at additive level of 0.2 % (mass fraction) and 0.1 % (mass fraction) and 0.02 % (mass fraction) were 87.0 % - 95.1 %, 90.6 % - 101.1 % and 91.2 % - 105 5 % respectively. The results of the repeatability tests on different days using a blended fertilizer, a compound fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this testing method is about 0.005 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average 2) (%)</th>
<th>Repeatability 4) (%)</th>
<th>Intermediate precision 5) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blended fertilizer</td>
<td>5</td>
<td>0.204</td>
<td>0.0006</td>
<td>0.3</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>5</td>
<td>0.0969</td>
<td>0.0006</td>
<td>0.6</td>
</tr>
<tr>
<td>Home garden-use mixed fertilizer</td>
<td>5</td>
<td>0.0103</td>
<td>0.0001</td>
<td>0.9</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) \( \times \) the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation

Table 2 Results and statistical analysis results from a collaborative study for the test method validation of biuret nitrogen method

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories 1)</th>
<th>Mean 2) (%)</th>
<th>s (_{r}) 4) (%)</th>
<th>RSD (_{r}) 5) (%)</th>
<th>s (_{R}) 6) (%)</th>
<th>RSD (_{R}) 7) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>9</td>
<td>0.00963</td>
<td>0.00030</td>
<td>3.1</td>
<td>0.00029</td>
<td>3.1</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>10</td>
<td>0.0201</td>
<td>0.0003</td>
<td>1.6</td>
<td>0.0007</td>
<td>3.4</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>12</td>
<td>0.114</td>
<td>0.013</td>
<td>11.7</td>
<td>0.017</td>
<td>15.3</td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>11</td>
<td>0.212</td>
<td>0.017</td>
<td>7.8</td>
<td>0.026</td>
<td>12.4</td>
</tr>
<tr>
<td>Nitrogen fertilizer</td>
<td>12</td>
<td>0.832</td>
<td>0.050</td>
<td>6.0</td>
<td>0.086</td>
<td>10.3</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (n = number of laboratories \( \times \) number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(5) **Flow sheet for testing method:** The flow sheet for biuret nitrogen in fertilizers is shown below:

![Flow sheet for biuret nitrogen in fertilizers](image)

- **Extraction**
- **Centrifugal separation**
- **Sample solution**
- **Measurement**

**Figure 1** Flow sheet for biuret nitrogen in fertilizers
(Extraction procedure (4.1.1) and measurement)

- **Extraction**
- **Centrifugal separation**
- **Sample solution**
- **Measurement**

**Figure 2** Flow sheet for biuret nitrogen in fertilizers
(Extraction procedure (4.1.2) and measurement)
**Reference:** The chromatogram of the standard solution of biuret nitrogen for the calibration curve preparation is shown below.

Reference diagram  HPLC chromatogram of the mixture standard solutions for the calibration curve preparation (10 mg/L for each)

Peak name
(1) Urea nitrogen  (2) Biuret nitrogen  (3) Dicyandiamide nitrogen  
(4) Guanidine nitrogen  (5) Guanylurea nitrogen

**Measurement conditions for HPLC**
Column: Asahipak ES-502C 7C  (7.5-mm inner diameter, 100-mm long, 9-μm particle diameter)  
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
5.11 Titanium
5.11.a ICP Optical Emission Spectrometry (1)

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 5.11.a-2017 or Ti.a-1.

Pretreat an analytical sample with nitric acid - sulfuric acid - perchloric acid, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the emission with titanium at a wavelength of 334.941 nm to obtain the titanium (Ti) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   c) Sulfuric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   d) Perchloric acid: A reagent of harmful metal analysis grade, microanalysis grade or equivalents.
   e) Hydrochloric acid: Hydrochloric acid for arsenic analysis specified in JIS K 8180, or of harmful metal analysis grade, microanalysis grade or equivalents.
   f) Titanium standard solution (Ti 1 mg/mL): A titanium standard solution (Ti 1 mg/mL) traceable to National Metrology.
   g) Titanium standard solution (Ti 0.1 mg/mL) (1): Transfer 10 mL of titanium standard solution (Ti 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
   h) Titanium standard solutions (Ti 0.1 µg - 20 µg/mL) for the calibration curve preparation (1): Transfer 1 mL - 20 mL of titanium standard solution (Ti 0.1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
   i) Blank test solution for the calibration curve preparation (1): Hydrochloric acid (1+23) used in the procedures in g) and h).

Note (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in h) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, in case of using the ICP-OES of an axial observation mode, it is recommended to prepare a titanium standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Instruments: Instruments are as shown below:
   a) ICP Optical Emission Spectrometry: A spectrophotometer specified in JIS K 0116
      1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity
      b) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 350 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to no less than 300 °C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
b) Add about 10 mL of nitric acid and about 5 mL of sulfuric acid, cover the tall beaker with a watch glass, and leave at rest overnight.

c) Heat mildly on a hot plate or sand bath at 170 °C - 220 °C for no less than 30 minutes. After bubbles cease to form, set the temperature of the hot plate or sand bath to no less than 300 °C, and heat until nitroxide (yellow-brown smoke) is no longer generated (2).

d) After standing to cool, add about 5 mL of perchloric acid.

e) Cover the tall beaker with a watch glass, and heat on a hot plate or sand bath at no less than 300 °C for 2 - 3 hours to digest (3).

f) Slightly move the watch glass (4), and keep on heating on the hot plate or sand bath to concentrate until the liquid volume becomes no more than 2 mL.

g) After standing to cool, add about 5 mL of hydrochloric acid (1+10) and about 20 mL of water, cover the tall beaker with a watch glass and heat mildly to dissolve.

h) After standing to cool, transfer to a 100-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper to make a sample solution.

i) As a blank test, conduct the procedures in b) - h) using another tall beaker to prepare the blank test solution.

Note (2) Oxidation of organic matters by perchloric acid progresses extremely rapidly and explosively. For that reason, add perchloric acid after fully degrading organic matters with nitric acid to avoid danger.

(3) When the white smoke of perchloric acid is generated, if the solution is colored such as black-brown or brown, stop heating immediately, and after standing to cool, add nitric acid, and heat again to degrade remaining organic matters.

(4) The watch glass can be removed.

Comment 2 When the analytical sample solidifies in the procedure in (4.1) b), moisten the analytical sample with a small amount of water as necessary in advance.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

Analytical line wavelength: 334.941 nm

b) Calibration curve preparation

1) Spray the titanium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 334.941 nm.

2) Prepare a curve for the relationship between the titanium concentration and the indicated value of the titanium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement

1) Transfer a predetermined amount of the sample solution (the equivalents of 0.01 mg - 2 mg as titanium) to a 100-mL volumetric flask.

2) Add 25 mL of hydrochloric acid (1+23) to the marked line.

3) Subject to the same procedure as in b) 1) to read the indicated value.

4) Obtain the titanium content from the calibration curve, and calculate titanium (Ti) in the analytical sample.
Comment 3 Additive recovery testing was conducted using an autoclaved lightweight concrete powdery fertilizer, a mixed phosphate fertilizer, a compound fertilizer, a slag manganese fertilizer, a solid fertilizer, a fluid compound fertilizer, a mixed compost compound fertilizer and a blended fertilizer respectively. As a result, the mean recovery at additive level of 0.01 % (mass fraction) - 0.5 % (mass fraction) were 92.9 % - 99.5 % respectively.

The results of the repeatability tests on different days using a mixed phosphate fertilizer and a compound fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.001 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Average&lt;sup&gt;2&lt;/sup&gt;</td>
<td>s&lt;sup&gt;4&lt;/sup&gt;</td>
</tr>
<tr>
<td>Mixed phosphate fertilizer</td>
<td>7</td>
<td>0.950</td>
<td>0.013</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>0.130</td>
<td>0.002</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) \times the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References
Flow sheet for testing method: The flow sheet for titanium in fertilizers is shown below:

1.00 - 2.00g analytical sample

200-mL - 300-mL tall beaker.

- A small amount of water, moisten the analytical sample (as appropriate)
- About 10 mL of nitric acid
- About 5 mL of sulfuric acid

Leave at rest overnight

Cover with a watch glass

Heat gently on a hot plate or a sand bath at 170 °C - 220 °C for no less than 30 minutes

Heat on a hot plate or sand bath at no less than 300 °C, until yellow-brown smoke no longer evolves.

Room temperature

- About 5 mL of perchloric acid

Cover with a watch glass, digest on a hot plate or a sand bath at 300 °C or more for 2-3 hours

Slightly move the watch glass, concentrate until it becomes no more than 2mL

Room temperature

- About 5 mL of hydrochloric acid (1+10)
- About 20 mL of water

Cover with a watch glass, and dissolve

Room temperature

Transfer

100-mL volumetric flask

- Water (up to the marked line)

Filtration

Type 3 filter paper

Sample solution

Figure 1 Flow sheet for titanium in fertilizers (Extraction procedure)

Sample solution

Aliquot (Predetermined)

100-mL volumetric flask

- About 25 mL of hydrochloric acid (1+23)
- Water (up to the marked line)

Measurement

ICP Optical Emission Spectrometer (334.941 nm)

Figure 2 Flow sheet for titanium in fertilizers (Measurement procedure)
5.11.b  ICP Optical Emission Spectrometry (2)

(1) Summary

The testing method is applicable to slag silicate fertilizers. This testing method is classified as Type D and its symbol is 5.11.b-2017 or Ti.b-1.

Melt an analytical sample with ammonium hydrogensulfate, introduce it to an ICP Optical Emission Spectrometer (“ICP-OES”) and measure the emission with titanium at a wavelength of 334.941 nm to obtain the titanium (Ti) in an analytical sample. In addition, the performance of this testing method is shown in Comment 2.

(2) Reagent, etc.: Reagents and water are as shown below.

b) Ammonium hydrogensulfate: A reagent of no less than 98 % (mass fraction) in purity.
c) Hydrochloric acid: Hydrochloric acid for arsenic analysis specified in JIS K 8180, or of harmful metal analysis grade, microanalysis grade or equivalents.
d) Titanium standard solution (Ti 1 mg/mL): A titanium standard solution (Ti 1 mg/mL) traceable to National Metrology.
e) Titanium standard solution (Ti 0.1 mg/mL) \(^{(1)}\): Transfer 10 mL of titanium standard solution (Ti 1 mg/mL) to a 100-mL volumetric flask and add hydrochloric acid (1+23) up to the marked line.
f) Titanium standard solutions (Ti 0.1 µg - 20 µg/mL) for the calibration curve preparation \(^{(1)}\): Transfer 1 mL - 20 mL of titanium standard solution (Ti 1 mg/mL) to 100-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) to the marked line.
g) Blank test solution for the calibration curve preparation \(^{(1)}\): Hydrochloric acid (1+23) used in the procedures in d), e) and f).

Note  (1) This is an example of preparation; prepare an amount as appropriate.

Comment 1 There are two modes to observe emission from an ICP-OES, a horizontal observation mode and an axial observation mode. The range of the concentration of the standard solution for the calibration preparation curve in f) applies to a horizontal observation mode. While an axial observation mode can observe a measurement component of low-level concentrations, it cannot gain the linearity of a calibration curve in the range of high-level concentrations. Therefore, in case of using the ICP-OES of an axial observation mode, it is recommended to prepare a titanium standard solution for the calibration curve in the concentration range which is suitable to a device used.

(3) Instruments: Instruments are as shown below:


1) Gas: Argon gas specified in JIS K 1105 of no less than 99.5 % (volume fraction) in purity

b) Hot plate: A hot plate whose surface temperature can be adjusted up to 400 ºC.

(4) Test procedure

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample and transfer to a 200-mL tall beaker.
b) Add about 10 g of ammonium hydrogensulfate \(^{(2)}\).
c) Heat on a hot plate at no less than 350 ºC and make the melted ammonium hydrogensulfate come into complete contact with the analytical sample \(^{(3)}\).
d) Cover with a watch glass, and heat at no less than 350 ºC for 1 hour.
e) After standing to cool, add about 25 mL of hydrochloric acid (1+5), cover the tall beaker with
a watch glass and heat mildly to dissolve.

f) After standing to cool, transfer to a 100-mL volumetric flask with water, add water up to the marked line, and filter with Type 3 filter paper.

g) Transfer 10 mL of the filtrate to another 100-mL volumetric flask and add hydrochloric acid (1+23) to the marked line to make a sample solution.

h) As a blank test, conduct the procedures in b) - f) using another tall beaker to prepare the blank test solution.

Note (2) Ammonium hydrogensulfate is corrosive. Therefore, use a dispensing spoon made of resin.

(3) After the ammonium hydrogensulfate melts, make it come into contact with the analytical sample by tilting the tall beaker or conducting a similar procedure.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0116 and as shown below. Specific measurement procedures are according to the operation method of the ICP Optical Emission Spectrometry used in measurement.

a) Measurement conditions for the ICP Atomic Emission Spectrometer: Set up the measurement conditions for the ICP Atomic Emission Spectrometer considering the following:

   Analytical line wavelength: 334.941 nm

b) Calibration curve preparation

1) Spray the titanium standard solution for the calibration curve preparation and blank test solution for the calibration curve preparation into inductively coupled plasma, and read the indicated value at a wavelength of 334.941 nm.

2) Prepare a curve for the relationship between the titanium concentration and the indicated value of the titanium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.

c) Sample measurement

1) Subject the sample solution to the same procedure as in b) 1) to read the indicated value.

2) Obtain the titanium content from the calibration curve, and calculate titanium (Ti) in the analytical sample.

Comment 2 Recovery testing was conducted to evaluate trueness using slag silicate fertilizers (2 samples). As a result, the average rate of recovery at additive level of 0.1 % (mass fraction) - 0.2 % (mass fraction) are 95.1 % - 98.2 %.

The results of the repeatability tests on different days using slag silicate fertilizers (2 samples) to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Average $^2)$</td>
<td>$s_r^4$</td>
</tr>
<tr>
<td>Slag silicate fertilizer 1</td>
<td>7</td>
<td>0.525</td>
<td>0.005</td>
</tr>
<tr>
<td>Slag silicate fertilizer 2</td>
<td>7</td>
<td>0.112</td>
<td>0.002</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $T$) $\times$ the number of duplicate testing (2)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

(5) **Flow sheet for testing method:** The flow sheet for titanium in fertilizers is shown below:

![Flow sheet for testing method](image)

Figure 1  Flow sheet for titanium in fertilizers (Extraction procedure)

![Flow sheet for measurement](image)

Figure 2  Flow sheet for titanium in fertilizers (Measurement procedure)
5.12 Sulfurous Acid
This method is according to 5.3 Sulfurous acid analysis in “The Official Methods of Analysis of Fertilizers 1992”.

References
6. Testing relating to the other limitations
6.1 Dicyandiamide nitrogen
6.1.a High-Performance Liquid Chromatography (1)

(1) Summary
The testing method is applicable to nitrolime and fertilizers containing nitrolime. This testing method is classified as Type B and its symbol is 6.1.a-2017 or Dd-N.a-1. Add methanol to an analytical sample to extract dicyandiamide (Dd), introduce it to a High-Performance Liquid Chromatograph (HPLC), isolate with an amino propyl silica gel column and measure at wavelength 215 nm to obtain dicyandiamide nitrogen (Dd-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent: Reagents are as shown below.
   a) Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.
   b) Methanol: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.
   c) Acetonitrile: A reagent of HPLC grade or equivalents.
   d) Dicyandiamide standard solution (1 mg/mL): Put 0.1 g of dicyandiamide [C₂H₄N₄] to a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of methanol to dissolve, transfer to a 100-mL volumetric flask and add the solvent up to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.
   e) Dicyandiamide standard solution (0.1 mg/mL): Put 10 mL of dicyandiamide standard solution (1 mg/mL) to a 100-mL volumetric flask and add methanol up to the marked line.
   f) Dicyandiamide standard solution (10 µg/mL - 50 µg/mL) for the calibration curve preparation: At the time of usage, put 5 mL - 25 mL of dicyandiamide standard solution (0.1 mg/mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.
   g) Dicyandiamide standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation: In the case of usage, transfer 2.5 mL - 25 mL of dicyandiamide standard solution for the calibration curve preparation (20 µg/mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) A reagent of no less than 98 % (mass fraction) in purity as dicyandiamide is commercially sold.

Comment 1 Dicyandiamide is commercially sold as dicyanodiamide by FUJIFILM Wako Pure Chemical Co., Ltd. and Kanto Chemical Co., Inc.

(3) Instruments: Instruments are as shown below:
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A column of 4 mm - 6 mm inner diameter and 150 mm - 250 mm long stainless steel column tube filled with silica gel, to which amino or amino propyl chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 30 °C - 45 °C.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 215 nm.
   b) Shaking apparatus
   c) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.

Comment 2 A column is sold under production names such as Hibar LiChrosorb NH₂, Inertsil
NH₂, Unison UK-Amino, Mighty sil NH₂, Shim-pack CLC-NH₂, Shodex NH-5A, Unisil Q NH₂, etc.

(4) Test procedures

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL - 200-mL ground-in stopper Erlenmeyer flask.

b) Immediately add 100 mL of methanol (3) and shake to mix by using a shaking apparatus for about 10 minutes.

c) After allowing to stand still, transfer a supernatant solution to a 1.5 mL ground-in stopper centrifugal precipitate tube (4).

d) Centrifuge at 8000 × g - 10000 × g for about five minutes (5).

e) 1 mL of the supernatant is used as the sample solution.

Note (3) Add methanol immediately as the determined value becomes higher than usual if it is left in air.

(4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

Comment 3 Instead of the procedures in (4.1) c) - e), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

1) Column: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino or amino propyl chemically bonds.

2) Column bath temperature: 30 °C - 40 °C

3) Eluent: Acetonitrile - methanol (6+1)

4) Flow rate: 1 mL/min

5) Detection unit: An absorptiometric detector, measurement wavelength: 215 nm

b) Calibration curve preparation

1) Inject 10 µL of respective dicyandiamide standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 215 nm, and obtain the peak area or the height.

2) Prepare a curve for the relationship between the concentration and the peak area or the height at wavelength 215 nm of the respective dicyandiamide standard solutions for the calibration curve preparation.

c) Sample measurement

1) Subject 10 µL of sample solution to the same procedure as in b) 1)

2) Obtain dicyandiamide (Dd) content from the calibration curve to calculate the concentration of dicyandiamide (Dd) in the analytical sample.

3) Calculate the dicyandiamide nitrogen (Dd-N) by the following formula.
Dicyandiamide nitrogen (Dd-N) (% (mass fraction)) in an analytical sample

\[ A \times (\frac{MW_1}{MW_2}) \]

\[ = A \times 0.6664 \]

\[ A: \] Dicyandiamide (Dd) (% (mass fraction)) in an analytical sample

\[ MW_1: \] 4 atomic weight of nitrogen (56.027)

\[ MW_2: \] Molecular weight of dicyandiamide (84.080)

**Comment 4**
Recovery testing was conducted using nitrolime (3 samples) and a blended fertilizer containing nitrolime (2 samples), as a result, the recovery of dicyandiamide at concentration level of 6 % (mass fraction) and 0.6 % (mass fraction) was 94.9 % - 105.1 % and 95.6 % - 103.5 % respectively.

Table 1 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction).

**Table 1** Results and analysis results from a collaborative study for the test method validation of dicyandiamide nitrogen

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (^2) (%) (^3)</th>
<th>(s_r) (^4) (%) (^3)</th>
<th>(RSD_r) (^5) (%)</th>
<th>(s_R) (^6) (%) (^3)</th>
<th>(RSD_R) (^7) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrolime 1</td>
<td>9</td>
<td>0.0321</td>
<td>0.0010</td>
<td>3.2</td>
<td>0.0012</td>
<td>3.8</td>
</tr>
<tr>
<td>Nitrolime 2</td>
<td>10</td>
<td>0.159</td>
<td>0.002</td>
<td>1.3</td>
<td>0.006</td>
<td>3.8</td>
</tr>
<tr>
<td>Nitrolime 3</td>
<td>11</td>
<td>0.245</td>
<td>0.002</td>
<td>0.7</td>
<td>0.008</td>
<td>3.3</td>
</tr>
<tr>
<td>Blended fertilizers 1</td>
<td>11</td>
<td>0.124</td>
<td>0.001</td>
<td>0.7</td>
<td>0.002</td>
<td>2.0</td>
</tr>
<tr>
<td>Blended fertilizers 2</td>
<td>11</td>
<td>0.410</td>
<td>0.007</td>
<td>1.6</td>
<td>0.008</td>
<td>1.9</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean (\(n = \) number of laboratories x number of samples (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

**References**


(5) Flow sheet for dicyandiamide nitrogen: The flow sheet for dicyandiamide nitrogen in nitrolime and fertilizers containing nitrolime is shown below:
Reference: HPLC chromatographs of dicyandiamide standard solution and sample solution (nitro lime) for the calibration curve preparation are shown below.

Reference diagram: HPLC chromatogram of dicyandiamide

1) Dicyandiamide standard solution (the equivalents of 100 ng (10 µg/mL, 10 µL) of dicyandiamide)
2) Sample solution (nitro lime)

Measurement conditions for HPLC
Column: Hibar LiChrosorb NH₂ (4.6-mm inner diameter, 25-cm long, 5-µm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
6.1.b High-Performance Liquid Chromatography (2)

(1) **Summary**
This testing method is applicable to fertilizers. In addition, nitrolime is excluded from the scope of application. This testing method is classified as Type B and its symbol is 6.1.b-2017 or Dd-N.b-1.

Add water to an analytical sample to extract dicyandiamide, introduce it to a High-Performance Chromatograph (HPLC) to isolate it with a weak acid ion-exchange column, and then measure at wavelength 190 nm to obtain dicyandiamide nitrogen (Dd-N) in an analytical sample. In addition, the performance of this testing method is shown in **Comment 6**.

Biuret nitrogen (B-N), urea nitrogen (U-N), guanidine urea (Gd-N) and guanylurea nitrogen (GU-N) can be simultaneously quantified by using this method. (Refer to **Comment 5**).

(2) **Reagent, etc.**: Reagents and water are as shown below.
   a) **Water**: Water of A3 specified in JIS K 0557.
   b) **Potassium dihydrogen phosphate**: A JIS Guaranteed Reagent specified in JIS K 9007 or a reagent of equivalent quality.
   c) **Phosphoric acid**: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.
   d) **Dicyandiamide nitrogen standard solution (Dd-N 2 mg/mL)** (1): Put 0.300 g of dicyandiamide [C$_2$H$_4$N$_4$] (2) in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
   e) **Dicyandiamide nitrogen standard solution for the calibration curve preparation (Dd-N 200 g/mL)** (1): Put 10 mL of biuret nitrogen standard solution (B-N 2 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   f) **Dicyandiamide nitrogen standard solution (Dd-N 50 µg/mL - 100 g/mL)** (1): Put 25 mL - 50 mL of dicyandiamide nitrogen standard solution (Dd-N 200 µg/mL) to 100-mL volumetric flasks and add water up to the marked line.
   g) **Dicyandiamide nitrogen standard solution for the calibration curve preparation (Dd-N 1 µg/mL - 50 µg/mL)** (1): At the time of usage, put 1 mL - 50 mL of dicyandiamide nitrogen standard solution (Dd-N 100 µg/mL) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

**Note** (1) This is an example of preparation; prepare an amount as appropriate.

(2) A reagent of no less than 98 % (mass fraction) in purity as dicyandiamide is commercially sold.

**Comment 1** Dicyandiamide is sold by Tokyo Chemical Industry Co., Ltd. In addition, dicyandiamide is commercially sold as dicyanodiamide by FUJIFILM Wako Pure Chemical Co., Ltd. and Kanto Chemical Co., Inc.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.
   a) **High-Performance Liquid Chromatograph (HPLC)**: HPLC specified in JIS K 0124 that satisfies following requirements.
      1) **Column**: A 7.5-mm inner diameter 100-mm long stainless steel column tube filled with weak acid ion-exchange resin.
      2) **Column bath**: A column bath whose temperature can be adjusted to 30 ºC - 45ºC.
      3) **Detection unit**: An absorptiometric detector that can measure at wavelength around 190 nm.
   b) **Magnetic stirrer**:
   c) **High speed centrifugal separator**: A centrifugal separator that can work at 8000 × g - 10000 × g.
Comment 2 A column is sold under the production name Asahipak ES-502C 7C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Powdery test sample
a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution (3) to a 1.5-mL ground-in stopper centrifugal precipitate tube (4).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (5) and use the supernatant as the extract.

Note (3) If there is a possibility that the dicyandiamide nitrogen (Dd-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.
(4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.
(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line (6) and transfer to a 1.5-mL ground-in stopper centrifugal precipitate tube (4).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (5) and use the supernatant as the extract.

Note (6) If there is a possibility that the dicyandiamide nitrogen (Dd-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of solution with water.

Comment 3 Instead of procedures in (4.1.1) c - d) or (4.1.2) c - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph: An example of measurement conditions is shown below. Set up the measurement conditions considering it:
1) Column: A weak acid ion-exchange resin column (7.5-mm inner diameter, 100-mm long, 5-µm - 10-µm particle diameter)
2) Column bath temperature: 40 °C
3) Eluent (1): Dissolve 3.92 g of potassium dihydrogenphosphate and 0.12 g of phosphoric acid in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.
4) Flow rate: 0.6 mL/min
5) **Injection volume**: 10 µL
6) **Detection unit**: An absorpiometric detector, measurement wavelength: 190 nm

**Comment 4** Eluent can be prepared as follows. Dissolve 19.6 g of potassium dihydrogenphosphate and 0.584 g of phosphoric acid with water to make 500 mL and store in a refrigerator. At the time of usage, dilute a predetermined volume of the solution by a factor of 10 and filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.

**b) Calibration curve preparation**
1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 190 nm and obtain the peak height.
2) Prepare a curve for the relationship between the dicyandiamide nitrogen (Dd-N) concentration and the peak height at wavelength 190 nm of the respective standard solutions for the calibration curve preparation.

c) **Sample measurement**
1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain the dicyandiamide nitrogen (Dd-N) content from the peak height using the calibration curve to calculate the dicyandiamide nitrogen (Dd-N) in the analytical sample.

**Comment 5** This testing method enables the simultaneous measurement of biuret nitrogen (B-N), urea nitrogen (U-N), dicyandiamide nitrogen (Dd-N), guanidine urea (Gd-N) and guanylurea nitrogen (GU-N). In that case, see 5.10.a Comment 5.

**Comment 6** Additive recovery testing was conducted using one brand of an acetaldehyde condensed urea fertilizer, a compound fertilizer, a blended fertilizer, a fluid compound fertilizer and a home garden-use mixed fertilizer respectively. As a result, the mean recovery at additive level of 3 % (mass fraction) and 1.5 % (mass fraction) and 0.3 % (mass fraction) were 96.3 % - 96.3 %, 94.5 % - 99.7 % and 88.9 % - 100.6 % respectively. The results of the repeatability tests on different days using a blended fertilizer, a compound fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction).
Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average (^2) ((%)^{3})</th>
<th>Repeatability ((%)^{4})</th>
<th>Intermediate precision ((%)^{5})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blended fertilizer</td>
<td>5</td>
<td>3.03</td>
<td>0.02</td>
<td>0.02</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>5</td>
<td>1.45</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>Home garden-use mixed fertilizer</td>
<td>5</td>
<td>0.145</td>
<td>0.001</td>
<td>0.003</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \((T)\) \times the number of duplicate testing \((2)\))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

Table 2 Results and analysis results from a collaborative study for the test method validation of dicyandiamide nitrogen

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories (^1)</th>
<th>Mean (^2) ((%)^{3})</th>
<th>(s) (^4) ((%)^{3})</th>
<th>(RSD) (^5) ((%))</th>
<th>(s) (^6) ((%)^{3})</th>
<th>(RSD) (^7) ((%))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>10</td>
<td>0.0464</td>
<td>0.0023</td>
<td>5.0</td>
<td>0.0148</td>
<td>31.9</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>12</td>
<td>0.206</td>
<td>0.011</td>
<td>5.1</td>
<td>0.031</td>
<td>15.2</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>12</td>
<td>1.69</td>
<td>0.05</td>
<td>2.8</td>
<td>0.12</td>
<td>7.0</td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>10</td>
<td>2.76</td>
<td>0.07</td>
<td>2.6</td>
<td>0.12</td>
<td>4.4</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean \((n = \text{number of laboratories} \times \text{number of samples} \,(2))\)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(5) **Flow sheet for testing method:** The flow sheet for dicyandiamide nitrogen in fertilizers is shown below:

![Flow sheet for dicyandiamide nitrogen in fertilizers](image)

1.00 g analytical sample (powdery)

200-mL ground-in stopper Erlenmeyer flask

← 100 mL of water

Extraction

Stir to mix, 10 minutes

Stand still

Centrifugal separation

Ground-in stopper centrifugal precipitate tube, 8000 ×g - 10000 ×g, 5 minutes

Sample solution

Supernatant

Measurement

Liquid Chromatograph

Figure 1 Flow sheet for dicyandiamide nitrogen in fertilizers (Extraction procedure (4.1.1) and measurement)

![Flow sheet for dicyandiamide nitrogen in fertilizers](image)

1.00 g analytical sample (fluid)

100-mL volumetric flask

← About 50 mL of water

Extraction

Shake to mix

← Water (up to the marked line)

Centrifugal separation

Ground-in stopper centrifugal precipitate tube, 8000 ×g - 10000 ×g, 5 minutes

Sample solution

Supernatant

Measurement

Liquid Chromatograph

Figure 2 Flow sheet for dicyandiamide nitrogen in fertilizers (Extraction procedure (4.1.2) and measurement)
Reference: Chromatogram of the standard solution for calibration curve preparation of dicyandiamide nitrogen is shown below.

Reference diagram  HPLC chromatogram of the mixture standard solutions (10 mg/L for each) for calibration curve preparation

Peak name
(1) Urea nitrogen  (2) Biuret nitrogen  (3) Dicyandiamide nitrogen  
(4) Guanidine nitrogen  (5) Guanylurea nitrogen

Measurement conditions for HPLC
Column: Asahipak ES-502C 7C (7.5-mm inner diameter, 100-mm long, 9-μm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
6.2 Chlorine
6.2.a Ion Chromatography

(1) Summary
The testing method is applicable to potassium sulfate, potassium bicarbonate, magnesium potassium sulfate, fish cakes powder, fish cakes and compost. This testing method is classified as Type D and its symbol is 6.2.a-2017 or Cl.a-1.

Add water to an analytical sample to extract chloride ions, introduce them to an Ion Chromatograph (IC) to isolate it with an ion exchange column, and then measure with an electric conductivity detector to obtain chlorine (Cl) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent, etc.: Reagents and water are as shown below.


b) 1 mol/L sodium carbonate solution: Solution for ion chromatography.

c) Phthalic acid: A reagent of no less than 98 % (mass fraction) in purity.

d) 6-aminohexanic acid (1): A reagent of no less than 97 % (mass fraction) in purity.

e) Phenylboronic acid: A reagent of no less than 97 % (mass fraction) in purity.

f) Chloride ion standard solution (Cl⁻ 1 mg/mL): A chloride ion standard solution (Cl⁻ 1000 mg/mL) traceable to National Metrology.

g) Chloride ion standard solution (Cl⁻ 100 µg/mL) (2): Put a predetermined amount of chloride ion standard solution (Cl⁻ 1 mg/mL) to a volumetric flask and add water up to the marked line.

h) Chloride ion standard solution (Cl⁻ 5 µg/mL - 50 µg/mL) for the calibration curve preparation (2): Put 5 mL - 50 mL of Chloride ion standard solution (Cl⁻ 100 µg/mL) to a 100-mL volumetric flask and add water up to the marked line.

i) Chloride ion standard solution (Cl⁻ 1 µg/mL - 2 µg/mL) for the calibration curve preparation (2): Put 5 mL - 10 mL of Chloride ion standard solution (Cl⁻ 20 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.

j) Eluent for suppressor method (3): Put 6.4 mL of 1 mol/L sodium carbonate solution to a 1000-mL volumetric flask and add water up to the marked line. Then filter with membrane type filter (pore size: no more than 0.5 μm) made of hydrophilic PTFE (2).

k) Eluent for non-suppressor method (2): Put 0.349 g of phthalic acid, 0.380 g of 6-aminohexanic acid and 0.732 g of phenylboronic acid to a 1000-mL volumetric flask and add about 500 mL of water to dissolve. Add water up to the marked line and filter with membrane type filter (pore size: no more than 0.5 μm) made of hydrophilic PTFE (3).

Note (1) It is also known as “6-amino-n-caproic acid”.
(2) This is an example of preparation; prepare an amount as appropriate.
(3) A solution concentrated by a factor of 10 can be prepared in advance and the solution can be diluted by a factor of 10 at each usage.

Comment 1 Instead of the chloride ion standard solution in (2), a chloride ion standard solution for the calibration curve preparation can be prepared by using a chloride ion standard solution (Cl⁻ 0.1 mg/mL) traceable to National Metrology.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) Magnetic stirrer;

b) Centrifugal separator: A centrifugal separator that can work at 1700 × g.

c) Ion Chromatograph (IC): IC specified in JIS K 0127 that satisfies following requirements.

1) Column: In case of a suppressor method, a 4-mm inner diameter 250-mm long 5-μm particle diameter column tube filled with poly vinyl alcohol porous particles, to which quaternary
ammonium group chemically bonds (4).
In case of non-suppressor method, a 4.6-mm inner diameter 100-mm long column tube filled with hydrophilic polymethacrylate-gel, to which quaternary ammonium group chemically bonds (5).

2) Column bath: A column bath whose temperature can be adjusted to 40 °C.
3) Suppressor: Cation exchange membrane or resin should be used.
4) Detection unit: An electric conductivity detector
d) Membrane filters: Pore size is no more than 0.45 μm, made of hydrophilic PTFE

Note (4) A column is commercially sold under the name Shodex IC SI-52 4E, etc.
(5) A column is commercially sold under the name Shodex IC NI-424, etc.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL- ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at 1700 × g centrifugal force for about five minutes (6) and use the supernatant as the extract.
e) Transfer a predetermined amount of the solution, and dilute exactly by a factor of 20 with water (7).
f) Filter with a membrane filter (pore size: no more than 0.45 μm) to make a sample solution.

Note (6) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.
(7) In case of exceeding the calibration curve, dilute by a factor of more than 20.

Comment 2 Instead of the procedures in (4.1.1) c) and d), it is allowed to filter with Type 3 filter paper and the filtrate can be the extract.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0127 and as shown below. Specific measurement procedures are according to the operation method of an Ion Chromatograph (IC) used in measurement.
a) Measurement conditions for Ion Chromatograph (IC): An example of measurement conditions is shown below. Set up the measurement conditions considering it:

aa) Suppressor method
1) Column: A polyvinyl alcohol porous particles column (4-mm inner diameter 250-mm long 5-μm particle diameter) to which quaternary ammonium group chemically bonds.
2) Column bath temperature: 40 °C
3) Eluent: Prepared in the procedure in (2) j).
4) Flow rate: 0.8 mL/min
5) Injection volume: 20 μL
6) Detection unit: An electric conductivity detector

ab) Non-suppressor method
1) Column: A hydrophilic polymethacrylate-gel column (4.6-mm inner diameter 100 mm long) to which quaternary ammonium group chemically bonds.
2) **Column bath temperature**: 40 °C  
3) **Eluent**: Prepared in the procedure in (2) k).  
4) **Flow rate**: 1.0 mL/min  
5) **Injection volume**: 20 µL  
6) **Detection unit**: An electric conductivity detector

**b) Calibration curve preparation**  
1) Inject 20 µL of respective standard solutions for the calibration curve preparation to an IC and record the chromatogram of electric conductivity to obtain peak area.  
2) Prepare a curve for the relationship between the concentration and the peak area of electric conductivity of respective standard solutions for the calibration curve preparation. Prepare a calibration curve when the sample is measured.

**c) Sample measurement**  
1) Subject 20 µL of sample solution to the same procedure as in b) 1)  
2) Obtain the chloride ion concentration from calibration curve by peak area to calculate the chlorine (Cl) in the analytical sample.

**Comment 3** Additive recovery testing was conducted with a suppressor method using samples that 1.8 % (mass fraction) – 33.4 % (mass fraction) of sodium chloride as chlorine is added to potassium sulfate, magnesium potassium sulfate, potassium bicarbonate, cow dung compost and fish cakes powder. As a result, the mean recovery at the chlorine additive level of 33.4 % (mass fraction), 10 % (mass fraction) - 13.4 % (mass fraction) and 1.8 % (mass fraction) - 9.1 % (mass fraction) are 100.8 %, 98.6 % - 101.1 % and 96.2 % - 103.2 % respectively. With a non-suppressor method, they are 100.2 %, 96.4 % - 97.2 % and 93.3 % - 101.4 %.

The results of the repeatability tests on different days using potassium sulfate, magnesium potassium sulfate, potassium bicarbonate, cow dung compost and fish cakes powder to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.1 % (mass fraction).
Table 1: Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^3$</th>
<th>Repeatability $^4$</th>
<th>Intermediate precision $^5$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T^{1)}$</td>
<td>$\times$</td>
<td>$s_{r}^{5)}$ (%)</td>
<td>$RSD_{r}^{5)}$ (%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>number of</td>
<td>$s_{I(T)}^{6)}$ (%)</td>
<td>$RSD_{I(T)}^{6)}$ (%)</td>
</tr>
<tr>
<td>&lt;Suppressor method&gt;</td>
<td></td>
<td>duplicate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pottasium sulfate</td>
<td>5</td>
<td>9.93</td>
<td>0.01</td>
<td>0.3</td>
</tr>
<tr>
<td>Fish cakes powder</td>
<td>5</td>
<td>6.13</td>
<td>0.03</td>
<td>0.5</td>
</tr>
<tr>
<td>&lt;Non-suppressor method&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pottasium sulfate</td>
<td>5</td>
<td>4.86</td>
<td>0.01</td>
<td>0.2</td>
</tr>
<tr>
<td>Magnesium pottasium sulfate</td>
<td>5</td>
<td>4.89</td>
<td>0.02</td>
<td>0.4</td>
</tr>
<tr>
<td>Pottasium bicarbonate</td>
<td>5</td>
<td>4.85</td>
<td>0.02</td>
<td>0.4</td>
</tr>
<tr>
<td>Cow dung compost</td>
<td>5</td>
<td>13.15</td>
<td>0.04</td>
<td>0.3</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) \times the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

References

5) Flow sheet for testing method: The flow sheet for chlorine in fertilizers is shown below:

![Flow sheet for chlorine in fertilizers](image-url)
Reference: IC chromatographs of sample solutions (magnesium potassium sulfate and fish cakes powder) are shown below.

(A) Chromatograph of magnesium potassium sulfate (Suppressor method)

(B) Chromatograph of magnesium potassium sulfate (Non-suppressor method)

(C) Chromatograph of fish cakes powder (Suppressor method)

(D) Chromatograph of fish cakes powder (Non-suppressor method)

Reference diagram: IC chromatogram of chloride ion (Peak: 1.chloride ion (Cl'))
6.2.b Silver nitrate method

(1) Summary
The test method is applicable to potassium sulfate, potassium bicarbonate and magnesium potassium sulfate. This testing method is classified as Type E and its symbol is 6.2.b-2017 or Cl.b-1.

Add water to an analytical sample to extract chloride ions and titrate (precipitate) with a 0.1 mol/L silver nitrate standard solution to obtain chlorine (Cl) in an analytical sample.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Nitric acid: A JIS Guaranteed (HNO3 60 % (mass fraction)) Reagent specified in JIS K 8541 or a reagent of equivalent quality.
   c) 0.1 mol/L silver nitrate solution:\(^{(1)}\): Weigh 17 g of silver nitrate specified in JIS K 8550 to a 2000-mL beaker, add 1000 mL of water to dissolve and store in a colored bottle.

Standardization: Heat a sodium chloride reference material for volumetric analysis specified in JIS K 8005 at 600 °C ± 25 °C for 1 hour, let it stand to cool in a desiccator, and then transfer about 1.5 g to a weighing dish, and weigh the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line to make the sodium chloride solution\(^{(1)}\). On each day to use a 0.1 mol/L silver nitrate solution, transfer 10 mL of the sodium chloride solution to a 200-mL Erlenmeyer flask, add a few drops of potassium chromate solution (5 g/100 mL) as an indicator, and titrate with a 0.1 mol/L silver nitrate solution until the color of the solution becomes reddish brown. Calculate the factor of a 0.1 mol/L silver nitrate solution by the following formula:

\[
\text{Factor } (f) \text{ of } 0.1 \text{ mol/L silver nitrate solution} = \frac{W_1 \times (A/100) \times (1/58.44) \times (V_1/V_2 \times (1000/V_3)) \times (1/C)}{W_1 \times A/V_3 \times (4/58.44)}
\]

- \(W_1\): Mass (g) of sodium chloride weighed
- \(A\): Purity (% (mass fraction)) of sodium chloride
- \(V_1\): Volume (10 mL) of sodium chloride solution transferred
- \(V_2\): Constant volume (250 mL) of sodium chloride solution
- \(V_3\): Volume (mL) of 0.1 mol/L silver nitrate solution needed for titration
- \(C\): Set concentration (0.1 mol/L) of 0.1 mol/L silver nitrate solution

d) Potassium chromate (5 g/ 100 mL)\(^{(1)}\): Dissolve 5 g of potassium chromate in 100 mL of water.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Magnetic stirrer:
   b) pH test paper: A pH test paper infiltrated with indicator and dried, which can measure the value from pH 1 to pH 11 and a color change chart with the pH interval value 1 is attached.

Comment 1 pH test paper is sold under the name UNIV Test Paper, etc.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL- ground-in stopper Erlenmeyer
flask.

b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.

c) Filter with Type 3 filter paper to make a sample solution.

(4.2) Measurement: Conduct measurement as shown below.

a) Transfer 25 mL of sample solution to a 200-mL tall beaker.

b) Confirm the pH of the solution with a pH-test paper. If the pH is basic, neutralize with nitric acid (1+10).

c) Add a few drops of potassium chromate solution (5 g/100 mL) to the sample solution as an indicator and titrate with a 0.1 mol/L silver nitrate solution until the color of the solution becomes reddish brown.

d) Calculate chlorine (Cl) by the following formula.

\[
\text{Chlorine (\% (mass fraction)) in the analytical sample} = \frac{V_4 \times C \times f \times (35.45)}{W_2 \times (100/1000) \times (V_5/ V_6)}
\]

\[
= \frac{V_4 \times f \times (35.45/25)}{W_2}
\]

\(V_4\): Volume (mL) of the 0.1 mol/L silver nitrate solution required for titration

\(C\): Set concentration (0.1 mol/L) of 0.1 mol/L silver nitrate solution

\(f\): Factor of 0.1 mol/L silver nitrate solution

\(V_5\): Liquid volume of water (100 mL) subjected to extraction in (4.1) b).

\(V_6\): Transferred volume (25 mL) of the sample solution subjected to titration in (4.2) a).

\(W_2\): Mass (g) of the analytical sample

References

(5) **Flow sheet for testing method**: The flow sheet for chlorine in potassium sulfate is shown below:

<table>
<thead>
<tr>
<th>1.00 g analytical sample</th>
<th>200-mL ground-in stopper Erlenmeyer flask</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>← 100 mL of water</td>
</tr>
<tr>
<td>Extraction</td>
<td>Magnetic stirrer, 10 minutes</td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 1** Flow sheet for chlorine in potassium sulfate (Extraction procedure)

<table>
<thead>
<tr>
<th>Sample solution</th>
<th>200-mL tall beaker</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aliquot (25 mL)</td>
<td>pH-test paper</td>
</tr>
<tr>
<td>pH confirmation</td>
<td>← Nitric acid (1+10) [neutralize in case of basic]</td>
</tr>
<tr>
<td></td>
<td>← Several drops of potassium chromate (5 g/100 mL)</td>
</tr>
<tr>
<td>Titration</td>
<td>0.1 mol/L silver nitrate solution</td>
</tr>
<tr>
<td></td>
<td>(until the solution becomes reddish brown)</td>
</tr>
</tbody>
</table>

**Figure 2** Flow sheet for chlorine in potassium sulfate (Measurement procedure)
6.3 Urea nitrogen
6.3.a Urease method

(1) Summary
The test method is applicable to fertilizers containing urea or urea compounds such as acetaldehyde condensed urea, etc. However, in some cases, it is not applicable to the fertilizers that contain such compounds as nitrolime that digests by heating. This testing method is classified as Type D and its symbol is 6.3.a-2017 or U-N.a-1.

Add water to an analytical sample to extract and add urease to a predetermined amount of the extract to hydrolyze urea into ammonium ion. Add sodium hydroxide to alkalize the solution and subject it to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and titrate (neutralize) surplus sulfuric acid using 0.1 mol/L - 0.2 mol/L sodium hydroxide solution and correct separately the titers of urease blank test and urease undigested blank test to obtain urea nitrogen (U-N) in the analytical sample. Or collect isolated ammonia with a boric acid solution and titrate (neutralize) the ammonium ion using 0.25 mol/L sulfuric acid and similarly correct to obtain ammonia nitrogen (A-N) in the analytical sample. In addition, the performance of this testing method is shown in Comment 11.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution \(^{(1)}\): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4 - 5 days. Transfer 5.5 mL - 11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line \(^{(1)}\). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (} f_1) \ = \ (W_1 \times A \times 0.01/97.095) \times (V_1/V_2) \times (1000/V_3) \times (1/C_1)
\]

\( W_1 \): Mass (g) of sulfamic acid sampled
\( A \): Purity (% (mass fraction)) of sulfamic acid
\( V_1 \): Volume (mL) of sulfamic acid solution transferred
\( V_2 \): Constant volume (250 mL) of sulfamic acid solution
\( V_3 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
\( C_1 \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Magnesium oxide: A JIS Guaranteed Reagent specified in JIS K 8432 or a reagent of equivalent quality.

c) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

d) 0.25 mol/L sulfuric acid \(^{(1)}(2)\): Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.
**Standardization**: Transfer a predetermined amount (3) of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, andtitrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

\[
\text{Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid}
\]

\[
= \frac{V_4}{V_5} 
\]

\[
\text{Factor of 0.25 mol/L sulfuric acid (f_2)}
\]

\[
= \frac{(f_1 \times C_1 \times V_4 / V_5)}{(C_2 \times 2)} 
\]

\[
V_4: \text{Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration}
\]

\[
V_5: \text{Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization}
\]

\[
C_1: \text{Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution}
\]

\[
C_2: \text{Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid}
\]

e) **Boric acid solution (40 g/L)**: Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

f) **Urease**: A reagent which completely digests 0.25 g of urea by 0.5 g of urease.

g) **Sodium hydroxide solution (5 g/L)** (1): Dissolve 5 g of sodium hydroxide specified in JIS K 8576 in water to make 1000 mL.

h) **Hydrochloric acid**: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.

i) **Bromothymol blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.

j) **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

k) **Methylene blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

l) **Methyl red–methylene blue mixture solution**: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

m) **Bromocresol green solution (0.5 g/100 mL)**: Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

n) **Methyl red–bromocresol green mixture solution**: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

**Note** (1) This is an example of preparation; prepare an amount as appropriate.  
(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).  
(3) 5 mL -10 mL  
(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.
Comment 2 0.25 mol/L sulfuric acid in (2) d) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.

Comment 3 Refined products of urease made from sword beans are commercially sold. The degree of activation may degrade even if stored in a refrigerator. Therefore, it is recommended to test similarly and confirm activation by using urea specified in JIS K 8731 in advance.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Extractor: Rotary shaker or magnetic stirrer as described below
   aa) Rotary shaker: A rotary shaker that can rotate a 500-mL volumetric flask upside down at 30 - 40 revolutions/min.
   ab) Magnetic stirrer:
   b) Steam distillation apparatus
   c) Distillation flask: A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.
   d) Water bath: Water bath that can be adjusted to 40 °C - 45 °C.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Rotary shaker
   a) Weigh 5.00 g of an analytical sample, and put it in a 500-mL volumetric flask.
   b) Add about 400 mL of water, and shake to mix at 30 - 40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

Comment 4 In the procedure in (4.1.1) a), it is also allowed to weigh 2.50 g of the analytical sample and transfer to a 250-mL volumetric flask.

Comment 5 The procedure in (4.1.1) is the same as the procedure in (4.1.1.1) in 6.3.c.

(4.1.2) Magnetic stirrer:
   a) Weigh 1.00 g of an analytical sample, and put it into a 200- mL ground-in stopper Erlenmeyer flask.
   b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
   c) Filter with Type 3 filter paper to make the extract.

Comment 6 The procedure in (4.1.2) is the same as the procedure in (4.1.1) in 6.3.c.

(4.2) Hydrolysis by urease: Conduct hydrolysis as shown below.
   a) Transfer a predetermined amount of the extract (the equivalents of 10 mg or more as U- N, the equivalents of 10 mg - 100 mg as N) to a 300-mL distillation flask.
   b) Add water to the solution to make about 50 mL.
   c) Add a few drops of methyl red solution (0.1 g/100 mL), and add a sodium hydroxide solution (5 g/L) or hydrochloric acid (1+200) until the color of the solution becomes light yellowish red (6).
   d) Add a sufficient amount of urease to digest urea in the extract (7) (8), seal tightly and heat in the water bath at 40 °C - 45 °C.
   e) Let it stand to cool to make a sample solution.
   f) As an extract blank test, conduct the procedures in a) (9) using another distillation flask to prepare the undigested test solution.
   g) As a urease blank test, conduct the procedures in b), d) and e) (8) (10) using another distillation
flask to prepare the blank test solution.

**Note**  (6) pH 5.6 - pH 5.8
(7) An example of the additive volume of urease is shown in Comment 10.
(8) Wash urease off with a small amount of water if it adheres to the surface of a vessel.
(9) Aliquot the same amount of extract as the prepared sample solution.
(10) Add the same amount of urease as the preparation of the sample solution used.

**4.3 Distillation:** Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.

a) Transfer a predetermined amount (11) of 0.25 mol/L sulfuric acid to an acceptor (12), add a few drops of methyl red - methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount (11) of boric acid solution (40 g/L) to an acceptor (12), add a few drops of methyl red - bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.

b) Add 2 g - 3 g of magnesium oxide to a distillation flask containing the sample solution (13), and connect this distillation flask to the steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

f) Subject the undigested test solution to the same procedure as in a) - e) to obtain the distillate from the undigested test solution.

g) Subject the blank test solution to the same procedure as in a) - e) to obtain the distillate from the blank test solution.

**Comment 7** Immediately conduct the procedure in (4.3) b) not to emit ammonia gas in a vessel.

**4.4 Measurement:** Conduct measurement as shown below.

**4.4.1** When 0.25 mol/L sulfuric acid is used in (4.3):

a) Titrante the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4).

b) Subject the distillate from the undigested test solution to the same procedure as in a) to titrate.

c) Subject the distillate from the blank test solution to the same procedure as in a) to titrate.

d) Calculate the urea nitrogen (U-N) in an analytical sample by the following formula:

\[ \text{Urea nitrogen (U-N) (% (mass fraction)) in the analytical sample} = \left( \frac{(B \times V_b - V_7) - (B \times V_b - V_5) - (B \times V_b - V_9) \times C_1 \times f_1 \times (V_{10}/V_{11}) \times (14.007/W_2) \times (100/1000)}{(B \times V_b - V_6 - V_7 - V_8 - V_9)} \times C_1 \times f_1 \times (V_{10}/V_{11}) \times (1.4007/W_2) \right) \]

\( B \): Volume of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

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$V_6$: Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.3) a

$V_7$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration in (4.4.1) a

$V_8$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration in (4.4.1) b

$V_9$: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration in (4.4.1) c

$C_1$: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

$f_1$: Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

$V_{10}$: Predetermined volume (mL) of the extract in (4.1.1) c) or additive volume of water in (4.1.2) b

$V_{11}$: Transferred amount (mL) of the extract subjected to hydrolysis in (4.2) a

$W_2$: Mass (g) of the analytical sample in (4.1.1) a) or (4.1.2) a)

(4.4.2) When a boric acid solution (40 g/L) is used in (4.3):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red.
b) Subject the distillate from the undigested test solution to the same procedure as in a) to titrate.
c) Subject the distillate from the blank test solution to the same procedure as in a) to titrate.
d) Calculate the urea nitrogen (U-N) in an analytical sample by the following formula:

\[
\text{Urea nitrogen (U-N) (\% (mass fraction)) in the analytical sample} = \left(\frac{V_{12} - V_{13} - V_{14}}{V_{12} - V_{13} - V_{14}}\right) \times \frac{0.25 \times 2 \times f_2 \times (14.007/W_2)}{2.8014/W_2} \times (100/1000)
\]

$V_{12}$: Volume (mL) of 0.25 mol/L sulfuric acid needed for titration in (4.4.2) a

$V_{13}$: Volume (mL) of 0.25 mol/L sulfuric acid needed for titration in (4.4.2) b

$V_{14}$: Volume (mL) of 0.25 mol/L sulfuric acid needed for titration in (4.4.2) c

$C_2$: Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

$f_2$: Factor of 0.25 mol/L sulfuric acid

$V_{10}$: Predetermined volume (mL) of the extract in (4.1.1) c) or additive volume of water (mL) in (4.1.2) b

$V_{11}$: Transferred amount (mL) of the extract subjected to hydrolysis in (4.2) a

$W_2$: Mass (g) of the analytical sample in (4.1.1) a) or (4.1.2) a)

Note (14) The endpoint is reached when the color changes from green to light red.

Comment 8 If it is hard to confirm the endpoint due to the carbon dioxide resulting from carbonate in the extract when magnesium oxide is used, it is recommended to boil the extract for 1-2 minute(s) after distilling and cool, and then titrate.

Comment 9 The titration procedures in (2) a) Standardization, (2) d) Standardization and (4.4) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

Comment 10 An example of the additive volume and titter of urease is shown below.

If the urea content can be estimated, calculate the volume of urea in the transferred volume of the extract in (4.2) a) by the following formula.

Estimated urea content (mg) in the transferred volume of the extract
\[
= \left(\frac{D_1}{100}\right) \times W_2 \times \left(\frac{V_{11}}{V_{10}}\right)
\]
Estimated urea content (mg) in the transferred volume of the extract
\[ = \frac{D_2}{100} \times \left( \frac{60.056}{(14.007 \times 2)} \right) \times W_2 \times \left( \frac{V_{11}}{V_{10}} \right) \]

\[ = \frac{D_2}{100} \times 2.1438 \times W_2 \times \left( \frac{V_{11}}{V_{10}} \right) \]

If the urea content cannot be estimated, subtract ammoniacal nitrogen and nitrate nitrogen from the total nitrogen of the permissible content or the display component content of the urea nitrogen in a urea compound to obtain nitrogen content. The obtained nitrogen content is calculated as around the maximum content of urea nitrogen (U-N). In this case, calculate the volume of urea in the transferred volume of the extract in (4.1.1) or (4.2) a) after the procedures in (4.1.1) by the following formula.

Comment 11 Additive recovery testing was conducted to evaluate trueness using samples prepared by adding urea to sulfur, a compound fertilizer, isobutyaldehyde condensed urea, a mixed fertilizer containing formaldehyde processed urea (one sample for each). As a result, the average rate of recovery at additive level of 1.58 % (mass fraction) - 39.96 % (mass fraction) are 98.7 % to 103.7 %

The results of the repeatability tests on different days using urea and a compound fertilizer containing UF (one sample for each), to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.4 % (mass fraction).
### Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$</th>
<th>Repeatability $^3$</th>
<th>Intermediate precision $^7$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T$ $^1$</td>
<td>Average (%)</td>
<td>$s_r$ (%)</td>
<td>RSD$^5$ (%)</td>
</tr>
<tr>
<td>Urea</td>
<td>5</td>
<td>43.17</td>
<td>0.36</td>
<td>0.8</td>
</tr>
<tr>
<td>Compound fertilizer containing UF</td>
<td>5</td>
<td>2.39</td>
<td>0.07</td>
<td>2.9</td>
</tr>
</tbody>
</table>

1) Gross average (the number of test days $(5)$ × duplicate analysis)
2) Average (the number of test days $(T)$ × the number of duplicate testing $(2)$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

### References


### Flow sheet for urea nitrogen testing method:

The flow sheets for urea nitrogen test method in fertilizers are shown below:

- 5.00 g Analytical sample
- Weigh into a 500-mL volumetric flask
- Shaking to mix
- Rotary shaker (30 - 40 revolutions/min), 1 hour
- Water (up to the marked line)
- Type 3 filter paper
- Filtration
- Extract

Figure 1-1  Flow sheet for urea nitrogen
(Extraction procedure (4.1.1))
1.00 g Analytical sample

→ 100 mL of water

Shaking to mix

Magnetic stirrer, 10 minutes

Filtration

Type 3 filter paper

Extract

Figure 1-1 Flow sheet for urea nitrogen
(Extraction procedure (4.1.2))

Extract

Aliquot
(predetermined volume)

300-mL digestion flask

→ Water (Until the solution reaches about 50 mL)
→ A few drops of methyl red solution (0.1 g/100 mL)
→ Hydrochloric acid (1+200) or sodium hydroxide solution (5 g/L)
    (until the color of the solution becomes light yellowish red)
→ Urease (a sufficient amount of urease to hydrolyze transferred urea)

40 °C - 50 °C, 1 hour

Heating

Standing to cool

Sample solution

→ 2 g - 3 g of magnesium oxide

Steam distillation apparatus

Receiver: 200-mL - 300-mL Erlenmeyer flask or 200-mL - 300-mL beaker. A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution

Steam distillation

Distillation rate: 5 mL/min - 7 mL/min

Stop distilling

120 mL - 160 mL distillate

→ Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)

Titration

0.1 mol/L - 0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or 0.25 mol/L sulfuric acid (until the solution becomes light red)

Figure 2  Flow sheet for urea nitrogen
(Hydrolysis by urease, distillation and measurement procedure)
6.3.b High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 6.3.b-2017 or U-N.b-1.

Add water to an analytical sample to extract urea, introduce it to a High-Performance Chromatograph (HPLC) to isolate it with a weak acid ion-exchange column, and then measure at wavelength 190 nm to obtain urea nitrogen (U-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

Biuret nitrogen (B-N), dicyandiamide nitrogen (Dd-N), guanidine urea (Gd-N) and guanylurea nitrogen (GU-N) can be simultaneously quantified by using this method. (Refer to Comment 4).

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Potassium dihydrogen phosphate: A JIS Guaranteed Reagent specified in JIS K 9007 or a reagent of equivalent quality.
   c) Phosphoric acid: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.
   d) Urea nitrogen standard solution (U-N 2 mg/mL) (1): Put 0.429 g of urea specified in JIS K 8731 in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
   e) Urea nitrogen standard solution for the calibration curve preparation (U-N 200 µg/mL): Put 10 mL of urea nitrogen standard solution (U-N 2 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   f) Urea nitrogen standard solution (U-N 50 µg/mL - 100 µg/mL): Put 25 mL - 50 mL of urea nitrogen standard solution (U-N 200 µg/mL) to 100-mL volumetric flasks and add water up to the marked line.
   g) Urea nitrogen standard solution for the calibration curve preparation (U-N 1 µg/mL - 50 µg/mL): At the time of usage, put 1 mL - 50 mL of urea nitrogen standard solution (U-N 0.1 mg/mL) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 7.5-mm inner diameter 100-mm long stainless steel column tube filled with weak acid ion-exchange resin.
      2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45ºC.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 190 nm.
   b) Magnetic stirrer:
   c) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.

Comment 1 A column is sold under the production name Asahipak ES-502C 7C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Powdery test sample
   a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL-ground-in stopper Erlenmeyer
flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution (2) to a 1.5-mL ground-in stopper centrifugal precipitate tube (3).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (4) and use the supernatant as the extract.

**Note** (2) If there is a possibility that the urea nitrogen (U-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

(3) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(4) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

(4.1.2) **Fluid test sample**
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line (5) and transfer to a 1.5-mL ground-in stopper centrifugal precipitate tube (3).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (4) and use the supernatant as the extract.

**Note** (5) If there is a possibility that the urea nitrogen (U-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

**Comment 2** Instead of procedures in (4.1.1) c) - d) or (4.1.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions is shown below. Set up the measurement conditions considering it:

1) **Column**: A weak acid ion-exchange resin column (7.5-mm inner diameter, 100-mm long, 5-µm - 10-µm particle diameter)
2) **Column bath temperature**: 40 °C
3) **Eluent** (1): Dissolve 3.92 g of potassium dihydrogenphosphate and 0.12 g of phosphoric acid in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.
4) **Flow rate**: 0.6 mL/min
5) **Injection volume**: 10 µL
6) **Detection unit**: An absorptiometric detector, measurement wavelength: 190 nm

**Comment 3** Eluent can be prepared as follows. Dissolve 19.6 g of potassium dihydrogenphosphate and 0.584 g of phosphoric acid with water to make 500 mL and store in a refrigerator. At the time of usage, dilute a predetermined volume of the
solution by a factor of 10 and filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.

b) **Calibration curve preparation**
1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 190 nm and obtain the peak height.
2) Prepare a curve for the relationship between the urea nitrogen (U-N) concentration and the peak height at wavelength 190 nm of the respective standard solutions for the calibration curve preparation.

c) **Sample measurement**
1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain the urea nitrogen (U-N) content from the peak height using the calibration curve to calculate the urea nitrogen (U-N) in the analytical sample.

Comment 4 This testing method enables the simultaneous measurement of biuret nitrogen (B-N), urea nitrogen (U-N), dicyandiamide nitrogen (Dd-N), guanidine urea (Gd-N) and guanylurea nitrogen (GU-N). In that case, see 5.10.a Comment 5.

When Asahipak ES-502C 7C is used as a column for HPLC to analyze urea nitrogen (U-N) in a mixed fertilizer containing formaldehyde processed urea (UF), the peak of the urea nitrogen (U-N) and the peak of the impurity components originating from UF are not separated, so that urea nitrogen (U-N) cannot be measured. In this case, the peak of the urea nitrogen (U-N) and the peak of the impurity components originating from UF are separated by using PRP-X200 as a column for HPLC instead to be able to analyze the urea nitrogen(U-N) in the compound fertilizer containing UF. However, when the PRP-X200 column is used as a column for HPLC, the simultaneous measurement of urea nitrogen (U-N) and biuret nitrogen (B-N), etc. is impossible.

Comment 5 Additive recovery testing was conducted using one brand of an acetaldehyde condensed urea fertilizer, a compound fertilizer, a blended fertilizer, a fluid compound fertilizer and a home garden-use mixed fertilizer respectively. As a result, the mean recovery at additive level of 6 % (mass fraction) and 3 % (mass fraction) and 0.6 % (mass fraction) were 98.3 % - 102.9 %, 98.9 % - 105.2 % and 92.3 % - 99.9 % respectively.

The results of the repeatability tests on different days using a blended fertilizer, a compound fertilizer and a home garden-use mixed fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of this testing method is about 0.03 % (mass fraction).

Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average(^2) (\bar{x}) (%)(^3)</th>
<th>Repeatability (s_r) (%) (^4)</th>
<th>(RSD_r) (%) (^5)</th>
<th>Intermediate precision (s_{I(T)}) (%) (^6)</th>
<th>(RSD_{I(T)}) (%) (^7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blended fertilizer</td>
<td>5</td>
<td>6.24 (\pm) 0.03 (%) (^3)</td>
<td>0.5</td>
<td>0.05 (\pm) 0.8 (%) (^5)</td>
<td>0.005 (\pm) 0.9 (%) (^7)</td>
<td></td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>5</td>
<td>3.01 (\pm) 0.03 (%) (^3)</td>
<td>0.7</td>
<td>0.04 (\pm) 1.4 (%) (^5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Home garden-use mixed fertilizer</td>
<td>5</td>
<td>0.315 (\pm) 0.003 (%) (^3)</td>
<td>0.9</td>
<td></td>
<td>0.005 (\pm) 0.9 (%) (^7)</td>
<td></td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days \(T\) \times the number of duplicate testing \(2\))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation

Table 2  Results and statistical analysis results from a collaborative study for the test method validation of urea nitrogen

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories (^1)</th>
<th>Mean(^2) (%) (^3)</th>
<th>(s_r) (%) (^4)</th>
<th>(RSD_r) (%) (^5)</th>
<th>(s_{R} ) (%) (^6)</th>
<th>(RSD_{R} ) (%) (^7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>8</td>
<td>0.296 (\pm) 0.011 (%) (^3)</td>
<td>3.6 (\pm) 0.012 (%) (^5)</td>
<td>4.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>10</td>
<td>0.589 (\pm) 0.015 (%) (^3)</td>
<td>2.6 (\pm) 0.024 (%) (^5)</td>
<td>4.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>10</td>
<td>3.08 (\pm) 0.04 (%) (^3)</td>
<td>1.1 (\pm) 0.06 (%) (^5)</td>
<td>2.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>10</td>
<td>6.03 (\pm) 0.11 (%) (^3)</td>
<td>1.7 (\pm) 0.20 (%) (^5)</td>
<td>3.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Compound fertilizer 5</td>
<td>10</td>
<td>46.5 (\pm) 0.6 (%) (^3)</td>
<td>1.4 (\pm) 1.3 (%) (^5)</td>
<td>2.8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean \(n = \text{number of laboratories} \times \text{number of samples (2)}\)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References

(5) **Flow sheet for testing method:** The flow sheet for urea nitrogen in fertilizers is shown below:

![Flow sheet for urea nitrogen in fertilizers](image)

**Figure 1** Flow sheet for urea nitrogen in fertilizers
(Extraction procedure (4.1.1) and measurement)

![Flow sheet for urea nitrogen in fertilizers](image)

**Figure 2** Flow sheet for urea nitrogen in fertilizers
(Extraction procedure (4.1.2) and measurement)
Reference: Chromatogram of the standard solution for calibration curve preparation of urea nitrogen is shown below.

Reference diagram 1  HPLC chromatogram of the mixture standard solutions (10 mg/L for each) for calibration curve preparation

Peak name
(1) Urea nitrogen  (2) Biuret nitrogen  (3) Dicyandiamide nitrogen
(4) Guanidine nitrogen  (5) Guanylurea nitrogen

Measurement conditions for HPLC
Column: Asahipak ES-502C 7C (7.5-mm inner diameter, 100-mm long, 9-μm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
Reference diagram 2  
HPLC chromatogram of the urea nitrogen standard solutions (20 mg/L) and a formaldehyde processed urea fertilizer for calibration curve preparation

Peak name
(1) Urea nitrogen  (2) Biuret nitrogen

Measurement conditions for HPLC
Column:  PRP-X200 (4.1-mm inner diameter, 150-mm long, 10-µm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
6.3.c  *p*-dimethylaminobenzaldehyde absorptiometry

(1) **Summary**

The testing method is applicable to fertilizers containing urea nitrogen. However, isobutyraldehyde condensed urea fertilizers, formaldehyde processed urea fertilizers, nitrolime, sludge fertilizers, etc., and special fertilizers are excluded. This testing method is classified as Type D and its symbol is 6.3.c-2018 or U-N.c-1.

Add water to an analytical sample to extract urea and measure the coloration formed by the reaction with dimethylaminobenzaldehyde by absorbance to obtain urea nitrogen (U-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) **Reagent, etc.**: Reagents and water are as shown below.

a) **Water**: Water of A3 specified in JIS K 0557.

b) **Coloring reagent solution** \(^{(1)}\): Dissolve 20 g of *p*-dimethylaminobenzaldehyde specified in JIS K 8101 in 1000 mL of ethanol (99.5) specified in JIS K 8101 and 100 mL of hydrochloric acid specified in JIS K 8180 and leave at rest overnight \(^{(2)}\).

c) **Urea nitrogen standard solution (U-N 2 mg/mL)** \(^{(1)}\): Put 0.429 g of urea specified in JIS K 8731 in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.

d) **Urea nitrogen standard solution for the calibration curve preparation (U-N 200 µg/mL)** \(^{(1)}\): Put 10 mL of urea nitrogen standard solution (U-N 2 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.

**Note** (1) This is an example of preparation; prepare an amount as appropriate.

(2) Store in an amber bottle.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.

a) **Spectrophotometer**: A spectrophotometer specified in JIS K 0115

b) **Magnetic stirrer**:

(4) **Test procedures**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it into a 200-mL Erlenmeyer flask.

b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.

c) After allowing to stand still, filter with Type 3 filter paper to make a sample solution.

**Comment 1** The procedure in (4.1) is the same as the procedure in (4.1.2) of 6.3.a.

**Comment 2** When the determination is affected by the coloring of the sample solution of (4.1) c), add about 0.5 g of active carbon and filter with Type 3 filter paper to make a sample solution from which the coloring disappears.

(4.2) **Coloring**: Conduct coloring as shown below.

a) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 mg - 5 mg as U-N) to a 50-mL volumetric flask.

b) Add 20 mL of coloring reagent solution, and further add water up to the marked line, and then leave at rest for about 30 minutes.

(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0115 and as shown below. Specific measurement procedures are according to the operation method of the spectrophotometer used for the measurement.
a) **Measurement conditions of spectrophotometer**: Set up the measurement conditions of spectrophotometer considering the following.
Detection wavelength: 450 nm

b) **Calibration curve preparation**
1) Transfer 2.5 mL - 25 mL of urea nitrogen standard solution (U-N 200 µg/mL) to 50-mL volumetric flasks step-by-step.
2) Conduct the same procedures as (4.2) b) to make U-N 0.5 mg/50 mL - 5 mg/50 mL of the urea nitrogen standard solutions for the calibration curve preparation.
3) Conduct the same procedures as 2) for another 50-mL volumetric flask to make the blank test solution for the calibration curve preparation.
4) Measure absorbance at a wavelength of 450 nm of the urea nitrogen standard solutions for the calibration curve preparation using the blank test solution for the calibration curve preparation as the control.
5) Prepare a curve for the relationship between the urea nitrogen (U-N) concentration and the absorbance of the urea nitrogen standard solutions for the calibration curve preparation.

c) **Sample measurement**
1) Regarding the solution in (4.2) b), measure absorbance by the same procedure as b) 4).
2) Obtain the urea nitrogen (U-N) content from the calibration curve to calculate the urea nitrogen (U-N) in the analytical sample.

**Comment 3** Additive recovery testing was conducted using a compound fertilizer, crustacea grade fertilizer powdery and a preparation sample. As a result, the mean recovery at additive level of 20 % (mass fraction), 10 % (mass fraction) and 3 % (mass fraction) were 100.0 % - 102.4 %, 100.5 % - 102.0 % and 98.0 % - 103.3 % as urea nitrogen (U-N) respectively.

The results of the repeatability tests on different days using urea, a designated blended fertilizer and a compound fertilizer to evaluate precision were analyzed by the one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.2 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Average&lt;sup&gt;2&lt;/sup&gt;</td>
<td>s&lt;sub&gt;r&lt;/sub&gt;&lt;sup&gt;4&lt;/sup&gt;</td>
</tr>
<tr>
<td>Urea</td>
<td>7</td>
<td>45.9</td>
<td>0.89</td>
</tr>
<tr>
<td>Designated blended fertilizer</td>
<td>7</td>
<td>7.45</td>
<td>0.16</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>11.12</td>
<td>0.02</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days T) x the number of duplicate testing (2)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
References

(5) Flow sheet for testing method: The flow sheet for urea nitrogen in fertilizers is shown below:

| 1.00 g Analytical sample (powdery) | 200-mL Erlenmeyer flask |
| ← 100 mL of water | Extract Stir for 10 minutes |
| Stand still | Filtration Type 3 filter paper |
| Sample solution |

Figure 1 Flow sheet for urea nitrogen in fertilizers (Extraction procedure)

| Sample solution |
| Sample solution |
| ← 20 mL of coloring reagent solution | ← Water (up to the marked line) |
| Leave at rest About 30 minutes | Measurement Spectrophotometer (450 nm) |

Figure 2 Flow sheet for urea nitrogen in fertilizers (Measurement procedure)
6.4 Guanidine nitrogen
6.4.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 6.4.a-2017 or Gd-N.a-1.

Add water to an analytical sample to extract guanidine, introduce it to a High-Performance Chromatograph (HPLC) to isolate it with a weak acid ion-exchange column, and then measure at wavelength 190 nm to obtain guanidine nitrogen (Gd-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

Biuret nitrogen (B-N), dicyandiamide nitrogen (Dd-N), urea nitrogen(U-N) and guanylurea nitrogen (GU-N) can be simultaneously quantified by using this method. (Refer to Comment 5).

(2) Reagent, etc.: Reagents and water are as shown below.


b) Potassium dihydrogen phosphate: A JIS Guaranteed Reagent specified in JIS K 9007 or a reagent of equivalent quality.

c) Phosphoric acid: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.

d) Guanidine nitrogen standard solution (Gd-N 2 mg/mL) \(^{(1)}\): Put 0.515 g of guanidine sulfate \([\text{C}_2\text{H}_6\text{N}_6\cdot\text{H}_2\text{SO}_4]\) \(^{(2)}\) in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.

e) Guanidine nitrogen standard solution for the calibration curve preparation (Gd-N 200 µg/mL) \(^{(1)}\): Put 10 mL of guanidine nitrogen standard solution (Gd-N 2 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.

f) Guanidine nitrogen standard solution (Gd-N 50 µg/mL - 100 µg/mL) \(^{(1)}\): Put 25 mL - 50 mL of guanidine nitrogen standard solution (Gd-N 200 µg/mL) to 100-mL volumetric flasks and add water up to the marked line.

Note \((1)\) This is an example of preparation; prepare an amount as appropriate.

\((2)\) A reagent of no less than 98 % (mass fraction) in purity as guanidine sulfate is commercially sold.

Comment 1 Guanidine sulfate is commercially sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Tokyo Chemical Industry Co., Ltd.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.

1) Column: A 7.5-mm inner diameter 100-mm long stainless steel column tube filled with weak acid ion-exchange resin.

2) Column bath: A column bath whose temperature can be adjusted to 30 °C - 45°C.

3) Detection unit: An absorptiometric detector that can measure at wavelength around 190 nm.

b) Magnetic stirrer:

c) High speed centrifugal separator: A centrifugal separator that can work at 8000 \(\times\) g - 10000 \(\times\) g.
Comment 2 A column is sold under the production name Asahipak ES-502C 7C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.

(4.1.1) Powdery test sample
a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution\(^{(3)}\) to a 1.5-mL ground-in stopper centrifugal precipitate tube\(^{(4)}\).
d) Centrifuge it at 8000 \(\times\) \(g\) - 10000 \(\times\) \(g\) centrifugal force for about five minutes\(^{(5)}\) and use the supernatant as the extract.

Note (3) If there is a possibility that the guanidine nitrogen (Gd-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

(4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 \(\times\) \(g\) - 10000 \(\times\) \(g\) centrifugal force.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water\(^{(6)}\), and shake to mix.
c) Add water up to the marked line and transfer it to a 1.5-mL ground-in stopper centrifugal precipitate tube\(^{(4)}\).
d) Centrifuge it at 8000 \(\times\) \(g\) - 10000 \(\times\) \(g\) centrifugal force for about five minutes\(^{(5)}\) and use the supernatant as the extract.

Note (6) If there is a possibility that the guanidine nitrogen (Gd-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

Comment 3 Instead of procedures in (4.1.1) c) - d) or (4.1.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5 µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions is shown below. Set up the measurement conditions considering it:

1) Column: A weak acid ion-exchange resin column (7.5-mm inner diameter, 100-mm long, 5-µm - 10-µm particle diameter)
2) Column bath temperature: 40 °C
3) Eluent\(^{(1)}\): Dissolve 3.92 g of potassium dihydrogenphosphate and 0.12 g of phosphoric acid in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than 0.5 µm) made of hydrophilic PTFE.
4) **Flow rate**: 0.6 mL/min
5) **Injection volume**: 10 µL
6) **Detection unit**: An absorptiometric detector, measurement wavelength: 190 nm

**Comment 4** Eluent can be prepared as follows. Dissolve 19.6 g of potassium dihydrogenphosphate and 0.584 g of phosphoric acid with water to make 500 mL and store in a refrigerator. At the time of usage, dilute a predetermined volume of the solution by a factor of 10 and filter with a membrane filter (aperture diameter: no more than 0.5 µm) made of hydrophilic PTFE.

b) **Calibration curve preparation**
1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 190 nm and obtain the peak height.
2) Prepare a curve for the relationship between the guanidine nitrogen (Gd-N) concentration and the peak height at wavelength 190 nm of the respective standard solutions for the calibration curve preparation.

c) **Sample measurement**
1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain the guanidine nitrogen (Gd-N) content from the peak height using the calibration curve to calculate the guanidine nitrogen (Gd-N) in the analytical sample.

**Comment 5** This testing method enables the simultaneous measurement of biuret nitrogen (B-N), urea nitrogen (U-N), dicyandiamide nitrogen (Dd-N), guanidine urea (Gd-N) and guanylurea nitrogen (GU-N). In that case, see 5.10.a **Comment 5**.

**Comment 6** Additive recovery testing was conducted using a preparation sample for a guanylurea fertilizer (one brand). As a result, the mean recovery at additive level of 3.71 % (mass fraction), 1.85 % (mass fraction) and 0.371 % (mass fraction) were 91.2 %, 94.0 % and 100.0 % respectively.

The results of the repeatability tests on different days using a guanylurea fertilizer to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability T</th>
<th>Average (%)</th>
<th>Repeatability s (%)</th>
<th>Intermediate precision RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guanylurea fertilizer</td>
<td>5</td>
<td>1.81</td>
<td>0.01</td>
<td>0.8</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days (T) * the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
Table 2  Results and analysis results from a collaborative study for the test method validation of guanidine nitrogen

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^2$ (%%)</th>
<th>$s_r$ $^4$ (%%)</th>
<th>$RSD_r$ $^5$ (%)</th>
<th>$s_R$ $^6$ (%%)</th>
<th>$RSD_R$ $^7$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>12</td>
<td>4.91</td>
<td>0.18</td>
<td>3.7</td>
<td>0.29</td>
<td>5.8</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>12</td>
<td>3.94</td>
<td>0.16</td>
<td>4.2</td>
<td>0.27</td>
<td>6.8</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>11</td>
<td>3.03</td>
<td>0.06</td>
<td>2.0</td>
<td>0.12</td>
<td>4.0</td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>11</td>
<td>2.05</td>
<td>0.05</td>
<td>2.6</td>
<td>0.09</td>
<td>4.2</td>
</tr>
<tr>
<td>Guanylurea fertilizer</td>
<td>11</td>
<td>5.13</td>
<td>0.21</td>
<td>4.0</td>
<td>0.19</td>
<td>3.6</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean ($n = number of laboratories \times number of samples (2)$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References


(5) **Flow sheet for testing method**: The flow sheet for guanidine nitrogen in fertilizers is shown below:

![Flow sheet for guanidine nitrogen in fertilizers](image)

Figure 1  Flow sheet for guanidine nitrogen in fertilizers  
(Extraction procedure (4.1.1) and measurement)

![Flow sheet for guanidine nitrogen in fertilizers](image)

Figure 2  Flow sheet for guanidine nitrogen in fertilizers  
(Extraction procedure (4.1.2) and measurement)
**Reference**: Chromatogram of the standard solution for calibration curve preparation of guanidine nitrogen is shown below.

Reference diagram  HPLC chromatogram of the mixture standard solutions (10 mg/L for each)

Peak name
1. Urea nitrogen  
2. Biuret nitrogen  
3. Dicyandiamide nitrogen  
4. Guanidine nitrogen  
5. Guanylurea nitrogen

Measurement conditions for HPLC
Column: Asahipak ES-502C 7C (7.5-mm inner diameter, 100-mm long, 9-μm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
6.5 Cold buffer solution soluble nitrogen (water-soluble nitrogen)

6.5.a Cold buffer solution method

(1) Summary
The test method is applicable to formaldehyde processed urea fertilizers. This testing method is classified as Type A (Def-M) and its symbol is 6.5.a-2017 or Buf(C)-N.a-1.

Add a phosphate solution (cold buffer solution) to an analytical sample to extract, add copper (II) sulfate pentahydrate, sulfuric acid and potassium sulfate, pretreat by the Kjeldahl method to change cold buffer solution soluble nitrogen to ammonium ion and add a sodium hydroxide solution to subject to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain cold buffer solution soluble nitrogen (water-soluble nitrogen) in the analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain cold buffer solution soluble nitrogen (water-soluble nitrogen) in the analytical sample.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL - 11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (f₁)} = \left( \frac{W₁ \times A \times 0.01/97.095}{V₁/V₂} \times \left( \frac{1000}{V₃} \right) \times \frac{1}{C₁} \right)
\]

- \(W₁\): Mass (g) of sulfamic acid sampled
- \(A\): Purity (% (mass fraction)) of sulfamic acid
- \(V₁\): Volume (mL) of sulfamic acid solution transferred
- \(V₂\): Constant volume (250 mL) of sulfamic acid solution
- \(V₃\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- \(C₁\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid (1)(2): Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

Standardization: Transfer a predetermined amount (3) of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L -0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium
hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\[ V_4 / V_5 \]  \quad ........ (1)

Factor of 0.25 mol/L sulfuric acid (f_2)

\[ f_2 = (f_1 \times C_1 \times V_4 / V_5) / (C_2 \times 2) \]  \quad ........ (2)

\[ V_4: \text{ Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration} \]
\[ V_5: \text{ Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization} \]
\[ C_1: \text{ Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution} \]
\[ C_2: \text{ Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid} \]

d) **Boric acid solution (40 g/L)**: Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

e) **Sodium hydroxide solution (200 g/L - 500 g/L)** (1): Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

f) **Bromothymol blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.

g) **Methyl red solution (0.1 g/100 mL)**: Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

h) **Methylene blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

i) **Methyl red–methylene blue mixture solution**: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

j) **Bromocresol green solution (0.5 g/100 mL)**: Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

k) **Methyl red–bromocresol green mixture solution**: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

l) **Phosphate solution**: Dissolve 3.63 g of potassium dihydrogen phosphate specified in JIS K 9007 and 5.68 g of disodium hydrogen phosphate specified in JIS K 9020 in 1000 mL of water (5). When it is used, adjust the liquid temperature at about 25 °C (cold buffer solution).

**Note**

(1) This is an example of preparation; prepare an amount as appropriate.

(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).

(3) 5 mL -10 mL

(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.

(5) pH 7.0 ± pH 0.2

**Comment 1** A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

**Comment 2** 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.
(3) **Apparatus and instruments**: Apparatus and instruments are shown below.
   a) **Rotary shaker**: A rotary shaker that can rotate a 250-mL volumetric flask upside down at 30 - 40 revolutions/min.
   b) **Steam distillation apparatus**
   c) **Digestion flask**: A Kjeldahl flask which can be connected to a steam distillation apparatus.

(4) **Test procedures**

(4.1) **Extraction**: Conduct extraction as shown below.
   a) Weigh 1.00 g of an analytical sample (6), and put it into a 250 mL- Erlenmeyer flask.
   b) Add about 200 mL of phosphate solution and shake to mix at the rate of 30 - 40 revolutions/min for about 30 minutes.
   c) Add water up to the marked line.
   d) Filter with Type 3 filter paper to make a sample solution.

**Note** (6) Prepare the test sample according to **Comment 3**.

**Comment 3** Crush a laboratory sample with a mortar or a pestle, etc. until it completely passes through an 850 µm aperture sieve.

**Comment 4** If there is no possibility for an analytical sample to hydrolyze, it is allowed to use water instead of a phosphoric acid solution.

**Comment 5** The temperature of a solution in the procedure in (4.1) b) - d) should be no more than 26 ºC.

(4.2) **Kjeldahl method**: Conduct digestion as shown below.
   a) Transfer a predetermined amount of the sample solution (the equivalents of 0.5 g or less as cold buffer solution soluble nitrogen) to a 300-mL distillation flask.
   b) Add 0.1 g of copper (II) sulfate pentahydrate (7) specified in JIS K 8962, and further add 5 mL of sulfuric acid, shake to mix and heat gently to evaporate moisture.
   c) After standing to cool, add 1 g of potassium sulfate specified in JIS K 8962 and heat to digest.
   d) Ignite further for 30 minutes.
   e) After standing to cool, add water while shaking to mix until liquid volume reaches about 30 mL and cool it to make the digestion solution.

**Note** (7) Crush into powder as appropriate.

(4.3) Distillation: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.
   a) Transfer a predetermined amount (8) of 0.25 mol/L sulfuric acid to an acceptor, (9) add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount (8) of boric acid solution (40 g/L) to an acceptor, (9) add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.
   b) Add a proper amount of sodium hydroxide solution (200 g/L - 500 g/L) (10), and immediately connect this digestion flask to the steam distillation apparatus.
   c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.
   d) Stop distilling when the distillate has reached 120 mL - 160 mL.
   e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.
Note  (8) 5 mL - 20 mL
(9) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).
(10) An amount sufficient to make the solution strong alkalinity. A blue color will appear.

(4.4) Measurement: Conduct measurement as shown below.

(4.4.1) When 0.25 mol/L sulfuric acid is used in (4.2):
a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4).
b) Calculate the cold buffer solution soluble nitrogen in the analytical sample by the following formula:

\[
\text{Cold buffer solution soluble nitrogen (% (mass fraction)) in the analytical sample} = \frac{B \times V_6 - V_7}{V_8/V_9} \times \frac{C_1 \times f_1 \times (V_8/V_9)}{(14.007/W_2) \times (100/1000)}
\]

\[
= \frac{B \times V_6 - V_7}{V_8/V_9} \times \frac{C_1 \times f_1 \times (V_8/V_9)}{(1.4007/W_2)}
\]

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>Volume of 0.1 mol/L -0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid</td>
</tr>
<tr>
<td>V_6</td>
<td>Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a</td>
</tr>
<tr>
<td>V_7</td>
<td>Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration</td>
</tr>
<tr>
<td>C_1</td>
<td>Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution</td>
</tr>
<tr>
<td>f_1</td>
<td>Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution</td>
</tr>
<tr>
<td>V_8</td>
<td>Predetermined volume (mL) of the extract in (4.1) c</td>
</tr>
<tr>
<td>V_9</td>
<td>Transferred amount (mL) of the extract subjected to the Kjeldahl method in (4.2) a</td>
</tr>
</tbody>
</table>

W_2: Mass (g) of the analytical sample

(4.4.2) When a boric acid solution (40 g/L) is used in (4.2):
a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red (11).
b) Calculate the cold buffer solution soluble nitrogen in the analytical sample by the following formula:

\[
\text{Cold buffer solution soluble nitrogen (% (mass fraction)) in the analytical sample} = \frac{V_{10} \times C_2 \times f_2 \times (V_{11}/V_{12}) \times (14.007/W_2)}{(100/1000)}
\]

\[
= \frac{V_{10} \times C_2 \times f_2 \times (V_{11}/V_{12})}{(2.8014/W_2)}
\]

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>V_{10}</td>
<td>Volume (mL) of 0.25 mol/L sulfuric acid needed for titration</td>
</tr>
<tr>
<td>C_2</td>
<td>Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid</td>
</tr>
<tr>
<td>f_2</td>
<td>Factor of 0.25 mol/L sulfuric acid</td>
</tr>
<tr>
<td>V_{11}</td>
<td>Predetermined volume (mL) of the extract in (4.1) c</td>
</tr>
<tr>
<td>V_{12}</td>
<td>Transferred amount (mL) of the extract subjected to the Kjeldahl method in (4.2) a</td>
</tr>
</tbody>
</table>

W_2: Mass (g) of the analytical sample

Note  (11) The endpoint is reached when the color changes from green to light red.

Comment 6 The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.4)
can be conducted by an automatic titrator. The setup of the titration program, the
determination parameter for the endpoint and vessels such as acceptors are according
to the specification and the operation method of the automatic titrator used.

References
1) Masayoshi KOSHINO: Second Revision of The Methods of Analysis of Fertilizers
(Details), p.67-68, Yokendo, Tokyo (1988)

(5) Flow sheet for cold buffer solution soluble nitrogen testing method: The flow sheets for
cold buffer solution soluble nitrogen testing method in formaldehyde processed urea
fertilizers are shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00 g analytical sample</td>
<td>Weigh into a 250-mL volumetric flask.</td>
</tr>
<tr>
<td>← 150 mL phosphate solution</td>
<td></td>
</tr>
<tr>
<td>Shaking to mix</td>
<td>Rotary shaker (30 - 40 revolutions/min), 1 hour</td>
</tr>
<tr>
<td>← small amount of water</td>
<td></td>
</tr>
<tr>
<td>Filtration</td>
<td>Type 3 filter paper</td>
</tr>
<tr>
<td>Extract</td>
<td></td>
</tr>
</tbody>
</table>

Figure 1 Flow sheet for cold buffer solution soluble nitrogen testing method
in formaldehyde processed urea fertilizers
(Extraction procedure)

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extract</td>
<td>300-mL digestion flask</td>
</tr>
<tr>
<td>Aliquot (predetermined volume)</td>
<td></td>
</tr>
<tr>
<td>← 0.1 g copper (II) sulfate pentahydrate</td>
<td></td>
</tr>
<tr>
<td>← 5 mL sulfuric acid</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>Gently</td>
</tr>
<tr>
<td>Standing to cool</td>
<td></td>
</tr>
<tr>
<td>← 1 g potassium sulfate</td>
<td></td>
</tr>
<tr>
<td>Heating</td>
<td>After digestion, ignite further for 30 minutes</td>
</tr>
<tr>
<td>Standing to cool</td>
<td></td>
</tr>
<tr>
<td>← Water (until liquid volume reaches about 30 mL)</td>
<td></td>
</tr>
<tr>
<td>Cooling</td>
<td></td>
</tr>
<tr>
<td>Digestion solution</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2 Flow sheet for cold buffer solution soluble nitrogen testing method
in formaldehyde processed urea fertilizers
(Kjeldahl method procedure)
Sodium hydroxide solution (200 g/L - 500 g/L)

Receiver: 200-mL - 300 - mL Erlenmeyer flask or beaker
A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution

Distillation rate: 5 mL/min - 7 mL/min

120 mL – 160 mL distillate

Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)

0.1 mol/L-0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or 0.25 mol/L sulfuric acid (until the solution becomes light red)

Figure 3 Flow sheet for cold buffer solution soluble nitrogen testing method in formaldehyde processed urea fertilizers
(Distillation and measurement procedure)
6.6 Heat buffer solution soluble nitrogen (hot-water soluble nitrogen)

6.6.a Heat buffer solution method

(1) Summary

The testing method is applicable to methylolurea polymerization fertilizers. This testing method is classified as Type A(Def-M) and its symbol is 6.6.a-2017 or Buf(H)-N.a-1.

Add a heat phosphate solution (heat buffer solution) to an analytical sample to elute heat buffer solution soluble nitrogen, add copper (II) sulfate pentahydrate, sulfuric acid and potassium sulfate, pretreat non-dissolved matter by Kjeldahl method to change heat buffer solution soluble nitrogen to ammonium ion and add a sodium hydroxide solution to subject to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain the nitrogen which is insoluble in a heat buffer solution in the analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain the nitrogen which is insoluble in a heat buffer solution in the analytical sample. Subtract the nitrogen which is insoluble in a heat buffer solution from the separately obtained total nitrogen (T-N) according to the method in 4.1.1 to calculate the nitrogen which is soluble in a heat buffer solution (hot water-soluble nitrogen).

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution \(^{(1)}\): Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4 - 5 days. Transfer 5.5 mL -11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line \(^{(1)}\). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
\text{Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (}f_1\text{)} = \left( W_1 \times A \times 0.01/97.095 \right) \times \left( V_1/V_2 \right) \times \left( 1000/V_3 \right) \times (1/C_1)
\]

\( W_1 \): Mass (g) of sulfamic acid sampled
\( A \): Purity (% (mass fraction)) of sulfamic acid
\( V_1 \): Volume (mL) of sulfamic acid solution transferred
\( V_2 \): Constant volume (250 mL) of sulfamic acid solution
\( V_3 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
\( C_1 \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid \(^{(1)}(2)\): Add about 14 mL of sulfuric acid to a beaker containing 100 mL of water in advance, stir well, and add water to make 1000 mL.

Standardization: Transfer a predetermined amount \(^{(3)}\) of 0.25 mol/L sulfuric acid to a 200-mL
- 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L -0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula (1). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula (2):

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\[ V_4/V_5 \] ............ (1)

Factor of 0.25 mol/L sulfuric acid \((f_2)\)

\[ (f_1 \times C_1 \times V_4/V_5)/(C_2 \times 2) \] ............ (2)

\( V_4 \): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\( V_5 \): Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization

\( C_1 \): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\( C_2 \): Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

d) Boric acid solution (40 g/L): Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

e) Catalyst \((5)\): Mix potassium sulfate specified in JIS K 8962 and copper (II) sulfate pentahydrate \((6)\) specified in JIS K 8983 in the ratio of 9 to 1.

f) Sodium hydroxide solution (200 g/L - 500 g/L) \((1)\): Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

g) Bromothymol blue solution (0.1 g/100 mL): Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.

h) Methyl red solution (0.1 g/100 mL): Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

i) Methylene blue solution (0.1 g/100 mL): Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

j) Methyl red–methylene blue mixture solution: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

k) Bromocresol green solution (0.5 g/100 mL): Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

l) Methyl red–bromocresol green mixture solution: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

m) Heat phosphate solution: Dissolve 3.63 g of potassium dihydrogen phosphate specified in JIS K 9007 and 5.68 g of disodium hydrogen phosphate specified in JIS K 9020 in 1000 mL of water \((7)\). When it is used, heat until it reaches boiling point (heat buffer solution).

Note

(1) This is an example of preparation; prepare an amount as appropriate.
(2) This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).
(3) 5 mL - 10 mL
(4) The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.
(5) A tablet is commercially available.
(6) Crush into powder as appropriate.
(7) pH 7.0 ± pH 0.2
Comment 1 A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

Comment 2 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Water bath: A water bath which can boil water.
   b) Steam distillation apparatus
   c) Digestion flask: A Kjeldahl flask which can be connected to a steam distillation apparatus.
   d) Distillation flask: A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1.00 g of an analytical sample (8) and transfer to a 300-mL tall beaker.
   b) Add 100 mL of heat phosphate solution and stir it gently.
   c) Cover the tall beaker with a watch glass, and heat in a boiling water bath while stirring at ten-minute intervals for about 30 minutes.
   d) Immediately filter with Type 3 filter paper, move the whole non-dissolved matter in a vessel on the filter paper with 100 mL of heat phosphate solution and wash the non-dissolved matter with hot water.

Note (8) Prepare the test sample according to Comment 3.

Comment 3 Crush a laboratory sample with a mortar or a pestle, etc. until it completely passes through an 850 µm aperture sieve.

(4.2) Kjeldahl method: Conduct digestion as shown below.
   a) Put the non-dissolved matter in (4.1) d) together with the filter paper into a 300-mL - 500-mL digestion flask.
   b) Add 5 g - 10 g of catalyst, and further add 20 mL - 40 mL of sulfuric acid, shake to mix and heat gently.
   c) After bubbles cease to form, heat until white smoke of sulfuric acid evolves.
   d) Ignite until organic matters are completely digested (9).
   e) After standing to cool, add a small amount of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water (10), and further mix by shaking.
   f) After standing to cool, add water up to the marked line to make the digestion solution.

Note (9) When the solution has finished changing color, heat further for no less than 2 hours.
(10) When the entire sample solution volume is used in measurement, it is not necessary to transfer it to a volumetric flask.

(4.3) Distillation: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.
   a) Transfer a predetermined amount (11) of 0.25 mol/L sulfuric acid to an acceptor (12), add a few drops of methyl red–methylen blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount (11) of boric acid solution (40 g/L) to an acceptor (12), add a few drops of methyl red–bromocresol green mixture solution, and

connect this acceptor to a steam distillation apparatus.

b) Transfer a predetermined amount of the digestion solution to a 300-mL distillation flask, add a proper amount of sodium hydroxide solution (200 g/L - 500 g/L) \(^{(13)}\), and immediately connect this distillation flask to the steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

\textbf{Note} \((11)\) 5 mL - 20 mL

\((12)\) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).

\((13)\) An amount sufficient to make the solution strong alkalinity. A blue color will appear.

\textbf{4.4 Measurement:} Conduct measurement as shown below.

\textbf{4.4.1} When 0.25 mol/L sulfuric acid is used in \((4.3)\):

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green \(^{(4)}\).

b) Calculate the heat buffer solution soluble nitrogen in the analytical sample by the following formula:

c) Subtract the nitrogen which is insoluble in a heat buffer solution from the separately obtained total nitrogen (T-N) according to the method in \(4.1.1\) to calculate the nitrogen which is soluble in a heat buffer solution \(^{(14)}\).

Heat buffer solution soluble nitrogen (% (mass fraction)) in the analytical sample

\[
= \left( B \times V_6 - V_7 \right) \times C_1 \times f_1 \times \left( V_8/V_9 \right) \times \left( 14.007/W_2 \right) \times \left( 100/1000 \right)
\]

\[
= \left( B \times V_6 - V_7 \right) \times C_1 \times f_1 \times \left( V_8/V_9 \right) \times \left( 1.4007/W_2 \right)
\]

\(B\): Volume of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\(V_6\): Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in \((4.2)\) a)

\(V_7\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration

\(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(f_1\): Factor of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

\(V_8\): Predetermined volume (mL) of the digestion solution in \((4.2)\) f)

\(V_9\): Transferred amount (mL) of the digestion solution subjected to distillation in \((4.3)\) b)

\(W_2\): Mass (g) of the analytical sample

\textbf{Note} \((14)\) The total nitrogen (T-N) and the nitrogen which is insoluble in a heat buffer solution use raw data without rounding numerical value.

\textbf{4.4.2} When a boric acid solution (40 g/L) is used in \((4.3)\):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red \(^{(15)}\).

b) Calculate the heat buffer solution soluble nitrogen in the analytical sample by the following formula:
c) Subtract the nitrogen which is insoluble in a heat buffer solution from the separately obtained total nitrogen (T-N) according to the method in 4.1.1 to calculate the nitrogen which is soluble in a heat buffer solution.

Heat buffer solution soluble nitrogen (% (mass fraction)) in the analytical sample

\[ V_{10} \times C_2 \times 2 \times f_2 \times \left( \frac{V_{11}}{V_{12}} \right) \times \left( \frac{14.007}{W_2} \right) \times \left( \frac{2.8014}{W_2} \right) \]

\[ V_{10} : \text{Volume (mL) of 0.25 mol/L sulfuric acid needed for titration} \]
\[ C_2 : \text{Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid} \]
\[ f_2 : \text{Factor of 0.25 mol/L sulfuric acid} \]
\[ V_{11} : \text{Predetermined volume (mL) of the digestion solution in (4.2) f} \]
\[ V_{12} : \text{Transferred amount (mL) of the digestion solution subjected to distillation in (4.3) b} \]
\[ W_2 : \text{Mass (g) of the analytical sample} \]

**Note** (15) The endpoint is reached when the color changes from green to light red.

**Comment 4** The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.4) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.

**References**


(5) **Flow sheet for heat buffer solution soluble nitrogen testing method**: The flow sheets for heat buffer solution soluble nitrogen testing method in methylolurea polymerization fertilizers are shown below:

![Flow sheet for heat buffer solution soluble nitrogen testing method](image)

Figure 1 Flow sheet for heat buffer solution soluble nitrogen testing method in methylolurea polymerization fertilizers

(Extraction procedure)
Figure 2  Flow sheet for heat buffer solution soluble nitrogen testing method in methylolurea polymerization fertilizers (Kjeldahl method procedure)

Figure 3  Flow sheet for heat buffer solution soluble nitrogen testing method in methylolurea polymerization fertilizers (Distillation and measurement procedure)
6.7 Activity coefficient of nitrogen
6.7.a Buffer solution method

(1) Summary
The testing method is applicable to formaldehyde processed urea fertilizers. This testing method is classified as Type A(Def-M) and its symbol is 6.7.a-2017 or AI-N.a-1.

Add water to an analytical sample to elute cold water-soluble nitrogen, add potassium sulfate copper (II) sulfate pentahydrate and sulfuric acid, pretreat non-dissolved matter by Kjeldahl method to change cold water-soluble nitrogen to ammonium ion and add a sodium hydroxide solution to subject to steam distillation. Collect isolated ammonia with 0.25 mol/L sulfuric acid and measure surplus sulfuric acid by (neutralization) titration using a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution to obtain cold water insoluble nitrogen in the analytical sample. Or collect isolated ammonia with a boric acid solution and measure ammonium ion by (neutralization) titration using 0.25 mol/L sulfuric acid to obtain cold water-soluble nitrogen in the analytical sample. Separately add a heat phosphate solution (heat buffer solution) to an analytical sample to elute the nitrogen which is soluble in a heat buffer solution and conduct the same procedures as above to obtain the nitrogen which is insoluble in a heat buffer solution in the analytical sample. Subtract the nitrogen which is insoluble in a heat buffer solution from cold water non-dissolved matter and calculate the activity coefficient of nitrogen by dividing the subtracted value by the cold water non-dissolved matter.

(2) Reagent: Reagents are as shown below.

a) 0.1 mol/L - 0.2 mol/L sodium hydroxide solution (1). Transfer about 30 mL of water to a polyethylene bottle, dissolve about 35 g of sodium hydroxide specified in JIS K 8576 by adding in small portions while cooling, seal tightly and leave at rest for 4-5 days. Transfer 5.5 mL - 11 mL of the supernatant to a ground-in stoppered storage container, and add 1000 mL of water.

Standardization: Dry sulfamic acid reference material for volumetric analysis specified in JIS K 8005 by leaving at rest in a desiccator at no more than 2 kPa for about 48 hours, then transfer about 2.5 g to a weighing dish, and measure the mass to the order of 0.1 mg.

Dissolve in a small amount of water, transfer to a 250-mL volumetric flask, and add water up to the marked line (1). Transfer a predetermined amount of the solution to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of bromothymol blue solution (0.1 g/100 mL) as an indicator, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes green. Calculate the factor of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution by the following formula:

\[
W_1 \times A \times 0.01/97.095 \times (V_1/V_2) \times (1000/V_3) \times (1/C_1)
\]

Where:
- \(W_1\): Mass (g) of sulfamic acid sampled
- \(A\): Purity (% (mass fraction)) of sulfamic acid
- \(V_1\): Volume (mL) of sulfamic acid solution transferred
- \(V_2\): Constant volume (250 mL) of sulfamic acid solution
- \(V_3\): Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- \(C_1\): Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution

b) Sulfuric acid: A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.

c) 0.25 mol/L sulfuric acid (1)(2): Add about 14 mL of sulfuric acid to a beaker containing 100
mL of water in advance, stir well, and add water to make 1000 mL.

**Standardization**: Transfer a predetermined amount \(^{(3)}\) of 0.25 mol/L sulfuric acid to a 200-mL - 300-mL Erlenmeyer flask, add a few drops of methyl red–methylene blue mixture solution, and titrate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green \(^{(4)}\). Calculate the volume of a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid by the following formula \(^{(1)}\). Or, calculate the factor of 0.25 mol/L sulfuric acid by the following formula \(^{(2)}\):

Volume (B) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid

\[
\frac{V_4}{V_5} = \frac{f_1 \times C_1 \times V_4}{(C_2 \times 2)} \quad \text{......... (1)}
\]

Factor of 0.25 mol/L sulfuric acid \((f_2)\)

\[
f_2 = \frac{(f_1 \times C_1 \times V_4 / V_5)}{(C_2 \times 2)} \quad \text{......... (2)}
\]

- **V_4**: Volume (mL) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution needed for titration
- **V_5**: Volume (mL) of 0.25 mol/L sulfuric acid subjected to standardization
- **C_1**: Set concentration (mol/L) of 0.1 mol/L - 0.2 mol/L sodium hydroxide solution
- **C_2**: Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid

**d) Boric acid solution (40 g/L)**: Dissolve 40 g of boric acid specified in JIS K 8863 in water to make 1000 mL.

**e) Catalyst \(^{(5)}\)**: Mix potassium sulfate specified in JIS K 8962 and copper (II) sulfate pentahydrate \(^{(6)}\) specified in JIS K 8983 in the ratio of 9 to 1.

**f) Sodium hydroxide solution (200 g/L - 500 g/L) \(^{(1)}\)**: Dissolve 100 g - 250 g of sodium hydroxide specified in JIS K 8576 in water to make 500 mL.

**g) Bromothymol blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of bromothymol blue specified in JIS K 8842 in 20 mL of ethanol (95) specified in JIS K 8102, add water to make 100 mL.

**h) Methyl red solution (0.1 g/100 mL)**: Dissolve 0.1 g of methyl red specified in JIS K 8896 in 100 mL of ethanol (95) specified in JIS K 8102.

**i) Methylene blue solution (0.1 g/100 mL)**: Dissolve 0.1 g of methylene blue specified in JIS K 8897 in 100 mL of ethanol (95) specified in JIS K 8102.

**j) Methyl red–methylene blue mixture solution**: To 2 volumes of methyl red solution (0.1 g/100 mL), add 1 volume of methylene blue solution (0.1 g/100 mL).

**k) Bromocresol green solution (0.5 g/100 mL)**: Dissolve 0.5 g of bromocresol green specified in JIS K 8840 in 100 mL of ethanol (95) specified in JIS K 8102.

**l) Methyl red–bromocresol green mixture solution**: To a methyl red solution (0.1 g/100 mL), add equal volume of bromocresol green solution (0.5 g/100 mL).

**m) Heat phosphate solution**: Dissolve 1.43 g of potassium dihydrogen phosphate specified in JIS K 9007 and 9.10 g of disodium hydrogen phosphate specified in JIS K 9020 in 1000 mL of water \(^{(7)}\). When it is used, heat until it reaches boiling point (heat buffer solution).

**Note**

1. This is an example of preparation; prepare an amount as appropriate.
2. This corresponds to the standard sulfuric acid solution 0.5 M (1/2 sulfuric acid) solution in the Official Methods of Analysis of Fertilizers (1992).
3. 5 mL - 10 mL
4. The endpoint is reached when the color becomes gray-green via dark blue from blue-purple.
5. A tablet is commercially available.
(6) Crush into powder as appropriate.
(7) pH 7.5 ± pH 0.2

Comment 1 A 0.1 mol/L - 0.2 mol/L sodium hydroxide solution in (2) a) can be replaced with a 0.1 mol/L sodium hydroxide solution or a 0.2 mol/L sodium hydroxide solution conforming to ISO/IEC 17025.

Comment 2 0.25 mol/L sulfuric acid in (2) c) can be replaced with 0.25 mol/L sulfuric acid conforming to ISO/IEC 17025.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Water bath: A water bath which can boil water.
   b) Steam distillation apparatus
   c) Digestion flask: A Kjeldahl flask which can be connected to a steam distillation apparatus.
   d) Distillation flask: A Kjeldahl flask or round bottom flask that can be connected to a steam distillation apparatus.

(4) Test procedures

(4.1) Extraction: Conduct extraction as shown below. Subject the non-dissolved matters of (4.1.1) f) and (4.1.2) d) to (4.2) Kjeldahl method respectively.

(4.1.1) Extraction by cold water
   a) Weigh 1.00 g of an analytical sample (8) and transfer to a 50-mL beaker.
   b) Add a small amount of ethanol specified in JIS K 8101 to moisten and add 20 mL of water at 25°C ± 2°C to stir.
   c) Leave at rest for 15 minutes while stirring at 5-minute intervals.
   d) Filter the supernatant with Type 2 filter paper
   e) Wash non-dissolved matter with water at 25°C ± 2°C five times and filter the supernatant.
   f) Transfer the whole non-dissolved matter on the filter paper using water at 25°C ± 2°C and wash the non-dissolved matter using water at the same temperature until filtrate reaches 250 mL.

Note (8) Prepare the test sample according to Comment 3.

Comment 3 Crush a laboratory sample with a mortar or a pestle, etc. until it completely passes through an 850 µm aperture sieve.

(4.1.2) Extraction by heat phosphate solution
   a) Weigh an analytical sample (8) equivalent to 0.12 g of cold non-resolved matter and put it in a 200-mL tall beaker.
   b) Add 100 mL of heat phosphate solution and stir it.
   c) Cover the tall beaker with a watch glass, and heat in a boiling water bath while stirring at ten-minute intervals for about 30 minutes.
   d) Immediately filter with Type 2 filter paper (9), transfer the whole non-dissolved matter in a vessel on the filter paper using boiling water and wash the non-dissolved matter using 100 mL of boiling water.

Note (9) If it takes 4 minutes or more for filtrating, conduct extracting anew according to Comment 4.

Comment 4 After conducting the procedures in (4.1.2) a) - c), add 1 g of diatomaceous earth to stir and conduct the procedure in (4.1.2) d).
(4.2) **Kjeldahl method**: Conduct digestion as shown below.

a) Put the non-dissolved matter in (4.1.1) f) and (4.1.2) d) together with the filter paper into respective 300-mL - 500-mL digestion flasks.

b) Add 5 g - 10 g of catalyst, and further add 20 mL - 40 mL of sulfuric acid, shake to mix and heat gently.

c) After bubbles cease to form, heat until white smoke of sulfuric acid evolves.

d) Ignite until organic matters are completely digested (10).

e) After standing to cool, add a small amount of water, mix well by shaking, transfer to a 250-mL - 500-mL volumetric flask with water (11), and further mix by shaking.

f) After cooling is complete, add water up to the marked line to make the digestion solution.

**Note** (10) When the solution has finished changing color, heat further for no less than 2 hours.

(11) When the entire sample solution volume is used in measurement, it is not necessary to transfer it to a volumetric flask.

(4.3) **Distillation**: Conduct distillation as shown below. Specific distillation procedures are according to the operation method of the steam distillation apparatus used in measurement.

a) Transfer a predetermined amount (12) of 0.25 mol/L sulfuric acid to an acceptor (13), add a few drops of methyl red–methylene blue mixture solution, and connect this acceptor to a steam distillation apparatus. Or, transfer a predetermined amount (12) of boric acid solution (40 g/L) to an acceptor (13), add a few drops of methyl red–bromocresol green mixture solution, and connect this acceptor to a steam distillation apparatus.

b) Transfer a predetermined amount of the digestion solution to a 300-mL distillation flask, add a proper amount of sodium hydroxide solution (200 g/L - 500 g/L) (14), and immediately connect this distillation flask to the steam distillation apparatus.

c) Send steam to the distillation flask to heat the solution in the distillation flask, and distill at a distillation rate of 5 mL/min - 7 mL/min.

d) Stop distilling when the distillate has reached 120 mL - 160 mL.

e) Wash the part of the steam distillation apparatus that came in contact with the solution in the acceptor with a small amount of water, and pool the washing with the distillate.

**Note** (12) 5 mL - 20 mL

(13) As an acceptor, use a 200-mL - 300-mL Erlenmeyer flask or a 200-mL - 300-mL beaker with which the distillate outlet of the steam distillation apparatus can be immersed in 0.25 mol/L sulfuric acid or a boric acid solution (40 g/L).

(14) An amount sufficient to make the solution strong alkalinity. A blue color will appear.

(4.4) **Measurement**: Conduct measurement as shown below.

(4.4.1) When 0.25 mol/L sulfuric acid is used in (4.3):

a) Titrate the distillate with a 0.1 mol/L - 0.2 mol/L sodium hydroxide solution until the color of the solution becomes gray-green (4).

b) Calculate the cold-water insoluble nitrogen \(N_1\) and the nitrogen which is insoluble in a heat buffer solution \(N_2\) in the analytical sample by the following formula (3):

c) Obtain the activity coefficient of nitrogen in the analytical sample by the following formula (4) (15).

\[
\text{Cold water insoluble nitrogen (} N_1 \text{) or the nitrogen which is insoluble in a heat buffer solution (} N_2 \text{) (\% (mass fraction)) in the analytical sample} \\
= (B \times V_6 - V_7) \times C_1 \times f_1 \times (V_6/V_5) \times \left(14.007/W_2\right) \times (100/1000)
\]
\[
(B \times V_6 - V_7) \times C_1 \times f_1 \times (V_8/V_9) \times (1.4007/W_2) \quad \cdots \cdots (3)
\]

**B:** Volume of 0.1 mol/L-0.2 mol/L sodium hydroxide solution equivalent to 1 mL of 0.25 mol/L sulfuric acid  
**V_6:** Volume (mL) of 0.25 mol/L sulfuric acid transferred to the acceptor in (4.2) a)  
**V_7:** Volume (mL) of 0.1 mol/L-0.2 mol/L sodium hydroxide solution needed for titration  
**C_1:** Set concentration (mol/L) of 0.1 mol/L-0.2 mol/L sodium hydroxide solution  
**f_1:** Factor of 0.1 mol/L-0.2 mol/L sodium hydroxide solution  
**V_8:** Predetermined volume (mL) of the digestion solution in (4.2) f)  
**V_9:** Transferred amount (mL) of the digestion solution subjected to distillation in (4.3) b)  
**W_2:** Mass (g) of the analytical sample

Activity coefficient of nitrogen (%)
\[
= ((N_1 - N_2)/N_1) \times 100 
\quad \cdots \cdots (4)
\]

**N_1:** Cold water insoluble nitrogen (% (mass fraction))  
**N_2:** Heat buffer solution insoluble nitrogen (% (mass fraction))

**Note** (15) Cold water insoluble nitrogen (N_1) or the nitrogen which is insoluble in a heat buffer solution (N_2) uses raw data without rounding numerical value.

(4.4.2) When a boric acid solution (40 g/L) is used in (4.3):

a) Titrate the distillate with 0.25 mol/L sulfuric acid until the color of the solution becomes light red (16).  
b) Calculate the cold-water insoluble nitrogen (N_1) and the nitrogen which is insoluble in a heat buffer solution (N_2) in the analytical sample by the following formula (5):
c) Obtain the activity coefficient of nitrogen in the analytical sample by the formula (4) in (4.4.1) (14).

Cold water insoluble nitrogen (N_1) or the nitrogen which is insoluble in a heat buffer solution (N_2) (% (mass fraction)) in the analytical sample
\[
V_{10} \times C_2 \times f_2 \times (V_{11}/V_{12}) \times (14.007/W_2) \times (100/1000) 
\quad = V_{10} \times C_2 \times f_2 \times (V_{11}/V_{12}) \times (2.8014/W_2) 
\quad \cdots \cdots (3)
\]

**V_{10}:** Volume (mL) of 0.25 mol/L sulfuric acid needed for titration  
**C_2:** Set concentration (0.25 mol/L) of 0.25 mol/L sulfuric acid  
**f_2:** Factor of 0.25 mol/L sulfuric acid  
**V_{11}:** Predetermined volume (mL) of the digestion solution in (4.2) f)  
**V_{12}:** Transferred amount (mL) of the digestion solution subjected to distillation in (4.3) b)  
**W_2:** Mass (g) of the analytical sample

**Note** (16) The endpoint is reached when the color changes from green to light red.

**Comment 5** The titration procedures in (2) a) Standardization, (2) c) Standardization and (4.4) can be conducted by an automatic titrator. The setup of the titration program, the determination parameter for the endpoint and vessels such as acceptors are according to the specification and the operation method of the automatic titrator used.
Flow sheet for the activity coefficient of nitrogen testing method: The flow sheets for the activity coefficient of nitrogen testing method in formaldehyde processed urea fertilizers are shown below:

Figure 1-1 Flow sheet for the activity coefficient of nitrogen testing method in formaldehyde processed urea fertilizers (Extraction procedure by cold water (4.1.1))

Figure 1-2 Flow sheet for the activity coefficient of nitrogen testing method in formaldehyde processed urea fertilizers (Extraction procedure by heat phosphate solution (4.1.2))
Subject Cold water non-dissolved mater or heat buffer solution non-dissolved matter to Kjeldahl method, distillation and measurement respectively

300-mL - 500-mL digestion flask

← 5 g - 10 g catalyst
← 20 mL - 40 mL sulfuric acid

Gently

After bubbles cease to form, ignite until organic matters are completely digested

A small amount of water

250-mL - 500-mL volumetric flask, water

Water (up to the marked line)

Figure 2 Flow sheet for the activity coefficient of nitrogen testing method in formaldehyde processed urea fertilizers (Kjeldahl method procedure)
Sodium hydroxide solution (200 g/L - 500 g/L)
Receiver: 200-mL - 300-mL Erlenmeyer flask or beaker
A predetermined amount of 0.25 mol/L sulfuric acid and a few drops of methyl red-methylene blue mixture solution, or boric acid solution (40 g/L), several drops of methyl red - bromocresol green mixture solution
Distillation rate: 5 mL/min - 7 mL/min
120 mL – 160 mL distillate
Water (wash the part of the distillation apparatus that came in contact with the solution in the receiver)
0.1 mol/L-0.2 mol/L sodium hydroxide solution (until the solution becomes gray-green), or 0.25 mol/L sulfuric acid (until the solution becomes light red)

Figure 3 Flow sheet for the activity coefficient of nitrogen testing method in formaldehyde processed urea fertilizers (Distillation and measurement procedure)
6.8 Initial elution rate
6.8.a Standing-in-water method

(1) Summary
The testing method is applicable to coated fertilizer. This testing method is classified as Type A(Def-E) and its symbol is 6.8.a-2017 or SDR.a-1.

An initial elution rate is a quick-acting component of coated fertilizers. Target components include total nitrogen (T-N), ammoniacal nitrogen (A-N), nitrate nitrogen (N-N), water-soluble phosphoric acid (W-P₂O₅), water-soluble potassium (W-K₂O) and water-soluble magnesia (W-MgO).

Add water to a laboratory sample, leave it at rest while maintaining in water at 30ºC for 24 hours and obtain the initial elution flow of a target component. Separately measure the corresponding component content by 4.1.1, 4.1.2, 4.1.3, 4.2.4, 4.3.3 or 4.6.3. Calculate an initial elution rate dividing the initial elution flow of a target component by the corresponding component content.

(2) Reagent: Reagents are as shown below.
   a) Reagent solutions for total nitrogen: Reagents corresponding to clauses in 4.1.1 when determining total nitrogen.
   b) Reagent solutions for ammoniacal nitrogen: Reagents corresponding to clauses in 4.1.2 when determining ammoniacal nitrogen.
   c) Reagent solutions for nitrate nitrogen: Reagents corresponding to clauses in 4.1.3 when determining nitrate nitrogen.
   d) Reagent solutions for water-soluble phosphoric acid: Reagents corresponding to clauses in 4.2.4 when determining water-soluble phosphoric acid.
   e) Reagent solutions for water-soluble potassium: Reagents corresponding to clauses in 4.3.3 when determining water-soluble potassium.
   f) Reagent solutions for water-soluble magnesia: Reagents corresponding to clauses in 4.6.3 when determining water-soluble magnesia.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
   a) Incubator: An incubator whose temperature is 30 ºC ± 1 ºC.
   b) Total nitrogen: Apparatus and instruments corresponding to clauses in 4.1.1 when determining total nitrogen.
   c) Ammonium nitrogen: Apparatus and instruments corresponding to clauses in 4.1.2 when determining ammoniacal nitrogen.
   d) Nitrate nitrogen: Apparatus and instruments corresponding to clauses in 4.1.3 when determining nitrate nitrogen.
   e) Water-soluble phosphoric acid: Apparatus and instruments corresponding to clauses in 4.2.4 when determining water-soluble phosphoric acid.
   f) Water-soluble potassium: Apparatus and instruments corresponding to clauses in 4.3.3 when determining water-soluble potassium.
   g) Water-soluble magnesia: Apparatus and instruments corresponding to clauses in 4.6.3 when determining water-soluble magnesia.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 12.5 g of an analytical sample, and put it into a 300 mL- ground-in stopper Erlenmeyer flask (1).
   b) Add 250 mL of water at 30°C ± 1°C, put it into an incubator at 30°C ± 1°C and leave it at rest for 24 hours (2).
   c) Filter with Type 3 filter paper (3) and shake to mix the filtrate to make a sample solution.
Note  
(1) Since no grinding is conducted and inhomogeneous laboratory samples are used, it is desirable to heighten the reliability of determined values by conducting tests using 3 - 5 laboratory samples.

(2) Since an initial elution flow is estimated to be higher than usual if the laboratory sample vibrates in water, water should be added gently. Do not shake to mix the sample solution until the filtering in c) is completed.

(3) Filter most of the solution so that non-dissolved matter remains in the Erlenmeyer flask.

(4.2) Measurement: Conduct the measurements of the initial elution flow of a target component as specified in respective clauses in a) - f). In addition, specific measurement procedure for each component is carried out according to a corresponding clause.

a) **Total nitrogen**: Transfer a predetermined amount of sample solution and quantitate total nitrogen according to respective clauses in 4.1.1 to make the initial elution flow.

b) **Ammonium nitrogen**: Transfer a predetermined amount of sample solution and quantitate ammoniacal nitrogen according to respective clauses in 4.1.2 to make the initial elution flow.

c) **Nitrate nitrogen**: Transfer a predetermined amount of sample solution and quantitate nitrate nitrogen according to respective clauses in 4.1.3 to make the initial elution flow.

d) **Water-soluble phosphoric acid**: Transfer a predetermined amount of sample solution and quantitate water-soluble phosphoric acid according to respective clauses in 4.2.4 to make the initial elution flow.

e) **Water-soluble potassium**: Transfer a predetermined amount of sample solution and quantitate water-soluble potassium according to respective clauses in 4.3.3 to make the initial elution flow.

f) **Water-soluble magnesia**: Transfer a predetermined amount of sample solution and quantitate water-soluble magnesia according to respective clauses in 4.6.3 to make the initial elution flow.

(5) **Calculation of initial elution rate**

a) Calculate an initial elution rate (%) by the following formula using the initial elution flow of a target component obtained in (4.2) and separately determine (4) the corresponding component content (5).

\[
\text{Initial elution rate} = \left( \frac{C_1}{C_2} \right) \times 100 \\
C_1: \text{Initial elution rate of a target component} \quad (\% \text{ (mass fraction)}) \\
C_2: \text{Corresponding component content} \quad (\% \text{ (mass fraction)})
\]

Note  
(4) Determine total nitrogen (T-N), ammoniacal nitrogen (A-N), nitrate nitrogen (N-N), water-soluble phosphoric acid (W-P\(_2\)O\(_5\), water-soluble potassium (W-K\(_2\)O) and water-soluble magnesia (W-MgO) in 4.1.1, 4.1.2, 4.1.3, 4.2.4, 4.3.3 or 4.6.3 using test samples prepared in 2.3 Preparation of test samples.

(5) Initial elution flow and corresponding component content use raw data without rounding numerical value.

References

(6) **Initial elution rate testing method**: The flow sheet for initial elution rate testing method in coated fertilizers is shown below:

```
| 12.5 g laboratory sample | 300 mL Erlenmeyer flask |
| ←250 mL water [30°C±1°C] | 30°C±1°C, 24 hours |
| Extract |  |
| Filtration | Type 3 filter paper |
| Determination of a target component | Initial elution rate of a target component |
```

Figure Flow sheet for initial elution rate testing method of coated fertilizers
6.9 Humic acid (Acid insoluble - alkali soluble component)  
6.9.a Gravimetric analysis  
(1) Summary  
The testing method is applicable to humic acid chloride fertilizers. This testing method is classified as Type A(Def-M) and its symbol is 6.9.a-2017 or H-acid.a-1. Add hydrochloric acid (1+9) to elute acid dissolved matter, filter non-dissolved matter and measure the mass of non-dissolved matter to obtain acid non-dissolved matter in the analytical sample. Separately add hydrochloric acid (1+9) to elute acid dissolved matter, add a sodium hydroxide solution (10 g/L) to non-dissolved matter to elute alkali dissolved matter, and filter the non-dissolved matter to obtain acid non-dissolved alkali non-dissolved matter in the analytical sample. Subtract acid non-dissolved alkali non-dissolved matter from acid dissolved matter to calculate humic acid (acid insoluble alkali soluble component).

(2) Reagent: Reagents are as shown below.  
a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.  
b) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.

(3) Instruments: Instruments are as shown below:  
a) Shaking apparatus  
b) Drying apparatus: A drying apparatus that can be adjusted to 105 °C - 110 °C.  
c) Crucible type glass filter: A crucible type glass filter 1G4 specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 105 °C - 110 °C in advance and measure the mass to the order of 1 mg.  
d) Ground-in stopperweighing bottles (1): Low-form weighing bottles, 50 mm × 30 mm, specified in JIS R 3503. Let it stand to cool in a desiccator after heating at 105 °C - 110 °C in advance and measure the mass to the order of 1 mg.

Note  
(1) Aluminum weighing dishes described in the Handbook of the Feed Analysis Standards -2009- can also be used.  
(2) A rotary shaker that can rotate a 100-mL ground-in stopper centrifugal precipitate tube upside down at 30 - 40 revolutions/min may also be used.

(4) Test procedures  
(4.1) Acid non-dissolved matter  
(4.1.1) Extraction: Conduct extraction as shown below.  
a) Weigh 1 g of an analytical sample to the order of 1 mg and put it in a 100-mL ground-in stopper centrifugal precipitate tube.  
b) Add 50 mL of hydrochloric acid (1+9) and shake to mix by using a shaking apparatus (2) for about 1 hour.  
c) Centrifuge at about 1700 × g for about five minutes (3) to remove supernatant (4).  
d) Add water to stir (5) and centrifuge at about 1700 × g for about five minutes (3) to remove supernatant (4).  
e) Repeat the procedure in d) 3 times.

Note  
(2) When using a rotary shaker, it should be adjusted to 30 - 40 revolutions/min.  
(3) 16.5-cm of radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.  
(4) Remove using Komagome pipet, etc.
(5) Stir with a glass rod, wash non-dissolved matter which may have adhered to the glass rod with water and add wash to a centrifugal precipitate tube.

(4.1.2) **Measurement**: Conduct measurement as shown below.

a) Transfer the whole non-dissolved matter in (4.1.1) e to a crucible type glass filter and filter under reduced pressure.

b) Put the non-dissolved matter together with the crucible type glass filter into a drying apparatus and heat at 105 °C - 110 °C for 3 hours.

c) As soon as heating is complete, move it into a desiccator and let it stand to cool.

d) After standing to cool, remove the crucible type glass filter from the desiccator and measure the mass to the order of 1 mg.

(4.2) **Acid non-dissolved - alkali non-dissolved matter**

(4.2.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 1 g of an analytical sample to the order of 1 mg and put it in a 100-mL ground-in stopper centrifugal precipitate tube.

b) Add 50 mL of hydrochloric acid (1+9) and shake to mix by using a shaking apparatus for about 1 hour.

c) Centrifuge at about 1700 × g for about five minutes to remove supernatant.

d) Add water to stir and centrifuge at about 1700 × g for about five minutes to remove supernatant.

e) Repeat the procedure in d) 3 times.

f) Add 50 mL of sodium hydroxide (10 g/L) and shake to mix by using a shaking apparatus for about 1 hour.

g) Centrifuge at about 1700 × g for about five minutes to remove supernatant.

h) Add water to stir and centrifuge at about 1700 × g for about five minutes to remove supernatant.

i) Repeat the procedure in h) 3 times.

(4.2.2) **Measurement**: Conduct measurement as shown below.

a) Put the non-dissolved matter together with a ground -in stopper weighing bottle into a drying apparatus and heat.

b) After standing to cool, transfer the non-dissolved matter into the ground -in stopper weighing bottle.

c) Put the non-dissolved matter together with a ground -in stopper weighing bottle into a drying apparatus and heat at 105 °C - 110 °C for 3 hours.

d) After heating, fit the stopper into the ground-in stoppered weighing bottle, and immediately transfer to a desiccator to let it stand to cool.

e) After standing to cool, remove the ground-in stoppered weighing bottle from the desiccator, and measure the mass to the order of 1 mg.

**Note** (6) Heat at the temperature to enable the procedure in (4.2.2) b).

(5) **Calculation of humic acid**

a) Calculate humic acid by the following formula

\[
\text{Humic acid (\% (mass fraction))} = \left( \frac{A_1}{W_1} \right) \times 100 - \left( \frac{A_2}{W_2} \right) \times 100 \quad \cdots \cdots \quad (1)
\]

\( A_1 \): Mass (g) of the acid non-dissolved matter determined in (4.1.2) d)
$W_1$: Mass (g) of the analytical sample sampled in (4.1.1)a

$A_2$: Mass (g) of the acid non-dissolved - alkali non-dissolved matter determined in (4.2.2) e

$W_2$: Mass (g) of the analytical sample sampled in (4.2.1)a

References

(6) **Humic acid testing method**: The flow sheet for humic acid testing method is shown below:

![Flow sheet for humic acid testing method](image)

Figure 1 Flow sheet for humic acid testing method in humic acid salt fertilizers  (Measurement procedure of acid non-dissolved matter (4.1))

Weigh to the order of 1 mg into a 100-mL ground-in stopper centrifugal precipitate tube.

$\leftarrow$ 50 mL hydrochloric acid (1+9)

Shaking, 1 hour

Ground-in stopper centrifugal precipitate tube

About $1700 \times g$, 5 minutes

Water

Ground-in stopper centrifugal precipitate tube

About $1700 \times g$, 5 minutes

Further repeat 3 times

$\leftarrow$ 50 mL sodium hydroxide (10 g/L)

Shaking, 1 hour

Ground-in stopper centrifugal precipitate tube

About $1700 \times g$, 5 minutes

Water

Ground-in stopper centrifugal precipitate tube

About $2000 \times g$, 5 minutes

Further repeat 3 times

Drying

Ground-in stopper weighing bottle

105 °C – 110 °C, 3 hours

Desiccator

Weigh to the order of 1 mg

Figure 2 Flow sheet for humic acid testing method in humic acid salt fertilizers

(Testing procedure of acid insoluble - alkali non-dissolved matter (4.2))
6.10 Sulfate
This method is according to 5.29.2 Sulfate analysis in “The Official Methods of Analysis of Fertilizers 1992”.

References


6.11 Carbon dioxide
This method is according to 5.20 Carbon dioxide analysis in “The Official Methods of Analysis of Fertilizers 1992”.

References
1) National Institute for Agro-Environmental Sciences, the Ministry of Agriculture, Forestry and Fisheries: The Official Methods of Analysis of Fertilizers 1992, p.121 - 123, Japan Fertilizers Analysis Association, Tokyo (1992)
7. Nitrification inhibitor
7.1 2-amino-4-chloro-6-methylpyrimidine (AM)
7.1.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers containing 2-amino-4-chloro-6-methylpyrimidine (AM). This testing method is classified as Type C and its symbol is 7.1.a-2017 or AM.a-1. Add methanol – water (1+1) to an analytical sample to extract 2-amino-4-chloro-6-methylpyrimidine, introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with an octadecyl silylation silica gel column, and measure at wavelength 295 nm to obtain 2-amino-4-chloro-6-methylpyrimidine (AM) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.
   c) Methanol: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.
   d) 2-amino-4-chloro-6-methylpyrimidine standard solution (1 mg/mL): Put 0.1 g of 2-amino-4-chloro-6-methylpyrimidine [C₆H₅ClN₃] in a weighing dish and measure the mass to the order of 0.1 mg. Add methanol - water (1+1) to dissolve, transfer to a 100-mL volumetric flask and add the same solvent up to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.
   e) 2-amino-4-chloro-6-methylpyrimidine standard solution (0.1 mg/mL): In the case of usage, transfer 10 mL of 2-amino-4-chloro-6-methylpyrimidine standard solution (1 mg/mL) to a 100-mL volumetric flask and add methanol - water (1+1) up to the marked line.
   f) 2-amino-4-chloro-6-methylpyrimidine standard solution (10 µg/mL - 50 µg/mL) for the calibration curve preparation: In the case of usage, transfer 5 mL - 25 mL of 2-amino-4-chloro-6-methylpyrimidine standard solution (0.1 mg/mL) for the calibration curve preparation to 50-mL volumetric flasks step-by-step and add methanol - water (1+1) up to the marked line.
   g) 2-amino-4-chloro-6-methylpyrimidine standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation: In the case of usage, transfer 2.5 mL - 25 mL of 2-amino-4-chloro-6-methylpyrimidine standard solution (20 µg/mL) for the calibration curve preparation to 50-mL volumetric flasks step-by-step and add methanol - water (1+1) up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) A reagent of no less than 98 % (mass fraction) in purity is commercially sold as 2-amino-4-chloro-6-methylpyrimidine.

Comment 1 2-amino-4-chloro-6-methylpyrimidine is commercially sold by FUJIFILM Wako Pure Chemical Co., Ltd. and Kanto Chemical Co., Inc.

(3) Instruments: Instruments are as shown below:
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube filled with silica gel, to which octadecyl chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 30 °C - 45 °C.
3) **Detection unit:** An absorptiometric detector that can measure at wavelength around 295 nm.

b) **Magnetic stirrer:**

c) **Centrifugal separator:** A centrifugal separator that can work at about 1700 × g.

d) **High speed centrifugal separator:** A centrifugal separator that can work at 8000 × g - 10000 × g.

e) **Acidic alumina cartridge column:** Link a 10-mL cylinder to a column (3) that is filled with 500 mg - 1 g of acidic alumina, put 3 mL of methanol and let it flow down.

**Note** (3) A cartridge with a 3-mL - 6-mL column filled with 500 mg - 1 g of silica gel can be used.

**Comment 2** A column is sold under production names such as Inertsil ODS, Mightysil RP-18, L-column ODS, Shim-pack VP-ODS, Silica C_{18}M 4D, Puresil C18, COSMOSIL 5C18-MS-II, etc.

**Comment 3** An acidic alumina cartridge is commercially sold under production names such as Bond Elut AL-A, Sep-Pak Alumina-A, Supelclean LC-Alumina-A.

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below:

a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.

b) Add 100 mL of methanol - water (1+1) and stir it by using a magnetic stirrer for about 30 minutes.

c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.

d) Centrifuge it at about 1700 × g centrifugal force for about five minutes (4) and use the supernatant (5) as the extract.

**Note** (4) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

(5) If there is a possibility that the 2-amino-4-chloro-6-methylpyrimidine in the sample solution exceeds the maximum limit of the calibration curve, dilute a predetermined amount of extract with methanol water (1+1).

(4.2) **Cleanup:** Conduct cleanup as shown below:

a) Transfer the extract to an acidic alumina cartridge column.

b) Dispose of about the first 3 mL of effluent and then transfer about the next 2 mL to a test tube.

c) Transfer the effluent to a 1.5-mL ground-in stopper centrifugal precipitate tube (6).

d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (7) and use the supernatant as the extract.

**Note** (6) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(7) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

**Comment 4** Instead of the procedures in (4.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of PTFE and the filtrate can be the sample solution.
Comment 5 The test is possible by the following procedures in the case of fertilizers not containing organic matters.
The procedures in (4.1) c) - d) and (4.2) a) - b) are omitted and “Transfer effluent” in (4.2) c) is replaced with the “After allowing to stand still, transfer supernatant” to operate.

(4.3) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:
1) Column: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which octadecyl chemically bonds.
2) Column bath temperature: 30 °C - 40 °C
3) Eluent: Methanol - water (4+6)
4) Flow rate: 1 mL/min
5) Detection unit: An absorptiometric detector, measurement wavelength: 295 nm

b) Calibration curve preparation
1) Inject 10 µL of respective 2-amino-4-chloro-6-methylpyrimidine standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 295 nm and obtain the peak area or height.
2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 295 nm of the respective 2-amino-4-chloro-6-methylpyrimidine standard solutions for the calibration curve preparation.

c) Sample measurement
1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain 2-amino-4-chloro-6-methylpyrimidine content from the calibration curve to calculate 2-amino-4-chloro-6-methylpyrimidine (AM) in the analytical sample.

Comment 6 A recovery testing was conducted using compound fertilizer (1 sample) and blended fertilizer (2 samples), as a result, the mean recovery rate of 2-amino-4-chloro-6-methylpyrimidine at concentration level of 1.0 % (mass fraction), 0.4 % (mass fraction) and 0.1 % (mass fraction) were 99.1 % - 100.5 %, 99.3 % - 101.6 % and 100.2 % - 100.7 %.
Additionally, the minimum limit of quantification of this testing method is about 0.005 % (mass fraction).

References
(5) **Flow sheet for 2-amino-4-chloro-6-methylpyrimidine:** The flow sheet for 2-amino-4-chloro-6-methylpyrimidine in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00 g analytical sample</td>
<td>200-mL of ground-in stopper Erlenmeyer flask</td>
</tr>
<tr>
<td>Extraction</td>
<td>100 mL of methanol-water (1+1)</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Stirring, 30 minutes</td>
</tr>
<tr>
<td>Cleanup</td>
<td>Ground-in stopper centrifugal precipitate tube, 1700 × g, 5 minutes</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Acidic alumina cartridge column</td>
</tr>
<tr>
<td>Sample solution</td>
<td>Ground-in stopper centrifugal precipitate tube, 8000 × g - 10000 × g, 5 minutes</td>
</tr>
<tr>
<td>Measurement</td>
<td>High Performance Liquid Chromatograph</td>
</tr>
</tbody>
</table>

Figure Flow sheet for 2-amino-4-chloro-6-methylprimidine (AM) in fertilizers.

**Reference:** HPLC chromatogram of 2-amino-4-chloro-6-methylpyrimidine (AM) standard solution for the calibration curve preparation is shown below.

![HPLC Chromatogram](image)

Reference diagram: HPLC chromatogram of 2-amino-4-chloro-6-methylpyrimidine (AM) standard solution

Measurement conditions for HPLC
- Column: MightySil RP-18 (4.6-mm inner diameter, 150-mm long, 5-µm particle diameter)
- 2-amino-4-chloro-6-methylpyrimidine standard solution (the equivalents of 100 ng)
- Other conditions are according to the example of HPLC measurement conditions in (4.3) a)
7.2 1-amidino-2-thiourea (ASU)
7.2.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers containing 1-amidino-2-thiourea (ASU). This testing method is classified as Type B and its symbol is 7.2.a-2017 or ASU.a-1.

Add water to an analytical sample to extract 1-amidino-2-thiourea, introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with an octadecyl silylation silica gel column, and measure at wavelength 262 nm to obtain 1-amidino-2-thiourea (ASU) in an analytical sample. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Methanol: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.
   c) 1-sodium hexasulfonate: A reagent of ion pair chromatography analysis grade or equivalents.
   d) Acetic acid: A reagent of HPLC grade or equivalents.
   e) 1-amidino-2-thiourea standard solution (1 mg/mL) (1): Put 0.1 g of 1-amidino-2-thiourea \([\text{C}_2\text{H}_6\text{N}_4\text{S}]\) (2) in a weighing dish and measure the mass to the order of 0.1 mg. Add water to dissolve, transfer to a 100-mL volumetric flask and add water up to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.
   f) 1-amidino-2-thiourea standard solution (0.1 mg/mL): In the case of usage, transfer 10 mL of 1-amidino-2-thiourea standard solution (1 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
   g) 1-amidino-2-thiourea standard solution (10 µg/mL - 50 µg/mL) for the calibration curve preparation: In the case of usage, transfer 5 mL - 25 mL of 1-amidino-2-thiourea standard solution (0.1 mg/mL) to 50-mL volumetric flasks step-by-step and add water up to the marked line.
   h) 1-amidino-2-thiourea standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation: In the case of usage, transfer 2.5 mL - 25 mL of 1-amidino-2-thiourea standard solution (20 µg/mL) for the calibration curve preparation to 50-mL volumetric flasks step-by-step and add water up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) A reagent of no less than 98 % (mass fraction) in purity is commercially sold as 1-amidino-2-thiourea.

Comment 1 1-amidino-2-thiourea is sold under the name guanylthiourea by Tokyo Chemical Industry Co., Ltd, and under the name amidino thiourea by Kanto Chemical Co., Inc.

(3) Instruments: Instruments are as shown below:
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube filled with silica gel, to which octadecyl chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 262 nm.
   b) Magnetic stirrer:
   c) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.
Comment 2 A column is sold under production names such as Inertsil ODS, Mightysil RP-18, L-column ODS, Shim-pack VP-ODS, Silica C18M 4D, Puresil C18, COSMOSIL 5C18-MS-II, etc.

(4) Test procedures
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.
   b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
   c) After allowing to stand still, transfer a supernatant solution to a 1.5 mL ground-in stopper centrifugal precipitate tube.
   d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes and use the supernatant as the extract.

Note (3) If there is a possibility that the 1-amidino-2-thiourea concentration in the sample solution exceeds the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

(4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(5) 7.2 cm - 8.9 cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

Comment 3 Instead of the procedures in (4.1) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:
   1) Column: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which octadecyl chemically bonds.
   2) Column bath temperature: 30 °C - 45 °C
   3) Eluent: Dissolve 0.94 g of sodium 1-hexasulfonic acid in 1000 mL of methanol water (2+8), adjust to pH 3.15 with acetic acid and filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.
   4) Flow rate: 1 mL/min
   5) Detection unit: An absorptiometric detector, measurement wavelength: 262 nm

b) Calibration curve preparation
   1) Inject 10 µL of respective 1-amidino-2-thiourea standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 262 nm and obtain the peak area or height.
   2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 262 nm of the respective 1-amidino-2-thiourea standard solutions for the calibration curve preparation.

c) Sample measurement
   1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain the 1-amidino-2-thiourea content from the calibration curve to calculate 1-amidino-2-thiourea (ASU) in the analytical sample.

Comment 4 A recovery testing with triplicates measurement was conducted using compound fertilizer (2 samples), as a result, the mean recovery rate of 1-amidino-2-thiourea at concentration level of 1.0 % (mass fraction), 0.5 % (mass fraction) and 0.25 % (mass fraction) were 99.0 % - 104.3 %, 97.7 % - 100.7 % and 99.7 % - 101.3 %.

Table 1 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of this testing method is about 0.005 % (mass fraction).

Table 1  Results and analysis results from a collaborative study for the test method validation of 1-amidino-2-thiourea (ASU)

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories (^{1)})</th>
<th>Mean (^{2)}) ((\text{%})^{3)})</th>
<th>(s_{r}^{4)}) ((\text{%})^{3)})</th>
<th>(RSD_{r}^{5)}) (\text{%})</th>
<th>(s_{R}^{6)}) ((\text{%})^{3)})</th>
<th>(RSD_{R}^{7)}) (\text{%})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>10</td>
<td>0.093</td>
<td>0.009</td>
<td>9.1</td>
<td>0.010</td>
<td>11.2</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>10</td>
<td>0.246</td>
<td>0.021</td>
<td>8.6</td>
<td>0.021</td>
<td>8.6</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>10</td>
<td>0.511</td>
<td>0.018</td>
<td>3.6</td>
<td>0.025</td>
<td>4.9</td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>10</td>
<td>0.759</td>
<td>0.039</td>
<td>5.1</td>
<td>0.040</td>
<td>5.3</td>
</tr>
<tr>
<td>Compound fertilizer 5</td>
<td>10</td>
<td>1.020</td>
<td>0.039</td>
<td>3.8</td>
<td>0.044</td>
<td>4.3</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean \((n = \text{number of laboratories} \times \text{number of samples (2)})\)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References


**Flow sheet for 1-amidino-2-thiourea:** The flow sheet for 1-amidino-2-thiourea in fertilizers is shown below:

- **1.00 g analytical sample**
- **200-mL ground-in stopper Erlenmeyer flask**
- ** ← 100 mL of water**
- **Extraction**
- **Stirring, 10 minutes**
- **Centrifugal separation**
- **Ground-in stopper centrifugal precipitate tube, 8000 × g - 10000 × g , 5 minutes**
- **Sample solution**
- **Measurement**
- **High Performance Liquid Chromatograph**

Reference diagram: HPLC chromatogram of 1-amidino-2-thiourea (ASU) standard solution

Reference: HPLC chromatogram of 1-amidino-2-thiourea (ASU) standard solution for the calibration curve preparation is shown below.

Measurement conditions for HPLC:
- Column: Mightysil RP-18 (4.6-mm inner diameter, 150-mm long, 5-μm particle diameter)
- 1-amidino-2-thiourea standard solution (the equivalents of 200 ng)
- Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
7.3  4-amino-1,2,4-triazole hydrochloride (ATC)
7.3.a High-Performance Liquid Chromatography

(1) **Summary**
This testing method is applicable to fertilizers containing 4-amino-1,2,4-triazole hydrochloride (ATC) but not containing organic matters. This testing method is classified as Type C and its symbol is 7.3.a-2017 or ATC.a-1.

Add methanol to an analytical sample to extract 4-amino-1,2,4-triazole hydrochloride, introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with an aminopropyl silica gel column, and measure at wavelength 220 nm to obtain 4-amino-1,2,4-triazole hydrochloride (ATC) in an analytical sample. In addition, the performance of this testing method is shown in **Comment 4**.

(2) **Reagent**: Reagents are as shown below.

a) **Methanol**: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.

b) **Methanol**: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.

c) **Acetonitrile**: Acetonitrile used in eluent of an HPLC is a reagent of HPLC analysis grade or equivalents.

d) **4-amino-1,2,4-triazole standard solution (1 mg/mL)**: Transfer 0.1 g of 4-amino-1,2,4-triazole [C₂H₄N₄] to a weighing dish and measure the mass to the order of 0.1 mg. Add methanol to dissolve, transfer to a 100-mL amber volumetric flask and add methanol to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.

e) **4-amino-1,2,4-triazole standard solution (0.1 mg/mL)**: In the case of usage, transfer 10 mL of 4-amino-1,2,4-triazole standard solution (1 mg/mL) to a 100-mL volumetric flask and add methanol up to the marked line.

f) **4-amino-1,2,4-triazole standard solution (10 µg/mL - 50 µg/mL) for the calibration curve preparation**: In the case of usage, transfer 5 mL - 25 mL of 4-amino-1,2,4-triazole standard solution (0.1 mg/mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.

g) **4-amino-1,2,4-triazole standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation**: In the case of usage, transfer 2.5 mL - 25 mL of 4-amino-1,2,4-triazole standard solution for the calibration curve preparation (20 µg/mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.

**Note**
(1) The solution contains 1.434 mg/mL as 4-amino-1,2,4-triazole hydrochloride.
(2) This is an example of preparation; prepare an amount as appropriate.
(3) A reagent of no less than 98 % (mass fraction) in purity is commercially sold as 4-amino-1,2,4-triazole.

**Comment 1** 4-amino-1,2,4-triazole is sold under production names such as 4-amino-1,2,4-triazole by FUJIFILM Wako Pure Chemical Co., Ltd., and Tokyo Chemical Industry Co., Ltd, and 4-amino-4H-1,2,4-triazole by Kanto Chemical Co., Inc.

(3) **Instruments**: Instruments are as shown below:

a) **High-Performance Liquid Chromatograph (HPLC)**: HPLC specified in JIS K 0124 that satisfies following requirements.

1) **Column**: A column of 4 mm - 6 mm inner diameter and 150 mm - 250 mm long stainless steel column tube filled with silica gel, to which amino or amino propyl chemically bonds.
2) **Column bath:** A column bath whose temperature can be adjusted to 30 °C - 45 ºC.

3) **Detection unit:** An absorptiometric detector that can measure at wavelength around 220 nm.

b) **Magnetic stirrer**

c) **High speed centrifugal separator:** A centrifugal separator that can work at 8000 × g - 10000 × g.

**Comment 2** A column is sold under production names such as Hibar LiChrosorb NH₂, Inertsil NH₂, Unison UK-Amino, Mightysil NH₂, Shim-pack CLC-NH₂, Shodex NH-5A, Unisil Q NH₂, etc.

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL- ground-in stopper Erlenmeyer flask.

b) Add 100 mL of methanol and stir it with using a magnetic stirrer for about 10 minutes.

c) After allowing to stand still, transfer a supernatant solution to a 1.5 mL- ground-in stopper centrifugal precipitate tube.

d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes and use the supernatant as the extract.

**Note** (4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

**Comment 3** Instead of the procedures in (4.1) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of PTFE and the filtrate can be the sample solution.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC):** An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

1) **Column:** A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino or amino propyl chemically bonds.

2) **Column bath temperature:** 30 °C - 40 °C

3) **Eluent:** Acetonitrile - methanol (9+1)

4) **Flow rate:** 1 mL/min

5) **Detection unit:** An absorptiometric detector, measurement wavelength: 220 nm

b) **Calibration curve preparation**

1) Inject 10 µL of respective 4-amino-1,2,4-triazole standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 220 nm and obtain the peak area or height.

2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 220 nm of the 4-amino-1,2,4-triazole standard solutions for the respective calibration curve preparation.

c) **Sample measurement**

1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain the amount of 4-amino-1,2,4-triazole from the calibration curve to calculate 4-amino-1,2,4-triazole in the analytical sample.

3) Calculate the 4-amino-1,2,4-triazole hydrochloride (ATC) by the following formula.

\[
\text{4-amino-1,2,4-triazole hydrochloride in the analytical sample (\% (mass fraction))} = A \times 1.434
\]

\(A\): 4-amino-1,2,4-triazole hydrochloride in the analytical sample (\% (mass fraction))

**Comment 4** A recovery testing was conducted using compound fertilizer (2 samples), as a result, the mean recovery rate of 4-amino-1,2,4-triazole hydrochloride at concentration level of 0.5 % (mass fraction), 0.3 % (mass fraction) and 0.2 % (mass fraction) were 100.2 % - 104.9 %, 100.8 % - 103.0 % and 100.7 % - 104.2 %.

Additionally, the minimum limit of quantification of this testing method is about 0.005 % (mass fraction).

**References**


**Flow sheet for 4-amino-1,2,4-triazole hydrochloride (ATC):** The flow sheet for 4-amino-1,2,4-triazole hydrochloride in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Extraction</th>
<th>1.00 g analytical sample ← 100 mL of Methanol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Centrifugal separation</td>
<td>Stirring, 10 minutes</td>
</tr>
<tr>
<td></td>
<td>Ground-in stopper centrifugal precipitate tube, 8000 × g - 10000 × g , 5 minutes</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
<tr>
<td>Measurement</td>
<td>High Performance Liquid Chromatograph</td>
</tr>
</tbody>
</table>

Figure Flow sheet for 4-amino-1,2,4-triazole hydrochloride (ATC) in fertilizers
7.4 N-2,5-dichlorophenyl succinamic acid (DCS)
7.4.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers containing N-2,5-dichlorophenyl succinamic acid (DCS) but not containing organic matters. This testing method is classified as Type C and its symbol is 7.4.a-2017 or DCS.a.

Add methanol - phosphoric acid (996+4) and water to an analytical sample to extract N-2,5-dichlorophenyl succinamic acid (DCS), introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with octadecyl silylation silica gel column, and measure at wavelength 246 nm to obtain N-2,5-dichlorophenyl succinamic acid (DCS) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent, etc.: Reagents and water are as shown below.


b) Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.

c) Methanol: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.

d) Phosphoric acid: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.

e) N-2,5-dichlorophenyl succinamic acid standard solution (1 mg/mL) (1): Transfer 0.1 g of N-2,5-dichlorophenyl succinamic acid \([C_{10}H_8Cl_2NO_3]\) to a weighing dish and measure the mass to the order of 0.1 mg. Add methanol to dissolve, transfer to a 100-mL volumetric flask and add methanol to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.

g) N-2,5-dichlorophenyl succinamic acid standard solution (0.1 mg/mL): In the case of usage, transfer 10 mL of N-2,5-dichlorophenyl succinamic acid standard solution (1 mg/mL) to a 100-mL volumetric flask and add methanol up to the marked line.

h) N-2,5-dichlorophenyl succinamic acid standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation: In the case of usage, transfer 2.5 mL - 25 mL of N-2,5-dichlorophenyl succinamic acid standard solution (0.1 mg/mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.

(3) Instruments: Instruments are as shown below:

a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.

1) Column: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube filled with silica gel, to which octadecyl chemically bonds.

2) Column bath: A column bath whose temperature can be adjusted to 30 °C - 45 °C.

3) Detection unit: An absorptiometrical detector that can measure at wavelength around 246 nm.

b) Magnetic stirrer

c) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.
Comment 1 A column is sold under production names such as Inertsil ODS, Mightysil RP-18, L-column ODS, Shim-pack VP-ODS, Silica C18M 4D, Puresil C_{18}, COSMOSIL 5C18-MS-II, etc.

(4) Test procedures

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.

b) Add 100 mL of methanol - phosphate (996+4) and stir it with a magnetic stirrer for about 30 minutes.

c) After allowing to stand still, transfer a supernatant solution \(^{(2)}\) to a 1.5-mL ground-in stopper centrifugal precipitate tube \(^{(3)}\).

d) Centrifuge it at 8000 \(\times\) g - 10000 \(\times\) g centrifugal force for about five minutes \(^{(4)}\) and use the supernatant as the extract.

Note \((2)\) If there is a possibility that the N-2,5-dichlorophenyl succinamic acid concentration in the sample solution exceeds the maximum limit of the calibration curve, dilute a predetermined amount of the outflow solution with methanol.

\(\text{(3)}\) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement

\(\text{(4)}\) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 \(\times\) g - 10000 \(\times\) g centrifugal force.

Comment 2 Instead of the procedures in (4.1) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-\(\mu\)m) made of PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

1) Column: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-\(\mu\)m particle diameter) to which octadecyl chemically bonds.

2) Column bath temperature: 30 °C - 40 °C

3) Eluent: Methanol - water \(^{(5)}\) (55 + 45)

4) Flow rate: 0.8 mL/min

5) Detection unit: An absorptiometric detector, measurement wavelength: 246 nm

Note \((5)\) Adjust the water used to pH 3 with phosphoric acid in advance.

b) Calibration curve preparation

1) Inject 10 \(\mu\)L of respective N-2,5-dichlorophenyl succinamic acid standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 246 nm and obtain the peak area or height.

2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 246 nm of the respective N-2,5-dichlorophenyl succinamic acid standard solution for the calibration curve preparation.

c) Sample measurement
1) Subject 10 μL of sample solution to the same procedure as in (a) 1)
2) Obtain the N-2,5-dichlorophenyl succinic acid content from the calibration curve to calculate N-2,5-dichlorophenyl succinic acid (DCS) in the analytical sample.

Comment 3 Recovery testing was conducted using compound fertilizer (2 samples) and blended fertilizer (1 sample), as a result, the mean recovery rate of N-2,5-dichlorophenyl succinic acid at concentration level of 0.4 % (mass fraction), 0.2 % (mass fraction) and 0.1 % (mass fraction) were 100.9 % - 101.4 %, 100.8 % - 101.4 % and 101.2 % - 103.4 %.
Additionally, the minimum limit of quantification of this testing method is about 0.005 % (mass fraction).

References

(5) Flow sheet for N-2,5-dichlorophenyl succinic acid (DCS): The flow sheet for N-2,5-dichlorophenyl succinic acid (DCS) in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Process</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00 g analytical sample</td>
<td>200-mL ground-in stopper Erlenmeyer flask ← 100 mL of methanol phosphoric acid (996+4)</td>
</tr>
<tr>
<td>Extraction</td>
<td>Stirring, 30 minutes</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Ground-in stopper centrifugal precipitate tube, 8000 × g - 10000 × g, 5 minutes</td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
<tr>
<td>Measurement</td>
<td>High Performance Liquid Chromatograph</td>
</tr>
</tbody>
</table>

Figure Flow sheet for N-2,5-dichlorophenyl succinic acid (DCS) in fertilizers.
Reference: HPLC chromatogram of N-2,5-dichlorophenyl succinamic acid (DCS) standard solution for the calibration curve preparation is shown below.

Reference diagram: HPLC chromatogram of N-2,5-dichlorophenyl succinamic acid

Measurement conditions for HPLC
- Column: Mightysil RP-18 (4.6-mm inner diameter, 150-mm long, 5-µm particle diameter)
- N-2,5-dichlorophenyl succinamic acid standard solution (the equivalents of 100 ng)
- Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
7.5  Dicyandiamide (Dd)
7.5.a  High-Performance Liquid Chromatography
(1)  Summary
This testing method is applicable to fertilizers containing dicyandiamide (Dd). This testing method is classified as Type B and its symbol is 7.5.a-2017 or Dd.a-1. Add water to an analytical sample, leave at rest for a little while and add methanol to extract dicyandiamide. After removing interfering substances with a silica gel cartridge column, introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with an aminopropyl silica gel column, and measure at wavelength 215 nm to obtain dicyandiamide (Dd) in an analytical sample. In addition, the performance of this testing method is shown in Comment 5.

(2)  Reagent, etc.: Reagents and water are as shown below.
  b)  Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.
  c)  Methanol: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.
  d)  Acetonitrile: A reagent of HPLC grade or equivalents.
  e)  Dicyandiamide standard solution (1 mg/mL) (1): Put 0.1 g of dicyandiamide \( \text{C}_2\text{H}_4\text{N}_4 \) (2) to a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of methanol to dissolve, transfer to a 100-mL volumetric flask and add the solvent up to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.
  f)  Dicyandiamide standard solution (0.1 mg/mL): At the time of usage, put 10 mL of dicyandiamide standard solution (1 mg/ mL) to a 100-mL volumetric flask and add methanol up to the marked line.
  g)  Dicyandiamide standard solution (10 µg/ mL - 50 µg/ mL) for the calibration curve preparation: At the time of usage, put 5 mL - 25 mL of dicyandiamide standard solution (0.1 mg/ mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.
  h)  Dicyandiamide standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation: In the case of usage, transfer 2.5 mL - 25 mL of dicyandiamide standard solution for the calibration curve preparation (20 µg/mL) to 50-mL volumetric flasks step-by-step and add methanol up to the marked line.
  i)  Sodium sulfate: A JIS Guaranteed Reagent specified in JIS K 8987 or a reagent of equivalent quality.

Note  (1) This is an example of preparation; prepare an amount as appropriate.
  (2) A reagent of no less than 98 % in purity is commercially sold as dicyandiamide.

Comment 1  Dicyandiamide is commercially sold as dicyanodiamide by FUJIFILM Wako Pure Chemical Co., Ltd. and Kanto Chemical Co., Inc.

(3)  Instruments: Instruments are as shown below:
  a)  High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
    1)  Column: A column of 4 mm - 6 mm inner diameter and 150 mm - 250 mm long stainless steel column tube filled with silica gel, to which amino or amino propyl chemically bonds.
    2)  Column bath: A column bath whose temperature can be adjusted to 30 ℃ - 40 ℃.
    3)  Detection unit: An absorptiometric detector that can measure at wavelength around 215 nm.
  b)  Shaking apparatus
  c)  Centrifugal separator: A centrifugal separator that can work at about 1700 × g.
d) **High speed centrifugal separator**: A centrifugal separator that can work at 8000 × g - 10000 × g.

e) **Silica gel cartridge column**: Link a 10-mL cylinder to the column (3) filled with 500 mg - 1 g of silica gel, add 3 mL of methanol to let it flow down.

**Note** (3) A cartridge with a 3-mL - 6-mL column filled with 500 mg – 1 g of silica gel can be used.

**Comment 2** A column is sold under production names such as Hibar LiChrosorb NH₂, Inertsil NH₂, Unison UK-Amino, Mightysil NH₂, Shim-pack CLC-NH₂, Shodex NH-5A, Unisil Q NH₂, etc.

**Comment 3** A silica gel cartridge column is commercially sold under production names such as Sep-Pak Plus Silica, InertSep Si.

(4) **Test procedures**

4.1 **Extraction**: Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.

b) Add 1 mL of water (4) and leave at rest for 5 minutes.

c) Add 100 mL of methanol and shake to mix with a shaking apparatus for about 10 minutes.

d) Add an appropriate amount of sodium sulfate (5).

e) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.

f) Centrifuge it at about 1700 × g centrifugal force for about five minutes (6) and use the supernatant (7) as the extract.

**Note** (4) Mix well until the whole sample comes in contact with water.

(5) About 5 g - 10 g.

(6) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

(7) If there is a possibility that the concentration of dicyandiamide in the sample solution exceeds the maximum limit of the calibration curve, dilute a predetermined amount of extract with methanol.

4.2 **Cleanup**: Conduct cleanup as shown below:

a) Transfer extract to a silica gel cartridge column.

b) Dispose of about the first 3 mL of effluent and then transfer about the next 2 mL to a test tube.

c) Transfer the effluent to a 1.5-mL ground-in stopper centrifugal precipitate tube (8).

d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (9) and use the supernatant as the extract.

**Note** (8) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(9) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

**Comment 4** Instead of the procedures in (4.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of PTFE and the filtrate can be the sample solution.
(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

1) **Column**: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter) to which amino or amino propyl chemically bonds.

2) **Column bath temperature**: 30 °C - 40 °C

3) **Eluent**: Acetonitrile - methanol (6+1)

4) **Flow rate**: 0.5 mL/min - 1 mL/min

5) **Detection unit**: An absorptiometric detector, measurement wavelength: 215 nm

b) **Calibration curve preparation**

1) Inject 10 µL of respective dicyandiamide standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 215 nm, and obtain the peak area or the height.

2) Prepare a curve for the relationship between the concentration and the peak area or the height at wavelength 215 nm of the respective dicyandiamide standard solutions for the calibration curve preparation.

c) **Sample measurement**

1) Subject 10 µL of sample solution to the same procedure as in b) 1)

2) Obtain dicyandiamide content from the calibration curve to calculate dicyandiamide (Dd) in the analytical sample.

**Comment 5** Recovery testing was conducted using inorganic compound fertilizer (2 samples) and organic compound fertilizer (3 samples), as a result, the mean recovery at the concentration level of 2 % (mass fraction) and 0.2% (mass fraction) were 101.2 % - 102.6 % and 98.4 % - 100.6 %.

Table 1 shows results and analysis results from a collaborative study for test method validation. Additionally, the minimum limit of quantification of this testing method is about 0.01 % (mass fraction).
Table 1  Results and analysis results from a collaborative study for the test method validation of dicyandiamide

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (^2)</th>
<th>(s_r) (^4)</th>
<th>(RSD_r) (^5)</th>
<th>(s_R) (^6)</th>
<th>(RSD_R) (^7)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>11</td>
<td>0.263</td>
<td>0.009</td>
<td>3.2</td>
<td>0.019</td>
<td>7.4</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>11</td>
<td>2.04</td>
<td>0.04</td>
<td>1.7</td>
<td>0.07</td>
<td>3.2</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>13</td>
<td>0.548</td>
<td>0.011</td>
<td>2.0</td>
<td>0.033</td>
<td>6.0</td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>12</td>
<td>0.423</td>
<td>0.013</td>
<td>3.2</td>
<td>0.022</td>
<td>5.2</td>
</tr>
<tr>
<td>Compound fertilizer 5</td>
<td>12</td>
<td>1.02</td>
<td>0.01</td>
<td>1.4</td>
<td>0.04</td>
<td>4.3</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Mean \((n = \text{number of laboratories} \times \text{number of samples (2)})\)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Reproducibility standard deviation
7) Reproducibility relative standard deviation

References
(5) **Flow sheet for dicyandiamide:** The flow sheet for dicyandiamide in fertilizers is shown

<table>
<thead>
<tr>
<th>1.00 g analytical sample</th>
<th>200-mL ground-in stopper Erlenmeyer flask</th>
</tr>
</thead>
<tbody>
<tr>
<td>← 1 mL of water</td>
<td>Leave at rest 5 minutes</td>
</tr>
<tr>
<td>← 100 mL of methanol</td>
<td>Shaking 10 minutes</td>
</tr>
<tr>
<td>← An adequate amount of sodium sulfate</td>
<td>Ground-in stopper centrifugal precipitate tube, 1700 ( \times g ), 5 minutes</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Cleanup Silica gel cartridge column</td>
</tr>
<tr>
<td></td>
<td>Centrifugal separation 1.5-mL ground-in stopper centrifugal precipitate tube, 8000 ( \times g ) - 10000 ( \times g ), 5 minutes</td>
</tr>
<tr>
<td></td>
<td>Sample solution</td>
</tr>
<tr>
<td></td>
<td>Measurement High Performance Liquid Chromatograph</td>
</tr>
</tbody>
</table>

Figure Flow sheet for dicyandiamide (Dd) in fertilizers
Reference: HPLC chromatogram of dicyandiamide standard solution for the calibration curve preparation and sample solution (compound fertilizer) are shown below.

1) Standard solution

2) Sample solution

Reference diagram

HPLC chromatogram of dicyandiamide (Dd)

1) Dicyandiamide standard solution (the equivalents of 100 ng (10 µg/mL, 10 µL) of dicyandiamide)
2) Sample solution (compound fertilizer)

Measurement conditions for HPLC
Column: Inertsil NH₂ (4.6-mm inner diameter, 250-mm long, 5-µm particle diameter)
Column oven temperature: 30 °C
Flow rate: 0.5 mL/min
Other conditions are according to the example of HPLC measurement conditions in (4.3) a)
7.6 2-sulfanilamide thiazole (ST)
7.6.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers containing 2-sulfanilamide thiazole (ST). This testing method is classified as Type C and its symbol is 7.6.a-2017 or ST.a-1.

Add methanol–water (1+1) to an analytical sample to extract 2-sulfanilamide thiazole, introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with an octadecyl silylation silica gel column, and measure at wavelength 285 nm to obtain 2-sulfanilamide thiazole (ST) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.
   c) Methanol: Methanol used in eluent of an HPLC is HPLC analysis grade or a reagent of equivalent quality.
   d) 2-sulfanilamide thiazole standard solution (1 mg/mL): Transfer 0.1 g of 2-sulfanilamide thiazole \([\text{C}_9\text{H}_9\text{N}_3\text{O}_2\text{S}_2]\) to a weighing dish and measure the mass to the order of 0.1 mg. Add water to dissolve, transfer to a 1000-mL volumetric flask and add methanol–water (1+1) to the marked line. Store in a refrigerator, and do not use after 6 months after preparation.
   e) 2-sulfanilamide thiazole standard solution (0.1 mg/mL): In the case of usage, transfer 10 mL of 2-sulfanilamide thiazole standard solution (100 µg/mL) to a 100-mL volumetric flask and add methanol–water (1+1) up to the marked line.
   f) 2-sulfanilamide thiazole standard solution (10 µg/mL - 50 µg/mL) for the calibration curve preparation: In the case of usage, transfer 5 mL - 25 mL of 2-sulfanilamide thiazole standard solution (0.1 mg/mL) to 50-mL volumetric flasks step-by-step and add methanol–water (1+1) up to the marked line.
   g) 2-sulfanilamide thiazole standard solution (1 µg/mL - 10 µg/mL) for the calibration curve preparation: In the case of usage, transfer 2.5 mL - 25 mL of 2-sulfanilamide thiazole standard solution for the calibration curve preparation (20 µg/mL) to 50-mL volumetric flasks step-by-step and add methanol–water (1+1) up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
   (2) A reagent of no less than 98 % (mass fraction) in purity is commercially sold as 2-sulfanilamide thiazole.

Comment 1 2-sulfanilamide thiazole is sold under the production name sulfathiazole by Tokyo Chemical Industry Co., Ltd, FUJIFILM Wako Pure Chemical Co., Ltd., and Kanto Chemical Co., Inc.

(3) Instruments: Instruments are as shown below:
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube filled with silica gel, to which octadecyl chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 285 nm.
   b) Magnetic stirrer:
   c) Centrifugal separator: A centrifugal separator that can work at about 1700 × g.
   d) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g -
10000 × g.
e) **Acidic alumina cartridge column:** Link a 10-mL cylinder to a column (3) that is filled with 500 mg - 1 g of acidic alumina, put 3 mL of methanol and let it flow down.

**Note** (3) A cartridge with a 3-mL - 6-mL column filled with 500 mg – 1 g of silica gel can be used.

**Comment 2** A column is sold under production names such as Inertsil ODS, Mighty sil RP-18, L-column ODS, Shim-pack VP-ODS, Silica C18M 4D, Puresil C18, COSMOSIL 5C18-MS-II, etc.

**Comment 3** An acidic alumina cartridge is commercially sold under production names such as Bond Elut AL-A, Sep-Pak Alumina-A, Supelclean LC-Alumina-A.

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below.
a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL- ground-in stopper Erlenmeyer flask.
b) Add 100 mL of methanol - water (1+1) and stir it by using a magnetic stirrer for about 15 minutes.
c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at 1700 × g centrifugal force for about five minutes (4) and use the supernatant (5) as the extract.

**Note** (4) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

(5) If there is a possibility that the 2-sulfanilamide thiazole concentration in the sample solution exceeds the maximum limit of the calibration curve, dilute a predetermined amount of the extract with methanol.

(4.2) **Cleanup:** Conduct cleanup as shown below:
a) Transfer the extract to an acidic alumina cartridge column.
b) Dispose of about the first 3 mL of effluent and then transfer about the next 2 mL to a test tube.
c) Transfer the effluent to a 1.5-mL ground-in stopper centrifugal precipitate tube (6).
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (7) and use the supernatant as the extract.

**Note** (6) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(7) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

**Comment 4** Instead of the procedures in (4.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of PTFE and the filtrate can be the sample solution.

**Comment 5** The test is possible by the following procedures in the case of fertilizers not containing organic matters.
The procedures in (4.1) c) - d) and (4.2) a) - b) are omitted and “Transfer effluent” in (4.2) c) is replaced with the “After allowing to stand still, transfer supernatant” to
operate.

(4.3) **Measurement**: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

1) **Column**: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-μm particle diameter) to which octadecyl chemically bonds.

2) **Column bath temperature**: 30 °C - 40 °C

3) **Eluent**: Methanol–water (2+8)

4) **Flow rate**: 1 mL/min

5) **Detection unit**: An absorptiometric detector, measurement wavelength: 285 nm

b) **Calibration curve preparation**

1) Inject 10 μL of respective 2-sulfanilamide thiazole standard solutions for the calibration curve preparation to an HPLC, record a chromatogram at wavelength 285 nm and obtain the peak area or height.

2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 285 nm of the respective 2-sulfanilamide thiazole standard solutions for the calibration curve preparation.

c) **Sample measurement**

1) Subject 10 μL of sample solution to the same procedure as in b) 1)

2) Obtain the 2-sulfanilamide thiazole content from the calibration curve to calculate 2-sulfanilamide thiazole (ST) in the analytical sample.

**Comment 6** Recovery testing was conducted using compound fertilizer (1 sample) and blended fertilizer (2 sample), as a result, the mean recovery rate of 2-sulfanilamide thiazole at the concentration level of 1.0 % (mass fraction), 0.4 % (mass fraction) and 0.1 % (mass fraction) were 101.2 % - 102.1 %, 99.6 % - 101.7 % and 99.4 % - 101.0 %.

Additionally, the minimum limit of quantification of this testing method is about 0.005 % (mass fraction).

**References**

(5) **Flow sheet for 2-sulfanilamide thiazole**: The flow sheet for 2-sulfanilamide thiazole in fertilizers is shown below:

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00 g analytical sample</td>
<td>200-mL ground-in stopper Erlenmeyer flask</td>
</tr>
<tr>
<td>Extraction</td>
<td>100 mL of methanol - water (1+1)</td>
</tr>
<tr>
<td>Stirring, 15 minutes</td>
<td></td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Ground-in stopper centrifugal precipitate tube,</td>
</tr>
<tr>
<td></td>
<td>1700 × g, 5 minutes</td>
</tr>
<tr>
<td>Cleanup</td>
<td>Acidic alumina cartridge column</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Ground-in stopper centrifugal precipitate tube,</td>
</tr>
<tr>
<td></td>
<td>8000 × g - 10000 × g, 5 minutes</td>
</tr>
<tr>
<td>Sample solution</td>
<td>High Performance Liquid Chromatograph</td>
</tr>
</tbody>
</table>

Figure Flow sheet for 2-sulfanilamide thiazole (ST) in fertilizers.

**Reference**: HPLC chromatogram of 2-sulfanilamide thiazole (ST) standard solution for the calibration curve preparation is shown below.

Reference diagram: HPLC chromatogram of 2-sulfanilamide thiazole (ST)

Measurement conditions for HPLC
- **Column**: Mightysil RP-18 (4.6-mm inner diameter, 150-mm long, 5-µm particle diameter)
- 2-sulfanilamide thiazole standard solution (the equivalents of 200 ng)
- Other conditions are according to the example of HPLC measurement conditions in (4.3) a)
8. Others
8.1 Melamine and its degradation products
8.1.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 8.1.a-2017 or Mel.a-1.
Extract melamine and its derivative substances (hereinafter referred to as “melamine derivations”) in organic matters and fertilizers containing organic matters with diethylamine–water–acetonitrile (1+4+5) and derivatize with BSTFA–TMCS (99+1) and then measure with a gas chromatography/mass spectrometer to obtain melamine deviations in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

Comment 1 The structural formula of melamine and its degradation products is shown in the figure 1. During the production process of melamine, a by-product that replaces “-NH\textsubscript{2}” of R1 - R3 with “-OH” is formed in some cases.

![Figure 1 Structural formula of melamine and its degradation products](image)

<table>
<thead>
<tr>
<th></th>
<th>R\textsubscript{1}</th>
<th>R\textsubscript{2}</th>
<th>R\textsubscript{3}</th>
<th>MW</th>
</tr>
</thead>
<tbody>
<tr>
<td>Melamine</td>
<td>NH\textsubscript{2}</td>
<td>NH\textsubscript{2}</td>
<td>NH\textsubscript{2}</td>
<td>126.12</td>
</tr>
<tr>
<td>Ammeline</td>
<td>OH</td>
<td>NH\textsubscript{2}</td>
<td>NH\textsubscript{2}</td>
<td>127.10</td>
</tr>
<tr>
<td>Ammelide</td>
<td>OH</td>
<td>OH</td>
<td>NH\textsubscript{2}</td>
<td>128.09</td>
</tr>
<tr>
<td>Cyanuric acid</td>
<td>OH</td>
<td>OH</td>
<td>OH</td>
<td>129.07</td>
</tr>
</tbody>
</table>

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Acetonitrile: A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8039 or a reagent of equivalent quality.
   c) Diethylamine: A guaranteed reagent or a reagent of equivalent quality.
   d) Pyridine (dehydration) (1): A reagent of organic synthesis grade of no less than 99.5 % (mass fraction) in purity and no more than 0.05 mg/mL in moisture or a reagent of equivalent quality.
   e) Derivatization reagent (2): Bis (trimethylsilyl) trifluoroacetamide–trimethylchlorosilane (99+1).
   f) Melamine derivations standard solution (0.5 mg/mL): Put about 0.05 g of melamine [C\textsubscript{3}H\textsubscript{6}N\textsubscript{6}] (3), ammeline [C\textsubscript{3}H\textsubscript{8}N\textsubscript{3}O] (3), ammellide [C\textsubscript{3}H\textsubscript{4}N\textsubscript{4}O\textsubscript{2}] (3) and cyanuric acid [C\textsubscript{3}H\textsubscript{3}N\textsubscript{3}O\textsubscript{3}] (3) into a weighing dish and measure the mass to the order of 0.1 mg. Dissolve in a small amount of diethylamine–water (1+4), transfer to a 100 mL volumetric flask respectively, and add the solvent up to the marked line.
   g) Mixture standard solution (50 µg/mL) (3): Put 5 mL of respective melamine derivations standard solutions (0.5 mg/mL) to 50-mL volumetric flasks and add diethylamine–water–acetonitrile (1+4+5) up to the marked line.

Note (1) After it is opened once, add a proper amount of sodium sulfate (anhydrous) and seal tightly to store.
(2) A mixed derivatization reagent is commercially sold under the name BSTFA–TMCS (99+1).
(3) The respective standard reagents of melamine, ammeline, ammelide and cyanuric acid

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are commercially sold.

**Comment 2** BSTFA–TMCS (99+1) is sold as 1-mL ampoule by SUPELCO. After it is opened once, use it on the same day.

**Comment 3** The standard reagent of melamine, ammeline, ammelide and cyanuric acid are sold by FUJIFILM Wako Pure Chemical Co., Ltd. and Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(3) **Instruments:** Instruments are as shown below:

a) **Gas Chromatograph/Mass Spectrometer (GC/MS):** GC/MS specified in JIS K 0123 that satisfies the following requirements.

1) **Gas Chromatograph**
   - (i) Sample injector: An injector that enables split less system.
   - (ii) Capillary column: A capillary column (0.25-mm - 0.32-mm inner diameter and 30-m long) made of fused silica. 5 % phenyl 95 % methyl polysiloxane chemically bonds to the inner surface of a capillary column with 0.25 µm thickness. The column is according to the specification of mass spectrometer.
   - (ii) Carrier gas: High purity helium of no less than 99.999 % (volume fraction) in purity.

2) **Mass Spectrometer**
   - (i) Ionization method: Electron-Impact ionization (EI) method
   - (ii) Ion detection method: Selected Ion Monitoring (SIM) method

b) **Ultrasonic generator:** An ultrasonic washer can be used.

c) **High speed centrifugal separator:** A centrifugal separator that can work at 8000 × g - 10000 × g.

d) **Concentrator:** A Centrifugal evaporator that can be adjusted to 70 ºC ± 2 ºC.

e) **Water bath:** Water bath that can be adjusted to 70 ºC ± 2 ºC.

**Comment 4** A capillary column is commercially sold under the names such as DB-5ms, Rtx-5ms, HP-5ms, SLB-5ms, BPX-5, CP-Sil 8CB low Bleed/MS and TC -5HT for GC/MS.

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 0.50 g of an analytical sample, and put it into a 200 mL - 200-mL ground-in stopper Erlenmeyer flask.

b) Add 160 mL - 200 mL of diethylamine–water–acetonitrile (1+4+5), and subject to ultra-sonication for about 30 minutes using an ultrasonic generator.

c) Transfer about 1.5 mL to a 1.5-mL ground-in stopper centrifugal precipitate tube (4), and centrifuge at 8000 × g - 10000 × g for about 5 minutes (5).

d) Transfer 1 mL of the supernatant solution to 5-mL - 50-mL volumetric flasks, add diethylamine–water–acetonitrile (1+4+5) up to the marked line to make the extract.

**Note** (4) Confirm that the tube is made of polypropylene, etc. to not affect testing results.

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

**Comment 5** Grind until it completely passes through a sieve of 500 µm aperture to prepare the test sample.

**Comment 6** Weigh 0.5 g of an analytical sample, extract with 200 mL of diethylamine–water–acetonitrile (1+4+5). If it is diluted 50 times in the procedure in d), the quantitative range of melamine derivations in the analytical sample is 0.2 %
(mass fraction) - 10 % (mass fraction). In the case of measuring melamine derivations which do not reach the quantitative range, make the dilution factor in the procedure in d) decrease. In addition, if the contents of melamine derivations exceed 10 % (mass fraction) respectively, the sampling volume of an analytical sample should be reduced.

(4.2) **Derivatization:** Conduct derivatization as shown below.

a) Transfer 0.2 mL of the extract to a 5-mL - 10-mL test tube with a screw stopper.

b) Place a test tube in a concentrator, concentrate under reduced pressure at 70 °C ± 2 °C and vaporize the solvent completely (6).

c) Add 0.3 mL of pyridine (dehydration) (1) and 0.2 mL of derivatization reagent (2) to the residue to mix, and then stopple to seal tightly.

d) After heating in a water bath at 70 °C ± 2 °C for about 45 minutes (7), let it stand to cool to make a sample solution (8).

Note (6) A spraying type concentrator can be used.

(7) If moisture remains after the procedure in b) or the reagent used in the procedure in c) contains moisture, the reaction of the derivatization in d) does not advance enough in some cases.

(8) If necessary, transfer the sample solution into a 1.5-mL ground-in stopper centrifugal precipitate tube (4) to centrifuge at 8000 × g - 10000 × g for about 5 minutes (5).

(4.3) **Measurement:** Conduct the measurement as indicated in JIS K 0123 and as shown below. Specific measurement procedures are according to the operation method of the gas chromatograph/ mass spectrometer used in measurement.

a) **Measurement conditions for the gas chromatograph/ mass spectrometer:** Set up the measurement conditions considering it:

1) **Gas Chromatograph**

(i) Sample injection method: split less injection method (1min)

(ii) Temperature of sample injector: 280 °C

(iii) Capillary column: A capillary column (0.25-mm - 0.32-mm inner diameter, 30-m long, 0.25 µm layer thickness) made of fused silica. 5 % phenyl 95 % methyl polysiloxane chemically bonds to the inner surface of the capillary column.

(iv) Temperature of column bath: 100 °C (1 min) → (15 °C /min) → 320 °C (3 min)

(v) Temperature of GC/MS coupling portion: 250 °C

(vi) Carrier gas: helium, flow rate: 1.5 mL/min

2) **Mass Spectrometer**

(i) Ionization method: Electron-Impact ionization (EI) method

(ii) Ionization voltage: 70 V

(iii) Temperature of ion source: 230 °C

(iv) Ion detection method: Selected Ion Monitoring (SIM) method

(v) Measurement of ion: Shown in table 1
b) **Calibration curve preparation**

1) Transfer 5 mL of mixture standard solution (50 µg/mL) to a 50-mL volumetric flask and add diethylamine–water–acetonitrile (1+4+5) up to the marked line to make the mixture standard solution (5 µg/mL).

2) Transfer 1 mL - 20 mL of the mixture standard solution (5 µg/mL) to 50 mL volumetric flasks step-by-step and add diethylamine–water–acetonitrile (1+4+5) up to the marked line to make the mixture standard solution (0.1 µg/mL - 2 µg/mL).

3) Conduct the procedures of (4.2) b) - d) for the mixture standard solution (0.1 µg/mL - 2 µg/mL) to make the mixture standard solution, the equivalents of 0.04 µg/mL - 0.8 µg/mL, for calibration curve.

4) Inject 1 µL of respective mixture standard solutions for calibration curve to GC/MS and record the chromatogram of ion (m/z) for determination and ion (m/z) for validation of materials subjected to measurement to obtain the peak area or height.

5) Calculate the peak area ratio or height ratio of ion (m/z) for determination and ion (m/z) for validation of respective materials subjected to measurement.

6) Prepare a curve for the relationship between the concentration of material subjected to measurement and the peak area or height of ion (m/z) for determination of respective mixtures for the calibration curve preparation.

c) **Sample measurement**

1) Subject 1 µL of the sample solution to the same procedure as in b) 4) - 5) (9).

2) Obtain the content of respective materials subjected to measurement from the calibration curve to calculate respective materials subjected to measurement in the analytical sample.

**Note** (9) Confirm that the ratio against the peak area ratio or height ratio of the standard solution is within the range of about ± 30 %. In addition, the peak area ratio or height ratio depends on the concentration.

**Comment 7** If the variation of sensitivity of melamine derivations is observed, conduct measurement by the following method of a) or b).

a) In the procedure in (4.3) c) 1), inject the sample solution into GC/MS predetermined times, and then correct the calibration curve according to (4.3) b) 4) - 6).

b) Add 2,6-diamino-4-chloropyrimidine (the equivalents of 0.5 µg) as an internal reference material to the sample solution and conduct the same procedures as (4.2) c) - d), (4.3) b) 4) - 6) and c) 1). However, prepare the calibration curve from the peak area ratio or height ratio of ion (m/z) for determination of respective materials subjected to measurement and internal reference material, and calculate the concentration of respective materials.
subjected to measurement in the analytical sample.

**Comment 8** Recovery testing of melamine derivations was conducted using soybean meal, fish meal, fish waste processed fertilizer, mixed organic fertilizer, blended fertilizer and compound fertilizer, as a result, the mean recovery rate at additive level of 10 % (mass fraction) and 0.2 % (mass fraction) were 92.1 % - 102.9 % and 90.3 % - 102.2 %.

Additionally, the minimum limit of quantification of melamine derivations of the test method is about 0.01 % (mass fraction).

**References**

(5) **Flow sheet for melamine derivations**: The flow sheet for melamine derivations in fertilizers is shown below.

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.50 g analytical sample</td>
<td>200-mL - 300-mL ground-in stopper Erlenmeyer flask</td>
</tr>
<tr>
<td>Extraction</td>
<td>160 mL - 200 mL of diethylamine - water - acetonitrile (1+4+5)</td>
</tr>
<tr>
<td></td>
<td>Ultrasonic extraction for 30 minutes</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Ground-in stopper centrifugal precipitation tube</td>
</tr>
<tr>
<td></td>
<td>8000 × g - 10000 × g, 5 minutes</td>
</tr>
<tr>
<td>Extract</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 1** Flow sheet for melamine derivations in fertilizers (Extraction procedure)

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extract</td>
<td>Transfer 1 mL to 5-mL - 50-mL volumetric flasks, add diethylamine - water - acetonitrile (1+4+5) up to the marked line</td>
</tr>
<tr>
<td>Dilution</td>
<td>5-mL - 10-mL test tube with a screw stopper</td>
</tr>
<tr>
<td>0.2 mL Aliquot</td>
<td></td>
</tr>
<tr>
<td>Concentration under reduced pressure / Drying up</td>
<td>Centrifugal evaporator (70 ºC ± 2 ºC)</td>
</tr>
<tr>
<td></td>
<td>← 0.3 mL of pyridine (dehydration)</td>
</tr>
<tr>
<td></td>
<td>← 0.2 mL of derivatization reagent (BSTFA−TMCS (99+1))</td>
</tr>
<tr>
<td></td>
<td>Stopple and seal</td>
</tr>
<tr>
<td>Derivatization</td>
<td>70 ºC ± 2 ºC, 45 minutes</td>
</tr>
<tr>
<td>Standing to cool</td>
<td></td>
</tr>
<tr>
<td>Sample solution</td>
<td></td>
</tr>
<tr>
<td>Measurement</td>
<td>Gas Chromatograph/Mass Spectrometer</td>
</tr>
</tbody>
</table>

**Figure 2** Flow sheet for melamine derivations in fertilizers (Derivatization and measurement procedure)
Reference: Total Ion Chromatogram (TIC) of GC/MS of mixture standard solution for calibration curve preparation of melamine derivations is shown below.

Figure 3 Total Ion Chromatogram (TIC) of GC/MS for melamine derivations

Measurement conditions of GC/MS

Capillary column: Rtx-5ms (0.25-mm inner diameter, 30-m long, 0.25 µm layer thickness)

Other conditions are according to the example of HPLC measurement conditions in (4.3) a)

Peak name of respective total ion chromatograms

a) Cyanuric acid   b) Ammelide
c) Ammeline       d) Melamine

Sample and amount injected into GC/MS

Injected sample: Mixture standard solutions (the equivalents of 2 µg/mL) for the calibration curve preparation of respective melamine derivations.

Injected amount: 1 µL (the equivalents of 2 ng of respective melamine derivations)
8.1.b
8.1.c High-Performance Liquid Chromatography (Fertilizers not containing organic matters)

(1) **Summary**
This testing method is applicable to fertilizers not containing organic matters. This testing method is classified as Type B and its symbol is 8.1.c-2017 or Mel.c-1.

Add hydrochloric acid (1+15) to an analytical sample, extract melamine and its degradation products (hereinafter referred to as “melamine derivations”), introduce them into a High-Performance Liquid Chromatograph (HPLC), isolate with a silica gel column to which carbamoyl chemically bonds, and measure at wavelength 214 nm to obtain melamine derivations in an analytical sample. In addition, the performance of this testing method is shown in **Comment 4**.

(2) **Reagent, etc.**:
Reagents and water are as shown below.

a) **Water**: Water of A3 specified in JIS K 0557.

b) **Acetonitrile**: A JIS Guaranteed Reagent specified in JIS K 8032 or a reagent of equivalent quality. In addition, acetonitrile used in eluent of an HPLC is a reagent of HPLC analysis grade.

c) **Hydrochloric acid**: A guaranteed reagent or a reagent of equivalent quality.

d) **Phosphate buffer solution**: Dissolve 0.237 g of disodium hydrogen-phosphate specified in JIS K 9020 and 0.520 g of sodium dihydrogenphosphate dihydrate specified in JIS K 9009 in water to make 1000 mL. If it is used for eluent of HPLC, filter with a membrane filter (pore size: no more than 0.5-µm) made of hydrophilic PTFE.

e) **Melamine derivations standard solution (0.5 mg/mL)**: Put about 0.05 g of melamine \(\text{C}_3\text{H}_6\text{N}_6\) (3), ammeline \(\text{C}_3\text{H}_5\text{N}_5\text{O}\) (3), ammelide \(\text{C}_3\text{H}_4\text{N}_4\text{O}_2\) (3) and cyanuric acid \(\text{C}_3\text{H}_3\text{N}_3\text{O}_3\) (3) into a weighing dish and measure the mass to the order of 0.1 mg. Dissolve them with a small amount of hydrochloric acid (1+15), transfer to 100 mL volumetric flasks respectively, and add the solutions up to the marked line.

f) **Mixture standard solution (50 µg/mL)**: Transfer 5 mL of respective melamine derivations standard solutions (0.5 mg/mL) to 50-mL volumetric flasks and add acetonitrile--phosphate buffer solutions (4+1) up to the marked line.

g) **Mixture standard solution for calibration curve preparation (1 µg/mL - 5 µg/mL)**: In the case of usage, transfer 1 mL - 5 mL of mixture standard solution (50 µg /mL) to 50-mL volumetric flasks step-by-step and add acetonitrile--phosphate buffer solutions (4+1) up to the marked line.

h) **Mixture standard solution for calibration curve preparation (0.05 µg/mL - 0.5 µg/mL)**: In the case of usage, transfer 2.5 mL - 25 mL of mixture standard solution (1 µg /mL) to 50-mL volumetric flasks step-by-step and add acetonitrile--phosphate buffer solutions (4+1) up to the marked line.

Note (1) This is an example of preparation; prepare an amount as appropriate.
(2) The phosphate buffer solution becomes pH 6.7 ± pH 0.2.
(3) The respective standard reagents of melamine, ammeline, ammelide and cyanuric acid are commercially sold.

**Comment 1** The standard reagent of melamine, ammeline, ammelide and cyanuric acid are sold by FUJIFILM Wako Pure Chemical Co., Ltd. and Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(3) **Instruments**:
Instruments are as shown below:

a) **High-Performance Liquid Chromatograph (HPLC)**: HPLC specified in JIS K 0124 that satisfies following requirements.

1) **Column**: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube
filled with silica gel, to which carbamoyl chemically bonds.

2) **Column bath:** A column bath whose temperature can be adjusted to 40 °C ± 1 °C.
3) **Detection unit:** An absorptiometric detector that can measure at wavelength around 214 nm.

b) **Ultrasonic generator:** An ultrasonic washer can be used.

c) **Centrifugal separator:** A centrifugal separator that can work at about 1700 × g.

d) **High speed centrifugal separator:** A centrifugal separator that can work at 8000 × g - 10000 × g.

**Comment 2** Column is sold under the production name TSKgel Amide-80, etc. A column which has actually isolated melamine, ammeline, ammelide and cyanuric acid should be used.

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 0.50 g of an analytical sample, and put it into a 200 mL- ground-in stopper Erlenmeyer flask.

b) Add 100 mL of hydrochloric acid (1+15) and conduct ultra-sonication for about 30 minutes using an ultrasonic generator.

c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.

d) Centrifuge it at 1700 × g centrifugal force for about five minutes (4) and use the supernatant as the extract.

e) Transfer 5 mL of the extract (5) into a 50-mL volumetric flask, and add an acetonitrile–phosphate buffer solution (4+1) up to the marked line to dilute.

f) Transfer dilution liquid to a 1.5-mL ground-in stopper centrifugal precipitate tube (6).

g) Centrifuge at 8000 × g - 10000 × g for about 5 minutes (7) to make supernatant as the sample solution

**Note** (4) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

(5) If there is a possibility that the concentration of melamine derivations in the sample solution exceeds the maximum limit of the calibration curb, the amount of a supernatant solution to be transferred should be1 mL - 2.5 mL.

(6) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement

(7) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

**Comment 3** Instead of the procedures in (4.1) f) - g), it is allowed to filter with a membrane filter (pore size: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC):** An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

1) **Column:** A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-µm particle diameter column) to which carbamoyl chemically bonds.
2) **Column bath temperature**: 40 °C ± 1 °C
3) **Eluent**: Acetonitrile - phosphate buffer solution (4+1)
4) **Flow rate**: 1 mL/min
5) **Detection unit**: An absorptiometric detector, measurement wavelength: 214 nm

**b) Calibration curve preparation**
1) Inject 10 µL of respective mixture standard solutions for calibration curve preparation into an HPLC, record a chromatogram at wavelength 214 nm and obtain the peak area or height.
2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 214 nm of respective mixture solutions for the calibration curve preparation.

**c) Sample measurement**
1) Subject 10 µL of sample solution to the same procedure as in **b) 1)**
2) Obtain the melamine derivations content from the calibration curve to calculate respective melamine derivations in the analytical sample.

**Comment 4** Recovery testing was conducted using 3 brands of nitrolime, 1 brand of compound fertilizer containing nitrolime, 2 brands of compound fertilizers not containing nitrolime, 1 brand of ammonium sulfate and 1 brand of urea, as a result, the recovery rate of melamine derivations at concentration level of 4 % (mass fraction) and 0.1 % (mass fraction) were 90.5 % - 106.3 % and 92.2 % - 107.0 %.

Table 1 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification of the test method is about 0.02 % (mass fraction) for melamine and cyanuric acid and about 0.01 % (mass fraction) for ammeline and ammelide. In the case of ammelide and cyanuric acid, the sufficient reproducibility was observed in the range of 0.188 % (mass fraction) - 1.10 % (mass fraction) and 0.105 % (mass fraction) - 1.15 % (mass fraction) respectively.
Table 1  Results and analysis results from a collaborative study for the test method validation of melamine derivations

<table>
<thead>
<tr>
<th>Agrichemical name</th>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (%)&lt;sup&gt;2&lt;/sup&gt;</th>
<th>s&lt;sub&gt;r&lt;/sub&gt;(%)&lt;sup&gt;3&lt;/sup&gt;</th>
<th>RSD&lt;sub&gt;r&lt;/sub&gt;(%)&lt;sup&gt;4&lt;/sup&gt;</th>
<th>s&lt;sub&gt;R&lt;/sub&gt;(%)&lt;sup&gt;5&lt;/sup&gt;</th>
<th>RSD&lt;sub&gt;R&lt;/sub&gt;(%)&lt;sup&gt;6&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Melamine</td>
<td>Nitrolime 1</td>
<td>9</td>
<td>2.83</td>
<td>0.04</td>
<td>1.4</td>
<td>0.12</td>
<td>4.3</td>
</tr>
<tr>
<td></td>
<td>Nitrolime 2</td>
<td>10</td>
<td>0.391</td>
<td>0.003</td>
<td>0.8</td>
<td>0.023</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer containing nitrolime</td>
<td>9</td>
<td>0.845</td>
<td>0.019</td>
<td>2.2</td>
<td>0.036</td>
<td>4.2</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer</td>
<td>11</td>
<td>0.198</td>
<td>0.005</td>
<td>2.6</td>
<td>0.012</td>
<td>6.2</td>
</tr>
<tr>
<td></td>
<td>Ammonium sulfate</td>
<td>10</td>
<td>0.0343</td>
<td>0.0015</td>
<td>4.5</td>
<td>0.0040</td>
<td>11.6</td>
</tr>
<tr>
<td>Ammeline</td>
<td>Nitrolime 1</td>
<td>9</td>
<td>1.60</td>
<td>0.02</td>
<td>1.3</td>
<td>0.06</td>
<td>3.8</td>
</tr>
<tr>
<td></td>
<td>Nitrolime 2</td>
<td>10</td>
<td>0.105</td>
<td>0.001</td>
<td>1.3</td>
<td>0.002</td>
<td>2.3</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer containing nitrolime</td>
<td>9</td>
<td>0.629</td>
<td>0.027</td>
<td>4.3</td>
<td>0.023</td>
<td>3.7</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer</td>
<td>11</td>
<td>0.195</td>
<td>0.004</td>
<td>2.1</td>
<td>0.009</td>
<td>4.5</td>
</tr>
<tr>
<td></td>
<td>Ammonium sulfate</td>
<td>10</td>
<td>0.0346</td>
<td>0.0013</td>
<td>3.7</td>
<td>0.0024</td>
<td>6.9</td>
</tr>
<tr>
<td>Ammeline</td>
<td>Nitrolime 1</td>
<td>9</td>
<td>1.10</td>
<td>0.02</td>
<td>2.1</td>
<td>0.08</td>
<td>7.6</td>
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<tr>
<td></td>
<td>Nitrolime 2</td>
<td>11</td>
<td>0.361</td>
<td>0.008</td>
<td>2.2</td>
<td>0.023</td>
<td>6.5</td>
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<tr>
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<td>Compound fertilizer containing nitrolime</td>
<td>9</td>
<td>0.188</td>
<td>0.004</td>
<td>2.2</td>
<td>0.014</td>
<td>7.5</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer</td>
<td>11</td>
<td>0.718</td>
<td>0.028</td>
<td>3.9</td>
<td>0.052</td>
<td>7.2</td>
</tr>
<tr>
<td></td>
<td>Ammonium sulfate</td>
<td>11</td>
<td>0.0345</td>
<td>0.0031</td>
<td>8.9</td>
<td>0.0056</td>
<td>16.1</td>
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<tr>
<td>Cyanuric acid</td>
<td>Nitrolime 1</td>
<td>9</td>
<td>1.15</td>
<td>0.06</td>
<td>4.8</td>
<td>0.09</td>
<td>7.7</td>
</tr>
<tr>
<td></td>
<td>Nitrolime 2</td>
<td>10</td>
<td>0.390</td>
<td>0.018</td>
<td>4.5</td>
<td>0.029</td>
<td>7.4</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer containing nitrolime</td>
<td>9</td>
<td>0.105</td>
<td>0.003</td>
<td>2.9</td>
<td>0.014</td>
<td>13.2</td>
</tr>
<tr>
<td></td>
<td>Compound fertilizer</td>
<td>9</td>
<td>0.788</td>
<td>0.026</td>
<td>3.2</td>
<td>0.054</td>
<td>6.8</td>
</tr>
<tr>
<td></td>
<td>Ammonium sulfate</td>
<td>10</td>
<td>0.0365</td>
<td>0.0015</td>
<td>4.2</td>
<td>0.0067</td>
<td>18.3</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Gross mean (\(n = \text{number of laboratories} \times \text{number of repeated tests (2)}\))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Reproducibility standard deviation
6) Reproducibility relative standard deviation
7) Mass fraction

References


5 Flow sheet for melamine derivations: The flow sheet for melamine derivations in fertilizers is shown below.

<table>
<thead>
<tr>
<th>0.50 g analytical sample</th>
<th>200-mL ground-in stopper Erlenmeyer flask ←100 mL of hydrochloric acid (1+15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extraction</td>
<td>Ultrasonic extraction for 30 minutes</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Ground-in stopper centrifugal precipitation tube 1700 × g, 5 minutes</td>
</tr>
<tr>
<td>Dilution</td>
<td>Transfer 5 mL to a 50-mL volumetric flask, add acetonitrile–phosphate buffer solution (4+1) up to the marked line</td>
</tr>
<tr>
<td>Centrifugal separation</td>
<td>Ground-in stopper centrifugal precipitation tube 8000 × g - 10000 × g, 5 minutes</td>
</tr>
<tr>
<td>Sample solution</td>
<td>High Performance Liquid Chromatograph</td>
</tr>
</tbody>
</table>

Figure Flow sheet for melamine derivations in fertilizers
Reference: HPLC chromatogram of the mixture standard solution for calibration curve preparation of melamine derivations is shown below.

Reference diagram HPLC chromatogram of melamine derivations

The names of materials for respective peaks:
1. cyanuric acid
2. ammelide
3. melamine
4. ammeline

Measurement conditions for HPLC:
- Column: TSKgel Amide-80 (4.6-mm inner diameter, 250-mm long, 5-µm particle diameter)
- Mixture standard solutions (respective equivalents of 10 ng (1 µg/mL, 10 µL)) for calibration curve preparation of respective melamine derivations
- Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
8.1.d High-Performance Liquid Chromatography (Fertilizers containing organic matters)

(1) Summary
This testing method is applicable to organic fertilizers and fertilizers containing organic matters. This testing method is classified as Type B and its symbol is 8.1.d-2017 or Mel.d-1.

Add water to an analytical sample to extract melamine, introduce it into a High-Performance Liquid Chromatograph (HPLC), isolate with a silica gel column to which carbamoyl chemically bonds, and measure at wavelength 214 nm to obtain melamine in an analytical sample. However, cyanuric acid, ammelide and ammeline which are melamine derivations are excluded from components subjected to measurement. In addition, the performance of this testing method is shown in Comment 4.

(2) Reagent, etc.: Reagents and water are as shown below.
   b) Acetonitrile: A JIS Guaranteed Reagent specified in JIS K 8032 or a reagent of equivalent quality. In addition, acetonitrile used in eluent of an HPLC is a reagent of HPLC analysis grade.
   c) Phosphate buffer solution \(^{(1)}\): Dissolve 0.237 g of disodium hydrogen-phosphate specified in JIS K 9020 and 0.520 g of sodium dihydrogenphosphate dihydrate specified in JIS K 9009 in water to make 1000 mL \(^{(2)}\). If it is used for eluent of HPLC, filter with a membrane filter (pore size: no more than 0.5-µm) made of hydrophilic PTFE.
   d) Melamine standard solution (0.5 mg/mL) \(^{(1)}\): Put 0.05 g of melamine \([C_3H_6N_6]\) \(^{(3)}\) to a weighing dish and measure the mass to the order of 0.1 mg. Dissolve it with a small amount of water, transfer to a 100 mL volumetric flask and add the solution up to the marked line.
   e) Melamine standard solution (50 µg/mL) \(^{(1)}\): Transfer 5 mL of melamine solution (0.5 mg/mL) to a 50-mL volumetric flask and add acetonitrile - phosphate buffer solutions (82+18) up to the marked line.
   f) Melamine standard solution for calibration curve preparation (1 µg/mL - 5 µg/mL): In the case of usage, transfer 1 mL - 5 mL of melamine standard solution (50 µg /mL) to 50-mL volumetric flasks step-by-step and add acetonitrile–phosphate buffer solutions (82+18) up to the marked line.
   g) Melamine standard solution for calibration curve preparation (0.05 µg/mL - 0.5 µg/mL): In the case of usage, transfer 2.5 mL - 25 mL of melamine standard solution (1 µg/ mL) to 50-mL volumetric flasks step-by-step, add acetonitrile–phosphate buffer solutions (82+18) up to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.
   (2) The phosphate buffer solution becomes pH 6.7 ± 0.2.
   (3) A standard reagent as melamine is commercially sold.

Comment 1 A standard reagent of melamine is sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(3) Instruments: Instruments are as shown below:
   a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
      1) Column: A 4-mm - 6-mm inner diameter 150-mm - 250-mm long stainless steel column tube filled with silica gel, to which carbamoyl chemically bonds.
      2) Column bath: A column bath whose temperature can be adjusted to 40 °C ± 1 °C.
      3) Detection unit: An absorptiometric detector that can measure at wavelength around 214 nm.
b) **Ultrasonic generator**: An ultrasonic washer can be used.

c) **Centrifugal separator**: A centrifugal separator that can work at about $1700 \times g$.

d) **High speed centrifugal separator**: A centrifugal separator that can work at $8000 \times g - 10000 \times g$.

**Comment 2** Column is sold under the production name TSKgel Amide-80, etc. A column which has actually isolated melamine, ammeline, ammelide and cyanuric acid should be used.

(4) **Test procedures**

(4.1) **Extraction**: Conduct extraction as shown below.

- a) Weigh 0.50 g of an analytical sample, and put it into a 200 mL ground-in stopper Erlenmeyer flask.
- b) Add 100 mL of water and conduct ultra-sonication for about 10 minutes using an ultrasonic generator.
- c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
- d) Centrifuge it at $1700 \times g$ centrifugal force for about 10 minutes (4) and use the supernatant as the extract.
- e) Transfer 5 mL of the extract (5) into a 50-mL volumetric flask, and add an acetonitrile–phosphate buffer solution (82+18) up to the marked line to dilute.
- f) Transfer dilution liquid to a 1.5-mL ground-in stopper centrifugal precipitate tube (6).
- g) Centrifuge at $8000 \times g - 10000 \times g$ for about 5 minutes (7) to make supernatant as the sample solution.

**Note** (4) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about $1700 \times g$ centrifugal force.

(5) If there is a possibility that the concentration of melamine derivations in the sample solution exceeds the maximum limit of the calibration curb, the amount of a supernatant solution to be transferred should be 1 mL - 2.5 mL.

(6) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(7) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about $8100 \times g - 10000 \times g$ centrifugal force.

**Comment 3** Instead of the procedures in (4.1) f) - g), it is allowed to filter with a membrane filter (pore size: no more than 0.5 μm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) **Measurement**: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of High-Performance Liquid Chromatograph (HPLC) used in measurement.

- a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC)**: An example of measurement conditions for a High-Performance Liquid Chromatograph (HPLC) is shown below. Set up the measurement conditions considering it:

  1) **Column**: A silica gel column (4-mm - 6-mm inner diameter, 150-mm - 250-mm long, 5-μm particle diameter column) to which carbamoyl chemically bonds.
  2) **Column bath temperature**: 40 °C ± 1 °C
  3) **Eluent**: Acetonitrile - phosphate buffer solution (82+18)
  4) **Flow rate**: 1 mL/min
5) **Detection unit:** An absorptiometric detector, measurement wavelength: 214 nm

b) **Calibration curve preparation**
1) Inject 10 µL of respective melamine standard solutions for calibration curve preparation into an HPLC, record a chromatogram at wavelength 214 nm and obtain the peak area or height.
2) Prepare a curve for the relationship between the concentration and the peak area or height at wavelength 214 nm of respective melamine solutions for the calibration curve preparation.

c) **Sample measurement**
1) Subject 10 µL of sample solution to the same procedure as in b) 1)
2) Obtain the melamine content from the calibration curve to calculate melamine in the analytical sample.

**Comment 4** Additive recovery testing was conducted to evaluate trueness using rape seed meal, soybean meal, compound fertilizer containing lime nitrogen and organic matter, compound fertilizer containing organic matter and blended fertilizer containing organic matter (1 brand for each). As a result, the mean recovery rate at additive level of 2 % (mass fraction), 0.4 % (mass fraction) and 0.1 % (mass fraction) were 94.6 % - 99.8 %, 92.4 % - 98.5 % and 93.1 % - 98.4 % respectively.

The results of the repeatability tests on different days using soybean meal and compound fertilizer containing organic matter, to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$</th>
<th>Repeatability</th>
<th>Intermediate precision</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$^1$</td>
<td>($^3$) (%)</td>
<td>$^4$</td>
<td>($^3$) (%)</td>
</tr>
<tr>
<td>Soybean meal</td>
<td>5</td>
<td>1.91</td>
<td>0.03</td>
<td>1.7</td>
</tr>
<tr>
<td>Compound fertilizer containing organic matter</td>
<td>5</td>
<td>0.100</td>
<td>0.001</td>
<td>1.4</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) × the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

**References**
(5) **Flow sheet for melamine derivations**: The flow sheet for melamine derivations in fertilizers is shown below.

![Flow sheet for melamine derivations]

Reference: HPLC chromatogram of the standard solution for calibration curve preparation of melamine is shown below.

![Reference diagram HPLC chromatogram of melamine]

**Measurement conditions for HPLC**
- **Column**: TSKgel Amide-80 (4.6-mm inner diameter, 250-mm long, 5-µm particle diameter)
- **Melamine standard solution** (the equivalents of 10 ng (1 µg/mL, 10 µL) for each)
- **Other conditions** are according to the example of HPLC measurement conditions in (4.2) a)
8.2 Clopyralid and its degradation products
8.2.a High-Performance Liquid Chromatography/Tandem Mass Spectrometry
(1: Simultaneous analysis of three components for clopyralid, etc.)

(1) Summary
This testing method is applicable to compost and composted sludge fertilizers. This testing method is classified as Type B and its symbol is 8.2.a-2017 or CLP.a-1.

Extract clopyralid, aminopyralid and picloram with methanol under alkaline condition, refine with a cleanup cartridge by taking advantage of characteristics that the behavior of elution varies between acidity and alkaline, and then measure with a High-Performance Liquid Chromatograph Tandem Mass Spectrometer to obtain clopyralid, aminopyralid and picloram in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

Comment 1 Structural formulas of clopyralid, aminopyralid and picloram are as shown in Figure 1.

![Clopyralid, Aminopyralid, and Picloram Structural Formulas](image)

Figure 1 Structural formula of clopyralid, aminopyralid and picloram

(2) Reagent, etc.: Reagents and water are as shown below.

a) Water: Water of A3 specified in JIS K 0557. Note that water of A4 should be used as the eluent which is introduced to LC-MS/MS.
b) Acetonitrile: A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8039 or a reagent of equivalent quality.
c) Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.
d) Methanol: Methanol used in eluent of LC-MS/MS is a reagent of LC-MS analysis grade or equivalents.
e) Sodium hydroxide: A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.
f) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
g) Ammonia solution: A JIS Guaranteed Reagent of 28 % (mass fraction) specified in JIS K 8085 or a reagent of equivalent quality.
h) Formic acid: A JIS Guaranteed Reagent specified in JIS K 8264 or a reagent of equivalent quality.
i) Ammonia solution (0.0028 % (mass fraction))\(^{(1)}\): Add 0.1 mL of ammonia solution to 1000 mL of water.
j) Respective agrichemical standard solutions (0.1 mg/mL)\(^{(1)}\): Put about 0.01 g of clopyralid \([C_6H_3Cl_2NO_2]\)\(^{(2)}\), aminopyralid \([C_6H_4Cl_2N_2O_2]\)\(^{(2)}\), picloram \([C_6H_3Cl_3N_2O_2]\)\(^{(2)}\) to weighing dishes and measure the mass to the order of 0.1 mg. Dissolve with a small amount of
acetonitrile, transfer to 100-mL volumetric flasks and add the solvent up to the marked line.

k) **Mixture standard solution (100 ng/mL)** (1): Dilute a predetermined amount of respective agrichemical standard solutions (0.1 mg/mL) with formic acid (1+1000) to prepare mixture standard solution (100 ng/mL).

l) **Mixture standard solution for calibration curve preparation (5 ng/mL - 50 ng/mL)** (1): In the case of usage, transfer 2.5 mL - 25 mL of mixture standard solution (100 ng/mL) to 50 mL volumetric flasks step-by-step, and add formic acid (1+1000) up to the marked line.

m) **Mixture standard solution for calibration curve preparation (0.5 ng/mL - 5 ng/mL)** (1): In the case of usage, transfer 2.5 mL - 25 mL of mixture standard solution (10 ng/mL) to 50 mL volumetric flasks step-by-step, and add formic acid (1+1000) up to the marked line.

**Note**  
(1) This is an example of preparation; prepare an amount as appropriate.  
(2) A standard reagent is commercially sold.

**Comment 2** Standard reagents of clopyralid, aminopyralid and picloram are sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.

a) **High-Performance Liquid Chromatograph/Mass Spectrometer (LC-MS/MS)**: LC-MS/MS specified in JIS K 0136 that satisfies the following requirements.

1) **High-Performance Liquid Chromatograph**
   (i) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.
   (ii) Column: A 2-mm - 3-mm inner diameter 50-mm - 150-mm long 1.6-µm - 2.2-µm particle diameter stainless steel column tube filled with silica gel to which octadecyl chemically bonds. The specification is according to the mass spectrometer specification.

2) **Mass Spectrometer**
   (i) Ionization method: Electro-Spray Ionization (ESI) method
   (ii) Ion detection method: Selected Reaction Monitoring

b) **Shaking apparatus**

c) **Manifold**

d) **Centrifugal separator**: A centrifugal separator that can work at about 1700 × g.

e) **High speed centrifugal separator**: A centrifugal separator that can work at 8000 × g - 10000 × g.

f) **Concentrator**: An evaporator that can adjust to 40 ºC ± 2 ºC.

g) **Copolymer cartridge column**: A divinylbenzene-N-vinylpyrrolidone copolymer mini column (200 mg)

**Comment 3** Column is sold under the production name ACQUITY UPLC HSS C18, etc.

**Comment 4** A copolymer cartridge is sold under the production names such as Oasis HLB 6cc (200 mg), Oasis PRiME HLB Plus Short Cartridge (225 mg).

(4) **Test procedures**

(4.1) **Extraction**: Conduct extraction as shown below.

a) Weigh 5.00 g of an analytical sample, and put it into a 200-mL - 300-mL ground-in stopper Erlenmeyer flask.

b) Add 1 mL of sodium hydroxide solution (40 g/L) and 99 mL of methanol (3), shake to mix with a shaking apparatus for about 30 minutes.

c) After allowing to stand still, transfer a supernatant solution to a 50-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at about 1700 \times g centrifugal force for about five minutes \((^4)\) and use the supernatant as the extract.

**Note** (3) It is also allowed to add 100 mL of sodium hydroxide solution (40 g/L)–methanol \([1+99]\)

\((^4)\) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about 1700 \times g centrifugal force.

**Comment 5** Grind until it completely passes through a sieve of 500 \(\mu m\) aperture to prepare the test sample.

\((^4.2\) Cleanup (1)\(^5\)): Conduct cleanup (1) as shown below.

a) Wash a cartridge column with about 5 mL of methanol and 5 mL of water in advance.

b) Place a 100-mL round-bottom flask \(^6\) under the cartridge column, transfer 5 mL or 7 mL \(^7\) of the extract to the cartridge column and allow the extract to overflow until the surface of the liquid reaches the top of the packing materials.

c) Add about 5 mL of sodium hydroxide solution (0.4 g/L)–methanol \((1+1)\) to the cartridge column 2 times and allow the liquid to overflow in the same manner in b).

d) Add 5 mL of methanol.

**Note** (5) Use a pressure reducing device in the procedure in \((^4.2\) and \((^4.3\) as appropriate.

(6) When pretreating many analytical samples, a free-standing type vessel that can contain a solution with a liquid volume of 20 mL may be used. In this case, instead of procedure d), put an effluent into a round-bottle flask, wash the vessel 2 times with 2.5 mL of methanol and add washing to the previous effluent.

(7) When using Oasis HLB 6cc (200 mg), add 5 mL of the extract 2 times.

\((^4.3\) Cleanup (2)\(^5\)): Conduct cleanup (2) as shown below.

a) Wash a new cartridge column with about 5 mL of acetonitrile and 5 mL of hydrochloric acid \((1+120)\) in advance.

b) After conducting vacuum concentration of the effluent in \((^4.2\) c\) until no more than 5 mL on a water bath at no more than 40 \(\circ C\), add 3 mL of hydrochloric acid \((1+11)\).

c) Put the concentrated effluent into the cartridge column and allow the effluent to overflow until the surface of liquid reaches the top of packing materials.

d) Wash a round-bottom flask with about 5 mL of hydrochloric acid \((1+120)\) 2 times and add washing into the cartridge successively.

e) Then, add about 5 mL of hydrochloric acid \((1+120)\)–acetonitrile \((9+1)\) and about 5 mL of water into the cartridge successively and allow the liquid to overflow.

f) Place a 5-mL volumetric flask under the cartridge column, add 4 mL of ammonia solution \((0.0028 \% \text{ mass fraction})\)–acetonitrile \((9+1)\) to the cartridge column and allow clopyralid, aminopyralid and picloram to elute quickly.

g) Add formic acid \((1+1000)\) up to the marked line \(^8\) and transfer it to 1.5-mL ground-in stopper centrifugal precipitate tube \(^9\).

h) Centrifuge it at 8000 \times g - 10000 \times g centrifugal force for about five minutes \(^{10}\) and use the supernatant as the sample solution.

**Note** (8) If there is a possibility that the clopyralid, aminopyralid and picloram concentration in the sample solution exceed the maximum limit of the calibration curve, dilute a predetermined amount of effluent with formic acid \((1+1000)\).

\((^9)\) The ground-in stopper centrifugal precipitate tube should be made of polypropylene,
etc. to not affect the measurement

(10) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about $8100 \times g$ - $10000 \times g$ centrifugal force.

(4.4) Measurement: Conduct the measurement as indicated in JIS K 0136 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph Mass Spectrometer used in measurement.

a) The measurement conditions of High-Performance Liquid Chromatograph/Mass spectrometer: An example of measurement conditions for a High-Performance Liquid Chromatograph/Mass Spectrometer is shown below. Set up the measurement conditions considering it:

1) High-Performance Liquid Chromatograph
(i) Column: A silica gel column (2-mm - 3-mm inner diameter, 50-mm - 150-mm long, 1.6-µm - 2.2-µm particle diameter column) to which octadecyl chemically bonds.
(ii) Flow rate: 0.2 mL/min - 0.5 mL/min
(iii) Eluent: A: Formic acid (1+1000) B: Methanol:
(iv) Gradient: 0 min (5 %B) $\rightarrow$ 5 min (60 %B) $\rightarrow$ 6 min (95 %B) $\rightarrow$ 7 min (5 %B)
(v) Temperature of column bath: 40 °C
(vi) Injection volume: 5 µL

2) Mass Spectrometer
(i) Ionization method: Electro-Spray Ionization (ESI) method
(ii) Mode: Positive
(iii) Capillary voltage: 1.0 kV
(iv) Ion source temperature: 120 °C
(v) Desolvation temperature: 400 °C
(vi) Cone voltage: Shown in table 1
(vii) Collision energy: Shown in table 1
(viii) Monitor ion: Shown in table 1

Table 1 Monitor ion, etc. of respective agrichemicals

<table>
<thead>
<tr>
<th>Substance name</th>
<th>Precursor ion (m/z)</th>
<th>Product ion (Determination) (m/z)</th>
<th>Product ion (Validation) (m/z)</th>
<th>Cone voltage (v)</th>
<th>Collision energy (Determination) (eV)</th>
<th>Collision energy (Validation) (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clopyralid</td>
<td>192</td>
<td>146</td>
<td>110</td>
<td>20</td>
<td>20</td>
<td>30</td>
</tr>
<tr>
<td>Aminipyralid</td>
<td>207</td>
<td>161</td>
<td>189</td>
<td>22</td>
<td>22</td>
<td>16</td>
</tr>
<tr>
<td>Picloram</td>
<td>241</td>
<td>195</td>
<td>223</td>
<td>28</td>
<td>22</td>
<td>16</td>
</tr>
</tbody>
</table>

b) Calibration curve preparation
1) Inject 5 µL of respective mixture standard solutions for calibration curve into the LC-MS/MS, record the chromatogram of ion (m/z) for determination and ion (m/z) for validation of clopyralid, aminopyralid and picloram and obtain respective peak areas.
2) Calculate the peak area ratio or height ratio of ion (m/z) for determination and ion (m/z) for validation of clopyralid, aminopyralid and picloram.
3) Prepare a curve for the relationship between the concentration of respective agrichemicals and the peak area of ion (m/z) for determination of respective mixture standard solutions for the
c) **Sample measurement**

1) Subject 5 µL of the sample solution to the same procedure as in b) 2) - 3)\(^{(11)}\).

2) Obtain the content of material subjected to measurement from the calibration curve to calculate the material subjected to measurement in the analytical sample.

**Note** (11) Confirm that the ratio against the peak area ratio or height ratio of the standard solution is within the range of about ± 30 %. In addition, the peak area ratio or height ratio depends on the concentration.

**Comment 6** Additive recovery testing of clopyralid, aminopyralid and picloram was conducted using cow dung compost (2 kinds), composted sludge fertilizers containing cow manure (2 kinds) and composted sludge fertilizers containing pig manure (1 kind), as a result, the mean recovery rates at additive level of 1000 µg/kg, 400 µg/kg and 40 µg/kg were 78.1 % - 90.0 %, 81.0 % - 117.6 % and 71.2 % - 101.3 % respectively. Table 2 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limits of quantification of clopyralid, aminopyralid and picloram of the test method are about 10 µg/kg respectively.
Table 2 Results and analysis results from a collaborative study for the test method validation of clopyralid and its degradation products

<table>
<thead>
<tr>
<th>Agrichemical name</th>
<th>Sample name</th>
<th>Number of laboratories&lt;sup&gt;1&lt;/sup&gt;</th>
<th>Mean&lt;sup&gt;2&lt;/sup&gt; (µg/kg)</th>
<th>s&lt;sup&gt;3&lt;/sup&gt; (µg/kg)</th>
<th>RSD&lt;sup&gt;4&lt;/sup&gt; (%)</th>
<th>s&lt;sup&gt;5&lt;/sup&gt; (µg/kg)</th>
<th>RSD&lt;sup&gt;6&lt;/sup&gt; (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clopyralid</td>
<td>Compost 1</td>
<td>10</td>
<td>128</td>
<td>6</td>
<td>4.5</td>
<td>21</td>
<td>16.4</td>
</tr>
<tr>
<td></td>
<td>Compost 2</td>
<td>10</td>
<td>835</td>
<td>41</td>
<td>4.9</td>
<td>100</td>
<td>11.9</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 1</td>
<td>9</td>
<td>16.2</td>
<td>1.7</td>
<td>10.6</td>
<td>5.2</td>
<td>31.8</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 2</td>
<td>10</td>
<td>89.6</td>
<td>11.3</td>
<td>12.6</td>
<td>11.3</td>
<td>12.6</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 3</td>
<td>10</td>
<td>339</td>
<td>28</td>
<td>8.3</td>
<td>28</td>
<td>8.3</td>
</tr>
<tr>
<td>Aminopyralid</td>
<td>Compost 1</td>
<td>8</td>
<td>324</td>
<td>15</td>
<td>4.5</td>
<td>39</td>
<td>12.0</td>
</tr>
<tr>
<td></td>
<td>Compost 2</td>
<td>8</td>
<td>21.2</td>
<td>5.2</td>
<td>24.7</td>
<td>6.4</td>
<td>30.3</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 1</td>
<td>7</td>
<td>5.39</td>
<td>1.41</td>
<td>26.2</td>
<td>2.22</td>
<td>41.2</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 2</td>
<td>10</td>
<td>701</td>
<td>146</td>
<td>20.8</td>
<td>263</td>
<td>37.6</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 3</td>
<td>9</td>
<td>59.5</td>
<td>8.9</td>
<td>15.0</td>
<td>16.6</td>
<td>28.0</td>
</tr>
<tr>
<td>Picloram</td>
<td>Compost 1</td>
<td>10</td>
<td>840</td>
<td>50</td>
<td>5.9</td>
<td>175</td>
<td>20.8</td>
</tr>
<tr>
<td></td>
<td>Compost 2</td>
<td>9</td>
<td>37.7</td>
<td>3.5</td>
<td>9.4</td>
<td>10.3</td>
<td>27.3</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 1</td>
<td>9</td>
<td>90.2</td>
<td>11.1</td>
<td>12.3</td>
<td>30.3</td>
<td>33.5</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 2</td>
<td>8</td>
<td>341</td>
<td>19</td>
<td>5.6</td>
<td>67</td>
<td>19.8</td>
</tr>
<tr>
<td></td>
<td>Composted sludge fertilizer 3</td>
<td>8</td>
<td>182</td>
<td>16</td>
<td>8.6</td>
<td>56</td>
<td>31.0</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis  
2) Gross mean (n = number of laboratories × number of repeated tests (2))  
3) Repeatability standard deviation  
4) Repeatability relative standard deviation  
5) Reproducibility standard deviation  
6) Reproducibility relative standard deviation

References


3) Kenji KOZUKA, Mayu OSHIMA, Yoshimi HASHIMOTO, Naoko TAMARU, and Yuji SHIRAI: Determination of Clopyralid, Aminopyralid and Picloram in Compost and
(5) **Flow sheet for clopyralid and its derivative substances**: The flow sheet for clopyralid and its derivative substances in compost and composted sludge fertilizer is shown below:

![Flow sheet for clopyralid and its derivative substances in compost and composted sludge fertilizers](image)

Figure 1  Flow sheet for clopyralid and its derivative substances in compost and composted sludge fertilizers (Extraction procedure)

![Flow sheet for clopyralid in compost and composted sludge fertilizers](image)

Figure 2  Flow sheet for clopyralid in compost and composted sludge fertilizers (Cleanup (1) and cleanup (2) and measurement procedure)
**References:** Selected Reaction Monitoring chromatograms of mixture standard solution for calibration curve preparation and sample solution (cow dung compost) are shown below.

Peak No. 1: Picloram  
No. 2: Aminopyralid  
No. 3: Clopyralid

Reference diagram: SRM chromatograms of respective agrichemicals  
Mixture standard solution (the equivalents of 200 pg as respective agrichemicals)

LC-MS/MS measurement conditions  
Column: ACQUITY UPLC HSS C18 (2.1-mm inner diameter, 100-mm long, 1.8-µm particle diameter)  
Other conditions are according to the examples of the measurement conditions in (4.4)  
a) LC-MS/MS.
8.2.b High-Performance Liquid Chromatography/Tandem Mass Spectrometry  
(2: microanalysis for clopyralid)

(1) **Summary**
This testing method is applicable to compost and composted sludge fertilizers. This testing method is classified as Type B and its symbol is 8.2.b-2018 or CLP.b-1.

Extract clopyralid in compost and composted sludge fertilizer with methanol under alkaline condition, refine with a cleanup cartridge and dichloromethane by taking advantage of characteristics that the behavior of elution varies between acidity and and then measure with a High-Performance Liquid Chromatograph Mass Spectrometer to obtain clopyralid in an analytical sample. In addition, the performance of this testing method is shown in Comment 8.

**Comment 1** Structural formula of clopyralid is as shown in Figure 1.

![Figure 1 Structural formula of clopyralid](image)

(2) **Reagent, etc.:** Reagents and water are as shown below.

- **a) Water:** Water of A3 specified in JIS K 0557. Note that water of A4 should be used as the eluent which is introduced to LC-MS/MS.
- **b) Acetonitrile:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8039 or a reagent of equivalent quality.
- **c) Methanol:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) or a reagent of equivalent quality.
- **d) Methanol:** Methanol used in eluent of LC-MS/MS is a reagent of LC-MS analysis grade or equivalents.
- **e) Sodium hydroxide:** A JIS Guaranteed Reagent specified in JIS K 8576 or a reagent of equivalent quality.
- **f) Hydrochloric acid:** A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
- **g) Ammonia solution:** A JIS Guaranteed Reagent of 28 % (mass fraction) specified in JIS K 8085 or a reagent of equivalent quality.
- **h) Formic acid:** A JIS Guaranteed Reagent specified in JIS K 8264 or a reagent of equivalent quality.
- **i) Formic acid:** Formic acid used in eluent of LC-MS/MS is a reagent of LC-MS analysis grade or equivalents.
- **j) Dichloromethane:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8117 or a reagent of equivalent quality.
- **k) Sulfuric acid:** A JIS Guaranteed Reagent specified in JIS K 8951 or a reagent of equivalent quality.
- **l) Acetone:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8040 or a reagent of equivalent quality.
- **m) Ammonia solution (0.0028 % (mass fraction))** (1): Add 0.1 mL of ammonia solution to 1000 mL of water.
- **n) Clopyralid standard solution (0.1 mg/mL)** (1): Put about 0.01 g of clopyralid [C₆H₃C₁₂NO₂]
(2) in a weighing dish and measure the mass to the order of 0.1 mg. Dissolve with a small amount of acetonitrile, transfer to 100-mL volumetric flasks and add the solvent up to the marked line.

o) **Clopyralid standard solution (250 ng/mL)** (1): Dilute a predetermined amount of clopyralid standard solutions (0.1 mg/mL) with formic acid (1+1000) to prepare clopyralid standard solution (250 ng/mL).

p) **Clopyralid standard solution for calibration curve preparation (5 ng/mL - 50 ng/mL)** (1): In the case of usage, transfer 2.5 mL - 25 mL of clopyralid standard solution (100 ng/mL) to 50 mL volumetric flasks step-by-step, and add formic acid (1+1000) up to the marked line.

q) **Clopyralid standard solution for calibration curve preparation (0.5 ng/mL - 5 ng/mL)** (1): In the case of usage, transfer 2.5 mL - 25 mL of clopyralid standard solution (10 ng/mL) to 50 mL volumetric flasks step-by-step, and add formic acid (1+1000) up to the marked line.

**Note** (1) This is an example of preparation; prepare an amount as appropriate.
(2) A standard reagent is commercially sold.

**Comment 2** A standard reagent of clopyralid is sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(3) **Apparatus and instruments**: Apparatus and instruments are shown below.

a) **High-Performance Liquid Chromatograph/Mass Spectrometer (LC-MS/MS)**: LC-MS/MS specified in JIS K 0136 that satisfies the following requirements.

1) **High-Performance Liquid Chromatograph**
   (i) Column bath: A column bath whose temperature can be adjusted to 30 °C - 45 °C.
   (ii) Column: A 2-mm - 3-mm inner diameter 50-mm - 150-mm long 1.6-µm - 2.2-µm particle diameter stainless steel column tube filled with silica gel to which octadecyl chemically bonds. The specification is according to the mass spectrometer specification.

2) **Mass Spectrometer**
   (i) Ionization method: Electro-Spray Ionization (ESI) method
   (ii) Ion detection method: Selected Reaction Monitoring

b) **Shaking apparatus**

c) **Manifold**

d) **Centrifugal separator**: A centrifugal separator that can work at 700 × g - 2000 × g.

e) **High speed centrifugal separator**: A centrifugal separator that can work at 8000 × g - 10000 × g.

f) **Concentrator**: An evaporator that can adjust to 40 °C ± 2 °C.

g) **Copolymer cartridge column**: A divinylbenzene-N-vinylpyrrolidone copolymer mini column (200 mg or 335 mg)

h) **Filter**: A funnel for filtering under reduced pressure (compatible filter diameter: 60 mm)

i) **Glass fiber filter paper**: A filter paper made of glass fiber (filter diameter: 60 mm) that can keep particle diameter 0.8 µm.

j) **Test tube mixer**: Vortex mixer

**Comment 3** Column is sold under the production name ACQUITY UPLC HSS C18, etc.
**Comment 4** A copolymer cartridge is sold under the production names such as Oasis HLB 6cc (225 mg), Oasis PRiME HLB Plus Short Cartridge (225 mg).
**Comment 5** A funnel for filtering under reduced pressure is sold under the production name KIRIYAMA Funnel SB-60, KIRIYAMA Funnel SU-60, etc.
**Comment 6** A glass fiber filter is sold under the production name Glass filter paper GFP-60, etc.
(4) Test procedures

(4.1) Extraction: Conduct extraction as shown below.

a) Weigh 5.00 g of an analytical sample to the order of 1 mg and put it in a 100-mL centrifugal precipitate tube with a screw cap.\(^{(3) (4)}\)

b) Add 50 mL of sodium hydroxide solution (40 g/L)–methanol [1+99] and shake to mix with a shaking apparatus for about 30 minutes.

c) Centrifuge at about \(1700 \times g\) for about 3 minutes \(^{(5)}\) and transfer the supernatant to a 100-mL Erlenmeyer flask.

d) Add 40 mL of sodium hydroxide solution (40 g/L)–methanol [1+99] to residue and shake to mix with a shaking apparatus for about 30 minutes.

e) Centrifuge at about \(1700 \times g\) for about five minutes \(^{(5)}\).

f) Filter supernatant in c) and e) under reduced pressure with a filter that places a glass fiber filter paper to a 100-mL short-neck volumetric flask as an acceptor \(^{(6)}\).

g) Wash the vessel and residue with a small amount of sodium hydroxide (40 g/L)–methanol [1+99] several times and transfer the washing to the previous filter to filter under pressure.

h) Add sodium hydroxide solution (40 g/L) - methanol [1+99] to the marked line to make the extract.

Note

(3) A vessel used for an extract procedure should be made of glass or polypropylene and it must enable vibration and centrifugation procedure.

(4) A 100-mL - 200-mL ground-in stopper or screw cap Erlenmeyer flask can also be used. In this case, however, suspension should be transferred to a ground-in stopper or screw cap centrifugal precipitate tube before the procedure c) and e).

(5) 16.5-cm of rotor radius and 3000 rpm of revolutions makes about \(1700 \times g\) centrifugal force.

(6) An Erlenmeyer flask can also be used. In this case, however, filtrate should be transferred to a 100-mL volumetric flask before the procedure h).

Comment

5 Grind until it completely passes through a sieve of 500 µm aperture to prepare the test sample.

(4.2) Cleanup (1) \(^{(7)}\): Conduct cleanup (1) as shown below.

a) Wash a cartridge column with about 5 mL of methanol and 5 mL of water in advance.

b) Place a 100-mL round-bottom flask \(^{(8)}\) under the cartridge column, transfer 9 mL \(^{(9)}\) of the extract to the cartridge column and allow the extract to overflow until the surface of the liquid reaches the top of the packing materials.

c) Add about 5 mL of sodium hydroxide solution (0.4 g/L)–methanol [1+1] to the cartridge column 2 times and allow the liquid to overflow in the same manner in b).

d) Add 5 mL of methanol.

Note

(7) Use a pressure reducing device in the procedure in (4.2) and (4.3) as appropriate.

(8) When making a pretreatment of many analytical samples, a free-standing type vessel that can contain a solution with a liquid volume of 20 mL may be used. In this case, instead of procedure d), put an effluent into a round-bottle flask, wash the vessel 2 times with 2.5 mL of methanol and add washing to the previous effluent.

(9) When using Oasis HLB 6cc (200 mg), add 5 mL of the extract 2 times.

(4.3) Cleanup (2) \(^{(7)}\): Conduct cleanup (2) as shown below.

a) Wash a new cartridge column with about 5 mL of acetonitrile and 5 mL of hydrochloric acid (1+120) in advance.
b) After conducting vacuum concentration of the effluent in (4.2) c) until no more than 5 mL on a water bath at no more than 40 °C, add 3 mL of hydrochloric acid (1 +11).

c) Put the concentrated effluent into the cartridge column and allow the effluent to overflow until the surface of liquid reaches the top of packing materials.

d) Wash a round-bottom flask with about 5 mL of hydrochloric acid (1+120) 2 times and add washing into the cartridge successively.

e) Then, add about 5 mL of hydrochloric acid (1+120)–acetonitrile (9+1) and about 5 mL of water into the cartridge and allow the liquid to overflow.

f) Place a 10-mL cone shaped centrifugal precipitate tube with a screw cap (10) under the cartridge column, add 4 mL of ammonia solution (0.0028 % (mass fraction))–acetonitrile [9+1] to the cartridge column and allow cleypralid to elute.

Note  (10) The part under 2 mL from the bottom forms a cone shape.

(4.4) Cleanup (3): Conduct cleanup (3) as shown below.

a) Add 0.1 mL of sodium hydroxide (40 g/L) to the elute in (4.3) f) and shake to mix by using a test tube mixer.

b) Add about 2 mL of dichloromethane, and shake to mix using a test tube mixer for about 30 minutes.

c) Centrifuge at about 740 × g for about five minutes (11) and remove a low layer by using a Pasteur pipet (12) or a syringe.

d) Repeat the procedure in b) - c) 1 time.

e) Add about 0.15 mL of sulfuric acid (1+2), and shake to mix using a test tube mixer.

f) Add about 2 mL of dichloromethane, and shake to mix using a test tube mixer for about 30 minutes (13).

g) Centrifuge at about 740 × g for about five minutes (11) and transfer a low layer to a 50mL round-bottle flask by using a Pasteur pipet (14) or a syringe.

h) Repeat the procedure in f) - g) 2 times. Note that lower layers are added to the same round-bottle flask.

i) Add 5 mL of acetone.

j) Concentrate under reduced pressure in a water bath of no more than 40 °C until most of the elute dries up and send a nitrogen gas to dry up.

k) Add 1 mL of formic acid (1+1000) and transfer it to a 1.5-mL ground-in stopper centrifugal precipitate tube (15).

l) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes (16) and use the supernatant as the sample solution.

Note  (11) 16.5-cm of rotor radius and 2000 rpm of revolutions makes about 740 × g centrifugal force. Confirm the permissible range of centrifugal force of a 10-mL cone shaped bottle centrifugal precipitate tube with a screw cap used.

(12) When using a Pasteur pipet, use the same Pasteur pipet through a series of procedures in c) - d).

(13) Disperse dichloromethane sufficiently. If vibration occurs while dichloromethane layer remains caked, the extraction efficiency of clopyralid deteriorates and measurements are affected.

(14) When using a Pasteur pipet, use the same Pasteur pipet through a series of procedures in g) - h). Do not use the Pasteur pipet used in Note (9).

(15) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement

(16) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g -
10000 $\times g$ centrifugal force.

(17) If there is a possibility that the clopyralid concentration in the sample solution exceed the maximum limit of the calibration curve, dilute a predetermined amount of effluent with formic acid (1+1000).

**Comment 6** Aa an alternative to the procedures in (4.4) k - l), filtration is allowed with a membrane filter (pore size: no more than 0.5-µm) made of hydrophilic PTFE or conduct centrifugal filtration with a centrifugal type filter unit (Ultrafree-MC PVDF membrane (0.22 µm), etc) and the filtrate can be the sample solution.

**Comment 7** When further concentration is required to ensure minimum quantitation, dissolve concentrated matters in the procedure in j) by adding acetone, transfer to a nitrogen concentration tube with the same solvent and send a nitrogen gas to dry up. Then add 0.2 mL of formic acid (1+1000) and conduct centrifugal filtration with a centrifugal type filter unit (Ultrafree-MC PVDF membrane (0.22 µm), etc) and the filtrate can be the sample solution. In this case, don not conduct the procedure in i).

**4.5) Measurement:** Conduct the measurement as indicated in JIS K 0136 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph Mass Spectrometer used in measurement.

a) **The measurement conditions of High-Performance Liquid Chromatograph/Mass spectrometer:** An example of measurement conditions for a High-Performance Liquid Chromatograph/Mass Spectrometer is shown below. Set up the measurement conditions considering it:

1) **High-Performance Liquid Chromatograph**
   (i) Column: A silica gel column (2-mm - 3-mm inner diameter, 50-mm - 150-mm long, 1.6-µm - 2.2-µm particle diameter column) to which octadecyl chemically bonds.
   (ii) Flow rate: 0.2 mL/min - 0.5 mL/min
   (iii) Eluent: A: Formic acid (1+1000) B: Methanol:
   (iv) Gradient: 0 min (5 %B) → 5 min (60 %B) → 6 min (95 %B) → 7 min (5 %B)
   (v) Temperature of column bath: 40 °C
   (vi) Injection volume: 5 µL

2) **Mass Spectrometer**
   (i) Ionization method: Electro-Spray Ionization (ESI) method
   (ii) Mode: Positive
   (iii) Capillary voltage: 1.0 kV
   (iv) Ion source temperature: 120 °C
   (v) Desolvation temperature: 400 °C
   (vi) Cone voltage: 20 V
   (vii) Collision energy: 20 eV for determination, 30 eV for validation
   (viii) Monitor ion: Precursor ion m/z 192
         Product ion m/z 146 for determination, m/z 110 for validation

b) **Calibration curve preparation**
1) Inject 5 µL of respective clopyralid standard solutions for calibration curve into the LC-MS/MS, record the chromatogram of ion (m/z) for determination and ion (m/z) for validation of clopyralid and obtain respective peak areas.
2) Calculate the peak area ratio or height ratio of ion (m/z) for determination and ion (m/z) for validation of clopyralid.
3) Prepare a curve for the relationship between the concentration of respective agrichemicals and the peak area of ion (m/z) for determination of respective mixture standard solutions for the
calibration curve preparation.

c) **Sample measurement**

1) Subject 5 µL of the sample solution to the same procedure as in b) 2) - 3) \(^{18}\).

2) Obtain the clopyralid content from the calibration curve to calculate clopyralid in the analytical sample.

**Note** \(^{18}\) Confirm that the ratio against the peak area ratio or height ratio of the standard solution is within the range of about ± 30%. In addition, the peak area ratio or height ratio may depend on the concentration.

**Comment 8** Additive recovery testing of clopyralid was conducted using cow dung compost (1 kinds), as a result, the mean recovery rates at additive level of 50 µg/kg, 10 µg/kg and 2 µg/kg were 78.9 %, 78.3 % and 71.5 % respectively. In addition, additive recovery testing of clopyralid was conducted using swine manure compost, poultry manure compost and composted sludge fertilizer and (1 sample for each), as a result, the mean recovery rates at additive level of 200µg/kg, 2 µg/kg and 80µg/kg were 88.6 %, 81.2 % and 94.2 % respectively.

Table 1 shows results and analysis results from a collaborative study for testing method validation.

Additionally, the minimum limit of quantification of clopyralid of this test method is about 2 µg/kg.

<table>
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<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (^{2}) (µg/kg)</th>
<th>(s_{r}^{3}) (µg/kg)</th>
<th>(RSD_{r}^{4}) (%)</th>
<th>(s_{R}^{5}) (µg/kg)</th>
<th>(RSD_{R}^{6}) (%)</th>
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<td>15</td>
<td>11.4</td>
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<td>15.3</td>
<td>0.40</td>
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<tr>
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<td>22.5</td>
<td>2.3</td>
<td>10.3</td>
<td>3.4</td>
<td>15.3</td>
</tr>
<tr>
<td>Poultry manure compost</td>
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<td>1.20</td>
<td>0.06</td>
<td>5.0</td>
<td>0.14</td>
<td>12.0</td>
</tr>
<tr>
<td>Composted sludge fertilizer</td>
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<td>48.1</td>
<td>1.2</td>
<td>2.5</td>
<td>5.6</td>
<td>11.6</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis
2) Gross mean (\(n = \) number of laboratories × number of repeated tests (2))
3) Repeatability standard deviation
4) Repeatability relative standard deviation
5) Reproducibility standard deviation
6) Reproducibility relative standard deviation

**References**

1) National Research and Development Agency: Institute for Agro-Environmental Sciences, National Agriculture and Food Research Organization: High-sensitive analysis
for clopyralid in cow dung compost (Reference method)

(5) **Flow sheet for clopyralid:** The flow sheet for clopyralid in compost is shown below:

![Flow sheet for clopyralid](Image)

**Figure 1** Flow sheet for clopyralid in compost

and composted sludge fertilizers (Extraction procedure)

Figure 2  Flow sheet for clomiprinal in compost and composted sludge fertilizers
(Cleanup (1) and cleanup (2) and measurement procedure)
Figure 2  Flow sheet for clopyralid in compost and composted sludge fertilizers (Cleanup (3) and measurement procedure)
**References:** Selected Reaction Monitoring chromatogram of clopyralid standard solution for calibration curve preparation is shown below

Reference diagram: SRM chromatograms of clopyralid
Clopyralid standard solution (the equivalents of 100 pg as clopyralid)

LC-MS/MS measurement conditions
1) Column: ACQUITY UPLC HSS C18 (2.1-mm inner diameter, 100-mm long, 1.8-µm particle diameter)
Other conditions are according to the examples of the measurement conditions in (4.4)
 a) LC-MS/MS.
8.3 Residue agrichemicals (multicomponent)

8.3.1 Residue agrichemicals multicomponent analysis (1)

8.3.1.a High-Performance Liquid Chromatography/Tandem Mass Spectrometry

(1) Compounds subjected to analysis


(2) Summary

This testing method is applicable to fluid home garden-use mixed fertilizer and fluid mixed fertilizer. This testing method is classified as Type B and its symbol is 8.3.1.a-2017 or AG-C-1.a-1.

Dissolve respective agricultural chemicals in fertilizers with acetonitrile and water, and extract. Refine by using 2 kinds of cleanup cartridge, and then measure with a High-Performance Liquid Chromatograph/Mass Spectrometer to obtain compounds subjected to analysis in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(3) Reagent, etc.: Reagents and water are as shown below.


b) Acetonitrile: A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8039 or a reagent of equivalent quality.

c) Methanol: A JIS Guaranteed Reagent specified in JIS K 8891 or a reagent of equivalent quality.

d) Methanol: Methanol used in eluent of an HPLC is a reagent of LC-MS analysis grade or equivalents.

e) Ethyl acetate: A JIS Guaranteed Reagent specified in JIS K 8361 or a reagent of equivalent quality.

f) Toluene: A JIS Guaranteed Reagent specified in JIS K 8680 or a reagent of equivalent quality.

g) Ammonium formate: A JIS Guaranteed Reagent (no less than 95 % (mass fraction) in purity) or a reagent of equivalent quality.

h) Ammonium formate solution (0.1 mol/L) (1): Add 6.306 g of ammonium formate into 1000 mL of water.

i) Ammonium formate solution (0.1 mmol/L) (1): Add 1 mL of ammonium formate solution (0.1 mol/L) into 1000 mL water.

j) Formic acid: A JIS Guaranteed Reagent specified in JIS K 8264 or a reagent of equivalent quality.

k) Formic acid solution (0.1 v/v %) (1): Add 1 mL of formic acid to 1,000 mL of water.

l) Acetonitrile formate solution (0.1 v/v %) (1): Add 1 mL of formic acid to 1,000 mL of acetonitrile.

m) Respective agricultural chemicals standard solutions (0.1 mg/mL) (1): Put about 0.01 g of abamectin \([C_{48}H_{72}O_{14}]\) (2), ivermectin \([C_{48}H_{74}O_{14}]\) (2), eprinomectin \([C_{50}H_{75}NO_{14}]\) (2), rotenone \([C_{23}H_{22}O_{6}]\) (2), piperonylbutoxide \([C_{19}H_{30}O_{3}]\) (2) and pyrethrin [pyrethrin I: \(C_{21}H_{28}O_{3}\) and pyrethrin II: \(C_{22}H_{28}O_{5}\)] (2) to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve with a small amount of methanol, transfer to a 100-mL volumetric flask and add the solvent up to the marked line. (However, 0.1 mg/mL of pyrethrin contains total of pyrethrin I/II)

n) Mixture standard solution (10 µg/mL): Transfer 10 mL of respective agricultural chemicals standard solutions to a 100-mL volumetric flask and add methanol up to the marked line.

o) Mixture standard solution (1000 ng/mL): Transfer 10 mL of mixture standard solution (10 µg/mL) to a 100-mL volumetric flask and add methanol up to the marked line.

p) Mixture standard solution for calibration curve preparation (50 ng/mL - 500 ng/mL): In
the case of usage, transfer 2.5 mL - 25 mL of mixture standard solution (1000 ng/mL) to 50-mL volumetric flasks step-by-step, and add methanol up to the marked line.

q) **Mixture standard solution for calibration curve preparation (5 ng/mL - 50 ng/mL):** In the case of usage, transfer 2.5 mL - 25 mL of mixture standard solution (100 ng/mL) to 50-mL volumetric flasks step-by-step, and add methanol up to the marked line.

**Note**

(1) This is an example of preparation; prepare an amount as appropriate.

(2) A standard reagent is commercially sold.

**Comment 1** A standard reagent of respective agricultural chemicals is sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(4) **Apparatus and instruments:** Apparatus and instruments are shown below.

a) **High-Performance Liquid Chromatograph/Mass Spectrometer (LC-MS/MS):**

LC-MS/MS specified in JIS K 0136 that satisfies the following requirements.

1) **High-Performance Liquid Chromatograph**

(i) **Column bath:** A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.

(ii) **Column:** A 2-mm - 3-mm inner diameter 50-mm - 150-mm long 1.6-µm - 3.0-µm particle diameter stainless steel column tube filled with silica gel to which octadecyl chemically bonds. The specification is according to the mass spectrometer specification (3).

2) **Mass Spectrometer**

(i) **Ionization method:** Electro-Spray Ionization (ESI) method

(ii) **Ion detection method:** Selected Reaction Monitoring

b) **Ultrasonic generator:** An ultrasonic washer can be used.

c) **Concentrator:** An evaporator whose temperature can be adjusted up to 40 ºC.

d) **Porous diatomaceous earth cartridge column:** A column that is filled with the porous diatomaceous earth (capacity: 5 mL) (4).

e) **Graphite carbon-NH$_2$ laminate cartridge column:** A 6-mL cylinder on which 500 mg of graphite carbon and 500 mg of aminopropyl silylation silica gel is laminated (5).

**Note**

(3) The column is sold under the names ACQUITY UPLC HSS C18, etc.

(4) The column is sold under the names Chem Elut (5 mL), etc.

(5) The column is sold under the names Envi-carb/LC-NH$_2$ (500 mg/500 mg, 6 mL), etc.

(5) **Test procedures**

(5.1) **Extraction:** Conduct extraction as shown below.

a) Put about 5.00 mL (6) of an analytical sample into a 10-mL volumetric flask.

b) Add 3 mL of acetonitrile to the same volumetric flask, and add water up to the marked line to shake to mix well.

c) Conduct ultra-sonication for about 30 minutes using an ultrasonic generator (7) to make the extract.

**Note**

(6) After measuring the specific gravity of sample, calculate the concentration of materials subjected to measurement in the analytical sample.

(7) Note that the volume of the solution may expand as a result of ultra-sonication. It is recommended to leave it at room temperature for a while when it expands.

**Comment 2** The specific gravity (density) can be calculated by placing a 10-mL volumetric flask
on an electric balance, aligning the scale to zero, transferring 5.00 mL of the analytical sample to the volumetric flask and reading the weighing value.

(5.2) Cleanup (1): Conduct cleanup (1) as shown below.

a) Put 5 mL of extract into a porous diatomaceous earth cartridge column and keep it in the column for about 5 minutes.

b) Place a 100-mL round-bottom flask under the same cartridge column, add about 5 mL of ethyl acetate into the same cartridge column 4 times successively and allow the solution to elute until the surface of the solution reaches the top of packing materials (8).

c) After conducting vacuum concentration of elute in a water bath of no more than 40 ºC until most of the elute dries up, send a nitrogen gas to dry up the elute (9), and add 2 mL of acetonitrile–toluene (3+1) to dissolve the residue.

Note  (8) Confirm the solution to elute before conducting the testing.

(9) There is a possibility for agricultural chemicals to vaporize if it is dried up excessively.

(5.3) Cleanup (2): Conduct cleanup (2) as shown below.

a) Wash the graphite carbon-NH$_2$ laminate cartridge column with about 10 mL of acetonitrile–toluene (3+1) in advance

b) Place a 100-mL round-bottom flask under the same cartridge column, put the solution in (5.2) c) to the same cartridge column, and allow the solution to overflow until the surface of the solution reaches the top of packing materials.

c) Wash the vessel with about 5 mL of acetonitrile–toluene (3+1) 5 times and add washing to the same cartridge successively to allow it to overflow.

d) After conducting vacuum concentration of elute in a water bath of no more than 40 ºC until most of the elute dries up, send a nitrogen gas to dry up the elute (10), and add 2 mL of methanol (11) to dissolve the residue. Transfer a predetermined amount of the solution precisely and dilute with methanol exactly by a factor of 5 to make the solution as the sample solution.

Note  (10) There is a possibility for agricultural chemicals to vaporize if it is dried up excessively.

(11) If there is a possibility that the concentration of agricultural chemicals in the sample solution exceeds the maximum limit of the calibration curve, dilute a predetermined amount of the sample solution with methanol.

(5.4) Measurement: Conduct the measurement as indicated in JIS K 0136 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph Mass Spectrometer used in measurement.

a) The measurement conditions of High-Performance Liquid Chromatograph/Mass spectrometer: An example of measurement conditions for a High-Performance Liquid Chromatograph/Mass Spectrometer is shown below. Set up the measurement conditions considering it:

1) High-Performance Liquid Chromatograph

   (i) Column: A silica gel column (2-mm - 3-mm inner diameter, 50-mm - 150-mm long, 1.6-µm - 3.0-µm particle diameter column) to which octadecyl chemically bonds.

   (ii) Flow rate: 0.2 mL/min - 0.5 mL/min

   (iii) Eluent: A: Ammonium formate solution (0.1 mmol/L)–formic acid solution (0.1 v/v%) [1+1]

   B: Acetonitrile formate solution (0.1 v/v%)

   (iv) Gradient: 0 min (50 %B) → 15 min (95 %B) → 20 min (98 %B) → 30 min (50 %B)
(v) Temperature of column bath: 40 °C
(vi) Injection volume: 5 µL

2) Mass Spectrometer
   (i) Ionization method: Electro-Spray Ionization (ESI) method
   (ii) Mode: Positive
   (iii) Capillary voltage: 3.0 kV
   (iv) Ion source temperature: 120 °C
   (v) Desolvation temperature: 400 °C
   (vi) Cone voltage: Shown in table 1
   (vii) Collision energy: Shown in table 1
   (viii) Monitor ion: Shown in table 1

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<th>Agrichemicals</th>
<th>Precursor ion (m/z)</th>
<th>Product ion (determination) (m/z)</th>
<th>Product ion (validation) (m/z)</th>
<th>Cone voltage (V)</th>
<th>Collision energy (eV)</th>
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</table>

b) Calibration curve preparation
1) Inject 5 µL of respective mixture standard solutions for calibration curve preparation into the LC-MS/MS, record the chromatogram of ion (m/z) for determination and ion (m/z) for validation of materials subjected to measurement.
2) Calculate the peak area ratio or height ratio of ion (m/z) for determination and ion (m/z) for validation of respective materials subjected to measurement.
3) Prepare a curve for the relationship between the concentration of material subjected to measurement and the peak area or height of ion (m/z) for determination of respective mixture solutions for the calibration curve preparation. Prepare a calibration curve when the sample is measured.

c) Sample measurement
1) Subject 5 µL of the sample solution to the same procedure as in b) 2) - 3)\(^{(12)}\).
2) Obtain the content of materials subjected to measurement from the calibration curve of the peak area or height to calculate materials subjected to measurement in the analytical sample.

Note \(^{(12)}\) Confirm that the ratio against the peak area ratio or height ratio of the standard solution is within the range of about ± 30 %. In addition, the peak area ratio or height ratio depends on the concentration.

(5.5) Calculation
Calculate the respective concentration of agricultural chemicals in the analytical sample by the following formula. in the analytical sample by the following formula:
Respective concentration of agricultural chemicals in the analytical sample (µg/kg)

\[ = \left( \frac{A \times B \times 10}{C} \right) \]

**A**: Concentration (ng/mL) of respective materials subjected to measurement in the final sample solution obtained from the calibration curve

**B**: Dilution factor in the case that the final sample solution is further diluted because it exceeds the upper limit of the calibration curve.

**C**: Specific gravity of the analytical sample (density) (g/mL)

**Comment 3** A recovery testing was conducted using fluid home garden-use mixed fertilizer (3 kinds) and fluid mixed fertilizer (2 kinds), as a result, the mean recovery at additive level of 4000 µg/kg and 400 µg/kg (However, 4000 µg/kg and 400 µg/kg of pyrethrin contain total of pyrethrin I/II) were 77.0 % - 104.5 % and 85.6 % - 107.9 % respectively.

Table 2 shows results and analysis results from a collaborative study for test method validation.

Additionally, the minimum limit of quantification for respective agrichemicals of the test method is about 10 µg/kg.
### Table 2  Analysis results of a collaborative study for the testing method validation of multicomponent analysis of Agrichemicals

<table>
<thead>
<tr>
<th>Agrichemicals</th>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^2$ (µg/kg)</th>
<th>Additive amount (µg/kg)</th>
<th>Recovery (%)</th>
<th>RSD $^3$ (%)</th>
<th>RSD $^4$ (%)</th>
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<tbody>
<tr>
<td>Abamectin</td>
<td>Home garden-use mixed fertilizer1</td>
<td>8</td>
<td>286.8</td>
<td>333.3</td>
<td>86.1</td>
<td>13.3</td>
<td>14.4</td>
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<td></td>
<td>Home garden-use mixed fertilizer2</td>
<td>8</td>
<td>358.9</td>
<td>416.7</td>
<td>86.1</td>
<td>13.4</td>
<td>14.8</td>
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<tr>
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<td>Home garden-use mixed fertilizer3</td>
<td>8</td>
<td>425.8</td>
<td>500.0</td>
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<td>8.6</td>
<td>11.6</td>
</tr>
<tr>
<td></td>
<td>Fluid mixed fertilizer1</td>
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<td>288.6</td>
<td>333.3</td>
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<td>Fluid mixed fertilizer2</td>
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<td>333.3</td>
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<td>10.1</td>
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<td>85.1</td>
<td>7.0</td>
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<td>500.0</td>
<td>87.2</td>
<td>5.8</td>
<td>7.4</td>
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</tbody>
</table>

1) Number of laboratories used in analysis
2) Gross mean (n = number of laboratories × number of repeated tests (2))
3) Repeatability (relative standard deviation)
4) Reproducibility (relative standard deviation)

Table 2 (Continued)

<table>
<thead>
<tr>
<th>Agrichemicals</th>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean $^2$ (µg/kg)</th>
<th>Additive amount (µg/kg)</th>
<th>Recovery (%)</th>
<th>RSD $^3$ (%)</th>
<th>RSD $^4$ (%)</th>
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<tr>
<td>Pyrethrin I</td>
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<td>160.7</td>
<td>186.0</td>
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<td>12.8</td>
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<td>221.0</td>
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<td>6.3</td>
<td>8.3</td>
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</tbody>
</table>
Flow sheet for simultaneous analysis of 6 kinds of agrichemicals: The flow sheet for simultaneous analysis of 6 kinds of agrichemicals in fertilizer is shown below.

Figure 1 Flow sheet for residue agrichemicals multicomponent analysis
((1) : simultaneous analysis of 6 kinds of agrichemicals) in fertilizers
(Extraction procedure)
Figure 2  Flow sheet for residue agrichemicals multicomponent analysis
((1) : simultaneous analysis of 6 kinds of agrichemicals) in fertilizers
(Cleanup (1) and cleanup (2) and measurement procedure)
Reference: Selected Reaction Monitoring chromatograms of mixture standard solution for calibration curve preparation and sample solution (fluid home garden-use mixed fertilizer) are shown below.

Peak No.1: AbamectinB1a
No.2: IvermectinB1a
No.3: Eprinomectin B1a
No.4: Rotenone
No.5: Piperonylbutoxide
No.6: Pyrethrin I
No.7: Pyrethrin II

1) Mixed standard solution 2) Sample solution

Reference diagram Selected Reaction Monitoring chromatograms of respective agricultural chemicals

1) Mixture standard solution (the equivalents of 2,500 pg as respective agrichemicals)
(For pyrethrin, the equivalents of 2,500 pg of the total of pyrethrin I/II)
2) Sample solution (fluid home garden-use mixed fertilizer, additive of the equivalents of 400 µg/kg in the sample)
(For pyrethrin, the equivalents of 400 µg/kg of the total of pyrethrin I/II)

LC-MS/MS measurement conditions
Column: ACQUITY UPLC HSS C18 (2.1-mm inner diameter, 100-mm long, 1.8-µm particle diameter)
Flow rate: 0.2 mL/min
Other conditions are according to the examples of the measurement conditions in (5.4)
a) LC-MS/MS.
8.3.2 Residue agrichemicals multicomponent analysis (2)

8.3.2.a Gas Chromatography

(1) **Compounds subjected to analysis** β-HCH (β-BHC), γ-HCH (γ-BHC), o,p′-DDD, p,p′-DDD, o,p′-DDE, p,p′-DDE, o,p′-DDT, p,p′-DDT, aldrin, endrin, dieldrin, trans-chlordane, cis-chlordane, trans-nonachlor, cis-nonachlor, heptachlor, heptachlor epoxide and hexachlorobenzene.

(2) **Summary**

This testing method is applicable to compost and straw, raw materials of compost. This testing method is classified as Type D and its symbol is 8.3.2.a-2017 or AG-C-2.a-1. Extract respective agricultural chemicals in fertilizers or raw materials with acetonitrile and water, refine by using a porous diatomaceous earth column, a gel permeation chromatograph and a synthetic magnesium silicate cartridge column, and then measure with an electron capture detector equipped gas chromatograph to obtain compounds subjected to analysis in an analytical sample. In addition, the performance of this testing method is shown in Comment 7.

(3) **Reagent, etc.:** Reagents and water are as shown below.

a) **Water:** Water of A3 specified in JIS K 0557.

b) **Acetonitrile:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8039 or a reagent of equivalent quality.

c) **Hexane:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8825 or a reagent of equivalent quality.

d) **Sodium chloride:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8039 or a reagent of equivalent quality.

e) **Cyclohexane:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) or a reagent of equivalent quality.

f) **Acetone:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8040 or a reagent of equivalent quality.

g) **Diethyl ether:** A reagent of agricultural chemicals residue/PCB testing grade (concentration: no less than 300) specified in JIS K 8357 or a reagent of equivalent quality.

h) **2,2,4-trimethylpentane:** A reagent of HPLC grade or a reagent of equivalent quality.

i) **Respective agricultural chemicals standard solutions (0.2 mg/mL)** (1): Put about 0.02 g of β-HCH (β-BHC) [C₆H₁₂Cl₆] (2), γ-HCH (γ-BHC) [C₆H₁₆Cl₂] (2), o,p′-DDD [C₁₂H₁₀Cl₄] (2), p,p′-DDD [C₁₂H₁₀Cl₄] (2), o,p′-DDE [C₁₂H₁₆Cl₂] (2), p,p′-DDE [C₁₂H₁₆Cl₂] (2), o,p′-DDT [C₁₄H₁₂Cl₄], endrin [C₁₃H₁₄Cl₄], dieldrin [C₁₃H₁₄Cl₄], trans-chlordane [C₁₀H₆Cl₆] (2), cis-chlordane [C₁₀H₆Cl₆] (2), trans-nonachlor [C₁₀H₆Cl₆] (2), cis-nonachlor [C₁₀H₆Cl₆] (2), heptachlor [C₁₀H₆Cl₇] (2), heptachlor epoxide [C₁₀H₆Cl₇] (2), and hexachlorobenzene [C₆Cl₆] (2) to a weighing dish, and measure the mass to the order of 0.1 mg. Dissolve them with 20 mL of acetone, transfer to 100 mL volumetric flasks respectively, and add 2,2,4-trimethylpentane up to the marked line.

j) **Mixture standard solution (1 µg/mL)** (1): Transfer 1 mL of respective agricultural chemicals standard solutions to a 100-mL volumetric flask and add 2,2,4-trimethylpentane-acetone (4+1) up to the marked line.

k) **Mixture standard solution for calibration curve preparation (0.02 µg/mL - 0.2 µg/mL)** (1): In the case of usage, transfer 1 mL - 10 mL of mixture standard solution (1 µg/mL) to 50 mL volumetric flasks step-by-step, and add 2,2,4-trimethylpentane-acetone (4+1) up to the marked line.

l) **Mixture standard solution for calibration curve preparation (0.005 µg/mL - 0.02 µg/mL)** (1): In the case of usage, transfer 2.5 mL - 10 mL of mixture standard solution (0.1 µg/mL) to 50 mL volumetric flasks step-by-step, and add 2,2,4-trimethylpentane-acetone (4+1) up to the marked line.
marked line.

**Note** (1) This is an example of preparation; prepare an amount as appropriate.
(2) A standard reagent is commercially sold.

**Comment 1** A standard reagent of respective agricultural chemicals is sold by FUJIFILM Wako Pure Chemical Co., Ltd., Kanto Chemical Co., Inc. and Hayashi Pure Chemical Industries., Ltd.

(4) **Apparatus and instruments:** Apparatus and instruments are shown below.

a) **Gas Chromatograph (GC):** GC specified in JIS K 0114 that satisfies the following requirements.

1) **Sample injector:** An injector that enables split less system.
2) **Capillary column:** A capillary column (0.25-mm inner diameter and 30-m long) made of fused silica. 14 % cyanopropylphenyl -86 % dimethyl polysiloxane chemically bonds to the inner surface of a capillary column with 0.25 μm thickness.
3) **Detection unit:** Electron capture detector (ECD)

b) **Gel permeation chromatograph (GPC):** Preparative liquid chromatograph specified in JIS K 0135 that satisfies the following requirements. No detector is required.

1) **Sample injector:** A sample injector that can inject 5 mL of sample solution.
2) **Column:** A 20-mm inner diameter 300-mm long stainless-steel column tube filled with styrendivynylbenzene copolymer system hard gel
3) **Guard column:** A 20-mm inner diameter 100-mm long stainless steel column tube filled with styrendivynylbenzene copolymer system hard gel
4) **Fraction collector:** A fraction collector that can set up a fraction to which agrichemical components elute.

c) **Shaking apparatus**
d) **Concentrator:** An evaporator whose temperature can be adjusted up to 40 ºC.
e) **Filter:** A funnel for filtering under reduced pressure (compatible filter diameter: 60 mm)
f) **Porous diatomaceous earth cartridge column:** A column that is filled with the porous diatomaceous earth (capacity: 20 mL).
g) **Synthetic magnesium silicate cartridge column:** A cartridge column that is filled with 910 mg of synthetic magnesium silicate.
h) **Membrane filters:** Made of PTFE (pore size is no more than 0.5 μm)

**Comment 2** Column for GC is sold under the production name DB-1701, Rtx-1701, SPB-1701, etc. A column which has actually isolated compounds subjected to analysis should be used.

**Comment 3** Gel permeation chromatograph (GPC) is a preparative liquid chromatograph that collects a fraction of a material subjected to measurement sieved and isolated by packing materials of the column for GPC according to the size of the molecule of the material. A column for GPC is sold under the production name Shodex CLNpak EV-2000 AC, etc. In addition, a guard column for GPC is sold under the production name Shodex CLNpak EV-G AC, etc.

**Comment 4** A funnel for filtering under reduced pressure is sold under the production name KIRIYAMA Funnel SB-60, KIRIYAMA Funnel SU-60, etc.

**Comment 5** A porous diatomaceous earth cartridge is commercially sold under production name Chem Elut (20 mL), etc.

**Comment 6** Synthetic magnesium silicate is commercially sold under the production names such as Sep-Pak Florisil Plus Long Cartridge (910 mg).
Comment 7 A membrane filter is sold under the production name HLC-DISK 25 Solvent system (pore size: 0.45 µm), DISMIC 25JP050, Millex FH (diameter: 25 mm, pore size: 0.45 µm), etc.

(5) Test procedures
(5.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample, and put it into a 200-mL ground-in stopper Erlenmeyer flask.
   b) Add 20 mL of acetonitrile–water (3:1) to moisten.
   c) After leaving at rest for ten minutes, add 100 mL of acetonitrile and shake to mix for about 30 minutes.
   d) Place a 300-mL round-bottom flask under the filter and filtrate the extract under reduced pressure with a filter paper (type 5 B)
   e) Conduct cleanup of the previous Erlenmeyer flask and residue with 50 mL of acetonitrile successively, similarly filtrate under reduced pressure and pool with the filtrate in d) to make the extract.

(5.2) Cleanup (1): Conduct cleanup (1) as shown below.
   a) Concentrate under reduced pressure in a water bath of no more than 40 ºC until most of the extract dries up.
   b) Add 20 mL of sodium chloride saturated solution, put into a porous diatomaceous earth cartridge column and leave it at rest for about 5 minutes.
   c) Place a 300-mL round-bottom flask under the same cartridge column, wash the vessel 3 times with about 20 mL of hexane to add the washing to the same cartridge column successively and allow the solution to elute until the surface of the solution reaches the top of packing materials.
   d) Further add about 60 mL of hexane into the same cartridge and allow the solution to elute until the surface of the solution reaches the top of packing materials.
   e) After conducting vacuum concentration of elute in a water bath of no more than 40 ºC until most of the elute dries up, send a nitrogen gas to dry up the elute (3), and add 10 mL of cyclohexane–acetone (4:1) to dissolve the residue.
   f) Filter with a membrane filter (pore size: no more than 0.5 µm).

Note (3) There is a possibility for agricultural chemicals to vaporize if it is dried up excessively.

(5.3) Cleanup (2): Conduct cleanup (2) as shown below.
   a) Inject 5 mL of the filtrate in (5.2) e) into a gel permeation chromatograph and transfer a fraction eluted from respective agrichemicals which are determined according to the procedure condition in b) into a 100-mL round-bottom flask.
   b) Procedure conditions for a gel permeation chromatograph (GPC): An example of procedure conditions for a gel permeation chromatograph (GPC) is shown below. Set up the measurement conditions considering it:
      1) Column: A styrene-divinylbenzene copolymer column (20-mm inner diameter, 300-mm long, 15-µm particle diameter)
      2) Guard column: A styrene-divinylbenzene copolymer column (20-mm inner diameter, 100-mm long, 15-µm particle diameter)
      3) Eluent: Cyclohexane-acetone (4:1)
      4) Flow rate: 5 mL/min
      5) Preparative fraction: 70 mL - 120 mL
   c) After conducting vacuum concentration of elute in a water bath of no more than 40 ºC until
most of the elute dries up, send a nitrogen gas to dry up the elute \(^{(3)}\), and add 2 mL of hexane to dissolve the residue.

\((5.4)\) **Cleanup (3):** Conduct cleanup (3) as shown below.

a) Conduct cleanup of a synthetic magnesium silicate cartridge column (910 mg) with about 5 mL of hexane.

b) Place a 50-mL round-bottom flask under the same cartridge column, put the solution in (5.3) to the same cartridge column, and allow the solution to overflow until the surface of the solution reaches the top of packing materials.

c) Wash the vessel with about 2 mL of hexane 2 times and add washing to the same cartridge successively to allow it to overflow.

d) Further add about 15 mL of hexane–diethyl ethel (9+1) into the same cartridge and allow respective materials subjected to measurement to elute.

e) After conducting vacuum concentration of the elute in a water bath of no more than 40 °C until most of the elute dries up, send a nitrogen gas to dry up the elute \(^{(3)}\). And add 1 mL of 2,2,4-trimethylpentane–acetone (4+1) \(^{(4)}\) to dissolve the residue, making the sample solution.

**Note** (4) If there is a possibility that the concentration of respective agricultural chemicals in the sample solution exceed the maximum limit of the calibration curve, dilute a predetermined amount of the sample solution with 2,2,4-trimethylpentane–acetone (4+1).

\((5.4)\) **Measurement:** Conduct the measurement as indicated in JIS K 0114 and as shown below. Specific measurement procedures are according to the operation method of a Gas Chromatograph used in measurement.

a) **The measurement conditions of Gas Chromatograph:** An example of measurement conditions for a Gas Chromatograph is shown below. Set up the measurement conditions considering it:

1) **Sample injection method:** split less injection method (1min)
2) **Temperature of sample injector:** 250 °C
3) **Capillary column:** A capillary column (0.25-mm inner diameter, 30-m long, 0.25 µm layer thickness) made of fused silica. 14 % cyanopropylphenyl–86 % dimethyl polysiloxane chemically bonds to the inner surface of the capillary column with 0.25 µm thickness.
4) **Column bath temperature:** 60 °C (1 min) → (20 °C/min) → 180 °C → (2 °C/min) → 260 °C → (5 °C/min) → 275 °C (1 min)
5) **Carrier gas:** helium, **Flow rate:** 1.5 mL/min
6) **Addition gas:** Nitrogen, **Flow rate:** 60 mL/min
7) **Detection unit:** Electron capture detector (ECD)
8) **Detector temperature:** 280 °C

b) **Calibration curve preparation**
1) Inject 1 µL of respective mixture standard solutions for calibration curve preparation into a GC, record a chromatogram and obtain the peak area or height.
2) Prepare a curve for the relationship between the concentration and the peak area or height of respective mixture solutions for the calibration curve preparation. Prepare a calibration curve when the sample is measured.

c) **Sample measurement**
1) Subject 1 µL of sample solution to the same procedure as in b) 1)
2) Obtain the content of materials subjected to measurement from the calibration curve of the
peak area or height to calculate materials subjected to measurement in the analytical sample.

Comment 8 Additive recovery testing of compounds subjected to analysis in compost was conducted, as a result, the mean recovery rates at additive level of 20 µg/kg and 50 µg/kg were 82.1 % - 118.1 % and 62.5 % - 120.2 % respectively. The results of the repeatability tests on different days using compost to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Note that α-HCH (α-BHC), δ-HCH (δ-BHC), and oxychlordane, which were subjected to simultaneous analysis, were excluded from the compounds subjected to analysis because sufficient recovery rates were not obtained.

Additionally, the minimum limit of quantification for respective agrochemicals of the test method is no more than 20 µg/kg.

References
(6) **Flow sheet for simultaneous analysis of chloride pesticides:** The flow sheet for simultaneous analysis of chloride pesticides in fertilizer is shown below.

**Figure 1** Flow sheet for residue agrichemicals multicomponent analysis
(2) simultaneous analysis of chloride agrichemicals in fertilizers
(Extraction procedure)

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td><strong>5.00 mL analytical sample</strong></td>
</tr>
<tr>
<td>2.</td>
<td>Leave at rest</td>
</tr>
<tr>
<td>3.</td>
<td>Shake to mix</td>
</tr>
<tr>
<td>4.</td>
<td>Filter under reduced pressure</td>
</tr>
<tr>
<td>5.</td>
<td>Extract</td>
</tr>
<tr>
<td>6.</td>
<td>Filter paper (type 5 B), 300-mL round-bottom flask (1)</td>
</tr>
<tr>
<td>7.</td>
<td>Pool washing with the filtrate</td>
</tr>
</tbody>
</table>

**Figure 2** Flow sheet for residue agrichemicals multicomponent analysis
(2) simultaneous analysis of chloride agrichemicals in fertilizers
(Cleanup (1) procedure)

<table>
<thead>
<tr>
<th>Step</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Extract</td>
</tr>
<tr>
<td>2.</td>
<td>Concentration under reduced pressure</td>
</tr>
<tr>
<td>3.</td>
<td>Cleanup (1)</td>
</tr>
<tr>
<td>4.</td>
<td>300-mL round-bottom flask (2)</td>
</tr>
<tr>
<td>5.</td>
<td></td>
</tr>
<tr>
<td>6.</td>
<td>Concentration under reduced pressure</td>
</tr>
<tr>
<td>7.</td>
<td>Exsiccation</td>
</tr>
<tr>
<td>8.</td>
<td>Filter</td>
</tr>
<tr>
<td>9.</td>
<td>Filtrate</td>
</tr>
</tbody>
</table>
Gel permeation chromatograph
Inject 5 mL of the filtrate

Transfer 70 mL - 120 mL of eluted fraction into a 100-mL round-bottom flask

40 °C
Nitrogen gas

→ About 2 mL of hexane

Clean up (2)
Synthetic magnesium silicate cartridge column

→ 50-mL round-bottom flask
→ Wash the 100-mL round-bottom flask 3 times with about 2 mL of hexane, add to the same cartridge column successively
→ About 15 mL of hexane-diethyl ethel (9+1)

Concentration under reduced pressure
Exsiccation

40 °C
Nitrogen gas

→ 1 mL of 2,2,4-trimethylpentane–acetone (4+1)

Sample solution

Measurement
GC (ECD)

Figure 3 Flow sheet for residue agrichemicals multicomponent analysis
((2) : simultaneous analysis of chloride agrichemicals) in fertilizers
(Cleanup (2) and cleanup (3) and measurement procedure)
8.4  Sodium
8.4.a Flame atomic absorption spectrometry

(1) Summary
This testing method is applicable to fertilizers containing organic matters. This testing method is classified as Type D and its symbol is 8.4.a-2017 or Na.a-1.

Pretreat an analytical sample with incineration and hydrochloric acid, spray into an acetylene–air flame, and measure the atomic absorption with sodium at a wavelength of 589.0 nm or 589.6 nm to obtain sodium (Na) in an analytical sample. In addition, the performance of this testing method is shown in Comment 3.

(2) Reagent: Reagents are as shown below.
   a) Hydrochloric acid: A JIS Guaranteed Reagent specified in JIS K 8180 or a reagent of equivalent quality.
   b) Sodium standard solution (Na 1 mg/mL) (1): Heat sodium chloride specified in JIS K 8150 at 600 °C ± 10 °C for about 1 hour, let it stand to cool in a desiccator, and weigh 2.542 g to a weighing dish. Dissolve in a small amount of water, transfer to a 1000-mL volumetric flask, and add water up to the marked line.
   c) Sodium standard solution (Na 0.1 mg/mL) (1): Transfer 20 mL of sodium standard solution (Na 1 mg/mL) to a 200-mL volumetric flask, and add hydrochloric acid (1+23) up to the marked line.
   d) Sodium standard solutions (Na 1 µg/mL - 10 µg/mL) for the calibration curve preparation (2): Transfer 2.5 mL - 25 mL of sodium standard solution (Na 0.1 mg/mL) to 250-mL volumetric flasks step-by-step, and add hydrochloric acid (1+23) up to the marked line (2).
   e) Blank test solution for the calibration curve preparation: Hydrochloric acid (1+23) used in the procedures in d) (3).

Note  (1) This is an example of preparation; prepare an amount as appropriate.
   (2) When using a device model which cannot degrade the sensitivity of the device by tilting the burner head, conduct the model appropriate dilution. (For example, 0.1 µg/mL - 4 µg/mL)
   (3) When preserving, use a container, which can be sealed tightly, made of material such as polypropylene, PTFE, etc. that sodium hardly elutes.

Comment 1 Instead of the sodium standard solution in (2) b), a sodium standard solution (Na 0.1 mg/mL 1 mg/mL or 10 mg/mL for Atomic Absorption Spectrometry traceable to National Metrology can also be used.

(3) Instruments: Instruments are as shown below:
   a) Flame atomic absorption spectrometer: An atomic absorption spectrometer specified in JIS K 0121.
      1) Light source: A sodium hollow cathode lamp
      2) Gas: Gas for heating by flame
         (i) Fuel gas: acetylene
         (ii) Auxiliary gas: Air sufficiently free of dust and moisture.
   b) Electric furnace: An electric furnace that can be adjusted to 550 °C ± 5 °C.
   c) Hot plate or sand bath: A hot plate whose surface temperature can be adjusted up to 250 °C. Adjust the amounts of gas and silica sand of a sand bath so that the sand bath temperature can be set to 250 °C.
(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
   a) Weigh 5.00 g of an analytical sample and transfer to a 200-mL - 300-mL tall beaker.
   b) Ignite at 550 °C ± 5 °C for no less than 4 hours to incinerate (4).
   c) Put the tall beaker in an electric furnace, and heat gently to char (4).
   d) After standing to cool, moisten the residue with a small amount of water, gradually add about 10 mL of hydrochloric acid, and further add water to make 20 mL.
   e) Cover the tall beaker with a watch glass, and heat on a hot plate or a sand bath to boil for about 5 minutes.
   f) After standing to cool, transfer the solution to a 250-mL - 500-mL volumetric flask with water.
   g) After standing to cool, transfer the solution to a 250-mL - 500-mL volumetric flask with water.
   h) Filter with Type 3 filter paper to make a sample solution.

Note (4) Example of charring and incineration procedure: After raising the temperature from room temperature to about 250 °C in 30 minutes to 1 hour, continue heating for about 1 hour and further raise to 550 °C in 1 to 2 hours.

Comment 2 The procedure in (4.1) is the same as the procedure in (4.1.2) in 4.2.1.a.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0121 and as shown below. Specific measurement procedures are according to the operation method of the atomic absorption spectrometer used in measurement.
   a) Measurement conditions for the atomic absorption spectrometer: Set up the measurement conditions for the atomic absorption spectrometer considering the following:
      Analytical line wavelength: 589.0 nm or 589.6 nm
   b) Calibration curve preparation
      1) Spray the sodium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation into a flame, and read the indicated value at a wavelength of 589.0 nm or 589.6 nm.
      2) Prepare a curve for the relationship between the sodium concentration and the indicated value of the sodium standard solutions for the calibration curve preparation and the blank test solution for the calibration curve preparation.
   c) Sample measurement
      1) Transfer a predetermined amount of the sample solution (the equivalents of 0.1 mg - 1 mg as Na) (5) to a 100-mL volumetric flask.
      2) Add hydrochloric acid (1+23) to the marked line.
      3) Subject to the same procedure as in b) 1) to read the indicated value.
      4) Obtain the sodium content from the calibration curve, and calculate the sodium (Na) in the analytical sample.

Note (5) Sample a predetermined amount of sample solution according to the device model in Note (2).

Comment 3 Additive recovery testing with triplicates measurement was conducted using fish caked powder, fish waste processed fertilizers, rape seed meal and its powder, composted sludge fertilizers and compost, as a result, the mean recovery rate at the additive concentration of sodium in the range of 1 % (mass fraction) - 10 % (mass fraction) was 97.0 % - 103 %.
The results of the repeatability tests on different days using fish caked powder (sample to which sodium chloride is added) and compost to evaluate precision were
analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Additionally, the minimum limit of quantification of this testing method is about 0.02 % (mass fraction).

Table 1 Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $T$ $(%)$</th>
<th>$s_r$ $(%)$</th>
<th>$RSD_r$ $(%)$</th>
<th>Intermediate precision $s_{I(T)}$ $(%)$</th>
<th>Intermediate relative standard deviation $RSD_{I(T)}$ $(%)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fish cakes powder</td>
<td>5</td>
<td>9.08</td>
<td>0.06</td>
<td>0.6</td>
<td>0.09</td>
<td>1.0</td>
</tr>
<tr>
<td>Compost</td>
<td>5</td>
<td>0.0973</td>
<td>0.0019</td>
<td>2.0</td>
<td>0.0037</td>
<td>3.8</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $T$)
3) Intermediate standard deviation $s_r$
4) Repeatability standard deviation $s_r$
5) Repeatability relative standard deviation $RSD_r$
6) Intermediate relative standard deviation $RSD_{I(T)}$
7) Intermediate relative standard deviation $RSD_{I(T)}$

References

(5) Flow sheet for sodium testing method: The flow sheet for sodium testing method in fertilizers is shown below:

- 5.00 g analytical sample
- Weigh into a 200-mL - 300-mL tall beaker.
- Charring
- Heat gently
- 550 ºC ± 5 ºC, no less than 4 hours
- Incineration
- Standing to cool
- Room temperature
- A small amount of water, moisten the residue
- About 10 mL of hydrochloric acid
- Water (up to about 20 mL)
- Heating
- Cover with a watch glass, and boil for 5 minutes.
- Standing to cool
- Room temperature
- Transfer
- 250-mL - 500-mL volumetric flask, water (up to the marked line)
- Water (up to the marked line)
- Filtration
- Type 3 filter paper
- Sample solution

Figure 1 Flow sheet for sodium in fertilizers (Extraction procedure)
100-mL volumetric flask, water

Atomic absorption spectrometer
(589.0 nm or 589.6 nm)

Figure 2  Flow sheet for sodium in fertilizers (Measurement procedure)
8.5 Guanylurea nitrogen
8.5.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type B and its symbol is 8.5.a-2017 or GU-N.a-1.

Add water to an analytical sample to extract guanylurea, introduce it to a High-Performance Chromatograph (HPLC) to isolate it with a weak acid ion-exchange column, and then measure at wavelength 190 nm to obtain guanylurea nitrogen (GU-N) in an analytical sample. In addition, the performance of this testing method is shown in Comment 6.

Biuret nitrogen (B-N), dicyandiamide nitrogen (Dd-N), urea nitrogen(U-N) and guanidine nitrogen (Gd-N) can be simultaneously quantified by using this method. (Refer to Comment 5).

(2) Reagent, etc.: Reagents and water are as shown below.
  b) Potassium dihydrogen phosphate: A JIS Guaranteed Reagent specified in JIS K 9007 or a reagent of equivalent quality.
  c) Phosphoric acid: A JIS Guaranteed Reagent specified in JIS K 9005 or a reagent of equivalent quality.
  d) Guanylurea nitrogen standard solution (GU-N 2 mg/mL): Put 0.540 g of guanylurea sulfate \([C_4H_12N_8O_2 \cdot H_2SO_4]\) (2) in a weighing dish and measure the mass to the order of 0.1 mg. Add a small amount of water to dissolve, then transfer to a 100-mL volumetric flask and add water up to the marked line.
  e) Guanylurea nitrogen standard solution (GU-N 200 g/mL): Put 10 mL of guanylurea nitrogen standard solution (GU-N 2 mg/mL) to a 100-mL volumetric flask and add water up to the marked line.
  f) Guanylurea nitrogen standard solution (GU-N 50 µg/mL - 100 µg/mL): Put 25 mL - 50 mL of guanylurea nitrogen standard solution (GU-N 200 µg/mL) to 100-mL volumetric flasks and add water up to the marked line.
  g) Guanylurea nitrogen standard solution for the calibration curve preparation (GU-N 1 µg/mL - 50 µg/mL): At the time of usage, put 1 mL - 50 mL of guanylurea nitrogen standard solution (GU-N 100 µg/mL) to 100-mL volumetric flasks step-by-step and add water up to the marked line.

Note  (1) This is an example of preparation; prepare an amount as appropriate.
  (2) A reagent of no less than 98 % (mass fraction) in purity as guanylurea sulfate is commercially sold.

Comment 1 Guanylurea sulfate is commercially sold by Kanto Chemical Co., Inc. and Tokyo Chemical Industry Co., Ltd.

(3) Apparatus and instruments: Apparatus and instruments are shown below.
  a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.
  1) Column: A 7.5-mm inner diameter 100-mm long stainless steel column tube filled with weak acid ion-exchange resin.
  2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45ºC.
  3) Detection unit: An absorptiometric detector that can measure at wavelength around 190 nm.
  b) Magnetic stirrer:
  c) High speed centrifugal separator: A centrifugal separator that can work at 8000 × g - 10000 × g.
Comment 2 A column is sold under the production name Asahipak ES-502C 7C.

(4) Test procedure
(4.1) Extraction: Conduct extraction as shown below.
(4.1.1) Powdery test sample
a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL- ground-in stopper Erlenmeyer flask.
b) Add 100 mL of water and stir it by using a magnetic stirrer for about 10 minutes.
c) After allowing to stand still, transfer a supernatant solution to a 1.5-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes and use the supernatant as the extract.

Note (3) If there is a possibility that the guanyourea nitrogen (GU-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with water.

(4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement.

(5) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

(4.1.2) Fluid test sample
a) Weigh 1.00 g of an analytical sample, and put it in a 100-mL volumetric flask.
b) Add about 50 mL of water, and shake to mix.
c) Add water up to the marked line and transfer to a 1.5-mL ground-in stopper centrifugal precipitate tube.
d) Centrifuge it at 8000 × g - 10000 × g centrifugal force for about five minutes and use the supernatant as the extract.

Note (6) If there is a possibility that the guanyourea nitrogen (GU-N) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of solution with water.

Comment 3 Instead of procedures in (4.1.1) c) - d) or (4.1.2) c) - d), it is allowed to filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) Measurement: Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) Measurement conditions for a High-Performance Liquid Chromatograph (HPLC): An example of measurement conditions is shown below. Set up the measurement conditions considering it:
1) Column: A weak acid ion-exchange resin column (7.5-mm inner diameter, 100-mm long, 5-µm - 10-µm particle diameter)
2) Column bath temperature: 40 °C
3) Eluent: Dissolve 3.92 g of potassium dihydrogenphosphate and 0.12 g of phosphoric acid in water to make 1000 mL. Filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.
4) **Flow rate**: 0.6 mL/min
5) **Injection volume**: 10 µL
6) **Detection unit**: An absorptiometric detector, measurement wavelength: 190 nm

**Comment 4** Eluent can be prepared as follows. Dissolve 19.6 g of potassium dihydrogenphosphate and 0.584 g of phosphoric acid with water to make 500 mL and store in a refrigerator. At the time of usage, dilute a predetermined volume of the solution by a factor of 10 and filter with a membrane filter (aperture diameter: no more than 0.5-µm) made of hydrophilic PTFE.

**b) Calibration curve preparation**
1) Inject 10 µL of respective standard solutions for the calibration curve preparation to an HPLC, record chromatogram at wavelength 190 nm and obtain the peak height.
2) Prepare a curve for the relationship between the guanylurea nitrogen (GU-N) concentration and the peak height at wavelength 190 nm of the respective standard solutions for the calibration curve preparation.

**c) Sample measurement**
1) Subject 10 µL of sample solution to the same procedure as in **b) 1)**
2) Obtain the guanylurea nitrogen (GU-N) content from the peak height using the calibration curve to calculate guanylurea nitrogen (GU-N) in the analytical sample.

**Comment 5** This testing method enables the simultaneous measurement of biuret nitrogen (B-N), urea nitrogen (U-N), dicyandiamide nitrogen (Dd-N), guanidine urea (Gd-N) and guanylurea nitrogen (GU-N). In that case, see **5.10.a Comment 5**.

**Comment 6** Additive recovery testing was conducted using a preparation sample for a guanylurea fertilizer (one brand). As a result, the mean recovery at additive level of 36.7 % (mass fraction), 35.2 % (mass fraction) and 33.4 % (mass fraction) were 103.8 %, 104.6 % and 105.6 % respectively.

The results of the repeatability tests on different days using a guanylurea fertilizer to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability. Table 2 shows results and analysis results from a collaborative study for testing method validation. Additionally, the minimum limit of quantification of this testing method is about 0.006 % (mass fraction).

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^{2)}$ (%$^{3)}$</th>
<th>Repeatability $^{4)}$</th>
<th>Intermediate precision $^{5)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T^{1)}$</td>
<td></td>
<td>$s_r$ ($^{4)}$ (%$^{3)}$)</td>
<td>$RSD_r$ ($^{5)}$ (%)</td>
</tr>
<tr>
<td>Guanylurea nitrogen</td>
<td>5</td>
<td>37.0</td>
<td>0.3</td>
<td>0.7</td>
</tr>
<tr>
<td>fertilizer</td>
<td></td>
<td></td>
<td>0.3</td>
<td>0.8</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days ($T$) $\times$ the number of duplicate testing (2))
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation
Table 2  Results and analysis results from a collaborative study for the test method validation of guanylurea nitrogen

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Number of laboratories</th>
<th>Mean (^2) (%)(^3)</th>
<th>(s_r) (^4) (%)(^3)</th>
<th>(RSD_r) (^5) (%)</th>
<th>(s_R) (^6) (%)(^3)</th>
<th>(RSD_R) (^7) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compound fertilizer 1</td>
<td>12</td>
<td>2.20</td>
<td>0.09</td>
<td>4.2</td>
<td>0.17</td>
<td>7.7</td>
</tr>
<tr>
<td>Compound fertilizer 2</td>
<td>11</td>
<td>4.38</td>
<td>0.07</td>
<td>1.5</td>
<td>0.19</td>
<td>4.3</td>
</tr>
<tr>
<td>Compound fertilizer 3</td>
<td>11</td>
<td>5.83</td>
<td>0.08</td>
<td>1.4</td>
<td>0.52</td>
<td>8.9</td>
</tr>
<tr>
<td>Compound fertilizer 4</td>
<td>12</td>
<td>7.43</td>
<td>0.43</td>
<td>5.7</td>
<td>0.78</td>
<td>10.5</td>
</tr>
<tr>
<td>Guanylurea nitrogen fertilizers</td>
<td>12</td>
<td>30.3</td>
<td>0.4</td>
<td>1.5</td>
<td>1.1</td>
<td>3.6</td>
</tr>
</tbody>
</table>

1) Number of laboratories used in analysis  
2) Mean (\(n = \) number of laboratories x number of samples (2))  
3) Mass fraction  
4) Repeatability standard deviation  
5) Repeatability relative standard deviation  
6) Reproducibility standard deviation  
7) Reproducibility relative standard deviation

References


(5) **Flow sheet for testing method:** The flow sheet for guanylurea nitrogen in fertilizers is shown below:

**Figure 1** Flow sheet for guanylurea nitrogen in fertilizers
(Extraction procedure (4.1.1) and measurement)

**Figure 2** Flow sheet for guanylurea nitrogen in fertilizers
(Extraction procedure (4.1.2) and measurement)
**Reference:** Chromatogram of the standard solution for calibration curve preparation of guanylurea nitrogen is shown below.

![HPLC Chromatogram](image)

**Reference diagram**  
HPLC chromatogram of the mixture standard solutions (10 mg/L for each)

**Peak name**
- (1) Urea nitrogen
- (2) Biuret nitrogen
- (3) Dicyandiamide nitrogen
- (4) Guanidine nitrogen
- (5) Guanylurea nitrogen

**Measurement conditions for HPLC**
- **Column:** Asahipak ES-502C 7C (7.5-mm inner diameter, 100-mm long, 9-μm particle diameter)
- Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
8.6 Uric acid
8.6.a High-Performance Liquid Chromatography

(1) Summary
This testing method is applicable to fertilizers. This testing method is classified as Type D and its symbol is 8.6.a-2018 or U-acid.a-2018.

Add phosphate solution (pH 8) to an analytical sample to extract uric acid, introduce it to a High-Performance Chromatograph (HPLC) to isolate it with a multi mode ODS (reverse phase+strong anion-exchange+strong cation-exchange+normal phase) column, and then measure at wavelength 290 nm to obtain uric acid (U-acid) in an analytical sample. The performance of this testing method is shown in Comment 5.

(2) Reagent, etc.: Reagents and water are as shown below.

a) Water: Water of A3 specified in JIS K 0557. Note that water of A4 should be used as the eluent which is introduced to an HPLC.

b) Potassium dihydrogenphosphate: A reagent specified in JIS K 9007 or a reagent of equivalent quality.

c) Disodium hydrogenphosphate: A reagent specified in JIS K 9020 or a reagent of equivalent quality.

d) Phosphate solution: Dissolve 9.073 g of potassium dihydrogenphosphate in water to make 1000 mL and dissolve 9.464 g of disodium hydrogenphosphate in water to make 1000 mL. Mix these solutions so that pH 8.0 ± 0.1 is reached.

e) Lithium carbonate solution: Dissolve 0.739 g of lithium carbonate (Li₂CO₃) of no less than 99 % (mass fraction) in purity in water to make 1000 mL.

f) Uric acid standard solution (U-acid 1 mg/mL): Put 0.100 g of uric acid in a weighing dish and measure the mass to the order of 0.1 mg. Dissolve it with a small amount of lithium carbonate solution, transfer to a 100 mL volumetric flask and add the solution up to the marked line.

g) Uric acid standard solution for the calibration curve preparation (U-acid 100 µg/mL): Put 10 mL of uric acid standard solution (U-acid 1 mg/mL) to a 100-mL volumetric flask and add phosphate solution up to the marked line.

h) Uric acid standard solution for the calibration curve preparation (U-acid 10 µg/mL - 50 µg/mL): Put 10 mL - 50 mL of uric acid standard solution (U-acid 100 µg/mL) to a 100-mL volumetric flask and add phosphate solution up to the marked line.

i) Uric acid standard solution for the calibration curve preparation (U-acid 0.1 µg/mL - 5 µg/mL): At the time of usage, put 1 mL - 50 mL of uric acid standard solution (U-acid 10 µg/mL) to 100-mL volumetric flasks step-by-step and add phosphate solution up to the marked line.

j) Ammonium acetate: A reagent specified in JIS K 8359 or a reagent of equivalent quality.

k) Methanol: A reagent of HPLC grade or a reagent of equivalent quality.

Note
(1) This is an example of preparation; prepare an amount as appropriate.

(3) Apparatus and instruments: Apparatus and instruments are shown below.

a) High-Performance Liquid Chromatograph (HPLC): HPLC specified in JIS K 0124 that satisfies following requirements.

1) Column: A 4.6-mm inner diameter 250-mm long stainless steel column tube filled with silica gel, to which octadecyl, ion-exchange group for strong acidity and strong basic anion-exchange group, chemically bond.

2) Column bath: A column bath whose temperature can be adjusted to 30 ºC - 45 ºC.

3) Detection unit: An absorptiometric detector that can measure at wavelength around 290 nm.
b) **Water bath:** Water bath that can be adjusted to 60 °C ± 2 °C.

c) **Magnetic stirrer**

d) **Centrifugal separator:** A centrifugal separator that can work at 1700 × g.

e) **High speed centrifugal separator:** A centrifugal separator that can work at 8000 × g - 10000 × g.

Comment 1 A column is sold under the production name Scherzo SS-C18.

(4) **Test procedures**

(4.1) **Extraction:** Conduct extraction as shown below.

a) Weigh 1.00 g of an analytical sample, and put it into a 200 mL-ground-in stopper Erlenmeyer flask.

b) Add 100 mL of phosphate solution (2) and heat for 30 minutes while shaking to mix (3) at every 10 minutes in a water bath at 60 °C ± 2 °C.

c) Immediately stir it by using a magnetic stirrer for about 10 minutes.

d) After allowing to stand still, transfer supernatant to a 15-mL or 50 mL-ground-in stopper centrifugal precipitate tube (4), and centrifuge at 1700 × g for about 5 minutes (5).

e) Transfer supernatant (6) to a 1.5-mL ground-in stopper centrifugal precipitate tube (4), and centrifuge at 8000 × g - 10000 × g for about 5 minutes (7) to make supernatant as a sample solution.

**Note**

(2) Use a silicone stopper instead of a glass stopper as the solution is heated.

(3) Steam easily expels silicon stoppers, so while lightly holding down the stopper from the top with your finger, shake it so that the water drops on the inside of the flask come down as much as possible. In addition, conduct this procedure before and after the heating procedure.

(4) The ground-in stopper centrifugal precipitate tube should be made of polypropylene, etc. to not affect the measurement

(5) 7.2 cm - 18.9-cm of rotor radius and 3000 rpm of revolutions makes about 1700 × g centrifugal force.

(6) If there is a possibility that the uric acid (U-acid) concentration in the sample solution will exceed the maximum limit of the calibration curve, dilute a predetermined amount of supernatant with phosphate solution.

(7) 7.2-cm - 8.9-cm of rotor radius and 10000 rpm of revolutions makes about 8100 × g - 10000 × g centrifugal force.

Comment 2 Instead of the procedures in (4.1) e), it is allowed to filter with a membrane filter (pore size: no more than 0.5-µm) made of hydrophilic PTFE and the filtrate can be the sample solution.

(4.2) **Measurement:** Conduct the measurement as indicated in JIS K 0124 and as shown below. Specific measurement procedures are according to the operation method of a High-Performance Liquid Chromatograph (HPLC) used in measurement.

a) **Measurement conditions for a High-Performance Liquid Chromatograph (HPLC):** An example of measurement conditions is shown below. Set up the measurement conditions considering it:

1) Column: A silica gel column, to which octadecyl, ion-exchange group for strong acidity and strong basic anion-exchange group chemically bond (4.6-mm inner diameter 250-mm long 3-µm particle diameter )

2) Column bath temperature: 40 °C
3) **Eluent** (1): Dissolve 1.54 g of ammonium acetate with water to make 1000 mL. Transfer 900 mL of the solution to mix with 100 mL of methanol. Filter with a membrane filter (aperture diameter: no more than 0.5-μm) made of hydrophilic PTFE.

4) **Flow rate**: 0.4 mL/min

5) **Injection volume**: 10 μL

6) **Detection unit**: An absorptiometric detector, measurement wavelength: 290 nm

**Comment 3** Eluent can be prepared as follows. Dissolve 15.4 g of ammonium acetate with water to make 1000 mL and store in a refrigerator. At the time of usage, dilute a predetermined volume of the solution by a factor of 10 to mix with methanol of volume ratio 1/9 and filter with a membrane filter (aperture diameter: no more than 0.5-μm) made of hydrophilic PTFE.

b) **Calibration curve preparation**

1) Inject 10 μL of respective standard solutions for calibration curve preparation into an HPLC, record a chromatogram at wavelength 290 nm and obtain the peak area or height.

2) Prepare a curve for the relationship between the uric acid (U-acid) concentration and the peak area or height at wavelength 290 nm of respective standard solutions for the calibration curve preparation.

c) **Sample measurement**

1) Subject 10 μL of sample solution to the same procedure as in b) 1)

2) Obtain the uric acid (U-A) content from the peak area or height using the calibration curve to calculate the uric acid (U-A) in the analytical sample.

**Comment 4** In addition to uric acid, allantoin and allantoic acid can be measured simultaneously by this measurement method (when using a Scherzo SS-C18 column). Note that the detection wavelength of allantoin and allantoic acid is 210 nm.

**Comment 5** Additive recovery testing was conducted using one brand of a compound fertilizer, a composted sludge fertilizer, a mixed compost fertilizer and compost. As a result, the mean recovery at additive level of 0.1 % (mass fraction), 0.01 % (mass fraction) and 0.005 % (mass fraction) were 92.4 % - 101.8 %, 85.3 % - 105.0 % and 92.5 % - 114.1 % respectively.

The results of the repeatability tests on different days using one brand of a compound fertilizer, a composted sludge fertilizer, a mixed compost fertilizer and compost to evaluate precision were analyzed by one-way analysis of variance. Table 1 shows the calculation results of intermediate precision and repeatability.

Additionally, the minimum limit of quantification of this testing method is about 0.0008 % (mass fraction).
Table 1  Analysis results of the repeatability tests on different days

<table>
<thead>
<tr>
<th>Name of sample</th>
<th>Test days of repeatability</th>
<th>Average $^2$</th>
<th>Repeatability $^5$</th>
<th>Intermediate precision $^7$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T$</td>
<td>$s_r$ $(%)$</td>
<td>$RSD_s$ $(%)$</td>
<td>$s_{I(T)}$ $(%)$</td>
</tr>
<tr>
<td>Compound fertilizer</td>
<td>7</td>
<td>0.0989</td>
<td>0.0006</td>
<td>0.6</td>
</tr>
<tr>
<td>Composted sludge fertilizer</td>
<td>7</td>
<td>0.0102</td>
<td>0.0001</td>
<td>0.7</td>
</tr>
<tr>
<td>Mixed compost fertilizer</td>
<td>7</td>
<td>0.09938</td>
<td>0.0004</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>0.00921</td>
<td>0.00005</td>
<td>0.9</td>
</tr>
<tr>
<td>Compost</td>
<td>7</td>
<td>0.1010</td>
<td>0.001</td>
<td>1.3</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>0.00966</td>
<td>0.00018</td>
<td>1.8</td>
</tr>
</tbody>
</table>

1) The number of test days conducting a duplicate test
2) Average (the number of test days $(T)$)
3) Mass fraction
4) Repeatability standard deviation
5) Repeatability relative standard deviation
6) Intermediate standard deviation
7) Intermediate relative standard deviation

(5) Flow sheet for testing method: The flow sheet for urea acid in fertilizers is shown below:

```
1.00 g analytical sample (powdery)
← 100 mL of phosphate solution
  Heating
  Extraction
  Centrifugal separation
  Centrifugal separation
  Sample solution
  Measurement

200-mL ground-in stopper Erlenmeyer flask
Heat at 60 ºC ± 2 ºC for 30 minutes while shaking to mix at every 10 minutes
Stir to mix, 10 minutes
Ground-in stopper centrifugal precipitate tube, 1700 × g, 10 minutes
Ground-in stopper centrifugal precipitate tube, 8000 × g - 10000 × g, 5 minutes
Supernatant
Liquid Chromatograph
```

Figure Flow sheet for urea acid in fertilizers
Reference: Chromatogram of the standard solution for calibration curve preparation of uric acid is shown below.

Reference diagram HPLC chromatogram of the uric acid standard solution (50 µg/mL) for calibration curve preparation

Peak name (↓) Uric acid
Measurement conditions for HPLC
Column: Scherzo SS-C18 (4.6-mm inner diameter, 250-mm long, 3-µm particle diameter)
Other conditions are according to the example of HPLC measurement conditions in (4.2) a)
Appendix. The procedure to validate characteristics of testing methods

(1) Purposes
This article explains the procedure to validate characteristics of testing methods which will be listed in the Testing Methods of Fertilizers. In addition, when testing institutes conduct a test which is not included in the Testing Methods of Fertilizers, a procedure to evaluate the validity of the test method should conform to a method stipulated in this article. Additionally, this article targets chemical testing methods. However, this article is not applicable to the extraction method of the content of effective figures (acid-, alkaline-, citrate- and water-soluble) in a powdery sample or a solidified fertilizer.

Comment 1
The contents of effective figures (acid-, alkaline-, citrate- and water-soluble) are stipulated in a notification of the Ministry of Agriculture, Forestry and Fisheries. In addition, the change of measurement conditions such as an extraction temperature may affect an observed value in some cases. Therefore, no changes will be implemented in the extraction method of the contents of effective figures in a powdery fertilizer and a solidified fertilizer for the present and the application of this article is limited to the change of a measurement method (including refining of extract, etc.).

(2) Definition of terminology
The definition of terminology in this article is as shown below.

a) Selectivity: Capability to accurately measure components subjected to analysis under the existence of materials which seem to exist in a sample.

b) Trueness: The degree of agreement between the mean obtained from multiple measurement results and the true value (1).

c) Precision: The degree of agreement among the independent measurement results which are repeatedly measured under the determined conditions.

d) Repeatability: The precision of the measurement results of analytical samples, which are regarded to be all identical, obtained under condition (repeatability conditions) that independent measurement results are measured in a short time, using the same method, in the same laboratory, by the same operator and with the same instrument.

e) Intermediate precision: The precision of a measurement result of analytical samples, which are regarded to be all identical, obtained under condition (intermediate conditions) that independent measurement results are measured, using the same method, in the same laboratory and in different factors (such as different time and a different operator).

f) Reproducibility: The precision of a measurement result of analytical samples, which are regarded to be all identical, obtained under condition (reproducibility conditions) that independent measurement results are measured, using the same method, in different laboratories, by different operators and with different instruments.

g) Minimum Limit of Quantification (LOQ): The quantifiable lowest volume or minimum concentration of a component subjected to analysis which is contained in an analytical sample.

h) Minimum Limit of Detection (LOD): The detectable lowest volume or minimum concentration of a component subjected to analysis which is contained in an analytical sample.

i) Reference material: A material which is uniform and stable enough for one or more prescribed properties, and is made suitable for the purpose of use in a measurement process.

j) Certified reference material: A reference material, whose values of one or more prescribed properties are characterized by a reasonable metrological procedure, having a certificate of
attestation on which the characteristics of prescribed properties and their uncertainty and metrological traceability are stated.

k) **Blank sample**: An analytical sample not containing components subjected to analysis \(^{(2)}\).

l) **Addition sample**: An analytical sample the content of whose components subjected to analysis is known, or an analytical sample to which reference materials are added \(^{(3)}\) \(^{(4)}\) or compounded \(^{(3)}\).

m) **Natural contamination sample**: A test sample prepared from fertilizers which naturally contain the components subjected to analysis such as harmful components.

n) **Distribution sample**: An analytical sample prepared from fertilizers \(^{(5)}\) which are manufactured in a fertilizer production factory, etc.

o) **Surrogate**: A material which is added to an analytical sample in order to conduct a pre-process operation, correct yields in respective steps of measurement procedure and confirm recovery, whose chemical structure is identical or similar to a target component.

p) **SN ratio**: Intensity ratio of a signal (response value) \(S\) originating from the analysis target and a signal (usually noise) \(N\) based on the other factors.

**Note**

(1) In reality, the certified value of a certified reference material, the chemical composition of a compound, the added content of a reference material, etc. and others.

(2) Reagents, etc. containing a target matrix can be used in the case that there is no distribution fertilizer used as a blank sample for a recovery test and the confirmation of the minimum limit of quantification, etc.

(3) Mix a component subjected to analysis with a mortar, etc. to sufficient uniformity

(4) In the case of adding a standard solution, vaporize the solvent sufficiently conducting measures such as letting it stand for one night.

(5) A fertilizer containing components subjected to analysis whose formation or form changed due to a chemical or physical process (a granulation process, etc.).

**References**

1) **JIS K 0211**: Technical terms for analytical chemistry (General part) (2013)

2) **JIS K 0214**: Technical terms for analytical chemistry (Chromatography part) (2013)

3) **JIS Q 0035**: Reference materials—General and statistical principles for certification (2008)


5) **JIS Z 8402-1**: Accuracy (trueness and precision) of measurement methods and results - Part 1: General principles and definitions (1999)


7) **ICH Harmonised Tripartite Guideline, Validation of Analytical Procedures: Text and Methodology Q2(R1)**, International Conference on Harmonisation of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH) (2005)

(3) **Validation method**

Test necessary items of (3.1) to (3.8) in a planned manner and estimate performance parameters from the obtained results.

Confirm whether the estimated values of performance parameters are suitable to target values (performance norm) respectively, and evaluate that the test method is validated if they are all suitable.
(3.1) Scope of application

As a result of a validation test in a single laboratory and a collaborative study, if the result is suitable up to reproducibility, the test method is evaluated as a validated test method as far as the kind of a fertilizer used in the test and the range of concentration are concerned. Therefore, a laboratory where the said test is conducted can use the performance (reproducibility, etc.) as a validated method through implementing internal quality control, etc.

As a result of a validation test in a single laboratory, if the result is suitable to trueness, repeatability and intermediate precision, etc., the test method is evaluated as a validated test method as far as the laboratory where the test was conducted and as far as the kind of a fertilizer used in the test and the range of concentration are concerned. Therefore, another laboratory which wants to introduce the test method is required to carry out the validation anew in an individual laboratory with the above test method.

(3.2) Selectivity

(3.2.1) Case of Chromatography

Conduct a procedure for a blank sample and confirm that there is no peak (interference peak) which affects the measurement of components subjected to analysis. In addition, in the case of the simultaneous measurement of multi components, confirm that adjacent peaks are sufficiently separated.

Note (6) Resolution \((R)\) should be 1.0 or more at minimum though 1.5 or more is preferable.

Comment 2 Resolution \((R)\) is used as a separation indicator of peaks. If Resolution \((R)\) is 1.5 or more, the adjacent two peaks are sufficiently separated and they do not affect a measurement, whether a peak height or a peak area is used. If Resolution \((R)\) is 1.0 or more, the adjacent two peaks are sufficiently separated and they do not affect a measurement, whether a peak height or a peak area is used.

Resolution \((R)\) can be obtained using a peak width by the formula (1a). In addition, if the peak is a normal distribution, it can be obtained using a peak width at half height by the formula (1b). With the data processing device of a chromatograph, the formula (1b) is often used to obtain Resolution \((R)\).

\[
\text{Resolution } (R) = \frac{t_2 - t_1}{\frac{1}{2} \times (W_1 + W_2)} \quad \cdots (1a)
\]

\[
\text{Resolution } (R) = \frac{1.18 \times (t_2 - t_1)}{(W_{1/2}^1 + W_{1/2}^2)} \quad \cdots (1b)
\]

\(t_1\): Retention time of Peak 1 \\
\(t_2\): Retention time of Peak 2 \\
\(W_1\): Peak width of Peak 1 \\
\(W_2\): Peak width of Peak 1 \\
\(W_{1/2}^1\): Peak width at half height of Peak 1 \\
\(W_{1/2}^2\): Peak width at half height of Peak 2
(3.2.2) Case of a method other than Chromatography

Conduct a procedure for a blank sample and confirm that there is no response which originates from other components than a component subjected to analysis and can be a factor of positive error of a quantification value.

Note (7) A test method such as Molecular absorption spectrometry, Atomic absorption spectrometry or Titration analysis which does not isolate with a measurement instrument.

(8) Absorbance, titer, etc.

References

1) AOAC Official Methods of Analysis Appendix K: Guidelines for Dietary Supplements and Botanicals, AOAC INTERNATIONAL (2012)
2) JIS K 0114: General rules for gas chromatography (2012)
3) JIS K 0124: General rules for high performance liquid chromatography (2011)

(3.3) Calibration curve

Measure respective standard solutions for the calibration curve preparation of the concentration or the content of level 6 to 8 a few times to make a figure plotting the obtained signals as a function of the concentration or the content subjected to analysis and evaluate its linearity visually using the figure.

If linearity is recognized, calculate the inclination ($b$) and the intercept ($a$) of a calibration curve, its confidence interval and the coefficient of determination ($r^2$) using a statistical method such as the calculation of a regression equation by the least square method. Moreover make the plot of residuals in respective levels.

Note (9) The blank test solution for the calibration curve preparation can be included.

(10) In order to avoid nonlinear confusion due to the variation of sensitivity, etc., conduct measurements randomly for each replicate determination.

(11) Absorbance, fluorescence intensity, peak height, peak area, etc.

(12) The difference between a signal obtained by measurement and a signal estimated using a regression equation.

Comment 3 It is recommended that the 95% confidence interval of an intercept ($a$) includes the origin (0).

Comment 4 Though it is usable if the coefficient of determination ($r^2$) is 0.99 or more, it is recommended that the coefficient of determination ($r^2$) is 0.999 or more for a precise analysis. If it is less than 0.99, use the equation of a higher order or study the conversion of a numerical value.

Comment 5 The mean of residuals is 0 and the residuals indicate a random pattern.

References

1) AOAC Official Methods of Analysis Appendix K, Guidelines for Dietary Supplements and Botanicals, AOAC INTERNATIONAL (2012)
(3.4) **Trueness**

As the estimation method of trueness, the methods are recommended in the following order: (1) Use of a certified reference material (3.4.1), (2) Comparison with an observed value by a validated method (3.4.2) and (3) Recovery test (3.4.3).

In addition, if a surrogate is used, it is recommended that a recovery is about 40% or more.

(3.4.1) **Use of a certified reference material**

With regard to a component which has matrixes similar to a fertilizer subjected to test and can use a certified reference material containing components subjected to measurement of the concentration in a measurement level, conduct repeatability testing using 3 or more analytical samples \((n)\) according to the test method of the certified reference material. As a result, the mean of the observed values should be within the warning level to the certified value (characteristic value) or the absolute value of the difference between the mean of the observed values and the certified value (characteristic value) should not exceed 2 times of the standard uncertainty composed of respective uncertainties of the mean of the observed values and the certified value \((13)\).

**Comment 6** A warning limit is given using the formula (2) which is obtained from a collaborative study for the characterization of a certified reference material.

\[
\text{A warning limit for a certified value } (\mu) = \mu \pm 2 \times \sqrt{\left(s_R^2 - s_r^2\right) + \frac{s_r^2}{n}} = \mu \pm 2 \times \sqrt{s_L^2 + \frac{s_r^2}{n}} \quad \cdots (2)
\]

- \(\mu\): Certified value
- \(s_R\): Reproducibility standard deviation in a collaborative study
- \(s_r\): Repeatability standard deviation in a collaborative study \((14)\)
- \(n\): The number of analytical samples to repeatability test
- \(s_L\): Pure between-laboratory standard deviation in a collaborative study

**Note** (13) The evaluation procedure of the difference between a measurement result and a certified value (characteristic value) is shown in Reference 1 Procedure to compare an observed value and a certified value.

(14) It may be expressed as within-laboratory standard deviation \((s_{W})\) in some cases.

(3.4.2) **Use of another validated test method**

For a component for which a certified reference material is not usable but another validated test method (hereinafter referred to as “a standard test method”) is applicable, confirm that the condition a) or b) is satisfied.

a) **In case 12 or more samples are available:** Conduct respective tests of 12 or more test samples composed of addition samples, natural contamination samples or distribution samples according to a new test method and a standard test method, create the correlation chart of
observed values with two methods for each sample and calculate the inclination \((b)\) and the intercept \((a)\) of a regression line, and a correlation coefficient \((r)\). Further confirm a prediction interval.

However, in case that the width between the minimum and the maximum observed value is small, conduct the paired samples \(t\)-test to confirm that a significant difference is not observed.

**Comment 7** It is recommended that the 95% confidence interval of an inclination \((b)\) includes 1, the 95% confidence interval of an intercept \((a)\) includes the origin (0) and the correlation coefficient \((r)\) is no less than 0.99.

b) **In case fewer samples are available:** With regard to 3 or more test samples of different concentration, conduct respective repeatability addition tests using 4 analytical samples according to the new test method and a standard test method, confirm the homoscedasticity of the results of 2 groups and conduct a \(t\)-test for each concentration to confirm that significant difference is not observed under the two-sided significant level of 5%.

(3.4.3) **In case neither certified reference material nor other validated test methods are usable**

For 3 or more test samples of different concentration, conduct respective repeatability tests using 3 analytical samples and evaluate by obtaining the recovery using the mean of the observed values.

The criteria of the trueness are shown in **Separate sheet: The target of trueness and the criteria of precision in respective concentration levels**.

**References**

1) AOAC Official Methods of Analysis Appendix K: Guidelines for Dietary Supplements and Botanicals, AOAC INTERNATIONAL (2012)


3) Linsinger, T.; Comparison of a measurement result with the certified value, European Reference Materials' application note 1, European Commission - Joint Research Centre Institute for Reference Materials and Measurements (IRMM) (2010)


(3.5) **Precision**

Evaluate reproducibility and repeatability by a collaborative study (3.5.1). Or evaluate an intermediate precision and repeatability by a repeatability test (3.5.2).

(3.5.1) **Reproducibility and repeatability by a collaborative study**

The number of laboratories to obtain effective data should be 8 or more \(^{(15)}\). Conduct undisclosed duplicate collaborative studies for 5 or more kinds of samples with different concentration. Obtain reproducibility and repeatability from the observed values \(^{(16)}\) to evaluate.

Criteria to evaluate these precisions are shown in **Separate sheet: The target of trueness and the criteria of precision in respective concentration levels**.

**Note** (15) In case the number of laboratories which have required facility/instruments is limited,
this should be 5 or more.

(16) The calculation method is shown in Reference 2: Calculation of reproducibility or intermediate precision and repeatability

(3.5.2) Intermediate precision and repeatability by a repeatability test in a single laboratory on different days

Conduct a duplicate test\(^{(17)}\) per test day for 5 to 7 days using two analytical samples of different concentration which is included in a normal range \(^{(18)}\). Obtain intermediate precision and repeatability from the observed values \(^{(19)}\) to evaluate.

Criteria to evaluate these precisions are shown in Separate sheet: The target of trueness and the criteria of precision in respective concentration levels.

Note \(^{(17)}\) The data of internal quality control can be used. 
\(^{(18)}\)It is not necessary for the same tester to conduct a test through 5 to 7 days. 
\(^{(19)}\)The calculation method is shown in Reference 2: Calculation of reproducibility or intermediate precision and repeatability

References

1) AOAC Official Methods of Analysis Appendix K: Guidelines for Dietary Supplements and Botanicals, AOAC INTERNATIONAL (2012)
3) AOAC Official Methods of Analysis Appendix D: Guidelines for Collaborative Study Procedures To Validate Characteristics of a Method of Analysis, AOAC INTERNATIONAL (2005)

(3.6) Minimum Limit of Quantification (LOQ)

Estimate Minimum Limit of Quantification according to (3.6.1) to (3.6.3). Prepare test samples which include the concentration estimated to be near the Minimum Limit of Quantification step by step as necessary. And conduct respective repeatability tests using 3 analytical samples and define the concentration of a prepared test sample as the Minimum Limit of Quantification, where the mean of the obtained values using the prepared test sample is suitable to the target value of trueness.

Comment 8 In case permissible content and equivalent level is 1.0 mg/kg or more, the Minimum Limit of Quantification (LOQ) of harmful components and restricted components, etc. should be no more than 1/5 of the permissible content and equivalent level. In case permissible content and equivalent level is no more than 1.0 mg/kg, the Minimum Limit of Quantification (LOQ) should be no more than 2/5 of the permissible content. Moreover, it is recommended that the Minimum Limit of Quantification of main components/major components and material components should be no more than 1/5 of minimum volume to be contained and the minimum content of a distribution fertilizer. In addition, in case the Minimum Limit of Quantification exceeds 1/5 of these minimum volumes, conduct the above-mentioned repeatability test, confirm the Minimum Limit of Quantification and state clearly the fact in the applicable range of a test method.

Comment 9 There are some methods to estimate Minimum Limit of Quantification. The methods differ depending on whether they are based on an instrument analysis or not and
depending on instruments used. A method different from the methods shown in (3.6.1) to (3.6.3) is allowed. However, the definition of a method and Minimum Limit of Quantification by the method should be clearly stated.

(3.6.1) Estimation method by a repeatability test

With regard to a test sample with concentration near Minimum Limit of Quantification, conduct a repeatability test using 7 to 10 analytical samples, obtain repeatability standard deviation and estimate Minimum Limit of Quantification (LOQ) in an analytical sample by the formula (3).

\[
\text{Estimated value of Minimum Limit of Quantification (LOQ)} = 10 \times s_r \quad \cdots (3)
\]

\(s_r\): Repeatability standard deviation

(3.6.2) Estimation method using a calibration curve

In case a calibration curve is linear, estimate Minimum Limit of Quantification (LOQ) in an analytical sample by the formula (4) using the standard deviation of the residuals of a calibration curve or estimated signals in concentration 0 and the inclination of a calibration curve.

\[
\text{Estimated value of Minimum Limit of Quantification (LOQ)} = \frac{10 \times s}{b} \quad \cdots (4)
\]

\(s\): The standard deviation of residuals. Or the standard deviation of signals in concentration 0, which are estimated from a regression line

\(b\): The inclination of a calibration curve

(3.6.3) Estimation method using an SN ratio

In a test method such as Chromatography, etc. which has a baseline noise, calculate from a concentration in an analytical solution whose SN ratio is 10 to 1 at the peak and estimate Minimum Limit of Quantification (LOQ) in an analytical sample.

References


(3.7) Minimum Limit of Detection (LOD)

Estimate Minimum Limit of Detection according to (3.7.1) to (3.7.3).

Comment 10 There are some methods to estimate Minimum Limit of Detection. The methods differ depending on whether they are based on an instrument analysis or not and depending on instruments used. A method different from the methods shown in (3.7.1) to (3.7.3) is allowed. However, the definition of a method and Minimum
Limit of Detection by the method should be clearly stated.

(3.7.1) Estimation method by a repeatability test
With regard to a test sample or a blank sample with concentration near Minimum Limit of Quantification, conduct repeatability tests using 7 to 10 analytical samples, obtain repeatability standard deviation and estimate Minimum Limit of Detection (LOD) in an analytical sample by the formula (5).

Estimated value of Minimum Limit of Detection (LOD) in an analytical sample

\[ 2 \times t(n - 1, 0.05) \times s_r \quad \cdots (5) \]

\( s_r \): Repeatability standard deviation
\( t(n - 1, 0.05) \): The Student value of Significance Level 5% (one side) (20)
\( n \): The number of analytical samples in a repeatability test

Note (20) In case of a repeatability test using 7 analytical samples, the value is 1.94. In case of using 10 analytical samples, the value is 1.83.

(3.7.2) Estimation method using a calibration curve
In case a calibration curve is linear, estimate Minimum Limit of Detection (LOD) in an analytical sample by the formula (6) using the standard deviation of the residuals of a calibration curve or estimated signals in concentration 0 and the inclination (b) of a calibration curve.

Estimated value of Minimum Limit of Detection (LOD)

\[ \frac{2 \times t(n - 2, 0.05) \times s}{b} \quad \cdots (6) \]

\( s \): The standard deviation of residuals. Or the standard deviation of signals in concentration 0, which are estimated from a regression line
\( b \): The inclination of a calibration curve
\( t(n - 2, 0.05) \): The Student value of Significance Level 5% (one side)
\( n \): The number of a measurement point on a calibration curve

(3.7.3) Estimation method using an SN ratio
In a test method such as Chromatography, etc. which has a baseline noise, calculate from a concentration in an analytical solution whose SN ratio is 3 to 1 at the peak and estimate Minimum Limit of Detection (LOD) in an analytical sample.

References
2) The notification by the Director of Evaluation and Licensing Division, Pharmaceutical and Food Safety Bureau, the Ministry of Health, Labour and Welfare: The text (operation method) on Bioanalytical Method Validation”, October 28, 1997, Iyaku-Shin
(3.8) **Robustness**

Robustness should be studied when an analysis method is developed, and the estimation method depends on the type of analysis method to be developed. Robustness expresses the reliability of an analysis method when its analysis conditions are intentionally changed. If an observed value tends to be easily affected by the variation of an analysis condition, it is necessary to consider a method to control an analysis condition appropriately or to state the fact as a precaution in a testing method. The evaluation of robustness enables the establishment of a series of parameters such as Resolution related to system conformance. Similarly, the confirmation of these parameters ensures that the validation of an analysis method is maintained in a daily analysis.

Typical variation factors are as follows.

(3.8.1) **Common variation factors**: Typical variation factors common to various kinds of test methods are as follows.

a) Extraction time, extraction temperature
b) Stability of a test solution in respective steps
c) Reagent’s grade

(3.8.2) **Variation factors in Chromatography, etc.**: Typical variation factors of measurements by Chromatography or refining by solid phase extraction are as follows.

a) Change of a column or a cartridge (A different lot or a different brand)
b) Influence by the variation of pH and composition of an eluent or a wash
c) Temperature
d) Flow rate
e) Influence of a matrix and effect of dilution

**References**


Reference 1: Procedure to compare an observed value and a certified value

Obtain the total mean (m) of the replication test results and the certified value (μ), and the absolute value (Δ_m) of the difference of the two values by the formula (R1.1). Next, obtain the standard uncertainty (u_{CRM}) of the certified value of a certified reference material by the formula (R1.2), and obtain the standard uncertainty (u_m) of the total mean by the formula (R1.3). Calculate the combined standard uncertainty (u_{C(Δ_m)}) of Δ_m by the formula (R1.4) using the obtained u_m and u_{CRM}. Further, calculate an expanded uncertainty (U_{Δ_m}) by the formula (R1.5) using the coverage factor (k = 2).

Compare Δ_m and U_{Δ_m} to confirm that the criterion (the formula (R1.6)) is satisfied, that is, Δ_m is no more than U_{Δ_m}.

The absolute value (Δ_m) of the difference of the total mean of repeatability test results and a certified value = |m − μ| \cdots (R1.1)

The standard uncertainty (u_{CRM}) of the certified value = \frac{U_{95\%}}{k_{CRM}} \cdots (R1.2)

The standard uncertainty of the measurement of a total mean (u_m) = \frac{s_r}{\sqrt{n}} \cdots (R1.3)

The combined standard uncertainty (u_{C(Δ_m)}) of Δ_m = \sqrt{u_m^2 + u_{CRM}^2} \cdots (R1.4)

The expanded uncertainty (U_{Δ_m}) of Δ_m = k_{C(Δ_m)} \times u_{C(Δ_m)} = 2 \times u_{C(Δ_m)} \cdots (R1.5)

Criterion Δ_m \leq U_{Δ_m} \cdots (R1.6)

m: The total mean of observed values
μ: A certified value
U_{95\%}: The expanded uncertainty of a certified value
k_{CRM}: The coverage factor of an expanded uncertainty of a standard reference material
s_r: Repeatability standard deviation
n: The number of repeatability test samples
k_{C(Δ_m)}: The coverage factor of an expanded uncertainty of Δ_m (k_{C(Δ_m)} = 2)
Reference 2: Calculation of reproducibility or intermediate precision and repeatability

(1) Structure of an observed value

An observed value \((x_{ij})\) in Table 1, as is shown in the formula (R2.1), consists of a true value \((\mu)\), a variation \((\beta)\) by a factor and a variation \((e)\) by an accidental error under repeatability conditions (hereinafter referred to as “an accidental error”). When \(p\) laboratories conduct a collaborative study in which respective laboratories conduct repeatability tests using \(n\) samples, the formula (R2.2) is introduced on the assumption that the distribution of \(\beta\) is equivalent to \(N(0, \sigma_L^2)\) which depends on a pure between-laboratory variation and the distribution of \(e\) is equivalent to \(N(0, \sigma_r^2)\) which depends on an accidental error. In addition, when the same laboratory conducts replicate tests for \(p\) days using \(n\) samples on respective test days, the formula (R2.3) is introduced on the assumption that the distribution of \(\beta\) is equivalent to \(N(0, \sigma(I)^2)\) which depends on test days variation (factor \(T\)) and the distribution of \(e\) is equivalent to \((0, \sigma_g^2)\) which depends on an accidental error.

\[
\text{Observed value } (x_{ij}) = \mu + \beta_i + e_{ij} \quad \cdots \quad (R2.1)
\]

\[
\text{Observed value } (x_{ij}) = \mu + N(0, \sigma_L^2) + N(0, \sigma_r^2) \quad \cdots \quad (R2.2)
\]

\[
\text{Observed value } (x_{ij}) = \mu + N(0, \sigma(T)^2) + N(0, \sigma_r^2) \quad \cdots \quad (R2.3)
\]

\(\mu\): True value

\(\beta_i\): Variation of a factor

\(e_{ij}\): Accidental error

\(N(0, \sigma_L^2)\): Normal distribution of \(\beta_i\) with the mean 0 and standard deviation \(\sigma_L\)

\(N(0, \sigma_r^2)\): Normal distribution of \(e_{ij}\) with the mean 0 and standard deviation \(\sigma_r\)

\(\sigma_L^2\): Pure between-laboratory variance

\(\sigma_r^2\): Repeatability variance

\(N(0, \sigma(T)^2)\): Normal distribution of \(\beta_i\) with the mean 0 and standard deviation \(\sigma(T)\)

\(\sigma(T)^2\): Test days variance
(2) Calculation procedure of reproducibility and repeatability of the results of a collaborative study

(2.1) Estimation of true value and variance

In an actual statistical analysis, a true value ($\mu$), a true and pure between-laboratory variance ($\sigma_L^2$) and a true repeatability variance ($\sigma_r^2$) are unknown. Therefore, they are replaced with estimated values obtained from the results of a collaborative study and are expressed as a mean ($m$), a pure between-laboratory variance ($\sigma_L^2$) and a repeatability variance ($\sigma_r^2$) respectively.

(2.2) One-way analysis of variance

Exclude ineffective observed values which have clearly objective reasons such as deviation from a protocol and malfunction of instruments from the report values by laboratories which participated in a collaborative study. Further exclude outliers by conducting Cochran’s test and Grubb’s test. And conduct one-way analysis of variance for the remaining results to obtain the unbiased variance ($V$) of respective variation factors in Table 2.

Table 2 Table of one-way analysis of variance

<table>
<thead>
<tr>
<th>Variation factor</th>
<th>Sum of squares</th>
<th>Degree of freedom</th>
<th>Unbiased variance ($V$)</th>
<th>Expectation of variance ($E(V)$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Between-laboratory ($L$)</td>
<td>$SS_L$</td>
<td>$p - 1$</td>
<td>$V_L$</td>
<td>$\sigma_r^2 + \sigma_L^2$</td>
</tr>
<tr>
<td>Accidental error ($e$)</td>
<td>$SS_e$</td>
<td>$p \times (n - 1)$</td>
<td>$V_e$</td>
<td>$\sigma_e^2$</td>
</tr>
</tbody>
</table>

Comment 1 It is possible to conduct one-way analysis of variance easily using a statistical program or a tool of a spreadsheet program. In this case, it should be noted that different terminologies may be used (Between-laboratory ($L$) → Between-group, Accidental error ($e$) → Within-group, Unbiased variance → Mean square, etc.).

Comment 2 Unbiased variance ($V$) is calculated by (Sum of squares)/ (Degree of freedom).

(2.3) The calculation of reproducibility and repeatability

The relation of the expectation of variance $E(V)$ of respective factors in Table 2 holds true. Therefore, calculate repeatability variance ($s_r^2$) and pure between-laboratory variance ($s_L^2$) by the formula (R2.4) and (R2.5), and further calculate reproducibility variance ($s_R^2$) by the formula (R2.6) (1) (2).

Repeatability variance \( (s_r^2) = V_r \) \( \cdots \) (R2.4)

Pure between – laboratory variance \( (s_L^2) = \frac{V_L - V_r}{n} \) \( \cdots \) (R2.5)

Reproducibility variance \( (s_R^2) = s_L^2 + s_r^2 \) \( \cdots \) (R2.6)

\( V_r \): The unbiased variance of a variation factor (accidental error (e))
in the table of one – way analysis of variance (Table 2)

\( V_L \): The unbiased variance of a variation factor (between – laboratories (L))
in the table of one – way analysis of variance (Table 2)

Calculate a repeatability standard deviation \( (s_r) \) and a reproducibility standard deviation \( (s_R) \) by the formula (R2.7) and (R2.8) using the obtained repeatability variance and reproducibility variance, and further calculate a repeatability relative standard deviation \( (RSD_r) \) and a reproducibility relative standard deviation \( (RSD_R) \) by the formula (R2.9) and (R2.10) \( ^{(2)}(3) \).

Repeatability standard deviation \( (s_r) = \sqrt{s_r^2} \) \( \cdots \) (R2.7)

Reproducibility standard deviation \( (s_R) = \sqrt{s_R^2} \) \( \cdots \) (R2.8)

Repeatability relative standard deviation \( (RSD_r, \%) = \frac{s_r}{m} \times 100 \) \( \cdots \) (R2.9)

Reproducibility relative standard deviation \( (RSD_R, \%) = \frac{s_R}{m} \times 100 \) \( \cdots \) (R2.10)

\( m \): the total mean ( ) of the effective data of collaborative study results

Note
1. In case \( V_L < V_r \), assume \( V_L = V_r \) (that is, the pure between-laboratory variance \( (s_L^2) = 0 \) in the formula (R2.5)) and let the formula (R2.6) form \( s_R^2 = s_r^2 \).
2. The rounding of a numerical value is not executed in the middle of the calculation.
3. The mean and the standard deviation are expressed rounding to the digit of the observed value. The relative standard deviation is expressed rounding to the first decimal place.

(3) Calculation procedure of intermediate precision and repeatability by the replicate test results on different days

3.1 Estimation of a true value and a variance

In an actual statistical analysis, a true value \( (\mu) \), a true test day variance \( (\sigma_{(T)}^2) \) and a true repeatability variance \( (\sigma_r^2) \) are unknown. Therefore, they are replaced with estimated values obtained from the replicate test results on different days and are expressed as a mean (m), test days variance \( (s_{(T)}^2) \) and a repeatability variance \( (s_r^2) \) respectively.

3.2 One-way analysis of variance

Conduct one-way analysis of variance for the replicate test results on different days to obtain the
unbiased variance \( (V) \) of respective variation factors in Table 3.

<table>
<thead>
<tr>
<th>Variation factor</th>
<th>Sum of squares</th>
<th>Degree of freedom</th>
<th>Unbiased variance</th>
<th>Expectation of variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test days ((T))</td>
<td>(SS_T)</td>
<td>(p - 1)</td>
<td>(V_T)</td>
<td>(\sigma_T^2 + n\times \sigma_T(T)^2)</td>
</tr>
<tr>
<td>Accidental error ((e))</td>
<td>(SS_r)</td>
<td>(p \times (n - 1))</td>
<td>(V_r)</td>
<td>(\sigma_r^2)</td>
</tr>
</tbody>
</table>

**Comment 3** It is possible to conduct one-way analysis of variance easily using a statistical program or a tool of a spreadsheet program. In this case, it should be noted that different terminologies may be used (Test days \((T)\) → between-group, Accidental error \((e)\) → within-group, Unbiased variance → Mean square, etc.).

**Comment 4** Unbiased variance \((V)\) is calculated by \((\text{Sum of squares})/ (\text{Degree of freedom})\).

### (3.3) The calculation of intermediate precision and repeatability

The relation of the expectation of variance \(E(V)\) of respective factors in Table 3 holds true. Therefore, calculate repeatability variance \((s_r^2)\) and test days variance \((s_T(T)^2)\) by the formula (R2.11) and (R2.12), and further calculate intermediate variance \((s_{I(T)}^2)\) by the formula (R2.13)\(^2\)\(^4\).

\[
\text{Repeatability variance } (s_r^2) = V_r \quad \cdots \quad (R2.11)
\]

\[
\text{Test days variance } (s_T(T)^2) = \frac{V_T - V_r}{n} \quad \cdots \quad (R2.12)
\]

\[
\text{Intermediate Variance } (s_{I(T)}^2) = s_T(T)^2 + s_r^2 \quad \cdots \quad (R2.13)
\]

\(V_r\): The unbiased variance of a variation factor (accidental error \((e)\))
  in the table of one-way analysis of variance (Table 3)

\(V_T\): The unbiased variance of a variation factor (test days \((T)\))
  in the table of one-way analysis of variance (Table 3)

Calculate a repeatability standard deviation \((s_r)\) and an intermediate standard deviation \((s_{I(T)})\) by the formula (R2.14) and (R2.15) using the obtained estimated values of repeatability variance and intermediate variance, and further calculate a repeatability relative standard deviation \((RSD_r)\) and an intermediate relative standard deviation \((RSD_{I(T)})\) by the formula (R2.16) and (R2.17)\(^2\)\(^3\).

\[
\text{Repeatability standard deviation } (s_r) = \sqrt{s_r^2} \quad \cdots \quad (R2.14)
\]

\[
\text{Intermediate standard deviation } (s_{I(T)}) = \sqrt{s_{I(T)}^2} \quad \cdots \quad (R2.15)
\]
Repeatability relative standard deviation \( (RSD_r, \%) = \frac{s_r}{m} \times 100 \quad \cdots \quad (R2.16) \)

Intermediate relative standard deviation \( (RSD_{I(T)}, \%) = \frac{s_I}{m} \times 100 \quad \cdots \quad (R2.17) \)

\( m \): Total mean of the replicate test results on different days

**Note** (4) In case \( V_T < V_r \) assume \( V_T = V_r \) (that is, the test days variance \( (s_{(T)}^2) \) in the formula \( (R2.12) = 0 \)) and let the formula \( (R2.13) \) form \( s_{(T)}^2 = s_r^2 \).

**Example of the calculation of intermediate precision and repeatability by the replicate test results on different days.**

An example of repeatability test results on different days of citric acid-soluble phosphoric acid using sample 1 and sample 2 containing phosphite is shown in Table 4. Conduct one-way analysis of variance for the test results of respective samples to obtain the unbiased variance \( (V) \) of respective variation factors (Table 5).

Examples of the calculation of intermediate precision and repeatability for the sample 1 and sample 2 using the formula \( (R2.11) \) to the formula \( (R2.17) \) are shown in Table 6-1 and 6-2. In addition, the results of respective standard deviation are expressed rounding to the digit of the observed value and the results of the respective relative standard deviations are expressed rounding to the first decimal place.

| Table 4 Example of repeatability test results on different days (mass fraction (%)) |
|---|---|---|---|---|---|---|---|
| Sample No. | Test day (factor) | | | | | |
| Sample 1 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 51.38 |
| | 51.20 | 52.15 | 51.00 | 51.35 | 51.35 | 51.38 | 51.28 |
| | 51.45 | 51.85 | 51.09 | 51.28 | 51.10 | 51.38 | 51.43 |
| Sample 2 | | | | | | | | 5.10 |
| | 5.18 | 4.90 | 5.01 | 5.15 | 5.14 | 5.13 | 5.21 |
| | 5.00 | 5.12 | 5.06 | 5.14 | 5.07 | 5.11 | 5.18 |

1) The mean is expressed rounding to the digit of the observed value.
Table 5

Table of one-way analysis of variance

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>Variation factor</th>
<th>Sum of squares</th>
<th>Degree of freedom</th>
<th>Unbiased variance (V)</th>
<th>Expectation of variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>Test days (T)</td>
<td>1.0570</td>
<td>6</td>
<td>0.17616</td>
<td>( \sigma_r^2 + 2 \times \sigma(T)^2 )</td>
</tr>
<tr>
<td></td>
<td>Accidental error (e)</td>
<td>0.1253</td>
<td>7</td>
<td>0.01789</td>
<td>( \sigma_r^2 )</td>
</tr>
<tr>
<td>Sample 2</td>
<td>Test days (T)</td>
<td>0.0478</td>
<td>6</td>
<td>0.00797</td>
<td>( \sigma_r^2 + 2 \times \sigma(T)^2 )</td>
</tr>
<tr>
<td></td>
<td>Accidental error (e)</td>
<td>0.0448</td>
<td>7</td>
<td>0.00640</td>
<td>( \sigma_r^2 )</td>
</tr>
</tbody>
</table>

Table 6-1 Calculation of intermediate precision and repeatability using the sample 1 replicate test results on different days

<table>
<thead>
<tr>
<th>Variation factor</th>
<th>Unit</th>
<th>Formula</th>
<th>Calculation</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Repeatability variance</td>
<td>( s_r^2 )</td>
<td>( V_r )</td>
<td>0.01789</td>
<td>0.01789</td>
</tr>
<tr>
<td>Repeatability standard deviation ( (s_r)^2 )</td>
<td>Mass fraction (%)</td>
<td>( \sqrt{s_r^2} )</td>
<td>( \sqrt{0.01789} )</td>
<td>0.13</td>
</tr>
<tr>
<td>Repeatability relative standard deviation ( (RSD_r)^3 )</td>
<td>%</td>
<td>( \frac{s_r}{m} \times 100 )</td>
<td>( \frac{0.1338}{51.38} \times 100 )</td>
<td>0.3</td>
</tr>
<tr>
<td>Test days variance ( (s(T))^2 )</td>
<td></td>
<td>( V_T - V_r )</td>
<td>( \frac{0.17616 - 0.01789}{2} )</td>
<td>0.07914</td>
</tr>
<tr>
<td>Intermediate variance ( (s(I))^2 )</td>
<td></td>
<td>( s(T)^2 + s_r^2 )</td>
<td>( 0.09714 + 0.01789 )</td>
<td>0.09703</td>
</tr>
<tr>
<td>Intermediate standard deviation ( (s(I))^2 )</td>
<td>Mass fraction (%)</td>
<td>( \sqrt{s(I)^2} )</td>
<td>( \sqrt{0.09703} )</td>
<td>0.31</td>
</tr>
<tr>
<td>Intermediate relative standard deviation ( (RSD(I))^3 )</td>
<td>%</td>
<td>( \frac{s(I)}{m} \times 100 )</td>
<td>( \frac{0.3115}{51.38} \times 100 )</td>
<td>0.6</td>
</tr>
</tbody>
</table>

1) The rounding of a numerical value is not executed in the middle of the calculation.

2) The standard deviation is expressed rounding to the digit of the observed value.

3) The relative standard deviation is expressed rounding to the first decimal place.
### Table 6-2 Calculation of intermediate precision and repeatability using the sample 2 replicate test results on different days

<table>
<thead>
<tr>
<th>Variation factor</th>
<th>Unit</th>
<th>Formula</th>
<th>Calculation</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Repeatability variance ((s_r^2))</td>
<td></td>
<td>(= V_r)</td>
<td>(= 0.00640)</td>
<td>0.00640</td>
</tr>
<tr>
<td>Repeatability standard deviation ((s_r))</td>
<td>Mass fraction (% (%))</td>
<td>(= \sqrt{s_r^2})</td>
<td>(= \sqrt{0.00640})</td>
<td>0.08</td>
</tr>
<tr>
<td>Repeatability relative standard deviation ((RSD_r))</td>
<td>Mass fraction (%)</td>
<td>(= \frac{s_r}{m} \times 100)</td>
<td>(= \frac{0.0800}{5.10} \times 100)</td>
<td>1.6</td>
</tr>
<tr>
<td>Test days variance ((s_T^2))</td>
<td></td>
<td>(= \frac{V_T - V_r}{n})</td>
<td>(= \frac{0.00797 - 0.00640}{2})</td>
<td>0.00078</td>
</tr>
<tr>
<td>Intermediate variance ((s_{IT}^2))</td>
<td></td>
<td>(= s_T^2 + s_r^2)</td>
<td>(= 0.00078 + 0.00640)</td>
<td>0.00718</td>
</tr>
<tr>
<td>Intermediate standard deviation ((s_{IT}))</td>
<td>Mass fraction (%)</td>
<td>(= \sqrt{s_{IT}^2})</td>
<td>(= \sqrt{0.00718})</td>
<td>0.08</td>
</tr>
<tr>
<td>Intermediate relative standard deviation ((RSD_{IT}))</td>
<td>Mass fraction (%)</td>
<td>(= \frac{s_{IT}}{m} \times 100)</td>
<td>(= \frac{0.0848}{5.10} \times 100)</td>
<td>1.7</td>
</tr>
</tbody>
</table>

Footnote: Refer to Table 6-1
Separate sheet: The target of trueness and the criteria of precision in respective concentration levels

The target of trueness (recovery) and the criteria of precision in respective concentration levels to evaluate Chromatography (1) and test methods other than Chromatography are shown in Table 1 and Table 2. The target of trueness is generally within the recovery of Table 1. As for precision, the permissible level may exceed them by a factor of 2.0.

**Note** (1) Gas chromatography, Gas Chromatography/Mass Spectrometry, High-Performance Liquid Chromatography, High-Performance Liquid Chromatography/Tandem Mass Spectrometry, Ion Chromatography, etc.

<table>
<thead>
<tr>
<th>Concentration level</th>
<th>Recovery (%)</th>
<th>Chromatography</th>
<th>Test methods other than Chromatography</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥25% (mass fraction)</td>
<td>90 - 108</td>
<td>98 - 102</td>
<td></td>
</tr>
<tr>
<td>≥10% (mass fraction)</td>
<td>90 - 108</td>
<td>97 - 103</td>
<td></td>
</tr>
<tr>
<td>≥1% (mass fraction)</td>
<td>85 - 110</td>
<td>96 - 104</td>
<td></td>
</tr>
<tr>
<td>≥0.1% (mass fraction)</td>
<td>85 - 110</td>
<td>94 - 106</td>
<td></td>
</tr>
<tr>
<td>≥100 mg/kg</td>
<td>80 - 115</td>
<td>92 - 108</td>
<td></td>
</tr>
<tr>
<td>≥10 mg/kg</td>
<td>70 - 120</td>
<td>90 - 110</td>
<td></td>
</tr>
<tr>
<td>≥1 mg/kg</td>
<td>70 - 120</td>
<td>85 - 115</td>
<td></td>
</tr>
<tr>
<td>≥100 μg/kg</td>
<td>70 - 120</td>
<td>85 - 115</td>
<td></td>
</tr>
<tr>
<td>≥10 μg/kg</td>
<td>70 - 120</td>
<td>80 - 120</td>
<td></td>
</tr>
<tr>
<td>&lt;10 μg/kg</td>
<td>60 - 125</td>
<td>75 - 125</td>
<td></td>
</tr>
</tbody>
</table>
### Table 2 Criteria of precision \(^1\) in respective concentration levels

<table>
<thead>
<tr>
<th>Concentration level</th>
<th>Chromatography</th>
<th>Test methods other than Chromatography</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Reproducibility relative standard deviation (%)</td>
<td>Intermediate relative standard deviation (%)</td>
</tr>
<tr>
<td>≧25 % (mass fraction)</td>
<td>8</td>
<td>6.5</td>
</tr>
<tr>
<td>≧10 % (mass fraction)</td>
<td>8</td>
<td>6.5</td>
</tr>
<tr>
<td>≧1 % (mass fraction)</td>
<td>8</td>
<td>6.5</td>
</tr>
<tr>
<td>≧0.1 % (mass fraction)</td>
<td>8</td>
<td>6.5</td>
</tr>
<tr>
<td>≧100 mg/kg</td>
<td>8</td>
<td>6.5</td>
</tr>
<tr>
<td>≧10 mg/kg</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td>≧1 mg/kg</td>
<td>16</td>
<td>13</td>
</tr>
<tr>
<td>≧100 μg/kg</td>
<td>22</td>
<td>18</td>
</tr>
<tr>
<td>≧10 μg/kg</td>
<td>22</td>
<td>18</td>
</tr>
<tr>
<td>&lt;10 μg/kg</td>
<td>22</td>
<td>18</td>
</tr>
</tbody>
</table>

\(^1\) As for precision, the permissible level may exceed them by a factor of 2.0.