

Edition 6.1 2023-02 CONSOLIDATED VERSION

INTERNATIONAL STANDARD

NORME INTERNATIONALE



Medical electrical equipment -

Part 2-2: Particular requirements for the basic safety and essential performance of high frequency surgical equipment and high frequency surgical accessories

Appareils électromédicaux -

Partie 2-2: Exigences particulières pour la sécurité de base et les performances essentielles des appareils d'électrochirurgie à courant haute fréquence et des accessoires d'électrochirurgie à courant haute fréquence





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CONTENTS

FOREWORD	4
INTRODUCTION	7
INTRODUCTION to Amendment 1	7
201.1 Scope, object and related standards	8
201.2 Normative references	9
201.3 Terms and definitions	10
201.4 General requirements	14
201.5 General requirements for testing of ME EQUIPMENT	15
201.6 Classification of ME EQUIPMENT and ME SYSTEMS	16
201.7 ME EQUIPMENT identification, marking and documents	16
201.8 Protection against electrical HAZARDS from ME EQUIPMENT	21
201.9 Protection against MECHANICAL HAZARDS of ME EQUIPMENT and ME SYSTEMS	37
201.10 Protection against unwanted and excessive radiation HAZARDS	37
201.11 Protection against excessive temperatures and other HAZARDS	38
201.12 Accuracy of controls and instruments and protection against hazardous outputs	39
201.13 HAZARDOUS SITUATIONS and fault conditions for ME EQUIPMENT	45
201.14 PROGRAMMABLE ELECTRICAL MEDICAL SYSTEMS (PEMS)	46
201.15 Construction of ME EQUIPMENT	46
201.16 ME SYSTEMS	51
201.17 Electromagnetic compatibility of ME EQUIPMENT and ME SYSTEMS	51
202 * ELECTROMAGNETIC DISTURBANCES – Requirements and tests	51
208 General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	52
Annexes	53
Annex AA (informative) Particular guidance and rationale	54
Annex BB (informative) ELECTROMAGNETIC DISTURBANCES created by HF SURGICAL EQUIPMENT	80
Bibliography	89
Index of defined terms used in this particular standard	91
Figure 201.101 – Symbol used with an EARTH REFERENCED PATIENT CIRCUIT	
Figure 201.102 - Symbol used with a HF ISOLATED PATIENT CIRCUIT	16
Figure 201.103 – Circuit suitable for testing compliance to 201.8.4.101	23
Figure 201.104 – Measurement of HF LEAKAGE CURRENT for EARTH REFERENCED PATIENT CIRCUITS and load between electrodes	26
Figure 201.105 – Measurement of HF LEAKAGE CURRENT for EARTH REFERENCED PATIENT CIRCUITS and a load resistance from ACTIVE ELECTRODE to earth	27
Figure 201.106 – Measurement of HF LEAKAGE CURRENT for HF ISOLATED PATIENT CIRCUITS	28
Figure 201.107 – Measurement of HF LEAKAGE CURRENT from a BIPOLAR ACCESSORY	29
Figure 201.108 – Test apparatus for anchorages of cords of ACTIVE ACCESSORY	36
Figure 201.109 – Measurement of output power – MONOPOLAR output	41
Figure 201.110 - Measurement of output power - BIPOLAR output	42

Figure 201.111 – Method of testing feedback from one active output to another in simultaneous activation	45
Figure AA.1 – Examples of various parts of an HF surgical ME SYSTEM	56
Figure AA.2 – Example of MONOPOLAR method of HF surgery using a NEUTRAL ELECTRODE	56
Figure AA.3 – Example of BIPOLAR method of HF surgery	57
Figure AA.4 – Crest factor vs. peak voltage	62
Figure AA.5 – Example of PATIENT circuit with NEUTRAL ELECTRODE referenced to earth at operating frequencies	66
Figure BB.1 – E-FIELD EMISSIONS test setup	83
Figure BB.2 – H-FIELD EMISSIONS test setup	84
Figure BB.3 – Conducted EMISSIONS test setup	85
Figure BB.4 – Unit ad hoc test	87
Figure BB.5 – Power cord ad hoc test	88
Figure BB.6 – Accessory cord ad hoc test	88
Table 201.101 – Colours of indicator lights and their meaning for HF SURGICAL EQUIPMENT	17
Table 201.102 – Maximum output powers in SINGLE FAULT CONDITIONS	44
Table 201.103 – Test currents by weight range	48
Table AA.1 – Summary of measured current and durations for 25 TUR procedures	75
Table AA.2 – Summary of measured currents and durations for general surgical procedures	76
Table BB.1 – Worst case EMISSIONS of spark gap type HF SURGICAL EQUIPMENT	86
Table BB.2 – Worst case EMISSIONS of non-spark gap (modern) HF SURGICAL EQUIPMENT	86

INTERNATIONAL ELECTROTECHNICAL COMMISSION

MEDICAL ELECTRICAL EQUIPMENT -

Part 2-2: Particular requirements for the basic safety and essential performance of high frequency surgical equipment and high frequency surgical accessories

FOREWORD

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This consolidated version of the official IEC Standard and its amendment has been prepared for user convenience.

IEC 60601-2-2 edition 6.1 contains the sixth edition (2017-03) [documents 62D/1427/FDIS and 62D/1442/RVD] and its amendment 1 (2023-02) [documents 62D/2010/FDIS and 62D/2021/RVD].

In this Redline version, a vertical line in the margin shows where the technical content is modified by amendment 1. Additions are in green text, deletions are in strikethrough red text. A separate Final version with all changes accepted is available in this publication.

IEC 60601-2-2:2017+AMD1:2023 CSV - 5 - © IEC 2023

International standard IEC 60601-2-2 has been prepared by IEC subcommittee 62D: Electromedical equipment, of IEC technical committee 62: Electrical equipment in medical practice.

This sixth edition constitutes a technical revision. This edition includes the following significant technical changes with respect to the previous edition:

- refinement and additions to the defined terms;
- additional separation of the requirements for HF surgical equipment and HF surgical accessories;
- a new requirement for adult neutral electrodes to be contact quality monitoring neutral electrodes;
- new requirements for devices that have or use a high current mode.

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

In this standard, the following print types are used:

- requirements and definitions: roman type;
- test specifications: italic 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 OF THE GENERAL STANDARD, IN THIS PARTICULAR STANDARD OR AS NOTED: SMALL CAPITALS.

In referring to the structure of this standard, the term

- "clause" means one of the seventeen numbered divisions within the table of contents, inclusive of all subdivisions (e.g. Clause 7 includes subclauses 7.1, 7.2, etc.);
- "subclause" means a numbered subdivision of a clause (e.g. 7.1, 7.2 and 7.2.1 are all subclauses of Clause 7).

References to clauses within this standard are preceded by the term "Clause" followed by the clause number. References to subclauses within this standard are by number only.

In this standard, the conjunctive "or" is used as an "inclusive or" so a statement is true if any combination of the conditions is true.

The verbal forms used in this standard conform to usage described in Clause 7 of the ISO/IEC Directives, Part 2. For the purposes of this standard, the auxiliary verb:

- "shall" means that compliance with a requirement or a test is mandatory for compliance with this standard;
- "should" means that compliance with a requirement or a test is recommended but is not mandatory for compliance with this standard;
- "may" is used to describe a permissible way to achieve compliance with a requirement or test.

An asterisk (*) as the first character of a title or at the beginning of a paragraph or table title indicates that there is guidance or rationale related to that item in Annex AA.

A list of all parts of the IEC 60601 series, published under the general title *Medical electrical equipment*, can be found on the IEC website.

The committee has decided that the contents of the base publication and its amendment will remain unchanged until the stability date indicated on the IEC web site under 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.

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INTRODUCTION

The minimum safety requirements specified in this particular standard are considered to provide for a practical degree of safety in the operation of HIGH FREQUENCY SURGICAL EQUIPMENT.

This particular standard amends and supplements IEC 60601-1:2005 and Amendment 1:2012, Medical electrical equipment – Part 1: General requirements for basic safety and essential performance, hereinafter referred to as the general standard (see 201.1.4).

The requirements are followed by specifications for the relevant tests.

A "Particular guidance and rationale" section giving some explanatory notes, where appropriate, about the more important requirements is included in Annex AA.

Clauses or subclauses for which there are explanatory notes in Annex AA are marked with an asterisk (*).

It is considered that a knowledge of the reasons for these requirements will not only facilitate the proper application of the standard but will, in due course, expedite any revision necessitated by changes in clinical practice or as a result of developments in technology. However, this annex does not form part of the requirements of this document.

INTRODUCTION to Amendment 1

The 6th Edition of IEC 60601-2-2 was published in 2017. This amendment is intended to align the standard to IEC 60601-1:2005/AMD2:2020. Additionally, this amendment is intended to address several issues reported from the national committees, including but not limited to:

- requirement for including the length of an accessory in the instructions for use;
- clarification of test setup for HF LEAKAGE CURRENTS;
- considering modes with high DUTY CYCLES above 45 % in the risk management;
- including text of the interpretation sheet 62D/1703/INF regarding the HIGH CURRENT MODE to Annex AA.

MEDICAL ELECTRICAL EQUIPMENT -

Part 2-2: Particular requirements for the basic safety and essential performance of high frequency surgical equipment and high frequency surgical accessories

201.1 Scope, object and related standards

Clause 1 of the general standard¹ applies, except as follows:

201.1.1 * Scope

Replacement:

This part of IEC 60601 applies to the BASIC SAFETY and ESSENTIAL PERFORMANCE of HF SURGICAL EQUIPMENT and HF SURGICAL ACCESSORIES as defined in 201.3.224 and 201.3.223.

HF SURGICAL EQUIPMENT having a RATED OUTPUT POWER not exceeding 50 W (for example for micro-COAGULATION, or for use in dentistry or ophthalmology) is exempt from certain of the requirements of this particular standard. These exemptions are indicated in the relevant requirements.

201.1.2 Object

Replacement:

The object of this particular standard is to establish particular BASIC SAFETY and ESSENTIAL PERFORMANCE requirements for HF SURGICAL EQUIPMENT and HF SURGICAL ACCESSORIES as defined in 201.3.224 and 201.3.223.

201.1.3 Collateral standards

Addition:

This particular standard refers to those applicable collateral standards that are listed in Clause 2 of the general standard and Clause 201.2 of this particular standard.

IEC 60601-1-2:2014 and IEC 60601-1-2:2014/AMD1:2020 and IEC 60601-1-8:2006, IEC 60601-1-8:2006/AMD1:2012 and IEC 60601-1-8:2006/AMD2:2020, apply as modified in Clauses 202 and 208 respectively. IEC 60601-1-3, IEC 60601-1-10, IEC 60601-1-11 and IEC 60601-1-12 do not apply. All other published collateral standards in the IEC 60601-1 series apply as published.

201.1.4 Particular standards

Replacement:

In the IEC 60601 series, particular standards may modify, replace or delete requirements contained in the general standard and collateral standards as appropriate for the particular

The general standard is IEC 60601-1:2005, IEC 60601-1:2005/AMD1:2012 and IEC 60601-1:2005/AMD2:2020, Medical electrical equipment – Part 1: General requirements for basic safety and essential performance.