# INTERNATIONAL STANDARD

# ISO 16000-20

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# Indoor air —

Part 20:

# Detection and enumeration of mouldsDetermination of total spore count

Air intérieur —

Partie 20: Détection et dénombrement des moisissures — Détermination du nombre total de spores





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## **Foreword**

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For an explanation on the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the WTO principles in the Technical Barriers to Trade (TBT) see the following URL: Foreword - Supplementary information

The committee responsible for this document is ISO/TC 146, *Air quality*, Subcommittee SC 6, *Indoor air*.

ISO 16000 consists of the following parts, under the general title *Indoor air*:

- Part 1: General aspects of sampling strategy
- Part 2: Sampling strategy for formaldehyde
- Part 3: Determination of formaldehyde and other carbonyl compounds in indoor air and test chamber air — Active sampling method
- Part 4: Determination of formaldehyde Diffusive sampling method
- Part 5: Sampling strategy for volatile organic compounds (VOCs)
- Part 6: Determination of volatile organic compounds in indoor and test chamber air by active sampling on Tenax TA sorbent, thermal desorption and gas chromatography using MS or MS-FID
- Part 7: Sampling strategy for determination of airborne asbestos fibre concentrations
- Part 8: Determination of local mean ages of air in buildings for characterizing ventilation conditions
- Part 9: Determination of the emission of volatile organic compounds from building products and furnishing — Emission test chamber method
- Part 10: Determination of the emission of volatile organic compounds from building products and furnishing — Emission test cell method
- Part 11: Determination of the emission of volatile organic compounds from building products and furnishing — Sampling, storage of samples and preparation of test specimens
- Part 12: Sampling strategy for polychlorinated biphenyls (PCBs), polychlorinated dibenzo-p-dioxins (PCDDs), polychlorinated dibenzofurans (PCDFs) and polycyclic aromatic hydrocarbons (PAHs)

- Part 13: Determination of total (gas and particle-phase) polychlorinated dioxin-like biphenyls (PCBs) and polychlorinated dibenzo-p-dioxins/dibenzofurans (PCDDs/PCDFs) Collection on sorbent-backed filters
- Part 14: Determination of total (gas and particle-phase) polychlorinated dioxin-like biphenyls (PCBs) and polychlorinated dibenzo-p-dioxins/dibenzofurans (PCDDs/PCDFs) Extraction, clean-up and analysis by high-resolution gas chromatography and mass spectrometry
- Part 15: Sampling strategy for nitrogen dioxide ( $NO_2$ )
- Part 16: Detection and enumeration of moulds Sampling by filtration
- Part 17: Detection and enumeration of moulds Culture-based method
- Part 18: Detection and enumeration of moulds Sampling by impaction
- Part 19: Sampling strategy for moulds
- Part 20: Detection and enumeration of moulds Determination of total spore count
- Part 21: Detection and enumeration of moulds Sampling from materials
- Part 23: Performance test for evaluating the reduction of formaldehyde concentrations by sorptive building materials
- Part 24: Performance test for evaluating the reduction of volatile organic compound (except formaldehyde) concentrations by sorptive building materials
- Part 25: Determination of the emission of semi-volatile organic compounds by building products Micro-chamber method
- Part 26: Sampling strategy for carbon dioxide (CO<sub>2</sub>)
- Part 27: Determination of settled fibrous dust on surfaces by SEM (scanning electron microscopy) (direct method)
- Part 28: Determination of odour emissions from building products using test chambers
- Part 29: Test methods for VOC detectors
- Part 30: Sensory testing of indoor air
- Part 31: Measurement of flame retardants and plasticizers based on organophosphorus compounds Phosphoric acid esters
- Part 32: Investigation of constructions on pollutants and other injurious factors Inspection

## The following parts are under preparation:

- Part 33: Determination of phthalates with GC-MS
- Part 34: Strategies for the measurement of airborne particles (PM 2,5 fraction)
- Part 35: Measurement of polybrominated diphenylether, hexabromocyclododecane and hexabromobenzene
- Part 36: Test method for the reduction rate of airborne bacteria by air purifiers using a test chamber

## Introduction

Mould is a common name for filamentous fungi from different taxonomic groups (Ascomycota, Zygomycota, and their anamorphic states former known as Deuteromycota or fungi imperfecti). They form a mycelium and spores by which they become visible macroscopically. Most spores are in the size range of 2 µm to 10 µm, some up to 30 µm and only few up to 100 µm. Spores of some mould genera are small and become airborne very easily (e.g. Aspergillus, Penicillium) while others are bigger and/or embedded in a slime matrix (e.g. Stachybotrys, Fusarium) and less mobile.

Mould spores are widely distributed in the outdoor environment and, therefore, occur in varying concentrations also indoors. Growth of moulds in indoor environments, however, has to be considered a hygienic problem because epidemiological studies have revealed that dampness and/or mould growth in homes and health problems affecting the occupants are closely related.

Harmonized methods for sampling, detection and enumeration of moulds including standards for sampling strategies are important for comparative assessment of mould problems indoors. Before doing any measurements a plan for the measurement strategy should be made.

This part of ISO 16000 describes methods for air sampling of mould spores for subsequent microscopic analysis.

This part of ISO 16000 is based on parts of VDI 4300 Part 10. [6]

## Indoor air —

## Part 20:

# **Detection and enumeration of moulds — Determination of total spore count**

## 1 Scope

This part of ISO 16000 specifies requirements for sampling of moulds from air. Following the instructions given, samples are obtained for microscopy to determine the total concentration of spores.

## 2 Terms and definitions

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

## 2.1

## cultivation

<air quality> growing of microorganisms on culture media

[SOURCE: ISO 16000-16:2008, 3.6]

## 2.2

## cut-off value

particle size (aerodynamic diameter) for which the sampling efficiency is 50 %

### 2.3

## filamentous fungus

fungus growing in the form of filaments of cells known as hyphae

[SOURCE: ISO 16000-16:2008, 3.3]

Note 1 to entry: The term "filamentous fungi" differentiates fungi with hyphal growth from yeasts.

## 2.4

## impaction

sampling of particles suspended in air by inertial separation on a solid surface

### 2.5

## microorganism

any microbial entity, cellular or non-cellular, capable of replication or of transferring of genetic material or entities that have lost these properties

[SOURCE: EN 13098:2000]

## 2.6

### mould

<air quality> filamentous fungi from several taxonomic groups; namely Ascomycota, Zygomycota, and their anamorphic states former known as Deuteromycota or fungi imperfecti

[SOURCE: ISO 16000-16:2008, 3.9]

Note 1 to entry: Moulds form different types of spores depending on the taxonomic group they belong to, namely conidiospores (conidia), sporangiospores, or ascospores.

## ISO 16000-20:2014(E)

## 2.7

## mycelium

branched hyphae network

[SOURCE: ISO/TS 10832:2009, 3.5]

## physical sampling efficiency

capacity of the sampler to collect particles with specific sizes suspended in air

Note 1 to entry: See Reference [7].

#### Principle of method 3

A defined air quantity is drawn through an impactor containing a sticky solid surface which can subsequently be used for microscopy. The particles in the air stream impact on the surface, due to their inertia, when the air flows bend to bypass the solid surface.

Airborne moulds are thereby collected directly on the sticky surface.

Physical sampling efficiency is influenced by the geometry of the slit, air velocity, and the adhesion capability of the surface.

The sampling device is constructed for the detection of particles in the size of mould spores (>1 µm to ca. 30 µm). To achieve this, the cut-off value of the sampling device should preferably be 1 µm or less and shall not be more than 2,6 µm.

Three main types of impactors are widely used and available commercially: samplers with replaceable slides and air velocity of ca. 30 L/min, e.g. PS 30 and MBASS30, samplers<sup>1)</sup> with replaceable slides and air velocity of ca. 15 L/min, e.g. Allergenco MK3<sup>2</sup>), and samplers with disposable cassettes and air velocity of ca. 15 L/min (see Annex A).

After sampling, the mould spores are counted under a microscope. No cultivation is performed. Therefore, the total spore concentration, including culturable and non-culturable spores can be determined.

## **Apparatus and materials**

Usual microbiological laboratory equipment, and in particular:

- **Stand**, for positioning the impactor at the sampling height needed. 5.1
- **Impactor**, with disposable slides or cassettes. 5.2
- 5.3 **Vacuum pump**, for ensuring a constant flow rate during continuous operation.
- Gas volume meter, for determining the gas volume sucked at the sampling head, in operating 5.4 cubic meters.
- 5.5 **Timer**, for presetting the time and duration of sampling.
- **Protective housing,** for protecting the impactor from harmful environmental conditions (optional, 5.6 mainly for outdoor use).

<sup>1)</sup> PS 30 and MBASS30 are examples of suitable products available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

Allergenco MK3 is an example of a suitable product available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

**5.7 Microscope**, equipped with 40× and 100× objectives for ca. 400× and 1000× magnification.

## 5 Reagents

## 5.1 General

All reagents and chemicals shall be of recognized quality "for microbiology" or better. Water used shall be distilled or of equivalent quality.

## 5.2 Lactophenol blue solution

The components of the staining solution are listed in Table 1.

WARNING — Lactophenol blue solution is toxic and can lead to adverse health reactions. Exposure through direct contact or inhalation has to be avoided.

Table 1 — Composition of staining solution

Component	Quantity
Cotton blue	0,5 g
Lactic acid (% by mass of 80 % to 85 %)	4,0 g
Phenol	4,0 g
Glycerol	8,0 g
Distilled water	100 ml

Add ingredients in 100 ml water and dissolve.

## 6 Measurement procedure

## 6.1 Sampling

Sampling is usually conducted at a height of 0,75 m to 1,5 m above ground. For special questions, other heights might be applicable. Take care when sampling at low heights that no settled house dust is sucked in the sampling device.

Prepare the required number of impactors and slides or cassettes in accordance with the measurement task and the measurement strategy.

NOTE If the concentration of spores cannot be anticipated, several volumes (e.g 50 L, 100 L, and 200 L) can be sampled and the most suitable sampling (enough spores, not overloaded) can then be used counting.

Check the equipment for completeness and functionality with a check list. Perform function control at regular intervals. Function control implies primarily the volumetric flow control (see <u>Clause 8</u>).

Use a sterile device containing the slides or cassettes for each measurement point. Alternatively, clean the slit with ethanol or isopropanol (70 %; volume fraction) and dry it afterwards (e.g. with compressed air).

Place the slides or cassettes in the impactors. Take care to avoid contamination.

Start the sampling device in accordance with the manufacturer's operating instructions. Multiple measurements using different sampling volumes are recommended. This is especially important when the level of the anticipated concentration of moulds is not known.

After sampling, remove the slides or cassettes from the sampling apparatus and pack them in sterile containers and/or plastic bags in order to avoid any secondary contamination. Subsequently, draw air

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through the impactor without slides for several minutes at the new sampling point prior to the sampling with the slide in place.

For disposable samplers, follow the instructions of the manufacturer.

Fill in a sampling protocol (see <u>Clause 9</u> and <u>Annex B</u>).

Transport samples to the laboratory (see 6.4) and analyse by direct microscopy (see 6.2).

## 6.2 Direct microscopy

The spores on the sticky surface are stained with e.g. lactophenol blue solution (see 5.2) or cotton blue in lactic acid and evaluated under the microscope at 400× and 1000× magnification.

To get an overview of the sample trace, it can be useful to make an initial view at 100× or 200× magnification.

If different volumes have been sampled, choose the most appropriate (enough spores, not overloaded) for counting.

Counting of the different spore types under the microscope is a difficult task which can only be performed by skilled and well trained personnel.

Lactophenol blue solution is toxic and has to be avoided whenever possible. Staining of spores with alternative staining solution has to be anticipated. Cotton blue in lactic acid does, however, not work with the PS 30 with MBASS30<sup>3)</sup> which was used for determining the performance characteristics (see Clause 10).

The slit impaction sampling method produces a sample trace approx. 1,6 cm long and approx. 1 mm wide on the glass slide which is evaluated by microscopy.

Normally, the slides are evaluated for the following spore types: Basidiospores, Ascospores, Cladosporium, type Aspergillus/Penicillium, Stachybotrys, Chaetomium, type Alternaria/Ulocladium, type Helminthosporium, Epicoccum, other spores, and mycelial fragments. Additional spore types which occur in unusual concentrations and can be assigned to a morphological type are likewise reported.

Basidiospores, ascospores, spores of the types Alternaria/Ulocladium, Helminthosporium, and Epicoccum do not usually originate from indoor sources and therefore provide an indication as to what extent the sample has been influenced by outdoor air (e.g. due to leaks around windows, mechanical disturbance of settled outdoor spores).

Large, readily recognizable spores, e.g. Stachybotrys or Chaetomium spores, are counted on the whole surface at 400× magnification in longitudinal direction of the sample trace. In this way, the complete loaded impaction surface can be evaluated within a reasonable time scale and hence, few spores determined per sample volume. The theoretical detection limit is one spore in the volume sampled.

For small spores of the type, e.g. Aspergillus/Penicillium, an additional detailed evaluation is made at 1000× magnification in the direction perpendicular to the sample trace. Detailed evaluations are very time-consuming so that only a small portion of the sample (usually approx. 10 % to 30 % of the total impaction surface, see <u>Clause 10</u>) can be evaluated. Consequently, the detection limit for small spores is higher than that for large spores. For a sample volume of 200 L and the evaluation of 10 transverse traverses, the theoretical detection limit is approx. 50 spores/m<sup>3</sup> (one spore present in the 10 traverses).

For quantification, preferably, 10 or more spores should be present in the microscopic area counted.

Particles like skin scales and other (mineral and organic) particles are not counted, but might be indicated in categories like low, medium, relatively high and high to have an indication of the influence of corresponding activities in the room.

<sup>3)</sup> PS 30 and MBASS30 are examples of suitable products available commercially from Umweltanalytik Holbach. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

## 6.3 Calculation and expression of results

The quantitative result is reported as concentration of the enumerated spore types and mycelial fragments per  $m^3$  of air. The total spore concentration is obtained by summating the concentrations of the individual spore types.

NOTE Spores present in aggregates are enumerated individually. It is, however, helpful to document the presence of aggregates in the analysis report so as to provide information on how many individual spores and how many aggregate spores were present, which might be needed for specific investigation objectives.

In the  $400\times$  evaluation (see 6.2), which is performed in the longitudinal direction of the sample trace, the entire sample volume is evaluated. For the calculation of the spore concentration per m<sup>3</sup> of air, the result per sample volume is multiplied by a corresponding factor (e.g. F = 5 for a sample volume of 200 L, see example 1).

In the detailed 1  $000 \times$  evaluation, which is performed in the direction perpendicular to the sample trace (see <u>6.2</u>), parameters like the slit geometry, the size of the objective visual field and the number of transverse traverses evaluated enter into the calculation. Results are rounded to two decimal numbers.

The concentration can be calculated according to Formula (1).

$$C_{\rm L} = \frac{L}{B} \cdot \frac{1}{V_{\rm G}} \cdot Z \tag{1}$$

where

 $C_{\rm L}$  is the concentration of the air sample in spores/m<sup>3</sup>;

L is the total length of the sample trace in mm;

*B* is the length of the evaluated area of the sample trace in mm;

 $V_{\rm G}$  is the sample volume in m<sup>3</sup>;

Z is the total spore count.

*B* is calculated according to Formula (2).

$$B = D \times Z_0 \tag{2}$$

where

*D* is the diameter of the visual field in mm;

 $Z_0$  is the number of evaluated transverse traverses.

EXAMPLE 1 20 spores of *Stachybotrys* were detected on the whole sampling surface. The sample volume was 200 L.

Result:  $20 \times 5 = 100 \text{ spores/m}^3 \text{ air} = 1,0 \times 10^2 \text{ spores/m}^3 \text{ air}$ 

EXAMPLE 2 A sample was collected using a slit impactor (16 mm long, 1,1 mm wide) and evaluated at 1 000× magnification. The visual field of the microscope and objective used has a diameter of 175  $\mu$ m. The sample volume was 200 L. 20 transverse traverses were evaluated with a count of 60 spores of the type Aspergillus/Penicillium.

$$C_{\rm L} = \frac{16}{0.175 \cdot 20} \cdot \frac{1}{0.2} \cdot 60 = 1371 \text{ spores/m}^3 \text{ air}$$

Result:  $1.4 \times 10^3$  spores/m<sup>3</sup> air

## 6.4 Transport and storage

Pack slides or cassettes into sterile containers and/or bags. Protect them from disturbing impacts (humidity, desiccation, heat, dust, etc.) and transport them to the laboratory for analysis. Process samples, preferably within one week after sampling. Store samples in the laboratory at room temperature protected from desiccation until further processing.

## 7 Quality assurance

The laboratory shall implement quality assurance measures to be documented and made available any time.

## 8 Calibration of flow rate, function control, and maintenance of the sampling system

Calibration of the sampling device shall be performed by means of a certified reference volume meter having a measurement accuracy of  $\leq \pm 5$  % expressed in operational cubic meters, referenced to ambient air conditions. The reference volume meter shall be connected to the air inlet of the sampling apparatus. The air inlet orifice of the reference apparatus shall unobstructed. After a successful adjustment of the flow rate, the display accuracy of the sampling apparatus has to be checked against the reference volume meter. The air volume sucked through the sampling apparatus for a duration of 30 min shall be indicated with an accuracy of  $\pm 1$  % compared with the reference volume meter.

The usual verification of the flow rate (function control) depends on the stability of the apparatus. A complete calibration shall be carried out prior to starting a new measurement campaign or following significant changes, e.g. when new or repaired equipment is used or after pump servicing. If the flow rate determined using the transfer standard deviates more than 2 % from the value required for correct operation of the inlet, the flow controller shall be adjusted according to the manufacturer's instructions. Make sure that the air flow is not fluctuating more than  $\pm 2$  % during sampling and that the time to reach the desired sampling velocity in the beginning of the sampling process is kept as short as possible to minimize the influence on the sample volume.

For some sampling devices, verification and adjustment of the nominal flow is not possible by the user but is conducted by the manufacturer in regular intervals. In this case, a constant flow between calibration intervals has to be guaranteed by the manufacturer and the apparatus has to have an internal control system preventing deviations from the nominal flow.

## 9 Sampling protocol

The samples shall be uniquely identified and labelled accordingly.

A sampling protocol shall be filled in for each sample before (or just after) sampling.

The protocol shall as minimum indicate

- a) a reference to this standard, i.e. ISO 16000-16,
- b) the name and address of the client,
- c) the measurement task,
- d) the type of sampling device used,
- e) the air volume sampled, and
- f) the sampling date, hour, and location of sampling.

## 10 Performance characteristics

Performance characteristics have been determined using the PS 30 with MBASS30<sup>4)</sup> by Umweltmykologie Dr. Dill & Dr. Trautmann GbR, Germany with a research grant of the Federal Environment Agency.

The physical sampling efficiency was determined at different air velocities (15 L/min, 22,5 L/min, 30 L/min, and 45 L/min) by filtration of the air at the outlet of the slit impactor to detect non impacted fungi. Sampling efficiency increased with increasing air velocity from 15 L/min to 30 L/min. No further increase was detected for an air velocity of 45 L/min. Sampling efficiency was higher for larger spores than for smaller spores. For Cladosporium, sampling efficiencies around 90 % (see Figure 1) were reached while for Aspergillus/Penicillium sampling efficiencies were only 30 % to 80 %. In further experiments, air velocities of 30 L/min were used. Comparable sampling efficiencies have been reported for the Air-O-Cell sampling head<sup>5)</sup> (see Reference [7]).

The slit impaction of the PM 30 sampler<sup>6)</sup> produces a trace approx. 1,6 cm long and 1,1 cm in diameter.

With a 1 000× magnification, about 94 microscopic fields lay within this 1,6 cm trace in longitudinal direction. The distribution of large (*Chaetomium* or *Stachybotrys*) and small spores (Aspergillus/Penicillium) at different concentrations was determined using pure cultures. At high concentrations, spores were found in all microscopic fields but very low numbers were detected in fields at the beginning and at the end of the trace (see Figures 2 and 3). At low concentrations of spores in the air sample, many microscopic fields did not contain any spores and the effect at the beginning and at the end of the trace was even more pronounced (see Figures 4 and 5). Counting of small spores is usually done with 1 000× magnification on a selected number of these microscopic fields. Because of the effects in the beginning and the end of the trace, these areas should not be included in the counting.

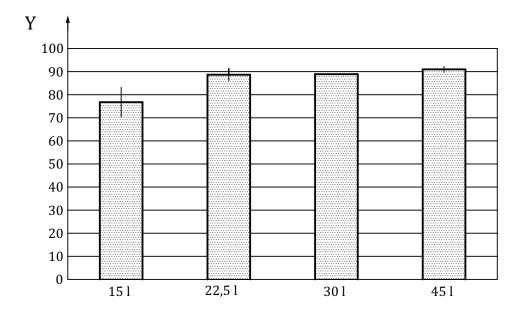
Further experiments were conducted to determine the measurement uncertainty if only some (5, 10, 20, or 30) of the 94 microscopic fields are examined and the results calculated from these counts. Measurement uncertainty decreased with increasing concentration of spores and increasing number of microscopic fields counted (see <u>Table 2</u>). To achieve a measurement uncertainty of below 10 %, at least 30, 20, 10, or 5 microscopic fields have to be counted with spore numbers of around 100, 200, 1 000, and 2 000 per 200 L air, respectively.

The suitability of the method under field conditions was tested by comparative measurements using PS30 and MBASS30<sup>4)</sup> from Umweltanalytik Holbach (see <u>Annex B</u>).

<sup>4)</sup> PS 30 and MBASS30 are examples of suitable products available commercially from Umweltanalytik Holbach. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

<sup>5)</sup> Air-O-Cell is an example of a suitable product available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

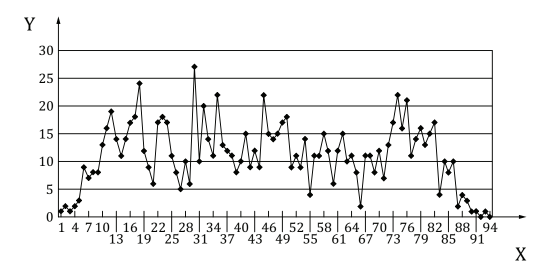
<sup>6)</sup> PM 30 is an example of a suitable product available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.



## Key

sampling efficiency in %

Figure 1 — Sampling efficiency for larger spores (Cladosporium) at different air velocities

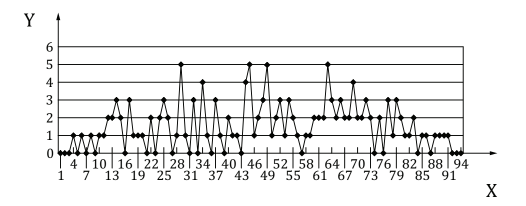


## Key

X transect number

number of spores

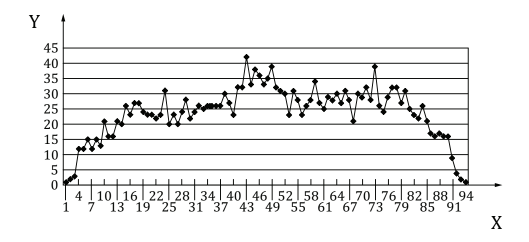
Figure 2 — Distribution of 1 038 spores of Stachybotrys chartarum in 94 transects along the sample trace



## Key

- X transect number
- Y number of spores

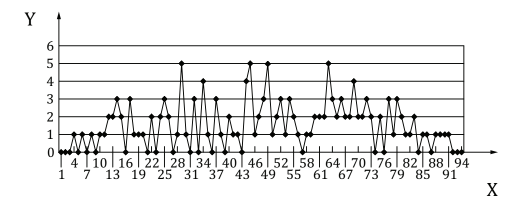
Figure 3 — Distribution of 2 250 spores type *Aspergillus/Penicillium* in 94 transects along the sample trace



## Key

- X transect number
- Y number of spores

Figure 4 — Distribution of 214 spores of *Chaetomium* in 94 transects along the sample trace



## Key

X transect number

number of spores

Figure 5 — Distribution of 144 spores type Aspergillus/Penicillium in 94 transects along the sample trace

Table 2 — Mean number of spores and standard deviation counted in different numbers of microscopic fields in four samples with different number of spores

		Concentration of spores in 200 L air		Concentration of spores in 200 L air		Concentration of spores in 200 L air		Concentration of spores in 200 L air	
Sample number	5 microscopic fields counted		10 microscopic fields counted		20 microscopic fields counted		30 microscopic fields counted		
	Mean	SD %	Mean	SD %	Mean	SD %	Mean	SD %	
1	133	34	122	24	120	14	126	9	
2	248	17	255	11	248	7	251	4	
3	1031	16	1 019	8	1 016	5	1 020	6	
4	2 239	6	2 283	4	2 260	3	2 277	2	

NOTE 1 Sample 1 and 4: Spores of Aspergillus or Penicillium.

NOTE 2 Sample 2: Chaetomium sp.

NOTE 3 Sample 3: Stachybotrys chartarum.

## Annex A

(informative)

# **Examples of impactors**

Table A.1 — Slit impactors for the determination of the total spore count in indoor air

Sampling system/ volumetric flow rate	Recommended sampling period <sup>5)</sup>	Sample volume	Cut-off (d <sub>50</sub> )	Sample preparation/ analysis method	Resulting evaluation range in spores (mycelial fragments)/m <sup>3</sup>
Sampler with replaceable slides, approx. 30 l/min e.g. PS30 and MBASS30 <sup>f</sup>	approx. 5 min to 7 min	0,15 m <sup>3</sup> to 0,2 m <sup>3</sup>	1,8 μm <sup>a b</sup>	Staining and light microscopy; enumeration of spore types (genera and/or genus groups)	50 to 100 000 <sup>c d</sup>
Sampler with replaceable slides approx. 15 l/min e.g. Allergenco MK3g	5 min to 10 min	0,075 m <sup>3</sup> to 0,15 m <sup>3</sup>	No data avail- able <sup>a b</sup>	Staining and light microscopy; enumeration of spore types (genera and /or genus groups)	50 to 100 000cd
Disposable cassette/ approx. 15 l/min e.g. Air-O-Cell® cassetteh, Allergenco-D cassettei	5 min to 10 min	0,075 m <sup>3</sup> to 0,15 m <sup>3</sup>	1,8 μm to 2,6 μm <sup>b c</sup>	Staining and light microscopy; enumeration of spore types (genera and /or genus groups)	50 to 100 000 <sup>c d</sup>

- a The collection efficiency depends on the selected sampling medium (adhesion, viscosity).
- b The collection efficiency depends on the configuration of the outer spore envelope (spore-to-sampling medium contact).
- The collection efficiency depends on the configuration of the disposable cassette.
- d The evaluation range applies to detailed evaluations. Here, the lower value is dependent upon the number of detailed evaluations (see 6.2 and 6.3); the upper value achievable in measurement practice varies greatly because spores, skin scales, other particles, etc. trapped on the impaction surface will restrict the adhesion of further spores. An overview evaluation of the complete sample trace, which makes only sense for spores with distinct morphological features (e.g. *Stachybotrys, Chaetomium*), allows the detection of a single spore on the entire impaction surface, i.e. the entire sample volume. Detection of single spores does, however, not enable a quantitative assessment.
- e Based on the sample volume and the flow velocity as recommended by the manufacturer.
- f PS 30 and MBASS30 are examples of suitable products available commercially from Umweltanalytik Holbach. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.
- g Allergenco MK3 is an example of a suitable product available commercially from Blewstone. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.
- h Air-O-Cell® is an example of a suitable product available commercially from Zefon. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.
- i Allergenco-D is an example of a suitable product available commercially from Environmental Monitoring Systems (EMS). This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

## Annex B

(normative)

# Sample exchange for method validation

The suitability of the sampling and analysis method described in this part of ISO 16000 was tested under field conditions in a small trial with four laboratories. Four samples were taken in parallel with four different samplers [all PS 30 with MBASS30<sup>7</sup>]. Each sample was consecutively analysed by two of the four participating laboratories. The results of the total spore concentrations obtained by sampling with different samplers and counted in different experienced laboratories were in good agreement (see Table B.1).

Table B.1 — Data of sample exchange

Lab	Sample number total spore count/m <sup>3</sup>									
number	1 A	1 B	2 A	2 B	3 A	3 B	4 A	4 B		
1		9,7 × 10 <sup>4</sup>		1,2 × 10 <sup>5</sup>		1,1 × 10 <sup>5</sup>		8,8 × 10 <sup>4</sup>		
2	8,1 × 10 <sup>4</sup>	9,7 × 10 <sup>4</sup>	9,2 × 10 <sup>4</sup>	8,6 × 10 <sup>4</sup>	1,1 × 10 <sup>5</sup>	7,1 × 10 <sup>4</sup>	8,5 × 10 <sup>4</sup>	7,4 × 10 <sup>4</sup>		
3	1,2 × 10 <sup>5</sup>		1,6 × 10 <sup>5</sup>		1,1 × 10 <sup>5</sup>		1,1 × 10 <sup>5</sup>			

<sup>7)</sup> PS 30 and MBASS30 are examples of suitable products available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by ISO of this product.

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