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Second edition 1996-02-15

Solid mineral fuels — Determination of carbon and hydrogen — Liebig method

iTeh STANDARD PREVIEW Combustibles minéraux solides — Dosage du carbone et de J'hydrogène — Méthode de Liebig

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ICA



Reference number ISO 625:1996(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 bodies casting. VIE W a vote.

(standards.iteh.ai) International Standard ISO 625 was prepared by Technical Committee ISO/TC 27, Solid mineral fuels, Subcommittee SC 5, Methods of analysis.

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This second edition cancels and replaces the first edition (ISO 625:1975), which has been technically revised.

Annex A of this International Standard is for information only.

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International Organization for Standardization

Introduction

An alternative method to that specified in this International Standard is given in ISO 609:1996, *Solid mineral fuels* — *Determination of carbon and hydrogen* — *High temperature combustion method.*

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Solid mineral fuels — Determination of carbon and hydrogen — Liebig method

1 Scope

This International Standard specifies a method of determining the total carbon and the total hydrogen in hard coal, brown coal and lignite, and coke, by the Liebig method. ISO 1015:1992, Brown coals and lignites — Determination of moisture content — Direct volumetric method.

ISO 1170:1977, Coal and coke — Calculation of analyses to different bases.

The results include the carbon in the carbonates and RD ISO 1988:1975, Hard coal — Sampling, the hydrogen combined in the moisture and in the water of constitution of silicates. A determination of SIISO 2309 1980, Coke — Sampling, moisture is carried out at the same time, and an ap-

propriate correction is applied to the hydrogen <u>Ivalue</u>5:1996 nation of moisture content — Indirect gravimetric dioxide may also be made and the total carbon value/iso-625-1996

corrected for the presence of mineral carbonates.

2 Normative references

The following standards contain provisions which, through reference in this text, constitute provisions of this International Standard. At the time of publication, the editions indicated were valid. All standards 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 331:1983, Coal — Determination of moisture in the analysis sample — Direct gravimetric method.

ISO 687:1974, Coke — Determination of moisture in the analysis sample.

ISO 925:1980, Solid mineral fuels — Determination of carbon dioxide content — Gravimetric method.

ISO 5069-2:1983, Brown coals and lignites — Principles of sampling — Part 2: Sample preparation for determination of moisture content and for general analysis.

3 Principle

A known mass of coal or coke is burnt in a current of oxygen in a tube impervious to gases, the products of the incomplete combustion being further burnt over copper oxide; all the hydrogen is converted to water and all the carbon to carbon dioxide. These products are absorbed by suitable reagents and determined gravimetrically. Oxides of sulfur are retained by lead chromate, chlorine by a silver gauze roll and oxides of nitrogen by granular manganese dioxide.

4 Reagents and materials

WARNING — Care should be exercised when handling reagents, many of which are toxic.

During the analysis, unless otherwise stated, use only reagents of recognized analytical grade and only distilled water or water of equivalent purity. **4.1 Magnesium perchlorate**, anhydrous, less than 1,2 mm in size and preferably within the size range 1,2 mm to 0,7 mm.

WARNING — Due regard must be taken of local regulations when disposing of exhausted magnesium perchlorate. Regeneration of magnesium perchlorate must not be attempted, owing to the risk of explosion.

4.2 Sodium hydroxide on an inert base, preferably of a coarse grading, for example 3,0 mm to 1,5 mm, but not finer than the grading 1,2 mm to 0,7 mm, and preferably of the self-indicating type.

4.3 Manganese dioxide, granular, 1,2 mm to 0,7 mm.

Manganese dioxide in the granular form and the size required can be prepared as follows.

Dissolve manganese sulfate in water and boil the solution. Make the solution alkaline with ammonium hydroxide and add solid ammonium persulfate, in small portions, to the boiling solution until precipied tation is complete. Filter through a hardened fast-filter paper, wash with water by decantation, then with did ar lute sulfuric acid and finally with water until acid-free.

Transfer the moist precipitate to a mortar and place in <u>ISO 62bree</u> furnaces. For the an oven until most of the water has evaporated, but standards stilled 7 in 4 clause 6 the powder is still damp. Press the mass into a scake ad12(0) lengths are appropriate: with a pestle, using firm pressure. Complete the drying, break up the cake cautiously and sieve to separate the 1,2 mm to 0,7 mm size.

4.4 Copper gauze, of mesh approximately 1 mm and 10 mm wide.

4.5 Copper oxide, wire form, chopped to particles approximately 3 mm long with a diameter of approximately 0,2 mm.

4.6 Lead chromate, fused, size range 2,4 mm to 1,2 mm.

4.7 Pure silver gauze, of mesh approximately 1 mm, made of wire approximately 0,3 mm in diameter.

4.8 Oxygen, hydrogen-free, preferably prepared from liquid air and not by electrolysis. Electrolytically prepared oxygen shall be passed over red-hot copper oxide before use to remove any trace of hydrogen.

4.9 Glass wool.

5.1 Analytical balance, capable of weighing to the nearest 0,1 mg.

5.2 Purification train, for absorbing water vapour and carbon dioxide from the oxygen used for the combustion. Assemble the train using a series of U-tubes containing the following reagents in the order stated, in the direction of flow:

- a) magnesium perchlorate (4.1) for absorbing water;
- b) sodium hydroxide on an inert base (4.2) for absorbing carbon dioxide;
- c) magnesium perchlorate for absorbing the water evolved in the reaction between carbon dioxide and sodium hydroxide.

The purification train shall be large enough to render frequent recharging unnecessary, even with continuous use.

5.3 Combustion assembly ds.iteh.ai)

with water until acid-free. te to a mortar and place in <u>ISO 62bree</u> furnaces. The combustion tube is heated by vater has evaporated, but standard described rin clause 6 the 1,25 mm combustion tube ress the mass into a scake ad120 lengths are appropriate:

- a) furnace No. 1 (to heat the boat and its contents to 925 °C) 250 mm;
- b) furnace No. 2 (to keep the entire copper oxide section of the tube heated to 800 °C) — 500 mm;
- c) furnace No. 3 (to cover the lead chromate and the roll of pure silver gauze and to heat the former to about 500 °C) 200 mm.

5.3.2 Combustion tube, of fused silica or suitable hard glass. The diameter of the tube shall be 12 mm to 15 mm. A suitable length is 1,25 m.

5.3.3 Combustion boat, of platinum, porcelain or fused silica, approximately 70 mm long.

5.4 Absorption train, for absorbing the water and carbon dioxide evolved by the combustion of the sample. Assemble the train using the following reagents in the order stated, in the direction of flow.

a) magnesium perchlorate (4.1) for absorbing the water evolved during the combustion;

- b) granular manganese dioxide (4.3) for absorbing oxides of nitrogen;
- c) magnesium perchlorate for absorbing the water evolved from the manganese dioxide;
- d) sodium hydroxide on an inert base (4.2) for absorbing carbon dioxide;
- e) magnesium perchlorate for absorbing the water produced in the reaction between carbon dioxide and sodium hydroxide.

Midvale tubes (figure 1), which provide a large area of reaction, are used for all the reagents except

manganese dioxide, which is contained in a guard tube (figure 2), providing a long contact time with minimum mass.

A typical absorption train, with details of the packing, is shown in figure 3. A is the absorber for water, B is a guard-tube absorber for oxides of nitrogen, and C absorbs any water evolved from the manganese dioxide. Carbon dioxide is absorbed in D, the magnesium perchlorate in the upper portion absorbing any water produced in the reaction between carbon dioxide and sodium hydroxide. A second carbon dioxide absorber, E, should be added as a precautionary measure.

Dimensions in millimetres

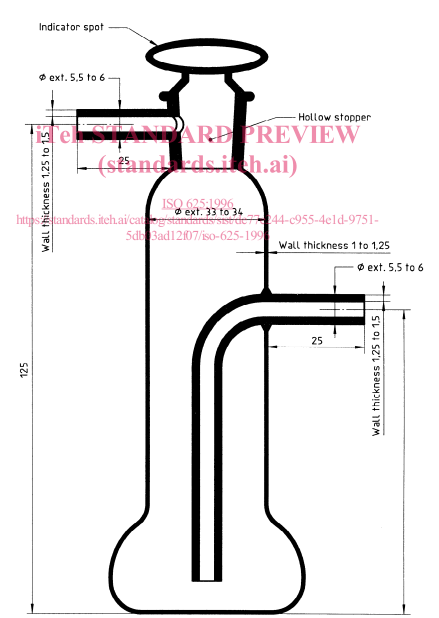


Figure 1 — Midvale tube

Dimensions in millimetres

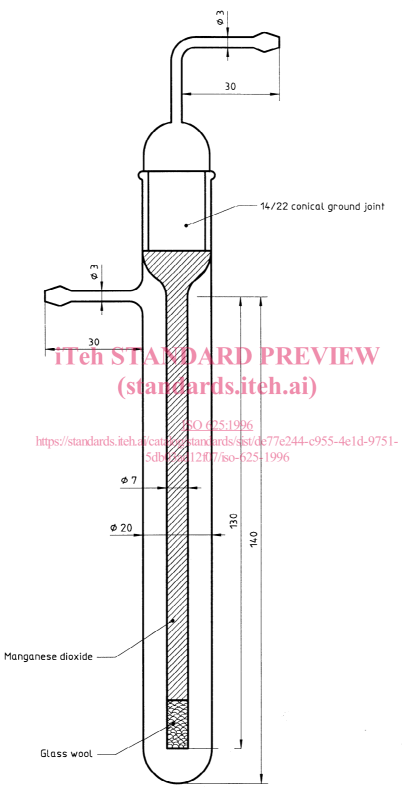
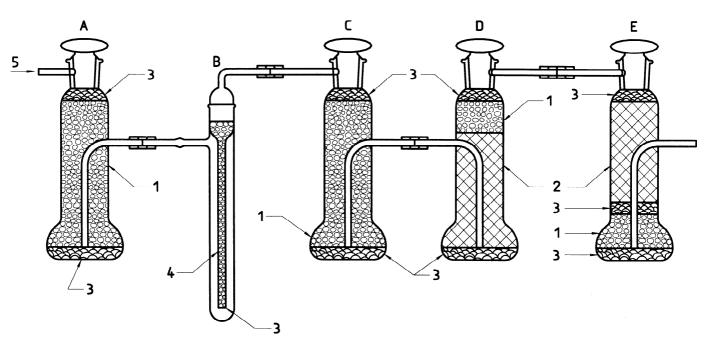


Figure 2 — Guard tube



Key

1 Magnesium perchlorate, 1,2 mm to 0,7 mm size

Manganese dioxide

2 Sodium hydroxide (see 4.2), 1,2 mm to 0,7 mm size 3 Glass wool

NOTE — In this illustration the optional second carbon dioxide absorber, E, is shown.

Figure 30.62 Absorption train

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Stoppered U-tubes may be used, if preferred, in place of the Midvale tubes.

Place glass wool (4.9), previously dried at 105 °C for 1 h, above and below the absorbents to prevent the carry-over of dust by the flow of oxygen, and to prevent the cracking of the Midvale tube by the heat of reaction.

If water is condensed in the first absorber, some nitrogen dioxide may dissolve in it and be considered as water. Because of the conversion factor from water to hydrogen, the error in the hydrogen determination thus caused is small, in the order of 0,05 % of hydrogen. This can only be avoided by heating the absorption tube to a temperature that is sufficiently high to prevent condensation of water.

NOTE 1 Oxides of nitrogen formed in the combustion would, in the absence of precautions, be absorbed by the sodium hydroxide and measured as carbon dioxide. The error in the carbon determination thus caused, in the order of 0,2 % of carbon, is substantially avoided by the use of a guard tube (see figure 2) in which the gases pass through an annular space to allow oxidation of nitrogen monoxide to

nitrogen dioxide which is absorbed by the manganese dioxide.

5.5 Oxygen flow-rate controller, a reducing valve on the oxygen cylinder together with a small needle valve and flowmeter, capable of measuring a flow of up to 100 ml/min, immediately before the purification train, is generally adequate. It may be useful to attach a bubbler device at the exit end of the assembled apparatus to give a visual indication of the rate of flow.

5.6 Heat-resistant stopper (acrylonitrile or chloroprene) for connecting the absorption train to the combustion tube.

5.7 Copper gauze roll, for constraining the reagents in the appropriate sections of the combustion tube (5.3.2). Roll the copper gauze into rolls 10 mm long and of sufficient diameter to ensure a close fit in the combustion tube.

5.8 Copper gauze spiral, through which passes a stout copper wire provided with a loop to facilitate removal from the combustion tube (5.3.2).