INTERNATIONAL STANDARD



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Medical suction equipment -

Part 2: Manually powered suction equipment iTeh STANDARD PREVIEW

Appareils d'aspiration médicaux) --

Partie 2: Appareils d'aspiration manuels ISO 10079-2:1992

https://standards.iteh.ai/catalog/standards/sist/62025224-ff21-44a7-b307-73adba4fa8da/iso-10079-2-1992



Contents

1	Scope	1
2	Normative reference	1
3	Definitions	1
4	Cleaning, disinfection and sterilization	3
5	Design requirements	3
6	Operational requirements	4
7	Physical requirements	5
8	Performance requirements for vacuum and flow	5
9	Resistance to environment	5
10	Marking	6
11	Information to be supplied by manufacturer iTeh STANDARD PR	EVIEW
Annexes (standards.iteh.ai)		
Α	Test methods	7
в	ISO 10079-2:1992 Table of typical range of volumes for collection containers for 55 specific uses 73adba4ia8da/iso-10079-2-199	224-ff21-44a7-b307- 10
С	Rationale statement	11

Page

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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 Inter-Teh Standard requires approval by at least 75% of the member bodies casting a vote.

(International Standard ISO 10079-2 was prepared by Technical Committee ISO/TC 121, Anaesthetic and respiratory equipment, Sub-Committee SC 8, Suction devices for hospital and emergency care use.

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- Part 1: Electrically powered suction equipment Safety requirements
- Part 2: Manually powered suction equipment
- Part 3: Suction equipment powered from vacuum or pressure source

Annex A forms an integral part of this part of ISO 10079. Annexes B and C are for information only.

Introduction

This International Standard, which has been prepared under the responsibility of Sub-Committee 8 of ISO/TC 121, comprises part 2 of the standard for medical suction equipment, and deals only with manually powered suction equipment. Part 1 deals with safety requirements for electrically powered equipment. Part 3 deals with suction equipment powered from a vacuum or pressure source.

This International Standard has been prepared in response to a need for a safety and performance standard for suction systems. Suction is used to clear the airway and remove unwanted material from body cavities. Suction is also used to assist drainage and decompress body cavities. Suction and vacuum systems are used widely both in health care facilities such as hospitals, for domiciliary care of patients who are nursed at home, and in emergency situations both outside hospitals in field conditions, and during transport in ambulances.

As far as possible, this International Standard has been written specifying performance requirements corresponding with those needed for effective and safe treatment of the patient. https://standards.itch.ai/catalog/standards/sist/62025224-ff21-44a7-b307-

Annex A gives test methods to be used to <u>Verify</u>(compliance/with the requirements given in this part of ISO 10079. Annex B gives a table of a typical range of volumes for collection containers for specific uses. Annex C gives a rationale statement for some requirements.

Medical suction equipment -

Part 2:

Manually powered suction equipment

1 Scope

This part of ISO 10079 specifies safety and performance requirements for manually powered medical suction equipment intended for oro-pharyngeal suction to establish and maintain the patency of the airway. It covers equipment operated by foot or by hand or both (see figure 1). Non-electrical suction equipment which may be integrated with electrical equipment is within the scope of this part. ISO 10079-2:1p)2breast pumps;

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powered suction equipment, whether mains electricity or battery-powered, which is dealt with in ISO 10079-1, nor to suction equipment powered from a vacuum or pressure source which is dealt with in ISO 10079-3, nor to the following:

- a) central power supply (by vacuum/compressed air generation), piping systems of vehicles and buildings, and wall connectors;
- b) catheter tubes, drains, curettes and suction tips;
- c) syringes;
- d) dental suction equipment;
- e) waste gas scavenging systems;
- f) laboratory suction;
- g) autotransfusion systems;
- h) passive urinary drainage;
- i) closed systems for wound drainage;
- j) gravity gastric drainage;
- k) orally operated mucous extractors;

- r) uterine aspiration;
- s) thoracic drainage.

2 Normative reference

The following standard contains provisions which, through reference in this text, constitute provisions of this part of ISO 10079. At the time of publication, the edition indicated was valid. All standards are subject to revision, and parlies to agreements based on this part of ISO 10079 are encouraged to investigate the possibility of applying the most recent edition of the standard indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.

suction equipment where the collection container

is downstream of the vacuum pump;

ISO 5356-1:1987, Anaesthetic and respiratory equipment — Conical connectors — Part 1: Cones and sockets.

3 Definitions

For the purposes of this part of ISO 10079, the following definitions apply. **3.1 collection container:** Container in which liquids or solid particles are collected.

3.2 collection container assembly: Collection container and its closure.

3.3 end piece: That part of the suction equipment applied to the patient. The end piece starts at the site where material is drawn in and ends at the first detachable connection.

3.4 exhaust opening: Port or ports through which exhaust is discharged.



NOTES

1 ISO 10079-1 applies to mains electricity and battery-powered suction equipment.

This part of ISO 10079 applies to manually powered suction equipment. ISO 10079-3 applies to suction equipment powered from a vacuum or pressure source.

2 Components illustrated are not necessarily required by this International Standard.

3 Suction equipment shown are only examples, and actual systems may consist of other arrangements and components not illustrated in the figure.

Figure 1 — Schematic drawing illustrating suction equipment

3.5 filter: Device for separation of particulate matter.

3.6 free air flow: Unrestricted flow of air through a nominated inlet.

high vacuum: Vacuum of at least 60 kPa¹⁾. 3.7

3.8 inlet: Part of a component through which fluids and/or solid particles enter.

3.9 intermediate tubing: Tubing between the collection container and the vacuum pump.

3.10 low Vacuum than vacuum: not more - 20 kPa.

3.11 manually powered (generated) vacuum: Local generation of vacuum by human effort with a hand or foot or both.

3.12 manually powered transportable suction equipment: Equipment in which vacuum is generated manually.

3.13 medium vacuum: Vacuum less than 60 kPa R and greater than - 20 kPa.

sterilized as recommended by the manufacturer. 3.14 outlet: Part of a component through which flu-ISO 10079-2:1992 ids and/or solid particles exit.

(standards.

https://standards.iteh.ai/catalog/standards/sis5/62 Design requirements

3.15 overfill protection: Prevention of liquid of solidiso-10079-2-1992 particles entering the intermediate tubing.

3.16 overfill protection device: Any device intended to prevent liquid or solid particles entering the intermediate tubing.

3.17 suction: Application of vacuum to remove fluids and/or solid particles.

3.18 suction equipment: Single self-contained unit or combination of units which generates or controls suction.

3.19 suction tubing: Tubing for conduction of fluids from the end piece to the collection container.

3.20 vacuum: Pressure less than atmospheric pressure, normally expressed as a difference from atmospheric pressure.

3.21 vacuum indicator: Device for displaying the level of vacuum.

3.22 vacuum pump: Powered device for generating vacuum.

3.23 vacuum source: Means of generating vacuum. The source may be integral with the suction equipment or be separate from the suction equipment.

Cleaning, disinfection and sterilization 4

The suction equipment shall meet the require-4.1 ments specified in 8.1 to 8.3 after those components which are subject to contamination and which are intended for re-use have been submitted to 30 cycles of cleaning, disinfection and/or sterilization as recommended by the manufacturer.

4.2 Any filters installed shall either be of the disposable type or be capable of being cleaned, disinfected and/or sterilized for re-use in accordance with 4.1.

4.3 Suction equipment incorporating a re-usable collection container assembly shall comply with the requirements specified in 8.1 to 8.3, as appropriate, before and after the collection container has been subjected to 30 cycles of cleaning, disinfection and/or sterilization as recommended by the manufacturer.

4.4 Suction tubing shall either be for single use or

be capable of being cleaned, disinfected and/or

Connectors 51

5.1.1 Collection container connectors

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The connectors for the suction tubing and the intermediate tubing to the vacuum source shall be designed to facilitate correct assembly or marked to indicate correct assembly when all parts are mated. Compliance shall be checked by inspection.

The construction of the connections has fre-NOTE 1 quently been a cause of spill-over into a vacuum pump. The use of mechanical fittings so as to ensure correct attachment is highly desirable.

5.1.2 Inside diameter of suction tubing connection

The inside diameter of the suction tubing connection (inlet port) shall be equal to or larger than the inside diameter of the largest tubing size recommended by the manufacturer.

5.1.3 Exhaust opening

It shall not be possible to connect suction tubing to the exhaust opening.

1) 1 kPa = 7,500 63 mmHg or 4,024 619 in H₂O or 10,197 16 cm H₂O or 19 hPa

5.2 Suction tubing

5.2.1 General

If supplied, suction tubing shall have an inside diameter of not less than 6 mm.

NOTE 2 Suction performance may be markedly affected by the length and diameter of the tubing between the collection container and the end piece.

When tested in accordance with A.1, suction tubing supplied with the equipment shall retain at least 50 % (0,5) of its inside diameter throughout its length.

5.2.2 Length of suction tubing for foot-operated suction equipment²⁾

The length of suction tubing shall be such that the end piece can be positioned at least 1,3 m above the floor when the foot-operated vacuum pump is on the floor in the operating position.

Operational requirements 6

6.1 Ease of operation

The suction equipment shall be designed to be op-ISO 1007except that excluded by the manufacturer as specierated by one person unaided.

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Dismantling and reassembly 6.2

Suction equipment intended to be dismantled by the user (for example, for cleaning) shall be designed to facilitate correct assembly or marked to indicate correct reassembly when all parts are mated. After dismantling, reassembly and testing in accordance with the manufacturer's instructions, the suction equipment shall meet the requirements specified in 8.1 to 8.3, as appropriate.

Mechanical shock 6.3

After suction equipment intended for field or transport use or both has been dropped in accordance with A.2, it shall meet the requirements specified in 8.1 to 8.3, as appropriate.

"Field use" of suction equipment is intended to NOTE 3 cover use in situations outside of the health care facility at the site of accidents or other emergencies. The use of suction equipment in these situations may expose the equipment to water including rain, dirt, uneven support, mechanical shock and extremes of temperature. "Transport use" of suction equipment is intended to cover situations outside of the health care facility such as in ambulances, cars or aeroplanes. Use of suction equipment in these situations may expose the equipment to uneven support, dirt, mechanical shock and a wider range of temperature than normally found in health care facilities

If the suction equipment can be operated outside of its carrying case, it shall meet the requirements specified in 8.1 to 8.3, as appropriate, after the individual parts of the suction equipment have been dropped in accordance with A.2 and reassembled.

Immersion in water 6.4

After suction equipment intended for field use has been dropped in its ready-for-use condition from a reservoir water into height of 1 m а $1 \text{ m} \times 1 \text{ m} \times 1 \text{ m}$, has been left in the water for 10 s and the water has been expelled for 7 s, it shall meet the requirements specified in 8.1 to 8.3, as appropriate.

6.5 Stability

Suction equipment operated by foot and intended for field or transport use or both shall meet the requirements specified in 8.1 to 8.3, as appropriate, when placed on a surface of 20° (0,35 rad) slope iTeh STANDA from the horizontal. Other manually powered suction equipment when operated 10° (0,17 rad) from its (standar normal orientation shall meet the requirements specified in 8.1 to 8.3, as appropriate, in any position

> 73adba4fa8da/iso-10079-2-1992 6.6 Overfill protection

6.6.1 Unless the suction equipment is intended to continue operating after overflow of liquids and solids, means shall be provided to prevent liquids entering the suction line downstream of the overfill protection device in normal use of the equipment, and, when tested in accordance with A.3, the volume collected in the collection container shall be not less than 90 % of the stated collection capacity.

6.6.2 Suction shall cease when the overfill protection device operates.

Vacuum indicators 6.7

6.7.1 Analogue displays shall have graduations not less than 2 mm apart, each graduation representing not more than 5 % of the full scale value.

6.7.2 Digital displays shall display vacuum at intervals of not greater than 2 % of the full scale value. The maximum vacuum for which the equipment is designed shall be marked prominently on the display case or immediately adjacent to it.

²⁾ See also annex C.

6.7.3 All markings on the vacuum indicator shall be legible to an operator having visual acuity, corrected if necessary, of at least 1,0, seated or standing 1 m from the vacuum indicator at an illuminance of 215 lx.

6.7.4 The full scale of analogue vacuum indicators shall be not more than 200 % of the maximum designed negative pressure of the suction equipment.

6.7.5 Vacuum indicators shall be accurate to within + 5 % of the full scale value.

Movement of a rotary analogue vacuum indicator should be counter-clockwise for an increase in vacuum

Physical requirements 7

7.1 Dimensions³⁾

Suction equipment is often combined with resuscitation equipment which may make it impossible to define a mass or dimensions for suction equipment. alone. In these circumstances, this clause may not apply but the mass and dimensions of all equipment intended for field use should be as small as possings ible.

Suction equipment intended for field use, including **Flow** 2015224-ff21-44a7-b307-8,2 any carrying case or frame, shall pass through a dimensions diversions dimensions having rectangular opening 600 mm × 300 mm.

7.2 Mass

The mass of suction equipment intended for field use, complete with its carrying case or frame and accessories, shall not exceed 6 kg.

Collection container 7.3

7.3.1 The inlet of the collection container shall have an inside diameter of not less than 6 mm and not less than the maximum inside diameter of the suction tubing recommended by the manufacturer. The inlet shall not be compatible with any conical connector specified in ISO 5356-1.

7.3.2 For suction equipment which is intended to continue operating when the collection container is full and is intended for field use, the volume of the collection container shall be not less than 200 ml. For all other suction equipment, including suction equipment intended for field and/or transport use, the usable volume of the collection container shall be not less than 500 ml.

7.3.3 For suction equipment not intended for field use, one or more collection containers recommended by the manufacturer and clearly visible in the position of normal use shall be used. The collection container shall be marked with its usable volume, expressed in millilitres. For collection containers having a capacity of 500 ml or greater, approximate indication of the volume of the contents shall be given by graduations at intervals of not less than 50 ml and not more than 250 ml.

7.3.4 The collection container shall not implode, crack or permanently deform when tested in accordance with A.4. Following this test, the suction equipment shall meet the requirements of 6.6.1, 6.6.2, 8.1, 8.2 and 8.3, as appropriate.

Performance requirements for vacuum 8 and flow⁴⁾

Vacuum 81

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When tested in accordance with A.5, suction equipment shall develop a vacuum of at least – 40 kPa within 10 s.

When tested in accordance with A.6, suction equipment shall evacuate 200 ml of simulated vomitus in not more than 10 s.

8.3 Free air flow

When tested in accordance with A.7, the peak free air flow shall be at least 0,33 l/s (20 l/min).

Resistance to environment 9

9.1 Operating conditions

When tested in accordance with A.8.2.1 and A.8.2.2, suction equipment intended for field and transport use shall meet the requirements specified in 8.1 to 8.3.

9.2 Storage

When tested in accordance with A.8.2.3 and A.8.2.4, suction equipment intended for field and transport use shall meet the requirements specified in 8.1 to 8.3.

³⁾ See also annex A.

⁴⁾ See also annex C.