

INTERNATIONAL STANDARD

NORME INTERNATIONALE

AMENDMENT 1
AMENDEMENT 1

Medical electrical equipment –
Part 2-45: Particular requirements for the basic safety and essential performance
of mammographic X-ray equipment and mammographic stereotactic devices

Appareils électromédicaux –
Partie 2-45: Exigences particulières pour la sécurité de base et les performances
essentiels des appareils de mammographie à rayonnement X et des appareils
mammographiques stéréotaxiques





THIS PUBLICATION IS COPYRIGHT PROTECTED

Copyright © 2015 IEC, Geneva, Switzerland

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either IEC or IEC's member National Committee in the country of the requester. If you have any questions about IEC copyright or have an enquiry about obtaining additional rights to this publication, please contact the address below or your local IEC member National Committee for further information.

Droits de reproduction réservés. Sauf indication contraire, aucune partie de cette publication ne peut être reproduite ni utilisée sous quelque forme que ce soit et par aucun procédé, électronique ou mécanique, y compris la photocopie et les microfilms, sans l'accord écrit de l'IEC ou du Comité national de l'IEC du pays du demandeur. Si vous avez des questions sur le copyright de l'IEC ou si vous désirez obtenir des droits supplémentaires sur cette publication, utilisez les coordonnées ci-après ou contactez le Comité national de l'IEC de votre pays de résidence.

IEC Central Office
3, rue de Varembe
CH-1211 Geneva 20
Switzerland

Tel.: +41 22 919 02 11
Fax: +41 22 919 03 00
info@iec.ch
www.iec.ch

About the IEC

The International Electrotechnical Commission (IEC) is the leading global organization that prepares and publishes International Standards for all electrical, electronic and related technologies.

About IEC publications

The technical content of IEC publications is kept under constant review by the IEC. Please make sure that you have the latest edition, a corrigenda or an amendment might have been published.

IEC Catalogue - webstore.iec.ch/catalogue

The stand-alone application for consulting the entire bibliographical information on IEC International Standards, Technical Specifications, Technical Reports and other documents. Available for PC, Mac OS, Android Tablets and iPad.

IEC publications search - www.iec.ch/searchpub

The advanced search enables to find IEC publications by a variety of criteria (reference number, text, technical committee,...). It also gives information on projects, replaced and withdrawn publications.

IEC Just Published - webstore.iec.ch/justpublished

Stay up to date on all new IEC publications. Just Published details all new publications released. Available online and also once a month by email.

Electropedia - www.electropedia.org

The world's leading online dictionary of electronic and electrical terms containing more than 30 000 terms and definitions in English and French, with equivalent terms in 15 additional languages. Also known as the International Electrotechnical Vocabulary (IEV) online.

IEC Glossary - std.iec.ch/glossary

More than 60 000 electrotechnical terminology entries in English and French extracted from the Terms and Definitions clause of IEC publications issued since 2002. Some entries have been collected from earlier publications of IEC TC 37, 77, 86 and CISPR.

IEC Customer Service Centre - webstore.iec.ch/csc

If you wish to give us your feedback on this publication or need further assistance, please contact the Customer Service Centre: csc@iec.ch.

A propos de l'IEC

La Commission Electrotechnique Internationale (IEC) est la première organisation mondiale qui élabore et publie des Normes internationales pour tout ce qui a trait à l'électricité, à l'électronique et aux technologies apparentées.

A propos des publications IEC

Le contenu technique des publications IEC est constamment revu. Veuillez vous assurer que vous possédez l'édition la plus récente, un corrigendum ou amendement peut avoir été publié.

Catalogue IEC - webstore.iec.ch/catalogue

Application autonome pour consulter tous les renseignements bibliographiques sur les Normes internationales, Spécifications techniques, Rapports techniques et autres documents de l'IEC. Disponible pour PC, Mac OS, tablettes Android et iPad.

Recherche de publications IEC - www.iec.ch/searchpub

La recherche avancée permet de trouver des publications IEC en utilisant différents critères (numéro de référence, texte, comité d'études,...). Elle donne aussi des informations sur les projets et les publications remplacées ou retirées.

IEC Just Published - webstore.iec.ch/justpublished

Restez informé sur les nouvelles publications IEC. Just Published détaille les nouvelles publications parues. Disponible en ligne et aussi une fois par mois par email.

Electropedia - www.electropedia.org

Le premier dictionnaire en ligne de termes électroniques et électriques. Il contient plus de 30 000 termes et définitions en anglais et en français, ainsi que les termes équivalents dans 15 langues additionnelles. Egalement appelé Vocabulaire Electrotechnique International (IEV) en ligne.

Glossaire IEC - std.iec.ch/glossary

Plus de 60 000 entrées terminologiques électrotechniques, en anglais et en français, extraites des articles Termes et Définitions des publications IEC parues depuis 2002. Plus certaines entrées antérieures extraites des publications des CE 37, 77, 86 et CISPR de l'IEC.

Service Clients - webstore.iec.ch/csc

Si vous désirez nous donner des commentaires sur cette publication ou si vous avez des questions contactez-nous: csc@iec.ch.

INTERNATIONAL STANDARD

NORME INTERNATIONALE

AMENDMENT 1
AMENDEMENT 1

Medical electrical equipment –
Part 2-45: Particular requirements for the basic safety and essential performance
of mammographic X-ray equipment and mammographic stereotactic devices

Appareils électromédicaux –
Partie 2-45: Exigences particulières pour la sécurité de base et les performances
essentielle des appareils de mammographie à rayonnement X et des appareils
mammographiques stéréotaxiques

INTERNATIONAL
ELECTROTECHNICAL
COMMISSION

COMMISSION
ELECTROTECHNIQUE
INTERNATIONALE

ICS 11.040.50

ISBN 978-2-8322-2785-5

Warning! Make sure that you obtained this publication from an authorized distributor.
Attention! Veuillez vous assurer que vous avez obtenu cette publication via un distributeur agréé.

FOREWORD

This amendment has been prepared by subcommittee 62B: Diagnostic imaging equipment, of IEC technical committee 62: Electrical equipment in medical practice.

This bilingual version (2015-07) corresponds to the English version, published in 2015-06.

The text of this amendment is based on the following documents:

CDV	Report on voting
62B/917/CDV	62B/954/RVC

Full information on the voting for the approval of this amendment can be found in the report on voting indicated in the above table.

The French version of this amendment has not been voted upon.

The committee has decided that the contents of this amendment and the base publication will remain unchanged until the stability date indicated on the IEC web site under "http://webstore.iec.ch" in the data related to the specific publication. At this date, the publication will be

- reconfirmed,
- withdrawn,
- replaced by a revised edition, or
- amended.

iTeh STANDARD PREVIEW
(standards.iteh.ai)

[IEC 60601-2-45:2011/AMD1:2015](https://standards.iteh.ai/catalog/standards/sist/697ae06c-f1e1-4ef6-ad00-0217a27da2e8/iec-60601-2-45-2011-amd1-2015)

[https://standards.iteh.ai/catalog/standards/sist/697ae06c-f1e1-4ef6-ad00-](https://standards.iteh.ai/catalog/standards/sist/697ae06c-f1e1-4ef6-ad00-0217a27da2e8/iec-60601-2-45-2011-amd1-2015)

NOTE The attention of National Committees is drawn to the fact that equipment MANUFACTURERS and testing organizations may need a transitional period following publication of a new, amended or revised IEC publication in which to make products in accordance with the new requirements and to equip themselves for conducting new or revised tests. It is the recommendation of the committee that the content of this publication be adopted for implementation nationally not earlier than 3 years from the date of publication.

INTRODUCTION TO THE AMENDMENT

This first amendment to the third edition of this particular standard has been prepared to provide a complete set of safety requirements for MAMMOGRAPHIC X-RAY EQUIPMENT, based on IEC 60601-1:2005 and IEC 60601-1:2005/AMD1:2012, IEC 60601-1-2:2014, and IEC 60601-1-3:2008 and IEC60601-1-3:2008/AMD1:2013. This particular standard addresses the system level of MAMMOGRAPHIC X-RAY EQUIPMENT and introduces equipment for MAMMOGRAPHIC TOMOSYNTHESIS.

FOREWORD

Replace, in the second paragraph, the reference to "IEC 60601-1-3 (2010)" by "IEC 60601-1-3 (2008)".

201.1 Scope, object and related standards

Replace, in footnote 1, the reference to "IEC 60601-1:2005" by "IEC 60601-1:2005 and IEC 60601-1:2005/AMD1:2012".

201.1.1 Scope

Add, in the first paragraph, after the term "MAMMOGRAPHIC X-RAY EQUIPMENT" the phrase "including equipment for MAMMOGRAPHIC TOMOSYNTHESIS,".

Replace, in the first dashed item of the third paragraph, the words "modes of operation" by "other than MAMMOGRAPHIC TOMOSYNTHESIS"

Add, after this first dashed item, the following new dashed item:

- CT SCANNERS covered by IEC 60601-2-44;

201.1.3 Collateral standards

Replace, in the first sentence of the second paragraph, the reference to "IEC 60601-1-2:2007" by "IEC 60601-1-2:2014"

Replace the reference to "IEC 60601-1-3:2008" by "IEC 60601-1-3:2008 and IEC 60601-1-3:2008/AMD1:2013"

Replace the second sentence of the second paragraph, including its footnote, with the following new sentence and footnote:

IEC 60601-1-8, IEC 60601-1-9, IEC 60601-1-10, IEC 60601-1-11 and IEC 60601-1-12 do not apply¹.

¹ IEC 60601-1-9:2007, *Medical electrical equipment – Part 1-9: General requirements for basic safety and essential performance – Collateral Standard: Requirements for environmentally conscious design*. IEC 60601-1-10:2007, *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*. IEC 60601-1-11:2010, *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*. IEC 60601-1-12:2004, *Medical Electrical Equipment – Part 1-12: General requirements for basic safety and essential performance – Collateral Standard: Requirements for medical electrical equipment and medical electrical systems used in the emergency medical services environment*.

201.2 Normative references

Replace the existing references to IEC 60601-1-2:2007 and IEC 60601-1-3:2008 by the following:

IEC 60601-1-2:2014, *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*

IEC 60601-1-3:2008/AMD1:2013

Delete the following normative reference:

IEC 61223-3-2:2007, *Evaluation and routine testing in medical imaging departments – Part 3-2: Acceptance tests – Imaging performance of mammographic X-ray equipment*

201.3 Terms and definitions

Replace, in the first paragraph, the reference “IEC 60601-1:2005, IEC 60601-1-3:2008” by “IEC 60601-1:2005 and IEC 60601-1:2005/AMD1:2012, IEC 60601-1-3:2008 and IEC 60601-1-3:2008/AMD1:2013”.

201.3.205

DIRECT FOCAL DISTANCE

<https://standards.iteh.ai/catalog/standards/sist/697ae06c-f1e1-4ef6-ad00-0317a27de2c8/iec-60601-2-45-2011-amd1-2015>

Replace the existing text of the definition by the following:

<X-ray mammography> shortest distance from the FOCAL SPOT to the axis of symmetry of the EFFECTIVE IMAGE RECEPTION AREA perpendicular to its chest wall edge for a specified position of the source

201.3.206

MAMMOGRAPHIC STEREOTACTIC DEVICE

Add an asterisk * in front of the term to read

*MAMMOGRAPHIC STEREOTACTIC DEVICE

Replace the existing text of the definition by the following:

device for mechanically guided placement of a needle or position marker based on radiographic images of an immobilized breast acquired at different known angles

Renumber the existing note as Note 1 and add the following new note:

NOTE 2 The purposes of such devices may be fine-needle aspiration, core biopsy, or pre-surgical localization.

Add the following new definitions:

201.3.210

MAMMOGRAPHIC TOMOSYNTHESIS

technique using MAMMOGRAPHIC X-RAY EQUIPMENT to produce multiple tomographic images reconstructed from multiple PROJECTIONS acquired over a total angular range of less than 180°

201.3.211

CONTRAST TO NOISE RATIO

CNR

physical quantity describing the ability to distinguish between various contrast objects of a digital image and the inherent noise within the image, defined as the difference of mean pixel values of the contrast objects and image background, and divided by the standard deviation of the image background pixel value

[SOURCE: IEC 61223-3-2:2007, definition 3.8]

201.4 General requirements

201.4.3 ESSENTIAL PERFORMANCE

201.4.3.101 *Additional ESSENTIAL PERFORMANCE requirements

Table 201.101 – Distributed ESSENTIAL PERFORMANCE requirements

Replace the subclause reference for "Accuracy of LOADING FACTORS" in the first row, by "203.6.4.3.102" <https://standards.iteh.ai/catalog/standards/sist/697ae06c-fl-e1-4ef6-ad00-0317a27de2c8/iec-60601-2-45-2011-amd1-2015>

201.4.101 Data recording

Add, after the final dashed item in the first paragraph, the following new dashed item and note:

- identification and version of image processing applied to ORIGINAL DATA and, in MAMMOGRAPHIC TOMOSYNTHESIS, identification and version of reconstruction processing applied.

NOTE An example for processed images are DICOM images for presentation.

201.7 ME EQUIPMENT identification, marking and documents

201.7.9 ACCOMPANYING DOCUMENTS

201.7.9.2 Instructions for use

Add the following new subclause:

201.7.9.2.17 *ME EQUIPMENT emitting radiation

This subclause of IEC 60601-1:2005 and IEC 60601-1:2005/AMD1:2012 does not apply.

201.7.9.3.101 Specification of X-RAY SOURCE ASSEMBLY and its position

Add, in item e), after the term "DIRECT FOCAL DISTANCE", the following new text:

and, in MAMMOGRAPHIC TOMOSYNTHESIS, for specified positions of the source

Add, in item f), after the term "DIRECT FOCAL DISTANCE" the following new text:

and, in MAMMOGRAPHIC TOMOSYNTHESIS, for specified positions of the source

Add, in item g) after the term "IMAGE RECEPTION AREA" the following new text:

and, in MAMMOGRAPHIC TOMOSYNTHESIS, for specified positions of the source

Add the following new items:

- h) in MAMMOGRAPHIC TOMOSYNTHESIS, the number of PROJECTIONS, and the geometric configuration for the acquisition of the PROJECTIONS;
- i) in MAMMOGRAPHIC TOMOSYNTHESIS, description of the distribution of x-ray LOADING FACTORS for the acquisition of the PROJECTIONS.

201.9 Protection against MECHANICAL HAZARDS of ME EQUIPMENT and ME SYSTEMS

201.9.2 HAZARDS associated with moving parts

Replace the existing title with the following:
IEC 60601-2-45:2011/AMD1:2015
<https://standards.iteh.ai/catalog/standards/sist/697ae06c-f1e1-4ef6-ad00-0317a27de2c8/iec-60601-2-45-2011-amd1-2015>

201.9.2 MECHANICAL HAZARDS associated with moving parts

201.9.2.1 General

Delete, in the second paragraph, the phrase "detect PATIENT contact and".

201.9.2.3 Other HAZARDS associated with moving parts

Replace the existing title with the following:

201.9.2.3 Other MECHANICAL HAZARDS associated with moving parts

201.9.2.101 * MAMMOGRAPHIC STEREOTACTIC DEVICE

Replace the existing title by the following:

201.9.2.101 * Three dimensional localization and interventional mammographic guidance

201.9.2.101.1 Positioning of X-RAY SOURCE ASSEMBLY for stereotactic imaging

Add after the first paragraph, before the compliance statement, the following new paragraph:

This subclause does not apply for MAMMOGRAPHIC TOMOSYNTHESIS.

201.9.2.101.3 Biopsy needle positioning accuracy of MAMMOGRAPHIC STEREOTACTIC DEVICES

Replace the existing title by the following:

201.9.2.101.3 Biopsy needle positioning accuracy

Delete, in the first paragraph, the word "stereotactic".

a) Test equipment

Delete, in the first sentence of the first paragraph and in the second and third sentences of the second paragraph, the word "stereotactic".

Delete, at the end of the second paragraph, the phrase "with the MAMMOGRAPHIC STEREOTACTIC DEVICE".

b) Test procedure

Replace, in the third sentence of the first paragraph the phrase "of the MAMMOGRAPHIC STEREOTACTIC DEVICE" by the word "SPECIFIED". Delete the two other instances of the word "stereotactic" in this sentence.

Delete in the second paragraph the phrase "with which the MAMMOGRAPHIC STEREOTACTIC DEVICE is".

iTeh STANDARD PREVIEW

Replace the second and third sentences of the third paragraph by the following new text: "Determine x, y, z positions of the test needle tips as specified by the MANUFACTURER for clinical use."

<http://standards.iteh.ai>

Delete, in the fourth sentence of the third paragraph, the phrase "by the MAMMOGRAPHIC STEREOTACTIC DEVICE".

201.10 Protection against unwanted and excessive radiation HAZARDS**201.10.1.2 ME EQUIPMENT intended to produce diagnostic or therapeutic X-radiation**

Replace, in the first paragraph, the reference to "IEC 60601-1-3:2008" by "IEC 60601-1-3:2008 and IEC 60601-1-3:2008/AMD1:2013".

201.12 Accuracy of controls and instruments and protection against hazardous outputs

Replace the existing text of this clause by the following:

Clause 12 of the general standard applies.

201.13 Hazardous situations and fault conditions

Replace the existing title of this clause by the following:

201.13 Hazardous situations and fault conditions for ME EQUIPMENT

202 Electromagnetic compatibility – Requirements and tests

Replace, in the first line, the reference to "IEC 60601-1-2:2007" by "IEC 60601-1-2:2014".

203 Radiation protection in diagnostic X-ray equipment

Replace, in the first line, the reference to "IEC 60601-1-3:2008" by "IEC 60601-1-3:2008 and IEC 60601-1-3:2008/AMD1:2013".

203.4 General requirements

203.4.1 Statement of compliance

Replace the reference to "IEC 60601-2-45:2011" by "IEC 60601-2-45:2015".

203.4.101.2 * LOADING TIME

Replace in Note 1 the reference to "IEC 60601-1-3:2008" by "IEC 60601-1-3:2008 and IEC 60601-1-3:2008/AMD1:2013".

<https://standards.iteh.ai/catalog/standards/sist/697ae06c-f1e1-4ef6-ad00-0317a27de2c8/iec-60601-2-45-2011-amd1-2015>

203.6.3 RADIATION dose and RADIATION QUALITY

203.6.3.1 Adjustment of RADIATION dose and RADIATION QUALITY

203.6.3.1.1 General requirements for the adjustment of RADIATION dose and RADIATION QUALITY

Replace, in the second paragraph, the reference to "IEC 60601-1-3:2008" by "IEC 60601-1-3:2008 and IEC 60601-1-3:2008/AMD1:2013".

203.6.3.1.2 Linearity of AIR KERMA over limited intervals of LOADING FACTORS

Add, at the end of the first dashed item, the following sentence:

For MAMMOGRAPHIC TOMOSYNTHESIS this lower value shall be the lowest CURRENT TIME PRODUCT setting available in a LOADING of a TOMOSYNTHESIS projection image acquisition series.

203.6.3.2 Reproducibility of the X-RADIATION output

Add, after the fourth dashed item, the following new dashed item:

- for MAMMOGRAPHIC TOMOSYNTHESIS a combination of X-RAY TUBE VOLTAGE specified by the MANUFACTURER and clinically justified, with the lowest CURRENT TIME PRODUCT setting available in a complete LOADING of a TOMOSYNTHESIS acquisition series

203.6.4.2.101 LOADING STATE in mammography

Add, after the third paragraph, before the compliance statement, the following new paragraph:

For MAMMOGRAPHIC TOMOSYNTHESIS operation the LOADING STATE shall encompass all acquired projections.

203.6.4.3 Indication of LOADING FACTORS and MODES of OPERATION

Add, at the beginning of the subclause, after the instruction "Addition:", the following new text:

For MAMMOGRAPHIC TOMOSYNTHESIS acquisition, which involves multiple IRRADIATIONS, LOADING FACTORS shall be provided after completion of the acquisition for each of these IRRADIATIONS.

NOTE An example of implementation of this requirement is through usage of DICOM objects.

203.6.4.3.102 Accuracy of LOADING FACTORS**203.6.4.3.102.1 General**

Replace, in the compliance statement, the reference to "203.6.4.3.104" by "203.6.4.3.103".

203.6.4.3.102.3 Accuracy of X-RAY TUBE CURRENT

Replace the existing text of this subclause by the following:

For operation of MAMMOGRAPHIC X-RAY EQUIPMENT where the X-RAY TUBE CURRENT can be independently selected, its value shall be accurate within $\pm 20\%$ of the INDICATED VALUE within the selectable range.

203.6.4.3.102.4 Accuracy of LOADING TIME

Replace the existing first paragraph by the following:

For operation of MAMMOGRAPHIC X-RAY EQUIPMENT where the LOADING TIME can be independently selected, the error on the value of the LOADING TIME shall not be greater than $\pm (10\% + 1 \text{ ms})$ for combinations of LOADING FACTORS representing the selectable range.

203.6.4.3.102.5 Accuracy of CURRENT TIME PRODUCT

Add at the end of this subclause the following new paragraph:

For MAMMOGRAPHIC TOMOSYNTHESIS operation the CURRENT TIME PRODUCT shall be the sum of the respective CURRENT TIME PRODUCTS of the individual projections.

203.6.4.3.103 Test conditions for the accuracy of loading factors**203.6.4.3.103.1 Accuracy and reproducibility of X-RAY TUBE VOLTAGE**

Replace the existing first paragraph by the following:

Measurements shall be made at 30 kV or at the X-RAY TUBE VOLTAGE specified by the MANUFACTURER if clinically justified. Measurements shall also be made at the lowest and highest selectable values of the X-RAY TUBE VOLTAGE and at the lowest, medium and highest selectable values of CURRENT TIME PRODUCT.

Replace, at the beginning of the third paragraph, the phrase "Calculate the average" by "Check each measured".

203.6.4.5 Dosimetric indications

Add, at the end of this subclause, the following new paragraph:

For MAMMOGRAPHIC TOMOSYNTHESIS the AVERAGE GLANDULAR DOSE to be indicated shall be the cumulative AVERAGE GLANDULAR DOSE over the entire tomographic acquisition.

203.6.5.3 AEC requirements for MAMMOGRAPHIC X-RAY EQUIPMENT provided with an integrated digital X-RAY IMAGE RECEPTOR

203.6.5.3.1 General requirements

Add, after the existing first paragraph, the following new paragraph:

For MAMMOGRAPHIC TOMOSYNTHESIS equipment this requirement may be assessed in projection images.

Replace, in the second sentence of the existing second paragraph, the phrase "magnification and (if applicable)" by "magnification, (if applicable) tomosynthesis and".

Replace, in the existing third paragraph the phrase "(e.g. magnification and stereotactic modes)" by "(e.g. magnification, tomosynthesis and stereotactic modes)".

Add the following new text at the end of the note:

and is included in 203.6.5.3.3

203.6.5.3.2 AEC reproducibility

a) Test method

Add, after the dashed list of item a), the following new paragraph:

For MAMMOGRAPHIC TOMOSYNTHESIS equipment the PIXEL values in a region of interest shall be assessed in projection images of the PHANTOM.

Add the following new subclause:

203.6.5.3.3 AEC thickness response

The response of the AEC to different breast thicknesses shall be evaluated by jointly assessing image quality, as measured by the CONTRAST TO NOISE RATIO, and the dose to the PATIENT, as characterized by the AVERAGE GLANDULAR DOSE, and comparing them to the specifications provided.

a) Test method

Measure the CONTRAST TO NOISE RATIO of radiograms and AVERAGE GLANDULAR DOSE of PHANTOMS made of breast tissue equivalent material, produced with the AUTOMATIC EXPOSURE CONTROL in operation. Determine these quantities, in all operational modes, for PHANTOM thicknesses from 20 mm to 70 mm in steps of 10 mm.