# INTERNATIONAL STANDARD



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# Copper sulfide concentrates — Determination of copper content — Titrimetric methods

# iTeh STANDARD PREVIEW

Concentrés de sulfuré de curvre — Dosage du cuivre — Méthodes titrimétriques

ISO 10258:1994 https://standards.iteh.ai/catalog/standards/sist/3b82b7c8-3de5-4c10-bfb1-6d6eb77b4eb6/iso-10258-1994



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- 6d6eb77b4eb6/iso-10258-19942 Derivation of precision equations С

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International Organization for Standardization

# Foreword

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ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75% of the member bodies casting

# iTeh STANDARD PREVIEW International Standard ISO 10258 was prepared by Technical Committee

International Standard ISO 10258 was prepared by Technical Committee (SO/TC 183) Copper lead and zinc ores and concentrates.

Annexes A and B form an integral part of this International Standard. Annex C is for information only.

https://standards.iteh.ai/catalog/standards/sist/3b82b7c8-3de5-4c10-bfb1-6d6eb77b4eb6/iso-10258-1994

# iTeh STANDARD PREVIEW (standards.iteh.ai)

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# **Copper sulfide concentrates** — Determination of copper content — Titrimetric methods

#### Scope 1

This International Standard specifies two titrimetric methods for the determination of the copper content of copper sulfide concentrates in the range 15 % (m/m) to 50 % (m/m), using sodium thiosulfate after separation (method 1) or without separation (method 2) of copper from interfering elements.

# 2 Normative references

residual iron, and excess potassium iodide is also The following standards contain provisions which, through reference in this text, constitute provisions 258:19added. Free iodine isolated by reaction between of this International Standard standard standards and and standards sister and a standard standard standard standard standard standards and a standard standard standard standard standard standards and a standard stand 

are subject to revision, and parties to agreements based on this International Standard are encouraged to investigate the possibility of applying the most recent editions of the standards indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.

ISO 385-1:1984. Laboratory glassware — Burettes — Part 1: General requirements.

ISO 648:1977, Laboratory glassware — One-mark pipettes.

ISO 1042:1983, Laboratory glassware — One-mark volumetric flasks.

ISO 4787:1984, Laboratory glassware — Volumetric glassware — Methods for use and testing of capacity.

ISO 9599:1991, Copper, lead and zinc sulfide concentrates — Determination of hygroscopic moisture in the analysis sample — Gravimetric method.

ISO Guide 35:1985, Certification of reference materials — General and statistical principles.

# 3 Principle

# 3.1 Method 1 (Long iodide method)

A test portion is decomposed in nitric and sulfuric acids, and arsenic, antimony and tin are removed by treatment with hydrobromic acid. Copper is separated from interfering elements by precipitation of copper iTeh STANDARI sulfide with sodium thiosulfate. The precipitate is dis-(standards.igen diffuside is added to eliminate interference of solved in nitric and sulfuric acids, ammonium hydro-

# 3.2 Method 2 (Short iodide method)

A test portion is decomposed in nitric and sulfuric acids, and arsenic, antimony and tin are removed by treatment with hydrobromic acid. Ammonium hydrogen difluoride is added to eliminate interference of iron, and excess potassium iodide is also added. Free iodine isolated by reaction between iodide ions and copper(II) ions is titrated with sodium thiosulfate using soluble starch as the indicator.

#### Reagents 4

During the analysis, use only reagents of recognized analytical grade and distilled water or water of equivalent purity.

**4.1 Copper metal**, minimum purity 99,99 %.

# 4.2 Potassium iodide.

4.3 Ammonium hydrogen difluoride.

# **4.4** Sulfuric acid, diluted 1 + 1.

Slowly add 500 ml of concentrated sulfuric acid ( $\rho_{20}$  1,84 g/ml) to 500 ml of water, while stirring and cooling.

# **4.5** Sulfuric acid, diluted 1 + 999.

Add 1 ml of dilute sulfuric acid (4.4) to 500 ml of water.

**4.6** Nitric acid, concentrated ( $\rho_{20}$  1,42 g/ml).

# **4.7** Nitric acid, diluted 1 + 1.

Slowly add 500 ml of concentrated nitric acid (4.6) to 500 ml of water.

**4.8 Hydrofluoric acid** ( $\rho_{20}$  1,14 g/ml).

4.9 Bromine.

4.10 Bromine water, saturated.

4.11 Hydrobromic acid ( $\rho_{20}$  1,50 g/ml).

# **4.12** Acetic acid, diluted 1 + 3.

Slowly add 25 ml of htglaćialndaabetib.ai/cacidg/standa  $(\rho_{20} 1,05 \text{ g/ml})$  to 75 ml of water. 6d6eb77b4eb6/

# 4.13 Nitration mixture.

Slowly add 250 ml of concentrated sulfuric acid ( $\rho_{20}$  1,84 g/ml) to 250 ml of concentrated nitric acid (4.6).

**4.14** Ammonium hydrogen difluoride, 250 g/l solution.

4.15 Sodium carbonate, 20 g/l solution.

**4.16 Sodium thiosulfate pentahydrate**, 200 g/l solution.

4.17 Potassium thiocyanate, 100 g/l solution.

# 4.18 Starch, 2 g/l solution.

Moisten 1 g of soluble starch with cold water, slowly pour into 500 ml of hot water while stirring, and boil for about 1 min.

#### 4.19 Ethanol.

### 4.20 Standard solutions.

NOTE 1 Standard solutions should be prepared at the same ambient temperature as that at which the determinations will be conducted.

**4.20.1 Sodium thiosulfate**, standard volumetric solution (20 g/l).

# 4.20.1.1 Preparation

Dissolve 20 g of sodium thiosulfate (pentahydrate) in 1 litre of freshly boiled and cooled water. Add 0,2 g of sodium carbonate, stir to dissolve and allow to stand for at least one day. Standardize this solution as specified in 4.20.1.2.

# 4.20.1.2 Standardization

Clean a piece of copper metal (4.1) by immersing it in warm dilute acetic acid (4.12). Wash the copper thoroughly with water followed by ethanol (4.19) and allow to dry in air. Weigh, into three separate 400 mil conical beakers to the nearest 0,1 mg, a mass of clean copper metal which approximates the copper tent in the test portion. Record these masses as  $m_1, m_2$  and  $m_3$ .

(standards.iteh.ai) Dissolve the copper using 10 ml of dilute nitric acid (4.7) followed by 5 ml of dilute sulfuric acid (4.4). Heat to evaporate to dryness. Add 40 ml of water, heat to etic ai/cacidg/standards/standard/standar

NOTE 2 The standardization factor of the standard volumetric solution varies with the volume of sample solution, mass of potassium iodide, mass of copper and temperature of solution. The same volume of solution and mass of potassium iodide as those used for the standardization should be used for the analysis of the test portion. The temperatures of standardization and determination should be essentially the same.

Calculate the standardization factors  $f_1$ ,  $f_2$  and  $f_3$  using the following equations:

$$f_1 = \frac{m_1}{V_1} \qquad \dots (1)$$

$$f_2 = \frac{m_2}{V_2} \qquad \dots (2)$$

$$f_3 = \frac{m_3}{V_3}$$
 (3)

Calculate, to four significant figures, the mean standardization factor f for the sodium thiosulfate standard

volumetric solution, provided that the range of the values of  $f_1$ ,  $f_2$  and  $f_3$  does not exceed  $10^{-5}$  gCu/ml. If this range is exceeded, repeat the standardization.

**4.20.2** Copper, standard solution (0,1 mg/ml).

Weigh, to the nearest 0,1 mg, 0,1 g of copper metal (4.1) into a 200 ml beaker, decompose with 10 ml of dilute nitric acid (4.7). Heat to remove nitrogen oxides, cool and add about 50 ml of water. Transfer to a 1 000 ml volumetric flask, fill up nearly to the mark with water, mix and cool to room temperature; then fill up exactly to the mark and mix again.

#### 5 Apparatus

Ordinary laboratory equipment and

5.1 Volumetric glassware, of class A complying with ISO 385-1, ISO 648 and ISO 1042, and used in accordance with ISO 4787.

5.2 Analytical balance, sensitive to 0,1 mg.

Platinum crucibles. 5.3

# 5.4 Atomic absorption spectrometer (AAS), with

a copper hollow cathode lamp.

ISO 10258:1994 Instrumental conditions: https://standards.iteh.ai/catalog/standards/sis/282Blanklefest10-bfb1-

6d6eb77b4eb6/iso-10258-1994

**iTeh STANDARI** 

Flame: air/acetylene

Wavelength: 324,7 nm.

5.5 Inductively coupled plasma (ICP) atomic emission spectrometer (optional).

#### Sample 6

# 6.1 Test sample

Prepare an air-equilibrated test sample in accordance with ISO 9599.

NOTE 3 A test sample is not required if predried test portions are to be used (see annex A).

# 6.2 Test portion

Taking multiple increments, extract a test portion from the test sample as specified in table 1 and weigh to the nearest 0,1 mg. At the same time as test portions are being weighed for analysis, weigh test portions for the determination of hygroscopic moisture in accordance with ISO 9599.

Alternatively, the method specified in annex A may be used to prepare predried test portions directly from the laboratory sample.

Table 1 — Recommended test portion mass
---

Copper conte	nt (presumed)	Mass of test portion
% ()	<i>m/m</i> )	g
≥	<	
15	25	0,8
25	50	0,4

# 7 Procedure

# 7.1 Number of determinations

Carry out the determinations at least in duplicate, as far as possible under repeatability conditions, on each test sample.

NOTE 4 Repeatability, conditions exist where mutually independent test results are obtained with the same method on identical test material in the same laboratory by (standards.ite same operator using the same equipment, within short intervals of time.

> Carry out a blank test in parallel with the analysis using the same quantities of all reagents but omitting the test portion. The purpose of the blank test in this method is to check the quality of reagents. If a significant blank titration value is obtained as a result of the blank test, check all reagents and rectify the problem.

# 7.3 Determination — method 1: Long iodide method

# 7.3.1 Decomposition of test portion

Transfer the test portion to a 400 ml conical beaker and moisten with 10 ml of water. Add 20 ml of dilute nitric acid (4.7), cover with a watch glass and heat for about 10 min at 60 °C to 70 °C. Add 10 ml of dilute sulfuric acid (4.4) and heat gradually to decompose the test portion.

After the completion of the initial reaction, rinse the underside of the watch glass with a minimum volume of water, collecting the washings in the conical beaker. Continue heating until strong white fumes are evolved, then cool.

If the residue appears dark (presence of carbon), slowly add a small amount of the nitration mixture (4.13) to the hot solution until the solution becomes colourless or bluish and heat until strong white fumes are evolved.

If decomposition of the deposited sulfur is insufficient, add 5 ml of nitric acid (4.6) and 1 ml of bromine (4.9), and heat until strong white fumes are evolved.

Carefully add 5 ml of water and 10 ml of hydrobromic acid (4.11) and heat until strong white fumes are evolved. Remove from the source of heat and cool. After addition of 5 ml of dilute sulfuric acid (4.4) and 10 ml of hydrobromic acid (4.11), heat until strong white fumes are evolved. Remove from the source of heat and cool.

Add 80 ml of water, warm to dissolve soluble salts, and heat until boiling. Filter through a medium porosity filter paper, wash well with hot water and collect the filtrate in a 400 ml conical beaker. Reserve the filter paper and residue for the determination of copper by flame atomic absorption spectrometry (FAAS) (as described in 7.3.5) unless it has been proven, through previous testing, that the copper in the sample is completely soluble using the initial dissolution.

to dryness. Use more nitration mixture if the residue appears dark. Continue heating strongly to destroy any elemental sulfur. After adding 10 ml of nitric acid (4.6) around the top of the beaker to rinse away the residual sulfur, add 2 ml of dilute sulfuric acid (4.4) and heat until strong white fumes are evolved. Remove from the heat source and cool. Add 40 ml of water, warm to dissolve the soluble salts and cool. Proceed to 7.3.4.

# 7.3.3 Dissolution of copper precipitate

Add 2 ml of dilute sulfuric acid (4.4) and 10 ml of nitric acid (4.6), heat slowly to decompose the precipitate and then evaporate to dryness. Continue heating strongly to destroy any elemental sulfur. After adding 10 ml of nitric acid (4.6) around the top of the beaker to rinse away the residual sulfur, add 2 ml of dilute sulfuric acid (4.4) and heat until strong white fumes are evolved. Remove from the source of heat and cool.

### 7.3.4 Titration RD PREVIEW

Add 40 ml of water, warm to dissolve the soluble Galts and cool the solution. Add sodium carbonate solution (4.15) until the copper precipitate appears,

ISO 102themad dilute acetic acid (4.12) until the copper prehttps://standards.iteh.ai/catalog/stand.cipitate/disappears/and/antexcess of 3 ml to 5 ml. Add 6d6eb77b4eb6/isom020f-10mmonium hydrogen difluoride solution

# 7.3.2 Separation of copper

Dilute the filtrate to 200 ml and heat to 70 °C to 90 °C, slowly add 40 ml of sodium thiosulfate solution (4.16) while stirring, to produce a yellow or yellowish brown emulsion. Heat gradually and continue boiling gently until the precipitate coagulates. Filter the solution through a medium porosity filter paper and wash the filter paper and precipitate with hot water. Retain the filtrate for FAAS measurements of copper (as described in 7.3.5).

Using water, rinse away the copper sulfide precipitate into the original conical beaker and decompose the remaining precipitate on the filter paper using drop by drop addition of bromine water (4.10) followed by nitric acid (4.6). Repeat this treatment as required, then wash well with hot water, collecting this solution in the beaker containing the main precipitate. Retain the filter paper for FAAS measurements of copper (as described in 7.3.5).

NOTE 5 Instead of using the above step, the following method can be used. Transfer the precipitate and filter paper into the original beaker, cover with a watch glass and add 30 ml of nitration mixture (4.13). Heat slowly to decompose the precipitate and the filter paper and evaporate

(4.14) and swirl.

Add 15 g of potassium iodide (4.2), swirl to dissolve, and immediately titrate with sodium thiosulfate standard volumetric solution (4.20.1). When the yellow brown iodine colour fades to a pale yellow, add 5 ml of starch solution (4.18) as the indicator.

#### NOTES

6 Instead of using the above step, the following method can be used. Add 3 g of potassium iodide (4.2), swirl to dissolve and immediately titrate with sodium thiosulfate standard volumetric solution (4.20.1). When the yellow brown iodine colour fades to a pale yellow, add 5 ml of starch solution (4.18) as the indicator and continue the titration until the colour of the solution becomes light blue. Then add 5 ml of potassium thiocyanate solution (4.17).

7 The presence of Ag, Bi, Hg and Pb may obscure the colour change. In this case, add the starch solution (4.18) earlier in the titration, when the solution is a light brown colour.

Continue the titration until the blue indicator colour just disappears. Record the volume V of sodium thiosulfate standard volumetric solution used in the titration.

# 7.3.5 FAAS determination of copper in the insoluble residue, filtrate and filter paper

# 7.3.5.1 Decomposition of the insoluble residue

Place the retained residue and the filter paper in a platinum crucible (5.3), dry and ignite at 750 °C to 800 °C. Allow the crucible to cool, add 5 ml of dilute sulfuric acid (4.4) and 5 ml to 10 ml of hydrofluoric acid (4.8), heat to evaporate almost to dryness and volatilize the silicon as silicon tetrafluoride. Dissolve with a small quantity of water and 1 ml of dilute sulfuric acid (4.4) by heating. Proceed to 7.3.5.3.

### 7.3.5.2 Decomposition of the precipitate remaining on the filter paper

Transfer the retained filter paper into a beaker and add 30 ml of nitration mixture (4.13). Heat to evaporate to dryness. If the residue appears dark (presence of carbon), repeat this step. Dissolve with a small quantity of water and 1 ml of dilute sulfuric acid (4.4) by heating. Proceed to 7.3.5.3.

# 7.3.5.3 Spectrometric measurement

and make up to the marks with water.

where

- is the mass, in grams, of copper in the in $m_{\Lambda}$ soluble residue, the precipitate remaining on the filter paper and the filtrate;
- $m_5$  is the mass, in micrograms, of copper in the test solution.

# 7.4 Determination — method 2: Short iodide method

# 7.4.1 Decomposition of the test portion

Transfer the test portion to a 400 ml conical beaker and moisten with 10 ml of water. Add 20 ml of dilute nitric acid (4.7), cover with a watch glass and heat for about 10 min at 60 °C to 70 °C. Add 10 ml of dilute sulfuric acid (4.4) and heat gradually to decompose the test portion.

After completion of the initial reaction, rinse the underside of the watch glass with a minimum volume iTeh STANDARI of water, collecting the washings in the conical (standards. beaker. Continue heating until strong white fumes are evolved, then cool.

<u>O 10258:1994</u> the residue appears dark (presence of carbon), Transfer the solutions prepared in 7.3.5.1. 7.3.5.2 and logs and add add add add add a small amount of the nitration mixture the retained filtrate from 7.3.2 into 666eb5994eb6/iso-10(2513)916 the hot solution until the solution becomes volumetric flask and make up to the mark with water. colourless or bluish and heat until strong white fumes are evolved. Prepare calibration solutions by adding, from a pipette

or a micro-burette, 0,0 ml, 0,50 ml, 1,00 ml, 1,50 ml, If decomposition of the deposited sulfur is insuf-2.00 ml and 3,00 ml of copper standard solution ficient, add 5 ml of nitric acid (4.6), 1 ml of bromine (4.20.2) into a series of 200 ml one-mark volumetric (4.9) and 2 ml of dilute sulfuric acid (4.5), and heat flasks, add 1 ml of dilute sulfuric acid (4.4) to each one until strong white fumes are evolved.

> Carefully add 5 ml of water, 10 ml of hydrobromic acid (4.11) and 5 ml of dilute sulfuric acid (4.4) and heat until strong white fumes are evolved. Remove from the source of heat and cool. Add 5 ml of dilute sulfuric acid (4.4) and 10 ml of hydrobromic acid (4.11), and heat until strong white fumes are evolved. Continue heating to evaporate to complete dryness and then cool.

> NOTE 9 If it has not been proven, through previous testing, that the copper in the sample is completely soluble using the initial dissolution described above, the following procedure should be carried out. Add 20 ml of water, warm to dissolve soluble salts, then heat until boiling. Filter through a medium-porosity filter paper, wash well with hot water collecting the filtrate and washings in a 400 ml conical beaker, and then heat to evaporate to dryness. Determine the copper content of the insoluble residue in accordance with 7.3.5.

Aspirate the test solution and the calibration solutions into the atomic absorption spectrometer (5.4) using an air/acetylene flame and a wavelength of 324,7 nm with background correction.

Prepare a calibration graph of masses of copper in the calibration solutions versus absorbances and read the mass, in micrograms, of copper in the test solution from the calibration graph.

NOTE 8 Alternatively, the ICP atomic emission spectrometer (5.5) can be used for the determination of copper at a wavelength of 324,7 mm.

Calculate the mass of copper in the residue and filtrate using the following equation:

$$m_4 = m_5 \times 10^{-6}$$
 ... (4)

### 7.4.2 Titration

Add 40 ml of dilute sulfuric acid (4.5), warm to dissolve the soluble salts and cool the solution. Add 3 g of ammonium hydrogen difluoride (4.3) to the test solution and swirl to dissolve.

Add 15 g of potassium iodide (4.2), swirl to dissolve and immediately titrate with sodium thiosulfate standard volumetric solution (4.20.1). When the yellow brown iodine colour fades to a pale yellow, add 5 ml of starch solution (4.18) as the indicator.

#### NOTES

10 Instead of using the above step, the following method can be used. Add 3 g of potassium iodide (4.2), swirl to dissolve and immediately titrate with sodium thiosulfate standard volumetric solution (4.20.1). When the yellow brown iodine colour fades to a pale yellow, add 5 ml of starch solution (4.18) as the indicator and continue the titration until the colour of the solution becomes light blue. Then add 5 ml of potassium thiocyanate solution (4.17).

11 The presence of Ag, Bi, Hg and Pb may obscure the colour change. In this c earlier in the titration, v colour.

Continue the titration until the blue indicator colour where where

just disappears. Record the volume V of sodium

thiosulfate standard volumetric solution used in the solution  $10258:19\overline{x}_{4}$  is the mean content of copper, expressed as https://standards.iteh.ai/catalog/standards/sist/3b8alpercentage bybribass, in the sample; titration. 6d6eb77b4eb6/iso-10258-1994

8 Expression of results

The copper content of the test portion  $w_{Cu}$ , expressed as a percentage by mass, is given by the following equation:

$$w_{\rm Cu} = \frac{\left[ (V \cdot f) + m_4 \right] \times 100}{m} \times \frac{100}{100 - H} \qquad \dots (5)$$

where

- is the volume, in millilitres, of sodium Vthiosulfate standard volumetric solution used;
- is the mean standardization factor, in grams fof copper per millilitre, for the sodium thiosulfate standard volumetric solution, calculated in 4.20.1.2;
- $m_A$  is the mass, in grams, of residual copper determined by FAAS, calculated in 7.3.5.3;
- is the mass, in grams, of the test portion; m

H is the hygroscopic moisture content, in percent, of the test portion (in the case of a predried test portion being used, H = 0).

Calculate the copper content of the test portion to the second decimal place.

# 9 Precision

#### 9.1 Expression of precision

The precision of this analytical method is expressed by the following equations:

#### Long iodide method

$s_{\rm r} = 0,000 \ 8 \ \overline{X} + 0,048 \ 5$	(6)
--	-----

$$s_{\rm L} = 0,004 \ 2 \ \overline{X} - 0,007 \ 7 \qquad \dots (7)$$

#### Short iodide method

case, add the starch solution (4.18)	$s_{\rm r} = 0,001 \ 4 \ \overline{X} + 0,028 \ 2$	(8)
when the solution is a Sight brown	ARD <sub>s</sub> P0,000 5 x F0,081 9	(9)
(standa)	rds itah ai)	

is the within-laboratory standard deviation, Sr. expressed as a percentage by mass of copper;

is the between-laboratories standard devi- $S_{\parallel}$ ation, expressed as a percentage by mass of copper:

NOTE 12 Additional information is given in annex C.

# 9.2 Method for obtaining the final result

(see annex B)

Calculate the following quantities from the duplicate results  $X_1$  and  $X_2$  and process according to the flowchart in annex B:

Mean of duplicates

$$\overline{X} = (X_1 + X_2)/2 \qquad \dots (10)$$

Within-laboratory standard deviation

Long iodide method

$$s_{\rm r} = 0,000 \ 8 \ \overline{X} + 0,048 \ 5 \qquad \dots \ (11)$$

Short iodide method

$$s_r = 0,001 \ 4 \ \overline{X} + 0,028 \ 2 \qquad \dots (12)$$

Repeatability limit

$$r = 2,8s_{\rm r} \qquad \dots (13)$$

#### 9.3 Precision between laboratories

The precision between laboratories is used to determine the agreement between the results reported by two (or more) laboratories. It is assumed that all the laboratories followed the same procedure.

Calculate the following quantities:

Mean of final results

... (14)  $\mu_{1,2} = (\mu_1 + \mu_2)/2$ 

Between-laboratories standard deviation

If E is less than or equal to P, the final results are in agreement.

#### 9.4 Check of trueness

The trueness of the analytical method can be checked by applying it to a certified reference material (CRM). The procedure is the same as that described in clause 7. When the precision has been confirmed, the final laboratory result can be compared with the certified value,  $A_{c}$ .

The following two possibilities exist:

$$|\mu_{\rm c} - A_{\rm c}| \leqslant C \qquad \dots (21)$$

If this condition exists, the difference between the reported result and the certified value is statistically insignificant.

$$|\mu_{\rm c} - A_{\rm c}| > C \qquad \dots (22)$$

If this condition exists, the difference between the reported result and the certified value is statistically significant

Long iodide method **iTeh STANDARD** in equations (21) and (22), the symbols have the fol-  

$$s_{L} = 0,004 \ 2 \ \mu_{1,2} - 0,007 \ 7$$
 (standards.ibwing meanings:

Short iodide method

 $\mu_{\rm c}$  is the final result, expressed as a percentage ISO 10258:1994  $s_{\rm L} = 0,0005 \ \mu_{\rm htps:}/9.081r_{\rm S.iteh.ai/catalog(16)lards/sist/3b82b7c8-3de5-4c10-bib1-$ 6d6eb77b4eb6/iso-10258-1994

Within-laboratory standard deviation

Long iodide method

$$s_r = 0,000 \ 8 \ \mu_{1,2} + 0,048 \ 5 \qquad \dots \ (17)$$

Short iodide method

$$s_r = 0,001 \ 4 \ \mu_{1,2} + 0,028 \ 2 \qquad \dots (18)$$

Permissible difference

$$P = 2.8\sqrt{s_{\rm L}^2 + (s_{\rm r}^2/2)}$$
 ....(19)

Range

$$E = |\mu_1 - \mu_2| \qquad \dots (20)$$

where

- $\mu_1$  is the final result, expressed as a percentage by mass of copper, reported by laboratory 1:
- $\mu_2$  is the final result, expressed as a percentage by mass of copper, reported by laboratory 2.

- A<sub>c</sub> is the certified value, expressed as a percentage by mass of copper, of the certified reference material:
- C is a quantity, expressed as a percentage by mass of copper, depending on the type of the certified reference material used, as defined in 9.4.1.

#### 9.4.1 Type of certified reference material (CRM) or reference material (RM)

The reference materials used for this purpose should be prepared and certified in accordance with ISO Guide 35.

#### 9.4.1.1 Reference material certified/characterized by an interlaboratory test programme

The quantity C (see 9.4), expressed as a percentage by mass of copper, is given by the following equation:

$$C = 2\sqrt{s_{\rm L}^2 + (s_{\rm r}^2/n) + S^2\{A_{\rm c}\}} \qquad \dots (23)$$

where