Elektrilised meditsiiniseadmed. Osa 2-28: Erinõuded meditsiinilises diagnoosimises kasutatavate röntgentorukoostude esmasele ohutusele ja olulistele toimimisnäitajatele

Medical electrical equipment - Part 2-28: Particular requirements for the basic safety and essential performance of X-ray tube assemblies for medical diagnosis



EESTI STANDARDI EESSÕNA

NATIONAL FOREWORD

Käesolev Eesti standard EVS-EN 60601-2-	This Estonian standard EVS-EN 60601-2-
28:2010 sisaldab Euroopa standardi EN 60601- 2-28:2010 ingliskeelset teksti.	28:2010 consists of the English text of the European standard EN 60601-2-28:2010.
	European standard EN 00001-2-20.2010.
Standard on kinnitatud Eesti Standardikeskuse	This standard is ratified with the order of
31.05.2010 käskkirjaga ja jõustub sellekohase	Estonian Centre for Standardisation dated
teate avaldamisel EVS Teatajas.	31.05.2010 and is endorsed with the notification published in the official bulletin of the Estonian
5.	national standardisation organisation.
5	
Euroopa standardimisorganisatsioonide poolt	Date of Availability of the European standard text
rahvuslikele liikmetele Euroopa standardi teksti kättesaadavaks tegemise kuopäev on	12.02.2010.
12.02.2010.	
3	
Standard on kättesaadav Eesti standardiorganisatsioonist.	The standard is available from Estonian
standardiorganisatsioonist.	Standardisation organisation.
CO CO	
ICS 11.040.55	12.02.2010. The standard is available from Estonian standardisation organisation.
	200
	Y K
	Č,
	<i>б</i> ,
	Ĩ,
	$\langle \gamma \rangle$
	-0
	U'

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.

Kui Teil on küsimusi standardite autorikaitse kohta, palun võtke ühendust Eesti Standardikeskusega: Aru 10 Tallinn 10317 Eesti; <u>www.evs.ee</u>; Telefon: 605 5050; E-post: <u>info@evs.ee</u>

Right to reproduce and distribute Estonian Standards belongs to the Estonian Centre for Standardisation

No part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying, without permission in writing from Estonian Centre for Standardisation.

If you have any questions about standards copyright, please contact Estonian Centre for Standardisation: Aru str 10 Tallinn 10317 Estonia; www.evs.ee; Phone: +372 605 5050; E-mail: info@evs.ee

EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM

EN 60601-2-28

May 2010

ICS 11.040.55

Supersedes EN 60601-2-28:1993

English version

Medical electrical equipment -Part 2-28 Particular requirements for the basic safety and essential performance of X-ray tube assemblies for medical diagnosis (IEC 60601-2-28:2010) Appareils éléctromédia Medizinische elektrische Geräte -Partie 2-28: Exigences particulières Teil 2-28: Besondere Festlegungen pour la sécurité de base für die Sicherheit einschließlich et les performances essentielles der wesentlichen Leistungsmerkmale des gaines équipées pour diagnostic von Röntgenstrahlern für die medizinische médical Diagnostik (CEI 60601-2-28:2010) (IEC 60601-2-28:2010) This European Standard was approved by CENELECon 2010-04-01. CENELEC members are bound to comply with the CEN/CENELEC Internal Regulations which stoulate the conditions for giving this European Standard the status of a national standard without any alteration. such national standards may be obtained on Up-to-date lists and bibliographical references concer application to the Central Secretariat or to any CENELEC memb This European Standard exists in three official versions (English French, German). A version in any other language made by translation under the responsibility of a CENELEC member 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, Croatia, 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 Management Centre: Avenue Marnix 17, B - 1000 Brussels

© 2010 CENELEC - All rights of exploitation in any form and by any means reserved worldwide for CENELEC members.

The text of document 62B/778/FDIS, future edition 2 of IEC 60601-2-28, prepared by SC 62B, Diagnostic imaging equipment, of IEC TC 62, Electrical equipment in medical practice, was submitted to the IEC-CENELEC parallel vote and was approved by CENELEC as EN 60601-2-28 on 2010-04-01.

This European Standard supersedes EN 60601-2-28:1993.

The second edition of this particular standard has been prepared to fit EN 60601-1:2006, which is referred to as the general standard.

When the first echion was developed, mainly X-RAY TUBE ASSEMBLIES holding a glass insert were considered and EM60601-1:1990 was in place. While the variety of modern X-RAY TUBE ASSEMBLIES and technologies has increased, the third edition of the general standard requires the MANUFACTURER to perform RISK MANAGEMENT. The technical modifications versus the first edition of EN 60601-2-28 account for these changes.

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

The following dates were 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) 2011-01-01
- latest date by which the national standard conflicting with the EN have to be withdrawn
 (dow) 2013-04-01

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.

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 congents, 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.

Annexes ZA and ZZ have been added by CENELEC.

The text of the Interprioral Standard IEC 60601-2-28:2010 was approved by CENELEC as a European standard without any pedification. **Endorsement notice**

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 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:						
Publication	Year	TIRE	<u>EN/HD</u>	Year		
Replacement:		nr.				
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	EN 60601-1-3 + corr. March	2008 2010		
Addition:						
IEC 60336	-	Medical electrical equipment - X-ray tube assemblies for medical diagnosis - Characteristics of focal spot	EN 60336	-		
IEC 60522	-	Determination of the permanent ditration of X-ray tube assemblies	EN 60522	-		
IEC 60613	2010	Electrical and loading characteristice of X-ray tube assemblies for medical diagnosis	EN 60613	2010		
IEC/TR 60788	2004	Medical electrical equipment - Glossary of defined terms	DI TILS	-		

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 requirements and other EC Directives may be applicable to the products falling within the scope of this standard.

re preme. Bard Comment is a preview generated by the second secon

CONTENTS

FOREWO	RD	3
201.1	Scope, object and related standards	5
201.2	Normative references	6
201.3	Terms and definitions	7
201.4	General requirements	7
201.5	General requirements for testing ME EQUIPMENT	7
201.6	Classification of ME EQUIPMENT and ME SYSTEMS	8
201.7	ME EQUIPMENT identification, marking and documents	8
201.8	Protection against electrical HAZARDS from ME EQUIPMENT	10
201.9	Protection a anst MECHANICAL HAZARDS of ME EQUIPMENT and ME SYSTEMS	11
201.10	Protection against unwanted and excessive radiation HAZARDS	12
201.11	Protection against excessive temperatures and other HAZARDS	12
201.12	Accuracy of controls and instruments and protection against hazardous outputs	13
201.13	HAZARDOUS SITUATIONS and fault conditions	13
201.14	PROGRAMMABLE ELECTRICA MEDICAL SYSTEMS (PEMS)	13
201.15	Construction of ME EQUIPMEN	13
201.16	ME SYSTEMS	13
201.17	Electromagnetic compatibility of MOQUIPMENT and ME SYSTEMS	13
203	RADIATION protection in diagnostic X RAY EQUIPMENT	13
Annexes		14
Annex AA	(informative) Test of X-RAY TUBE ASSEMBLIES for pressure-related RISKS	15
Index of d	lefined terms used in this particular standates for pressure-related RISKS	17

metated by FLS

MEDICAL ELECTRICAL EQUIPMENT -

Part 2-28: Particular requirements for the basic safety and essential performance of X-ray tube assemblies 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 X-RAY TUBE ASSEMBLIES and to components thereof:

- hereafter referred to as ME EQUEMENT;
- intended for medical diagnosis and imaging.

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 o ME EQUIPMENT and to ME SYSTEMS, as relevant.

NOTE This International Standard is also applicable to the X-RAY TUBE ASSEMBLY aspects of X-RAY SOURCE ASSEMBLIES and X-RAY TUBE HEADS.

201.1.2 Object

Replacement:

The object of this particular standard is to establish particular BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for X-RAY TUBE ASSEMBLIES for medical diagnosis.

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 60601-1-3 applies as modified in Clause 203. IEC 60601-1-2, IEC 60601-1-6, IEC 60601-1-8, IEC 60601-1-9, IEC 60601-1-10 and IEC 60601-1-11 do not apply. All other published collateral standards in the IEC 60601-1 series apply as published.

NOTE 101 IEC 60601-1-2 does not apply because RISKS for the X-RAY TUBE ASSEMBLY outside the system may only be indicative of RISKS for the system due to the difference in electromagnetic environment.

NOTE 102 IEC 60601-1-6 and IEC 60601-1-8 do not apply because X-RAY TUBE ASSEMBLIES are not operated as a stand-alone device.

NOTE 103 X-RAY TUBE ASSEMBLIES are not in the scope of IEC 60601-1-10 and IEC 60601-1-11.

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