



Edition 3.0 2010-03

INTERNATIONAL STANDARD

NORME INTERNATIONALE



Medical electrical equipment -

Part 2-33: Particular requirements for the basic safety and essential performance of magnetic resonance equipment for medical diagnosis

Appareils électromédicaux –

Partie 2-33: Exigences particulières pour la sécurité de base et les performances essentielles des appareils à résonance magnétique utilisés pour le diagnostic médical





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IEC 60601-2-33

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CONTENTS

FOREWORD				
INTRODU	CTION	7		
201.1	Scope, object and related standards	8		
201.2	Normative references	9		
201.3	Terms and definitions	10		
201.4	General requirements	15		
201.5	General requirements for testing of ME EQUIPMENT	15		
201.6	01.6 Classification of ME EQUIPMENT and ME SYSTEMS			
201.7	ME EQUIPMENT identification, marking and documents	16		
201.8	Protection against electrical HAZARDS from ME EQUIPMENT	27		
201.9	Protection against mechanical HAZARDS of ME EQUIPMENT and ME SYSTEMS	28		
201.10	Protection against unwanted and excessive radiation HAZARDS	28		
201.11	Protection against excessive temperatures and other HAZARDS	28		
201.12	Accuracy of controls and instruments and protection against hazardous outputs	29		
201.13	HAZARDOUS SITUATIONS and fault conditions	47		
201.14	PROGRAMMABLE ELECTRICAL MEDICAL SYSTEM (PEMS)	47		
201.15	Construction of ME EQUIPMENT	47		
201.16	ME SYSTEMS	47		
201.17	Electromagnetic compatibility of ME EQUIPMENT and ME SYSTEMS	47		
202	Electromagnetic compatibility – Requirements and tests	48		
Annexes .		48		
Annex D (informative) Symbols on marking				
Annex AA (informative) Particular guidance and rationale				
Bibliograp	vhy	96		
Index of d	efined terms used in this particular standard	104		
Figure 20	1.101 – Gradient waveform and EFFECTIVE STIMULUS DURATION	11		
Figure 201.102 – Limits for cardiac and peripheral nerve stimulation				
Figure 201.103 – Reduction of WHOLE BODY SAR limits at high temperatures				
Figure 201.104 – Volume for determining the spatial maximum of gradient output43				
Figure 201.105 – Volume for determining the <i>B</i> ₁ stray field				
Figure 201.D.101 – Signs indicating a transmit only RF coil, transmit / receive RF coil and a receive only RF coil				
Figure AA	1 – Static magnetic fields: flow potentials and retardation	68		
Figure AA	2 – Experimental data on PNS threshold of human volunteers in WHOLE EQUIPMENT	83		
Figure AA nerve stim	3 – Double logarithmic plot of experimental threshold values for peripheral nulation	84		
Figure AA stimulus c	A.4 – Response value $R(t)$ generated by convolution of a rectangular IB/dt and a nerve impulse response function $n(t-\theta)$	88		
Figure AA for a trape	1.5 – Gradient waveform G, stimulus waveform dB/dt and response value R, ezoid EPI waveform starting at $t = 0$	89		

Figure AA.7 – Threshold value of d <i>B/dt</i> for a sinusoid gradient waveform, as function of the number of half periods in the waveform	Figure AA.6 – Threshold values d <i>B</i> /d <i>t</i> for two gradient waveforms, plotted against EFFECTIVE STIMULUS DURATION	.89
Figure AA.8 – SAR limits for the exposed mass of a PATIENT 93 Fable 201.101 – List of symbols. 15 Fable 201.102 – Rheobase values per type of gradient system 32 Fable 201.103 – Weight factors for summation of the maximum output <i>O₁</i> per 34 Fable 201.104 – Temperature limits 34 Fable 201.105 – SAR limits for volume transmit coils 35 Fable 201.106 – SAR limits for local transmit coils 36 Fable 201.101 – Examples of warning signs and prohibitive signs) 49 Fable AA.1 – Static field occupational standards 67	Figure AA.7 – Threshold value of dB/dt for a sinusoid gradient waveform, as function of the number of half periods in the waveform	.90
Fable 201.101 – List of symbols 15 Fable 201.102 – Rheobase values per type of gradient system 32 Fable 201.103 – Weight factors for summation of the maximum output O _I per 34 Fable 201.104 – Temperature limits 34 Table 201.105 – SAR limits for volume transmit coils 35 Fable 201.106 – SAR limits for local transmit coils 36 Fable 201.010 – Examples of warning signs and prohibitive signs): 49 Fable 201.010 – Examples of warning signs and prohibitive signs): 67 Fable AA.1 – Static field occupational standards 67	Figure AA.8 – SAR limits for the exposed mass of a PATIENT	.93
Fable 201.101 – List of symbols 15 Fable 201.102 – Rheobase values per type of gradient system 32 Fable 201.103 – Weight factors for summation of the maximum output O _i per 34 Fable 201.104 – Temperature limits 34 Fable 201.105 – SAR limits for volume transmit coils 35 Fable 201.106 – SAR limits for local transmit coils 36 Fable 201.010 – SAR limits for local transmit coils 36 Fable 201.010 – SAR limits for local transmit coils 36 Fable 201.010 – SAR limits for local transmit coils 36 Fable 201.010 – SAR limits for local transmit coils 36 Fable 201.010 – SAR limits for local transmit coils 36 Fable 201.010 – SAR limits for local transmit coils 37 Fable AA.1 – Static field occupational standards 67		
Fable 201.102 – Rheobase values per type of gradient system 32 Fable 201.103 – Weight factors for summation of the maximum output O _j per 34 SRADIENT UNIT 34 Fable 201.104 – Temperature limits. 34 Fable 201.105 – SAR limits for volume transmit coils 35 Fable 201.106 – SAR limits for local transmit coils 36 Fable 201.0101 – Examples of warning signs and prohibitive signs): 49 Table AA.1 – Static field occupational standards 67	Table 201.101 – List of symbols	. 15
Fable 201.103 – Weight factors for summation of the maximum output O _j per 34 SRADIENT UNIT 34 Fable 201.104 – Temperature limits 34 Fable 201.105 – SAR limits for volume transmit coils 35 Fable 201.106 – SAR limits for local transmit coils 36 Fable 201.107 – Examples of warning signs and prohibitive signs): 49 Fable 201.01 – Examples of warning signs and prohibitive signs): 49 Fable AA.1 – Static field occupational standards 67	Table 201.102 – Rheobase values per type of gradient system	. 32
Fable 201.104 - Temperature limits 34 Fable 201.105 - SAR limits for volume transmit coils 35 Fable 201.106 - SAR limits for local transmit coils 36 Fable 201.010 - Examples of warning signs and prohibitive signs): 49 Fable 201.01 - Examples of warning signs and prohibitive signs): 49 Fable AA.1 - Static field occupational standards 67	Table 201.103 – Weight factors for summation of the maximum output O_i per	31
Fable 201.105 – SAR limits for volume transmit coils	Table 201 104 - Temperature limits	34
Table 201.100 – SAR limits for local transmit coils	Table 201 105 – SAR limits for volume transmit coils	35
Fable 201.D.101 – Examples of warning signs and prohibitive signs):	Table 201 106 – SAR limits for local transmit coils	36
Fable AA.1 – Static field occupational standards	Table 201 D 101 – Examples of warning signs and prohibitive signs).	.30 49
is a orall in the state of the	Table AA 1 – Static field occupational standards	.40
S A Drouie and a second and a second		.01
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INTERNATIONAL ELECTROTECHNICAL COMMISSION

MEDICAL ELECTRICAL EQUIPMENT -

Part 2-33: Particular requirements for the basic safety and essential performance of magnetic resonance equipment for medical diagnosis

FOREWORD

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International standard IEC 60601-2-33 has been prepared by IEC subcommittee 62B: Diagnostic imaging equipment, of IEC technical committee 62: Electrical equipment in medical practice.

This third edition cancels and replaces the second edition published in 2002, its Amendment 1 (2005) and Amendment 2 (2007) and constitutes a technical revision. This third edition of IEC 60601-2-33 is based on the second amendment to Edition 2. It has also been adapted to the third edition of IEC 60601-1 (2005), with technical modifications being introduced where appropriate.

The text of this particular standard is based on the following documents:

FDIS	Report on voting
62B/777/FDIS	62B/782/RVD

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

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

In this standard, 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 standard, 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 standard are preceded by the term "Clause" followed by the clause number. References to subclauses within this particular standard are by number only.

In this standard, 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 standard conform to usage described in Annex H of the ISO/IEC Directives, Part 2. For the purposes of this standard, the auxiliary verb:

- "shall" means that compliance with a requirement or a test is mandatory for compliance with this standard;
- "should" means that compliance with a requirement or a test is recommended but is not mandatory for compliance with this standard;
- "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 60601 series, published under the general title: *Medical electrical equipment*, can be found on the IEC website.

The committee has decided that the contents of this amendment and the base publication will remain unchanged until the stability date indicated on the IEC web site under "http://webstore.iec.ch" in the data related to the specific publication. At this date, the publication will be

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

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* INTRODUCTION

This particular standard is written at a moment in which the technical evolution of MR EQUIPMENT is in rapid progress and the scientific foundation of its safe use is still expanding.

This International Standard addresses technical aspects of the medical diagnostic MR SYSTEM and the MR EQUIPMENT therein related to the safety of PATIENTS examined with this system, the safety of the MR WORKER involved with its operation and the safety of the MR WORKER involved with the development, manufacturing, installation, and servicing of the MR SYSTEM. Where limits of electromagnetic fields (EMF) exposure of PATIENTS and MR WORKERS are stated, these limits do not imply that such levels of exposure can be assumed to be acceptable for workers in other professional settings and for the population at large. The limits provide a sensible balance between RISKS for the PATIENTS and MR WORKERS and benefits for the PATIENTS.

Organizational aspects of safety are the task of the RESPONSIBLE ORGANIZATION. This task includes adequate training of staff, rules of access to the MR SYSTEM, qualification of staff for decisions that are related to safety, definition of medical responsibility and specific requirements for personnel following from that responsibility when the PATIENT is in or near the MR SYSTEM.

Examples of such organizational aspects are:

- operation in FIRST LEVEL CONTROLLED OPERATING MODE;
- emergency procedures for resuscitation of the PATIENT who is in the MR SYSTEM;
- emergency procedures after a QUENCH of the superconductive magnet when present;
- set-up and maintenance of a protocol for screening the PATIENT for contraindications or for conditions that may affect acceptable exposure;
- rules for ROUTINE MONITORING and for MEDICAL SUPERVISION of the PATIENT during the exam.
- rules to minimize and to limit the exposure of MR WORKERS to EMF.

Extensive rationale is provided in Annex AA for some of the definitions and requirements in order to provide the user of this standard with a reasonably complete access to the source material that was used in support of the considerations during drafting.

The relationship of this particular standard with IEC 60601-1 and the collateral standards is explained in subclauses 201.1.3 and 201.1.4.

The introduced EMF exposure limits required in this standard for an MR WORKER will never exceed those allowed for PATIENTS All exposure limits allowed for a PATIENT and for an MR WORKER are expected to protect them against negative health effects and unacceptable RISKS.

For the exposure to static magnetic fields, subjective short-term physiological and sensory effects are expected. These influence the well being of the MR WORKER marginally and only during or shortly after exposure.

For the exposure to GRADIENT OUTPUT and RF transmit fields, normally no short-term physiological and sensory effects are expected for MR WORKERS.

In addition no experimental or theoretical basis for cumulative biological effects in humans, resulting from exposure at the allowed levels has been generally accepted.

The requirements for acoustic noise exposure are different for PATIENTS and MR WORKERS.

MEDICAL ELECTRICAL EQUIPMENT -

Part 2-33: Particular requirements for the basic safety and essential performance of magnetic resonance equipment for medical diagnosis

201.1 Scope, object and related standards

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

201.1.1 Scope

Replacement:

This International Standard applies to the BASIC SAFETY and ESSENTIAL PERFORMANCE of MR EQUIPMENT and MR SYSTEMS, hereafter referred to also as ME EQUIPMENT.

This standard does not cover the application of MR EQUIPMENT beyond the INTENDED USE.

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.

The standard does not formulate ESSENTIAL PERFORMANCE requirements related to INTERVENTIONAL MR EXAMINATIONS.

201.1.2 Object

Replacement:

The object of this particular standard is to establish particular BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for MR EQUIPMENT to provide protection for the PATIENT and the MR WORKER.

NOTE This standard presumes that the MR WORKERS are properly medically screened, and properly trained and instructed in their duties.

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.

IEC 60601-1-2:2007 applies as modified in Clause 202. IEC 60601-1-3 and IEC 60601-1-10 do not apply. All other published collateral standards in the IEC 60601-1 series apply as published.

201.1.4 Particular standards

Replacement:

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

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 is 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 standard 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 60601-1-3 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 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 3.139, additional definitions in this standard 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 standard" 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 beginning on page 96.

Clause 2 of the general standard applies except as follows:

Replacement:

IEC 60601-1-2:2007, Medical electrical equipment – Part 1-2: General requirements for basic safety and essential performance – Collateral standard: Electromagnetic compatibility – Requirements and tests

Addition:

IEC 60601-1:2005, Medical electrical equipment – Part 1: General requirements for basic safety and essential performance

NEMA MS 4:2006, Acoustic noise measurements procedure for diagnostic magnetic resonance imaging devices

NEMA MS 8:2008, Characterization of the specific absorption rate (SAR) for magnetic resonance imaging systems

201.3 Terms and definitions

For the purposes of this document, the terms and definitions given in IEC 60601-1:2005 and the following apply:

NOTE An index of defined terms is found beginning on page 104. A list of symbols used in the document is provided in Table 201.101.

Addition:

* 201.3.201

B1RMS

root mean square (rms) of B_{1} the radio frequency magnetic induction

$$B_{1} \text{RMS} = \sqrt{\frac{\int_{0}^{t_{x}} (B_{1}(t))^{2} dt}{t_{x}}}$$

where t is time, and t_x is the evaluation time, and is estimated at the RF transmit coil centre.

201.3.202

COMPLIANCE VOLUME

PATIENT accessible space in which compliance of GRADIENT OUTPUT is inspected

In MR EQUIPMENT with a cylindrical WHOLE BODY MAGNET, the COMPLIANCE VOLUME is a cylinder with its axis coinciding with the magnet axis and with a radius of 0,20 m. and with a length equal to the gradient coil

In MR EQUIPMENT with a TRANSVERSE FIELD MAGNET and a WHOLE BODY GRADIENT SYSTEM, the COMPLIANCE VOLUME is a cylinder aligned with the patient's axis, of length equal to the gradient coil diameter, and a diameter of 0,40 m or equal to the distance between the poles of the magnet, whichever is less.

In all other MR EQUIPMENT the COMPLIANCE VOLUME is the volume where any part of a PATIENT body can be properly located according to the INTENDED USE of the MR EQUIPMENT.

201.3.203 CONTROLLED ACCESS AREA area to which access is controlled for safety reasons

201.3.204 CORE TEMPERATURE mean temperature of the body core