SVENSK STANDARD SS-EN 60601-2-65



Fastställd 2013-02-15

Utgåva 1 Sida 1 (1+43) Ansvarig kommitté SEK TK 62

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Elektrisk utrustning för medicinskt bruk – Säkerhet och väsentliga prestanda – Del 2-65: Särskilda fordringar på tandröntgenutrustning för intra-oral bildtagning

Medical electrical equipment – Part 2-65: Particular requirements for the basic safety and essential performance of dental intra-oral X-ray equipment

Som svensk standard gäller europastandarden EN 60601-2-65:2013. Den svenska standarden innehåller den officiella engelska språkversionen av EN 60601-2-65:2013.

Nationellt förord

Europastandarden EN 60601-2-65:2013

består av:

- europastandardens ikraftsättningsdokument, utarbetat inom CENELEC
- IEC 60601-2-65, First edition, 2012 Medical electrical equipment Part 2-65: Particular requirements for the basic safety and essential performance of dental intra-oral X-ray equipment

utarbetad inom International Electrotechnical Commission, IEC.

Standarden ska användas tillsammans med SS-EN 60601-1, utgåva 2, 2006.

Standarden ersätter delvis tidigare fastställd svensk standard SS-EN 60601-2-7, utgåva 1,1998 och SS-EN 60601-2-32, utgåva 1, 1995.

ICS 11.040.50

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EUROPEAN STANDARD

EN 60601-2-65

NORME EUROPÉENNE EUROPÄISCHE NORM

January 2013

ICS 11.040.50

Supersedes EN 60601-2-7:1998 (partially), EN 60601-2-32:1994 (partially)

English version

Medical electrical equipment Part 2-65: Particular requirements for the basic safety and essential performance of dental intra-oral X-ray equipment

(IEC 60601-2-65:2012)

Appareils électromédicaux -Partie 2-65: Exigences particulières pour la sécurité de base et les performances essentielles des appareils à rayonnement X dentaires intra-oraux (CEI 60601-2-65:2012) Medizinische elektrische Geräte -Teil 2-65: Besondere Festlegungen für die Sicherheit einschließlich der wesentlichen Leistungsmerkmale von intraoralen zahnärztlichen Röntgeneinrichtungen (IEC 60601-2-65:2012)

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Up-to-date lists and bibliographical references concerning such national standards may be obtained on application to the CEN-CENELEC Management Centre or to any CENELEC member.

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CENELEC

European Committee for Electrotechnical Standardization Comité Européen de Normalisation Electrotechnique Europäisches Komitee für Elektrotechnische Normung

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Ref. No. EN 60601-2-65:2013 E

Foreword

The text of document 62B/889/FDIS, future edition 1 of IEC 60601-2-65, prepared by IEC/SC 62B "Diagnostic imaging equipment" of IEC/TC 62 "Electrical equipment in medical practice" was submitted to the IEC-CENELEC parallel vote and approved by CENELEC as EN 60601-2-65:2013.

The following dates are fixed:

- latest date by which the document has (dop) 2013-07-24 to be implemented at national level by publication of an identical national standard or by endorsement
- latest date by which the national standards conflicting with the document have to be withdrawn

This document supersedes EN 60601-2-7:1998 (PART) and EN 60601-2-32:1994 (PART).

EN 60601-2-65:2013 includes the following significant technical changes with respect to EN 60601-2-7:1998 and EN 60601-2-32:1994:

2015-10-24

Within its specific scope, the clauses of EN 60601-2-65:2012 supersede and replace those of EN 60601-2-7:1998 and EN 60601-2-32:1994.

This standard is to be read in conjunction with EN 60601-1:2006.

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.

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

Endorsement notice

The text of the International Standard IEC 60601-2-65:2012 was approved by CENELEC as a European Standard without any modification.

In the official version, for Bibliography, the following notes have to be added for the standards indicated:

IEC 60601-2-7:1998	NOTE	Harmonised as EN 60601-2-7:1998 1) (not modified).
IEC 60601-2-28:2010	NOTE	Harmonised as EN 60601-2-28:2010 (not modified).
IEC 60601-2-32:1994	NOTE	Harmonised as EN 60601-2-32:1994 $^{\mathrm{1})}$ (not modified).
IEC 60601-2-43:2010	NOTE	Harmonised as EN 60601-2-43:2010 (not modified).
IEC 60601-2-44:2009	NOTE	Harmonised as EN 60601-2-44:2009 (not modified).
IEC 60601-2-45:2011	NOTE	Harmonised as EN 60601-2-45:2011 (not modified).
IEC 60601-2-54:2009	NOTE	Harmonised as EN 60601-2-54:2009 (not modified).
IEC 60601-2-63	NOTE	Harmonised as EN 60601-2-63.

¹⁾ Superseded by EN 60601-2-54:2009 (IEC 60601-2-54:2009, not modified).

Annex ZA

(normative)

Normative references to international publications with their corresponding European publications

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. 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 international publication has been modified by common modifications, indicated by (mod), the relevant EN/HD applies.

Annex ZA of EN 60601-1:2006 applies, except as follows:

<u>Publication</u>	<u>Year</u>	<u>Title</u>	EN/HD	<u>Year</u>			
In Annex ZA of EN 60601-1:2006 replace IEC 60601-1-2 and IEC 60601-1-3 by:							
IEC 60601-1-2 (mod)	2007	Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility - Requirements and tests	EN 60601-1-2 + corr. March	2007 2010			
IEC 60601-1-3	2008	Medical electrical equipment - Part 1-3: General requirements for basic safety and essential performance - Collatera Standard: Radiation protection in diagnostic X-ray equipment	EN 60601-1-3 + corr. March I	2008 2010			
Add to Annex ZA of EN 60601-1:2006 the following new references:							
IEC 60336	-	Medical electrical equipment - X-ray tube assemblies for medical diagnosis - Characteristics of focal spots	EN 60336	-			
IEC/TR 60788	2004	Medical electrical equipment - Glossary of defined terms	-	-			
IEC 62220-1	2003	Medical electrical equipment - Characteristics of digital X-ray imaging devices - Part 1: Determination of the detective quantum efficiency	EN 62220-1	2004			

CONTENTS

INTRO	DUCTION	5
201.1	Scope, object and related standards	6
201.2	Normative references	8
201.3	Terms and definitions	8
201.4	General requirements	10
201.5	General requirements for testing of ME EQUIPMENT	10
201.6	Classification of ME EQUIPMENT and ME SYSTEMS	10
201.7	ME EQUIPMENT identification, marking and documents	11
201.8	Protection against electrical HAZARDS from ME EQUIPMENT	13
201.9	Protection against MECHANICAL HAZARDS of ME EQUIPMENT and ME SYSTEMS	14
201.10	Protection against unwanted and excessive radiation HAZARDS	14
201.11	Protection against excessive temperatures and other HAZARDS	14
201.12	Accuracy of controls and instruments and protection against hazardous outputs	15
201.13	HAZARDOUS SITUATIONS and fault conditions	15
201.14	PROGRAMMABLE ELECTRICAL MEDICAL SYSTEMS (PEMS)	15
201.15	Construction of ME EQUIPMENT	15
201.16	ME SYSTEMS	15
201.17	Electromagnetic compatibility of ME EQUIPMENT and ME SYSTEMS	15
202	Electromagnetic compatibility – Requirements and tests	15
203	Radiation protection in diagnostic X-ray equipment	16
Annexe	s	26
	C (informative) Guide to marking and labeling requirements for ME EQUIPMENT SYSTEMS	27
Annex A	AA (informative) Particular guidance and rationale	28
	BB (informative) Identification of parts of dental X-RAY INTRA-ORAL SYSTEMS in to defined terms in this standard	35
Bibliogr	aphy	37
Index o	f defined terms used in this particular standard	40
Figure /	AA.1 – AIR KERMA during IRRADIATION with direct current X-RAY GENERATOR	29
-	AA.2 – AIR KERMA during IRRADIATION with ONE-PEAK X-RAY GENERATOR	31
	AA.3 – Waveform of long IRRADIATION TIME X-RADIATION from a AK X-RAY GENERATOR	32
Figure I	BB.1 – Structure of DENTAL INTRA-ORAL X-RAY EQUIPMENT	35
Figure I	BB.2 – Parts of DENTAL INTRA-ORAL X-RAY EQUIPMENT	36
	01.101 – List of potential ESSENTIAL PERFORMANCE to be considered by CTURER in the RISK MANAGEMENT PROCESS	10
Table 2	01.С.101 – Marking on the outside of ме еquiрмент or its parts	27
Table 2	01.C.102 – Subclauses requiring statements in ACCOMPANYING DOCUMENTS	27

INTRODUCTION

This particular standard has been prepared to provide, based on IEC 60601-1:2005 and its collaterals, a complete set of BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for DENTAL INTRA-ORAL X-RAY EQUIPMENT. While the previously existing standards for such equipment were dedicated to components and subsystems, this particular standard addresses the system level of DENTAL INTRA-ORAL X-RAY EQUIPMENT. Components and their functions are addressed as far as necessary.

The minimum safety requirements specified in this particular standard are considered to provide for a practical degree of safety in the operation of DENTAL INTRA-ORAL X-RAY EQUIPMENT.

The minimum safety requirements for DENTAL EXTRA-ORAL X-RAY EQUIPMENT are specified in a separate particular standard IEC 60601-2-63 to simplify and improve the readability

This particular standard amends and supplements IEC 60601-1 (third edition, 2005): *Medical electrical equipment – Part 1: General requirements for safety and essential performance*, hereinafter referred to as the general standard.

Within its specific scope, the clauses of this particular standard supersede and replace those of IEC 60601-2-7, *Medical electrical equipment* – Particular requirements for the safety of high-voltage generators of diagnostic X-ray generators. Requirements particular to DENTAL X-RAY-EQUIPMENT which were included in previous editions of the collateral standard IEC 60601-1-3 or the particular standard IEC 60601-2-28, IEC 60601-2-7 or IEC 60601-2-32 have been extracted and moved into this particular standard.

All requirements addressing integrated X-RAY TUBE ASSEMBLIES are covered by this particular standard. Therefore IEC 60601-2-28 does not apply to equipment in the scope of this International Standard.

MEDICAL ELECTRICAL EQUIPMENT -

Part 2-65: Particular requirements for the basic safety and essential performance of dental intra-oral X-ray equipment

201.1 Scope, object and related standards

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

201.1.1 Scope

Replacement:

This International Standard applies to the BASIC SAFETY and ESSENTIAL PERFORMANCE of DENTAL INTRA-ORAL X-RAY EQUIPMENT and its main components, hereafter also called ME EQUIPMENT.

The scope of this standard is restricted to X-RAY EQUIPMENT where the X-RAY TUBE ASSEMBLY contains the HIGH-VOLTAGE TRANSFORMER ASSEMBLY.

DENTAL EXTRA-ORAL X-RAY EQUIPMENT is excluded from the scope of this standard

NOTE 1 The X-RAY GENERATOR in DENTAL INTRA-ORAL X-RAY EQUIPMENT always comprises an X-RAY MONOBLOCK ASSEMBLY. Therefore in this particular standard the concept of X-RAY TUBE ASSEMBLY is replaced by that of X-RAY MONOBLOCK ASSEMBLY.

NOTE 2 Main components may be for instance the X-RAY MONOBLOCK ASSEMBLY and an ELECTRONIC X-RAY IMAGE RECEPTOR.

NOTE 3 Photostimulated phosphor plates and their readers (hardware and software) are excluded from the scope of this particular standard, since they have no electrical APPLIED PARTS in the PATIENT ENVIRONMENT, and are not ME EQUIPMENT.

ME EQUIPMENT and ME SYSTEMS in the scope of IEC 60601-2-63, IEC 60601-2-44, IEC 60601-2-54, IEC 60601-2-45 or IEC 60601-2-43 are excluded from the scope of this particular standard. The scope of this International Standard also excludes RADIOTHERAPY SIMULATORS and equipment for bone or tissue absorption densitometry. Excluded from the scope is also ME EQUIPMENT intended to be used for DENTAL RADIOSCOPY.

Within its specific scope, the clauses of this particular standard supersede and replace those of IEC 60601-2-7, Medical electrical equipment – Particular requirements for the safety of high-voltage generators of diagnostic X-ray generators and of IEC 60601-2-32, Medical electrical equipment – Particular requirements for the safety of associated equipment of X-ray equipment.

NOTE 4 Requirements for X-ray generators and for associated equipment, which were previously specified in IEC 60601-2-7 and IEC 60601-2-32, have been included in either IEC 60601-1:2005 (Ed3) or in this particular standard. Therefore IEC 60601-2-7 and IEC 60601-2-32 are not part of the IEC 60601-1 3rd edition scheme for Dental Intra-Oral X-ray equipment.

All requirements addressing integrated X-RAY TUBE ASSEMBLIES are covered by this particular standard. Therefore IEC 60601-2-28 does not apply to ME EQUIPMENT in the scope of this International Standard.

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

201.1.2 Object

Replacement:

The object of this particular standard is to establish particular BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for ME EQUIPMENT for DENTAL INTRA-ORAL RADIOGRAPHY.

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 and IEC 60601-1-3 apply as modified in Clauses 202 and 203 respectively. IEC 60601-1-8, IEC 60601-1- 10^{2}) and IEC 60601-1- 11^{3}) do not apply. All other published collateral standards in the IEC 60601-1 series apply as published.

NOTE OPERATORS of DENTAL INTRA-ORAL X-RAY EQUIPMENT are used to audible signals as required in this particular standard rather than to the concepts of IEC 60601-1-8. Therefore IEC 60601-1-8 does not apply.

201.1.4 Particular standards

Replacement:

In the IEC 60601 series, particular standards may modify, replace or delete requirements contained in the general standard or 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.

²⁾ IEC 60601-1-10, 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

³⁾ IEC 60601-1-11, 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

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 37.

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

IEC 60601-1-3:2008, Medical electrical equipment – Part 1-3: General requirements for basic safety and essential performance – Collateral Standard: Radiation protection in diagnostic X-ray equipment

Addition:

IEC 60336, Medical electrical equipment – X-ray tube assemblies for medical diagnosis – Characteristics of focal spots

IEC/TR 60788:2004, Medical electrical equipment – Glossary of defined terms

IEC 62220-1:2003, Medical electrical equipment – Characteristics of digital X-ray imaging devices – Part 1: Determination of the detective quantum efficiency