Elektrilised meditsiiniseadmed. Osa 1-2: Üldnõuded esmasele ohutusele ja olulistele toimimisnäitajatele. Kollateraalstandard: Elektromagnetiline ühilduvus. Nõuded ja katsetused

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



EESTI STANDARDI EESSÕNA

NATIONAL FOREWORD

Käesolev Eesti standard EVS-EN 60601-1- 2:2007 sisaldab Euroopa standardi EN 60601- 1-2:2007 ingliskeelset teksti.	This Estonian standard EVS-EN 60601-1-2:2007 consists of the English text of the European standard EN 60601-1-2:2007.
Standard on kinnitatud Eesti Standardikeskuse 23.11.2007 käskkirjaga ja jõustub sellekohase teate avaldamisel EVS Teatajas.	This standard is ratified with the order of Estonian Centre for Standardisation dated 23.11.2007 and is endorsed with the notification published in the official bulletin of the Estonian national standardisation organisation.
Euroopa standardimisorganisatsioonide poolt rahvuslikele liikmetele Euroopa standardi teksti kättesaadavaks tegemise kuupäev on 15.07.2007.	Date of Availability of the European standard text 15.07.2007.
Standard on kättesaadav Eesti Standardiorganisatsioonist.	The standard is available from Estonian standardisation organisation.
кор ICS 11.040.01, 33.100.10, 33.100.20	The senerated by The

Standardite reprodutseerimis- ja levitamisõigus kuulub Eesti Standardikeskusele

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

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EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM

EN 60601-1-2

July 2007

ICS 11.040.01; 33.100.10; 33.100.20

Supersedes EN 60601-1-2:2001 + A1:2006

English version

Medical electrical equipment -Part 1-2: General requirements for basic safety and essential performance ateral standard: Electromagnetic compatibility -**Requirements and tests** (IEC 60601-1-2:2007, modified) Medizinische elektrische Geräte -Appareils électromédicau Partie 1-2: Exigences générale Teil 1-2: Allgemeine Festlegungen pour la sécurité de base für die Sicherheit einschließlich et les performances essentielles der wesentlichen Leistungsmerkmale -Norme collatérale: Ergänzungsnorm: Compatibilité électromagnétique Elektromagnetische Verträglichkeit -Anforderungen und Prüfungen Exigences et essais (CEI 60601-1-2:2007, modifiée) (IEC 60601-1-2:2007, modifiziert) This European Standard was approved by CENELEC 02007-04-11. CENELEC 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 Central Secretariat or to any CENELEC member This European Standard exists in three official versions (English, German). A version in any other language made by translation under the responsibility of a CENELEC monoton ber into its own language and notified to the Central Secretariat has the same status as the official versions. CENELEC members are the national electrotechnical committees of Austria Belgium, Bulgaria, Cyprus, the Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, the Netherlands, Norway, Poland, Portugal, Romania Slovakia, Slovenia, Spain, Sweden, Switzerland and the United Kingdom. CENELEC European Committee for Electrotechnical Standardization Comité Européen de Normalisation Electrotechnique Europäisches Komitee für Elektrotechnische Normung Central Secretariat: rue de Stassart 35, B - 1050 Brussels

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Foreword

The text of document 62A/560/FDIS, future edition 3 of IEC 60601-1-2, prepared by SC 62A, Common aspects of electrical equipment used in medical practice, of IEC TC 62, Electrical equipment in medical practice, was submitted to the IEC-CENELEC parallel vote and was approved by CENELEC as EN 60601-1-2 on 2007-04-11.

The following date was fixed:

 latest date by which the EN has to be implemented at national level by publication of an identical national standard or by endorsement

(dop) 2008-02-01

This European Standard supersedes EN 60601-1-2:2001 and its amendment A1:2006. However, EN 60601-1-2:2001 mains valid until all the parts 2 that are used in conjunction with it have been withdrawn. No date of withdrawal of conflicting national standards (dow) has therefore been fixed. However, when Part 1 2 is used for appliances not covered by a part 2, EN 60601-1-2:2001 is not to be used after 2009-09-12.

This European Standard has been prepared under a mandate given to CENELEC by the European Commission and the European Free Trade Association and covers essential requirements of EC Directive 93/42/EEC. See Annex ZZ.

This European Standard constitutes collateral standard to EN 60601-1:2006, hereafter referred to as the general standard.

This EN 60601-1-2 was revised to structurally align it with EN 60601-1:2006 and to implement the decision of IEC SC 62A that the clause numbering structure of collateral standards written to EN 60601-1:2006 would adhere to the form specified in ISO/IEC Directives, Part 2:2004. The principle technical changes are in Clause 4, which now recognizes that there is a general requirement for a risk management process in EN 60601-1:2006.

In the 60601 series of publications, collateral standards specify general requirements for safety applicable to

- a subgroup of MEDICAL ELECTRICAL EQUIPMENT (e.g. radiological equipment); or
- a specific characteristic of all MEDICAL ELECTRICAL EQUIPMENT, not fully addressed in the general standard (e.g. alarm systems).

In this collateral standard the following print types are used:

- requirements and definitions: in roman type;
- test specifications: in 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 COLLATERAL STANDARD OR AS NOTES: IN SMALL CAPITALS.

NOTE Defined terms are not printed in SMALL CAPITALS in Table 1 through Table 8, in the tables in Annex C and in statements required to appear in the technical description or instructions for use because they are intended for the OPERATOR or RESPONSIBLE ORGANIZATION, who may not be familiar with the defined terms of EN 60601 standards.

In referring to the structure of this standard, the term

- "clause" means one of the six numbered divisions within the table of contents, inclusive of all subdivisions (e.g. Clause 6 includes Subclauses 6.1, 6.2, etc.);
- "subclause" means a numbered subdivision of a clause (e.g. 6.1, 6.2 and 6.2.1 are all subclauses of Clause 6).

References to clauses within this standard are preceded by the term "Clause" followed by the clause number. References to subclauses within this 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 text of the International Standard IEC 60601-1-2:2007 was approved by CENELEC as a European Standard with agreed common modifications as given below.

COMMON MODIFICATIONS

Annex A, Subclause 4.2:

Replace the last sentence by:

These HAZARDS shall be considered in the RISK MANAGEMENT PROCESS.

Bibliography			
Add the following notes fo	or the star	ndards indicated:	
CISPR 11 + A1 + A2	NOTE	Harmonized as EN 55011:2007 (modified) and A2:2007 (not modified).	
CISPR 24	NOTE	Harmonized as EN 55024:1998 (modified).	
IEC 61000-4-2 + A1 + A2	NOTE	Harmonized as EN 61000-4-2:1995 + A1:1998 + A2:2001 (not modified).	
IEC 61000-4-3	NOTE	Harmonized as EN 61000-4-3:2006 (not modified).	
IEC 61000-4-4	NOTE	Halonized as EN 61000-4-4:2004 (not modified).	
IEC 61000-4-6 + A1 + A2	NOTE	Harmonized as EN 61000-4-6:2007 (not modified).	
IEC 61326-1	NOTE	Harmonized SEN 61326-1:2006 (not modified).	
ISO 14971	NOTE	Harmonized as Exercise 14971:2000 (not modified).	
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		(A)	
		6.	
		Z	

⁽n)

Annex ZA

(normative)

Normative references to international publications with their corresponding European publications

The following referenced documents are indispensable for the application 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 When an internapplies.	ational publ	lication has been modified by common modifications, ind	icated by (mod), the relev	/ant EN/HD
Publication IEC 60417	Data base	<u>Title</u> Graphical symbols for use on equipment	<u>EN/HD</u> -	<u>Year</u> -
IEC 60601-1	2005	Part 1: General requipment - Part 1: General requirements for basic safet and essential performance	EN 60601-1	2006
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	EN 60601-1-8	2007
IEC 61000-3-2	_ 1)	Electromagnetic compatiblity (EMC) - Part 3-2: Limits - Limits for harmonic current emissions (equipment input current \leq 16 A per phase)	EN 61000-3-2	2006 ²⁾
IEC 61000-3-3	_ 1)	Electromagnetic compatibility (EMP) Part 3-3: Limits - Limitation of voltage changes, voltage fluctuations and flicter in public low-voltage supply systems, for equipment with rated current \leq 16 A per phase and not subject to conditional connection	EN 61000-3-3 + corr. July	1995 ²⁾ 1997
IEC 61000-4-2	_ 1)	Electromagnetic compatibility (EMC) - Part 4-2: Testing and measurement techniques - Electrostatic discharge immunity test	EN 6400-4-2	1995 ²⁾
IEC 61000-4-3	_ 1)	Electromagnetic compatibility (EMC) - Part 4-3: Testing and measurement techniques - Radiated, radio-frequency, electromagnetic field immunity test	EN 61000-4-3	2006 ²⁾

¹⁾ Undated reference.

 $^{^{\}rm 2)}$ Valid edition at date of issue.

Publication	<u>Year</u>	<u>Title</u>	<u>EN/HD</u>	Year
IEC 61000-4-4	_ 1)	Electromagnetic compatibility (EMC) - Part 4-4: Testing and measurement techniques - Electrical fast transient/burst immunity test	EN 61000-4-4	2004 ²⁾
IEC 61000-4-5	_ 1)	Electromagnetic compatibility (EMC) - Part 4-5: Testing and measurement techniques - Surge immunity test	EN 61000-4-5	2006 ²⁾
IEC 61000-4-6 + A1 + A2	2003 2004 2006	Electromagnetic compatibility (EMC) - Part 4-6: Testing and measurement techniques - Immunity to conducted disturbances, induced by radio-frequency fields	EN 61000-4-6	2007
IEC 61000-4-8	-00	Electromagnetic compatibility (EMC) - Part 4-8: Testing and measurement techniques - Power frequency magnetic field immunity test	EN 61000-4-8	1993 ²⁾
IEC 61000-4-11	_ 1)	Electromagnetic compatibility (EMC) - Part 4 11: Testing and measurement techniques - Voltage dips, short interruptions and voltage variations immunity tests	EN 61000-4-11	2004 ²⁾
CISPR 11 (mod)	_ 1)	Industrial scientific and medical (ISM) radio-frequency equipment - Electromagnetic disturbance characteristics - Linus and methods of measurement	EN 55011	2007 ²⁾
CISPR 14-1	_ 1)	Electromagnetic compatibility - Requirements for household appliances, electric tools and similar apparatus - Part 1: Emission	EN 55014-1	2006 ²⁾
CISPR 15	_ 1)	Limits and methods of measurement of radio disturbance characteristics of electrica lighting and similar equipment	EN 55015 I	2006 ²⁾
CISPR 16-1-2	_ 1)	Specification for radio disturbance and immunity measuring apparatus and methods - Part 1-2: Radio disturbance and immunity measuring apparatus - Ancillary equipment - Conducted disturbances	EN 55016-1-2	2004 ²⁾
CISPR 22 (mod)	_ 1)	Information technology equipment - Radio disturbance characteristics - Limits and methods of measurement	EN 5502	2006 ²⁾

Annex ZZ

(informative)

Coverage of Essential Requirements of EC Directives

This European Standard has been prepared under a mandate given to CENELEC by the European Commission and the European Free Trade Association and within its scope the standard covers all relevant essential requirements as given in Annex I of the EC Directive 93/42/EEC.

Compliance with this standard provides one means of conformity with the specified essential requirements of the Directive concerned.

WARNING: Other equirements and other EC Directives may be applicable to the products falling within the scope of this standard.

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IEC 60601-1-2

Edition 3.0 2007-03

INTERNATIONAL STANDARD NORME INTERIONALE Medical electrical equipment – 🔗 Part 1-2: General requirements for the size safety and essential performance -Collateral standard: Electromagnetic pompatibility – Requirements and tests Appareils électromédicaux – Partie 1-2: Exigences générales pour la ségurité de base et les performances Appareils électromédicaux -essentielles – Norme collatérale: Compatibilité électromagnétique – **Exigences et essais**



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IEC 60601-1-2

Edition 3.0 2007-03

INTERNATIONAL STANDARD



0 Medical electrical equipment -Part 1-2: General requirements for basic safety and essential performance – Collateral standard: Electromagnetic Compatibility – Requirements and tests

TIONALE

Appareils électromédicaux –

Partie 1-2: Exigences générales pour la securité de base et les performances essentielles - Norme collatérale: Compatibilité électromagnétique -Exigences et essais

INTERNATIONAL ELECTROTECHNICAL COMMISSION

COMMISSION ELECTROTECHNIQUE **INTERNATIONALE**



ICS 11.040.01; 33.100.10; 33.100.20

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INTERNATIONAL ELECTROTECHNICAL COMMISSION

MEDICAL ELECTRICAL EQUIPMENT –

Part 1-2: General requirements for basic safety and essential performance -**Collateral standard:** Electromagnetic compatibility – Requirements and tests

FOREWORD

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International standard IEC 60601-1-2 has been prepared by IEC subcommittee 62A: Common aspects of electrical equipment used in medical practice, of IEC technical committee 62: Electrical equipment in medical practice.

This third edition constitutes a collateral standard to IEC 60601-1: Medical electrical equipment – Part 1: General requirements for safety and essential performance hereafter referred to as the general standard.

This document cancels and replaces the second edition of IEC 60601-1-2, and constitutes a technical revision.

This edition of IEC 60601-1-2 was revised to structurally align it with the 2005 edition of IEC 60601-1 and to implement the decision of IEC subcommittee 62A that the clause numbering structure of collateral standards written to IEC 60601-1:2005 would adhere to the form specified in ISO/IEC Directives, Part 2:2004. The principle technical changes are in Clause 4, which now recognizes that there is a general requirement for a risk management process in IEC 60601-1:2005.

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

λ	FDIS	Report on voting
Ś.	62A/560/FDIS	62A/567/RVD

Full information on the voting for the approval of this 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 the 60601 series of publications, collateral standards specify general requirements for safety applicable to:

- a subgroup of MEDICAL ELECTRICAL EQUIPMENT (e.g. radiological equipment); or
- a specific characteristic of all MEDICAL ELECTRICAL EQUIPMENT, not fully addressed in the general standard (e.g. alarm system).

In this collateral standard, the following print types are used:

- requirements and definitions: roman type.
- test specifications: italic type.
- informative material appearing outside of tables, such 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 COLLATERAL STANDARD OR AS NOTED: SMALL CAPITALS.

NOTE Defined terms are not printed in SMALL CAPITALS in Table 1 through Table 8, in the tables in Annex C and in statements required to appear in the technical description or instructions in use because they are intended for the OPERATOR OF RESPONSIBLE ORGANIZATION, who may not be familiar with the connect terms of IEC 60601 standards.

In referring to the structure of this standard, the term

- "clause" means one of the six numbered divisions within the table of contents, inclusive of all subdivisions (e.g. Clause 6 includes 6.1, 6.2, etc.);
- "subclause" means a numbered subdivision of a clause (e.g. 6.1, 62 and 6.2.1 are all subclauses of Clause 6).

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

Clauses, subclauses, items and definitions for which a rationale is provided in informative Annex A are marked with an asterisk (*).

A list of all parts of the IEC 60601 series, under the general title: Medical electrical equipment, can be found on the IEC website.

al part it can be the interance reserved to th The committee has decided that the contents of this publication will remain unchanged until the maintenance result date indicated on the IEC web site under "http://webstore.iec.ch" in the data related to the pecific publication. At this date, the publication will be

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INTRODUCTION

The need for establishing specific ELECTROMAGNETIC COMPATIBILITY standards for MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS is well recognized.

In particular, the existence of ELECTROMAGNETIC EMISSION standards is essential for the protection of:

- safety services;
- other MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS;
- electrical equipment that is not ME EQUIPMENT (e.g. computers);
- telecommunications (e.g. radio/TV, telephone, radio-navigation).

Of even more importance, the existence of ELECTROMAGNETIC IMMUNITY standards is essential to assure safety of DEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS. ELECTROMAGNETIC COMPATIBILITY (see Definition 3.4) differs from other aspects of safety covered by IEC 60601-1 because the electromagnetic phenomena exist, with varying degrees of severity, in the normal use environment of all MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS and by definition the equipment must "perform satisfactorily" within its intended environment in order to establish ELECTROMAGNETIC COMPATIBILITY. This means that the conventional single fault approach to safety is not appropriate for application to ELECTROMAGNETIC COMPATIBILITY standards. The ELECTROMAGNETIC DISTURBANCE environment can be compared to ambient temperature, humidity and atmospheric pressure. MEDICAL ELECTRICAL EQUIPMENT and MEDICAL EDECTRICAL SYSTEMS may experience environmental conditions within the expected range at any time, and for extended periods of time. As with atmospheric pressure and humidity, the OPERATOR of the MEDICAL ELECTRICAL EQUIPMENT or MEDICAL ELECTRICAL SYSTEM may not be aware of ambient levels on a continuous basis. The IMMUNITY TEST LEVELS specified in this collateral standard (IEC 60601 TEST LEVELS) represent the range found in the general medical use environment. Therefore, under these conditions, the performance of the MEDICAL ELECTRICAL EQUIPMENT or MEDICAL ELECTRICAL SYSTEM would also be expected to be normal.

MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS are used in the practice of medicine because they provide needed FUNCTIONS. IT MEDICAL ELECTRICAL EQUIPMENT or a MEDICAL ELECTRICAL SYSTEM does not provide its needed FUNCTION, because of a lack of IMMUNITY to events expected in the normal use environment, this interferes with the practice of medicine and cannot be considered an acceptable situation.

This edition recognizes that there is a shared responsibility between MANUFACTURERS, RESPONSIBLE ORGANIZATIONS and OPERATORS to ensure that MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS are designed and operated as intended. The MEDICAL ELECTRICAL EQUIPMENT or MEDICAL ELECTRICAL SYSTEM MANUFACTURER'S responsibility is to design and manufacture to meet the requirements of this collateral standard and to disclose information to the RESPONSIBLE ORGANIZATION or OPERATOR so that a compatible ELECTROMAGNETIC ENVIRONMENT can be maintained in order that the MEDICAL ELECTRICAL EQUIPMENT or MEDICAL ELECTRICAL SYSTEM will perform as intended. O

Because the practice of medicine involves many specialities, there will by necessity be MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS that are designed to perform a variety of FUNCTIONS. Some FUNCTIONS involve, for example, measurement of signals from a PATIENT that are of very low levels when compared to ELECTROMAGNETIC NOISE levels that can be coupled into MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS during the ELECTROMAGNETIC IMMUNITY testing specified in this collateral standard. Because of the proven benefits of many such MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS, this collateral standard allows the IMMUNITY TEST LEVELS to be lowered, provided there is sufficient justification based on physical, technological or physiological limitations. In this case, the MANUFACTURER is required to disclose the levels at which the MEDICAL ELECTRICAL EQUIPMENT or MEDICAL ELECTRICAL SYSTEM meets the performance requirements of this collateral standard and to specify the characteristics of the ELECTROMAGNETIC use environment and how this environment is established, in which the MEDICAL ELECTRICAL EQUIPMENT or MEDICAL ELECTRICAL SYSTEM will perform as intended.

This collateral standard also recognizes that for certain environments, higher IMMUNITY LEVELS may be required. Research necessary to determine how to identify the environments that may require higher IMMUNITY LEVELS, as well as what the levels should be, is in progress.

Finally, this collateral standard recognizes that for LIFE-SUPPORTING MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS, higher levels of IMMUNITY are necessary in order to establish a broader safety margin, even for use in the general medical use environment. Therefore, this collateral standard specifies additional requirements for LIFE-SUPPORTING MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS.

This collateral standard is based on existing IEC standards prepared by subcommittee 62A, technical committee 77 (electromagnetic compatibility between electrical equipment including networks) and CISPR (International special committee on radio interference).

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The ELECTROMAGNETIC COMPATIBILITY requirements specified by this collateral standard are generally applicable to MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS as defined in 3.63 and 3.64 in the general standard. For certain types of MEDICAL ELECTRICAL EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS, these to uirements may need to be modified by the special requirements of a particular standard. Writers of particular standards are encouraged to refer to Annex E for guidance in the application of this collateral standard.

the special requirements of a particular standard. Writers of particular standards encouraged to refer to Annex E for guidance in the application of this collateral standard.

MEDICAL ELECTRICAL EQUIPMENT -

Part 1-2: General requirements for basic safety and essential performance – Collateral standard: Electromagnetic compatibility – Requirements and tests

1 Scope, object and related standards

1.1 * Scope

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

This collateral standard apples to ELECTROMAGNETIC COMPATIBILITY of ME EQUIPMENT and ME SYSTEMS.

1.2 Object

The object of this collateral standard is to specify general requirements and tests for ELECTROMAGNETIC COMPATIBILITY of ME EQUIPMENT and ME SYSTEMS. They are in addition to the requirements of the general standard and serve as the basis for particular standards.

1.3 Related standards

1.3.1 IEC 60601-1

For ME EQUIPMENT and ME SYSTEMS, this collateral standard complements IEC 60601-1.

When referring to IEC 60601-1 or to this collateral sendard, either individually or in combination, the following conventions are used:

- "the general standard" designates IEC 60601-1 alone;
- "this collateral standard" designates IEC 60601-1-2 alone; C
- "this standard" designates the combination of the general standard and this collateral standard.

1.3.2 Particular standards

A requirement in a particular standard takes priority over the corresponding equirement in this collateral standard.

2 Normative references

The following referenced documents are indispensable for the application of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

IEC 60417, Graphical symbols for use on equipment

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

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

IEC 61000-3-2, Electromagnetic compatibility (EMC) – Part 3-2: Limits – Limits for harmonic current emissions (equipment input current \leq 16 A per phase)

IEC 61000-3-3, Electromagnetic compatibility (EMC) – Part 3-3: Limits – Limitation of voltage fluctuations and Hicker in low-voltage supply systems for equipment with rated current \leq 16 A

IEC 61000-4-2, *Electromagnetic compatibility (EMC) – Part 4-2: Testing and measurement techniques – Electrostatic discharge immunity test*

IEC 61000-4-3, Electromagnetic compatibility (EMC) – Part 4-3: Testing and measurement techniques – Radiated, regio-frequency, electromagnetic field immunity test

IEC 61000-4-4, Electromagnetic compatibility (EMC) – Part 4-4: Testing and measurement techniques – Electrical fast transient/burst immunity test

IEC 61000-4-5, Electromagnetic compatibility (EMC) – Part 4-5: Testing and measurement techniques – Surge immunity test

IEC 61000-4-6:2003, Electromagnetic compatibility (EMC) – Part 4-6: Testing and measurement techniques – Immunity to conducted disturbances, induced by radio-frequency fields ¹)

Amendment 1 (2004) Amendment 2 (2006)

IEC 61000-4-8, Electromagnetic compatibility (ENC) – Part 4-8: Testing and measurement techniques – Power frequency magnetic field immedity test

IEC 61000-4-11, Electromagnetic compatibility (ENC) – Part 4-11: Testing and measuring techniques –Voltage dips, short interruptions and voltage variations immunity tests

CISPR 11, Industrial, scientific and medical (ISM) radio-frequency equipment – Electromagnetic disturbance characteristics – Limits and methods of measurement

CISPR 14-1, Electromagnetic compatibility – Requirements for pousehold appliances, electric tools and similar apparatus – Part 1: Emission

CISPR 15, Limits and methods of measurement of radio disturbance characteristics of electrical lighting and similar equipment

CISPR 16-1-2, Specification for radio disturbance and immunity measuring apparatus and methods – Part 1-2: Radio disturbance and immunity measuring apparatus – Ancillary equipment – Conducted disturbances

CISPR 22, Information technology equipment – Radio disturbance characteristics – Limits and methods of measurement

There exists a consolidated edition 2.2 (2006) that includes IEC 61000-4-6 (2003) and its Amendment 1 (2004) and Amendment 2 (2006).