

---

---

**Compressed air —**

**Part 7:**

**Test method for viable microbiological  
contaminant content**

*Air comprimé —*

**STANDARD PREVIEW**  
**(standards.iteh.ai)**

*Partie 7: Méthode d'essai pour la détermination de la teneur en  
polluants microbiologiques viables*

ISO 8573-7:2003

<https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003>



Reference number  
ISO 8573-7:2003(E)

**PDF disclaimer**

This PDF file may contain embedded typefaces. In accordance with Adobe's licensing policy, this file may be printed or viewed but shall not be edited unless the typefaces which are embedded are licensed to and installed on the computer performing the editing. In downloading this file, parties accept therein the responsibility of not infringing Adobe's licensing policy. The ISO Central Secretariat accepts no liability in this area.

Adobe is a trademark of Adobe Systems Incorporated.

Details of the software products used to create this PDF file can be found in the General Info relative to the file; the PDF-creation parameters were optimized for printing. Every care has been taken to ensure that the file is suitable for use by ISO member bodies. In the unlikely event that a problem relating to it is found, please inform the Central Secretariat at the address given below.

**iTeh STANDARD PREVIEW**  
**(standards.iteh.ai)**

ISO 8573-7:2003

<https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003>

© ISO 2003

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office  
Case postale 56 • CH-1211 Geneva 20  
Tel. + 41 22 749 01 11  
Fax + 41 22 749 09 47  
E-mail [copyright@iso.org](mailto:copyright@iso.org)  
Web [www.iso.org](http://www.iso.org)

Published in Switzerland

# Contents

Page

<b>Foreword</b> .....	<b>iv</b>
<b>1 Scope</b> .....	<b>1</b>
<b>2 Normative references</b> .....	<b>1</b>
<b>3 Terms and definitions</b> .....	<b>1</b>
<b>4 Method for verifying presence of viable micro-organisms by partial flow sampling</b> .....	<b>2</b>
<b>5 Operating conditions</b> .....	<b>2</b>
<b>6 Determination of viable, colony-forming organisms</b> .....	<b>3</b>
<b>7 Test report statement</b> .....	<b>3</b>
<b>Annex A (informative) Determination of viable microbiological particle content in compressed air — Sample test report</b> .....	<b>4</b>
<b>Annex B (normative) Quantitative sampling method</b> .....	<b>5</b>
<b>Annex C (informative) Sampling endotoxins</b> .....	<b>7</b>
<b>Annex D (informative) Preparation of Petri dish with culturable media</b> .....	<b>8</b>
<b>Bibliography</b> .....	<b>9</b>

<https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003>  
 (standards.iteh.ai)

## 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 8573-7 was prepared by Technical Committee ISO/TC 118, *Compressors, pneumatic tools and pneumatic machines*, Subcommittee SC 4, *Quality of compressed air*.

ISO 8573 consists of the following parts, under the general title *Compressed air*:

- *Part 1: Contaminants and purity classes*
- *Part 2: Test methods for aerosol oil content* [ISO 8573-7:2003](https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003)
- *Part 3: Test methods for measurement of humidity*
- *Part 4: Test methods for solid particle content*
- *Part 5: Test methods for oil vapour and organic solvent content*
- *Part 6: Test methods for gaseous contaminant content*
- *Part 7: Test method for viable microbiological contaminant content*
- *Part 8: Test methods for solid particle content by mass concentration*
- *Part 9: Test methods for liquid water content*

# Compressed air —

## Part 7:

## Test method for viable microbiological contaminant content

### 1 Scope

This part of ISO 8573 specifies a test method for distinguishing viable, colony-forming, microbiological organisms (e.g. yeast, bacteria, endotoxins) from other solid particles which may be present in compressed air. One of a series of standards aimed at harmonizing air contamination measurements, it provides a means of sampling, incubating and determining the number of microbiological particles. The test method is suitable for determining purity classes in accordance with ISO 8573-1, and is intended to be used in conjunction with ISO 8573-4 when there is need to identify solid particles that are also viable, colony-forming units.

### 2 Normative references

The following referenced documents are indispensable for the application of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 8573-1, *Compressed air — Part 1: Contaminants and purity classes*  
<https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003>

ISO 8573-4, *Compressed air — Part 4: Test methods for solid particle content*

### 3 Terms and definitions

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

#### 3.1

##### **microbiological organisms**

particles characterized by their ability to form viable colonies

NOTE These can be identified as bacteria, yeast or fungi.

#### 3.2

##### **number of viable micro-organisms**

number of micro-organisms having a potential for metabolic activity

#### 3.3

##### **culturable number**

number of micro-organisms, single cells or aggregates able to form colonies on a solid nutrient medium

#### 3.4

##### **colony-forming unit**

##### **CFU**

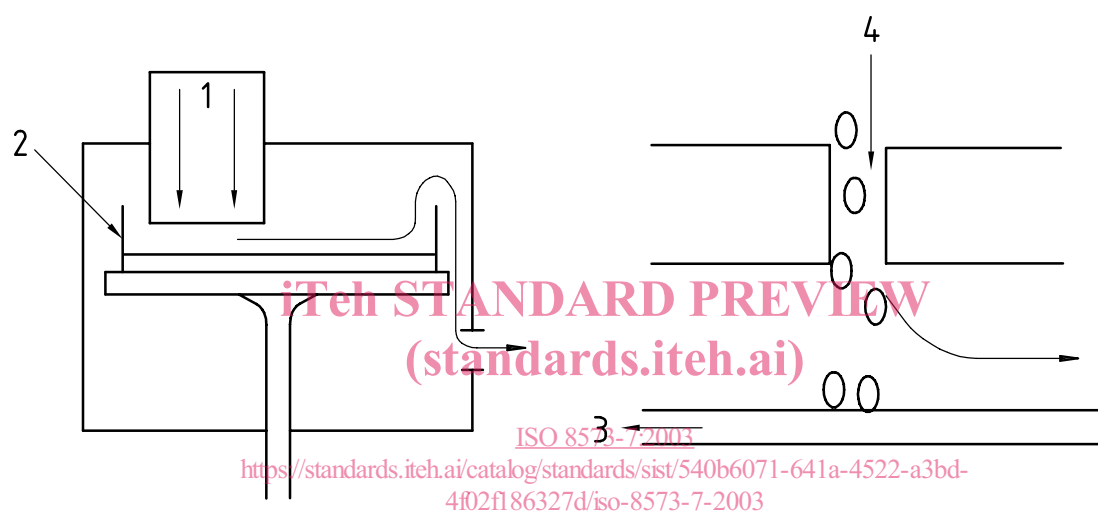
unit by which the culturable number is expressed

#### 4 Method for verifying presence of viable micro-organisms by partial flow sampling

The method for verifying the presence of viable micro-organisms is to expose an agar nutrient to the compressed air sample. A quantitative assessment may be made by the method given in Annex B. See Annex D for details on the preparation of an agar plate with a culturable media.

For partial-flow sampling, a slit-sampler, a type of impaction air tester (see Figure 1), shall be used together with the method given in ISO 8573-4. Isokinetic sampling of the air shall be carried out and reduced until it is within the range of the sampler as identified by the manufacturer. Pressure reduction to atmospheric conditions and flow measurements shall be performed in order to establish compatibility with the manufacturer's recommendations or in accordance with ISO 8573-4. Where the flow is known, the time for the exposure of the agar media to the compressed air sample shall be recorded.

To assist in discriminating non-microbiological from microbiological particles, measurements shall be taken within 4 h.



#### Key

- 1 air intake
- 2 rotating Petri dish with agar
- 3 air outlet
- 4 air

**Figure 1 — Slit-sampler**

It is necessary to eliminate, as far as possible, the influence of liquids on particle size and number so that a correct reading can be obtained. The influence of water shall not be reduced by heating or drying of air, where this might otherwise have been appropriate, in order to avoid influencing the viability of microbiological organisms.

The influence of liquids other than water shall be given due consideration.

#### 5 Operating conditions

Actual operating conditions shall be described in the test report (see Annex A).

## 6 Determination of viable, colony-forming organisms

After incubation of the sample on the agar nutrient (see B.3), the surface shall be visually examined to confirm the presence of viable, colony-forming micro-organisms.

## 7 Test report statement

A statement shall be made in the test report, supplementary to the statement in accordance with ISO 8573-4 for solid particles, providing confirmation that there are viable, colony-forming microbiological particles among the solid particles.

This phrase “Declared sterility of the compressed air in accordance with ISO 8573-1”, shall be followed by

- “Sterile” or “non-sterile”,
- date of sampling,
- date of measurements, and
- location.

Annex A presents a sample test report.

**iTeh STANDARD PREVIEW**  
**(standards.iteh.ai)**

ISO 8573-7:2003

<https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003>

## Annex A (informative)

### Determination of viable microbiological particle content in compressed air — Sample test report

Once the solid particle content in accordance with ISO 8573-4 has been established, a tabulated test report (see Figure A.1) is used to identify those particles present as viable microbiological CFUs in a sample of air taken from the compressed air system under investigation.

NOTE For information on agar media, see B.3

[Value of actual, average measured value (see Annex B) for ...]	
Microbiological organism	CFU/m <sup>3</sup> given at reference conditions <sup>a</sup>
Bacteria	100
Yeast	14
Fungi	No indication
Endoro-bacteria	50
Pressure to which the measurement refers	... MPA [= bar (e)]
[Statement regarding the applicable uncertainty ... (see Clause 7)]	
Date of calibration record	yyyy/mm/dd
<sup>a</sup> Reference conditions: <a href="https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003">https://standards.iteh.ai/catalog/standards/sist/540b6071-641a-4522-a3bd-4f02f186327d/iso-8573-7-2003</a> Temperature 20 °C; Pressure 0,1 MPa (= 1 bar). Relative humidity does not affect volume in this application.	

Figure A.1 — Sample test report



## Annex B (normative)

### Quantitative sampling method

#### B.1 Sampling with slit-sampler (see Figure 1)

##### B.1.1 Principle

The principle of capturing micro-organisms with a slit-sampler (impaction air tester) is both simple and reliable. Air from a compressed air installation is channelled through a specially designed connecting link and accelerated through a narrow slit towards a moist agar surface. The micro-organisms, due to their weight, are flung into the agar surface, whereas the air molecules are deflected. Suitably incubated, they multiply into colonies, which are counted on the assumption that one micro-organism gives rise to one colony.

The slit-sampler can be used for bacteria, yeast or fungi and, with special methods, for viruses and bacteriophages. As a large agar surface (e.g. 140 mm Petri dish) rotates under a radially positioned slit (0,5 mm), a large number of colonies, i.e. organisms, can be counted.

##### B.1.2 Aseptic techniques

The sampling methodology is covered by the adoption of aseptic techniques. The use of a disinfecting agent such as 70 % ethanol is recommended. In periods when the slit-sampler is not in use (stored) precautions shall be taken to avoid the growth of micro-organisms in the equipment. All operations in which the test equipment is to be opened should be carried out with the minimum of delay in order to avoid possible ingress of contaminants from the local environment. Precautions should also be taken against the effects of draughts.

#### B.2 Sampling procedure

The following procedure shall be used for sampling.

- a) Sterilize all sampling equipment by disinfecting the equipment, including tubes and hoses, with a suitable cleaning agent immediately before use.
- b) Allow a test sample to pass through the sampling equipment and associated tubes and hoses without the Petri dish and agar. This is done to allow evaporation of the disinfecting agent and to adjust the slit-sampler.
- c) Perform a blind test before, and after, the actual measurement by carrying out steps d) to f) without starting the slit-sampler. The dishes used shall not subsequently show growth.
- d) Take a 14 cm Petri dish with agar. The Petri dish shall have a label fixed to the bottom with traceability information (date, time of start, test site address, code, etc). Indicate the starting position and the direction of rotation.
- e) Ensure that the air inlet and level indicator of the slit-sampler are turned up. Lift the lid of the slit-sampler and ensure that the plate holder is placed correctly in relation to the micro-switch. Wipe the internal sides of the slit-sampler with a disinfecting pad.
- f) Insert the Petri dish in the slit-sampler, which should be exposed so that the radial line is situated directly under the air inlet slit. Remove the lid and store it in a sterile plastic bag.