

SLOVENSKI STANDARD SIST ISO 9682-1:2000

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Iron ores -- Determination of manganese content -- Part 1: Flame atomic absorption spectrometric method

iTeh STANDARD PREVIEW

Minerais de fer -- Dosage du manganèse an Partie 1: Méthode par spectrométrie d'absorption atomique dans la flamme

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Iron ores

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INTERNATIONAL STANDARD

ISO 9682-1

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Iron ores — Determination of manganese content —

Part 1: iTeh SFlame atomic absorption spectrometric method (standards.iteh.ai)

Minerais de fer Dosage du manganèse -

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Reference number ISO 9682-1:1991(E)

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. 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 VIEW bodies casting a vote.

International Standard ISO 9682-1 was prepared by Technical Committeel) ISO/TC 102, *Iron ores*, Sub-Committee SC 2, *Chemical analysis*.

ISO 9682 consists of the following parts, under the general little from gress-564a-4b59-ac84-— Determination of manganese content: 3a7td467475fsist-iso-9682-1-2000

- Part 1: Flame atomic absorption spectrometric method
- Part 2: Periodate spectrophotometric method (At present available under reference ISO 3886:1986)

Annex A forms an integral part of this part of ISO 9682. Annexes B and C are for information only.

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Iron ores — Determination of manganese content —

Part 1:

Flame atomic absorption spectrometric method

Scope 1

This part of ISO 9682 specifies a flame atomic absorption spectrometric method for the determination of the manganese content of iron ores.

Decomposition of a test portion by one of the fol-This method is applicable to manyanese contents lowing methods. between 0,01 % (m/m) and 3 % (m/m) in natural iron ores, iron ore concentrates and aggiomerates CIS. itch ai) of including sinter products.

ISO 7764:1985, Iron ores - Preparation of predried test samples for chemical analysis.

3 Principle

the test portion in sodium carbonate/sodium tetraborate flux and dissol-

SIST ISO 9682-1:200tion of the cooled melt in hydrochloric acid. https://standards.iteh.ai/catalog/standards/sist/db193c54-564a-4b59-ac84-

3a7fd467475f/sist-iso-9b)82Treatment with hydrochloric acid, with the ad-

2 Normative references

The following standards contain provisions which, through reference in this text, constitute provisions of this part of ISO 9682. At the time of publication, the editions indicated were valid. All standards are subject to revision, and parties to agreements based on this part of ISO 9682 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 648:1977, Laboratory glassware — One-mark pipettes.

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

ISO 3081:1986, Iron ores - Increment sampling -Manual method.

ISO 3082:1987, Iron ores - Increment sampling and sample preparation — Mechanical method.

ISO 3083:1986, Iron ores - Preparation of samples - Manual method.

dition of a small amount of nitric acid. Evaporation to dehydrate silica, addition of hydrochloric acid, dilution and filtration. Ignition of the residue and removal of silica by evaporation with hydrofluoric and sulfuric acids. Fusion with sodium carbonate and sodium tetraborate and dissolution of the cooled melt in the filtrate.

Aspiration of the solution obtained in either case into the flame of an atomic absorption spectrometer using a dinitrogen oxide-acetylene flame.

Comparison of the absorbance values obtained with those obtained from the calibration solutions.

4 Reagents

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

4.1 **Sodium carbonate** (Na₂CO₃), anhydrous.

Sodium tetraborate (Na₂B₄O₇), anhydrous. 4.2

4.3 Hydrochloric acid, ρ 1,16 g/ml to 1,19 g/ml.

4.4 Hydrochloric acid, ρ 1,16 g/ml to 1,19 g/ml, diluted 1 + 1.

4.5 Nitric acid, ρ 1,4 g/ml.

4.6 Hydrofluoric acid, ρ 1,13 g/ml, 40 % (m/m), or ρ 1,19 g/ml, 48 % (m/m).

4.7 Sulfuric acid, ρ 1,84 g/ml.

Sulfuric acid, ρ 1,84 g/ml, diluted 1 + 1. 4.8

4.9 Background solution.

Dissolve 12 g of metallic iron (wire or powder) or 17 g of iron oxide [in both cases with manganese content less than 0,003 % (m/m)], in 100 ml of hydrochloric acid (4.3) by heating in a covered 1000 ml beaker. If metallic iron is used, oxidize by adding nitric acid (4.5) drop by drop. Cool and add 300 ml of hydrochloric acid (4.3). Dissolve 32 g of sodium carbonate (4.1) and 16 g of sodium tetraborate (4.2) in 300 ml of water, add carefully to the iron solution and heat to remove carbon dioxide. Cool, transfer to a 1000 ml one-mark volumetric flask, dilute to volume with water and mix.

4.10 Manganese standard solutions.

Apparatus 5

Ordinary laboratory apparatus, including one-mark pipettes and one-mark volumetric flasks complying with the specifications of ISO 648 and ISO 1042 respectively, and

5.1 Platinum or suitable platinum alloy crucibles. of minimum capacity 25 ml.

5.2 Muffle furnace, to provide a minimum temperature of 1020 °C.

5.3 Combined magnetic stirrer/hotplate.

5.4 Stirring bars, polytetrafluoroethylene (PTFE)coated, 10 mm long.

NOTE 2 Items 5.3 and 5.4 are required only when decomposition by alkaline fusion is used.

5.5 Atomic absorption spectrometer, equipped with a dinitrogen oxide-acetylene burner.

The atomic absorption spectrometer used in this method will be satisfactory if it meets the following criteria.

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a) Minimum sensitivity - the absorbance of the SIST ISO 9682-1,2000 c and a manganese calibration solution (4.11) is at least 0.27.

4.10.1 Manganese stock solution, 500aug Mn/mbtalog/standards

manganese Dissolve 0,500 g of high purity [minimum purity 99,9 % (m/m)] in 25 ml of hydrochloric acid (4.4) in a covered tall-form beaker, while heating. When dissolution is complete, cool, transfer the solution to a 1000 ml one-mark volumetric flask, dilute to volume with water and mix.

4.10.2 Manganese standard solution, 50 µg Mn/ml.

Transfer 50 ml of stock solution (4.10.1) to a 500 ml one-mark volumetric flask, dilute to volume with water and mix.

4.11 Manganese calibration solutions.

To a series of 100 ml one-mark volumetric flasks. transfer 0 ml; 5,0 ml; 10,0 ml; 20,0 ml; 30,0 ml respectively of manganese standard solution (4.10.2). Add 25,0 ml of background solution (4.9) to each flask, dilute to volume with water and mix.

NOTE 1 The range of manganese concentrations that can be covered may vary from one instrument to another. Attention should be paid to the minimum performance criteria described in 5.5. For instruments with high sensitivity, the solution of highest concentration shown above may be deleted from the series and an additional solution included at the lower end, e.g. 2,0 ml of manganese standard solution (4.10.2).

- ^{5f/sist}by-Graph-linearity the slope of the calibration graph covering the top 20 % of the concentration range (expressed as a change in absorbance) is not less than 0,7 of the value of the slope for the bottom 20 % of the concentration range determined in the same way.
 - c) Minimum stability --- the standard deviation of the absorbance of the most concentrated calibration solution and that of the zero calibration solution, each being calculated from a sufficient number of repetitive measurements, are less than 1,5 % and 0,5 % respectively of the mean value of the absorbance of the most concentrated calibration solution.
 - NOTES

3 The use of a strip-chart recorder and/or digital readout device is recommended to evaluate criteria a), b) and c) and for all subsequent measurements.

4 Instrument parameters may vary with each instrument. The following parameters were successfully used in several laboratories and they can be used as guidelines. Solutions were aspirated into a dinitrogen oxide-acetylene flame of a premix burner.

Hollow cathode lamp, mA	5
Wavelength, nm	279,5

Dinitrogen oxide flow rate, I/min 7,5

Acetylene flow rate, I/min 4,0

In systems where the values shown above for gas flow rates do not apply, the ratio of the gas flow rates may still be a useful guideline.

6 Sampling and samples

6.1 Laboratory sample

For analysis, use a laboratory sample of minus 100 μ m particle size which has been taken in accordance with ISO 3081 or ISO 3082 and prepared in accordance with ISO 3082 or ISO 3083. In the case of ores with significant contents of combined water or oxidizable compounds, use a particle size of minus 160 μ m.

NOTE 5 A guideline on significant contents of combined water and oxidizable compounds is incorporated in ISO 7764.

Thoroughly mix the laboratory sample and, taking

6.2 **Preparation of predried test samples**

7.4 Blank test and check test

In each run, one blank test and one analysis of a certified reference material of the same type of ore shall be carried out in parallel with the analysis of the ore sample(s) under the same conditions. A predried test sample of the certified reference material shall be prepared as specified in 6.2.

NOTE 7 The certified reference material should be of the same type as the sample to be analysed and the properties of the two materials should be sufficiently similar to ensure that in both cases no significant changes in the analytical procedure will become necessary.

When the analysis is carried out on several samples at the same time, the blank value may be represented by one test, provided that the procedure is the same and the reagents used are from the same reagent bottles.

When the analysis is carried out on several samples of the same type of ore at the same time, the analytical value of one certified reference material may be used.

multiple increments, extract a test sample in such a RD7.5 Determination way that it is representative of the whole contents of the container. Dry the test sample and site 7.5.1 Decomposition of the test portion 105 °C \pm 2 °C, as specified in ISO 7764. (This is the predried test sample.)

7 Procedure

SIST ISO 9682-13f)the decomposition is to be based on fusion, prohttps://standards.iteh.ai/catalog/standards/sist ceed3 asl-specified/-in 87.5.1.1. If it is to be based on 3a7fd467475f/sist-iso-96acid-attack, proceed as specified in 7.5.1.2.

7.1 Number of determinations

Carry out the analysis at least in duplicate in accordance with annex A, independently, on one predried test sample.

NOTE 6 The expression "independently" means that the second and any subsequent result is not affected by the previous result(s). For this particular analytical method, this condition implies that the repetition of the procedure shall be carried out either by the same operator at a different time or by a different operator including, in both cases, appropriate recalibration.

7.2 Safety precautions

WARNING — Follow the manufacturer's instructions for igniting and extinguishing the dinitrogen oxideacetylene flame to avoid possible explosion hazards. Wear tinted safety glasses whenever the burner is in operation.

7.3 Test portion

Taking several increments, weigh, to the nearest 0,0002 g, approximately 0.5 g of the predried test sample obtained in accordance with 6.2.

7.5.1.1 Alkali fusion

Place 0,80 g of sodium carbonate (4.1) in a platinum or suitable platinum alloy crucible (5.1), transfer the test portion (7.3) to the crucible and mix well using a platinum or stainless-steel rod. Add 0,40 g of sodium tetraborate (4.2) and repeat the mixing using the metal rod. Place the crucible in a muffle furnace (5.2), with the temperature maintained at 1 020 °C to 1 040 °C, for 30 min.

Remove the crucible and gently swirl the melt as it solidifies. Allow to cool, then place a PTFE-coated stirring bar (5.4) in the crucible and place the crucible on its side in a 150 ml low-form beaker. Add 20 ml of hydrochloric acid (4.4) and 30 ml of water. Cover and heat while stirring on a magnetic stirrer/hotplate (5.3) until dissolution of the melt is complete.

NOTES

8 The crucible may need to be rolled to ensure complete dissolution of the fused melt.

9 As an alternative to a magnetic stirrer/hotplate, an ultrasonic bath may be used to facilitate dissolution of the fused melt.

Remove the crucible and stirrer and rinse, cool the solution and transfer to a 100 ml one-mark volumetric flask. Dilute to volume with water and mix. (This is the test solution.)

7.5.1.2 Acid attack

Transfer the test portion (7.3) to a 250 ml tall-form beaker. Add 20 ml of hydrochloric acid (4.3), cover with a watch-glass and heat for about 1 h on a hotplate adjusted to give a temperature of 100 $^{\circ}$ C in a test beaker containing a similar volume and depth of sulfuric acid (4.7).

NOTE 10 When the amount of insoluble residue is high, continue heating on a higher temperature zone of the hotplate without boiling the solution.

Add 1 ml of nitric acid (4.5) and 0,2 ml of sulfuric acid (4.8), and digest for 15 min. Remove the watchglass and evaporate the solution to dryness. Heat the salts for 30 min on a hotplate adjusted as before D to give a temperature of 100 °C to 115 °C in sulfuric acid. Add 20 ml of hydrochloric acid (4.4), cover the beaker with a watch-glass and warm for several minutes. Add 30 ml of water and heat to boiling.

Cool, wash the watch-glass hand/sthelawaitsh of the stand beaker, and filter the solution through a medium 475 fish texture filter paper containing filter pulp (0,3 g to 0,4 g dry mass) into a 150 ml low-form beaker. Carefully remove all adhering particles with a rubber-tipped glass rod or piece of moistened filter paper, and transfer to the filter, washing the paper with water until it is visibly free from iron. Reserve the filtrate. Transfer the filter paper and residue to a platinum or suitable platinum alloy crucible (5.1) and start evaporating the filtrate to a final volume of approximately 70 ml.

Dry and ignite the filter paper and residue in the platinum or suitable platinum alloy crucible (5.1) at approximately 800 °C. Cool, add 3 drops of sulfuric acid (4.8) and 2 ml of hydrofluoric acid (4.6) or, in the case of ores of high silica content, 5 ml of hydrofluoric acid. Evaporate slowly to expel silica and continue heating to remove the sulfuric acid. Ignite at approximately 800 °C for several minutes and cool. Add 0,4 g of sodium tetraborate (4.2) and 0,8 g of sodium carbonate (4.1) to the residue, mix and heat at 1000 °C for 15 min in a muffle furnace (5.2) or over a pressurized air burner for a time sufficient to produce a clear melt.

Cool the melt and place the crucible on its side in the beaker containing the reserved filtrate evaporated to approximately 70 ml. Heat the solution until the fused salts are completely dissolved from the crucible. Remove and rinse the crucible, cool the solution and transfer to a 100 ml one-mark volumetric flask. Dilute to volume with water and mix. (This is the test solution.)

7.5.2 Treatment of the test solution

If the manganese content in the test sample is between 0.01% (m/m) and 0.3% (m/m), use the test solution, obtained in either 7.5.1.1 or 7.5.1.2, directly without dilution. For manganese contents between 0.2% (m/m) and 3% (m/m), transfer an appropriate aliquot of the test solution (see table 1) to a 100 ml one-mark volumetric flask, add the amount of back-

Cool, wash the watch-glass hand sthe wallsh of the solution solution (4.9), indicated in table 1, dilute to beaker, and filter the solution through a medium 475f sist solution water and mix. (This is the diluted test to dure filter paper containing filter pulp (0.2 g to solution.)

If a diluted test solution has been prepared, prepare a corresponding diluted blank test solution as follows.

Transfer a corresponding amount of the blank test solution, and the amount of background solution (4.9) specified in table 1, to a 100 ml one-mark volumetric flask. Dilute to volume with water and mix. (This is the diluted blank test solution.)

Expected content, w _{Mn} , of Mn in sample	Aliquot	Volume of background solution (4.9) to be added
% (m/m)	ml	ml
0,01 to 0,3	No dilution	0
0,2 to 0,6	50,0	12,5
0,5 to 1,5	20,0	20
1,0 to 3	10,0	22,5
0,2 to 0,6	50,0	12,5
0,5 to 1,5	20,0	20
1,0 to 3	10,0	22,5

Table 1 — Dilution guide for test and blank test solutions

NOTE – The overlap in concentration ranges allows the content obtained to differ slightly from the content expected. When in doubt, use the first range specified.

7.5.3 Adjustment of the atomic absorption spectrometer

Set the wavelength of manganese (279.5 nm) to obtain minimum absorbance and adjust the readout to zero absorbance. Light the dinitrogen oxideacetylene flame, observing the safety precautions of 7.2. After 10 min of preheating the burner, aspirate the calibration solution of highest manganese content (4.11) and carefully adjust the fuel flow and burner position to obtain maximum absorbance.

Check that the conditions for zero absorbance were maintained while aspirating water and again aspirate the calibration solution of highest manganese content to establish that the absorbance reading is not drifting. Set the reading for water to zero absorbance.

7.5.4 Atomic absorption measurements

Aspirate the calibration solutions (4.11) and the test or diluted test solution in order of increasing concentration, starting with the zero calibration solution and the blank test or diluted blank test solution. When stable responses have been obtained, record RD the readings in absorbance units. Aspirate water RD between each solution and repeat the measure-ments twice ments twice.

NOTE 13 If any curvature obtained approaches the limit specified in 5.5 b), replot the graph using uncorrected values for all solutions and establish the concentration of the zero calibration solution from the intercept of the graph on the negative side of the concentration axis. Add this value to the nominal concentration values of the calibration solutions and replot the graph to pass through the origin. Determine from the graph the concentration of manganese, in micrograms per millilitre, in the blank test and test solutions respectively, and correct the concentration of the test solution or diluted test solution with the concentration of the blank test or diluted blank test as appropriate.

Expression of results 8

8.1 Calculation of manganese content

The manganese content, w_{Mn} , as a percentage by mass, is calculated to five decimal places for manganese contents below 0,5 % (m/m) and to four decimal places for manganese contents between 0,5 % (m/m) and 3 % (m/m), from the equation

$$w_{\rm Mn} = \frac{\rho_{\rm Mn}}{m \times 100} \times \rm DF \qquad \dots (1)$$

NOTES

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where

 ρ_{Mn}

11 The reagents used for the apreparation of the start don's the start of the start calibration solution should have a low enough in an gane se-iso-9682-1-2000 content for the absorbance value not to significantly exceed 0,01 with high performance instruments (absorbance for the highest calibration solution above 0,6) and 0,005 with instruments of average to low performance (approaching the criterion in 5.5).

12 If, with an instrument of high sensitivity, the solution of highest manganese concentration has been deleted from the calibration series (see 4.11), it may also be necessary to change the dilution guide in table 1 (7.5.2). In this case, move the values for the volumes of aliquots and background solution to the line above for each range of manganese content, using for the range of 1 % (m/m)to 3 % (m/m) Mn a 5,0 ml aliquot with 23,8 ml of background solution.

Correct the absorbance values obtained for the calibration solutions by subtracting the absorbance of the zero calibration solution and prepare a calibration graph by plotting the net absorbance values against micrograms of manganese per millilitre. If the graph is substantially linear, subtract the absorbance obtained for the blank test or diluted blank test from the absorbance obtained for the test solution or diluted test solution as appropriate and, using the graph, convert the net absorbance value of the test solution or diluted test solution to micrograms of manganese per millilitre.

is the concentration, in micrograms per millilitre, of manganese in the test solution or diluted test solution;

is the mass, in grams, of the test portion; DF is the dilution factor calculated from

100/V, when a dilution has been made using V ml of test solution. When no dilution has been made, DF = 1.

8.2 General treatment of results

8.2.1 Repeatability and permissible tolerance

The precision of this analytical method is expressed by the following regression equations¹⁾:

for alkali fusion

$r = 0,0^{\circ}$	16 8 X + 0,004 8	 (2)

- $P = 0,031 \ 8 \ X + 0,006 \ 2$...(3)
- $\sigma_r = 0,005 \ 9 \ X + 0,001 \ 7$...(4)

$$\sigma_{\rm L} = 0,010 \ 4 \ X + 0,001 \ 9 \qquad \dots (5)$$

for acid attack

$$r = 0.019 \ 0 \ X + 0.004 \ 0 \qquad \dots$$
(6)
 $P = 0.074 \ 5 \ X - 0.007 \ 5$

¹⁾ Additional information is given in annexes B and C.