### SLOVENSKI PREDSTANDARD

### oSIST prEN 60601-2-22:2005

december 2005

Medicinska električna oprema – 2-22. del: Posebne zahteve za varnost in bistvene lastnosti kirurške, terapevtske in diagnostične laserske opreme

Medical electrical equipment - Part 2-22: Particular requirements for the safety and essential performance of surgical, therapeutic and diagnostic laser equipment

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Secretary: Gerald Glen Secrétaire:					
Also of interest to the following committe Intéresse également les comités suivant		Supersedes document Remplace le document 76/287/CD, 76/300/CC			
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Titre : tps://standards.iteh.ai/catalog/stand		Title : IEC 60601-2-22 Ed.3: MEDICAL ELECTRICAL EQUIPMENT – Part 2: Particular requirements for the safety and essential performance of surgical, therapeutic and diagnostic laser equipment			
Note d'introduction		Introductory note IEC TC76/WG4 foreword to the CDV only. This draft incorporates all changes due to the comments received prior to the WG4 Valbella meeting in October 2004 and observed at the meeting. The document was 76/287/60601-2- 22/comments. This document was distributed by IEC as 76/300/CC. The old numbering of clauses and Definitions were retained in order to facilitate reviewing. The Numbering will be brought in an order editorially in the FDIS stage.			
		March 2005, The convener The French version will be circulated later.			
ATTENTION		March 2005, The convener			
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CDV soumis en parallèle au		March 2005, The convener The French version will be circulated later. ATTENTION			

IEC.

### INTERNATIONAL ELECTROTECHNICAL COMMISSION

### MEDICAL ELECTRICAL EQUIPMENT

# Part 2: Particular requirements for the safety and essential performance of surgical, therapeutic and diagnostic laser equipment

### FOREWORD

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International Standard IEC 60601-2-22 has been prepared by technical committee No. 76: Optical radiation safety and laser equipment of IEC.

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

FDIS	Report on voting		
XX/XX/FDIS	XX/XX/RVD		

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

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

The committee has decided that the contents of this publication will remain unchanged until the maintenance result date<sup>1)</sup> 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
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<sup>1)</sup> 

The National Committees are requested to note that for this publication the maintenance result date is ....

In this Particular Standard, the following print types are used:

- requirements and definitions: in Arial type;
- NOTES: in smaller Arial type;

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- compliance: in italic type;

- TERMS USED THROUGHOUT THIS PARTICULAR STANDARD WHICH HAVE BEEN DEFINED IN CLAUSE 3 AS WELL AS THOSE DEFINED IN IEC 60601-1 AND IEC 60825-1: SMALL CAPITALS

### INTRODUCTION

This Particular Standard amends and supplements IEC 60601-1 (third edition, xxxx, see Note): *Medical Electrical Equipment – Part 1: General requirements for safety and essential performance.* 

This Standard also refers to IEC 60825-1 (2001).

The requirements of this Standard have to be taken as the minimum to comply with, in order to achieve a reasonable level of safety and reliability during operation and application of medical laser equipment.

Clauses or subclauses for which there are explanatory notes in annex AA: Rationale 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 result of developments in technology. However, this annex does not form part of the requirements of this Standard.

WG4 note for IEC: This document has been prepared by IEC/TC76 Working Group 4 using the Committee Drafts of IEC 60601-1 3<sup>rd</sup> edtion, 62A/389/CDV (Rockville) and 62A/449/CDV (Valbella). SISTEN 60601-2-22:2013

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TC76/WG4 has produced a table listing the clause and subclause numbering of the second and third editions (CD) which follows below and is included in this Part-2 standard. The table should only facilitate navigating and is not meant to be finally published.

When the third edition of 60601-1 is issued as an FDIS, this standard will be reviewed for possible changes.

Old clause number	New clause number	Old clause number	New clause number	Old clause number	New clause number
Appendix L	2	19.3	8.7.3	52.4.101	13.1.101
2.x.x	3.1xxx	32	10.4/10.6	52.5.9	13.2.101
3	4	36	17	55	16.4
3.6	13.2.101	45	9.7	56.11	8.10.4.101

60601-2-22/CD\	(© IEC	_ {	5 –		
6	7	49	11.8	56.101	10.4.g)
6.1	7.2	50.2	12.1	57.10	8.9
6.3	7.4	51.2	12.4.4	57.101	8.9.101
6.8	7.9	51.5	12.4.6	59.101	15.101
6.8.2	7.9.2	51.101	12.4.101		

This note together with the table are for review purposes only. They will be omitted from the final publication.

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### MEDICAL ELECTRICAL EQUIPMENT

# Part 2: Particular requirements for the safety and essential performance of surgical, therapeutic and diagnostic laser equipment

### SECTION ONE – GENERAL

The clauses and subclauses of this section of the General Standard apply except as follows:

### 1 Scope, object and related standards

This clause of the General Standard applies except as follows:

1.1 Scope

Addition:

This Particular International Standard applies to laser equipment for either surgical, therapeutic, medical diagnostic, cosmetic, or veterinary APPLICATIONS, intended for its use on humans or animals, classified as a CLASS 3B or CLASS 4 LASER PRODUCT according to 3.21 and 3.22 in IEC 60825-1, hereinafter referred to as laser equipment.

Throughout this document light emitting diodes (LED) are included whenever the word "laser" is used.

NOTE – Laser equipment classified as a CLASS 1, 1M, 2, 2M or CLASS 3R LASER PRODUCT, is covered by IEC 60825-1 and IEC 60601-1.

#### <u>IST EN 60601-2-22:2013</u>

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The object of this Particular Standard is to specify particular requirements for the safety of laser equipment classified as a CLASS 3B or CLASS 4 LASER PRODUCT.

NOTE - Laser classification (IEC 60825-1) is not to be confused with electrical classification (IEC 60601-1)

#### 1.3 Particular Standards

Addition:

This Particular Standard refers to IEC 60601-1, 3rd edition (xxxx).

For brevity, Part 1 of IEC 60601-1 is referred to in this Particular Standard either as the "General Standard" or as the "General Requirement(s)".

The numbering of clauses and subclauses of this Particular Standard corresponds to that of the General Standard. The changes to the text of the General Standard are specified by the use of the following words:

"Replacement" means that the clause or subclause of the General Standard is replaced completely by the text of this Particular Standard.

"Addition" means that the text of this Particular Standard is additional to the requirements of the General Standard.

"Amendment" means that the clause or subclause of the General Standard is amended as indicated by the text of this Particular Standard.

Subclauses or figures which are additional to those of the General Standard are numbered starting from 101, additional annexes are lettered AA, BB, etc., and additional items *aa*), *bb*), etc.

The term "this Standard" is used to make reference to the General Standard and this Particular Standard taken together.

Where there is no corresponding clause or subclause in this Particular Standard, the clause or subclause of the General Standard, although possibly not relevant, applies without modification. Where it is intended that any part of the General Standard, although possibly relevant, is not to be applied, a statement to that effect is given in this Particular Standard.

Concerning LASER RADIATION safety of laser equipment, IEC 60825-1 applies, except that the relevant requirements are specified, changed or amended in this Particular Standard.

Clauses and subclauses of the General Standard and IEC 60825-1, which are not applicable to laser equipment for medical applications, are not necessarily indicated as "not applicable".

#### 2. Normative references

Addition:

IEC 60601-1: xxxx, Medical electrical equipment – Part 1: General requirements for safety and essential performance

IEC 60664-1: 1992, Insulation coordination for equipment within low-voltage systems – Part 1: Principles, requirements and tests

IEC 60664-3: 1992, Insulation coordination for equipment within low-voltage systems – Part 3: Use of coatings to achieve insulation coordination of printed board assemblies

IEC 60825-1, Ed. 1.2: 2001-08, Safety of laser products – Part 1: Equipment classification, requirements and user's guide

IEC 60947-3: 1990, Low-voltage switchgear and controlgear – Part 3: Switches, disconnectors, switch-disconnectors and fuse-combination units

#### 3. Terminology and definitions

This clause of the General Standard applies except as follows:

Additional definitions:

3.1001 ACCESSIBLE EMISSION LIMIT (AEL)

ACCESSIBLE EMISSION LIMIT for Class 1 (1M, 2, 2M, 3R, 3B) lasers. See 3.2 and tables 1 through 4 of IEC 60825-1.

3.1002 AIMING BEAM

Beam of optical radiation, producing a visible AIMING BEAM spot, intended for indication of the anticipated point of impact of the WORKING BEAM.

3.1003 AIMING BEAM POSITION

Area of impact of the AIMING BEAM within the WORKING AREA.

3.1004 AIMING LASER

LASER emitting an AIMING BEAM.

3.1005 APERTURE

See 3.8 of IEC 60825-1.

3.1006 BEAM DELIVERY SYSTEM

Optical system which guides the LASER RADIATION from its origin to a LASER APERTURE.

3.1007 CLASS 1 (1M, 2, 2M, 3R, 3B, 4) LASER PRODUCT

Laser equipment, incorporating a LASER as defined in 3.17 through 3.22 and 3.40 of IEC 60825-1.

### 3.1008 EMERGENCY LASER STOP STAND and siteh.ai)

Hand- or foot-actuated device intended to stop the LASER OUTPUT immediately in case of emergency.

3.1009 LASER EMISSION INDICATOR SIST EN 60601-2-22:2013

ttps://standards.iteh.ai/catalog/standards/sist/c7858878-5ac5-4dcf-89a6-1c479336170d/sist-en-60601-2-22-2013 Visible and/or audible signal which indicates that the WORKING BEAM is being emitted through any APERTURE.

3.1010 LASER ENERGY

RADIANT ENERGY of the WORKING BEAM, incident on the WORKING AREA, see 3.68 of IEC 60825-1.

3.1012 LASER OUTPUT

Either LASER POWER or LASER ENERGY.

3.1013 LASER POWER

RADIANT POWER of the WORKING BEAM, incident on the WORKING AREA, see 3.70 of IEC 60825-1.

3.1014 LASER READY INDICATOR

Visible means which indicates that the laser equipment is in the READY condition.