ISO 80601-2-80:2023(E) (Ed 2)

ISO/TC 121/SC 3/

Secretariat: ANSI

2024-04-<mark>03</mark>02

Medical Equipment — Part 2-80: Particular requirements for basic safety and essential performance of ventilatory support equipment for ventilatory insufficiency

Appareils électromédicaux — Partie 2-80: Exigences particulières pour la sécurité de base et les performances essentielles des équipements d'assistance ventilatoire en cas d'insuffisance ventilatoire

(https://standards.iteh.ai)
Document Preview

FDIS stage

Warning for WDs and CDs

This document is not an ISO International Standard. It is distributed for review and comment. It is subject to change without notice and may not be referred to as an International Standard.

Recipients of this draft are invited to submit, with their comments, notification of any relevant patent rights of which they are aware and to provide supporting documentation.

Edited DIS - MUST BE USED FOR FINAL DRAFT

ISO-<u>/FDIS</u> 80601-2-80:2023(E2024(en))

© ISO 2023, Published in Switzerland

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

- __ISO copyright office
- Ch. de Blandonnet 8 CP 401
- CH-1214 Vernier, Geneva, Switzerland
- ___Tel. + 41 22 749 01 11
- Fax + 41 22 749 09 47
- copyright@iso.org
- _www.iso.org

iTeh Standards (https://standards.iteh.ai) Document Preview

ISO/FDIS 80601-2-80

Contents

Introduction	ix
201.1 Scope, object and related standards	1
201.2 Normative references	4
201.3 Terms and definitions	5
201.4 General requirements	24
201.5 General requirements for testing of ME equipment	<u></u> 28
201.6 Classification of ME equipment and ME systems	
201.7 ME equipment identification, marking and documents	30
201.8 Protection against electrical hazards from ME equipment	37
201.9 Protection against mechanical hazards of ME equipment and ME systems	38
201.10 Protection against unwanted and excessive radiation hazards	 40
201.11 Protection against excessive temperatures and other hazards	 40
201.12 Accuracy of controls and instruments and protection against hazardous outputs	 44
201.13 Hazardous situations and fault conditions for ME equipment	 57
201.14 Programmable electrical medical systems (PEMS)	 58
201.15 Construction of ME equipment	 59
201.17 Electromagnetic compatibility of ME equipment and ME systems	 60
201.101 Gas connections.	 61
201.102 Requirements for the VBS and accessories	 64
201.103 Spontaneous breathing during loss of power supply	 65
201.104 Indication of duration of operation	 66
201.105 Functional connection	 66
201.106 Display loops	 67
201.108 Ventilatory support equipment security	 68
202 Electromagnetic disturbances — Requirements and tests	 68
206 Usability	 69
208 General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	 71
211 Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	 73
Annex C (informative) Guide to marking and labelling requirements for ME equipment and ME systems	 74
Annex D (informative) Symbols on marking	 81
© ISO 2023 – All rights reserved iii	

ISO-<u>/FDIS</u>80601-2-80:2023(E2024(en)

Annex AA (informative) Particular guidance and rationale	 83
Annex BB (informative) Data interface requirements	 105
Annex CC (informative) Reference to the IMDRF essential principles and labelling guidance	.s 112
Annex DD (informative) Reference to the essential principles Error! Bookmark not do	efined.
Foreword	<u></u> vii
Introduction	
201.1 Scope, object and related standards	<u></u> 1
201.2 Normative references	<u>4</u>
201.3 Terms and definitions	
201.4 General requirements	<u></u> 24
201.5 General requirements for testing of ME equipment	<u></u> 28
201.6 Classification of ME equipment and ME systems	<u></u> 30
201.7 ME equipment identification, marking and documents	
201.8 Protection against electrical hazards from ME equipment	
201.9 Protection against mechanical hazards of ME equipment and ME systems	<u></u> 38
201.10 Protection against unwanted and excessive radiation hazards	
201.11 Protection against excessive temperatures and other hazards	
201.12 Accuracy of controls and instruments and protection against hazardous outputs	
201.13 Hazardous situations and fault conditions for ME equipment	<u></u> 57
201.14 Programmable electrical medical systems (PEMS)(PEMS)	
201.15 Construction of ME equipment	<u></u> 59
201.17 Electromagnetic compatibility of ME equipment and ME systems	
201.101 Gas connections	
201.102 Requirements for the VBS and accessories	<u></u> 64
201.103 Spontaneous breathing during loss of power supply	
201.104 Indication of duration of operation	
201.105 Functional connection	
201.106 Display loops	<u></u> 67
201.107 Ventilatory support equipment security	
202 Electromagnetic disturbances — Requirements and tests	68
206 Usability	
208 General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	
211 Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	<u>1</u> 73
Annex C (informative) Guide to marking and labelling requirements for ME equipment and ME systems	<u></u> 74

ISO-<u>/FDIS</u> 80601-2-80:2023(E2024(en)

Annex D (informative) Symbols on marking81
Annex AA (informative) Particular guidance and rationale
AA.1 General guidance83
AA.2 Rationale for particular clauses and subclauses
Annex BB (informative) Data interface requirements105
BB.1 Background and purpose
BB.2 Data definition106
Annex CC (informative) Reference to the IMDRF essential principles and labelling guidances. 112
Bibliography116
<u>Terminology — Alphabetized index of defined terms</u> 119
Figures Figure 201.101 — Typical test setup for volume-control and pressure-control inflation-type accuracy
Tables iTeh Standards
Table 201.101— Distributed essential performance requirements23
Table 201.102 — Test conditions for acoustic tests
Table 201.103 — Examples of permissible combinations of temperature and relative humidity38
Table 201.104 — Volume-control inflation-type type test settings
Table 201.105 — Pressure-control inflation-type type test settings45
Table 201.C.101 — Marking on the outside of ventilatory support equipment, its parts or accessories
Table 201.C.102 — Accompanying documents, general69
Table 201.C.103 — Instructions for use70
Table 201.C.104 — Technical description
Table 201.D.1.101 — Additional symbols on marking
Table BB.101 — Parameters and units of measurement98
Table BB.102 — Equipment Identification98
Table BB.103 — Usage monitoring99
Table BB.104 — Equipment settings99
Table BB.105 — Ventilation monitoring
Table BB.106 — Ventilatory support equipment alarm limits
Table BB.107 — Event information
Table BB.108 — Service monitoring
© ISO 2023 – All rights reserved v

ISO-<u>/FDIS</u> 80601-2-80:2023(E2024(en)

Table C	:C.1 —	Corresp e	ondence	between	this do	cument	and the	IMDRF	essential	principl	es 1	.04
Table C	CC.2 —	Corresp e	ondence	between	this do	cument	and the	IMDRF	labelling	, principl	les 1	06
Table E	D.1 —	Corresp	ondence	betweer	this d	ocument	and the	e essent	ial princi	ples	1	07

iTeh Standards (https://standards.iteh.ai) Document Preview

ISO/FDIS 80601-2-80

Foreword

ISO (the International Organization for Standardization) and IEC (the International Electrotechnical Commission) form the specialized system for worldwide standardization. National bodies that are members of ISO or IEC participate in the development of International Standards through technical committees established by the respective organization to deal with particular fields of technical activity. ISO and IEC technical committees collaborate in fields of mutual interest. Other international organizations, governmental and non-governmental, in liaison with ISO and IEC, also take part in the work.

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 document 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 or www.iso.org/directives<

ISO and IEC draw attention to the possibility that the implementation of this document may involve the use of (a) patent(s). ISO and IEC take no position concerning the evidence, validity or applicability of any claimed patent rights in respect thereof. As of the date of publication of this document, ISO and IEC had not received notice of (a) patent(s) which may be required to implement this document. However, implementers are cautioned that this may not represent the latest information, which may be obtained from the patent database available at www.iso.org/patents and https://patents.iec.ch. ISO and IEC shall not be held responsible for identifying any or all such patent rights.

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. In the IEC, see www.iso.org/iso/foreword.html. In the IEC, see www.iso.org/iso/foreword.html.

This document was prepared jointly by Technical Committee ISO/TC 121, *Anaesthetic and respiratory equipment*, Subcommittee SC 3, *Lung ventilators and related equipment*, and Technical Committee IEC/TC 62, *Medical equipment, software, and systems*, Subcommittee SC D, *Particular medical equipment, software, and systems*, in collaboration with the European Committee for Standardization (CEN) Technical Committee CEN/TC 215 *Respiratory and anaesthetic equipment*, in accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).

This second edition cancels and replaces the first edition (ISO 80601-2-80:2018), which has been technically revised.

The main changes are as follows:

- alignment with IEC 60601-1:2005+AMD1:2012+AMD2:2020, IEC 60601-1-2:2014+AMD1:2020
 IEC 60601-1-6:2010+AMD1:2013+AMD2:2020, IEC 60601-1-8:2006+AMD1:2012+AMD2:2020
 and IEC 60601-1-11:2015+AMD1:2020;
- reformatted according to most recent Central Secretariat editing rules;
- clarified maximum limited pressure requirements;
- clarified high airway pressure alarm condition requirements;

ISO-/FDIS 80601-2-80:2023(E2024(en)

- added requirements for *ventilatory support equipment system recovery*; and
- harmonization with ISO 20417, where appropriate.

A list of all parts in the ISO 80601 series and the IEC 80601 series can be found on the ISO and IEC websites.

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 and www.iec.ch/national-committees.

iTeh Standards (https://standards.iteh.ai) Document Preview

ISO/FDIS 80601-2-80

Introduction

This document specifies requirements for *ventilatory support equipment* that is intended for use in the *home healthcare environment* for *patients* who are not dependent for *ventilation* for their life support. *Ventilatory support equipment* is frequently used in locations where *supply mains* is not reliable. *Ventilatory support equipment* is often supervised by non-healthcare personnel (*lay operators*) with varying levels of training. *Ventilatory support equipment* conforming with this document can be used elsewhere (i.e. in healthcare facilities).

Varying levels of ventilatory support is often used for *patients* who have stable ventilatory needs and in some cases, changing needs as their disease worsens. This document addresses *patients* who typically have severe enough respiratory function to prohibit certain activities that the *patient* might normally pursue, and to interfere with daily living, occurring in association with measurements of respiratory mechanics or gas exchange that are markedly abnormal. This is best characterised by *lung* functions worse than^[36]

- $FEV_1/FVC^1 < 70$ %, or
- $FEV_1 < 50 \%$ predicted

where

FEV₁ is the forced expiratory volume in 1 s, and

FVC is the forced vital capacity.

Examples of diseases that require ventilatory support are:

- moderate to severe Chronic Obstructive Pulmonary Disease (COPD);
- moderate Amyotrophic Lateral Sclerosis (ALS)^[44];
- severe bronchopulmonary dysplasia; and
- muscular dystrophy.

Ventilatory support equipment intended for this group of *patients* typically can require *technical alarm conditions* in the event that *essential performance* is absent. The most fragile of these *patients* would likely experience injury, but not serious injury or death, with the loss of this *artificial ventilation*. For these *patients*, it is likely that ventilatory support is needed during waking hours while *patients* are moving inside or outside the home in order to facilitate mobility and functional independence in the activities of daily living.

In this document, the following print types are used:

- requirements and definitions: roman type;
- terms defined in Clause 3 of the general standard², in this document or as noted: italic type; and
- informative material appearing outside of tables, such as notes, examples and references: in smaller type.
 Normative text of tables is also in a smaller type.

This is also known as the Tiffeneau-Pinelli index.

The general standard is IEC 60601-1:2005+AMD1:2012+AMD2:2020, Medical electrical equipment — Part 1: General requirements for basic safety and essential performance.

ISO-<u>/FDIS</u> 80601-2-80:2023(E2024(en))

In referring to the structure of this document, the term:

- "clause" means one of the five numbered divisions within the table of contents, inclusive of all subdivisions (e.g. Clause 201 includes subclauses 201.7, 201.8, etc.); and
- "subclause" means a numbered subdivision of a clause (e.g. 201.7, 201.8 and 201.9 are all subclauses of Clause 201).

References to clauses within this document are preceded by the term "Clause" followed by the clause number. References to subclauses within this particular document are by number only.

In this document, the conjunctive "or" is used as an "inclusive or" so a statement is true if any combination of the conditions is true.

For the purposes of this document, the auxiliary verb:

- "shall" indicates a requirement;
- "should" indicates a recommendation;
- "may" indicates a permission;
- "can" indicates a possibility or capability; and
- "must" is used to express an external constraint.

Annex C contains a guide to the *marking* and labelling requirements in this document.

Annex D contains a summary of the *symbols* referenced in this document.

Requirements in this document have been decomposed so that each requirement is uniquely delineated. This is done to support automated requirements tracking.

ISO/FDIS 80601-2-80

Medical electrical equipment

Part 2-80:

Particular requirements for basic safety and essential performance of ventilatory support equipment for ventilatory insufficiency

201.1 Scope, object and related standards

IEC 60601-1:2005+AMD1:2012+AMD2:2020, Clause 1, applies, except as follows:

201.1.1 Scope

Replacement:

NOTE 1 There is guidance or rationale for this subclause contained in Clause AA.2.

This document applies to the *basic safety* and *essential performance* of *ventilatory support equipment*, as defined in 201.3.302, for *ventilatory insufficiency*, as defined in 201.3.302, hereafter also referred to as *ME equipment*, in combination with its *accessories*:

- intended for use in the *home healthcare environment*;
 - NOTE 2 In the *home healthcare environment*, the *supply mains* driving the *ventilatory support equipment* is often not reliable.
 - NOTE 3 Such ventilatory support equipment can also be used in professional health care facilities.
- intended for use by a lay operator;
- intended for use with *patients* who have *ventilatory insufficiency* or failure, the most fragile of which would likely experience injury with the loss of this *artificial ventilation*;
- intended for transit-operable use; and
- not intended for *patients* who are dependent on *artificial ventilation* for their immediate life support.

EXAMPLE 1 *Patients* with moderate to severe chronic obstructive pulmonary disease (COPD), moderate amyotrophic lateral sclerosis (ALS), severe bronchopulmonary dysplasia or muscular dystrophy.

Ventilatory support equipment is not considered to use a physiologic closed-loop control system unless it uses a physiological patient variable to adjust the artificial ventilation therapy settings.

This document is also applicable to those *accessories* intended by their *manufacturer* to be connected to the *ventilator breathing system* of *ventilatory support equipment* for *ventilatory insufficiency*, where the characteristics of those *accessories* can affect the *basic safety* or *essential performance* of the *ventilatory support equipment* for *ventilatory insufficiency*.

ISO-<u>/FDIS</u> 80601-2-80:2023(E2024(en)

EXAMPLE 2 Breathing sets, *connectors*, water traps, expiratory valve, *humidifier*, *breathing system filter*, external electrical power source, *distributed alarm system*.

If a clause or subclause is specifically intended to be applicable to *ME equipment* only, or to *ME systems* only, the title and content of that clause or subclause will say so. If that is not the case, the clause or subclause applies both to *ME equipment* and to *ME systems*, as relevant.

Hazards inherent in the intended physiological function of *ME equipment* or *ME systems* within the scope of this document are not covered by specific requirements in this document except in IEC 60601-1:2005+AMD1:2012+AMD2:2020, 7.2.13 and 8.4.1.

NOTE 4 Additional information can be found in IEC 60601-1:2005+AMD1:2012+AMD2:2020, 4.2.

NOTE 5 See ISO/TR 21954 for guidance on the selection of the appropriate *ventilator* for a given *patient*.

This document does not specify the requirements for:

- *ventilators* or *accessories* for *ventilator-dependent patients* intended for critical care applications, which are given in ISO 80601-2-12;
- *ventilators* or *accessories* intended for anaesthetic applications, which are given in ISO 80601-2-13;
- ventilators or accessories intended for the emergency medical services environment, which are given in ISO 80601-2-84;
- *ventilators* or *accessories* intended for *ventilator-dependent patients* in the *home healthcare environment*, which are given in ISO 80601-2-72;
- ventilatory support equipment or accessories intended for ventilatory impairment, which are given in ISO 80601-2-79;
- sleep apnoea therapy ME equipment, which are given in ISO 80601-2-70;
- high-frequency jet *ventilators* (HFJVs), which are given in ISO 80601-2-87;
- high-frequency oscillatory ventilators (HFOVs)^[20];
- respiratory high flow equipment, which are given in ISO 80601-2-90;
 - NOTE 6 *Ventilatory support equipment* can incorporate high-flow therapy operational mode, but such a mode is only for spontaneously breathing *patients*.
- user-powered resuscitators, which are given in ISO 10651-4;
- gas-powered emergency resuscitators, which are given in ISO 10651-5;
- oxygen therapy constant flow ME equipment; and
- cuirass or "iron-lung" *ventilation* equipment.

201.1.2 Object

IEC 60601-1:2005+AMD1:2012+AMD2:2020, 1.2 is replaced by:

The object of this document is to establish particular *basic safety* and *essential performance* requirements for *ventilatory support equipment,* for *ventilatory insufficiency,* as defined in 201.3.302, and its *accessories*.

ISO-<u>/FDIS</u> 80601-2-80:2023(E2024(en))

Accessories are included because the combination of the *ventilatory support equipment* and the *accessories* need to be adequately safe. Accessories can have a significant impact on the *basic safety* or *essential performance* of the *ventilatory support equipment*.

NOTE 1 This document has been prepared to address the relevant International Medical Device Regulators Forum (IMDRF) *essential principles*^[31] and labelling^[32] guidances as indicated in Annex CC.

NOTE 2 This document has been prepared to address the relevant *essential principles of safety and performance* of ISO 16142-1:2016 as indicated in Annex DD.

NOTE 3 This document has been prepared to address the relevant general safety and performance requirements of European regulation (EU) $2017/745^{[33]}$.

201.1.3 Collateral standards

IEC 60601-1:2005+AMD1:2012+AMD2:2020, 1.3 applies with the following addition:

This document refers to those applicable collateral standards that are listed in Clause 2 of the general standard and Clause 201.2 of this document.

201.1.4 Particular standards Teh Standards

IEC 60601-1:2005+AMD1:2012+AMD2:2020, 1.4 is replaced by:

In the IEC 60601 series, particular standards define *basic safety* and *essential performance* requirements, and may modify, replace or delete requirements contained in the general standard, including the collateral standards as appropriate for the particular *ME equipment* under consideration.

A requirement of a particular standard takes priority over the general standard.

For brevity, IEC 60601-1:2005+AMD1:2012+AMD2:2020 is referred to in this particular document as the general standard. Collateral standards are referred to by their document number.

The numbering of clauses and subclauses of this document corresponds to that of the general standard with the prefix "201" (e.g. 201.1 in this document addresses the content of Clause 1 of the general standard) or applicable collateral standard with the prefix "2xx", where xx is the final digits of the collateral standard document number (e.g. 202.4 in this document addresses the content of Clause 4 of the IEC 60601-1-2 collateral standard, 211.10 in this document addresses the content of Clause 10 of the IEC 60601-1-11 collateral standard, etc.). The changes to the text of the general standard are specified by the use of the following words:

"Replacement" means that the clause or subclause of the general standard or applicable collateral standard is replaced completely by the text of this document.

"Addition" means that the text of this document is additional to the requirements of the general standard or applicable collateral standard.

"Amendment" means that the clause or subclause of the general standard or applicable collateral standard is amended as indicated by the text of this document.

Subclauses, figures or tables that are additional to those of the general standard are numbered starting from 201.101. However, due to the fact that definitions in the general standard are numbered 3.1 through 3.154, additional definitions in this document are numbered beginning from 201.3.201. Additional annexes are lettered AA, BB, etc., and additional items aa), bb), etc.

ISO-/FDIS 80601-2-80:2023(E2024(en)

Subclauses, figures or tables which are additional to those of a collateral standard are numbered starting from 20x, where "x" is the number of the collateral standard, e.g. 202 for IEC 60601-1-2, 203 for IEC 60601-1-3, etc.

The term "this document" is used to make reference to the IEC 60601-1:2005+AMD1:2012+AMD2:2020, any applicable collateral standards and this document taken together.

Where there is no corresponding clause or subclause in this particular document, the clause or subclause of the general standard or applicable collateral standard, although possibly not relevant, applies without modification; where it is intended that any part of the general standard or applicable collateral standard, although possibly relevant, is not to be applied, a statement to that effect is given in this document.

201.2 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.

IEC 60601-1:2005+AMD1:2012+AMD2:2020, Clause 2, applies, except as follows:

Replacement:

ISO 15223-1:2021, Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements

IEC 61672-1:2013, Electroacoustics — Sound level meters — Part 1: Specifications

Addition:

Document Preview

ISO 3744:2010, Acoustics — Determination of sound power levels and sound energy levels of noise sources using sound pressure — Engineering methods for an essentially free field over a reflecting plane

ISO 4871:1996, Acoustics — Declaration and verification of noise emission values of machinery and equipment

ISO 5356-1:2015, Anaesthetic and respiratory equipment — Conical connectors — Part 1: Cones and sockets

ISO 5359:2014+AMD1/Amd 1:2017, Anaesthetic and respiratory equipment — Low-pressure hose assemblies for use with medical gases

ISO 5367:2023, Anaesthetic and respiratory equipment — Breathing sets and connectors

ISO 7396-1:2016+AMD1/Amd 1:2017, Medical gas pipeline systems — Part 1: Pipeline systems for compressed medical gases and vacuum

ISO 9360-1:2000, Anaesthetic and respiratory equipment — Heat and moisture exchangers (HMEs) for humidifying respired gases in humans — Part 1: HMEs for use with minimum tidal volumes of 250 ml

ISO 9360-2:2001, Anaesthetic and respiratory equipment — Heat and moisture exchangers (HMEs) for humidifying respired gases in humans — Part 2: HMEs for use with tracheostomized patients having minimum tidal volumes of 250 ml