EVS-EN ISO 80601-2-80:2019

Medical electrical equipment - Part 2-80: Particular requirements for basic safety and essential performance of ventilatory support equipment for ventilatory insufficiency (ISO 80601-2-80:2018), (corrected version 11.2019)



EESTI STANDARDI EESSÕNA

NATIONAL FOREWORD

3	
See Eesti standard EVS-EN ISO 80601-2-80:2019 sisaldab Euroopa standardi EN ISO 80601-2-80:2019 ingliskeelset teksti.	This Estonian standard EVS-EN ISO 80601-2-80:2019 consists of the English text of the European standard EN ISO 80601-2-80:2019.
Standard on jõustunud sellekohase teate avaldamisega EVS Teatajas	This standard has been endorsed with a notification published in the official bulletin of the Estonian Centre for Standardisation.
Euroopa standardimisorganisatsioonid on teinud Euroopa standardi rahvuslikele liikmetele kättesaadavaks 18.09.2019.	Date of Availability of the European standard is 18.09.2019.
Standard on kättesaadav Eesti Standardikeskusest.	The standard is available from the Estonian Centre for Standardisation.

Tagasisidet standardi sisu kohta on võimalik edastada, kasutades EVS-i veebilehel asuvat tagasiside vormi või saates e-kirja meiliaadressile <u>standardiosakond@evs.ee</u>.

ICS 11.040.10

Standardite reprodutseerimise ja levitamise õigus kuulub Eesti Standardikeskusele

Andmete paljundamine, taastekitamine, kopeerimine, salvestamine elektroonsesse süsteemi või edastamine ükskõik millises vormis või millisel teel ilma Eesti Standardikeskuse kirjaliku loata on keelatud.

Kui Teil on küsimusi standardite autorikaitse kohta, võtke palun ühendust Eesti Standardikeskusega: Koduleht <u>www.evs.ee</u>; telefon 605 5050; e-post <u>info@evs.ee</u>

The right to reproduce and distribute standards belongs to the Estonian Centre for Standardisation

No part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying, without a written permission from the Estonian Centre for Standardisation.

If you have any questions about copyright, please contact Estonian Centre for Standardisation:

Homepage www.evs.ee; phone +372 605 5050; e-mail info@evs.ee

EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM

EN ISO 80601-2-80

September 2019

ICS 11.040.10

Supersedes EN ISO 10651-6:2009

English Version

Medical electrical equipment - Part 2-80: Particular requirements for basic safety and essential performance of ventilatory support equipment for ventilatory insufficiency (ISO 80601-2-80:2018)

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 (ISO 80601-2-80:2018) Medizinische elektrische Geräte - Teil 2-80: Besondere Festlegungen für die grundlegende Sicherheit und die wesentlichen Leistungsmerkmale von Heimbeatmungsgeräten zur Atemunterstützung von Patienten mit Atmungsinsuffizienz (ISO 80601-2-80:2018)

This European Standard was approved by CEN on 28 July 2019.

This European Standard was corrected and reissued by the CEN-CENELEC Management Centre on 30 October 2019.

CEN members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this European Standard the status of a national standard without any alteration. Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN-CENELEC Management Centre or to any CEN member.

This European Standard exists in three official versions (English, French, German). A version in any other language made by translation under the responsibility of a CEN member into its own language and notified to the CEN-CENELEC Management Centre has the same status as the official versions.

CEN members are the national standards bodies of Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and United Kingdom.



EUROPEAN COMMITTEE FOR STANDARDIZATION COMITÉ EUROPÉEN DE NORMALISATION EUROPÄISCHES KOMITEE FÜR NORMUNG

CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels

© 2019 CEN All rights of exploitation in any form and by any means reserved worldwide for CEN national Members.

European foreword

The text of ISO 80601-2-80:2018 has been prepared by Technical Committee ISO/TC 121 "Anaesthetic and respiratory equipment" of the International Organization for Standardization (ISO) and has been taken over as EN ISO 80601-2-80:2019 by Technical Committee CEN/TC 215 "Respiratory and anaesthetic equipment" the secretariat of which is held by BSI.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by March 2020, and conflicting national standards shall be withdrawn at the latest by March 2020.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN shall not be held responsible for identifying any or all such patent rights.

This document supersedes l'EN ISO 10651-6:2009.

According to the CEN-CENELEC Internal Regulations, the national standards organizations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and the United Kingdom.

Endorsement notice

The text of ISO 80601-2-80:2018 has been approved by CEN as EN ISO 80601-2-80:2019 without any modification.

Contents

Foreword	vi
Introduction	.viii
201 1 Scone object and related standards	1
201.1 Scope, object and related standards	1 1
201.1.2 Object	2
201112 Object:	2
201.1.4 Particular standards	3
201.2 Normative references	3
201.3 Terms and definitions	5
201.4 General requirements	7
201.4.3 Essential performance	7
201.4.3.101 * Additional requirements for ESSENTIAL PERFORMANCE	7
201.4.6 * ME EQUIPMENT OF ME SYSTEM parts that contact the PATIENT	7
201.4.11.101 * Additional requirements for pressurized gas input	8
201 5 Conoral requirements for testing of ME FOULDMENT	Q
201.5.101 * Additional requirements for the general requirements for testing of	9
	9
201.5.101.1 Ventilatory support equipment test conditions	9
201.5.101.2 * Gas nowrate and leakage specifications	9
201.5.101.3 * VENTILATORY SUPPORT EQUIPMENT LESUING ETTORS	9
201.6 Classification of ME EQUIPMENT and ME SYSTEMS	10
201.6.101 * Additional requirements for classification of ME EQUIPMENT and ME SYSTEMS	10
201.7 ME EQUIPMENT identification, marking and documents	10
201.8 Protection against electrical HAZARDS from ME EQUIPMENT	17
201.9 Protection against mechanical hazards of ME EQUIPMENT and ME SYSTEMS	17
201.10 Protection against unwanted and excessive radiation HAZARDS	19
201.11 Protection against excessive temperatures and other HAZARDS	19
201.11.7 Biocompatiblity of ME EQUIPMENT and ME SYSTEMS	20
201.11.8 Interruption of the power supply/SUPPLY MAINS to ME EQUIPMENT	21
201.11.8.101 Additional requirements for interruption of the power supply/supply mains to ME EQUIPMENT ALARM CONDITION	21
201.12 Accuracy of controls and instruments and protection against bagardous outputs	22
201.12 Accuracy of controls and instruments	43
201.12.1 Accuracy of controlled breath type	
201.12.1.101 Volume-controlled breath type	
20112.1.102 Tressure controlled breach type	
20112.11100 Other Dreath types imment	29
20112.2.101 Osubility of the equipment introduction of the second se	29
201.12.4.101 * Measurement of AIRWAY PRESSURE	29
201.12.4.102 Measurement of expired volume	31
201.12.4.103 * Maximum limited pressure protection device	31
201.12.4.104 Hypoventilation ALARM CONDITION	31
201.12.4.105 * High leakage ALARM CONDITION	31
201.12.4.106 * O_2 rebreathing	32
201.12.101 * Protection against accidental adjustments	32

201.13 Hazardous situations and fault conditions for ME EQUIPMENT	33
201.14 PROGRAMMABLE ELECTRICAL MEDICAL SYSTEMS (PEMS)	34
201.15 Construction of ME EQUIPMENT	34
201.15.101 Mode of operation	34
201.15.102 Pre-use check	34
201.16 ME SYSTEMS	34
201.17 Electromagnetic compatibility of ME EQUIPMENT and ME SYSTEMS	35
201.101 Gas connections	35
201.101.1 VBS connectors	35
201.101.1.1 General	35
201.101.1.2 Other named ports	35
201.102 Requirements for the VBS and ACCESSORIES	36
201.102.1 * General	36
201.102.2 Labelling	37
201.102.3 Breathing sets	37
201.102.4 * Humidification	37
201.102.4.1 HUMIDIFIER	37
201.102.4.2 HEAT AND MOISTURE EXCHANGER (HME)	3 / 27
201.102.5 DREATHING SYSTEM FILTERS (BSF)	37
201.105 Spontaneous breating uning loss of power suppry	37
201.105 * Indication of duration of operation	38
201 106 Functional connection	38
201.106 1 General	38
201.106.2 * Connection to an electronic health record	39
201.106.3 * Connection to a distributed alarm system	39
201.106.4 Connection for remote control	39
201.107 Display loops	39
201.107.1 Pressure-volume loops	39
201.107.2 Flow-volume loops	39
201.108 Power supply cords	40
201.109 Ventilatory support equipment security	40
202 Electromagnetic disturbances — Requirements and tests	40
206 Usability	41
208 General requirements, tests and guidance for alarm systems in medical electrical	
equipment and medical electrical systems	43
211 Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	44
Annex C (informative) Guide to marking and labelling requirements for ME EQUIPMENT and	45
Anney D (informative) Symbols on marking	57
Annex AA (informative) Particular guidance and rationale	
Anney BB (informative) Data interface requirements	60
	09
Annex LL (informative) Reference to the ESSENTIAL PRINCIPLES	76

Bibliography				
2.				
J.				
0				
	4			
	2			
	S.			
	Ū.			
	Ŷ	A		
	1			
		Q,		
		4		
		CL.		
			0	
			2	
			4	
				6.
				L
				50
				0.

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 complying with this document can be used elsewhere (i.e. in healthcare facilities).

Varying levels of ventilatory support are needed 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^[3]

- FEV₁/FVC² < 70 %, or
- FEV₁ < 50 % predicted

where

FEV₁ is the forced expiratory volume in 1 s, and

FVC is the forced vital capacity.

Examples of diseases that require ventilation support are severe Chronic Obstructive Pulmonary Disease (COPD), Amyotrophic Lateral Sclerosis (ALS)^[4], 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.

This publication has been drafted in accordance with the ISO/IEC Directives, Part 2.

In this document, the following print types are used:

- requirements and definitions: roman type;
- test specifications: italic type;
- 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.
- TERMS DEFINED IN CLAUSE 3 OF THE GENERAL STANDARD³, IN THIS DOCUMENT OR AS NOTED: SMALL CAPITALS.

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.);

² This is also known as the Tiffeneau-Pinelli index.

³ The general standard is IEC 60601-1:2005+AMD1:2012, Medical electrical equipment — Part 1: General requirements for basic safety and essential performance.

— "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.

The verbal forms used in this document conform to usage described in ISO/IEC Directives, Part 2. For the purposes of this document, the auxiliary verb:

- "shall" means that compliance with a requirement or a test is mandatory for compliance with this document:
- "should" means that compliance with a requirement or a test is recommended but is not mandatory for compliance with this document;
- "may" is used to describe permission (e.g. a permissible way to achieve compliance with a requirement or test);
- "can" is used to describe a possibility or capability;
- "must" is used express an external constraint.

An asterisk (*) as the first character of a title or at the beginning of a paragraph or table title indicates that there is guidance or rationale related to that item in Annex AA.

The ISO and IEC 80601 family of documents are also parts of the IEC 60601 family of documents.

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, Clause 1, applies, except as follows:

201.1.1 * Scope

Replacement:

This document applies to the BASIC SAFETY and ESSENTIAL PERFORMANCE of VENTILATORY SUPPORT EQUIPMENT, as defined in 201.3.205, for VENTILATORY INSUFFICIENCY, as defined in 201.3.204, hereafter also referred to as ME EQUIPMENT, in combination with its ACCESSORIES:

- intended for use in the HOME HEALTHCARE ENVIRONMENT;
- 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;
- 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.

NOTE 1 In the HOME HEALTHCARE ENVIRONMENT, the SUPPLY MAINS is often not reliable.

NOTE 2 Such VENTILATORY SUPPORT EQUIPMENT can also be used in non-critical care applications of professional health care facilities.

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.

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, 7.2.13 and 8.4.1.

NOTE 3 Additional information can be found in IEC 60601-1:2005+AMD1:2012, 4.2.

This document does not specify the requirements for:

- VENTILATORS OF 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^[5];
- VENTILATORS or ACCESSORIES intended for the emergency medical services environment, which are given in ISO 80601-2-84^{[6]4}, the future replacement for ISO 10651-3^[7];
- VENTILATORS OF ACCESSORIES intended for VENTILATOR-DEPENDENT PATIENTS in the HOME HEALTHCARE ENVIRONMENT, which are given in ISO 80601-2-72;
- VENTILATORY SUPPORT EQUIPMENT OF ACCESSORIES intended for VENTILATORY IMPAIRMENT, which are given in ISO 80601-2-79^[1];
- sleep apnoea therapy ME EQUIPMENT, which are given in ISO 80601-2-70^[8];
- continuous positive airway pressure (CPAP) ME EQUIPMENT;
- high-frequency jet VENTILATORS (HFJVs);
- high-frequency oscillatory VENTILATORS (HFOVs)^[9];
- oxygen therapy constant flow ME EQUIPMENT;
- cuirass or "iron-lung" ventilation equipment.

This document is a particular standard in the IEC 60601 and IEC/ISO 80601 series of documents.

201.1.2 Object

Replacement:

The object of this document is to establish particular BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for VENTILATORY SUPPORT EQUIPMENT, as defined in 201.3.205, and its ACCESSORIES.

NOTE ACCESSORIES are included because the combination of the VENTILATORY SUPPORT EQUIPMENT and the ACCESSORIES needs to be adequately safe. ACCESSORIES can have a significant impact on the BASIC SAFETY OF ESSENTIAL PERFORMANCE of the VENTILATORY SUPPORT EQUIPMENT.

201.1.3 Collateral standards

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.

⁴ Under preparation. Stage at the time of publication: ISO/DIS 80601-2-84:2017.

IEC 60601-1-2:2014, IEC 60601-1-6:2010+AMD1:2013, IEC 60601-1-8:2006+AMD1:2012 and IEC 60601-1-11:2015 apply as modified in Clauses 202, 206, 208 and 211 respectively. IEC 60601-1-3:2008^[10] does not apply. All other published collateral standards in the IEC 60601-1 series apply as published.

201.1.4 Particular standards

Replacement:

In the IEC 60601 series, particular standards may modify, replace or delete requirements contained in the general standard, including the collateral standards as appropriate for the particular ME EQUIPMENT under consideration, and may add other BASIC SAFETY OR ESSENTIAL PERFORMANCE requirements.

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

For brevity, IEC 60601-1:2005+AMD1:2012 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 3.147, 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.

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 general standard, any applicable collateral standards and this particular 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.

NOTE 1 The way in which these referenced documents are cited in normative requirements determines the extent (in whole or in part) to which they apply.

NOTE 2 Informative references are listed in the Bibliography.

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

Replacement:

IEC 60601-1-2:2014, Medical electrical equipment — Part 1-2: General requirements for basic safety and essential performance — Collateral standard: Electromagnetic disturbances — Requirements and tests

IEC 60601-1-6:2010, Medical electrical equipment — Part 1-6: General requirements for basic safety and essential performance — Collateral standard: Usability +Amendment 1:2013⁵

IEC 60601-1-8:2006, Medical electrical equipment — Part 1-8: General requirements for basic safety and essential performance — Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems+Amendment 1:2012⁶

IEC 60601-1-11:2015, Medical electrical equipment — Part 1-11: General requirements for basic safety and essential performance — Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment

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

Addition:

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 4135:2001, Anaesthetic and respiratory equipment - Vocabulary

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 5367:2014, Anaesthetic and respiratory equipment — Breathing sets and connectors

ISO 7000:2014, Graphical symbols for use on equipment — Registered symbols

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

ISO 8836:2014, Suction catheters for use in the respiratory tract

ISO 9000:2015, Quality management systems — Fundamentals and vocabulary

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

⁵ There exists a consolidated edition 3.1(2013) including IEC 60601-1-6:2010 and its Amendment 1:2013.

⁶ There exists a consolidated edition 2.1(2012) including IEC 60601-1-8:2006 and its Amendment 1:2012.

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

ISO 15223-1:2016, Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1: General requirements

ISO 16142-1:2016, Medical devices — Recognized essential principles of safety and performance of medical devices — Part 1: General essential principles and additional specific essential principles for all non-IVD medical devices and guidance on the selection of standards

ISO 17510:2015, Medical devices — Sleep apnoea breathing therapy — Masks and application accessories

ISO 17664:2017, *Processing of health care products* — *Information to be provided by the medical device manufacturer for the processing of medical devices*

ISO 18562-1:2017, Biocompatibility evaluation of breathing gas pathways in healthcare applications — Part 1: Evaluation and testing within a risk management process

ISO 23328-1:2003, Breathing system filters for anaesthetic and respiratory use — Part 1: Salt test method to assess filtration performance

ISO 23328-2:2002, Breathing system filters for anaesthetic and respiratory use — Part 2: Non-filtration aspects

ISO 80369-1:2010, Small-bore connectors for liquids and gases in healthcare applications — Part 1: General requirements

ISO 80369-7:2016, Small-bore connectors for liquids and gases in healthcare applications — Part 7: Connectors for intravascular or hypodermic applications

ISO 80601-2-12:—⁷, Medical electrical equipment —Part 2-12: Particular requirements for basic safety and essential performance of critical care ventilators

ISO 80601-2-72:2015, Medical electrical equipment — Part 2-72: Particular requirements for basic safety and essential performance of home healthcare environment ventilators for ventilator-dependent patients

ISO 80601-2-74:2017, Medical electrical equipment — Part 2-74: Particular requirements for basic safety and essential performance of respiratory humidifying equipment

IEC 62366-1:2015, Medical devices — Part 1: Application of usability engineering to medical devices

EN 15986:2011, Symbol for use in the labelling of medical devices — Requirements for labelling of medical devices containing phthalates

201.3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 4135:2001, ISO 7396-1:2016, ISO 8836:2014, ISO 9000:2015, ISO 9360-1:2000, ISO 16142-1:2016, ISO 17510:2015, ISO 17664:2017, ISO 18562-1:2017, ISO 23328-2:2002, IEC 60601-1:2005+AMD1:2012, IEC 60601-1-2:2014,

⁷ To be published. Stage at time of publication ISO/DIS 80601-2-12:2017.