

SLOVENSKI STANDARD SIST ISO 9556:1997

01-maj-1997

Jeklo in železo - Ugotavljanje celotnega deleža ogljika - Metoda infrardeče absorpcije po zgorevanju v indukcijski peči

Steel and iron -- Determination of total carbon content -- Infrared absorption method after combustion in an induction furnace

iTeh STANDARD PREVIEW

Aciers et fontes -- Dosage du carbone total -- Méthode par absorption dans l'infrarouge après combustion dans un four à induction

SIST ISO 9556:1997

Ta slovenski standard je istoveten z: 1989

ICS:

71.080.01	Organske kemikalije na splošno	Organic chemicals in general
77.080.01	Železne kovine na splošno	Ferrous metals in general

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INTERNATIONAL STANDARD

ISO 9556

First edition 1989-07-15

Corrected and reprinted 1989-11-15

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Reference number ISO 9556 : 1989 (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 approval before their acceptance as International Standards by the ISO Council. They are approved in accordance with ISO procedures requiring at IEW least 75 % approval by the member bodies voting.

standards.iteh.ai) International Standard ISO 9556 was prepared by Technical Committee ISO/TC 17, Steel. SIST ISO 9556:1997

https://standards.iteh.ai/catalog/standards/sist/25d846c0-675d-47ee-8883-Annexes A, B and C of this International Standard are for information only 56-1997

Steel and iron — Determination of total carbon content — Infrared absorption method after combustion in an induction furnace

1 Scope

This International Standard specifies an infrared absorption method after combustion in an induction furnace for the determination of the total carbon content in steel and iron.

The method is applicable to carbon contents between 0,003 % (m/m) and 4.5 % (m/m).

Reagents

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

4.1 Water, free from carbon dioxide.

Boil water for 30 min, cool to room temperature and bubble with oxygen (4.2) for 15 min. Prepare just before use.

2 Normative references, (Teh STANDAR 4,2 Poxygen, 99,5,% (m/m) minimum.

The following standards contain provisions which, through An oxidation catalyst [copper(II) oxide or platinum] tube heated reference in this text, constitute provisions of this International US to a temperature above 450 °C must be used prior to a purify-Standard. At the time of publication, the editions indicated ing unit (see annex C), when the presence of organic conwere valid. All standards are subject to revision, and parties to 955(taminants is suspected in the oxygen. agreements based on this International Standard are encouraged to investigate the possibility of applying the most recent sist-is:4:356Pure7iron, of known low carbon content less than editions of the standards listed below. Members of IEC and ISO maintain registers of currently valid International Standards.

ISO 377: 1985, Wrought steel - Selection and preparation of samples and test pieces.

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 5725: 1986, Precision of test methods - Determination of repeatability and reproducibility for a standard test method by inter-laboratory tests.

Principle 3

Combustion of a test portion with accelerator at a high temperature in a high-frequency induction furnace in a current of pure oxygen. Transformation of carbon into carbon dioxide and/or carbon monoxide.

Measurement by infrared absorption of the carbon dioxide and/or carbon monoxide carried by a current of oxygen.

0,0010% (m/m).

4.4 Suitable solvent, appropriate for washing greasy or dirty test samples, for example, acetone.

4.5 Magnesium perchlorate [Mg(ClO₄)₂], particle size: from 0,7 mm to 1,2 mm.

4.6 **Barium carbonate**

Dry barium carbonate (minimum assay: 99,5 %) at 105 °C to 110 °C for 3 h and cool in a desiccator before use.

4.7 Sodium carbonate

Dry anhydrous sodium carbonate (minimum assay: 99,9 %) at 285 °C for 2 h and cool in a desiccator before use.

4.8 Accelerator: copper, tungsten-tin mixture or tungsten of known low carbon content less than 0,001 0 % (m/m).

4.9 Sucrose, standard solution, corresponding to 25 g of C per litre.

Weigh, to the nearest 1 mg, 14,843 g of sucrose (C12H22O11) (analytical standards grade) previously dried at 100 °C to 105 °C for 2,5 h and cooled in a desiccator.

Dissolve in about 100 ml of water (4.1), transfer to a 250 ml one-mark volumetric flask quantitatively, dilute to the mark with water (4.1) and mix.

1 ml of this standard solution contains 25 mg of C.

4.10 Sodium carbonate, standard solution, corresponding to 25 g of C per litre.

Weigh, to the nearest 1 mg, 55,152 g of sodium carbonate (4.7), dissolve in about 200 ml of water (4.1), transfer to a 250 ml one-mark volumetric flask quantitatively, dilute to the mark with water (4.1) and mix.

1 ml of this standard solution contains 25 mg of C.

4.11 Inert ceramic (attapulgus clay) impregnated with sodium hydroxide, particle size: from 0,7 mm to 1,2 mm.

Apparatus 5

During the analysis, unless otherwise stated, use only ordinary laboratory apparatus.

All volumetric glassware shall be class AC in accordance with ISO 385-1, ISO 648 or ISO 1042 as appropriate.

standa The apparatus required for combustion in a high-frequency induction furnace and the subsequent infrared absorption samples to be analysed prior to setting up for analysis. measurement of the evolved carbon dioxide and/or carbon /sist/25d846c0-675d-47ee-8883monoxide may be obtained commercially from it abai mote of tandards Flush oxygen through the apparatus and adjust the instrument manufacturers. Follow the manufacturer's instructions for the 00a/six controls to give a zero reading. operation of the instrument.

Features of commercial instruments are given in annex C.

5.1 Micropipette, 100 µl, limit of error shall be less than 1 μl.

5.2 Tin capsule, about 6 mm in diameter, 18 mm in height, 0,3 g in mass and approximately 0,4 ml in volume, of known low carbon content less than 0,001 0 % (m/m).

Ceramic crucible, capable of withstanding combustion 5.3 in an induction furnace.

Ignite crucibles in an electric furnace in air or in a current of oxygen for not less than 2 h at 1 100 °C and store in a desiccator before use.

NOTE - For the determination of low carbon contents it is advisable to ignite crucibles at 1 350 °C in a current of oxygen.

6 Sampling

Carry out sampling in accordance with ISO 377 or appropriate national standards for iron.

Procedure

SAFETY INSTRUCTIONS - The risks related to combustion analysis are mainly burns in pre-igniting the ceramic crucibles and in the fusions. Use crucible tongs at all times and suitable containers for the used crucibles. Normal precautions for handling oxygen cylinders shall be taken. Oxygen from the combustion process shall be removed effectively from the apparatus since a high concentration of oxygen in a confined space can present a fire hazard.

7.1 General operating instructions

Purify the oxygen supply using tubes packed with the inert ceramic (attapulgus clay) impregnated with sodium hydroxide (4.11) and magnesium perchlorate (4.5), and maintain a quiescent flow rate whilst on standby. Maintain a glass wool filter or a stainless steel net as a dust collector. Clean and change as necessary. The furnace chamber, pedestal post and filter trap shall be cleaned frequently to remove oxide build-up.

Allow each item of equipment to stabilize for the time recommended by the equipment manufacturers when the main supply is switched on after being out of action for any length of

time. REVIEW

After cleaning the furnace chamber and/or changing filters or after the equipment has been inoperative for a period, stabilize the apparatus by burning several samples of similar type to the

If the instrument used provides a direct reading in percentage of carbon, adjust the instrument reading for each calibration range as follows.

Select the certified reference material with a carbon content close to the maximum carbon content in the calibration series, measure the carbon content of the certified reference material in the manner specified in 7.4.

Adjust the reading of the instrument to the certified value.

NOTE - This adjustment shall be made before the calibration as specified in 7.5. It cannot replace or correct the calibration.

7.2 Test portion

Degrease the test sample by washing in a suitable solvent (4.4). Evaporate the last traces of the washing liquid by heating.

Weigh, to the nearest 1 mg, approximately 1 g of the test sample for carbon contents less than 1.0 % (m/m) and approximately 0,5 g for carbon contents greater than 1,0 % (m/m).

NOTE - The mass of the test portion may be dependent on the type of instrument used.

7.3 Blank test

Prior to the determination, carry out the following blank tests in duplicate.

Transfer the tin capsule (5.2) (see note 1) to the ceramic crucible (5.3), press the tin capsule lightly against the bottom of the crucible. Add the same quantity of pure iron (4.3) as that of the test portion (7.2), and the same quantity of the accelerators (4.8) (see note 2) as will be added to the test portion.

Treat the crucible and contents as specified in the second and third paragraphs of 7.4.

Obtain the reading of the blank tests and convert it to milligrams of carbon by means of the calibration graph (7.5).

The blank value is obtained by subtracting the mass of carbon in the pure iron used (4.3) from the mass of carbon in the blank tests.

The mean blank value (\overline{m}_1) is calculated from the two blank values (see note 3).

NOTES

1 In cases where the calibration graphs of 7.5.1 or 7.5.2 are applied, use the following prepared capsule.

Prepare the tin capsule; using the micropipette (5.1), transfer 100 µl of water (4.1) to a tin capsule (5.2) and dry at 90 °C for 2 h.

2 The quantity of accelerators will depend on the individual characteristics of the instrument and the type of material being analysed. The amount used shall be sufficient for complete combustion.

3 The mean blank value and the difference between the two blank ands/siportion. values shall both not exceed 0,01 mg of carbon. If these values are abnormally high, investigate and eliminate the source of contamination.

7.4 Determination

Transfer one tin capsule (5.2) to the ceramic crucible (5.3), press the tin capsule lightly against the bottom of the ceramic crucible, add the test portion (7.2) and cover with the appropriate mass of the accelerators (4.8) (see note 2 of 7.3).

Place the ceramic crucible and contents on the pedestal post, raise to the combustion position and lock the system. Operate the furnace in accordance with the manufacturer's instructions.

At the end of the combustion and measuring cycle, remove and discard the crucible, and record the analyser reading.

7.5 Establishment of the calibration graph

7.5.1 Samples having carbon contents between 0,003 % (m/m) and 0,01 % (m/m)

7.5.1.1 Preparation of the calibration series

Transfer the volumes of sucrose standard solution (4.9) or sodium carbonate standard solution (4.10) indicated in table 1 to five 250 ml one-mark volumetric flasks. Dilute to the mark with water (4.1) and mix.

Using the micropipette (5.1), transfer 100 μl of each of the diluted solutions to five tin capsules (5.2) and dry at 90 oC for 2 h.

Cool to room temperature in a desiccator.

Table	1
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Volume of the standard solution [(4.9) or (4.10)] ml	Mass of carbon in the diluted solution, per millilitre mg	Mass of carbon taken in the tin capsule mg	Carbon content in the test portion % (<i>m</i> / <i>m</i>)
0*)	0	0	0
1,0	0,10	0,010	0,001
2,0	0,20	0,020	0,002
5,0	0,50	0,050	0,005
10,0	1,00	0,100	0.010

7.5.1.2 Measurements

Transfer the tin capsule containing sucrose or sodium carbonate to the ceramic crucible (5.3), press the tin capsule lightly against the bottom of the ceramic crucible, add 1,000 g of pure iron (4.3) and cover with the same quantity of the accelerators (4.8) (see note 2 of 7.3) as will be added to the test

Treat the crucible and contents as specified in the second and third paragraphs of 7.4.

7.5.1.3 Plotting the calibration graph

Obtain the net reading by subtracting the reading of the zero member from that of each member of the calibration series.

Prepare a calibration graph by plotting the net reading against milligrams of carbon for each member of the calibration series.

7.5.2 Samples having carbon contents between 0,01 % (m/m) and 0,1 % (m/m)

7.5.2.1 Preparation of the calibration series

Transfer the volumes of sucrose standard solution (4.9) or sodium carbonate standard solution (4.10) indicated in table 2 to five 50 ml one-mark volumetric flasks. Dilute to the mark with water (4.1) and mix.

Using the micropipette (5.1), transfer 100 μl of each of the diluted solutions to five tin capsules (5.2) and dry at 90 °C for 2 h.

Cool to room temperature in a desiccator.

Volume of the standard solution [(4.9) or (4.10)]	Mass of carbon in the diluted solution, per millilitre	Mass of carbon taken in the tin capsule	Carbon content in the test portion
ml	mg	mg	% (<i>m/m</i>)
0*)	0	0	0
2,0	1,0	0,10	0,010
4,0	2,0	0,20	0,020
10,0	5,0	0,50	0,050
20,0	10,0	1,00	0,100
*) Zero member			

Table 2

7.5.2.2 Measurements

As specified in 7.5.1.2.

7.5.2.3 Plotting the calibration graph

As specified in 7.5.1.3.

7.5.3 Samples having carbon contents between 0,1 % (m/m) and 1,0 % (m/m)

7.5.3.1 Preparation of the calibration series

Weigh, to the nearest 0,1 mg, the masses of barium carbonate (4.6) or sodium carbonate (4.7) indicated in table 3 and transfer to five tin capsules (5.2).

Table 3

Mass of the reference material mg		Mass of carbon taken	(Carbonse5) content in
Barium carbonate (4.6)	Sodium carbonate (4.7)	in the tin capsule mg	the test portion % (m/m)
0*)	0*)	0	0
16,4	8,8	1,0	0,10
32,9	17,7	2,0	0,20
82,1	44,1	5,0	0,50
164,3 88,2		10,0	1,00

7.5.3.2 Measurements

Transfer the tin capsule containing barium carbonate or sodium carbonate to the ceramic crucible (5.3), press the tin capsule lightly against the bottom of the ceramic crucible, add 1,000 g of pure iron (4.3) and cover with the same quantity of the accelerators (4.8) (see note 2 of 7.3) as will be added to the test portion.

Treat the crucible and contents as specified in the second and third paragraphs of 7.4.

7.5.3.3 Plotting the calibration graph

As specified in 7.5.1.3.

7.5.4 Samples having carbon contents between 1,0 % (m/m) and 4,5 % (m/m)

7.5.4.1 Preparation of the calibration series

Weigh, to the nearest 0,1 mg, the masses of barium carbonate (4.6) or sodium carbonate (4.7) indicated in table 4 and transfer to five tin capsules (5.2).

 $\mathsf{NOTE}-\mathsf{If}$ the weighed barium carbonate cannot be transferred to the tin capsule, it may be placed directly on the bottom of the ceramic crucible.

Table	4
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	erence material	Mass of carbon taken	Carbon content in	
Barium carbonate (4.6)	Sodium carbonate (4.7)	in the tin capsule mg	the test portion % (<i>m/m</i>)	
0*)	0*)	0	0	
82,1	44,1	5,0	1,0	
164,3	88,2	10,0	2,0	
246,4	132,3	15,0	3,0	
369,7 198,6		22,5	4,5	
*) Zero member				

7.5.4.2 Measurements

Transfer the tin capsule containing barium carbonate or sodium carbonate to the ceramic crucible (5.3), press the tin capsule lightly against the bottom of the ceramic crucible, add 0,500 g of pure iron (4,3) and cover with the same quantity of the accelerators (4,8) (see note 2 of 7.3) as will be added to the test portion.

SIST ISO 9576at the crucible and contents as specified in the second and ards.iteh.ai/catalog/standardthird/paragraphs/off/2-47ee-8883-

0a/sist-iso-9556-1997 7.5.4.3 Plotting the calibration graph

As specified in 7.5.1.3.

8 Expression of results

8.1 Method of calculation

Convert the analyser reading of the test portion to milligrams of carbon (m_0) by means of the calibration graph (7.5).

The carbon content, expressed as a percentage by mass, $w_{\rm c}$ (%), is given by the equation:

$$\frac{(m_0 - \bar{m}_1)}{m \times 103} \times 100$$
$$\frac{(m_0 - \bar{m}_1)}{10m}$$

where

=

 m_0 is the mass, expressed in milligrams of carbon, in the test portion;

 \bar{m}_1 is the mass, expressed in milligrams of carbon, in the blank test (7.3);

m is the mass, in grams, of the test portion (7.2).

8.2 Precision

A planned trial of this method was carried out by 22 laboratories, at 12 levels of carbon, each laboratory making three determinations of carbon content at each level (see notes 1 and 2).

The test samples used and mean results obtained are listed in table A.1.

The results obtained were treated statistically in accordance with ISO 5725.

The data obtained showed a logarithmic relationship between carbon content and repeatability (r) and reproducibility (R and R_w) of the test results (see note 3) as summarized in table 5. The graphical representation of the figures is given in annex B.

				The test report shall include the following information:
Carbon content % (m/m)	Repeatability	Reprod <i>R</i>	ucibility	a) all information necessary for the identification of the sample, the laboratory and the date of the analysis:
70 (111/111)	/	ĸ	R_{w}	
0,003	0,000 53	0,001 19	0,000 77	
0,005	0,000 69	0.001 60	0.001 02	b) the method used, by reference to this International
0,01	0,000 99	0,002 40	0,001 50	Standard;
0,02	0,001 42	0,003 59	0,002 20	
0,05	0,002 29	0,006 12 0	0,003 65	D c) the results, and the form in which they are expressed;
0,1	0,003 29	0,009 17 🔍	0,005 36	
0,2	0,004 72	0,013 7	0,007 85	d any-unusual features noted during the determination;
0,5	0,007 62	0,023 4 【	502,013021	ds.iten.al)
1,0	0,011 0	0,035 1	0,019 1	e) any operation not specified in this International Stan-
2,0	0,015 7	0,052 6	0,028 0	ossectordard, or any optional operation which may have influenced
4,5	0,024 0	0,084 4	0,043 8T ISC	19110-1997
	http	na.//standarda ita	h allaata la alstand	the results.

Table 5

NOTES

1 Two of the three determinations were carried out under repeatability conditions as defined in ISO 5725, i.e. one operator, same apparatus, identical operating conditions, same calibration, and a minimum period of time.

2 The third determination was carried out at a different time (on a different day) by the same operator as in note 1 using the same apparatus with a new calibration.

3 From the two values obtained on day 1 the repeatability (r) and reproducibility (R) were calculated using the procedure specified in ISO 5725. From the first value obtained on day 1 and the value obtained on day 2, the within-laboratory reproducibility (R_w) was calculated.

9 Test report

The test report shall include the following information:

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