

SLOVENSKI STANDARD SIST ISO 778:2002

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Paper, board and pulp -- Determination of copper

Papier, carton et pâte - Détermination de la teneur en cuivre EW

(standards.iteh.aj Ta slovenski standard je istoveten z: ISO 778:2001

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INTERNATIONAL STANDARD

ISO 778

Second edition 2001-06-01

Paper, board and pulp — Determination of copper

Papier, carton et pâte - Détermination de la teneur en cuivre

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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.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 3.

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 a vote.

Attention is drawn to the possibility that some of the elements of this International Standard may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights.

International Standard ISO 778 was prepared by Technical Committee ISO/TC 6, Paper, board and pulps.

This second edition cancels and replaces the first edition (ISO 778:1982), of which it constitutes a technical revision.

The previous edition of this International Standard included the photometric procedure as well as the procedure based on atomic absorption spectroscopy. The photometric procedure has been deleted, as it is now seldom used. The scope has been enlarged to include paper and board in addition to pulp.

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Introduction

This International Standard corresponds to ISO 777^[1] and ISO 779^[2] in order to make it possible to perform the final measurement of all three elements on the same solution.

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Paper, board and pulp — Determination of copper

WARNING — The method specified in this International Standard involves the use of some hazardous chemicals and of gases that can form explosive mixtures with air. Care must be taken to ensure that the relevant precautions are observed.

1 Scope

This International Standard specifies the procedure for the determination of copper by atomic absorption spectrometry or by plasma emission spectrometry.

It is applicable to all types of paper, board and pulp.

It specifies a method to determine the acid-soluble part of the incineration residue, i.e. that part of the ignition residue obtained after incineration which is soluble in hydrochloric acid. If the residue is completely soluble, the result obtained by the procedure specified in this International Standard is taken as the total amount of copper in the sample. The limit of determination is normally 0,1 mg/kg of paper, board or pulp, but it depends on the spectrometer used.

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2 Normative references

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The following normative documents contain provisions which, through reference in this text, constitute provisions of this International Standard. For dated references, subsequent amendments to, or revisions of, any of these publications do not apply. However, parties to agreements based on this International Standard are encouraged to investigate the possibility of applying the most recent editions of the normative documents indicated below. For undated references, the latest edition of the normative document referred to applies. Members of ISO and IEC maintain registers of currently valid International Standards.

- ISO 186, Paper and board Sampling to determine average quality.
- ISO 287, Paper and board Determination of moisture content Oven-drying method.
- ISO 638, Pulps Determination of dry matter content.
- ISO 1762, Paper, board and pulps Determination of residue (ash) on ignition at 525 °C.
- ISO 7213, Pulps Sampling for testing.

3 Term and definition

For the purposes of this International Standard, the following term and definition applies.

3.1

mass fraction of copper

amount of the element copper in the solution obtained after incineration of the specimen at 525 °C and treating the residue with 6 mol/l hydrochloric acid as specified in this International Standard

4 Principle

A test portion is incinerated at 525 °C and the residue is treated with 6 mol/l hydrochloric acid. The test solution is aspirated into an acetylene/dinitrogen monoxide or acetylene/air flame and the mass fraction of copper is determined by one of the following procedures:

- measurement of the absorption of the 324,7 nm line emitted by a copper hollow-cathode lamp; or
- measurement of the absorption of the 324,7 nm line emitted by plasma emission spectrometry.

5 Reagents and materials

Use only chemicals of recognized analytical grade and only distilled or deionized water.

5.1 Hydrochloric acid, about 6 mol/l.

Dilute 500 ml of hydrochloric acid (density 1,19 g/ml) in 500 ml of water.

- 5.2 Nitric acid, density 1,4 g/ml.
- 5.3 Copper stock solution, 100 mg/l of Cu.

Dissolve 100 mg of pure electrolytic copper in the smallest quantity possible of nitric acid (density 1,4 g/ml). Boil in order to expel nitrous fumes and allow to cool. Transfer the solution completely to a 1 000 ml volumetric flask. Dilute with water to the mark and mix.h STANDARD PREVIEW

1 ml of this stock solution contains 0,10 mg of Cundards.iteh.ai)

5.4 Copper standard solution, 10 mg/l of Cu. SIST ISO 778:2002

Transfer 100 ml of the copper stock solution (5.3) to a 1000 ml volumetric flask and add 200 ml of hydrochloric acid (5.1). Dilute with water to the mark and mix.

1 ml of this standard solution contains 0,01 mg of Cu.

Commercially available, certified standard solutions may be used.

5.5 Acetylene gas and/or dinitrogen monoxide gas, of a grade suitable for atomic absorption spectrometry.

WARNING — Acetylene gas forms explosive mixtures with air.

5.6 Appropriate gas for the plasma spectrometer (6.4). Argon is usually recommended as a carrier gas.

6 Apparatus and equipment

Ordinary laboratory equipment. Clean all equipment in 0,1 mol/l hydrochloric acid.

- **6.1** Filter paper, ash free, particle retention 20 μ m to 25 μ m.
- 6.2 Dishes, of platinum or quartz.

6.3 Atomic absorption spectrometer, with a burner for dinitrogen monoxide/acetylene or air/acetylene and with a hollow-cathode lamp for copper.

NOTE A multi-element lamp may be used.

6.4 Inductively coupled plasma spectrometer.

7 Sampling and preparation of sample

If the analysis is being made to evaluate a lot of paper, board or pulp, the sample shall be selected in accordance with ISO 186 or ISO 7213, as relevant. If the analysis is made on another type of sample, report the source of the sample, and, if possible, the sampling procedure. Select the specimens so that they are representative of the sample received. A sufficient amount of sample shall be collected to allow for at least duplicate determinations. Avoid cut edges, punched holes and other parts where metallic contamination may have occurred.

Prepare a test specimen by tearing at least 50 g of small pieces from various parts of the sample. This amount is sufficient for the duplicate determinations as prescribed in clause 8.

8 Procedure

8.1 Incineration and dissolution of the residue

Carry out the procedure in duplicate.

Air-dry the specimen in the laboratory atmosphere until it reaches moisture equilibrium.

Determine the moisture content on a separate air-dried portion as specified in ISO 287 or ISO 638, as relevant. Weigh this portion at the same time as the test portion used for incineration.

Carry out ashing of the test portion as described in ISO 1762. A test portion of 10 g to 20 g is recommended, because the copper content of paper, board and pulp is normally very low.

Carry out the dissolution of the ash under a tune bood. To avoid splattering, carefully moisten the ash with water and add 5 ml of hydrochloric acid (5.1) to the dish. Evaporate to dryness on a boiling-water bath or equivalent device. Repeat this procedure.

For samples with a high carbonate content, more than 10 ml of acid $(2 \times 5 \text{ ml})$ may be needed, for example 20 ml $(2 \times 10 \text{ ml})$.

Add 2,5 ml of the hydrochloric acid (5.1) in order to dissolve the dry residue. If necessary, heat the dish covered by a watch glass for a few minutes.

Using the filter paper (6.1), filter the contents of the dish into a 25 ml volumetric flask. To ensure that the transfer is complete, add another portion of 2,5 ml of acid to the dish and heat again. Filter this last portion of acid into the main portion in the volumetric flask with the aid of some water. Fill up to the mark and mix. This is the test solution.

8.2 Blank

Run a blank with the same quantity of each of the chemicals as those added to the incineration residue but without any residue.

9 Preparation of calibration solutions

It is important that the acid concentration is the same in the calibration solutions as in the test solution, since the acid concentration influences the signal.

From the copper standard solution (5.4), prepare at least three calibration solutions, and in addition one zero solution, for the construction of the calibration graph. (The zero solution is similar to the calibration solutions, but contains no added copper. Do not confuse it with the blank.)

NOTE Not more than two calibration solutions are needed for plasma emission spectrometry.