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Anaesthetic and respiratory equipment — Heat and moisture exchangers for use in humidifying respired gases in humans

iTeh STANDARD PREVIEW

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*Matériel d'anesthésie et de réanimation respiratoire — Échangeurs de
chaleur et d'humidité utilisés pour humidifier les gaz respirés par les
êtres humains*

ISO 9360:1992

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

International Standard ISO 9360 was prepared by Technical Committee ISO/TC 121, *Anaesthetic and respiratory equipment*, Sub-Committee SC 3, *Lung ventilators and related equipment*.

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Annex A of this International Standard is for information only.
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Introduction

Heat and moisture exchangers (HMEs) are used to raise the water content and the temperature of gas delivered to the respiratory tract of patients. They are primarily intended for use with tracheotomized or intubated patients, independently or as a part of a breathing system.

The gases generally available for medical use lack sufficient moisture to be physiologically acceptable to the respiratory tract. HMEs capture the exhaled heat and moisture and transfer them to the inspired gases.

Although heat and moisture exchangers have been used for many years, the introduction of HMEs utilizing primarily non-metallic components and hygroscopic additives or hydrophobic material have prompted the development of this International Standard.

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Anaesthetic and respiratory equipment — Heat and moisture exchangers for use in humidifying respired gases in humans

1 Scope

This International Standard specifies minimum performance and safety requirements for heat and moisture exchangers (HMEs) intended for humidification of respired gases in humans, and describes test methods for their evaluation.

2 Normative references

The following standards contain provisions which, through reference in this text, constitute provisions of this International Standard. At the time of publication, the editions indicated were valid. All standards are subject to revision, and parties to agreements based on this International Standard are encouraged to investigate the possibility of applying the most recent editions of the standards indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.

ISO 4135:1979, *Anaesthesiology — Vocabulary*.

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

ISO 5356-2:1987, *Anaesthetic and respiratory equipment — Conical connectors — Part 2: Screw-threaded weight-bearing connectors*.

ISO 7000:1989, *Graphical symbols for use on equipment — Index and synopsis*.

IEC 601-1:1988, *Medical electrical equipment — Part 1: General requirements for safety*.

3 Definitions

For the purposes of this International Standard, the definitions given in ISO 4135 and the following definitions apply.

3.1 identification mark; identification number: Symbols, numbers or lettering marked on a device from which the manufacturer/user derives information concerning its production (such as material batch or date of manufacture).

3.2 HME: Device intended to retain a portion of the expired moisture and heat, and return it to the patient's respiratory tract during inspiration.

3.3 HME patient port: That port of the HME which is connected to the patient's respiratory tract.

3.4 HME machine; atmospheric end port: That port of the HME which is connected to the patient connection port of a breathing system or is open to ambient air.

3.5 HME moisture output: Total amount of water, in milligrams per litre, of inspired gas leaving the HME patient port, under specified test conditions.

4 Symbols and abbreviations

The principal symbols and abbreviations used in this International Standard are given in table 1. Additional symbols are explained in the relevant context.

5 General requirements and recommendations

5.1 Patient port connector

The connector at the patient port shall be a 15 mm female conical connector as specified in ISO 5356-1:1987.

The connector at the patient port may also have a 22 mm male conical connector as specified in ISO 5356-1:1987.

If the HME incorporates an accessory port, that port shall not accept the 15 mm or 22 mm connectors specified in ISO 5356-1:1987 or ISO 5356-2:1987.

Other ports intended to accept breathing attachments, if present, shall be 15 mm male and/or 22 mm female conical connectors as specified in ISO 5356-1:1987.

If the HME incorporates a gas-scavenging port, that port shall be either a 19 mm or 30 mm male conical connector as specified in ISO 5356-1:1987.

5.2 Gas leakage

When tested according to 6.6, the leakage from HMEs intended to be used at elevated intermittent or continuous pressures shall not exceed 25 ml/m

at a pressure of 30 hPa (30 cm H₂O). (See also annex A.)

5.3 Pressure drop

When tested according to 6.7, the pressure drop across the HME shall not exceed 5 hPa (5 cm H₂O). (See also annex A.)

5.4 Packaging

5.4.1 HMEs supplied sterile and intended for single use shall be individually packaged.

5.4.2 The type of container used shall be such as to ensure that once opened, the container cannot be easily resealed, and that it shall be obvious that the container has been opened.

Each HME should be packed in a single container, the materials of which should not have detrimental effects on the contents. The material and design of this container should be such as to ensure

- minimal risk of contamination of the contents from opening and removal from the container;

- adequate protection of the contents during normal handling, transit and storage.

6 Test methods

The apparatus and test methods specified in 6.1 to 6.7.5 are not intended to exclude the use of other measuring devices or methods yielding results of an accuracy equal to or greater than those specified. In case of dispute, the methods given in this International Standard shall be the reference methods.

Table 1 — Symbols and abbreviations

| Symbol | Definition | Unit |
|-----------|--|----------|
| HME | Heat and moisture exchanger | |
| E_w | HME moisture output | mg/l |
| m_1 | Initial mass of the patient model before testing with the HME | g or mg |
| m_2 | Final mass of the patient model after testing with the HME | g or mg |
| m_3 | Initial mass of the patient model before testing without the HME | g or mg |
| m_4 | Final mass of the patient model after testing without the HME | g or mg |
| f | Frequency | |
| bpm | breaths per minute | |
| C | Compliance | ml/hPa |
| R | Resistance | hPa/l/s |
| I:E ratio | Inspiratory: expiratory ratio | |
| r.h. | Relative humidity | per cent |
| V_t | Tidal volume | ml |

6.1 Temperatures and pressures (see also annex A)

6.1.1 The ambient temperatures (defined as temperature t_A in figure 1, zone 1) for the duration of the test shall be $23\text{ °C} \pm 2\text{ °C}$. Barometric pressure shall be stated, as shall the temperature at which the measurements were taken.

6.1.2 Temperature t_B in figure 1, zone 3, shall be high enough to eliminate condensation in rubber bags, valves and tubing.

A suggested temperature is $37\text{ °C} \pm 3\text{ °C}$.

6.1.3 The water-bath temperature, t_G , in figure 1 shall be regulated to give a maximum temperature t_F measured at the HME patient port of $34\text{ °C} \pm 1\text{ °C}$ when averaged over 20 expirations.

6.1.4 The temperature in the inspiratory flow, t_H , in figure 1 shall be $23\text{ °C} \pm 2\text{ °C}$.

6.1.5 The response time for probes measuring t_A , t_B and t_H in figure 1 shall be 10 s or less for 50 % of the actual value. The response time in flowing air at points E and F shall be 0,1 s or less for 50% of the temperature cycle.

6.2 Test gas and apparatus

6.2.1 The test gas shall be air having a humidity not exceeding 0,88 mg/l, equivalent to a dew point of -20 °C at atmospheric pressure.

6.2.2 Gas flow measuring equipment shall be calibrated to an accuracy of $\pm 5\%$ of the reading in the range 1 l/min to 100 l/min.

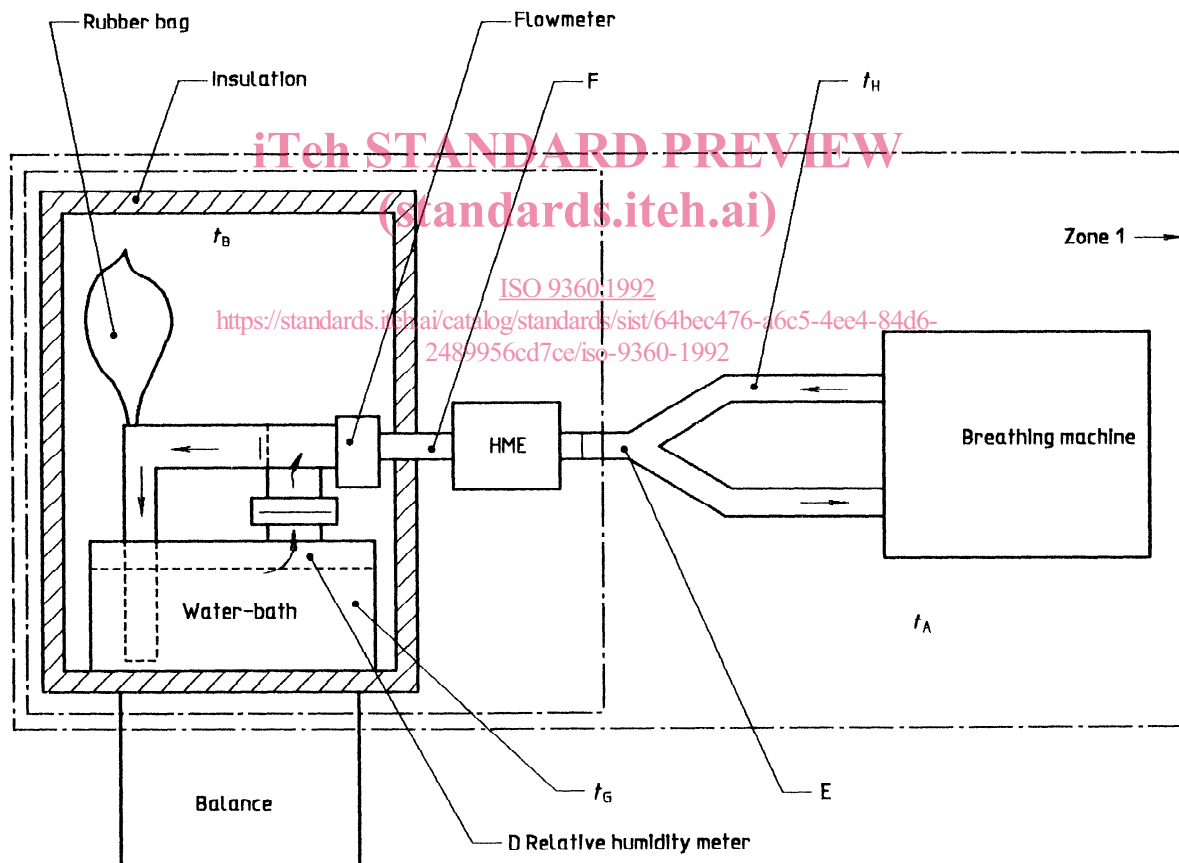


Figure 1 — Test set-up

6.2.3 Test apparatus consisting of a temperature-controlled water-bath with a means of providing compliance and resistance values as specified in table 2 shall be used (see figure 1). The inspiratory and expiratory flows shall be directed via one-way valves into separate pathways. Provision shall be made to measure the flow through the HME. Temperature probes shall be mounted at points A, B, E, F, G and H (where these letters refer also to subscripts of points at which temperature-measuring probes are inserted). The temperatures at these points shall be recorded. The temperature probe at point F shall be connected to a control system in order to regulate the water-bath temperature, t_G . Temperature probes at points E and F shall be mounted within 10 mm of the HME patient port. The length of tubing between the HME and the ventilator shall not exceed 1 m, and the length of tubing between the HME and the point of separation of the inspiratory and expiratory gas flow pathways shall not exceed 10 cm. The inspiratory and expiratory limbs shall be isolated by a uni-directional valve. A relative humidity probe shall be mounted at point D. A flowmeter shall be connected at point F. The capacity of the bag shall be greater than the tidal volumes given in table 2.

6.2.4 The HME shall be connected to the test apparatus and a ventilator.

For HMEs intended only for use during spontaneous breathing, a suitable adaptor with minimal dead space should be used.

6.2.5 The ventilator shall deliver the minute volumes at the frequencies shown in table 2. The flow profile of the ventilator during the measurements shall be stated in the test report.

NOTE 1 It is recognized that the flow profile may influence the efficiency of the HME; it is therefore assumed that the ventilator used in the test apparatus, as shown in figure 1, is capable of delivering as constant a flow as possible during inspiration.

6.2.6 The weighing equipment used shall have an accuracy of $\pm 0,1$ g or better in the range of mass to be measured.

6.2.7 Pressure drop measuring equipment shall consist of a differential pressure gauge with an accuracy equal to or better than ± 10 kPa (0,1 cm H₂O) for the test as shown in figure 2.

6.3 Measurement of HME moisture output

6.3.1 Principle

The test apparatus, comprising the patient model and the HME under test, is connected to a flow gen-

erator, for example a ventilator. After it has operated for approximately 60 min, a steady state is achieved in the test system. During inspiration, the dry gas at ambient temperature passes through the HME, thereby absorbing accumulated heat and moisture. During expiration, the expired gas, which now is assumed to be saturated with water vapour, passes through the HME, in the reverse direction, so that heat and moisture are retained in the HME.

6.3.2 Procedure

6.3.2.1 Connect the HME to the patient model and the ventilator.

6.3.2.2 Record temperatures t_A , t_B and t_H .

6.3.2.3 Adjust the ventilator to give one of the test conditions in the combinations listed in table 2 within the HME's operating range as specified by the manufacturer.

6.3.2.4 Adjust the water-bath temperature to give a maximum temperature at point F of $34\text{ °C} \pm 1\text{ °C}$ during expiration.

6.3.2.5 Verify that the relative humidity at point D is 100 % during the entire breathing cycle.

Care should be taken to choose a humidity-measuring instrument suitable for measurements in the region of 100 % r.h. at the test temperatures.

6.3.2.6 Verify that the humidity at point H is less than 0,88 mg/l (equivalent to a dew point of -20 °C).

6.3.2.7 Let the equipment run for 60 min ± 5 min to precondition the HME and stabilize the test system.

6.3.2.8 Disconnect the HME, seal the ports, and weigh the patient model.

6.3.2.9 Connect the HME and operate the test apparatus for 1 h. Record the temperatures at points E and F continuously.

NOTE 2 This temperature will vary, and typical variations are shown in figure 3.

6.3.2.10 Disconnect the HME such that any condensation that has occurred is retained *in situ* within the patient model and then weigh the patient model.

Table 2 — Test conditions

| Intended for V_t of | Test conditions | V_t ml | f bpm | Minimum volume l/m | I:E ratio | C | R |
|-----------------------|-----------------|-------------|------------|-----------------------|-----------|-----|-----|
| > 500 ml | 1 | 1 000 | 20 | 20 | 1:2 | 50 | 5 |
| | 2 | 1 000 | 10 | 10 | 1:2 | 50 | 5 |
| | 3 | 500 | 20 | 20 | 1:2 | 50 | 5 |
| 51 ml to 500 ml | 1 | 500 | 20 | 10 | 1:2 | 50 | 5 |
| | 2 | 250 | 20 | 5 | 1:2 | 10 | 20 |
| < 50 ml | 1 | 50 | 40 | 2 | 1:1 | 10 | 20 |
| | 2 | 25 | 40 | 1 | 1:1 | 10 | 20 |

Dimensions in millimetres

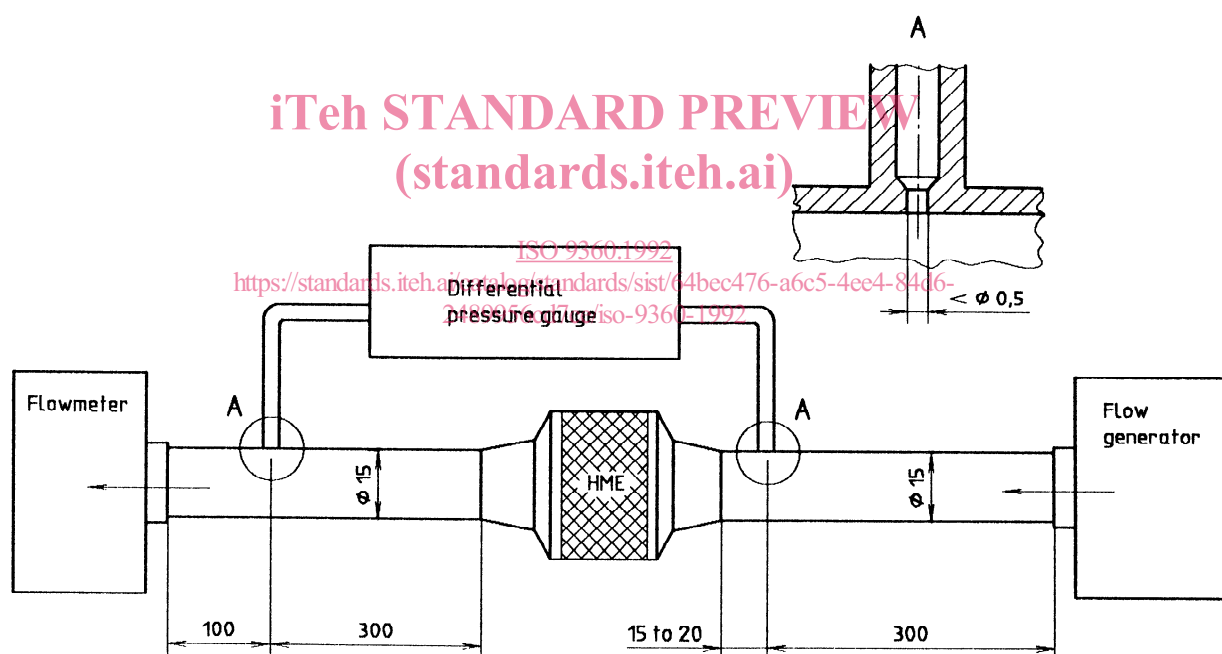


Figure 2 — Pressure drop measuring set-up