
Medicinska električna oprema - 2-37. del: Posebne varnostne zahteve za ultrazvočno medicinsko diagnostično in nadzorno opremo

Medical electrical equipment - Part 2-37: Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment

Medizinische elektrische Geräte - Teil 2-37: Besondere Festlegungen für die Sicherheit einschließlich der wesentlichen Leistungsmerkmale von Ultraschallgeräten für die medizinische Diagnose und Überwachung

Appareils électromédicaux - Partie 2-37: Exigences particulières pour la sécurité de base et les performances essentielles des appareils de diagnostic et de surveillance médicaux à ultrasons

Ta slovenski standard je istoveten z: EN 60601-2-37:2008/A11:2011

ICS:

11.040.55	Diagnostična oprema	Diagnostic equipment
17.140.50	Elektroakustika	Electroacoustics

SIST EN 60601-2-37:2008/A11:2012 **en,fr,de**

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EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN 60601-2-37/A11

October 2011

ICS 11.040.55; 17.140.50

English version

**Medical electrical equipment -
Part 2-37: Particular requirements for the basic safety and essential
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CENELEC

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

Management Centre: Avenue Marnix 17, B - 1000 Brussels

Foreword

This document (EN 60601-2-37:2008/A11:2011) has been prepared by CLC/TC 62 “Electrical equipment in medical practice”.

The following dates are fixed:

- latest date by which this document has (dop) 2012-10-01
to be implemented at national level by
publication of an identical national
standard or by endorsement
- latest date by which the national (dow) 2014-10-01
standards conflicting with this
document have to be withdrawn

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Replace Annex ZZ of EN 60601-2-37:2008 by:

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 except as follows:

- Essential Requirement 6a
- Essential Requirement 7.4
- Essential Requirement 7.5 paragraph 2 & 3
- Essential Requirement 13.6 (q)

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.

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