EAST AFRICAN STANDARD

Meat and meat products — Determination of total ash

EAST AFRICAN COMMUNITY

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Foreword

Development of the East African Standards has been necessitated by the need for harmonizing requirements governing quality of products and services in East Africa. It is envisaged that through harmonized standardization, trade barriers which are encountered when goods and services are exchanged within the Community will be removed.

In order to meet the above objectives, the EAC Partner States have enacted an East African Standardization, Quality Assurance, Metrology and Test Act, 2006 (EAC SQMT Act, 2006) to make provisions for ensuring standardization, quality assurance, metrology and testing of products produced or originating in a third country and traded in the Community in order to facilitate industrial development and trade as well as helping to protect the health and safety of society and the environment in the Community.

East African Standards are formulated in accordance with the procedures established by the East African Standards Committee. The East African Standards Committee is established under the provisions of Article 4 of the EAC SQMT Act, 2006. The Committee is composed of representatives of the National Standards Bodies in Partner States, together with the representatives from the private sectors and consumer organizations. Draft East African Standards are circulated to stakeholders through the National Standards Bodies in the Partner States. The comments received are discussed and incorporated before finalization of standards, in accordance with the procedures of the Community.

Article 15(1) of the EAC SQMT Act, 2006 provides that “Within six months of the declaration of an East African Standard, the Partner States shall adopt, without deviation from the approved text of the standard, the East African Standard as a national standard and withdraw any existing national standard with similar scope and purpose”.

East African Standards are subject to review, to keep pace with technological advances. Users of the East African Standards are therefore expected to ensure that they always have the latest versions of the standards they are implementing.

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Introduction

In the preparation of this East African Standard, the following sources were consulted extensively:


Codex Alimentarius website: http://www.codexalimentarius.net/mrls/pestdes/jsp/pest_q-e.jsp

USDA Foreign Agricultural Service website: http://www.mrldatabase.com

USDA Agricultural Marketing Service website: http://www.ams.usda.gov/AMSv1.0/Standards


European Union: http://ec.europa.eu/sanco_pesticides/public

Assistance derived from these sources is hereby acknowledged.
INTERNATIONAL STANDARD

ISO

936

Second edition
1998-08-01

Meat and meat products — Determination of total ash

Viande et produits à base de viande — Dosage des cendres totales
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 a vote.

International Standard ISO 936 was prepared by Technical Committee ISO/TC 34, Agricultural food products, Subcommittee SC 6, Meat and meat products.

This second edition cancels and replaces the first edition (ISO 936:1978), which has been technically revised.

Annex A of this International Standard is for information only.
Meat and meat products — Determination of total ash

1 Scope

This International Standard specifies a method for the determination of the total ash from all kinds of meat and meat products, including poultry.

2 Normative reference

The following standard contains provisions which, through reference in this text, constitute provisions of this International Standard. At the time of publication, the edition indicated was 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 edition of the standard indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.


3 Definition

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

3.1 total ash from meat and meat products
mass of the residue obtained after incineration at a temperature of (550 ± 25) °C under the operating conditions specified in this International Standard, divided by the mass of the test portion

NOTE The mass fraction of ash is usually expressed as a percentage.

4 Principle

A test portion is dried, carbonized and then incinerated at (550 ± 25) °C. After cooling, the mass of the residue is determined.

5 Reagents

Use only reagents of recognized analytical grade, unless otherwise specified.

5.1 Water, complying with at least grade 3 in accordance with ISO 3696.
5.2 Hydrogen peroxide, 30 %.

6 Apparatus

Usual laboratory apparatus and, in particular, the following.

6.1 Mechanical or electrical equipment capable of homogenizing the laboratory sample.

This includes a high-speed rotational cutter, or a mincer fitted with a plate with apertures not exceeding 4,0 mm in diameter.

6.2 Dish, flat-bottomed, made of porcelain, quartz or metal (e.g. nickel, platinum, stainless steel) or of other material unaffected by the conditions of the test, with a diameter of at least 60 mm, and inclined walls of height at least 25 mm.

6.3 Muffle furnace, electrically heated and equipped with a programmable time-temperature controller, and capable of being maintained at (550 ± 25) °C.

6.4 Desiccator, containing an efficient desiccant.

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

6.6 Drying oven, capable of being maintained at (103 ± 2) °C (if the muffle furnace has no time-temperature controller).

6.7 Electric hot-plate or a gas flame (if the muffle furnace has no time-temperature controller).

7 Sampling

Sampling is not part of the method specified in this International Standard. A recommended sampling method is given in ISO 3100-1 [1].

It is important that the laboratory receive a sample which is truly representative and has not been damaged or changed during transport or storage.

Start from a representative sample of at least 200 g. Store the sample in such a way that deterioration and change in composition are prevented.

8 Preparation of test sample

Homogenize the laboratory sample with the appropriate equipment (6.1). Take care that the temperature of the sample material does not rise above 25 °C. If a mincer is used, pass the sample at least twice through the equipment.

Fill a suitable airtight container with the prepared sample. Close the container and store in such a way that deterioration and change in composition are prevented. Analyse the sample as soon as practicable, but always within 24 h after homogenization.
9 Procedure

NOTE If it is required to check whether the repeatability limit (see 11.2) is met, carry out two single determinations in accordance with 9.1 to 9.3.

9.1 Test portion

Heat the dish (6.2) for 20 min in the muffle furnace (6.3) set at 550 °C.

Allow the dish to cool in the desiccator (6.4) to room temperature and weigh \( m_0 \) on the analytical balance (6.5) to the nearest 0.1 mg.

Transfer 1.5 g to 2 g of the prepared test sample (see clause 8) to the dish. Spread it out evenly and without delay weigh \( m_1 \) the dish again to the nearest 0.1 mg.

NOTE If deemed appropriate by the analyst, having regard to the nature of the laboratory sample, up to 5 g of the prepared test sample may be taken. This should be noted in the test report.

If the muffle furnace is provided with a time-temperature controller, proceed in accordance with 9.2. If not, proceed in accordance with 9.3.

9.2 Determination applying muffle furnace with time-temperature controller

Place the dish with its contents in the cool muffle furnace (6.3) and gradually raise the temperature of the furnace over 5 h to 6 h to \((550 \pm 25) \, ^{\circ}\text{C}\). Continue the ashing at \((550 \pm 25) \, ^{\circ}\text{C}\) until the ash has a grey-white appearance.

Remove the dish from the muffle furnace and allow to cool in the desiccator (6.4) to room temperature.

CAUTION - Avoid loss of ash when transferring the dish with the ash from the furnace to the desiccator and from the desiccator to the analytical balance.

Inspect the ash.

If the ash is still black, treat it with a few drops of hydrogen peroxide (5.2) or water (5.1) and repeat the procedure described in this subclause.

If the ash has a grey-white appearance, weigh on the analytical balance (6.5), to the nearest 0.1 mg, the dish with its contents \( m_2 \). Proceed in accordance with clause 10.

9.3 Determination applying muffle furnace without time-temperature controller

Place the dish with its contents for 1 h in the drying oven (6.6) set at 103 °C.

Remove the dish from the oven and place it on an electric hot-plate or over a gas flame (6.7). Heat progressively until the substance carbonizes with the evolution of smoke. Carefully continue carbonization until smoke evolution ceases. The sample material shall neither ignite nor burn with a flame.

Transfer the dish to the cool muffle furnace (6.3) and raise the temperature to \((550 \pm 25) \, ^{\circ}\text{C}\).

After 4 h, remove the dish with its contents from the muffle furnace and allow to cool in the desiccator (6.4) to room temperature.

CAUTION - Avoid loss of ash when transferring the dish with the ash from the furnace to the desiccator and from the desiccator to the analytical balance.
Inspect the ash.

If the ash is still black, treat it with a few drops of hydrogen peroxide (5.2) or water (5.1) and repeat the procedure described in this subclause.

If the ash has a grey-white appearance, weigh on the analytical balance (6.5), to the nearest 0.1 mg, the dish with its contents ($m_2$).

## 10 Calculation

Calculate the mass fraction of ash of the test sample using the equation:

$$w_a = \frac{(m_2 - m_0)}{(m_1 - m_0)} \times 100\%$$

where

- $w_a$ is the mass fraction of ash, as a percentage, of the test sample;
- $m_0$ is the mass, in grams, of the empty dish;
- $m_1$ is the mass, in grams, of the dish with the test portion;
- $m_2$ is the mass, in grams, of the dish with the ash.

Report the result rounded to the nearest 0.01 %.

## 11 Precision

### 11.1 Interlaboratory test

The precision of the method was established by an interlaboratory test carried out in accordance with ISO 5725 [2]$^1$.

The results of the interlaboratory test have been published (see reference [6]). The values derived from this test may not be applicable to concentration ranges and matrices other than those given.

### 11.2 Repeatability

The absolute difference between two independent single test results, obtained using the same method on identical test material in the same laboratory by the same operator using the same equipment within a short interval of time, will in not more than 5 % of cases exceed the repeatability limit $r$ given by the equation:

$$r = 0.0990\% + 0.00933\overline{w}$$

where

- $r$ is the repeatability limit, as a percentage;
- $\overline{w}$ is the mean of both results, as a percentage.

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$^1$ ISO 5725:1986 (now withdrawn) was used to obtain the precision data.
11.3 Reproducibility

The absolute difference between two single test results, obtained using the same method on identical test material in different laboratories with different operators using different equipment, will in not more than 5 % of cases exceed the reproducibility limit $R$ given by the equation:

$$R = 0.138 \% + 0.0046 \bar{w}$$

where

$R$ is the reproducibility limit, as a percentage;

$\bar{w}$ is the mean of both results, as a percentage.

12 Test report

The test report shall specify:

— all information necessary for the complete identification of the sample;
— the sampling method used, if known;
— the test method used, with reference to this International Standard;
— all operating details not specified in this International Standard, or regarded as optional, together with details of any incidents which may have influenced the test result(s);
— the test result(s) obtained; or
— if the repeatability has been checked, the final quoted result obtained.
Annex A
(informative)

Bibliography


