
**Unalloyed steel — Determination of low
carbon content —**

Part 3:

Infrared absorption method after combustion
in an electric resistance furnace (with
preheating)

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Acier non allié — Détermination des faibles teneurs en carbone —

*Partie 3: Méthode par absorption dans l'infrarouge après combustion dans
un four électrique à résistances (avec préchauffage)*

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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.

The main task of technical committees is to prepare International Standards. In exceptional circumstances a technical committee may propose the publication of a Technical Report of one of the following types:

- type 1, when the required support cannot be obtained for the publication of an International Standard, despite repeated efforts;
- type 2, when the subject is still under technical development or where for any other reason there is the future but not immediate possibility of an agreement on an International Standard;
- type 3, when a technical committee has collected data of a different kind from that which is normally published as an International Standard ("state of the art", for example).

Technical Reports of types 1 and 2 are subject to review within three years of publication, to decide whether they can be transformed into International Standards. Technical Reports of type 3 do not necessarily have to be reviewed until the data they provide are considered to be no longer valid or useful.

ISO/TR 15349-3, which is a Technical Report of type 2, was prepared by Technical Committee ISO/TC 17, *Steel*, Subcommittee SC 1, *Methods of determination of chemical composition*.

This document is being issued in the Technical Report (type 2) series of publications (according to subclause G.3.2.2 of part 1 of the ISO/IEC Directives, 1995) as a "prospective standard for provisional application" in the field of determination of carbon content in steel because there is an urgent need for guidance on how standards in this field should be used to meet an identified need.

This document is not to be regarded as an "International Standard". It is proposed for provisional application so that information and experience of its use in practice may be gathered. Comments on the content of those documents should be sent to the ISO Central Secretariat.

A review of this Technical Report (type 2) will be carried out not later than three years after its publication with the options of: extension for another three years; conversion into an International Standard; or withdrawal.

ISO/TR 15349 consists of the following parts, under the general title *Unalloyed steel — Determination of low carbon content*:

- *Part 1: Infrared absorption method after combustion in an electric resistance furnace (by peak separation)*
- *Part 2: Infrared absorption method after combustion in an induction furnace (with preheating)*
- *Part 3: Infrared absorption method after combustion in an electric resistance furnace (with preheating)*

Annex A of this part of ISO/TR 15349 is for information only.

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Unalloyed steel — Determination of low carbon content —

Part 3:

Infrared absorption method after combustion in an electric resistance furnace (with preheating)

1 Scope

This part of ISO 15349 describes an infrared absorption method after preheating and combustion in an electric resistance furnace for the determination of the low carbon content in unalloyed steel.

The method is applicable to carbon contents between 0,000 3 % (m/m) and 0,010 % (m/m).

2 Normative references

The following standards contain provisions which, through references in this text, constitute provisions of this part of ISO 15349. 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 15349 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 1042:1998, *Laboratory glassware — One-mark volumetric flasks*.

ISO 3696:1987, *Water for analytical laboratory use — Specification and test methods*.

ISO 14284:1996, *Steel and iron — Sampling and preparation of samples for the determination of chemical composition*.

ISO/TR 15349-1:1998, *Unalloyed steel — Determination of low carbon content — Part 1: Infrared absorption method after combustion in an electric resistance furnace (by peak separation)*.

ISO 15349-2:—¹⁾, *Unalloyed steel — Determination of low carbon content — Part 2: Infrared absorption method after combustion in an induction furnace (with preheating)*.

3 Principle

Preheating of a test portion at low temperature and combustion of the test portion with accelerator at a high temperature in an electric resistance furnace in a current of pure oxygen. Transformation of carbon into carbon dioxide and/or carbon monoxide.

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

¹⁾ To be published.

4 Reagents

During the analysis, unless otherwise stated, use only reagents of recognized analytical grade and only grade 3 water as specified in ISO 3696.

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.

4.2 Oxygen, 99,95 % (m/m) minimum.

An oxidation catalyst [copper(II)oxide or platinum] tube heated to a temperature above 450 °C must be used prior to a purifying unit, when the presence in the oxygen of organic contaminants is suspected.

4.3 Suitable solvent, appropriate for washing greasy or dirty test samples; e.g. acetone.

4.4 Accelerator, copper plate (see note 1) or granular tin (see note 2) of known very low carbon content less than 0,000 1 % (m/m).

NOTE 1 Copper plate (about 0,1 g/plate) should be used after the following treatment: heat the copper plate at 450 °C to 600 °C for 10 min in a current of oxygen and cool in a desiccator without grease. This treatment shall be carried out just before use.

NOTE 2 Granular tin should be used after the following treatment: rinse the granular tin in hydrochloric acid (ρ about 1,19 g/ml, diluted 1 + 1) for 5 min using an ultrasonic wave washer, wash it thoroughly with water and dry. Stock it in a clean glass bottle with ground-in stopper.

4.5 Standard substances

4.5.1 Sucrose, standard solution

Weigh, to the nearest 0,1 mg, the masses of sucrose, indicated in table 1, (analytical standard grade) previously dried at 100 °C to 105 °C for 2,5 h then cooled in a desiccator and transfer to seven 100 ml beakers.

Add 30 ml of water (4.1) to dissolve, transfer to seven 100 ml one-mark volumetric flasks quantitatively, dilute to the mark with water (4.1) and mix.

Table 1 — Standard solution series of sucrose

Standard solution reference number	Mass of sucrose g	Corresponding mass of carbon added μg	Carbon content in 1 g of the test portion % (m/m)
1	0 ¹⁾	0	0
2	0,010 0	4,21	0,000 42
3	0,025 0	10,53	0,001 05
4	0,060 0	25,26	0,002 53
5	0,120 0	50,53	0,005 05
6	0,180 0	75,79	0,007 58
7	0,240 0	101,1	0,010 11

1) Zero member.

4.5.2 Calcium carbonate

Dry calcium carbonate [minimum assay 99,9 % (m/m)] at 180 °C for 1 h and cool in a desiccator before use.

4.6 Magnesium perchlorate [$\text{Mg}(\text{ClO}_4)_2$], particle size: from 0,7 mm to 1,2 mm.

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

5 Apparatus

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

All laboratory glassware shall be class A, in accordance with ISO 1042 as appropriate.

The apparatus required for combustion in an electric resistance furnace and subsequent infrared absorption measurement of the evolved carbon dioxide and/or carbon monoxide from steel may be obtained commercially, for example, Trace carbon analyser Model EMIA — U610, HORIBA, LTD²⁾.

Follow the manufacturer's instructions for the operation of the instrument.

5.1 Ceramic boat, 80 mm × 13,5 mm × 10 mm

Ignite a ceramic boat in an electric resistance furnace in a current of oxygen for not less than 2 h at 1 200 °C and store in a desiccator without grease.

Just before use, ignite the ceramic boat at 1 300 °C for several minutes and use the hot boat.

5.2 Quartz boat, 30 mm × 9 mm × 8 mm

Rinse a quartz boat in the washing solution [mix nine volumes of nitric acid (ρ about 1,40 g/ml, diluted 1 + 6) and one volume of hydrogen peroxide (30 % V/V)] for 2 h to 3 h, wash it thoroughly with water and dry. Stock in a desiccator without grease.

Just before use, place the quartz boat on the ceramic boat, ignite it at 1 200 °C for 10 min and cool in a desiccator without grease.

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

6 Sampling and preparation of the test samples

Sampling and preparation of the samples shall be carried out in accordance with ISO 14284.

Prepare the chip size of test sample between 0,75 mm and 2,0 mm.

7 Procedure

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

²⁾ EMIA — U610, HORIBA is the tradename of a product supplied by HORIBA LTD. This information is given for the convenience of users of this part of ISO/TR 15349 and does not constitute an endorsement by ISO of the product named. Equivalent products may be used if they can be shown to lead to the same results.

7.1 Preparation of the apparatus

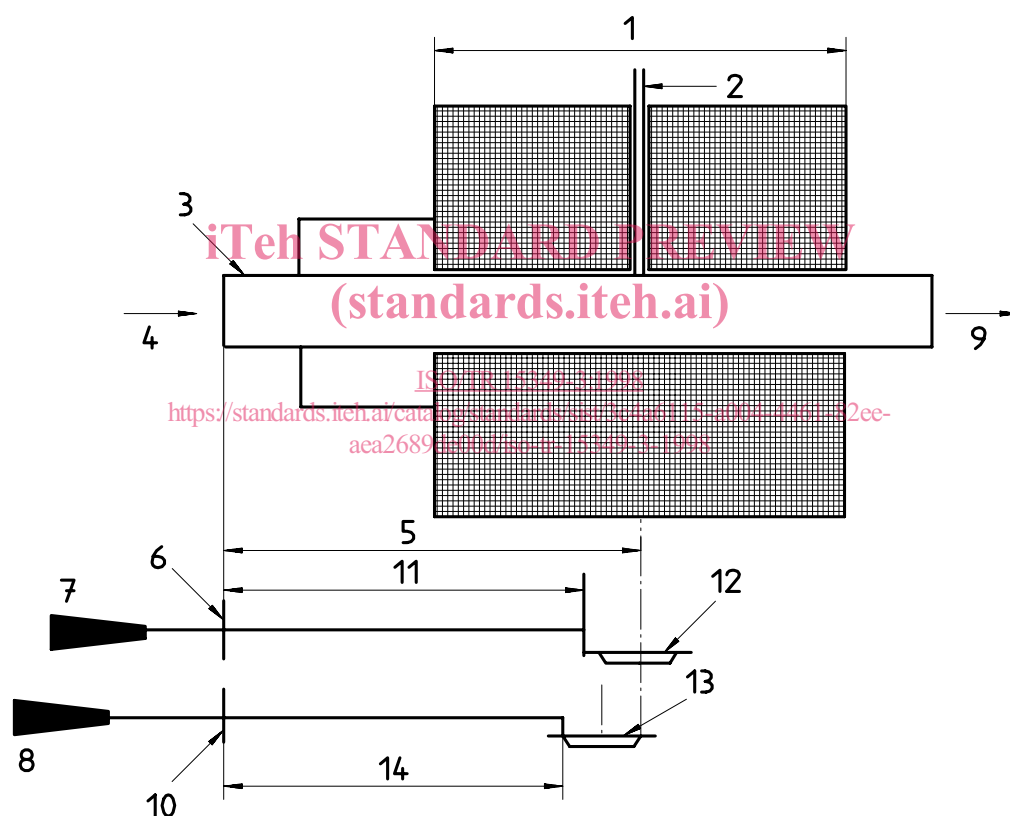
Raise the temperature of the combustion furnace to 1 300 °C.

After having verified the highest temperature (more than 1 200 °C) at the centre of the heating zone in the combustion tube by use of a thermocouple, measure the distance, L , from the inlet of combustion tube to the centre of the heating zone in millimetres. Prepare two rigid nickel bars. Fix the stopper of a rigid nickel bar, A at the point where $L = (\text{the length of the ceramic boat used})/2$ (mm) distance from the bar end. This is used to set the ceramic boat containing the test portion or calcium carbonate (see note below and figure 1).

NOTE 1 e.g., for model EMIA — U610, L is 290 mm and boat length is 80 mm. In this case the stopper is 250 mm distant from the bar end.

Fix the stopper of another rigid nickel bar, B at the point where $L = \text{the length of the ceramic boat used (mm)}$ distance from the bar end. This is used to set the ceramic boat containing sucrose (see note below and figure 1).

NOTE 2 e.g., for model EMIA — U610, L is 290 mm and boat length is 80 mm. In this case the stopper is 210 mm distant from the bar end.



Key:

- | | |
|----------------------|---|
| 1 Heating zone | 8 Rigid nickel bar B |
| 2 Thermocouple | 9 Evolved gases carried by oxygen to measuring system |
| 3 Combustion tube | 10 Stopper B |
| 4 Oxygen | 11 $L - (\text{length of ceramic boat}) / 2$ mm |
| 5 L mm | 12 Ceramic boat (test portion or calcium carbonate) |
| 6 Stopper A | 13 Ceramic boat containing quartz boat (sucrose) |
| 7 Rigid nickel bar A | 14 $L - (\text{length of ceramic boat})$ mm |

Figure 1 — Combustion furnace

Purify the oxygen supply using tubes packed with the inert ceramic (4.7) (attapulugus clay) impregnated with sodium hydroxide and magnesium perchlorate (4.6), and maintain a quiescent flowrate whilst on standby. Maintain a glass wool filter. Clean and charge as necessary.

When the main supply is switched on after being out of action for any length of time, allow each item of equipment to stabilize for the time recommended by the equipment manufacturers.

After changing filters or after the equipment has been inoperative for a period, stabilize the apparatus by burning several samples of similar type to the samples to be analysed prior to setting up for analysis.

Flush oxygen through the apparatus and adjust the instrument controls to give a zero reading.

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.

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

7.2 Test portion

Degrease the test sample by washing in a suitable solvent (4.3). Evaporate the last traces of the washing liquid by heating.

Weigh, to the nearest 0,1 mg, approximately 1,0 g of the test sample.

7.3 Blank test

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

Add the 0,50 g of the accelerators (4.4) to the ceramic boat.

Treat the boat and contents as described in the second and third paragraph of 7.4.

Obtain the reading of the blank tests and convert it to micrograms, to the nearest 0,1 μg , of carbon by means of the calibration graph (see 7.5).

The mean blank value is calculated from the two blank values to the nearest 0,1 μg .

The mean blank value and the difference between the two blank values shall both not exceed 1,0 μg of carbon. If these values are abnormally high, investigate and eliminate the source of contamination.

7.4 Determination

7.4.1 Pre-treatment of the test portion

Transfer the test portion (7.1) to the middle zone of ceramic boat (5.1).

Place the ceramic boat containing the test portion in the muffle or wire-wound furnace (5.6) heated to $420\text{ }^{\circ}\text{C} \pm 10\text{ }^{\circ}\text{C}$ for 5 min to 10 min.

Remove the ceramic boat containing the test portion from the muffle or wire-wound furnace and immediately add the appropriate mass of the accelerator (4.4) to the ceramic boat containing the test portion.

7.4.2 Combustion of the test portion

Immediately after pre-treatment place the ceramic boat and contents in the inlet of the combustion tube, using the rigid nickel bar A, insert the boat until it is stopped and lock the system. (In this case the centre of boat is at the centre of heating zone — see figure 1). Operate the instrument in accordance with the manufacturer's instructions.