

INTERNATIONAL STANDARD

NORME INTERNATIONALE

Good refurbishment practices for medical imaging equipment

Bonnes pratiques de reconditionnement pour les appareils d'imagerie médicale





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CONTENTS

FOREWORD.....	3
INTRODUCTION.....	5
1 Scope.....	6
2 Normative references	6
3 Terms and definitions	7
4 General requirements for REFURBISHMENT of USED MEDICAL IMAGING EQUIPMENT	9
4.1 Quality management system	9
4.2 Resource management	9
4.3 Corrective and preventive action.....	9
4.4 Customer complaints	9
4.5 Production and service provision	9
4.6 Control of nonconforming PRODUCT	10
4.7 Post-market surveillance PROCESS	10
4.8 Document control.....	10
4.9 Purchasing.....	10
4.10 Control of design and design changes	10
4.11 RISK management PROCESS	10
5 Specific requirements for good REFURBISHMENT practice	11
5.1 General.....	11
5.2 Selection of MEDICAL IMAGING EQUIPMENT for REFURBISHMENT	11
5.3 Evaluating market access requirements	11
5.4 Preparation for REFURBISHMENT, disassembly, packing, and transport	11
5.5 Planning	11
5.6 Installation of software and hardware to ensure the safety of the MEDICAL IMAGING EQUIPMENT.....	12
5.7 Performance and safety test	12
5.8 Packing, transport, and installation of refurbished MEDICAL IMAGING EQUIPMENT.....	12
5.9 Record of REFURBISHMENT	12
5.10 REFURBISHMENT label	12
Annex A (informative) Cross reference list of the contents of IEC 63077 versus ISO 13485	13
Bibliography.....	15
Index of defined terms used in this document	16
Table A.1 – Cross reference list of the contents of IEC 63077 versus ISO 13485.....	13

INTERNATIONAL ELECTROTECHNICAL COMMISSION

**GOOD REFURBISHMENT PRACTICES
FOR MEDICAL IMAGING EQUIPMENT**

FOREWORD

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International Standard IEC 63077 has been prepared by subcommittee 62B: Diagnostic imaging equipment, of IEC technical committee 62: Electrical equipment in medical practice.

This first edition cancels and replaces the second edition of IEC PAS 63077 published in 2016. This edition constitutes a technical revision.

This edition includes the following significant technical changes with respect to IEC PAS 63077:2016:

- a) the scope was delineated more clearly;
- b) an informative cross reference list of IEC 63077 vs ISO 13485 (Annex A) was added;
- c) smaller corrections were performed.

The text of this International Standard is based on the following documents:

FDIS	Report on voting
62B/1149/FDIS	62B/1155/RVD

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

This document has been drafted in accordance with the ISO/IEC Directives, Part 2.

In this document, the following print types are used:

- requirements and definitions: roman 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: SMALL CAPITALS.

The committee has decided that the contents of this document will remain unchanged until the stability date indicated on the IEC website under "<http://webstore.iec.ch>" in the data related to the specific document. At this date, the document will be

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

INTRODUCTION

This document specifies requirements for a quality management system that can be used by organizations involved in REFURBISHMENT of MEDICAL IMAGING EQUIPMENT.

The requirements defined in this document can be used by MANUFACTURERS or organizations providing REFURBISHMENT. Organizations providing REFURBISHMENT can voluntarily choose to conform to the requirements of this document or can be required by contract with the MANUFACTURER of the MEDICAL IMAGING EQUIPMENT to conform.

Several jurisdictions have regulatory requirements regarding refurbished MEDICAL IMAGING EQUIPMENT e.g. regarding the import and making refurbished MEDICAL IMAGING EQUIPMENT available. These regulatory requirements differ from nation to nation and region to region. The organizations involved in REFURBISHMENT of MEDICAL IMAGING EQUIPMENT should understand how the regulatory requirements in the several jurisdictions will be interpreted and may be met by applying this document.

In some jurisdictions a definition of the term remanufacturer is available. This document does not cover the topic of how organizations are acting in the role of a remanufacturer.

This document can also be used by internal and external parties, including certification bodies, to assess the organization's ability to meet requirements applicable for the REFURBISHMENT of MEDICAL IMAGING EQUIPMENT.

It is emphasized that the requirements specified in this document are complementary to other International Standards such as on quality management system and on RISK management.

There is a wide variety of medical equipment with different requirements on REFURBISHMENT. Therefore, this document only applies to named groups of MEDICAL IMAGING EQUIPMENT. These groups are defined in Clause 1 Scope.

GOOD REFURBISHMENT PRACTICES FOR MEDICAL IMAGING EQUIPMENT

1 Scope

This document describes and defines the PROCESS of REFURBISHMENT of USED MEDICAL IMAGING EQUIPMENT and applies to the restoring of USED MEDICAL IMAGING EQUIPMENT to a condition of safety and performance comparable to that of new MEDICAL IMAGING EQUIPMENT i.e. MEDICAL IMAGING EQUIPMENT that was not in use. This restoration includes actions such as REPAIR, REWORK, software/hardware updates, and the replacement of worn parts with original parts. This document enumerates the actions, that are performed, and the manner consistent, with relevant specifications and service procedures required to ensure that the REFURBISHMENT of MEDICAL IMAGING EQUIPMENT is done without changing the finished MEDICAL IMAGING EQUIPMENT's performance, safety specifications, or INTENDED USE according to its original or applicable valid registration.

The MEDICAL IMAGING EQUIPMENT and systems covered by this document include:

- X-RAY EQUIPMENT;
- X-RAY EQUIPMENT for RADIOSCOPICALLY GUIDED INTERVENTIONAL PROCEDURES;
- X-RAY EQUIPMENT FOR COMPUTED TOMOGRAPHY;
- MAGNETIC RESONANCE EQUIPMENT;
- ULTRASONIC DIAGNOSTIC EQUIPMENT;
- GAMMA CAMERAS;
- PLANAR WHOLEBODY IMAGING EQUIPMENT;
- equipment for SINGLE PHOTON EMISSION COMPUTED TOMOGRAPHY (SPECT);
- SPECT/CT hybrid systems, combining a GAMMA CAMERA with X-RAY EQUIPMENT FOR COMPUTED TOMOGRAPHY (CT);
- POSITRON EMISSION TOMOGRAPHS (PET);
- PET/CT hybrid systems combining a POSITRON EMISSION TOMOGRAPH with X-RAY EQUIPMENT FOR COMPUTED TOMOGRAPHY (CT);
- PET/MRI hybrid systems combining a POSITRON EMISSION TOMOGRAPH with MAGNETIC RESONANCE EQUIPMENT; and
- other combinations of the MEDICAL IMAGING EQUIPMENT or systems listed above.

This document does not apply to endoscopic equipment, funduscopy equipment, radiation therapy equipment, nor associated systems.

2 Normative references

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.

ISO 13485:2016, *Medical devices – Quality management systems – Requirements for regulatory purposes*

ISO 14971:2007, *Medical devices – Application of risk management to medical devices*

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- IEC Electropedia: available at <http://www.electropedia.org/>
- ISO Online browsing platform: available at <http://www.iso.org/obp>

3.1

EXPECTED SERVICE LIFE

time period specified by the MANUFACTURER during which the medical electrical equipment or medical electrical system is expected to remain safe for use (i.e. maintain basic safety and essential performance)

Note 1 to entry: Maintenance can be necessary during the EXPECTED SERVICE LIFE.

[SOURCE: IEC 60601-1:2005/AMD1:2012, 3.28]

3.2

INTENDED USE

INTENDED PURPOSE

use for which a PRODUCT, PROCESS, or service is intended according to the specifications, instructions and information provided by the MANUFACTURER

Note 1 to entry: INTENDED USE should not be confused with NORMAL USE. While both include the concept of use as intended by the MANUFACTURER, INTENDED USE focuses on the medical purpose while NORMAL USE incorporates not only the medical purpose, but maintenance, transport, etc. as well.

[SOURCE: IEC 60601-1:2005/AMD1:2012, 3.44]

3.3

MANUFACTURER

natural or legal person with responsibility for the design, manufacture, packaging, labelling, assembling, or adapting MEDICAL IMAGING EQUIPMENT, regardless of whether these operations are performed by that person or on that person's behalf by a third party

Note 1 to entry: Adapting includes making substantial modifications to MEDICAL IMAGING EQUIPMENT already in use.

[SOURCE: IEC 60601-1:2005/AMD1:2012, 3.55, modified – The term MEDICAL IMAGING EQUIPMENT is replacing ME EQUIPMENT or ME SYSTEM in the definition and in the Note to entry, and three Notes to entry have been deleted.]

3.4

MEDICAL IMAGING EQUIPMENT

medical electrical equipment that provides images for clinical applications

Note 1 to entry: See IEC 60601-1:2005, 3.63 for a definition of MEDICAL ELECTRICAL EQUIPMENT.

3.5

NORMAL USE

operation, including routine inspection and adjustments by any OPERATOR, and stand-by, according to the instructions for use

Note 1 to entry: NORMAL USE should not be confused with INTENDED USE. While both include the concept of use as intended by the MANUFACTURER, INTENDED USE focuses on the medical purpose while NORMAL USE incorporates not only the medical purpose, but maintenance, transport, etc. as well.

[SOURCE: IEC 60601-1:2005/AMD1:2012, 3.71]