

# SLOVENSKI STANDARD SIST EN ISO 13366-1:1998 01-avgust-1998

# Mleko - Ugotavljanje števila somatskih celic - 1. del: Mikroskopska metoda (ISO 13366-1:1997)

Milk - Enumeration of somatic cells - Part 1: Microscopic method (ISO 13366-1:1997)

Milch - Zählung somatischer Zellen - Teil 1: Mikroskopisches Verfahren (ISO 13366-1:1997)

Lait - Dénombrement des cellules somatiques - Partie 1: Méthode au microscope (ISO (standards.iteh.ai) 13366-1:1997)

Ta slovenski standard je istoveten z: ogstan EN ISO 13366-1:1997-228-53aa13288226/sist-en-iso-13366-1-1998

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#### EUROPEAN STANDARD

#### EN ISO 13366-1

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#### NORME EUROPÉENNE

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# EUROPÄISCHE NORM

ICS 07.100.30; 67.100.10

Descriptors: see ISO document

English version

### Milk - Enumeration of somatic cells - Part 1: Microscopic method (ISO 13366-1:1997)

Lait - Dénombrement des cellules somatiques -Partie 1: Méthode au microscope (ISO 13366-1:1997) Milch - Zählung somatischer Zellen - Teil 1: Mikroskopisches Verfahren (ISO 13366-1:1997)

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# 53aa13288226/sist-en-CEN-1-1998

European Committee for Standardization Comité Européen de Normalisation Europäisches Komitee für Normung

Central Secretariat: rue de Stassart,36 B-1050 Brussels

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# Foreword

The text of the International Standard ISO 13366-1:1997 has been prepared by Technical Committee ISO/TC 34 "Agricultural food products" in collaboration with Technical Committee CEN/TC 302 "Milk and milk products - Methods of sampling and analysis", the secretariat of which is held by NNI.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by December 1997, and conflicting national standards shall be withdrawn at the latest by December 1997.

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# **Endorsement notice**

The text of the International Standard ISO 13366-1:1997 was approved by CEN as a European Standard without any modification.

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# INTERNATIONAL STANDARD

# ISO 13366-1

First edition 1997-06-15

# Milk — Enumeration of somatic cells —

# Part 1: Microscopic method

Lait — Dénombrement des cellules somatiques —

Partie 1: Méthode au microscope iTeh STANDARD PREVIEW (standards.iteh.ai)

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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.

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 13366-1 was prepared by Technical Committee ISO/TC 34, Agricultural food products, in collaboration with the International Dairy Federation (IDF) and AOAC INTERNATIONAL, and will also be published by these organizations.

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IEW

ISO 13366 consists of the following parts, under the general title *Milk* <u>47-fa7c-4b47-a2a8-</u> Enumeration of somatic cells: 53aa13288226/sist-en-iso-13366-1-1998

- Part 1: Microscopic method
- Part 2: Electronic particle counter method
- Part 3: Fluoro-opto-electronic method

Annex A of this part of ISO 13366 is for information only.

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# Milk — Enumeration of somatic cells —

# Part 1:

Microscopic method

WARNING — The use of this standard may involve hazardous materials, operations and equipment. This standard does not purport to address all of the safety problems associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and to determine the applicability of regulatory limitations prior to use.

# 1 Scope iTeh STANDARD PREVIEW

This part of ISO 13366 specifies a method for counting somatic cells in both raw and chemically preserved milk. The method is suitable for preparing standard test samples and for calibrating mechanized and automatic cell-counting procedures.

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# 2 Definition

For the purposes of this part of ISO 13366, the following definition applies.

2.1 somatic cells: Those cells with nuclei, that is, all leucocytes and epithelial cells.

# **3** Principle

Spreading of a test portion of the milk to be examined over a slide to form a film. Drying and staining of the film and subsequent counting of the stained cells using a microscope. Multiplication of the number of cells counted in a defined area by a working factor to give the number of cells per millilitre.

# 4 Reagents

WARNING — Tetrachloroethane is poisonous. Ethidium bromide is toxic. The preparation and application of the dye solution shall be carried out in a fume cupboard. Use gloves for protection.

Use only reagents of recognized analytical grade, unless otherwise specified, and distilled or deionized water or water of equivalent purity.

### 4.1 Dye solution

#### 4.1.1 Composition

Ethanol 95 % ( <i>V/V</i> )	54.0 ml
Tetrachloroethane	40,0 ml
Methylene blue	0,6 g
Acetic acid, glacial	6,0 ml

NOTE — As an alternative, tetrachloroethane may be replaced by the same amount of trichloroethane. Instead of methylene blue, ethidium bromide can be used (see ISO 13366-3).

#### 4.1.2 Preparation

Mix the ethanol and tetrachloroethane in a bottle. Heat in the water bath (5.1) set at 65 °C. Add the methylene blue and mix carefully. Cool in a refrigerator to 4 °C and then add the glacial acetic acid. Pass the solution through an appropriate filter (5.3) into an airtight bottle and store it as such. If necessary, filter again before use.

# **5** Apparatus

Usual laboratory equipment and, in particular, the following.

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- **5.1 Water bath**, capable of being maintained at a temperature of  $65 \degree C \pm 5 \degree C$ .
- **5.2** Water bath, capable of being maintained at a temperature of  $35 \degree C \pm 5 \degree C$ .

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5.3 Filter, resistant to the solvents used, with a pore/size of between 10 μm and 12 μm or less.

**5.4** Microscope, with magnification of between  $\times$  500 and  $\times$  1 000.

If ethidium bromide is used, the microscope shall have fluorescence equipment.

5.5 Microsyringe, of capacity 0,01 ml, with maximum tolerance of 2 %.

**5.6** Slides, marked with the outline of a shape of a film of dimensions 20 mm  $\times$  5 mm, or a standard slide and a template of dimensions 20 mm  $\times$  5 mm.

**5.7** Hot-plate, capable of being maintained at a temperature of 40 °C  $\pm$  10 °C.

**5.8** Fan, hairdryer type.

# 6 Sampling

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

Sampling is not part of the method specified in this International Standard. A recommended sampling method is given in ISO 707<sup>[1]</sup>.

6.2 If automatic samplers are used, they shall be properly tested.

6.3 Prior to testing or preservation, samples should be stored at a temperature of between 2 °C and 6 °C.

**6.4** Samples not to be tested within 6 h after sampling shall be preserved by the addition of boric acid. The final concentration of boric acid shall not exceed 0,6 g per 100 ml of sample. Store such samples at a temperature of between 2 °C and 6 °C for no longer than 24 h.

# 7 Preparation of test sample

Heat the test sample in the water bath (5.2) set at a temperature of 35 °C. Mix the test sample carefully and cool to the temperature at which the microsyringe has been calibrated, for example to 20 °C.

# 8 Procedure

Prepare and count from each test sample at least two films. Clean the slides (5.6), for example with ethanol. Dry them with dust-free paper, flame and cool.

### 8.1 Test portion and preparation of film

Use the microsyringe (5.5) to take 0,01 ml of the prepared test sample (clause 7). Carefully clean the outside of the syringe which has been in contact with the sample. Place the test portion on a clean slide with the outline of the shape ( $20 \text{ mm} \times 5 \text{ mm}$ ) (5.6). Then fill in this area as evenly as possible with the test portion. Dry the film on a level hot-plate (5.7) until completely dry. Better results can be obtained by drying the films at ambient temperature for several hours.

Dip the dried film on the slide in the dye solution (4.1) for 10 min. Complete drying with the fan (5.8) if required. Then dip the film in tap water until all surplus dye is washed away. Dry again and store with protection against dust.

#### 8.2 Determination

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Using the microscope (5.4), count the cell nuclei in the film (at least 400). These are clearly recognizable and at least half should be visible in the microscope field. Count nuclei in vertical strips in the middle third of the film. Avoid counting strips selected exclusively from the peripheral areas of the film.

Check at least once a month the proper preparation of the films, and hence the reliability of the results, by counting different parts of the film.

# 9 Calculation and expression of results

**9.1** The length of the strips to be counted is 5 mm each. The width of a strip corresponds to the diameter of the microscope field. With a test portion of 0,01 ml of sample, calculate the working factor,  $w_f$ , using the following equation:

$$w_{\rm f} = \frac{20 \times 100}{d \times b}$$

where

- d is the numerical value of the diameter, in millimetres, of the microscope field;
- *b* is the number of strips counted completely.