



SLOVENSKI STANDARD

oSIST prEN IEC 60601-2-44:2026

01-januar-2026

Medicinska električna oprema - 2-44. del: Posebne zahteve za osnovno varnost in bistvene lastnosti rentgenske opreme za računalniško tomografijo

Medical electrical equipment - Part 2-44: Particular requirements for the basic safety and essential performance of X-ray equipment for computed tomography

Appareils électromédicaux - Partie 2-44: Exigences particulières pour la sécurité de base et les performances essentielles des équipements à rayonnement X de tomomodensitométrie

Ta slovenski standard je istoveten z: prEN IEC 60601-2-44:2025

[oSIST prEN IEC 60601-2-44:2026](https://standards.iteh.ai/catalog/standards/sist/ae2e34b5-c8ee-4627-835d-618dc9a53a9c/osist-pr-en-iec-60601-2-44-2026)

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ICS:

11.040.50	Radiografska oprema	Radiographic equipment
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oSIST prEN IEC 60601-2-44:2026	en,fr,de
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62B/1400/CDV

COMMITTEE DRAFT FOR VOTE (CDV)

PROJECT NUMBER:

IEC 60601-2-44 ED4

DATE OF CIRCULATION:

2025-11-21

CLOSING DATE FOR VOTING:

2026-02-13

SUPERSEDES DOCUMENTS:

62B/1376/CD, 62B/1384A/CC

IEC SC 62B : MEDICAL IMAGING EQUIPMENT, SOFTWARE, AND SYSTEMS

SECRETARIAT:

Germany

SECRETARY:

Ms Regina Geierhofer

OF INTEREST TO THE FOLLOWING COMMITTEES:

SC 62A, SC 62C

HORIZONTAL FUNCTION(S):

ASPECTS CONCERNED:

Electromagnetic Compatibility, Safety

☒ SUBMITTED FOR CENELEC PARALLEL VOTING☐ NOT SUBMITTED FOR CENELEC PARALLEL VOTING**Attention IEC-CENELEC parallel voting**

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TITLE:

Medical electrical equipment - Part 2-44: Particular requirements for the basic safety and essential performance of X-ray equipment for computed tomography

PROPOSED STABILITY DATE: 2029

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