

SLOVENSKI STANDARD oSIST prEN ISO 80601-2-87:2020

01-marec-2020

Medicinska električna oprema - 2-87. del: Posebne zahteve za osnovno varnost in bistvene lastnosti visokofrekvenčnega ventilatorja (ISO/DIS 80601-2-87:2020)

Medical electrical equipment - Part 2-87: Particular requirements for basic safety and essential performance of high-frequency ventilators (ISO/DIS 80601-2-87:2020)

Medizinische elektrische Geräte - Teil 2-87: Besondere Festlegungen an die Sicherheit und die wesentlichen Leistungsmerkmale von Hochfrequenzbeatmungsgeräten (ISO/DIS 80601-2-87:2020)

(standards.iteh.ai)

Appareils électromédicaux - Partie 2-87: Titre manque (ISO/DIS 80601-2-87:2020) <u>oSIST prEN ISO 80601-2-87:2020</u>

https://standards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-

Ta slovenski standard je istoveten z. pren 150 80601-2-87

ICS:

11.040.10 Anestezijska, respiratorna in Anaesthetic, respiratory and reanimacijska oprema reanimation equipment

oSIST prEN ISO 80601-2-87:2020 en

oSIST prEN ISO 80601-2-87:2020

iTeh STANDARD PREVIEW (standards.iteh.ai)

oSIST prEN ISO 80601-2-87:2020 https://standards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-7845635478ef/osist-pren-iso-80601-2-87-2020

DRAFT INTERNATIONAL STANDARD ISO/DIS 80601-2-87

ISO/TC **121**/SC **3**

Secretariat: ANSI

Voting begins on: **2020-01-17**

Voting terminates on:

2020-04-10

Medical electrical equipment —

Part 2-87:

Particular requirements for basic safety and essential performance of high-frequency ventilators

ICS: 11.040.10

iTeh STANDARD PREVIEW (standards.iteh.ai)

oSIST prEN ISO 80601-2-87:2020 https://standards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-7845635478ef/osist-pren-iso-80601-2-87-2020

This document is circulated as received from the committee secretariat.

THIS DOCUMENT IS A DRAFT CIRCULATED FOR COMMENT AND APPROVAL. IT IS THEREFORE SUBJECT TO CHANGE AND MAY NOT BE REFERRED TO AS AN INTERNATIONAL STANDARD UNTIL PUBLISHED AS SUCH.

IN ADDITION TO THEIR EVALUATION AS BEING ACCEPTABLE FOR INDUSTRIAL, TECHNOLOGICAL, COMMERCIAL AND USER PURPOSES, DRAFT INTERNATIONAL STANDARDS MAY ON OCCASION HAVE TO BE CONSIDERED IN THE LIGHT OF THEIR POTENTIAL TO BECOME STANDARDS TO WHICH REFERENCE MAY BE MADE IN NATIONAL REGULATIONS.

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.

This draft is submitted to a parallel vote in ISO and in IEC.

ISO/CEN PARALLEL PROCESSING



Reference number ISO/DIS 80601-2-87:2020(E)

iTeh STANDARD PREVIEW (standards.iteh.ai)

oSIST prEN ISO 80601-2-87:2020 https://standards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-7845635478ef/osist-pren-iso-80601-2-87-2020



COPYRIGHT PROTECTED DOCUMENT

© ISO 2020

All rights reserved. Unless otherwise specified, or required in the context of its implementation, 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 CP 401 • Ch. de Blandonnet 8 CH-1214 Vernier, Geneva Phone: +41 22 749 01 11 Fax: +41 22 749 09 47 Email: copyright@iso.org Website: www.iso.org

Published in Switzerland

Contents

2	201.1	Scope, object and related standards	1
3	201.2	Normative references	4
4	201.3	Terms and definitions	7
5	201.4	General requirements	8
6	201.5	General requirements for testing of ME equipment	10
7	201.6	Classification of ME equipment and ME systems	11
8	201.7	ME equipment identification, marking and documents	11
9	201.8	Protection against electrical hazards from ME equipment	18
10	201.9	Protection against mechanical hazards of ME equipment and ME systems	18
11	201.10	Protection against unwanted and excessive radiation hazards	22
12	201.11	Protection against excessive temperatures and other hazards	22
13 14	201.12	Accuracy of controls and instruments and protection against hazardous outputs	26
15	201.13	Hazardous situations and fault conditions for ME equipment	40
16	201.14	Programmable electrical medical systems (PEMS)	42
17	201.15	Construction of ME equipment ARD PREVIEW	
18	201.16	ME systems (standards.iteh.ai)	45
19	201.17	Electromagnetic compatibility of <i>ME equipment</i> and <i>ME systems</i>	45
20	201.101	Gas connections oSIST prEN ISO 80601-2-87:2020 https://standards.iteh.a/catalog/standards/sist/8a63d826-b514-44c9-aeb5-	46
21	201.102	Requirements for the HFV breathing system and accessories	48
22	201.103	Spontaneous breathing during loss of power supply	49
23	201.104	Indication of duration of operation	50
24	201.105	Functional connection	50
25	201.106	Display loops	51
26	201.107	Timed high frequency oscillation pause	51
27	202	Electromagnetic disturbances – Requirements and tests	51
28	206	Usability	52
29 30	208	General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	54
31	Annexes		57
32 33	Annex C	(informative) Guide to marking and labelling requirements for <i>ME equipment and ME systems</i>	58
34	Annex D	(informative) Symbols on marking	63
35	Annex AA	A (informative) Particular guidance and rationale	66
36	Annex BI	3 (informative) Data interface requirements	97
37	Annex CO	C (informative) Reference to the essential principles	103
38	Annex DI	O (informative) Reference to the general safety and performance	
39		requirements	
40		E (informative) Terminology — alphabetized index of defined terms	
	© ISO/IEC	2020 - All rights reserved	iii

41	Bibliography	114
42		
43	Figures	
44	Figure 201.101 — Typical closed suctioning test setup	21
45 46	Figure 201.102 — Performance test setup for an HFV with a patient-connection port	28
47 48	Figure 201.103 —Performance test setup for an HFV without a patient-connection port	29
49	Figure AA.1 — Representative Pressure waveform in an HFV	67
50 51	Figure AA.2 — Breathing system leakage flowrate limits as a function of pressure as specified in ISO 80601-2-12 [1] and ISO 80601-2-13 [3]	93
52		
53	Tables	
54	Table 201.101 — Distributed essential performance requirements	8
55	Table 201.102 — Test conditions for acoustic tests	19
56 57	Table 201.103 — Examples of permissible combinations of temperature and relative humidity	22
58	Table 201.104 — Test conditions for mean airway pressure and other tests	30
59	Table 201.105 — Tracheal tube parameters for <i>HFV</i> performance tests	30
60	Table 201.C.101 — Marking on the outside of a ventilator, its parts or accessories https://standards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-Table 201.C.102 — Accompanying documents, general 22-87-2020	58
61		
62	Table 201.C.103 — Instructions for use	
63	Table 201.C.104 — Technical description	
64	Table 201.D.2.101 — Additional symbols on marking	
65	Table AA.1 — Calculated conductance values by <i>patient</i> weight range	
66	Table BB.101 — Parameters and units of measurement	
67	Table BB.102 — Equipment identification	
68	Table BB.103 — Usage monitoring	
69	Table BB.104 — Equipment settings	
70	Table BB.105 — Ventilation monitoring	
71	Table BB.106 — Ventilator alarm limits	
72	Table BB.107 — Event information	101
73	Table BB.108 — Service monitoring	102
74	Table CC.1 — Correspondence between this document and the <i>essential principles</i>	103
75 76	Table DD.1 — Correspondence between this document and the general safety and performance requirements	106
77		

Foreword

78

- ISO (the International Organization for Standardization) is a worldwide federation of 79 national standards bodies (ISO member bodies). The work of preparing International 80 81 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 82 to be represented on that committee. International organizations, governmental and non-83 governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with 84 the International Electrotechnical Commission (IEC) on all matters of electrotechnical 85 standardization. 86
- 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. 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.

 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.

 (Standards.iteh.ai)
- This document was prepared by Technical Committee ISO/TC 121, Anaesthetic and respiratory equipment, Subcommittee ISCI3, Lung-ventilators and related equipment and Technical Committee/IEC/TC62, Electrical equipment in medical practice, Subcommittee 62D: Electric equipment. 7845635478efosist-pren-iso-80601-2-87-2020
- This is the first edition of ISO 80601-2-87.
- This publication has been drafted in accordance with the ISO/IEC Directives, Part 2.

105

Introduction

106

- In this document, the following print types are used:
- 108 Requirements and definitions: roman type
- Instructions, test specifications and terms defined in Clause 3 of the general standard, in
 this document or as noted: 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
- In referring to the structure of this document, the term
- "clause" means one of the four numbered divisions within the table of contents, inclusive of all subdivisions (e.g. Clause 201 includes subclauses 201.7, 201.8, etc.);
- "subclause" means a numbered subdivision of a clause (e.g. 201.7, 201.8 and 201.9 are all subclauses of Clause 201).
- 118 References to clauses within this document are preceded by the term "Clause" followed by the clause number. References to subclauses within this 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. ARD PREVIEW
- 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:

oSIST prEN ISO 80601-2-87:2020

- "shall" means hthat conformance with dad requirements or 42 stests is mandatory for conformance with this document of sixty pren-iso-80601-2-87-2020
- "should" means that conformance with a requirement or a test is recommended but is
 not mandatory for conformance with this document;
- "may" is used to describe permission (e.g. a permissible way to achieve conformance with a requirement or test);
- "can" is used to describe 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.
- 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.

- Medical Electrical Equipment Part 2-87: Particular
- requirements for basic safety and essential performance
- of high-frequency critical care ventilators
- 139 201.1 Scope, object and related standards
- 140 Clause 1 of the general standard applies, except as follows:
- 141 **201.1.1** * Scope
- 142 Replacement:
- This document applies to the *basic safety* and *essential performance* of a *high-frequency* ventilator (HFV) in combination with its accessories, hereafter referred to as ME equipment:
- intended for use in an environment that provides specialized care for *patients* whose conditions can be life-threatening and who can require comprehensive care and constant monitoring in a *professional healthcare facility*;
- NOTE 1 For the purposes of this document, such an environment is referred to as a critical care environment. *High-frequency ventilators* for this environment are considered life-sustaining.
- NOTE 2 For the purposes of this document, such a high-frequency ventilator can provide transport within a professional healthcare facility (i.e. be a transit-operable ventilator).
- NOTE 3 A high-frequency ventilator intended for use in transport within a professional healthcare facility is not considered as an ventilator intended for the emergency medical services environment.

 https://standards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-
- 154 intended to be operated by a healthcare professional operator;
- intended for those *patients* who need differing levels of support from *artificial* ventilation including ventilator-dependent patients; and
- capable of providing more than 150 *inflations*/min.
- 158 There are three principal designations of *HFV*:
- high frequency percussive *ventilation* (HFPV, with a typical *HFV frequency* of (60 to 1000) *HFV inflations*/min);
- high frequency jet *ventilation* (HFJV, with a typical *HFV frequency* of (100 to 1500) *HFV inflations*/min); and
- high frequency oscillatory *ventilation* (HFOV, with a typical *HFV frequency* of (180 to 1200) *HFV inflations*/min and typically having an active *expiratory phase*).
- Additionally, *HFV* designations can be combined together or with *ventilation* at *rates* less than 150 *inflations*/min.

© ISO/IEC 2020 - All rights reserved

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

- * A high-frequency ventilator is not considered to utilize physiologic closed loop-control system unless it uses a physiological patient variable to adjust the ventilation therapy
- 169 settings.
- This document is also applicable to those *accessories* intended by their *manufacturer* to be
- 171 connected to an HFV breathing system, or to a high-frequency ventilator, where the
- characteristics of those accessories can affect the basic safety or essential performance of
- the *high-frequency* ventilator.
- 174 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
- 177 relevant.
- 178 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 7.2.13 and 8.4.1 of IEC 60601-1:2005.
- NOTE 4 Additional information can be found in 4.2 of IEC 60601-1:2005+AMD1:2012.
- This document is not applicable to *ME equipment* that is intended solely to augment the
- *ventilation* of spontaneously breathing *patients* within a *professional healthcare facility*.
- This document does not specify the requirements for: REVIEW
- non-high-frequency ventilators or accessories which provide conventional ventilation for use in critical care environments, which are given in ISO 80601-2-12;.
- NOTE 5 An HFV can incorporate conventional critical care ventilator operational modes, in which case
- ISO 80601-2-12 is applicable to those modes. 7845635478ef/osist-pren-iso-80601-2-87-2020
- 189 ventilators or accessories intended for anaesthetic applications, which are given in 190 $ISO 80601-2-13 [3]^2$;
- ventilators or accessories intended for the emergency medical services environment, which are given in ISO 80601-2-84, the future replacement for ISO 10651-3 [4];
- NOTE 6 An *HFV* can incorporate *EMS ventilator* capability.
- *ventilators* or *accessories* intended for *ventilator-dependent patients* in the *home* healthcare environment, which are given in ISO 80601-2-72 [5];
- 196 ventilators or accessories intended for home-care ventilatory support devices, which are given in ISO 80601-2-79 [6] and ISO 80601-2-80 [7], the replacements for
- 198 ISO 10651-6 [8];
- sleep apnoea breathing therapy *ME equipment*, which are given in ISO 80601-2-70 [9];
- 200 continuous positive airway pressure (CPAP) ME equipment;
- 201 oxygen therapy constant flow *ME equipment*; and

2

² Figures in square brackets refer to the Bibliography.

- 202 cuirass or "iron-lung" ventilation equipment.
- 203 This document is a particular standard in the IEC 60601 and IEC/ISO 80601 series of
- 204 documents.

205 **201.1.2 Object**

- 206 Replacement:
- The object of this document is to establish particular basic safety and essential performance
- requirements for a *high-frequency ventilator*, as defined in 201.3.201, and its *accessories*.
- NOTE 1 Accessories are included because the combination of the high-frequency ventilator and the
- accessories needs to be adequately safe. Accessories can have a significant impact on the basic safety or
- 211 essential performance of a high-frequency ventilator.
- 212 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 CC.
- NOTE 3 This document has been prepared to address the relevant general safety and performance
- requirements of European regulation (EU) 2017/745 [10] as indicated in Annex DD.

216 **201.1.3 Collateral standards**

- 217 Amendment (add after existing text): NDARD PREVIEW
- 218 This document refers to those applicable collateral standards that are listed in Clause 2 of
- the general standard and 201.2 of this document.
- 220 IEC 60601-1-2, IEC 60601-1-6 and IEC 60601-1-8 apply as modified in Clauses 202, 206
- 221 and 208 respectively. IEC 60601-1-3 [11], IEC 60601-1-9 [12], IEC 60601-1-11 and
- IEC 60601-1-12 [13] do not apply. All other published collateral standards in the
- IEC 60601-1 series apply as published.

224 201.1.4 Particular standards

- 225 Replacement:
- 226 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
- 228 the particular ME equipment under consideration, and may add other basic safety or
- 229 essential performance requirements.
- 230 A requirement of a particular standard takes priority over IEC 60601-1:2005 or the
- 231 collateral standards.
- 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 those of the
- 235 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"
- 237 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,
- 239 208.4 in this document addresses the content of Clause 4 of the IEC 60601-1-8 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 IEC 60601-1:2005 or the 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
- 245 IEC 60601-1:2005 or the applicable collateral standard.
- "Amendment" means that the clause or subclause of IEC 60601-1:2005 or the 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 or figures that 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
- 255 IEC 60601-1-2, 203 for IEC 60601-1-3 [11], etc.
- 256 The term "this document" is used to make reference to the general standard, any
- 257 applicable collateral standards and this particular document taken together.
- 258 Where there is no corresponding clause or subclause in this document, the clause or
- subclause of IEC 60601-1:2005+AMD1:2012 or the applicable collateral standard,
- although possibly not relevant, applies without modification; where it is intended that any
- part of IEC 60601-1:2005+AMD1:2012 or the applicable collateral standard, although
- 262 possibly relevant, is not to be applied, a statement to that effect is given in this particular
- 263 document.

oSIST prEN ISO 80601-2-87:2020

201.2 Normative references log/standards/sist/8a63d826-b514-44c9-aeb5-

- 7845635478ef/osist-pren-iso-80601-2-87-2020

 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
- 268 (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.
- 272 Clause 2 of the general standard applies, except as follows:
- 273 Replacement:
- ISO 7000:2014, Graphical symbols for use on equipment Registered symbols
- 275 ISO 7010:2011+AMD1:2012+AMD2:2012+AMD3:2012+AMD4:2013+AMD5:2014
- +AMD6:2014+AMD7:2016+AMD8:2017, Graphical symbols Safety colours and safety
- 277 signs Registered safety signs
- 278 ISO 15223-1:2016, Medical devices Symbols to be used with medical device labels,
- labelling and information to be supplied Part 1: General requirements
- IEC 60601-1-2:2014, Medical electrical equipment Part 1-2: General requirements for
- 281 basic safety and essential performance Collateral standard: Electromagnetic
- 282 disturbances Requirements and tests

- IEC 60601-1-6:2010+AMD1:2013, Medical electrical equipment Part 1-6: General
- requirements for basic safety and essential performance Collateral standard: Usability
- IEC 60601-1-8:2006+AMD1:2012, 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
- 288 medical electrical systems
- IEC 61672-1:2013, Electroacoustics Sound level meters Part 1: Specifications
- IEC 62304:2006+AMD1:2015, Medical device software Software life cycle processes
- 291 Addition:
- ISO 32:1977, Gas cylinders for medical use Marking for identification of content
- 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
- 295 over a reflecting plane
- 296 ISO 4871:1996, Acoustics Declaration and verification of noise emission values of
- 297 machinery and equipment
- ISO 5356-1:2015, Anaesthetic and respiratory equipment Conical connectors Part 1:
- 299 Cones and sockets iTeh STANDARD PREVIEW
- ISO 5359:2014, Anaesthetic and respiratory equipment Low-pressure hose assemblies for
- 301 use with medical gases
- ISO 5367:2014, Angesthetic and respiratory equipment -- Breathing sets and connectors
- 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
- 307 ISO 14937:2009, Sterilization of health care products General requirements for
- characterization of a sterilizing agent and the development, validation and routine control
- of a sterilization process for medical devices
- 310 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
- 313 standards
- 314 ISO 17510:2015, Medical devices -- Sleep apnoea breathing therapy -- Masks and application
- 315 *accessories*
- 316 ISO 17664:2017, Processing of health care products -- Information to be provided by the
- medical device manufacturer for the processing of medical devices
- 318 ISO 18562-1:2017, Biocompatibility evaluation of breathing gas pathways in healthcare
- applications-- Part 1: Evaluation and testing within a risk management process
- ISO 19223:2019, Lung ventilators and related equipment -- Vocabulary and semantics

- ISO 23328-1:2003, Breathing system filters for anaesthetic and respiratory use: Part 1:
- 322 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 80369-1:2018, Small-bore connectors for liquids and gases in healthcare applications -
- 326 Part 1: General requirements
- ISO 80601-2-12:— ³ (Ed 2), Medical electrical equipment Part 2-12: Particular
- requirements for the basic safety and essential performance of critical care ventilators
- ISO 80601-2-55:2018 (Ed 2), Medical electrical equipment Part 2-55: Particular
- requirements for the basic safety and essential performance of respiratory gas monitors
- ISO 80601-2-74:2017, Medical electrical equipment Part 2-74: Particular requirements
- 332 for the basic safety and essential performance of respiratory humidifying equipment
- ISO $80601-2-84:\frac{4}{100}$, Medical electrical equipment Part 2-84: Particular requirements for
- basic safety and essential performance of emergency and transport ventilators
- IEC 60068-2-27:2008, Environmental testing Part 2-27: Tests Test Ea and guidance:
- 336 Shock
- IEC 60068-2-31:2008, Environmental testing Part 2-31: Tests + Test Ec: Rough handling
- 338 shocks, primarily for equipment-type specimens
- (standards.iteh.ai)

 IEC 60068-2-64:2008, Environmental testing Part 2-64: Tests Test Fh: Vibration,
- broadband random and guidance SIST prEN ISO 80601-2-87:2020
- 341 IEC 60529:1989+AMD1:1999+AMD2:2013, Degrees of protection provided by enclosures
- 342 (*IP Code*)
- 343 IEC 60601-1:2005+AMD1:2012, Medical electrical equipment Part 1: General
- 344 requirements for basic safety and essential performance
- IEC 60601-1-10:2007, Medical electrical equipment Part 1-10: General requirements for
- 346 basic safety and essential performance Collateral Standard: Requirements for the
- 347 development of physiologic closed-loop controllers
- IEC 60601-1-11:2015, Medical electrical equipment Part 1-11: General requirements for
- basic safety and essential performance Collateral Standard: Requirements for medical
- 350 electrical equipment and medical electrical systems used in the home healthcare
- 351 environment
- 352 IEC 60601-1-12:2014, Medical electrical equipment Part 1-12: General requirements for
- basic safety and essential performance Collateral Standard: Requirements for medical
- 354 electrical equipment and medical electrical systems intended for use in the emergency
- 355 medical services environment
- 356 IEC 60601-2-2:2009, Medical electrical equipment Part 2-2: Particular requirements for
- the basic safety and essential performance of high frequency surgical equipment and high
- 358 frequency surgical accessories

³ Under preparation. Stage at the time of publication: ISO FDIS 80601-2-12:2019.

⁴ Under preparation. Stage at the time of publication: ISO FDIS 80601-2-84:2019.

- 359 IEC 62366-1:2015, Medical devices Part 1: Application of usability engineering to
- 360 medical devices
- IEC 62570:2014, Standard practice for marking medical devices and other items for safety
- in the magnetic resonance environment
- EN 15986:2011, Symbols for medical devices containing phthalates

364 **201.3** Terms and definitions

- For the purposes of this document, the terms and definitions given in ISO 7010:2011,
- 366 ISO 7396-1:2016, ISO 8836:2014, ISO 9000:2015, ISO 16142-1:2016, ISO 17510:2015,
- 367 ISO 17664:2017, ISO 18562-1:2017, ISO 19223:2019, ISO 23328-2:2002,
- 368 IEC 60601-1:2005+AMD1:2012, IEC 60601-1-2:2014, IEC 60601-1-6:2010,
- 369 IEC 60601-1-8:2006+AMD1:2012, IEC 60601-1-10:2007, IEC 60601-1-11:2015,
- 370 IEC 60601-1-12:2014, IEC 60601-2-2:2009, IEC 62304:2006+AMD1:2015,
- 371 IEC 62366-1:2015, ISO 80601-2-12:—, ISO 80601-2-74:2017, ISO 80601-2-84:— and the
- 372 following apply.
- ISO and IEC maintain terminological databases for use in standardization at the following
- 374 addresses:
- 375 IEC Electropedia: available at http://www.electropedia.org/
- ISO Online browsing platform: available at http://www.iso.org/obp
- NOTE An alphabetized index of defined terms is found Annex EE
- 378 **201.3.201** <u>oSIST prEN ISO 80601-2-87:2020</u>
- 379 high-frequency ventilatorards.iteh.ai/catalog/standards/sist/8a63d826-b514-44c9-aeb5-
- 7845635478ef/osist-pren-iso-80601-2-87-2020
- 381 *ME equipment* intended to provide *ventilation* of the *lungs* of the *patient* when connected
- to the airway of the *patient* using a *rate* greater than 150 *inflations*/min
- Note 1 to entry: Inflation rates are specified as per minute solely when differentiating from conventional-
- rate ventilation. All normative requirements regarding HFV are written using inflation rates per second.
- 385 **201.3.202**
- 386 HFV breathing system
- pathways through which gas flows to or from the *HFV* and to or from the *patient*
- 388 **201.3.203**
- 389 **HFV inflation**
- periodic *ventilator* action intended to increase the volume of gas in the *lungs*
- 391 **201.3.204**
- 392 HFV volume
- volume of gas delivered through the *patient-connection port* or at the distal outlet of the
- jet system during an *HFV inflation*
- Note 1 to entry: The effective volume delivered to the *lung* can be significantly smaller than the *HFV volume*.
- The leakage of uncuffed tracheal tubes and even small changes in resistance or compliance of the
- respiratory system (e.g. by secretion in the airways, through the use of a different *HFV breathing system* or
- tracheal tube) can change the volume delivered to the *lung*.
- Note 2 to entry: The achievable *HFV volume* depends characteristically on the *HFV frequency*. In general,
- lower *HFV frequencies* permit higher *HFV volumes*.