INTERNATIONAL **STANDARD**

ISO 8784-1

> Second edition 2005-07-01

Pulp, paper and board — Microbiological examination —

Part 1:

Total count of bacteria, yeast and mould based on disintegration

iTeh STANDARD PREVIEW
Pâte, papier et carton — Analyse microbienne

Strartie 1. Dénombrement total des bactéries, levures et moisissures basé sur la désintégration

ISO 8784-1:2005

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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 2.

The main task of technical committees is to prepare International Standards. 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 document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights.

ISO 8784-1 was prepared by Technical Committee ISO/TC 6, *Paper, board an pulps*, Subcommittee SC 2, *Test methods and quality specifications for paper and board*.

This second edition cancels and replaces the first edition (ISO 8784-1:1987), which has been technically revised. The first edition was only applicable to paper and board and only the bacterial colonies were determined. The second edition is also applicable to pulp (dry market pulp), and yeast and mould as well as bacteria are determined. The incubation conditions have been changed in the second edition (37 °C, 48 h for bacteria, 30 °C, 5 days for yeast and mould) compared to the conditions stated in the first edition (30 °C, 72 h). https://standards.iteh.ai/catalog/standards/sist/99abc9af-7d0a-479e-8b02-

ISO 8784 consists of the following parts, under the general title Pulp, paper and board — Microbiological examination:

- Part 1: Total count of bacteria, yeast and mould based on disintegration
- Part 2: Surface method for enumeration of microbes on paperboard

Introduction

This part of ISO 8784, which deals with the microbiological examination of dry market pulp, paper and paperboard, is broadly based on ISO 4833^[1], although the conditions are not identical: However, it provides specific amplification where necessary. It is intended for the estimation of colony-forming units, CFU, without any attempt to isolate species of particular public-health significance.

Because of the exacting techniques required in aseptic procedures, reproducible results can only be secured by skilled microbiological technicians. In addition, health risks may arise from the employment of inadequately trained staff.

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Pulp, paper and board — Microbiological examination —

Part 1:

Total count of bacteria, yeast and mould based on disintegration

Scope

This part of ISO 8784 specifies a method for determining the total number of colony-forming units of bacteria, yeast and moulds in dry market pulp, paper and paperboard after desintegration. The enumeration relates to specific media.

This part of ISO 8784 is applicable to most kinds of paper and paperboard, especially those grades intended to come into contact with foodstuffs.

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Normative references (standards.iteh.ai)

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The following referenced documents are indispensable for the application of this document. For dated references, only the edition cited applies SFor undated references, the latest edition of the referenced document (including any amendments) applies tandards/sist/99abc9af-7d0a-479e-8b02-

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 7213, Pulps — Sampling for testing

Terms and definitions

For the purposes of this document, the following terms and definitions apply.

3.1

microbiological count

number of colony-forming units (CFU) formed in a standard culture medium, after incubation under the conditions specified

Principle

Preparation of culture plates from specified dilutions of a suspension of the sample on a specified culture medium. For the determination of bacteria, aerobic incubation of the culture plates for 48 h at 37 °C. For the determination of yeast and mould, aerobic incubation of the culture plates for 5 days at 30 °C. At the end of the incubation time, counting the CFU on the plates.

From the number of colonies counted, the number of CFU per gram of the sample is calculated.

Substrates and diluents

All substrates and diluents shall be appropriately sterilized. When preparing the culture medium, make sure that the ingredients are completely dissolved prior to dispensing into suitable containers and sterilization.

5.1 **Nutrient substrate**

5.1.1 Tryptone Glucose Extract Agar (TGEA)

Use TGEA for bacterial counts. Composition per litre:

Beef extract	3,0	g
Tryptone	5,0	g
Dextrose (d-glucose)	1,0	g
Agar	15,0	g
pH of the ready-made medium	7,0	•

If TGEA is not available, Plate Count Agar (PCA) may be substituted. The use of PCA shall be stated in the test report. The composition of the PCA medium is the same as TGEA, except that beef extract (3 g) is substituted with 2,5 g yeast extract.

5.1.2 Potato Dextrose Agar (PDA)

Use PDA for yeast and mould counts. Composition per litre: 1Teh STANDARD PREVIEW

Potatoes, infusion from 200 g (standards.iteh.ai) 20 g Dextrose

15 g Agar

pH of the ready-made medium 5,6 ISO 8784-1:2005

https://standards.iteh.ai/catalog/standards/sist/99abc9af-7d0a-479e-8b02-Acidification of PDA: Acidify the medium with drops of 10 % sterile tartaric acid to reach a pH of 3,5 \pm 0,1. After adding the tartaric acid to the medium, mix without foaming, and prepare poured plates as usual. Do not heat the medium after the acid has been added, since heating in the acid state will hydrolyse the agar and destroy its solidifying properties.

It is frequently desirable in making yeast and mould counts to inhibit bacterial growth by acidifying the PDA medium, and it is recommended that the reaction of the medium be reduced to pH (3.5 ± 0.1) subsequent to sterilization. The growth of bacteria can alternatively be inhibited by addition of a bactericide. The use of a bactericide shall be stated in the test report.

The Tryptone Glucose Extract Agar (TGEA) and the Potato Dextrose Agar (PDA) are commercially available NOTE in dehydrated form. When using the dehydrated medium, follow the instructions printed on the container.

Standard diluent: Ringer's solution (see A.1) is preferred, although other isotonic solutions may be used. Ringer's tablets are commercially available.

To facilitate the release of cells from the fibres, it is recommended to add 20 µl of Tween 80 (see A.2) per litre to the Ringer's solution prior to sterilization.

Equipment

All equipment in direct contact with the sample or the diluents shall be sterilized. Use ordinary microbiological laboratory equipment, and the following.

Colony-counting equipment, fitted with a lens having a magnifying power of at least 1,5 times. The use of an additional lens in conjunction with the lens on the colony counter may be necessary to increase the magnification to 8 to 10 times, to facilitate the counting of pin-point bacterial colony-forming units.

- **6.2 Disintegrators**, with metal or glass jars of about 500 ml capacity, fitted with a high-speed impeller near the bottom and fitted with a cap or lid; or other suitable disintegrator which ensures disintegration of the sample. Place an aluminium foil hood over the cap of each disintegrator jar prior to sterilization.
- **6.3** Incubator, capable of maintaining temperatures of (30 ± 1) °C and (37 ± 1) °C.
- **6.4** Petri dishes, having a diameter of 150 mm.
- **6.5** Pipettes, of suitable wide-mouth type.
- **6.6 Aluminium foil, envelopes, or self-closing plastic bags**, for sampling. Samples can be wrapped as such in aluminium foil, in ready-to-use sterile envelopes of different sizes or self-closing plastic bags, of which all are commercially available.

NOTE Petri dishes having a diameter of 90 mm may be used. However, the size 150 mm is recommended to make it easier to separate the colonies from the material.

7 Sampling

Make sure that the sampling procedure is performed aseptically.

If the sample is to represent a lot of paper or paperboard, the sampling shall be in accordance with ISO 186^[1]. If the sample is to represent a lot of pulp, the sampling shall be in accordance with ISO 7213. From each unit of dry market pulp to be sampled, discard several top sheets from each bale to eliminate surface contamination. From each unit of paper or paperboard to be sampled, cut away several top layers and discard them to eliminate surface contamination. Use a sterile knife and cut a bunch of sheets (see the third paragraph of this clause). Discard the top sheet. To Salte 1. 21

In other cases, sample a sufficient number of units so that the test material is representative of the paper or paperboard or the dry market pulp to be tested. In all sampling and examination procedures, make sure that the test material taken is representative of the sample received.

Ideally, a sample should contain at least five sheets, each of them having a minimum size of $200 \text{ mm} \times 250 \text{ mm}$ of dry market pulp, paper or paperboard (at least 3 sheets for testing and 2 protective sheets).

After sampling, wrap the unexposed test material in suitable wrapping material (6.6).

8 Preparation of the test material

Preferably, the procedure is conducted in a sterile hood. Weighing takes place outside the hood. Unwrap the test material under aseptic conditions and remove the protective sheets without touching the test sheets.

8.1 Determination of dry-matter content

If the result is to be reported on a dry-mass basis, determine the dry-matter content of the test material, x, in accordance with ISO 287 or ISO 638, as relevant.

If the result is to be reported on an "as received"-mass basis, the determination of dry-matter content shall be omitted (also, see Note 1 to 11.1).

8.2 Weighing

Place a closed Petri dish (6.4) on the pan of the balance and determine its tare mass.

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