

STN	Zdravotnícke elektrické prístroje Časť 2-3: Osobitné požiadavky na základnú bezpečnosť a nevyhnutné prevádzkové vlastnosti krátkovlnných terapeutických prístrojov Zmena A2	STN EN 60601-2-3/A2 36 4800
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Medical electrical equipment - Part 2-3: Particular requirements for the basic safety and essential performance of short-wave therapy equipment

Táto norma obsahuje anglickú verziu európskej normy.
This standard includes the English version of the European Standard.

Táto norma bola oznámená vo Vestníku ÚNMS SR č. 11/24

STN EN 60601-2-3 z decembra 2015 sa bez tejto zmeny A2 môže používať do 31. 7. 2027.

Obsahuje: EN 60601-2-3:2015/A2:2024, IEC 60601-2-3:2012/AMD2:2022

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Úrad pre normalizáciu, metrológiu a skúšobníctvo Slovenskej republiky, 2024
Slovenská technická norma a technická normalizačná informácia je chránená zákonom č. 60/2018 Z. z. o technickej normalizácii v znení neskorších predpisov.

EUROPEAN STANDARD
NORME EUROPÉENNE
EUROPÄISCHE NORM

EN 60601-2-3:2015/A2

September 2024

ICS 11.040.60

English Version

**Medical electrical equipment - Part 2-3: Particular requirements
for the basic safety and essential performance of short-wave
therapy equipment
(IEC 60601-2-3:2012/AMD2:2022)**

Appareils électromédicaux - Partie 2-3: Exigences
particulières pour la sécurité de base et les performances
essentielles des appareils de thérapie à ondes courtes
(IEC 60601-2-3:2012/AMD2:2022)

Medizinische elektrische Geräte - Teil 2-3: Besondere
Festlegungen für die Sicherheit einschließlich der
wesentlichen Leistungsmerkmale von Kurzwellen-
Therapiegeräten
(IEC 60601-2-3:2012/AMD2:2022)

This amendment A2 modifies the European Standard EN 60601-2-3:2015; it was approved by CENELEC on 2024-07-31. CENELEC members are bound to comply with the CEN/CENELEC Internal Regulations which stipulate the conditions for giving this amendment 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 CEN-CENELEC Management Centre or to any CENELEC member.

This amendment 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 CEN-CENELEC Management Centre has the same status as the official versions.

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

CEN-CENELEC Management Centre: Rue de la Science 23, B-1040 Brussels

EN 60601-2-3:2015/A2:2024 (E)**European foreword**

The text of document 62D/1846/CDV, future edition 3 of IEC 60601-2-3/AMD2, prepared by SC 62D "Particular medical equipment, software, and systems" of IEC/TC 62 "Medical equipment, software, and systems" was submitted to the IEC-CENELEC parallel vote and approved by CENELEC as EN 60601-2-3:2015/A2:2024.

The following dates are fixed:

- latest date by which the document has to be implemented at national (dop) 2025-05-01 level by publication of an identical national standard or by endorsement
- latest date by which the national standards conflicting with the (dow) 2027-07-31 document have to be withdrawn

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Any feedback and questions on this document should be directed to the users' national committee. A complete listing of these bodies can be found on the CENELEC website.

Endorsement notice

The text of the International Standard IEC 60601-2-3:2012/AMD2:2022 was approved by CENELEC as a European Standard without any modification.

Annex ZA (normative)

Normative references to international publications with their corresponding European publications

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements 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 1 Where an International Publication has been modified by common modifications, indicated by (mod), the relevant EN/HD applies.

NOTE 2 Up-to-date information on the latest versions of the European Standards listed in this annex is available here: www.cencenelec.eu.

Add:

<u>Publication</u>	<u>Year</u>	<u>Title</u>	<u>EN/HD</u>	<u>Year</u>
IEC 60601-1	2005	Medical electrical equipment - Part 1: General requirements for basic safety and essential performance	EN 60601-1	2006
-	-		+ AC	2010
+ A1	2012		+ A1	2013
-	-		+ AC	2014
-	-		+ A12	2014
+ A2	2020		+ A2	2021
-	-		+ AC	2022
-	-		+ A13	2024
IEC/TR 60878	2015	Graphical symbols for electrical equipment in medical practice	-	-

koniec náhľadu – text ďalej pokračuje v platenej verzii STN