

INTERNATIONAL STANDARD

NORME INTERNATIONALE



Medical electrical equipment –
Part 2-78: Particular requirements for basic safety and essential performance of
medical robots for rehabilitation, assessment, compensation or alleviation

Appareils électromédicaux – [IEC 80601-2-78:2019](#)
Partie 2-78: Exigences particulières pour la sécurité de base et les performances
essentiels des robots médicaux dédiés à la rééducation, l'évaluation, la
compensation ou l'atténuation



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NORME INTERNATIONALE



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CONTENTS

FOREWORD.....	4
INTRODUCTION.....	6
201.1 Scope, object and related standards	7
201.2 Normative references.....	9
201.3 Terms and definitions.....	9
201.4 General requirements	13
201.5 General requirements for testing of ME EQUIPMENT	14
201.6 Classification of ME EQUIPMENT and ME SYSTEMS.....	14
201.7 ME EQUIPMENT identification, marking and documents	14
201.8 Protection against electrical HAZARDS from ME EQUIPMENT.....	15
201.9 Protection against MECHANICAL HAZARDS of ME EQUIPMENT and ME SYSTEMS.....	15
201.10 Protection against unwanted and excessive radiation HAZARDS	21
201.11 Protection against excessive temperatures and other HAZARDS	21
201.12 Accuracy of controls and instruments and protection against hazardous outputs	21
201.13 HAZARDOUS SITUATIONS and fault conditions for ME EQUIPMENT	22
201.14 PROGRAMMABLE ELECTRICAL MEDICAL SYSTEMS (PEMS).....	22
201.15 Construction of ME EQUIPMENT	22
201.16 ME SYSTEMS	24
201.17 ELECTROMAGNETIC COMPATIBILITY of ME EQUIPMENT and ME SYSTEMS	24
202 ELECTROMAGNETIC DISTURBANCES – Requirements and tests	24
206 USABILITY	24
208 * General requirements, tests and guidance for alarm systems in MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS	25
210 * Process requirements for the development of PHYSIOLOGIC CLOSED-LOOP CONTROLLERS	25
211 * Requirements for MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS used in the HOME HEALTHCARE ENVIRONMENT	26
Annexes	27
Annex A (informative) General guidance and rationale.....	27
Annex AA (informative) Particular guidance and rationale.....	28
Annex BB (informative) Guidance and examples of SITUATION AWARENESS.....	57
Bibliography.....	71
Index of defined terms used in this particular standard.....	74
Figure AA.1 – Relationship of the terms used to describe equipment, accessories or equipment parts	29
Figure AA.2 – Endsley's model of SITUATION AWARENESS (based on [10], drawn by Dr. Peter Lankton, May 2007).....	33
Figure AA.3 – Model of user-medical device interaction	34
Figure AA.4 – RACA ROBOT shared control system block diagram: control by PATIENT and RACA ROBOT	38
Figure AA.5 – RACA ROBOT shared control system block diagram: control by PATIENT, OPERATOR and RACA ROBOT	39

Figure AA.6 – RACA ROBOT shared control system block diagram: control by PATIENT and RACA ROBOT, and control modulation by OPERATOR	40
Figure AA.7 – WALKING RACA ROBOT using motion-related biosignal input.....	41
Figure AA.8 – System block diagram of a WALKING RACA ROBOT using a motion-related biosignal as input.....	41
Figure AA.9 – RACA ROBOT that is an arm exoskeleton for REHABILITATION that applies a PATIENT-cooperative shared control strategy	42
Figure AA.10 – System block diagram of an arm exoskeleton for REHABILITATION that applies a PATIENT-cooperative shared control strategy	43
Figure AA.11 – Cane-type RACA ROBOT for REHABILITATION of walking.....	44
Figure AA.12 – System block diagram of a cane-type RACA ROBOT	44
Figure AA.13 – Example of ROBOT arm type RACA ROBOT for lower extremities	45
Figure AA.14 – Example of ROBOT arm type RACA ROBOT for upper extremities	46
Figure AA.15 – Example of exoskeleton type RACA ROBOT for upper extremities	47
Figure AA.16 – Example of exoskeleton type RACA ROBOT for knee joint.....	49
Figure AA.17 – Example of soft artificial muscle-type RACA ROBOT for knee joint	50
Figure AA.18 – Example of exoskeleton-type WALKING RACA ROBOT	51
Figure AA.19 – Example of RACA ROBOT for balance control	52
Figure AA.20 – Example of a body-weight support-type RACA ROBOT with gait following function.....	54
Figure BB.1 – All the proximate causes of loss of SITUATION AWARENESS [19]	58
Figure BB.2 – Relationship between SITUATION AWARENESS, the RISK MANAGEMENT PROCESS (ISO 14971:2007) and the USABILITY ENGINEERING PROCESS (IEC 62366-1:2015)	60
Figure BB.3 – Relationship between GDTA and RISK MANAGEMENT and USABILITY ENGINEERING PROCESSES.....	63
Figure BB.4 – WALKING exoskeleton RACA ROBOT	69
Table 201.101 – List of potential ESSENTIAL PERFORMANCES.....	14
Table 19 – MECHANICAL HAZARDS covered by this clause.....	16
Table 201.102 – Overview of different stopping procedures	16
Table 28 – Mechanical strength test applicability	23
Table 1 – Mechanical strength test applicability, non-TRANSIT-OPERABLE	26
Table 2 – Mechanical strength test applicability, TRANSIT-OPERABLE	26
Table AA.1 – Correlation mapping between Figure AA.2 and Figure AA.3.....	35
Table BB.1 – Example of using GDTA in BB.5.2	64
Table BB.2 – Example of using GDTA in BB.5.3	66
Table BB.3 – Example of using GDTA in BB.5.4	68
Table BB.4 – Example of using GDTA in BB.5.5	70

INTERNATIONAL ELECTROTECHNICAL COMMISSION

MEDICAL ELECTRICAL EQUIPMENT –

Part 2-78: Particular requirements for basic safety and essential performance of medical robots for rehabilitation, assessment, compensation or alleviation

FOREWORD

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International standard IEC 80601-2-78 has been prepared by IEC subcommittee 62D: Electromedical equipment, of IEC Technical Committee 62: Electrical equipment in medical practice, and ISO Technical Committee 299: Robotics.

This publication is published as a double logo standard.

The text of this International Standard is based on the following documents:

FDIS	Report on voting
62D/1676/FDIS	62D/1688/RVD

Full information on the voting for the approval of this International Standard can be found in the report on voting indicated in the above table.

This document 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 STANDARD, IN THIS PARTICULAR STANDARD OR AS NOTED: SMALL CAPITALS.

In referring to the structure of this document, the term

- "clause" means one of the seventeen numbered divisions within the table of contents, inclusive of all subdivisions (e.g. Clause 7 includes subclauses 7.1, 7.2, etc.);
- "subclause" means a numbered subdivision of a clause (e.g. 7.1, 7.2 and 7.2.1 are all subclauses of Clause 7).

References to clauses within this document are preceded by the term "Clause" followed by the clause number. References to subclauses within this particular standard 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 Clause 7 of the 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 a permissible way to achieve compliance with a requirement or test.

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.

A list of all parts of the IEC 80601 and IEC 60601 International Standard, published under the general title *Medical electrical equipment*, can be found on the IEC website.

The committee has decided that the contents of this document will remain unchanged until the stability date indicated on the IEC website under "<http://webstore.iec.ch>" in the data related to the specific document. At this date, the document will be

- reconfirmed,
- withdrawn,
- replaced by a revised edition, or
- amended.

NOTE The attention of users of this document is drawn to the fact that equipment manufacturers and testing organizations may need a transitional period following publication of a new, amended or revised IEC publication in which to make products in accordance with the new requirements and to equip themselves for conducting new or revised tests. It is the recommendation of the committees that the content of this publication be adopted for implementation nationally not earlier than 3 years from the date of publication.

IMPORTANT – The 'colour inside' logo on the cover page of this publication indicates that it contains colours which are considered to be useful for the correct understanding of its contents. Users should therefore print this document using a colour printer.

INTRODUCTION

This part of IEC 80601 International Standard was written at a time when technical evolution of MEDICAL ROBOTS was in rapid progress and the scientific foundation of safe use was still being expanded.

This document is the result of work that began in ISO/TC 184/SC 2/WG 7 in October 2006 on personal care ROBOTS, to address an emerging type of MEDICAL ROBOT that was used outside of an industrial environment. That group was working on a new standard, ISO 13482, which was published as an International Standard (IS) in 2014. While initially focused on non-medical applications, WG 7 recognized that work was needed on medical devices utilizing robotic technology. In October 2009, ISO/TC 184/SC 2 established a WG 7, *Study Group (SG) on Medical care robots*, comprised of experts from Canada, France, Germany, Japan, Korea, Romania, Switzerland, UK and USA.

The work of ISO/TC 184/SC 2/WG 7 SG cumulated in a proposal to form a Joint Working Group (JWG 9) with IEC/TC 62/SC 62A focusing on MEDICAL ELECTRICAL EQUIPMENT using robotic technology. This JWG began developing a technical report (IEC TR 60601-4-1) dealing with degree of autonomy. While developing this document, a particular standard was deemed required for REHABILITATION type ROBOTS. This led to the creation of a Joint Working Group 36 (MEDICAL ROBOTS for REHABILITATION) in April, 2015 within IEC/TC 62/SC 62D to develop particular requirements of SAFETY of MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS for REHABILITATION type ROBOTS. ISO/TC 184/SC 2 has since been promoted to ISO/TC 299, and JWG 9 has merged with JWG35 and 36 to form JWG 5 (MEDICAL ROBOT Safety) on the ISO side. This proposal was approved from both IEC and ISO and work began.

(standards.iteh.ai)

The minimum safety requirements specified in this particular standard are presented to provide for an acceptable degree of BASIC SAFETY and ESSENTIAL PERFORMANCE for MEDICAL ROBOTS that physically interact with a PATIENT with an IMPAIRMENT to support or perform REHABILITATION, ASSESSMENT, COMPENSATION or ALLEVIATION related to the PATIENT'S MOVEMENT FUNCTIONS .

The requirements are followed by particular specifications for the relevant tests.

MEDICAL ELECTRICAL EQUIPMENT –

Part 2-78: Particular requirements for basic safety and essential performance of medical robots for rehabilitation, assessment, compensation or alleviation

201.1 Scope, object and related standards

Clause 1 of the general standard¹ applies, except as follows:

201.1.1 Scope

Replacement:

This part of IEC 80601 applies to the general requirements for BASIC SAFETY and ESSENTIAL PERFORMANCE of MEDICAL ROBOTS that physically interact with a PATIENT with an IMPAIRMENT to support or perform REHABILITATION, ASSESSMENT, COMPENSATION or ALLEVIATION related to the PATIENT'S MOVEMENT FUNCTIONS, as intended by the MANUFACTURER.

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.

NOTE See also 4.2 of the general standard.

This particular standard does not apply to <https://www.iso.org/standards/sist/17a468e6-beea-48d5-8f7f-dfc0bc969fd5/iec-80601-2-78-2019>

- external limb prosthetic devices (use ISO 22523),
- electric wheelchairs (use ISO 7176 (all parts)),
- diagnostic imaging equipment (e.g. MRI, use IEC 60601-2-33), and
- personal care ROBOTS (use ISO 13482).

201.1.2 Object

Replacement:

The object of this particular standard is to establish particular BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for MEDICAL ROBOTS that physically interact with a PATIENT with an IMPAIRMENT, to support or perform REHABILITATION, ASSESSMENT, COMPENSATION or ALLEVIATION related to the PATIENT'S MOVEMENT FUNCTIONS.

201.1.3 Collateral standards

Addition:

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

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

IEC 60601-1-2:2014, IEC 60601-1-6:2010 and IEC 60601-1-6:2010/AMD1:2013, IEC 60601-1-8:2006 and IEC 60601-1-8:2006/AMD1:2012, IEC 60601-1-10:2007 and IEC 60601-1-10:2007/AMD1:2013, and IEC 60601-1-11:2015 apply as modified in Clauses 202, 206, 208, 210 and 211 respectively. IEC 60601-1-3 and IEC 60601-1-12 do 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 and collateral standards as appropriate for the particular ME EQUIPMENT under consideration, and may add other BASIC SAFETY and ESSENTIAL PERFORMANCE requirements.

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

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

The numbering of clauses and subclauses of this particular standard 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 "20x" where x is the final digit(s) of the collateral standard document number (e.g. 202.4 in this particular standard addresses the content of Clause 4 of the 60601-1-2 collateral standard, 203.4 in this particular standard addresses the content of Clause 4 of the IEC 60601-1-3 collateral standard, etc.). The changes to the text of the general standard are specified by the use of the following words:

[IEC 80601-2-78:2019](https://standards.iteh.ai/catalog/standards/sist/17a468e6-beea-48d5-8f7f-69055-21e)

<https://standards.iteh.ai/catalog/standards/sist/17a468e6-beea-48d5-8f7f-69055-21e>

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

"*Addition*" means that the text of this particular standard 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 particular standard.

Subclauses, figures or tables which 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 standard taken together.

Where there is no corresponding clause or subclause in this particular standard, 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 particular standard.

201.2 Normative references

NOTE Informative references are listed in the Bibliography.

Clause 2 of the general standard 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*
IEC 60601-1-6:2010/AMD1:2013

IEC 60601-1-8:2006, *Medical electrical equipment – Part 1-8: General requirements for safety and essential performance – Collateral Standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems*
IEC 60601-1-8:2006/AMD1:2012

ISO 14971:2007, *Medical devices – Application of risk management to medical devices*

Addition:

iTeh STANDARD PREVIEW

IEC 60601-1-10:2007, *Medical electrical equipment – Part 1-10: General requirements for basic safety and essential performance – Collateral Standard: Requirements for the development of physiologic closed-loop controllers*
IEC 60601-1-10:2007/AMD1:2013

<https://standards.itech.ai/catalog/standards/sist/17a468e6-beea-48d5-8f7f-dfc0bc969fd5/iec-80601-2-78-2019>

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 62366-1:2015, *Medical devices – Part 1: Application of usability engineering to medical devices*

ISO 22523:2006, *External limb prostheses and external orthoses – Requirements and test methods*

201.3 Terms and definitions

For the purposes of this document, the terms and definitions given in IEC 60601-1:2005 and IEC 60601-1:2005/AMD1:2012 apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- IEC Electropedia: available at <http://www.electropedia.org/>
- ISO Online browsing platform: available at <http://www.iso.org/obp>

NOTE An Index of defined terms is found beginning on page 74.

Amendment:

201.3.63
*** MEDICAL ELECTRICAL EQUIPMENT**
ME EQUIPMENT

Addition:

Note 1 to entry: See Figure AA.1.

201.3.65
MOBILE

Addition:

Note 1 to entry: MOBILE also includes equipment intended to support the movement of a PATIENT from one location to another.

201.3.144
*** BODY-WORN**

Replacement:

PORTABLE equipment whose INTENDED USE includes operation while being worn by a PATIENT or attached to a PATIENT'S clothing

Addition:

201.3.201
***ACTUATED APPLIED PART**

STANDARD PREVIEW
 (Standard not for sale)

subcategory of APPLIED PART that is intended to provide actively controlled physical interactions with the PATIENT that are related to the PATIENT'S MOVEMENT FUNCTIONS, to perform a CLINICAL FUNCTION of a RACA ROBOT

[IEC 80601-2-78:2019](http://www.iec.ch/iec-80601-2-78-2019)

Note 1 to entry: "Actively controlled" as used above is intended to mean controlled by the RACA ROBOT, including shared control with the PATIENT or OPERATOR.

<http://www.iec.ch/iec-80601-2-78-2019>

Note 2 to entry: Actively controlled physical interactions include position control, force control, impedance control, admittance control, or any other controls that regulate the interaction between a RACA ROBOT and the PATIENT.

Note 3 to entry: Each ACTUATED APPLIED PART is part of an actuation system according to 9.8.1 of the general standard.

201.3.202
ALLEVIATION

treatment to ease symptoms due to an IMPAIRMENT of a PATIENT

Note 1 to entry: An example of ALLEVIATION is physical therapy performed by a RACA ROBOT to reduce pain or other secondary effects of an IMPAIRMENT.

201.3.203
*** ASSESSMENT**

procedure to quantify or to aid in the qualification of the level of IMPAIRMENT of a PATIENT.

Note 1 to entry: The term "ASSESSMENT" should not be confused with the term "RISK ASSESSMENT"

Note 2 to entry: ASSESSMENT in this definition focuses on the level of IMPAIRMENT of body functions and structures and not of activity limitations or participation restrictions ([1]², Chapter 4).

Note 3 to entry: ASSESSMENT can be distinguished from both measurement and diagnosis. Measurement typically refers to the physiological parameter being measured (e.g. PATIENT hand position by a RACA ROBOT) and is typically related to, or is the direct output of a transducer. Those measurements are then used to quantify an IMPAIRMENT (e.g., proprioception, planning of a movement, etc). Because IMPAIRMENTS are not the same as the underlying pathology, but are the manifestations of an underlying pathology, an ASSESSMENT that quantifies the IMPAIRMENT

² Numbers in square brackets refer to the Bibliography.

can then be used by a clinician to support a diagnosis that identifies the pathology, or to monitor the progression of a pathology or recovery from an injury. See Annex AA for an example.

201.3.204

CLINICAL FUNCTION

clinically significant operation that the ME EQUIPMENT or ME SYSTEM is intended to perform, as specified by the MANUFACTURER

Note 1 to entry: CLINICAL FUNCTION is generally a subset of the INTENDED USE of the ME EQUIPMENT or ME SYSTEM and is related to the PATIENT.

Note 2 to entry: In the context of this document, a clinically significant operation refers to:

- support of the diagnosis of a PATIENT with an IMPAIRMENT,
- treatment, mitigation, or monitoring of an IMPAIRMENT of a PATIENT.

[SOURCE: IEC TR 60601-4-1:2017, 3.6, modified – In the definition, the words "medical operation" have been replaced by "clinically significant operation", and Note 2 to entry has been added.]

201.3.205

COMPENSATION

mitigation of IMPAIRMENT of a PATIENT through support of body structures or through support or replacement of body functions

Note 1 to entry: COMPENSATION could be provided by, for example, externally powered orthoses. COMPENSATION does not include the improvement of MOVEMENT FUNCTIONS related to an IMPAIRMENT, which is defined as REHABILITATION.

201.3.206

EMERGENCY STOP

manually initiated interruption of operation intended to stop the RACA ROBOT to prevent HARM

<https://standards.iteh.ai/catalog/standards/sist/17a468e6-beea-48d5-8f7f-426a5a1e300e-1-2019>

Note 1 to entry: EMERGENCY STOP is meant as a last resort when no other means to reduce RISK are available.

Note 2 to entry: EMERGENCY STOP does not allow an automatic restart, in contrast to a PROTECTIVE STOP.

Note 3 to entry: The activation of an EMERGENCY STOP does not necessarily shut down all the power in a RACA ROBOT. Refer to 9.2.4 of the general standard for further clarification.

201.3.207

* IMPAIRMENT

problem in body function or structure, such as significant deviation or loss

Note 1 to entry: "Problems" here is used to refer to a (negative) deviation from generally accepted population normality levels in the biomedical status of the body and its functions. Notification of the constituents of IMPAIRMENT is undertaken primarily by those qualified to judge functioning according to the generally accepted normality levels. An IMPAIRMENT usually follows a disease or accident, but also includes birth defects.

Note 2 to entry: This document only concerns ME EQUIPMENT or ME SYSTEM, meaning that it addresses IMPAIRMENT of a PATIENT and not normal age-related IMPAIRMENTS.

[SOURCE: ISO 9999:2016, 2.11, modified – The term is here in singular instead of plural. The notes to entry have been added.]

201.3.208

MEDICAL ROBOT

ROBOT intended to be used as ME EQUIPMENT or ME SYSTEM

[SOURCE: IEC TR 60601-4-1:2017, 3.20]

201.3.209**MOVEMENT FUNCTION**

function consisting of one or more of sensory, neuromusculoskeletal or movement-related body functions that comprise PATIENT motor control

Note 1 to entry: "Sensory", "neuromusculoskeletal" and "movement-related" come from the body functions classification presented in [1].

Note 2 to entry: A RACA ROBOT does not need to address all of these aspects.

201.3.210*** PROTECTIVE STOP**

interruption of operation automatically initiated by the RACA ROBOT, that allows a cessation of motion for BASIC SAFETY and ESSENTIAL PERFORMANCE purposes, which could allow the PROGRAMMABLE ELECTRICAL MEDICAL SYSTEM (PEMS)/PROGRAMMABLE ELECTRONIC SUBSYSTEM (PESS) to facilitate a restart

Note 1 to entry: See also ISO 8373:2012, 5.17.

Note 2 to entry: A PROTECTIVE STOP is defined in contrast to an EMERGENCY STOP or a normal stop; clarification is provided in Table 201.102.

Note 3 to entry: "Automatically initiated" means that the RACA ROBOT is detecting a potentially HAZARDOUS SITUATION and as a result of this detection the RACA ROBOT then initiates a stopping procedure

Note 4 to entry: The defined term is related to FALLBACK MODE defined in IEC 60601-1-10:2007, 3.11.

Note 5 to entry: Even during the cessation of motion, the actuators are not necessarily deactivated. In certain conditions, the actuators may hold the PATIENT or PATIENT'S limbs in a stable position.

201.3.211*** REHABILITATION**

treatment to improve MOVEMENT FUNCTIONS related to an IMPAIRMENT of a PATIENT

<https://standards.iteh.ai/catalog/standards/sist/17a468e6-beea-48d5-8f7f-dfbb591cc050/iec-80601-2-78-2019>

EXAMPLE Relearning of motor control of the upper or lower extremities, restoration of muscle strength and endurance.

Note 1 to entry: REHABILITATION can be carried out in situations where a PATIENT has an IMPAIRMENT due to an accident or disease or congenital conditions (e.g. cerebral palsy) or can be used to slow down the expected loss of body functions due to neurodegenerative diseases (e.g. Parkinson's, multiple sclerosis).

Note 2 to entry: This definition differs from the World Health Organization (WHO) definition and covers only a subset of that definition, in order to agree with the scope of this document.

201.3.212**REHABILITATION, ASSESSMENT, COMPENSATION AND ALLEVIATION ROBOT****RACA ROBOT**

MEDICAL ROBOT intended by its MANUFACTURER to perform REHABILITATION, ASSESSMENT, COMPENSATION or ALLEVIATION comprising an ACTUATED APPLIED PART

201.3.213***ROBOT**

programmed actuated mechanism with a degree of autonomy, moving within its environment, to perform intended tasks

201.3.214*** SITUATION AWARENESS**

OPERATOR'S perception, comprehension and projection of a RACA ROBOT'S behaviour in its environment

Note 1 to entry: SITUATION AWARENESS allows the OPERATOR of the RACA ROBOT to react appropriately and timely to the RACA ROBOT'S behaviour both under NORMAL USE and in the case of a SINGLE FAULT CONDITION.

Note 2 to entry: SITUATION AWARENESS has a direct link to USABILITY ENGINEERING PROCESS. However, IEC 62366-1:2015 relates to SITUATION AWARENESS through the use of the concepts of "perception" and "cognition". Annex AA provides additional information.

201.3.215*** WALKING**

MOBILE equipment that, once installed and placed into service, is intended to move from one location to another, by making reciprocating motion having intermittent contact with the travel surface and the RACA ROBOT

EXAMPLE A powered exoskeleton for the lower limbs is an example of a WALKING RACA ROBOT that is intended by the MANUFACTURER to ambulate the PATIENT.

Note 1 to entry: WALKING is a new subcategory of TRANSPORTABLE ME EQUIPMENT and ME SYSTEM as shown in Figure AA.1.

201.4 General requirements

Clause 4 of the general standard applies, except as follows:

201.4.2.3 Evaluating RISK

Additional subclauses before 4.2.3.1:

201.4.2.3.101 General

In performing a RISK MANAGEMENT PROCESS, the degree of autonomy of the RACA ROBOT shall be taken into consideration.

NOTE 1 IEC TR 60601-4-1 provides considerations on the effects that the degree of autonomy could have on the RISK MANAGEMENT PROCESS. For example, RACA ROBOTS could accomplish a task fully autonomously with less RISK than an OPERATOR performing the same intended task, or vice versa.

NOTE 2 Degree of autonomy has the potential for increasing performance, adding functionality or improving USABILITY of RACA ROBOTS. However, a higher degree of autonomy can increase the complexity of the system and thus can increase the number and complexity of HAZARDOUS SITUATIONS. It is important the MANUFACTURER takes this complexity into account in its RISK MANAGEMENT PROCESS. As one or more PESS provide the autonomy, these PESS deserve specific scrutiny.

NOTE 3 The role of the OPERATOR can differ depending on their tasks and interactions with the RACA ROBOT. The SITUATION AWARENESS of the OPERATOR can become more important for RACA ROBOTS with a higher degree of autonomy.

Compliance is checked by inspection of the RISK MANAGEMENT FILE.

201.4.2.3.102 * RISK MANAGEMENT and SITUATION AWARENESS

The MANUFACTURER shall identify the HAZARDS or HAZARDOUS SITUATIONS that could occur following a loss of SITUATION AWARENESS in the RISK MANAGEMENT PROCESS, when an OPERATOR is needed to supervise a task of, or interact with, a RACA ROBOT to reduce RISK to an acceptable level. The MANUFACTURER shall include the necessary information in the ACCOMPANYING DOCUMENTS.

Compliance shall be checked by inspection of the RISK MANAGEMENT FILE and, where applicable, of the ACCOMPANYING DOCUMENTS.

IEC 62366-1:2015, 5.5 requires HAZARD-RELATED USE SCENARIOS to be evaluated using a SUMMATIVE EVALUATION. Because of the subjective nature of evaluating SITUATION AWARENESS, it can be considered sufficient to perform a FORMATIVE EVALUATION of this HAZARD-RELATED USE SCENARIO instead.

NOTE Additional guidance can be found in Annex BB.

201.4.3 ESSENTIAL PERFORMANCE

Addition: