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Second edition 2022-01

Pore size distribution and porosity of solid materials by mercury porosimetry and gas adsorption —

Part 2: **Analysis of nanopores by gas adsorption**

Distribution des dimensions des pores et porosité des matériaux solides par porosimétrie au mercure et par adsorption de gaz —

Partie 2: Analyse des nanopores par adsorption de gaz

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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.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT), see www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 24, *Particle characterization including sieving*, Subcommittee SC 4, *Particle characterization*.

This second edition cancels and replaces ISO 15901-2:2006 and ISO 15901-3:2007, which have been technically revised. It also incorporates the Technical Corrigendum ISO 15901-2:2006/Cor.1:2007.

The main changes compared to the previous edition are as follows:

- the analysis of nanopores by gas adsorption which combines the characterization of both microand mesopores is now addressed;
- the classification of adsorption isotherms and hysteresis loops has been updated.

A list of all parts in the ISO 15901 series can be found on the ISO website.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

Introduction

In general, different types of pores may be pictured as apertures, channels, or cavities within a solid body or as the space (i.e. interstices or voids) between solid particles in a bed, compact or aggregate. Porosity is a term which is often used to indicate the porous nature of solid material and is more precisely defined as the ratio of the volume of accessible pores and voids to the total volume occupied by a given amount of the solid. According to the 2015 IUPAC recommendations^[1], nanopores are defined as pores with internal widths of equal or less than 100 nm and are divided into several subgroups dependent on their pore width:

- pores with width greater than about 50 nm are called macropores;
- pores of widths between 2 nm and 50 nm are called mesopores;
- pores with width of about 2 nm and less are called micropores;

Further, IUPAC suggested a subclassification of micropores into supermicropores (pore width 0,7 nm to 2 nm), and ultramicropores (pore width < 0,7 nm). In addition to the accessible pores, a solid may contain closed pores which are isolated from the external surface and into which fluids are not able to penetrate. The characterization of closed pores, i.e. cavities with no access to an external surface, is not covered in this document.

Porous materials may take the form of fine or coarse powders, compacts, extrudates, sheets or monoliths. Their characterization usually involves the determination of the pore size distribution as well as the total pore volume or porosity. For some purposes it is also necessary to study the pore shape and interconnectivity, and to determine the internal and external surface area.

Porous materials have great technological importance, e.g. in the context of the following:

- a) controlled drug release;
- b) hcatalysis; hcatalysis; hcatalog/standards/sist/fc10be6d-f321-4a41-9951-2857a71d01ba/iso-
- c) gas separation;
- d) filtration including sterilization;
- e) materials technology;
- f) environmental protection and pollution control;
- g) natural reservoir rocks;
- h) building material properties;
- i) polymer and ceramic industries.

It is well established that the performance of a porous solid (e.g. its strength, reactivity, permeability or adsorbent power) is dependent on its pore structure. Many different methods have been developed for the characterization of pore structure. The choice of the most appropriate method depends on the application of the porous solid, its chemical and physical nature and the range of pore size.

Different methods for the characterization of nanopores are available, including spectroscopy, electron and tunnel microscopy and sorption methods. In view of the complexity of most porous solids, it is not surprising that the results obtained are not always in agreement and that no single technique can be relied upon to provide a complete picture of the pore structure. Among these, mercury porosimetry (see ISO 15901-1) and gas adsorption are popular ones because by combining both it is possible to assess a wide range of pore sizes from below 0,5 nm up to 400 μ m. While mercury porosimetry is the standard technique for macropore analysis, gas adsorption techniques allow to assess pores up to approximately 100 nm. In this case, physical adsorption can be conveniently used, is not destructive, and is not that cost intensive as compared to some of the above-mentioned methods. Particularly, with regard to the

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application of microporous material as specific sorbents, molecular sieves and carriers for catalysts and biological active material, the field-proven methods of gas sorption are of special value.

The measuring techniques of the method described in this document are similar to those described in ISO 9277 for the measurement of gas adsorption at low temperature. However, in order to assess the full range of pore sizes including microporosity, adsorption experiments have to be performed over a wide range of pressures from the ultralow pressure range (e.g. turbomolecular pump vacuum) up to atmospheric pressure (0,1 MPa).

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Pore size distribution and porosity of solid materials by mercury porosimetry and gas adsorption —

Part 2:

Analysis of nanopores by gas adsorption

1 Scope

This document describes a method for the evaluation of porosity and pore size distribution by physical adsorption (or physisorption). The method is limited to the determination of the quantity of a gas adsorbed per unit mass of sample as a function of pressure at a controlled, constant temperature [1]-[9]. Commonly used adsorptive gases for physical adsorption characterization include nitrogen, argon, krypton at the temperatures of liquid nitrogen and argon (77 K and 87 K respectively) as well as CO_2 (at 273 K). Traditionally, nitrogen and argon adsorption at 77 K and 87 K, respectively, allows one to assess pores in the approximate range of widths 0,45 nm to 50 nm, although improvements in temperature control and pressure measurement allow larger pore widths to be evaluated. CO_2 adsorption at 273 K – 293 K can be applied for the microporous carbon materials exhibiting ultramicropores. Krypton adsorption at 77 K and 87 K is used to determine the surface area or porosity of materials with small surface area or for the analysis of thin porous films.

The method described is suitable for a wide range of porous materials. This document focuses on the determination of pore size distribution from as low as 0,4 nm up to approximately 100 nm. The determination of surface area is described in ISO 9277. The procedures which have been devised for the determination of the amount of gas adsorbed may be divided into two groups:

- those which depend on the measurement of the amount of gas removed from the gas phase, i.e. manometric (volumetric) methods; 15901-2-2022
- those which involve the measurement of the uptake of the gas by the adsorbent (i.e. direct determination of increase in mass by gravimetric methods).

In practice, static or dynamic techniques can be used to determine the amount of gas adsorbed. However, the static manometric method is generally considered the most suitable technique for undertaking physisorption measurements with nitrogen, argon and krypton at cryogenic temperatures (i.e. 77 K and 87 K, the boiling temperature of nitrogen and argon, respectively) with the goal of obtaining pore volume and pore size information. This document focuses only on the application of the manometric method.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements 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 3165, Sampling of chemical products for industrial use — Safety in sampling

ISO 8213, Chemical products for industrial use — Sampling techniques — Solid chemical products in the form of particles varying from powders to coarse lumps

ISO 9277, Determination of the specific surface area of solids by gas adsorption — BET method

ISO 14488, Particulate materials — Sampling and sample splitting for the determination of particulate properties

3 Terms and definitions

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

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at https://www.iso.org/obp
- IEC Electropedia: available at https://www.electropedia.org/

3.1

adsorbate

adsorbed gas

3.2

adsorption

enrichment of the adsorptive at the external and accessible internal surfaces of a solid

3.3

adsorptive

gas or vapour to be adsorbed

3.4

adsorbent

solid material on which adsorption occurs

3.5

adsorption isotherm

relationship between the amount of gas adsorbed and the equilibrium pressure of the gas at constant temperature

3.6

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adsorbed amount

amount of gas adsorbed at a given pressure, *p*, and temperature, *T*

3.7

equilibrium adsorption pressure

pressure of the adsorptive in equilibrium with the adsorbate

3.8

monolayer amount

amount of the adsorbate that forms a monomolecular layer over the surface of the adsorbent

3.9

monolayer capacity

volumetric equivalent of monolayer amount expressed as gas at standard conditions of temperature and pressure (STP)

3.10

nan opore

pore with width of 100 nm or less

3.11

macropore

pore with width greater than about 50 nm

3.12

mesopore

pore with width between approximately 2 nm and 50 nm

3.13

micropore

pore with width of about 2 nm or less

3.14

supermicropore

pore with width between approximately 0,7 nm and 2 nm

3.15

ultramicropore

pore with width of approximately < 0,7 nm

3.16

physisorption

weak bonding of the adsorbate, reversible by small changes in pressure or temperature

3.17

pore size

pore width, i.e. diameter of cylindrical pore or distance between opposite walls of slit

3.18

pore volume

volume of pores as determined by stated method

3.19

relative pressure an STANDARD PRRV RW

ratio of the equilibrium adsorption pressure, p, to the saturation vapour pressure, p_0 , at analysis temperature

3.20

saturation vapour pressure

vapour pressure of the bulk liquefied adsorptive at the temperature of adsorption

3.21

volume adsorbed

volumetric equivalent of the amount adsorbed, expressed as gas at standard conditions of temperature and pressure (STP), or expressed as the adsorbed liquid volume of the adsorbate

4 Symbols

For the purposes of this document, the following symbols apply, together with their units. All specific dimensions are related to sample mass, in grams.

Symbol	Description	Unit
$A_{\rm a}$	Kirkwood-Mueller constant of adsorptive	J·cm ⁶
$A_{\rm s}$	Kirkwood-Mueller constant of adsorbent	J·cm ⁶
$a_{\rm s}$	specific surface area	m ² ·g ⁻¹
$a_{\rm s,ref}$	specific surface area of reference sample	m ² ·g ⁻¹
$\alpha_{\rm a}$	polarizability of adsorptive	cm ³
$\alpha_{\rm s}$	normalized adsorption	1 ^a
α_{s^*}	polarizability of adsorbent	cm ³
c	speed of light	m·s ^{−1}
$d_{\rm a}$	diameter of an adsorptive molecule	nm
d_{HS}	diameter of hard spheres	nm

^a According to ISO 80000-1, the coherent SI unit for any quantity of dimension one (at present commonly determined "dimensionless") is the unit one, symbol 1.

Symbol	Description	Unit
d_p	pore diameter (cylindrical pore)	nm
$d_{\rm s}$	diameter of an adsorbent molecule	nm
d_0	$d_0 = (d_s + d_a)/2$, distance between adsorptive and adsorbent molecules	nm
$\varepsilon_{\rm ff}/k_{\rm B}$	well depth parameter of gas-gas Lennard Jones potential	K
$\varepsilon_{\rm sf}/k_{\rm B}$	well depth parameter of gas-solid Lennard Jones potential	K
k_{B}	Boltzmann constant (1,381 × 10^{-23})	J K ^{−1}
1	nuclei-nuclei pore width	nm
m_a^*	mass adsorbed	g
$m_{\rm e}$	mass of an electron	kg
$N_{\rm A}$	Avogadro's constant (6,022 × 10 ²³)	mol ⁻¹
N_a	number of atoms per unit area (m ²) of monolayer	m ⁻²
$N_{\rm s}$	number of atoms per unit area (m²) of adsorbent	m ⁻²
$n_{\rm a}$	specific amount adsorbed	mol∙g ⁻¹
P	pressure of the adsorptive in equilibrium with the adsorbate	Pa
p_0	saturation vapour pressure of the adsorptive	Pa
p/p_0	relative pressure of the adsorptive	1 ^a
R	ideal gas constant (8,314)	Jmol ⁻¹ K ⁻¹
$ ho_{ m g}$	gas density The same of the sa	g·cm ⁻³
$ ho_{ m g,STP}$	gas density at STP (273,15 K; 101,3 kPa)	g·cm ⁻³
ρ_1	liquid density	g·cm ⁻³
σ	distance between two molecules at zero interaction energy	nm
$\sigma_{ m ff}$	distance parameter of gas-gas Lennard Jones potential	nm
$\sigma_{ m sf}$	distance parameter of gas-solid Lennard Jones potential	nm
T http	temperature iteh.ai/catalog/standards/sist/fc10be6d-f321-4a41-9951-2857a71d0	kba/iso-
$T_{\rm cr}$	critical temperature 15901-2-2022	K
t	statistical layer thickness	nm
$V_{\rm a}$	specific volume of the adsorbate	cm ³ ·g ⁻¹
$V_{\rm g}$	specific adsorbed gas volume at STP (273,15 K; 101,3 kPa)	cm ³ ·g ⁻¹
$V_{\rm micro}$	micropore volume	cm ³ ·g ⁻¹
W	pore width (slit pore)	nm
$\chi_{\rm a}$	diamagnetic susceptibility of adsorptive	cm ³
$\chi_{\rm s}$	diamagnetic susceptibility of adsorbent	cm ³

 $^{^{}a}$ According to ISO 80000-1, the coherent SI unit for any quantity of dimension one (at present commonly determined "dimensionless") is the unit one, symbol 1.

5 Principles

5.1 General

Physisorption is a general phenomenon and occurs whenever an adsorbable gas (the adsorptive) is brought into contact with the surface of a solid (the adsorbent). The forces involved are the van der Waals forces. Physisorption in porous materials is governed by the interplay between the strength of fluid-wall and fluid-fluid interactions as well as the effects of confined pore space on the state of fluids in narrow pores. The effect of pore width on the interaction potential is demonstrated schematically in Figure $1^{[8]}$.

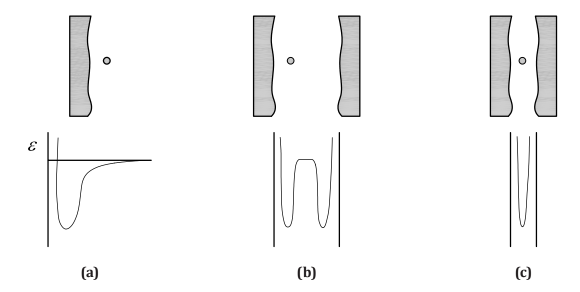


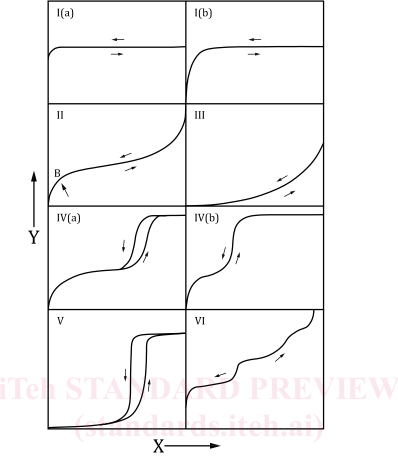
Figure 1 — Schematic illustration of adsorption potential, ϵ , on (a) planar, nonporous surface; (b) mesopore; (c) micropore

The interplay between the strength of attractive adsorptive-adsorbent interactions and the effect of confinement (as controlled by pore size/geometry) affect the shape of adsorption isotherm, as demonstrated in the IUPAC classification of adsorption isotherms^[1] as shown in Figure 2.

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Figure 2 — Standard isotherm types (IUPAC 2015)

Reversible Type I isotherms are given by microporous solids having relatively small external surfaces (e.g. some activated carbons, molecular sieve zeolites and certain porous oxides). Because the pore size is similar to the molecule diameter, the choice of the gas is decisive, i.e. the size of the gas molecule controls the accessibility of the pores, and hence affects the obtained porosity information. A Type I isotherm is concave to the relative pressure (i.e. p/p_0 axis) and the amount adsorbed approaches a limiting value. This limiting uptake is governed by the accessible micropore volume rather than by the internal surface area. A steep uptake at very low p/p_0 is due to enhanced adsorbent-adsorptive interactions in narrow micropores (micropores of molecular dimensions), resulting in micropore filling at very low p/p_0 . Type I(a) isotherms are given by microporous materials having mainly narrow micropores (of width ≤ 1 nm), which includes ultramicropores). A significant portion of the micropores is indicated by a large and steep increase of the isotherm near its origin and subsequent bending to a plateau. Type I(b) isotherms are found with materials having pore size distributions over a broader range including wider micropores (including supermicropores).

Type II isotherms are typically produced by solids which are non-porous or macroporous. Point B is often taken as indicative of the completion of the monolayer capacity.

Type III isotherms are distinguished by a convexity towards the relative pressure axis. These isotherms are found when weak gas-solid interactions occur on non-porous or macroporous solids (e.g. water adsorption on carbon surfaces).

Type IV isotherms are found for mesoporous solids. Type IV isotherms are given by mesoporous adsorbents (e.g. many oxide gels, industrial adsorbents and mesoporous molecular sieves). The