

SLOVENSKI STANDARD SIST ISO 7520:1997

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Feronikelj - Določanje kobalta - Plamenska atomska absorpcijska spektrometrična metoda

Ferronickel -- Determination of cobalt content -- Flame atomic absorption spectrometric method

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Ferro-nickel -- Dosage du cobalt st Méthode par spectrométrie d'absorption atomique dans la flamme

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International Standard



INTERNATIONAL ORGANIZATION FOR STANDARDIZATION MEX CHARCON OF A PRANCING A CTANDAPT A CARDINAL ORGANISATION INTERNATIONALE DE NORMALISATION

Ferronickel — Determination of cobalt content — Flame atomic absorption spectrometric method

Ferro-nickel — Dosage du cobalt — Méthode par spectrométrie d'absorption atomique dans la flamme

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Descriptors : nickel, ferronickel, chemical analysis, determination of content, cobalt, atomic absorption method.

ISO 7520:199

Foreword

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.

Draft International Standards adopted by the technical committees are circulated to the member bodies for approval before their acceptance as International Standards by the ISO Council. They are approved in accordance with ISO procedures requiring at least 75 % approval by the member bodies voting, 'ANDARD PREVIEW

International Standard ISO 7520 was prepared by Technical Committee ISO/TC 155, Nickel and nickel alloys.

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Users should note that all International Standards undergo revision from time to time and that any reference made herein to any other threin attornational Standard implies lits - 8b6e-4db9-9e03latest edition, unless otherwise stated. 5c8533a9c43a/sist-iso-7520-1997

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SIST ISO 7520:1997

Ferronickel — Determination of cobalt content — Flame atomic absorption spectrometric method

1	Scope and field of application	4.4	Perchloric acid, $\rho_{20} = 1,61 \text{ g/ml} [72 \% (m/m)].$		
This International Standard specifies a flame atomic absorption spectrometric method for the determination of the cobalt content of ferronickel in the range 0,025 to 2,5 % (m/m) .		4.5	Hydrofluoric acid, $\varrho_{20} = 1,14$ g/ml, diluted 1 + 1.		
		WARNING — Hydrofluoric acid is extremely irritating and corrosive to skin and mucous membranes, producing			
2	References	cont	severe skin burns which are slow to heal. In case of skin contact wash well with water and seek medical advice.		
ISC	0 385/1, Laboratory glassware T Burettes T Part 1: General R	4.6	Lanthanum, solution, containing 200 g of La per litre.		
ISC	0 648, Laboratory glassware – One-mark pipettes.	Weig and	h 250 g of lanthanum chloride hexahydrate (LaCl ₃ ·6H ₂ O) ransfer to a 600 ml beaker. Add 25 ml of hydrochloric acid		
ISO 1042, Laboratory glassware – One-mark volumetric flasks.			(4.1) and 300 ml of water. Stir to complete dissolution. Filter, if <u>Stinec</u> essary, into a 500 ml one-mark volumetric flask, make up to		
IS(rep	0 5725, Precision of testpriethods d s in Determination of ds/s beatability and reproducibility by inter-laboratory tests 43a/sist-iso	ist be d -7520-	nark4with6water/and/mix. 1997		
IS(Dii	0 6352, Ferronickel — Determination of nickel content — methylglyoxime gravimetric method.	4.7 and	Nickel plus iron , matrix solution, containing 12 g of Ni 28 g of Fe per litre.		
3 Dis	Principle ssolution of a test portion in a nitric acid-hydrochloric acid	4.7 .1 less 50 m the i	Weigh 12,0 g of high purity nickel powder [containing than 0,001 % (m/m) cobalt] into an 800 ml beaker. Add I of water and 50 ml of nitric acid ($\varrho_{20} = 1,41$ g/ml). When nitial reaction subsides, stir and heat to complete dissolu-		
mi aci elii by a v	ixture. Precipitation of silica by dehydration in perchloric id. Removal of silica by filtration. Addition of lanthanum for mination of potential interferences. Determination of cobalt atomic absorption spectrometry in an air-acetylene flame at wavelength of 240,7 nm.	4.7.2 than of hy acid	2. Weigh 28,0 g of high purity iron powder [containing less 0,001 % (m/m) cobalt] into an 800 ml beaker. Add 100 ml vdrochloric acid diluted 1 + 1. Carefully add 50 ml of nitric ($g_{20} = 1,41$ g/ml) and heat to complete dissolution and		
NC use	DTE — This analysis can be carried out on solutions which have been ed for the determination of nickel by ISO 6352.	oxida 4.7.3 iron flask	ation of iron. Dilute to about 250 ml. Carefully combine the nickel solution (4.7.1) with the solution (4.7.2). Filter into a 1 000 ml one-mark volumetric , make up to the mark with water and mix.		
4	Reagents				
Du	uring the analysis, use only reagents of recognized analytical	4.8 Cor	Cobalt , standard solution, corresponding to 0,500 g of ther litre.		

Weigh, to the nearest 0,001 g, 0,500 g of high purity [99,9 % (m/m) Co, minimum] cobalt powder, transfer to a 600 ml beaker and add 40 ml of nitric acid (4.3). Heat to complete dissolution, boil gently to expel oxides of nitrogen, cool and transfer to a 1 000 ml one-mark volumetric flask containing 160 ml of nitric acid (4.3). Make up to the mark with water and mix.

1 ml of this standard solution contains 0,500 mg of Co.

grade and only distilled water or water of equivalent purity.

4.2 Hydrochloric acid, $\varrho_{20} = 1,19$ g/ml, diluted 1+9.

4.1 Hydrochloric acid, $\varrho_{20} = 1,19$ g/ml.

4.3 Nitric acid, $\rho_{20} = 1,41$ g/ml, diluted 1+1.

5 Apparatus

Ordinary laboratory glassware, and

5.1 Atomic absorption spectrometer, equipped with a laminar flow burner for an air-acetylene flame and a cobalt hollow cathode lamp.

5.2 Burette, of capacity 50 ml, graduated in divisions of 0,1 ml, in accordance with ISO 385/1, class A.

5.3 Glass beakers, of capacity 600 ml, clean, unetched and flat bottomed.

5.4 Pipettes, of capacities 25 and 50 ml, in accordance with ISO 648, class A.

5.5 Volumetric flasks, of capacities 250; 500; and 1 000 ml, in accordance with ISO 1042, class A.

5.6 Polytetrafluoroethylene (PTFE) beaker, of capacity 600 ml, for samples with a high silicon content.

iTeh STAND 6 Sampling and samples

6.1 Sampling and preparation of the laboratory sample shall be carried out by normal agreed procedures or, in case of dispute, by the relevant International Standard.

6.2 The laboratory sample normally is in the form of millings, c43a/sist-iso-7520-1997 drillings or granules and no further preparation of the sample is necessarv.

6.3 If it is suspected that the laboratory sample is contaminated with oil or grease from the milling or drilling process, it shall be cleaned by washing with high purity acetone and drying in air.

6.4 If the laboratory sample contains particles or pieces of widely varying sizes, the test portion should be obtained by riffling.

7 Procedure

WARNING - Fuming perchloric acid is a powerful oxidant and can cause an explosive mixture when in contact with organic materials. All evaporations should be done in fume cupboards suitable for use with perchloric acid.

7.1 Test portion

Weigh, to the nearest 0,001 g, 3,9 to 4,1 g of the laboratory sample and transfer to a glass beaker (5.3).

7.2 Blank test

Carry out a blank test in parallel with the determination, following the same procedure and using the same quantities of all the reagents.

7.3 Preparation of test solution

7.3.1 Dissolve the test portion (7.1) by adding 25 ml of water followed by 50 ml of nitric acid (4.3). Cover the beaker with a watch-glass and heat gently, if necessary, to complete dissolution.

NOTE – For ferronickel samples containing more than 1 % (m/m)silicon, use a polytetrafluoroethylene beaker (5.6). Attack the test portion by adding successively 25 ml of water, 40 ml of nitric acid (4.3) and 10 ml of hydrochloric acid (4.1). To obtain complete dissolution of the sample, add, at the end of effervescence, 10 ml of hydrofluoric acid (4.5) and 40 ml of perchloric acid (4.4). Heat until evolution of fumes of perchloric acid. Allow to cool and transfer the solution quantitatively to a glass beaker (5.3). Heat at 260 °C until abundant white fumes are obtained. Reflux at this temperature for 20 min and proceed as directed in 7.3.2, "Remove the beaker ...".

7.3.2 When the metal is dissolved, add 40 ml of perchloric acid (4.4) and heat at 260 °C until abundant white fumes are obtained. Reflux at this temperature for 20 min. Remove the beaker from the hotplate and allow to cool. Add 20 ml of hydrochloric acid (4.1) and 200 ml of warm water. Filter off the silica using a medium porosity filter paper, collecting the filtrate in a 1 000 ml one-mark volumetric flask. Rinse the beaker and wash the silica precipitate three times with hydrochloric acid diluted 1 + 9 and four times with warm water. Discard the silica precipitate. Add 50 ml of lanthanum solution (4.6) to the filtrate, make up to the mark with water and mix thoroughly (test solution A).

7.4/c Preparation of calibration solutions https://standards.iteh.ai/catalog/stand

7.4.1 Set A

7.4.1.1 Transfer six 50,0 ml volumes of the nickel plus iron matrix solution (4.7) to six 150 ml beakers.

7.4.1.2 Add, using a burette (5.2), 0; 1,0; 2,0; 3,0; 5,0; and 10.0 ml respectively of the cobalt standard solution (4.8). Add 20 ml of perchloric acid (4.4) and heat just to the liberation of white fumes. Cool, add 50 ml of water and transfer quantitatively into a 500 ml one-mark volumetric flask containing 10 ml of hydrochloric acid (4.1). Add 25 ml of lanthanum solution (4.6), make up to the mark with water and mix.

7.4.2 Set B

7.4.2.1 Transfer six 5 ml volumes of the nickel plus iron matrix solution (4.7) to six 150 ml beakers.

7.4.2.2 See 7.4.1.2.

7.5 Calibration and determination

7.5.1 Expected cobalt contents 0,025 to 0,25 % (m/m)

Carry out the determination on test solution A (7.3.2) and the set of calibration solutions A (7.4.1).

7.5.1.1 Set the required instrument parameters of the atomic absorption spectrometer according to the manufacturer's instructions using a cobalt hollow cathode lamp at a wavelength of 240,7 nm.

7.5.1.2 Light the burner and adjust the air and acetylene flows to obtain a slightly oxidizing clear non-luminescent flame while aspirating water.

7.5.1.3 Aspirate test solution A (7.3.2) into the flame and note the absorbance reading.

7.5.1.4 Aspirate water and check that the reading returns to zero.

7.5.1.5 Aspirate the set of calibration solutions A (7.4.1) in order of increasing instrument response and note the absorbance readings. Flush the system with water between each reading.

7.5.1.6 Aspirate the blank test solution and note the reading.

7.5.1.7 Repeat the measurement of the test solution, blank and calibration solutions to check the repeatability of the readings.

NOTES

1 Poor repeatability may indicate an instrument malfunction or a variation in the aspiration rate among the solutions arcatalog/standards/si

2 Ensure that the measurement corresponding to the test sample falls between two calibration solutions containing cobalt as close as possible to that being determined.

7.5.2 Expected cobalt content 0,25 to 2,5 % (m/m)

7.5.2.1 Transfer, using a pipette, 25,0 ml of test solution A (7.3.2) into a 250 ml one-mark volumetric flask. Add 12 ml of lanthanum solution (4.6), 10 ml of perchloric acid (4.4) and 5 ml of hydrochloric acid (4.1). Make up to the mark with water and mix (test solution B).

7.5.2.2 Dilute the blank test (7.2) as directed in 7.5.2.1.

7.5.2.3 Proceed with the determination as directed in 7.5.1 using test solution B (7.5.2.1), blank test (7.5.2.2) and the set of calibration solutions B (7.4.2).

7.6 Preparation of calibration graphs

Plot the instrument reading for each of the calibration solutions less the reading for the zero member against the concentration of cobalt in the calibration solution.

NOTE — Some instruments may be adjusted to give a read-out directly in concentration of the analyte. A graph of instrument response versus concentration should be plotted to check the linearity and validity of the readings.

8 Expression of results

8.1 Calculation

8.1.1 Determine the concentration of cobalt in the test solution and in the blank from the calibration graph.

8.1.2 The cobalt content, expressed as a percentage by mass, is given by the formula

$$\frac{q(\varrho_1 - \varrho_0)}{m} \times 10^{-4} \times f$$

where

V

 ϱ_1 is the concentration, expressed in milligrams per litre, of cobalt in the test solution;

 ϱ_0 is the concentration, expressed in milligrams per litre, of cobalt in the blank test;

m is the mass, in grams, of the test portion;

V is the volume, in millilitres, of the test solution;

f is the dilution factor of the test solution (f = 10 for test solution B).

8.2 Precision

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This International Standard was subjected to a limited interlaboratory test programme involving seven laboratories in four countries. Eight samples of commercial ferronickel covering the range 21 to 41 % (m/m) nickel and 0,6 to 1,3 % (m/m) cobalt were analysed.

Repeatability and reproducibility were calculated according to ISO 5725 with the results given in the table. It is expected that r and R will be correspondingly lower for lower cobalt levels.

Table — nesults of statistical analysis	Table –	Results	of	statistical	analysis
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Nickel Content [% (m/m)] Cobalt content [% (m/m)]	20 to 25 0,7	25 to 40 0,7 to 1,3
Standard deviation — within laboratory, s_w — between laboratories, s_b	0,021 0,013	0,044 0,028
Repeatability, $r = 2,83 \sqrt{s_w^2}$	0,06	0,12
Reproducibility , $R = 2,83 \sqrt{s_w^2 + s_b^2}$	0,07	0,15

9 Special cases

If the test solution from the determination of nickel by the dimethylglyoxime gravimetric method is used, an appropriate aliquot shall be taken and lanthanum added for the determination of cobalt following the principles of 7.5.2.

10 Test report

The test report shall include the following information:

- a) the reference of the method used;
- b) the results of the analysis;
- c) the number of independent replications;
- d) any unusual features noted during the analysis;

e) any operation not included in this International Standard or regarded as optional.