





IEC 60601-1 Medical electrical equipment

Part 1: General requirements for basic safety and essential performance

 Report Reference No......
 T223-0554/19

 Date of issue......
 2019-09-23

 Total number of pages......
 231 pages

CB Testing Laboratory.....: SIQ Ljubljana

SIQ Ljubljana is accredited by Slovenian Accreditation with accreditation number

LP-009 in the field of testing

Address Tržaška cesta 2, SI-1000 Ljubljana, Slovenia

Applicant's name...... GlobTek, Inc.

Test specification:

Standard IEC 60601-1:2005 (Third Edition) + CORR. 1:2006 + CORR. 2:2007 +

A1:2012

(or IEC 60601-1: 2012 reprint)

Test procedure: CB Scheme

Non-standard test method.....: N/A

Test Report Form No.....: IEC60601 1K

 Test Report Form Originator.....:
 UL(US)

 Master TRF......
 2015-11

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Trade Mark.....:

Manufacturer:

Model/Type reference ...:

Ratings:

Power supply for medical use

GlobTek, Inc.

GlobTek, Inc.

186 Veterans Drive, Northvale, NJ-07647, New Jersey, USA

GT-500160-30

Input: 100-240 V~; 50-60 Hz; 1,6-0,7 A; Class I

Output: 30 Vdc; 2 A



Testing procedure and testing location:	
	SIQ Ljubljana
Testing location/ address:	Tržaška cesta 2, SI-1000 Ljubljana, Slovenia
Associated CB Testing Laboratory:	
Testing location/ address:	
Tested by (name, function, signature):	Gregor Cesar Osov Gregor
Approved by (name, function, signature):	Janez Vidmar Jan Vid
Testing procedure: CTF Stage 1:	
Testing location/ address:	
Tested by (name, function, signature):	
Approved by (name, function, signature):	The second section of the second seco
☐ Testing procedure: CTF Stage 2:	
Testing location/ address:	
Tested by (name, function, signature):	0.000
Witnessed by (name, function, signature) .:	
Approved by (name, function, signature):	
☐ Testing procedure: CTF Stage 3:	
☐ Testing procedure: CTF Stage 4:	
Testing location/ address:	
Tested by (name, function, signature):	Luca made de la constante de l
Witnessed by (name, function, signature).:	
Approved by (name, function, signature):	
Supervised by (name, function, signature):	



List of Attachments (including a total number of pages in each attachment):

- 1. Test Report: 151 pages
- 2. National Differences Enclosure No. 1: 25 pages
- 3. European Differences Enclosure No. 1a: 37 pages
- 4. Photo Documentation Enclosure No. 2: 6 pages
- 5. Schematics, layouts, user manual and transformer drawings Enclosure No. 3: 12 pages

Summary of testing

Tests performed (name of test and test clause):

4.11 Power Input

7.1.2 Legibility of marking

7.1.3 Durability of marking

8.5.4 Working voltage Measurement

8.6.4. Impedance and current- carrying capability of protective earth connections

8.7.4.5 Earth leakage current

8.7.4.6 Touch Current

8.8.3 Dielectric Strength test of solid insulation materials with safety functions

8.8.4.1 Resistance to heat - Ball pressure test of thermoplastic parts

11.1 Excessive temperatures in ME EQUIPMENT

13.2 Single Fault conditions

15.3.2 Push test

15.3.3 Impact test

15.3.4.2 Drop test - portable ME Equipment

15.3.6 Mould-stress relief test

15.5.1.2 Transformer short circuit

15.5.1.3 Transformer overload

Evaluation of voltage limiting components in SELV circuits

Testing location:

SIQ Ljubljana

Tržaška cesta 2

SI-1000 Ljubljana, Slovenia

Revision 1.0:

SIQ Ljubljana

Mašera-Spasićeva ulica 10, SI-1000 Ljubljana, Slovenia



Summary of compliance with National Differences

List of countries addressed:

• IEC 60601-1: 2005 + A1: 2012

No national differences for IEC 60601-1: 2005 + A1: 2012 declared.

• IEC 60601-1: 2005

List of countries addressed:

- US NATIONAL DIFFERENCES to IEC 60601-1 Third edition National standard ANSI/AAMI ES60601-1: 2005
- CANADA NATIONAL DIFFERENCES to IEC 60601-1 Third edition National standard CAN/CSA-C22.2 No. 60601-1:14
- SWITZERLAND NATIONAL DIFFERENCES to IEC 60601-1 Third edition National standard SN EN 60601-1:06
- JAPAN NATIONAL DIFFERENCES to IEC 60601-1 Third edition National standard: JIS T0601-1:2012

☐ The product fulfils the requirements of EN 60601-1:2006 + A1:2013 + A12:2014.



Copy of marking plate

The artwork below may be only a draft. The use of certification marks on a product must be authorized by the respective NCBs that own these marks.



POWER SUPPLY FOR MEDICAL USE

MODEL: GT-500160-30

INPUT: 100-240V ~ 50-60Hz 1.6-0.7A

OUTPUT: 30VDC === 2A







RoHS O € (



S/N:

RoHS000158101/07

MADE IN CHINA



GENERAL INFORMATION	
Test item particulars (see also Clause 6):	
Classification of installation and use:	Portable equipment (desk-top version)
Device type (component/sub-assembly/ equipment/ system):	Component level power supply for desktop use
Intended use (Including type of patient, application location):	EUT is intended to provide power to medical devices with isolation grade 2xMOOP
Mode of operation:	Continuous operation
Supply connection	Appliance coupler
Accessories and detachable parts included:	No accessories and detachable parts included
Other options include:	No other options included
Testing	
Date of receipt of test item(s):	2012-06-28
	Revision 1.0:
	2019-07-23
Dates tests performed:	From 2012-07-02 to 2012-09-03
	Revision 1.0:
	From 2019-07-24 to 2019-07-26
Possible test case verdicts:	
- test case does not apply to the test object:	N/A
- test object does meet the requirement:	Pass (P)
- test object was not evaluated for the requirement:	N/E (collateral standards only)
- test object does not meet the requirement:	Fail (F)
Abbreviations used in the report:	
- normal condition: N.C.	- single fault condition: S.F.C.
- means of Operator protection: MOOP	- means of Patient protection: MOPP
General remarks:	
Before starting to use the TRF please read carefully the 4 in on how to complete the new version "K" of TRF for IEC for "(See Attachment #)" refers to additional information appended "(See appended table)" refers to a table appended to the report The tests results presented in this report relate only to the object This report shall not be reproduced except in full without the write test equipment must be kept on file and available for reviated the data and/or information provided in the attachment. Throughout this report a comma / point is used as the	to the report. t. tt tested. itten approval of the testing laboratory. iew. ents to this report.



Manufacturer's Declaration per sub-clause 4.2.5 of I	ECEE 02:2012
The application for obtaining a CB Test Certificate includes more than one factory location and a declaration from the Manufacturer stating that the sample(s) submitted for evaluation is (are) representative of the products from each factory has been provided	✓ Yes☐ Not applicable
When differences exist; they shall be identified in th	e General product information section.
Name and address of factory (ies):	1) GlobTek Inc.
	186 Veterans Drive,
	Northvale, NJ-07647, New Jersey, USA
	2) GlobTek (Suzhou) Co., Ltd.
	Building 4, No. 76, Jinling East Road,
	Suzhou Industrial Park, Jiangsu CN-215021, China



General product information:

Equipment under test (Model: GT-500160-30) is switch mode power supply, provided with an appliance inlet (IEC 60320) for input power connection and non-detachable output cables with low-voltage-plug.

Units are desk-top type power supplies. Enclosure consists of two parts which are fixed together by 2 screws (cannot be open without the use of a tool). There are no openings in enclosure.

Units are designed for a max. ambient temperature of 40°C.

The unit shall not be used for use in an oxygen rich environment.

The unit it is not intended to be use with flammable anaesthetics and not intended for use in conjunction with flammable agents.



Essential performance shall be determined within the end medical equipment; however for this medical power supply essential performance is considered MOOP. MOOP is tested within this test report.

- 1. The unit desktop power supply with appliance inlet used for connection to the mains.
- 2. The unit provides internally two fuses in both mains supply leads.
- 3. Power supply unit was evaluated only for Means of Operator Protection:
 - 2 x MOOP between primary and secondary circuit
 - 2 x MOOP between primary and external plastic enclosure surface
 - 1 x MOOP between primary circuit and protective earth
- 4. Secondary output circuit is separated from mains by reinforced insulation and rated SELV. The output does not provide hazard energy level.
- 5. Power supply is provided with safety instructions and technical specifications.
- 6. The power supply is rated as class I (provided in fully plastic enclosure) construction.
- 7. The transformers T1 provides reinforced insulation. This transformer is built up to fulfil the requirement of insulation class B. See also list of safety critical components for details.
- 8. The equipment has been evaluated for use in a Pollution Degree 2 and overvoltage category II environment and a maximum altitude of <2000m.
- 9. Power supply unit is provided with plastic enclosure made by non-flammable material min. UL94 V-1. See also list of safety critical components for details.
- 10. <u>Approval within the end product:</u> Leakage current measurement for the whole medical system is subject of end product evaluation. Earth Leakage current and touch current were measured within this investigation and relate to the power supply only.
- 11. The temperature shall be measured within the end product when power supply is used under other conditions as tested within this test report.
- 12. Power supply cord is not part of the investigation.

Approval within the end product:

- Leakage current measurement should be verified with the unit built into the end product.
- EMC testing has to be performed together with the end medical product.
- Cleaning shall be considered during end product investigation.

Technical Considerations:

Scope of Power Supply evaluation defers the following clauses to be determined as part of the end product investigation:

- Clause 7.5 (Safety Signs),
- Clause 7.9 (Accompanying Documents),
- Clause 9 (ME Hazard), except 9.1 and 9.3 are evaluated,
- Clause 10 (Radiation),
- Clause 14 (PEMS),
- Clause 16 (ME Systems)



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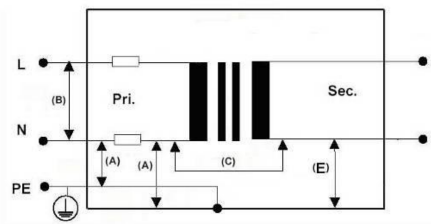
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		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

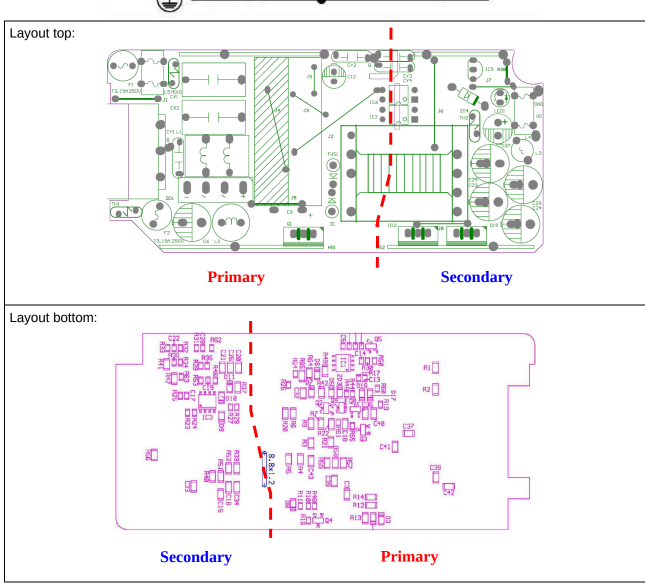
		History sheet	
Date	Report Number	Change	Revision No.
2012-09-11	T223-0207/12	Initial Test Report issued.	_
2019-09-23	T223-0554/19	Test report updated from IEC 60601-1:2005 (3rd Edition) to IEC 60601-1:2005 (3rd Edition) + A1:2012.	1.0
		No changes on the product since initial approval.	
		After review, the following tests were considered required:	
		- Clause 8.7: Leakage current (measured with frequency-non-weighted measuring device)	
		- RMF review	
		Power supply unit additionally evaluated for the latest European differences EN 60601-1:2006 + A1:2013 +A12:2014 and National Differences to IEC 60601-1: 2005 + AM1: 2012.	
		Added new user manual in Enclosure No. 3.	
		Updated table 8.10. List of critical components.	



		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

INSULATION DIAGRAM







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		1 490 10 01 201	1 topolities 1	220 000 17 10
		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

	E: INSULATION								Р
	tion degree				PD2				_
	voltage categor				2				_
Altitu	de			: Up to	2.000 mete	ers			_
	Additional details on parts considered as applied parts				one	Areas for details)			_
Area	Number and type of Means of Protection:	СТІ	Working		Required creepage	Required clearance (mm)	Measured creepage	Measured clearance	Remarks
	MOOP, MOPP		V _{rms}	V_{pk}	(mm)	(11111)	(mm)	(mm)	
A1	1 x MOOP	IIIb	250	354	2,5	2,0	4,0	4,0	Between L, N and PE (near appliance inlet) on PCB
A2	1 x MOOP	IIIb	250	354	2,5	2,0	4,0	2,5	Between diode bridge and PE
A3	1 x MOOP	IIIb	266	574	2,7	2,3	4,9	4,9	Between L, N and PE (near TransformerT 1) on PCB
B1	1 x MOOP	IIIb	250	354	3,0	1,6	7,0	7,0	Measured on main board PCB between L and N before fuse.
B2	1 x MOOP	IIIb	250	354	L to N afte	r fuse.			
					Shall be ve	erified via sł	nort-circuitir	ng.	
C1	2 x MOOP	IIIb	264	560	5,4	4,4	8,1	8,0	Between Primary to Secondary (on PCB)
C2	2 x MOOP	IIIb	264	560	5,4	4,4	9,0	9,0	Between Primary to Secondary on Transformer T1
C3	2 x MOOP	IIIb	223	367	5,0	4,0	7,8	7,8	Between Primary to Secondary on optical insulator IC2



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		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

C4	2 x MOOP	IIIb	201	340	5,0	4,0	7,8	7,8	Between Primary to Secondary on optical insulator IC6
C5	2 x MOOP	IIIb	250	354	5,0	4,0	See *)	5,6	Between Primary to accessible part via opening in enclosure

Supplementary Information:

Transformer T1: Core is considered as floating. Primary and secondary winding is made of copper wire. Min. two layers of insulation foil between primary and secondary winding provided for rainforced insulation.

Tubing provided on all primary and secondary windings to achieve sufficient creapage and clearance distance between primary and secondary.

Electric strenght test (3000 Vac) was done on insulaing foil of the Transformer T1.

*) Min. clearance distance measured between inside enclosure and primary heat sink is 1,3 mm. Three pins of heat sink are fixed on PCB.





INSULATION DIAGRAM CONVENTIONS and GUIDANCE:

A measured value must be provided in the value columns for the device under evaluation. The symbol > (greater than sign) must not be used. Switch-mode power supplies must be re-evaluated in the device under evaluation therefore N/A must not be used with a generic statement that the component is certified.

Insulation diagram is a graphical representation of equipment insulation barriers, protective impedance and protective earthing. If feasible, use the following conventions to generate the diagram:

- All isolation barriers are identified by letters between separate parts of diagram, for example separate transformer
- windings, optocouplers, wire insulation, creepage and clearance distances.
- Parts connected to earth with large dots are protectively earthed. Other connections to earth are functional
- Applied parts are extended beyond the equipment enclosure and terminated with an arrow.
- Parts accessible to the operator only are extended outside of the enclosure, but are not terminated with an arrow.





	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict

4	GENERAL REQUIREMENTS		Р
4.1	Requirements of this standard applied in NORMAL USE and reasonably foreseeable misuse		Р
4.2	RISK MANAGEMENT PROCESS FOR ME EQUIPMENT OR ME	SYSTEMS	Р
4.2.2	General requirement for RISK MANAGEMENT - PROCESS complies with ISO14971 (2007):	See Appended RM Results Table 4.2.2.	Р
4.2.3	Evaluating RISK		Р
4.2.3.1	a) Compliance with the standard reduces residual risk to an acceptable level		Р
	b) Manufacturer has defined risk acceptability criteria in the RISK MANAGEMENT PLAN:	RISK MANAGEMENT PLAN Document:	Р
		GT-RMPLAN2019-001	
	c) When no specific technical requirements provided manufacturer has determined HAZARDS or HAZARDOUS SITUATIONS exists.		Р
	- HAZARDS or HAZARDOUS SITUATIONS have been evaluated using the RISK MANAGEMENT PROCESS.		Р
4.2.3.2	MANUFACTURER has addressed HAZARDS or HAZARDOUS SITUATIONS not specifically addressed in the IEC 60601-1 series.	Manufacturer addressed all hazards and hazardous situations during risk analysis.	P
4.3	Performance of clinical functions necessary to achieve intended use or that could affect the safety of the ME EQUIPMENT or ME SYSTEM were identified during RISK ANALYSIS.	Power supply unit is not end medical product; therefore no clinical functions specified by the manufacturer.	N/A
	- Performance limits were identified in both NORMAL CONDITION and SINGLE FAULT CONDITION.		N/A
	- Loss or degradation of performance beyond the limits specified by the MANUFACTURER were evaluated		N/A
	- Functions with unacceptable risks are identified as ESSENTIAL PERFORMANCE:	Power supply unit is not end medical product; therefore no clinical functions specified by the manufacturer.	N/A
	- RISK CONTROL measures implemented		N/A
	- Methods used to verify the effectiveness of RISK CONTROL measures implemented		N/A
4.4	EXPECTED SERVICE LIFE stated in RISK MANAGEMENT FILE:	The expected service life is specified by the manufacturer: 5 years.	Р
4.5	Alternative RISK CONTROL methods utilized:	No equivalent safety used.	N/A



	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict
	RESIDUAL RISK resulting from the alternative RISK CONTROL measures or tests is acceptable and comparable to RESIDUAL RISK resulting from application of this standard:	No equivalent safety used.	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5) Alternative means based scientific data or clinical opinion or comparative studies:		N/A
4.6	RISK MANAGEMENT PROCESS identifies parts that can come into contact with PATIENT but not defined as APPLIED PARTS, subjected to the requirements for APPLIED PARTS, except for Clause 7.2.10:	See Appended Insulation Diagram Table and RM Results Table 4.6 No such parts.	N/A
	MANUFACTURER assesses the risk of accessible parts coming into contact with the patient: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	No such parts. No RM considered necessary	N/A
	Assessment identified the APPLIED PART TYPE requirements:	Type B requirements considered for power supply output and power supply accessible enclosure.	Р
4.7	ME EQUIPMENT remained SINGLE FAULT SAFE, or the RISK remained acceptable as determined by Clause 4.2:	Short circuit or open circuit of relevant single components performed. See Table 13.2: Single fault conditions in accordance with 13.2.2 to 13.2.13, inclusive.	P
	MANUFACTURER RISK ANALYSIS was used to determine failures to be tested:	RISK ANALYSIS reference: EL5	Р
	(ISO 14971 CI. 4.2-4.4) Failure of any one component at a time that could result in a HAZARDOUS SITUATION, including those in 13.1, simulated physically or theoretically:	(ISO 14971 Cl. 4.2-4.4) See appended Table 13.2 for simulated physical test.	P
4.8	All components and wiring whose failure could result in a HAZARDOUS SITUATION used according to their applicable ratings, unless specified:	All components are suitable and used within their rating. See appended table 8.10.	Р
	Components and wiring exception in the standard or by RISK MANAGEMENT PROCESS		Р
	RISK MANAGEMENT PROCESS assesses components to identify components where the failure results in a HAZARDOUS SITUATION for components used outside their ratings:	RMF Reference to specific RISKS: (ISO 14971 CI)	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)		
	MANUFACTURER identified components where the failure results in a HAZARDOUS SITUATION:	All components are suitable and used within their ratings.	N/A

Р

Ρ

N/A

N/A

Ρ

Power supply unit is suitable

for connection to supply mains.

Not hand-held ME Equipment.

See appended Table 4.11

100-240 Vac



4.10.1

4.10.2

4.11

	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict
	Components determined to be acceptable where used as a MEANS OF PROTECTION:	All components are suitable and used within their ratings.	N/A
	Reliability of components used as MEANS OF PROTECTION assessed for conditions of use in ME EQUIPMENT, and they complied with one of the following		Р
	a) Applicable safety requirements of a relevant IEC or ISO standard	Approved critical components used.	Р
	b) Requirements of this standard applied in the absence of a relevant IEC or ISO standard		Р
4.9	A COMPONENT WITH HIGH-INTEGRITY CHARACTERISTICS provided and selected appropriately:		N/A
	RISK MANAGEMENT FILE includes an assessment to determine if the failure of components results in unacceptable RISK:	No such components provided. No RM considered necessary.	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)		
	Components identified and required to be COMPONENTS WITH HIGH INTEGRITY CHARACTERISTIC:	See Table 8.10 b	Р
4.10	Power supply		Р

ME EQUIPMENT is suitable for connection to

Maximum rated voltage for ME EQUIPMENT

- 500 V for all other ME EQUIPMENT and ME

intended to be connected to SUPPLY MAINS:

indicated power source (select applicable):

- 250 V for HAND-HELD ME EQUIPMENT (V)

- 250 V d.c. or single-phase a.c., or 500 V poly-

phase a.c. for ME EQUIPMENT and ME SYSTEMS with a RATED input ≤ 4 kVA (V)

Steady-state measured input of ME EQUIPMENT or

ME SYSTEM at RATED voltage or voltage range and at operating settings indicated in instructions for use didn't exceed marked rating by more than 10%

5	GENERAL REQUIREMENTS FOR TESTING ME	GENERAL REQUIREMENTS FOR TESTING ME EQUIPMENT	
5.1	Test not performed when analysis indicated condition being tested was adequately evaluated by other tests or methods:	Type test performed according to all applicable clauses of standard IEC 60601-1:2005 + A1:2012.	Р

SYSTEMS

Power input



	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict
	RISK MANAGEMENT FILE identifies combinations of simultaneous independent faults that could result in a HAZARDOUS SITUATION. (ISO 14971 Cl. 4.2-4.4)	Type test performed according to all applicable clauses of standard IEC 60601-1:2005 + A1:2012. No RM considered necessary.	N/A
5.3	Tests conducted within the environmental conditions specified in technical description		Р
	Temperature (°C), Relative Humidity (%):	0-40 °C Up to 90 % RH.	_
	Atmospheric Pressure (kPa):	620 hPa to 1060 hPa	_
5.5	a) Supply voltage during tests was the least favourable of the voltages specified in 4.10.2 or voltages marked on ME EQUIPMENT (V)::		Р
	b) ME EQUIPMENT marked with a RATED frequency range tested at the least favourable frequency within the range (Hz):	50-60Hz	Р
	c) ME EQUIPMENT with more than one RATED voltage, both a.c./ d.c. or both external power and INTERNAL ELECTRICAL POWER SOURCE tested in conditions (see 5.4) related to the least favourable voltage, nature of supply, and type of current:	Supply voltage: 100-240 Vac Only AC supply voltage used for supplying power supply unit.	P
	d) ME EQUIPMENT intended for only d.c. supply connection tested with d.c. and influence of polarity considered:	AC supply voltage only.	N/A
	e)ME EQUIPMENT tested with alternative ACCESSORIES and components specified in ACCOMPANYING DOCUMENTS to result in the least favourable conditions:	No accessories provided.	N/A
	f) ME EQUIPMENT connected to a separate power supply as specified in instructions for use	Mains operated equipment.	Р
5.7	ME EQUIPMENT or parts thereof affected by climatic conditions were set up completely, or partially, with covers detached and subjected to a humidity preconditioning prior to tests of Clauses 8.7.4 and 8.8.3	Complete unit was subjected to humidity preconditioning treatment.	P
	ME EQUIPMENT heated to a temperature between	Revision 1.0:	_
	T and T + 4°C for at least 4 h and placed in a humidity chamber and ambient within 2 °C of T in range of +20°C to +32°C for indicated time	2 days (48h): T = 27,5°C RH= 95,1 %	
5.9	Determination of APPLIED PARTS and ACCESSIBLE PARTS		P
5.9.1	APPLIED PARTS identified by inspection and reference to ACCOMPANYING DOCUMENTS:	No APPLIED PARTS provided.	N/A
5.9.2	ACCESSIBLE PARTS		Р



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	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict
5.9.2.1	Accessibility determined using standard test finger of Fig. 6	See Appended Table 5.9.2 Power supply unit is provided with plastic enclosure without openings (to cover all internal parts).	Р
5.9.2.2	Test hook of Fig. 7 inserted in all openings of ME EQUIPMENT and pulled with a force of 20 N for 10 s	No openings provided.	N/A
5.9.2.3	Conductive parts of actuating mechanisms of electrical controls accessible after removal of handles, knobs, levers and the like regarded as ACCESSIBLE PARTS	No actuating mechanism.	N/A
	Conductive parts of actuating mechanisms not considered ACCESSIBLE PARTS when removal of handles, knobs, required use of a TOOL:	No actuating mechanism.	N/A

6.2	CLASSIFICATION OF ME EQUIPMENT AND ME SYSTEMS		Р
	CLASS I ME EQUIPMENT, externally powered	Power supply unit is Class I equipment.	Р
	CLASS II ME EQUIPMENT, externally powered		N/A
	INTERNALLY POWERED ME EQUIPMENT	Equipment is not internally powered equipment.	N/A
	EQUIPMENT with means of connection to a SUPPLY MAINS complied with CLASS I or CLASS II ME EQUIPMENT requirements when so connected, and when not connected to SUPPLY MAINS with INTERNALLY POWERED ME EQUIPMENT requirements		N/A
	TYPE B APPLIED PART	No applied parts provided.	N/A
	TYPE BF APPLIED PART		N/A
	TYPE CF APPLIED PART		N/A
	DEFIBRILLATION-PROOF APPLIED PARTS		N/A
6.3	ENCLOSURES classified according to degree of protection against ingress of water and particulate matter as per IEC 60529	No IP protection	N/A
6.4	ME EQUIPMENT or its parts intended to be sterilized classified according to method(s) of sterilization in instructions for use:	No such parts.	N/A
6.5	ME EQUIPMENT and ME SYSTEMS intended for use in an OXYGEN RICH ENVIRONMENT classified for such use and complied with 11.2.2	Power supply not investigated for OXYGEN RICH ENVIRONMENT.	N/A
6.6	CONTINUOUS OF Non-CONTINUOUS OPERATION:	The equipment is intended for continuous operation.	Р



IEC 60601-1			
Clause	Requirement + Test	Result - Remark	Verdict

7	ME EQUIPMENT IDENTIFICATION, MARKING, A	ND DOCUMENTS	Р
7.1.2	Legibility of Markings Test for Markings specified in Clause 7.2-7.6:	See Appended Table 7.1.2	Р
7.1.3	Required markings can be removed only with a TOOL or by appreciable force, are durable and remain CLEARLY LEGIBLE during EXPECTED SERVICE LIFE OF ME EQUIPMENT IN NORMAL USE	See appended Tables 7.1.3 and 8.10	Р
7.2	Marking on the outside of ME EQUIPMENT or ME EQ	NUIPMENT parts	Р
7.2.1	At least markings in 7.2.2, 7.2.5, 7.2.6, 7.2.10, and 7.2.13 were applied when size of EQUIPMENT, its part, an ACCESSORY, or ENCLOSURE did not permit application of all required markings	See attached copy of Marking Plate All required markings are provided on the marking plate.	Р
	Remaining markings fully recorded in ACCOMPANYING DOCUMENTS	All necessary information are provided on the marking plate.	Р
	Markings applied to individual packaging when impractical to apply to ME EQUIPMENT	Packaging not part of the investigation.	N/A
	Single use item marked:	EUT is not intended for single use.	N/A
7.2.2	ME EQUIPMENT marked with:		Р
	- the name or trademark and contact information of the MANUFACTURER		Р
	- a MODEL OR TYPE REFERENCE	See attached copy of Marking Plate	Р
	- a serial number or lot or batch identifier; and		Р
	- the date of manufacture or use by date		Р
	Detachable components of the ME EQUIPMENT not marked; misidentification does not present an unacceptable risk, or	Detachable mains plugs not marked, misidentification does not present an unacceptable risk.	P
	RISK MANAGEMENT FILE includes an assessment of the RISKS relating to misidentification of all detachable parts:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.4)	(ISO 14971 Cl)	
	Detachable components of the ME EQUIPMENT are marked with the name or trademark of the MANUFACTURER, and	Standard plug used. Misidentification does not result in an unacceptable risk, there no markings required.	N/A
	- a MODEL OR TYPE REFERENCE		N/A
	Software forming part of a PEMS identified with a unique identifier:	No software incorporated.	N/A
7.2.3	Symbol 11 on Table D.1 used, optionally, advice to OPERATOR to consult ACCOMPANYING DOCUMENTS	Symbol not required. EUT is medical in power supply unit.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Safety sign 10 on Table D.2) used, advising OPERATOR that ACCOMPANYING DOCUMENTS must be consulted		N/A
7.2.4	Accessories marked with name or trademark and contact information of their MANUFACTURER, and:	No accessories provided.	N/A
	- with a MODEL or TYPE REFERENCE		N/A
	- a serial number or lot or batch identifier		N/A
	- the date of manufacture or use by date		N/A
	Markings applied to individual packaging when not practical to apply to ACCESSORIES		N/A
7.2.5	ME EQUIPMENT and ME SYSTEM intended to receive power from other equipment, provided with one of the following	Mains operated equipment.	N/A
	- the name or trademark of the manufacturer of the other electrical equipment and type reference marked adjacent to the relevant connection point; or		N/A
	 Table D.2, safety sign No. 10 adjacent to the relevant connection point and listing of the required details in the instructions for use; or 		N/A
	 Special connector style used that is not commonly available on the market and listing of the required details in the instructions for use. 		N/A
7.2.6	Connection to the Supply Mains		Р
	Marking appearing on the outside of part containing SUPPLY MAINS connection and, adjacent to connection point	Marking plate provided on the surface of the plastic enclosure.	Р
	For PERMANENTLY INSTALLED ME EQUIPMENT, NOMINAL supply voltage or range marked inside or outside of ME EQUIPMENT	EUT is not permanently installed equipment.	N/A

OPERATOR that ACCOMPANYING DOCUMENTS must be consulted		
ACCESSORIES marked with name or trademark and contact information of their MANUFACTURER, and:	No accessories provided.	N/A
- with a MODEL or TYPE REFERENCE		N/A
- a serial number or lot or batch identifier		N/A
- the date of manufacture or use by date		N/A
Markings applied to individual packaging when not practical to apply to ACCESSORIES		N/A
ME EQUIPMENT and ME SYSTEM intended to receive power from other equipment, provided with one of the following	Mains operated equipment.	N/A
- the name or trademark of the manufacturer of the other electrical equipment and type reference marked adjacent to the relevant connection point; or		N/A
 Table D.2, safety sign No. 10 adjacent to the relevant connection point and listing of the required details in the instructions for use; or 		N/A
 Special connector style used that is not commonly available on the market and listing of the required details in the instructions for use. 		N/A
Connection to the Supply Mains		Р
Marking appearing on the outside of part containing SUPPLY MAINS connection and, adjacent to connection point	Marking plate provided on the surface of the plastic enclosure.	Р
For PERMANENTLY INSTALLED ME EQUIPMENT, NOMINAL supply voltage or range marked inside or outside of ME EQUIPMENT	EUT is not permanently installed equipment.	N/A
- RATED supply voltage(s) or RATED voltage	100-240 Vac	Р
range(s) with a hyphen (-) between minimum and maximum voltages (V, V-V):	Markings provided on the marking plate.	
Multiple RATED supply voltages or multiple RATED supply voltage ranges are separated by (V/V):		N/A
- Nature of supply and type of current:	Symbol " ~ " provided near rated supply voltage.	Р
Symbols 1-5, Table D.1 (used for same parameters:	Symbol No. 1 from table D.1 used for input voltage.	Р
	Symbol No. 4 from table D.1 used for output voltage.	
- RATED supply frequency or RATED frequency	50-60 Hz	Р
	ACCESSORIES marked with name or trademark and contact information of their MANUFACTURER, and	ACCESSORIES marked with name or trademark and contact information of their MANUFACTURER, and



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Clause	Requirement + Test	Result - Remark	Verdict

	- Symbol 9 of Table D.1 used for CLASS II ME EQUIPMENT:	Class I equipment.	N/A
7.2.7	RATED input in amps or volt-amps, (A, VA):	Rated input expressed in Amperes:	Р
		1,6-0,7 A	
		(specified for lower and upper voltage range)	
	RATED input in amps or volt-amps, or in watts when power factor exceeds 0.9 (A, VA, W):	Rated input expressed in Amperes.	Р
		Rated input expressed in Amperes:	Р
	the range or ranges when the range(s) is/are greater than ± 10 % of the mean value of	1,6-0,7 A	
	specified range (A, VA,W):	(specified for lower and upper voltage range)	
	Input at mean value of range marked when range limits do not differ by more than 10 % from mean value (A, VA, W):		N/A
	Marking includes long-time and most relevant momentary volt-ampere ratings when provided, each plainly identified and indicated in ACCOMPANYING DOCUMENTS (VA):		N/A
	Marked input of ME EQUIPMENT provided with means for connection of supply conductors of other electrical equipment includes RATED and marked output of such means (A, VA, W):		N/A
7.2.8	Output connectors		Р
7.2.8.2	Output connectors are marked, except for MULTIPLE SOCKET-OUTLETS or connectors intended for specified ACCESSORIES or equipment	Rating of the output provided on the marking plate.	Р
	Rated Voltage (V), Rated Current (A):	See attached copy of Marking Plate.	_
	Rated Power (W), Output Frequency (Hz):	DC output voltage.	_
7.2.9	ME EQUIPMENT or its parts marked with the IP environmental Code per IEC 60529 according to classification in 6.3 (Table D.3, Code 2), marking optional for ME EQUIPMENT or parts rated IPX0:	Ordinary equipment (IP20).	N/A
7.2.10	Degrees of protection against electric shock as classified in 6.2 for all APPLIED PARTS marked with relevant symbols:	No APPLIED PARTS in power supply.	N/A
	TYPE B APPLIED PARTS with symbol 19 of Table D.1		N/A
	TYPE BF APPLIED PARTS with symbol 20 of Table D.1:		N/A
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Clause	Requirement + Test	Result - Remark	Verdict
	TYPE CF APPLIED PARTS with symbol 21 of Table D.1:		N/A
	DEFIBRILLATION-PROOF APPLIED PARTS marked with symbols 25-27 of Table D.1:		N/A
	Proper symbol marked adjacent to or on connector for APPLIED PART:		N/A
	Safety sign 2 of Table D.2 placed near relevant outlet:		N/A
	An explanation indicating protection of ME EQUIPMENT against effects of discharge of a cardiac defibrillator depends on use of proper cables included in instructions for use:		N/A
7.2.11	ME EQUIPMENT suitable for CONTINUOUS OPERATION	No marking provided, therefore Power supply unit is suitable for continuous operation.	P
	DUTY CYCLE for ME EQUIPMENT intended for non- CONTINUOUS OPERATION appropriately marked to provide maximum "on" and "off" time:	EUT is designed for continuous operation.	N/A
7.2.12	Type and full rating of a fuse marked adjacent to ACCESSIBLE fuse-holder	Fuse is not accessible by the operator.	N/A
	Fuse type:		_
	Voltage (V) and Current (A) rating:		_
	Operating speed (s) and Breaking capacity:		_
7.2.13	Physiological effects – safety sign and warning statements	Equipment does not produce physiological effects.	N/A
	Nature of HAZARD and precautions for avoiding or minimizing the associated RISK described in instructions for use	RMF Reference to specific RISKS: (ISO 14971 CI)	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.3)	Equipment does not produce physiological effects. No RM considered necessary.	
7.2.14	HIGH VOLTAGE TERMINAL DEVICES on the outside of ME EQUIPMENT accessible without the use of a TOOL marked with symbol 24 of Table D.1	No high voltage terminal devices provided.	N/A
7.2.15	Requirements for cooling provisions marked:	Not provided.	N/A
7.2.17	Packaging marked with special handling instructions for transport and/or storage:	No special handling requirements.	N/A
	Permissible environmental conditions marked on outside of packaging:	Power supply is not end medical product, shall be considered during end medical product approval.	N/A
	Packaging marked with a suitable safety sign indicating premature unpacking of ME EQUIPMENT could result in an unacceptable RISK:		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	RISK MANAGEMENT FILE includes the assessment to determine premature unpacking of ME EQUIPMENT or its parts could result in an unacceptable RISK: (ISO 14971 Cl. 4.2-4.4, 5, 6.3-6.4)	RMF Reference to specific RISKS: (ISO 14971 CI)	N/A
	Packaging of sterile ME EQUIPMENT or ACCESSORIES marked sterile and indicates the methods of sterilization		N/A
7.2.18	RATED maximum supply pressure from an external source marked on ME EQUIPMENT adjacent to each input connector, and:	No pressure used.	N/A
	- the RATED flow rate also marked		N/A
7.2.19	Symbol 7 of Table D.1 marked on FUNCTIONAL EARTH TERMINAL:	No functional earth terminal provided.	N/A
		EUT is Class I equipment provided with protective earthing.	
7.2.20	Removable protective means marked to indicate the necessity for replacement when the function is no longer needed:	Equipment does not provide alternative applications.	N/A
7.2.21	MOBILE ME EQUIPMENT marked with its mass including its SAFE WORKING LOAD in kilograms:	No mobile equipment.	N/A
7.3	Marking on the inside of ME EQUIPMENT or ME EQUIP	PMENT parts	Р
7.3.1	Maximum power loading of heating elements or lamp-holders designed for use with heating lamps marked near or in the heater (W):	No heating elements provided.	N/A
	A marking referring to ACCOMPANYING DOCUMENTS provided for heating elements or lamp-holders designed for heating lamps that can be changed only by SERVICE PERSONNEL using a TOOL		N/A
7.3.2	Symbol 24 of Table D.1, or safety sign No.3 of Table D.2 used to mark presence of HIGH VOLTAGE parts:	No HIGH VOLTAGE parts within the equipment.	N/A
7.3.3	Type of battery and mode of insertion marked:	No battery provided.	N/A
	An identifying marking provided referring to instructions in ACCOMPANYING DOCUMENTS for batteries intended to be changed only by SERVICE PERSONNEL using a TOOL:		N/A
	A warning provided indicating replacement of lithium batteries or fuel cells when incorrect replacement would result in an unacceptable RISK		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	RISK MANAGEMENT FILE includes an assessment to determine the replacement of lithium batteries or fuel cells leads to an unacceptable RISK if replaced incorrectly:	RMF Reference to specific RISKS: (ISO 14971 CI) Neither batteries nor battery	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.3)	compartments incorporated. No RM considered necessary.	
	ACCOMPANYING DOCUMENTS contain a warning indicating the replacement of lithium batteries or fuel cells by inadequately trained personnel could result in a HAZARD		N/A
7.3.4	Fuses, replaceable THERMAL CUT-OUTS and OVER-CURRENT RELEASES, accessible by use of a TOOL Identified	Inside the EUT a reference to the F1 and F2 provided near fuses on the PCB.	Р
	Voltage (V) and Current (A) rating:	T3.15A250V	_
	Operating speed(s), size & breaking capacity.:	During single fault conditions, no safety hazard occurs in case of primary fuse opened.	-
7.3.5	PROTECTIVE EARTH TERMINAL marked with symbol 6 of Table D.1	Protective earth terminal provided as part of the approved appliance inlet.	P
		Symbol number 6 of Table D.1 provided on the appliance inlet enclosure.	
	Markings on or adjacent to PROTECTIVE EARTH TERMINALS not applied to parts requiring removal to make the connection, and remained visible after connection made		P
7.3.6	Symbol 7 of Table D.1 marked on FUNCTIONAL EARTH TERMINALS	No functional earthing provided.	N/A
7.3.7	Terminals for supply conductors marked adjacent to terminals:	Mains supply terminals are part of the approved appliance inlet in accordance with IEC 60320-1.	N/A
		Marking by means of L and N is therefore not applicable.	
	Terminals for supply connections are not marked, the RISK MANAGEMENT FILE includes an assessment of the RISKS resulting from misconnections	RMF Reference to specific RISKS: (ISO14971 CI)	N/A
	Terminal markings included in ACCOMPANYING DOCUMENTS when ME EQUIPMENT too small to accommodate markings		N/A
	Terminals exclusively for neutral supply conductor in PERMANENTLY INSTALLED ME EQUIPMENT marked with Code 1 of Table D.3		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Marking for connection to a 3-phase supply, complies with IEC 60445		N/A
	Markings on or adjacent to electrical connection points not applied to parts requiring removal to make connection, and remained visible after connection made		N/A
7.3.8	"For supply connections, use wiring materials suitable for at least X °C" or equivalent, marked at the point of supply connections	No special requirements for detachable power supply cords. Power supply cord is not part of the investigation.	N/A
	Statement not applied to parts requiring removal to make the connection, and CLEARLY LEGIBLE after connections made		N/A
7.4	Marking of controls and instruments		N/A
7.4.1	The "on" & "off" positions of switch to control power to ME EQUIPMENT or its parts, including mains switch, marked with symbols 12 and 13 of Table D.1 or		N/A
	- indicated by an adjacent indicator light, or		N/A
	- indicated by other unambiguous means		N/A
	The "on/off" positions of push button switch with bi-stable positions marked with symbol 14 of Table D.1, and		N/A
	- status indicated by adjacent indicator light		N/A
	status indicated by other unambiguous means		N/A
	The "on/off" positions of push button switch with momentary on position marked with symbol 15 of Table D.1 or		N/A
	- status indicated by adjacent indicator light		N/A
	- status indicated by other unambiguous means		N/A
7.4.2	Different positions of control devices/switches indicated by figures, letters, or other visual means	No control devices provided.	N/A
	RISK MANAGEMENT FILE identifies controls where a change in setting during NORMAL USE results in an unacceptable RISK: (ISO 14971 Cl. 4.2-4.4, 5, 6.2, 6.3)	RMF Reference to specific RISKS: List of controls: (ISO14971 Cl) No control devices/switches provided. No RM considered necessary.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Controls provided with an associated indicating device when change of setting of a control could result in an unacceptable RISK to PATIENT in NORMAL USE:		N/A
	 or an indication of direction in which magnitude of the function changes 		N/A
	Control device or switch that brings the ME EQUIPMENT into the "stand-by" condition marked with symbol IEC 60417-5009	No stand-by condition.	N/A
7.4.3	Numeric indications of parameters on ME EQUIPMENT expressed in SI units according to ISO 80000-1 except the base quantities listed in Table 1 expressed in the indicated units		N/A
	ISO 80000-1 applied for application of SI units, their multiples, and certain other units		N/A
	All Markings in Sub-clause 7.4 complied with tests and criteria of 7.1.2 and 7.1.3:	See Appended Tables 7.1.2 and 7.1.3.	N/A
7.5	Safety signs		N/A
	Safety sign with established meaning used	No safety signs provided.	N/A
	RISK MANAGEMENT PROCESS identifies markings used to convey a warning, prohibition or mandatory action that mitigate a RISK not obvious to the OPERATOR	RMF Reference to specific RISK & Marking: Safety Sign Used: (ISO 14971 Cl) No safety signs provided. No	N/A
	Affirmative statement together with safety sign placed in instructions for use if insufficient space on ME EQUIPMENT	RM considered necessary.	N/A
	Specified colours in ISO 3864-1 used for safety signs:		N/A
	Safety notices include appropriate precautions or instructions on how to reduce RISK(S)		N/A
	Safety signs including any supplementary text or symbols described in instructions for use		N/A
	- and in a language acceptable to the intended OPERATOR		N/A
7.6	Symbols		N/A
7.6.1	Meanings of symbols used for marking described in instructions for use:	Symbols shall be described within the end product documentation.	N/A
7.6.3	Symbols used for controls and performance conform to the IEC or ISO publication where symbols are defined, as applicable		N/A
7.7	Colours of the insulation of conductors		Р



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Clause	Requirement + Test	Result - Remark	Verdict
7.7.1	PROTECTIVE EARTH CONDUCTOR identified by green and yellow insulation	EUT is Class I equipment. Protective earth conductor provided from approved appliance inlet to PCB (soldered on PCB) green/yellow colour of insulation.	P
7.7.2	Insulation on conductors inside ME EQUIPMENT forming PROTECTIVE EARTH CONNECTIONS identified by green and yellow at least at terminations		N/A
7.7.3	Green and yellow insulation identify only following conductors:	Protective earth conductor provided from approved appliance inlet to PCB (soldered on PCB).	N/A
	- PROTECTIVE EARTH CONDUCTORS		N/A
	- conductors specified in 7.7.2		N/A
	- POTENTIAL EQUALIZATION CONDUCTORS		N/A
	- FUNCTIONAL EARTH CONDUCTORS		N/A
7.7.4	Neutral conductors of POWER SUPPLY CORDS are "light blue"	Power supply cord is not part of the investigation.	N/A
7.7.5	Colours of conductors in POWER SUPPLY CORDS in accordance with IEC 60227-1 or IEC 60245-1		N/A
7.8	Indicator lights and controls		Р
7.8.1	Red indicator lights used only for Warning	No red indicator light provided.	N/A
	Yellow indicator lights used only for Caution	No yellow indicator light provided.	N/A
	Green indicator lights used only for Ready for use	Green indicator LED provided meaning power supply is operating and delivering required SELV voltage on the output.	P
	Other colours: Meaning other than red, yellow, or green (colour, meaning):	No other colours used.	N/A
7.8.2	Red used only for emergency control		N/A
7.9	ACCOMPANYING DOCUMENTS		Р
7.9.1	ME EQUIPMENT accompanied by documents containing instructions for use, and a technical description	Safety instruction and technical specification provided by the manufacturer.	Р
	ACCOMPANYING DOCUMENTS identify ME EQUIPMENT by the following, as applicable:		Р
	- Name or trade-name of MANUFACTURER and contact information for the RESPONSIBLE ORGANIZATION can be referred to	Provided within safety instruction and technical specification.	Р
	- MODEL OF TYPE REFERENCE	GT-500160-30	Р



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Clause	Requirement + Test	Result - Remark	Verdict
	When ACCOMPANYING DOCUMENTS provided electronically, USABILITY ENGINEERING PROCESS includes instructions as to what is required in hard copy or as markings on ME EQUIPMENT	Documents are provided in hard copy.	N/A
	ACCOMPANYING DOCUMENTS specify special skills, training, and knowledge required of OPERATOR or RESPONSIBLE ORGANIZATION and environmental restrictions on locations of use		N/A
	ACCOMPANYING DOCUMENTS written at a level consistent with education, training, and other needs of individuals for whom they are intended	Technical specification is intended for the end product development personal.	Р
' .9.2	Instructions for use include the required inform	ation	Р
7.9.2.1	- use of ME EQUIPMENT as intended by the MANUFACTURER:		Р
	- frequently used functions,		N/A
	- known contraindication(s) to use of ME EQUIPMENT		N/A
	- parts of the ME EQUIPMENT that are not serviced or maintained while in use with the patient	No such parts.	N/A
	- name or trademark and address of the MANUFACTURER		Р
	- MODEL OR TYPE REFERENCE		Р
	Instruction for use included the following when the PATIENT is an intended OPERATOR:	End product consideration. Power supply unit is not end medical product.	N/A
	- the PATIENT is an intended OPERATOR		N/A
	- warning against servicing and maintenance while the ME EQUIPMENT is in use		N/A
	- functions the PATIENT can safely use and, where applicable, which functions the PATIENT cannot safely use; and		N/A
	-maintenance the PATIENT can perform		N/A
	Classifications as in Clause 6, all markings per Clause 7.2, and explanation of safety signs and symbols marked on ME EQUIPMENT	Provided in technical specification.	Р
	Instructions for use are in a language acceptable to the intended operator	English language evaluated.	Р
'.9.2.2	Instructions for use include all warning and safety notices	Warning and safety notices shall be provided within the end product documentation.	N/A
	Warning statement for CLASS I ME EQUIPMENT included	Class I equipment.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Warnings regarding significant RISKS of reciprocal interference posed by ME EQUIPMENT during specific investigations or treatments	End product consideration. Power supply unit is not end medical product.	N/A
	Information on potential electromagnetic or other interference and advice on how to avoid or minimize such interference	Warning shall be provided within the end product documentation depending on the type of the end product.	N/A
	Warning statement for ME EQUIPMENT supplied with an integral MULTIPLE SOCKET-OUTLET provided	No MULTIPLE SOCKET-OUTLET provided.	N/A
	The RESPONSIBLE ORGANIZATION is referred to this standard for the requirements applicable to ME SYSTEMS		N/A
7.9.2.3	Statement on ME EQUIPMENT for connection to a separate power supply provided in instructions	Mains operated equipment.	N/A
7.9.2.4	Warning statement for mains- operated ME EQUIPMENT with additional power source not automatically maintained in a fully usable condition indicating the necessity for periodic checking or replacement of power source	Mains operated equipment. No additional power source incorporated.	N/A
	RISK MANAGEMENT FILE assesses the RISK resulting from leakage of batteries:	Specific RISKS: (ISO 14971 CI)	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.3)	Neither batteries nor battery compartments incorporated. RM considered necessary.	
	Where the RISK is unacceptable, the IFU includes a warning to remove the battery if the ME EQUIPMENT is not likely to be used for some time:	No batteries provided.	N/A
	Specifications of replaceable INTERNAL ELECTRICAL POWER SOURCE when provided:	No internal replaceable power source provided. Power supply unit is direct plug-in equipment.	N/A
	Warning indicating ME EQUIPMENT must be connected to an appropriate power source when loss of power source would result in an unacceptable RISK:	Power supply unit is not end medical product; shall be considered during end medical product approval.	N/A
7.9.2.5	Instructions for use include a description of ME EQUIPMENT, its functions, significant physical and performance characteristics together with the expected positions of OPERATOR, PATIENT, or other persons near ME EQUIPMENT IN NORMAL USE	medical product; shall be considered during end medical product approval.	N/A
	Information provided on materials and ingredients PATIENT OF OPERATOR is exposed to		N/A



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Clause	Requirement + Test	Result - Remark	Verdict	

	Restrictions specified on other equipment or NETWORK/DATA COUPLINGS, other than those forming part of an ME SYSTEM, to which a SIGNAL INPUT/OUTPUT PART may be connected	No signal input/signal output parts provided.	N/A
	APPLIED PARTS specified	No applied parts provided.	N/A
7.9.2.6	Information provided indicating where the installation instructions may be found or information on qualified personnel who can perform the installation		N/A
7.9.2.7	Instructions provided indicating not to position ME EQUIPMENT to make it difficult to operate the disconnection device	Appliance coupler used as disconnecting device. The following note provided within safety instruction: The socket-outlet shall be installed near the equipment and shall be easily accessible.	Р
7.9.2.8	Necessary information provided for OPERATOR to bring ME EQUIPMENT into operation	Power supply unit is part of the investigation.	N/A
7.9.2.9	Information provided to operate ME EQUIPMENT		N/A
	Meanings of figures, symbols, warning statements, abbreviations and indicator lights described in instructions for use		N/A
7.9.2.10	A list of all system messages, error messages, and fault messages provided with an explanation of messages including important causes and possible action(s) to be taken to resolve the problem indicated by the message		N/A
7.9.2.11	Information provided for the OPERATOR to safely terminate operation of ME EQUIPMENT	Termination of the ME equipment shall be investigated within the end product.	N/A
7.9.2.12	Information provided on cleaning, disinfection, and sterilization methods, and applicable parameters that can be tolerated by ME EQUIPMENT parts or ACCESSORIES specified	Power supply unit is not end medical product; shall be considered during end medical product approval.	N/A
	Components, ACCESSORIES or ME EQUIPMENT marked for single use, except when required by MANUFACTURER to be cleaned, disinfected, or sterilized prior to use	No such parts.	N/A
7.9.2.13	Instructions provided on preventive inspection, calibration, maintenance and its frequency	Power supply unit is maintenance free.	N/A
	Information provided for safe performance of routine maintenance necessary to ensure continued safe use of ME EQUIPMENT		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Parts requiring preventive inspection and maintenance to be performed by SERVICE PERSONNEL identified including periods of application		N/A
	Instructions provided to ensure adequate maintenance of ME EQUIPMENT containing rechargeable batteries to be maintained by anyone other than SERVICE PERSONNEL		N/A
7.9.2.14	A list of ACCESSORIES, detachable parts, and materials for use with ME EQUIPMENT provided	No such parts.	N/A
	Other equipment providing power to ME SYSTEM sufficiently described	Mains operated equipment.	N/A
7.9.2.15	Disposal of waste products, residues, etc., and of ME EQUIPMENT and ACCESSORIES at the end of their EXPECTED SERVICE LIFE are identified in the instruction for use:	Provided within user instructions.	P
7.9.2.16	Instructions for use include information specified in 7.9.3 or identify where it can be found (e.g. in a service manual)		N/A
7.9.2.17	Instruction for use for ME EQUIPMENT emitting radiation for medical purposes, indicate the nature, type, intensity and distribution of this radiation	Equipment does not emit radiation for medical purposes.	N/A
7.9.2.18	The instructions for use for ME EQUIPMENT or ACCESSORIES supplied sterile indicate that they have been sterilized and the method of sterilization	No parts supplied sterile.	N/A
	The instructions for use indicate the necessary instructions in the event of damage to the sterile packaging, and where appropriate, details of the appropriate methods of resterilization		N/A
7.9.2.19	The instructions for use contain a unique version identifier:	Version: GT-500160 series REV 1.0	Р
7.9.3	Technical description		Р
7.9.3.1	All essential data provided for safe operation, transport, storage, and measures or conditions necessary for installing ME EQUIPMENT, and preparing it for use		P
Technical description separable from instruction information, as follows		ns for use contains required	Р
	all applicable classifications in Clause 6, warning and safety notices, and explanation of safety signs marked on ME EQUIPMENT		P
	- a brief description of the ME EQUIPMENT, how the ME EQUIPMENT functions and its significant physical and performance characteristics; and	Provided within technical specifications.	Р
	a unique version identifier::		Р



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Clause	Requirement + Test		Result - Remark	Verdict

	MANUFACTURER'S optional requirements for minimum qualifications of SERVICE PERSONNEL documented in technical description		N/A
7.9.3.2	The technical description contains the following required information		N/A
	-type and full rating of fuses used in SUPPLY MAINS external to PERMANENTLY INSTALLED ME EQUIPMENT:	Power supply unit is not permanently installed equipment.	N/A
	- a statement for ME EQUIPMENT with a non- DETACHABLE POWER SUPPLY CORD IF POWER SUPPLY CORD IS replaceable by SERVICE PERSONNEL, and	Appliance coupler provided on the EUT.	N/A
	- instructions for correct replacement of interchangeable or detachable parts specified by MANUFACTURER as replaceable by SERVICE PERSONNEL, and	No replaceable or detachable parts provided.	N/A
	RISK MANAGEMENT FILE includes an assessment to determine if replacement of components results in any unacceptable RISKS:	RMF Reference to specific RISKS: (ISO 14971 Cl.)	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	No replaceable components specified by the manufacturer. No RM considered necessary.	
	- warnings identifying nature of HAZARD when replacement of a component could result in an unacceptable RISK, and when replaceable by SERVICE PERSONNEL all information necessary to safely replace the component	No operator replaceable components.	N/A
7.9.3.3	Technical description indicates, MANUFACTURER will provide circuit diagrams, component part lists, descriptions, calibration instructions to assist to SERVICE PERSONNEL in parts repair		N/A
7.9.3.4	Means used to comply with requirements of 8.11.1 clearly identified in technical description		N/A

8	PROTECTION AGAINST ELECTRICAL HAZARDS FROM ME EQUIPMENT		Р
8.1	Limits specified in Clause 8.4 not exceeded for ACCESSIBLE PARTS and APPLIED PARTS in NORMAL or SINGLE FAULT CONDITIONS		P
	RISK MANAGEMENT FILE identifies conductors and connectors where breaking free results in a HAZARDOUS SITUATION: (ISO 14971 CI. 4.3)	RMF Reference to specific RISKS: EL7 (ISO 14971 Cl. 4.3)	P
8.2	Requirements related to power sources		N/A
8.2.1	Connection to a separate power source		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	When ME EQUIPMENT specified for connection to a separate power source other than SUPPLY MAINS, separate power source considered as part of ME EQUIPMENT or combination considered as an ME SYSTEM	Equipment is intended for connection to the supply mains. No separate power source specified.	N/A
	Tests performed with ME EQUIPMENT connected to separate power supply when one specified		N/A
	When a generic separate power supply specified, specification in ACCOMPANYING DOCUMENTS examined		N/A
8.2.2	Connection to an external d.c. power source		N/A
	No HAZARDOUS SITUATION as described in 13.1 developed when a connection with wrong polarity made for ME EQUIPMENT from an external d.c. source	Equipment is intended for connections to the AC mains.	N/A
	ME EQUIPMENT connected with correct polarity maintained BASIC SAFETY and ESSENTIAL PERFORMANCE		N/A
	Protective devices that can be reset by anyone without a TOOL returns to NORMAL CONDITION on reset		N/A
8.3	Classification of APPLIED PARTS		N/A
	a) APPLIED PART specified in ACCOMPANYING DOCUMENTS as suitable for DIRECT CARDIAC APPLICATION is TYPE CF	No Applied Parts provided	N/A
	b) An APPLIED PART provided with a PATIENT CONNECTION intended to deliver electrical energy or an electrophysiological signal to or from PATIENT IS TYPE BF OR CF APPLIED PART		N/A
	c) An APPLIED PART not covered by a) or b) is a TYPE B, BF, or CF		N/A
8.4	Limitation of voltage, current or energy		N/A
8.4.2	ACCESSIBLE PARTS and APPLIED PARTS		Р
	a) Currents from, to, or between PATIENT CONNECTIONS did not exceed limits for PATIENT LEAKAGE CURRENT & PATIENT AUXILIARY CURRENT:	No Applied Parts provided.	N/A
	ACCESSIBLE PARTS did not exceed limits for TOUCH CURRENT	See appended Table 8.7	Р
		Leakage currents measured on the secondary side outputs of the power supply.	
	c) Limits specified in b) not applied to parts when probability of a connection to a PATIENT, directly or through body of OPERATOR, is negligible in NORMAL USE, and the OPERATOR is appropriately instructed	Power supply unit is not end medical product. The likelihood of the current flowing through body of OPERATOR to be determined in end-product evaluation.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Voltage to earth or to other ACCESSIBLE PARTS did not exceed 42.4 V peak a.c. or 60 V d.c. for above parts in NORMAL or single fault condition (V a.c. or d.c.):	See appended Table 8.4.2	P
	Energy did not exceed 240 VA for longer than 60 s or stored energy available did not exceed 20 J at a potential of 2 V or more (VA or J):	See appended Table 8.4.2	Р
	d) Voltage and energy limits specified in c) above also applied to the following:		N/A
	- internal parts touchable by test pin in Fig 8 inserted through an opening in an ENCLOSURE; and	No such parts.	N/A
	- internal parts touchable by a metal test rod with a diameter of 4 mm and a length 100 mm, inserted through any opening on top of ENCLOSURE or through any opening provided for adjustment of pre-set controls by RESPONSIBLE ORGANIZATION in NORMAL USE using a TOOL	No such parts.	N/A
	Test pin or the test rod inserted through relevant openings with minimal force of no more than 1 N	No openings provided. Plastic enclosure without openings provided.	N/A
	Test rod inserted in every possible position through openings provided for adjustment of pre-set controls that can be adjusted in NORMAL USE, with a force of 10 N	No pre-set controls provided.	N/A
	Test repeated with a TOOL specified in instructions for use		N/A
	Test rod freely and vertically suspended through openings on top of ENCLOSURE	No top openings provided.	N/A
	e) Devices used to de-energize parts when an ACCESS COVER opened without a TOOL gives access to parts at voltages above levels permitted by this Clause comply with 8.11.1 for mains isolating switches and remain effective in SINGLE FAULT CONDITION		N/A
	A TOOL is required when it is possible to prevent the devices from operating		N/A
8.4.3	Worst case voltage between pins of plug and between either supply pin and ENCLOSURE did not exceed 60 V one sec after disconnecting the plug of ME EQUIPMENT or its parts (V):	See appended Table 8.4.3	N/A
	When voltage exceeded 60 V, calculated or measured stored charge didn't exceed 45 μC:	See appended Table 8.4.3	N/A
8.4.4	Residual voltage of conductive parts of capacitive circuits, having become accessible after ME EQUIPMENT was de-energized after removal of ACCESS COVERS, didn't exceed 60V or calculated stored charge didn't exceed 45µC:	See appended Table 8.4.4	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	A device manually discharging capacitors used when automatic discharging was not possible and ACCESS COVERS could be removed only with aid of a TOOL		N/A
	Capacitor(s) and connected circuitry marked with symbol 24 of Table D.1, and manual discharging device specified in technical description:		N/A
8.5	Separation of parts		Р
8.5.1	MEANS OF PROTECTION (MOP)		Р
8.5.1.1	Two MEANS of PROTECTION provided for ME EQUIPMENT to prevent APPLIED and other ACCESSIBLE PARTS from exceeding limits in 8.4	2 x MOOP provided between primary and accessible parts. 1 x MOOP provided between	Р
	Varnishing, enamelling, oxidation, and similar protective finishes and coatings with sealing compounds re-plasticizing at temperatures expected during operation and sterilization disregarded as MEANS OF PROTECTION	primary and PE.	N/A
	Components and wiring forming a MEANS OF PROTECTION comply with 8.10		Р
8.5.1.2	MEANS OF PATIENT PROTECTION (MOPP)		N/A
	Solid insulation forming a MEANS OF PATIENT PROTECTION complied with dielectric strength test:	See appended Table 8.8.3	N/A
	CREEPAGE and CLEARANCES forming a MEANS OF PATIENT PROTECTION complied with Table 12		N/A
	PROTECTIVE EARTH CONNECTIONS forming a MEANS OF PATIENT PROTECTION complied with Cl. 8.6		N/A
	Y1 or Y2 capacitor complying with standard IEC 60384-14 considered one MEANS OF PATIENT PROTECTION:	See appended Tables 8.8.3 and 8.10.	N/A
	Single Y1 capacitor used for two MEANS OF PATIENT PROTECTION when the working voltage is less than 42,4 V peak a.c. or 60 V d.c:		N/A
	Two capacitors used in series, each RATED for total WORKING VOLTAGE across the pair and have the same NOMINAL capacitance		N/A
	Voltage _{Total Working} (V) and C _{Nominal} (μF):		_
8.5.1.3	MEANS OF OPERATOR PROTECTION (MOOP)		Р
	Solid insulation forming a MEANS OF OPERATOR PROTECTION complied with:		Р
	- dielectric strength test:		Р
	- requirements of IEC 60950-1 for INSULATION CO-ORDINATION		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	CREEPAGE and CLEARANCES forming a MEANS OF OPERATOR PROTECTION complied with:		Р
	- limits of Tables 13 to 16 (inclusive); or		Р
	- requirements of IEC 60950-1 for INSULATION CO-ORDINATION		N/A
	PROTECTIVE EARTH CONNECTIONS forming a MEANS OF OPERATOR PROTECTION complied with Cl. 8.6		Р
	- or with requirements and tests of IEC 60950-1 for protective earthing:		N/A
	A Y2 (IEC 60384-14) capacitor is considered one MEANS OF OPERATOR PROTECTION:	See appended Tables 8.8.3 and 8.10	Р
		Min. Y2 capacitor connected between primary and PE.	
		See list of critical components for details.	
	A Y1 (IEC 60384-14) capacitor is considered two MEANS OF OPERATOR PROTECTION:	See appended Tables 8.8.3 and 8.10	N/A
		No bridging capacitors between primary and secondary circuit provided.	
	Two capacitors used in series each RATED for total WORKING VOLTAGE across the pair and have the same NOMINAL capacitance		N/A
	Voltage _{Total Working} (V) and C _{Nominal} (μF):		_
	Points and applied parts at which impedances of components, CREEPAGE, CLEARANCES, PROTECTIVE EARTH CONNECTIONS or insulation, prevent ACCESSIBLE PARTS from exceeding limits in 8.4 were examined whether a failure at any of these points is to be regarded as a NORMAL or SINGLE FAULT CONDITION		N/A
	A MEANS OF PROTECTION protecting APPLIED PARTS, or parts identified by 4.6 as parts subject to the same requirements, considered MEANS OF PATIENT PROTECTION:		N/A
	A MEANS OF PROTECTION protecting other parts considered MEANS OF OPERATOR PROTECTION:		N/A
.5.2	Separation of PATIENT CONNECTIONS		N/A
.5.2.1	PATIENT CONNECTIONS OF F-TYPE APPLIED PART separated from all other parts by equivalent to one MEANS OF PATIENT PROTECTION for a WORKING VOLTAGE equal to the MAX. MAINS VOLTAGE:	Component power supply, no PATIENT CONNECTIONS provided.	N/A
	Separation requirement not applied between multiple functions of a single F-TYPE APPLIED PART		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	PATIENT CONNECTIONS treated as one APPLIED PART in the absence of electrical separation between PATIENT CONNECTIONS of same or another function		N/A
	MANUFACTURER has defined if multiple functions are to be considered as all within one APPLIED PART or as multiple APPLIED PARTS:		N/A
	Classification as TYPE BF, CF, or DEFIBRILLATION-PROOF applied to one entire APPLIED PART		N/A
	LEAKAGE CURRENT tests conducted per 8.7.4:	See appended Table 8.7	N/A
	Dielectric strength test conducted per 8.8.3:	See appended Table 8.8.3	N/A
	CREEPAGE and CLEARANCES measured:	Refer to Insulation Diagram	N/A
	A protective device connected between PATIENT CONNECTIONS of an F-TYPE APPLIED PART and ENCLOSURE to protect against excessive voltages did not operate below 500 V r.m.s		N/A
8.5.2.2	PATIENT CONNECTIONS of a TYPE B APPLIED PART not PROTECTIVELY EARTHED are separated by one MEANS OF PATIENT PROTECTION from metal ACCESSIBLE PARTS not PROTECTIVELY EARTHED:		N/A
	except when metal ACCESSIBLE PART is physically close to APPLIED PART and can be regarded as a part of APPLIED PART; and		N/A
	RISK that metal ACCESSIBLE PART will make contact with a source of voltage or LEAKAGE CURRENT above permitted limits is acceptably low		N/A
	LEAKAGE CURRENT tests conducted per 8.7.4:	See appended Table 8.7	N/A
	Dielectric strength test conducted per 8.8.3:	See appended Table 8.8.3	N/A
	Relevant CREEPAGE and CLEARANCES measured		N/A
	RISK MANAGEMENT FILE includes an assessment of the RISK of metal ACCESSIBLE PARTS contacting a source of voltage or LEAKAGE CURRENT above the limits: (ISO 14971 Cl. 4.2-4.4, 5)	RMF Reference to specific RISKS: (ISO 14971 Cl)	N/A
8.5.2.3	A connector on a PATIENT lead or PATIENT cable to or cable remote from PATIENT, with conductive patient connections by one MEANS OF PATIENT PROVOLTAGE equal to MAXIMUM MAINS VOLTAGE	art not separated from all	N/A
	- cannot be connected to earth or hazardous voltage while the PATIENT CONNECTIONS are in contact with PATIENT		N/A
	- conductive part of connector not separated from all PATIENT CONNECTIONS did not come into contact with a flat conductive plate of not less than 100 mm diameter		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	- CLEARANCE between connector pins and a flat surface is at least 0.5 mm		N/A
	- conductive part pluggable into a mains socket protected from making contact with parts at MAINS VOLTAGE by insulation with a CREEPAGE DISTANCE of at least 1.0 mm, a 1500 V dielectric strength and complying with 8.8.4.1		N/A
	 required test finger did not make electrical contact with conductive part when applied against access openings with a force of 10 N, 		N/A
	Test finger test (10 N):	See appended Table 5.9.2	N/A
	Except when RISK MANAGEMENT PROCESS includes an assessment of RISKS resulting from contact with objects other than mains sockets or flat surfaces: (ISO 14971 Cl. 4.2-4.4, 5)	RMF Reference to specific RISKS: (ISO 14971 CI) See appended Table 5.9.2	N/A
8.5.4	WORKING VOLTAGE		Р
	- Input supply voltage to ME EQUIPMENT was RATED voltage or voltage within RATED range resulting in highest measured value (V):	240 Vac	P
	- WORKING VOLTAGE for d.c. voltages with superimposed ripple was average value when peak-to-peak ripple less than 10% of average value or peak voltage when peak-to-peak ripple exceeding 10% of average value (V)::		N/A
	- Working voltage for each MEANS OF PROTECTION forming DOUBLE INSULATION was voltage DOUBLE INSULATION, as a whole, subjected to (V):	See Insulation Diagram and Insulation Table	Р
	- Intentional or accidental earthing of PATIENT regarded as a NORMAL CONDITION for WORKING VOLTAGE involving a PATIENT CONNECTION not connected to earth	No APPLIED PARTS in power supply.	N/A
	- WORKING VOLTAGE between PATIENT CONNECTIONS of an F-TYPE APPLIED PART and ENCLOSURE was highest voltage appearing across insulation in NORMAL USE including earthing of any part of APPLIED PART (V)::	No APPLIED PARTS in power supply.	N/A
	- WORKING VOLTAGE for DEFIBRILLATION-PROOF APPLIED PARTS determined disregarding possible presence of defibrillation voltages	No APPLIED PARTS in power supply.	N/A
	- WORKING VOLTAGE was equal to resonance voltage in case of motors provided with capacitors between the point where a winding and a capacitor are connected together and a terminal for external conductors (V)::	No motors provided.	N/A
8.5.5	DEFIBRILLATION-PROOF APPLIED PARTS	No DEFIBRILLATION-PROOF APPLIED PARTS provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
8.5.5.1	Classification "DEFIBRILLATION-PROOF APPLIED PART" applied to one APPLIED PART in its entirety		N/A
	Isolation of PATIENT CONNECTIONS of a DEFIBRILLATION-PROOF APPLIED PART from other parts of ME EQUIPMENT accomplished as follows:		N/A
	a) No hazardous electrical energies appear during a discharge of cardiac defibrillator:	See appended Table 8.5.5.1a	N/A
	b) ME EQUIPMENT complied with relevant requirements of this standard, providing BASIC SAFETY and ESSENTIAL PERFORMANCE following exposure to defibrillation voltage, and recovery time stated in ACCOMPANYING DOCUMENTS:	See appended Table 8.5.5.1b	N/A
3.5.5.2	Means provided to limit energy delivered to a 100 Ω load:	See appended Table 8.5.5.2	N/A
3.6	Protective and functional earthing and potential	equalization of ME EQUIPMENT	Р
3.6.1	Requirements of 8.6.2 to 8.6.8 applied	EUT is Class I equipment.	Р
	Parts complying with IEC 60950-1 for protective earthing and serving as MEANS OF OPERATOR PROTECTION but not PATIENT PROTECTION exempted from requirements of 8.6.2 to 8.6.8		N/A
3.6.2	PROTECTIVE EARTH TERMINAL is suitable for connection to an external protective earthing system by a PROTECTIVE EARTH CONDUCTOR in a POWER SUPPLY CORD and a suitable plug or by a FIXED PROTECTIVE EARTH CONDUCTOR	Protective earth terminal on the power supply provided by means of approved appliance inlet.	Р
	Clamping means of PROTECTIVE EARTH TERMINAL of ME EQUIPMENT for FIXED supply conductors or POWER SUPPLY CORDS comply with 8.11.4.3, and cannot be loosened without TOOL	Approved appliance inlet used.	N/A
	Screws for internal PROTECTIVE EARTH CONNECTIONS completely covered or protected against accidental loosening from outside:		N/A
	Earth pin of APPLIANCE INLET forming supply connection to ME EQUIPMENT regarded as PROTECTIVE EARTH TERMINAL		Р
	PROTECTIVE EARTH TERMINAL not used for mechanical connection between different parts of ME EQUIPMENT or securing components not related to protective or functional earthing		N/A
3.6.3	PROTECTIVE EARTH CONNECTION not used for a moving part,	No moving part provided.	N/A
	except when MANUFACTURER demonstrated in RISK MANAGEMENT FILE connection will remain reliable during EXPECTED SERVICE LIFE	RMF Reference to proof of reliability: (ISO 14971 CI)	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	\	



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Clause	Requirement + Test	Result - Remark	Verdict
8.6.4	a) PROTECTIVE EARTH CONNECTIONS carried fault currents reliably and without excessive voltage drop:	See appended Table 8.6.4	Р
	b) Allowable TOUCH CURRENT and PATIENT LEAKAGE CURRENT in SINGLE FAULT CONDITION were not exceeded, when impedance of PROTECTIVE EARTH CONNECTIONS exceeded values in 8.6.4 a) and Table 8.6.4, due to limited current capability of relevant circuits:	See appended Table 8.6.4 & Clause 8.7 Touch current in SFC remains below 500uA.	P
8.6.5	Surface coatings		N/A
	Poorly conducting surface coatings on conductive elements removed at the point of contact		N/A
	Coating not removed when requirements for impedance and current-carrying capacity met		N/A
8.6.6	Plugs and sockets		Р
	PROTECTIVE EARTH CONNECTION where connection between SUPPLY MAINS and ME EQUIPMENT or between separate parts of ME EQUIPMENT made via a plug and socket was made before and interrupted after supply connections	Approved appliance coupler used.	P
	- applied also where interchangeable parts are PROTECTIVELY EARTHED		N/A
8.6.7	Terminal for connection of a POTENTIAL EQUALIZATION CONDUCTOR		N/A
	- Terminal is accessible to OPERATOR with ME EQUIPMENT in any position of NORMAL USE	No potential equalization terminal provided.	N/A
	-accidental disconnection avoided in NORMAL USE		N/A
	- Terminal allows conductor to be detached without a TOOL		N/A
	- Terminal not used for a PROTECTIVE EARTH CONNECTION		N/A
	- Terminal marked with symbol 8 of Table D.1		N/A
	 Instructions for use contain information on function and use of POTENTIAL EQUALIZATION CONDUCTOR together with a reference to requirements of this standard 		N/A
	Power supply cord does not incorporate a POTENTIAL EQUALIZATION CONDUCTOR		N/A
8.6.8	FUNCTIONAL EARTH TERMINAL not used to provide a PROTECTIVE EARTH CONNECTION	No functional earth terminal provided.	N/A
8.6.9	Class II ME EQUIPMENT		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Third conductor of POWER SUPPLY CORD connected to protective earth contact of MAINS PLUG provided with CLASS II ME EQUIPMENT with isolated internal screens used as functional earth connection to the screen's FUNCTIONAL EARTH TERMINAL, coloured green and yellow	Class I equipment.	N/A
	ACCOMPANYING DOCUMENTS include a statement that the third conductor in the POWER SUPPLY CORD is only a functional earth.		N/A
	Two MEANS OF PROTECTION provided between insulation of internal screens and all internal wiring connected to them and ACCESSIBLE PARTS		N/A
8.7	LEAKAGE CURRENTS and PATIENT AUXILIARY CURREN	ΓS	Р
8.7.1	a) Electrical isolation providing protection against electric shock limits currents to values in 8.7.3:	See appended Tables 8.7	Р
	b) Specified values of EARTH LEAKAGE, TOUCH, PATIENT LEAKAGE, and PATIENT AUXILIARY CURRENTS applied in combination of conditions in appended Table 8.7	See appended Tables 8.7	Р
8.7.2	Allowable values specified in 8.7.3 applied under SINGLE FAULT CONDITIONS of 8.1 b), except		Р
	 where insulation used in conjunction with a PROTECTIVE EARTH CONNECTION, insulation short circuited only under conditions in 8.6.4 b) 		N/A
	 the only SINGLE FAULT CONDITION for EARTH LEAKAGE CURRENT was interruption of one supply conductor at a time 		Р
	LEAKAGE CURRENTS and PATIENT AUXILIARY CURRENT not measured in SINGLE FAULT CONDITION of short circuiting of one constituent part of DOUBLE INSULATION		N/A
	SINGLE FAULT CONDITIONS not applied at same time as special test conditions of MAXIMUM MAINS VOLTAGE ON APPLIED PARTS and non-PROTECTIVELY EARTHED parts of ENCLOSURE	No applied parts provided.	N/A
8.7.3	Allowable Values		Р
	a) Allowable values in 8.7.3 b), c), and d)	See appended Table 8.7	Р
	measured based on, and are relative to currents in Fig 12 a), or by a device measuring frequency contents of currents as in Fig 12 b.:	Leakage current meter with frequency characteristics as specified on Fig. 12 b) used.	
	b) Allowable values of PATIENT LEAKAGE and AUXILIARY CURRENTS are according to Tables 3 & 4, and values of a.c. are relative to currents having a frequency not less than 0.1Hz	No applied parts provided.	N/A
	c) Touch current did not exceed 100 µA in NORMAL CONDITION and 500 µA in SINGLE FAULT CONDITION (I _{TNC} , I _{TSFC}):	See appended Table 8.7	Р



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Clause	Requirement + Test	Result - Remark	Verdict
	d) EARTH LEAKAGE CURRENT did not exceed 5 mA in NORMAL CONDITION and 10 mA in SINGLE FAULT CONDITION (I _{ENC} , I _{ESFC}):	See appended Table 8.7	Р
	Higher values of EARTH LEAKAGE CURRENT permitted for PERMANENTLY INSTALLED ME EQUIPMENT connected to a supply circuit supplying only this ME EQUIPMENT according to local regulations or IEC 60364-7-710:	Not permanently installed equipment.	N/A
	e) LEAKAGE CURRENTS, regardless of waveform and frequency, did not exceed 10 mA r.m.s. in NORMAL or in SINGLE FAULT CONDITION (measured with a non-frequency-weighted device:	See appended Table 8.7	N/A
	f) LEAKAGE CURRENTS flowing in a FUNCTIONAL EARTH CONDUCTOR in a non-PERMANENTLY INSTALLED ME EQUIPMENT are 5 mA in NORMAL CONDITION, 10 mA in SINGLE FAULT CONDITION:	No functional earth conductor provided.	N/A
8.7.4	LEAKAGE and PATIENT AUXILIARY CURRENTS measurements:	See appended Table 8.7	Р
8.8	Insulation		Р
8.8.1	Insulation relied on as MEANS OF PROTECTION, including REINFORCED INSULATION subjected to testing		Р
	Insulation exempted from test (complies with clause 4.8)	IEC approved optocouplers IC2 and IC6 provided between primary and secondary circuit.	Р
	Insulation forming MEANS OF OPERATOR PROTECTION and complying with IEC 60950-1 for INSULATION CO-ORDINATION not tested as in 8.8		Р
8.8.2	Distance through solid insulation or use of thin sheet material		Р
	Solid insulation forming SUPPLEMENTARY or REINFORCED INSULATION for a PEAK WORKING VOLTAGE greater than 71 V provided with:		Р
	a) 0.4 mm, min, distance through insulation, or	IEC approved optocouplers IC2 and IC6 provided between primary and secondary circuit.	Р
	b) does not form part of an ENCLOSURE and not subject to handling or abrasion during NORMAL USE, and comprised of:		Р
	- at least two layers of material, each passed the appropriate dielectric strength test:	See appended Table 8.8.3	N/A
	- or three layers of material, for which all combinations of two layers together passed the appropriate dielectric strength test:	See appended Table 8.8.3 3 layers of insulation between primary and secondary windings.	Р
	Dielectric strength test for one or two layers was same as for one MEANS OF PROTECTION for SUPPLEMENTARY INSULATION		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Dielectric strength test for one or two layers was same as for two MEANS OF PROTECTION for REINFORCED INSULATION	3.000 Vac for 2 layers of insulation tape.	Р
	BASIC, SUPPLEMENTARY, and REINFORCED INSULATION required between windings of wound components separated by interleaved insulation complying with a) or b), or both, except when		N/A
	c) Wire with solid insulation, other than solvent based enamel, complying with a)		N/A
	d) Wire with multi-layer extruded or spirally wrapped insulation complying with b) and complying with Annex L		N/A
	e) Finished wire with spirally wrapped or multi- layer extruded insulation, complying with Annex L		Р
	BASIC INSULATION: minimum two wrapped layers or one extruded layer		N/A
	- SUPPLEMENTARY INSULATION: minimum two layers, wrapped or extruded		N/A
	- REINFORCED INSULATION: minimum three layers, wrapped or extruded		N/A
	In d) and e), for spirally wrapped insulation with CREEPAGE DISTANCES between layers less than in Table 12 or 16 (Pollution Degree 1) depending on type of insulation, path between layers sealed as a cemented joint in 8.9.3.3 and test voltages of TYPE TESTS in L.3 equal 1.6 times of normal values		N/A
	Protection against mechanical stress provided where two insulated wires or one bare and one insulated wire are in contact inside wound component, crossing at an angle between 45° and 90° and subject to winding tension:	No contact between 45° and 90°.	N/A
	Finished component complied with routine dielectric strength tests of 8.8.3	See appended Table 8.8.3	N/A
	Tests of Annex L not repeated since material data sheets confirm compliance	See Table 8.10 and Material Information Attachment	N/A
3.8.3	Dielectric Strength		Р
	Solid insulating materials with a safety function withstood dielectric strength test voltages:	See appended Table 8.8.3	Р
3.8.4	Insulation other than wire insulation		Р
3.8.4.1	Resistance to heat retained by all insulation and insulating partition walls during EXPECTED SERVICE LIFE OF ME EQUIPMENT	Approved materials used.	Р
	ME EQUIPMENT and design documentation examined:	See enclosed documentation.	Р



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Clause	Requirement + Test	Result - Remark	Verdict
	RISK MANAGEMENT FILE examined in conjunction with resistance to moisture, dielectric strength, and mechanical strength tests	Approved components provided with adequate safety insulation. See appended Table 8.10. No RM considered necessary.	Р
	Satisfactory evidence of compliance provided by manufacturer for resistance to heat:		N/A
	Tests conducted in absence of satisfactory evidence for resistance to heat:		N/A
	a) ENCLOSURE and other external parts of insulating material, except insulation of flexible cords and parts of ceramic material, subjected to ball-pressure test using Fig 21 apparatus:	See appended Table 8.8.4.1	N/A
	b) Parts of insulating material supporting uninsulated parts of MAINS PART subjected to ball-pressure test in a), except at 125 °C ± 2 ° C or ambient indicated in technical description ±2°C plus temperature rise determined during test of 11.1 of relevant part, if higher (°C):	See appended Table 8.8.4.1	N/A
	Test not performed on parts of ceramic material, insulating parts of commutators, brush-caps, and similar, and on coil formers not used as REINFORCED INSULATION		N/A
8.8.4.2	Resistance to environmental stress		N/A
	Insulating characteristics and mechanical strength of all MEANS OF PROTECTION not likely to be impaired by environmental stresses including deposition of dirt resulting from wear of parts within EQUIPMENT, potentially reducing CREEPAGE and CLEARANCES below 8.9	Power supply is fully enclosed in plastic enclosure. No dust or deposition of dirt as a result of wear occurs inside the EUT.	N/A
	Ceramic and similar materials not tightly sintered, and beads alone not used as SUPPLEMENTARY OF REINFORCED INSULATION	No such materials used for insulation.	N/A
	Insulating material with embedded heating conductors considered as one MEANS OF PROTECTION but not two MEANS OF PROTECTION		N/A
	Parts of natural latex rubber aged by suspending samples freely in an oxygen cylinder containing commercial oxygen to a pressure of 2.1 MPa ± 70 kPa, with an effective capacity of at least 10 times volume of samples	Rubber not used for insulation.	N/A
	There were no cracks visible to naked eyes after samples kept in cylinder at 70 °C ± 2 °C for 96h, and afterwards, left at room temperature for at least 16h		N/A
8.9	CREEPAGE DISTANCES and AIR CLEARANCES		Р
8.9.1.1	CREEPAGE DISTANCES and AIR CLEARANCES are equal to or greater than values in Tables 12 to 16 (inclusive):	Refer to Insulation Diagram	Р



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Clause	Requirement + Test	Result - Remark	Verdict
8.9.1.15	CREEPAGE DISTANCES and AIR CLEARANCES for DEFIBRILLATION-PROOF APPLIED PARTS are 4 mm or more to meet 8.5.5.1	No DEFIBRILLATION-PROOF APPLIED PARTS provided.	N/A
8.9.2	a) Short circuiting of each single one of CREEPAGE DISTANCES and CLEARANCES in turn did not result in a HAZARDOUS SITUATION, min CREEPAGE and CLEARANCES not applied:	See appended Table 8.9.2. Sufficient creepage and clearance distances provided between parts of opposite polarity before mains fuses.	N/A
		Short circuit performed after primary fuses. No hazardous situation. See appended table 13.2.	
3.9.3	Spaces filled by insulating compound		N/A
8.9.3.1	Only solid insulation requirements applied where distances between conductive parts filled with insulating compound	No parts filled with insulation compounds.	N/A
	Thermal cycling, humidity preconditioning, and dielectric strength tests		N/A
8.9.3.2	For insulating compound forming solid insulation between conductive parts, a single sample subjected to thermal cycling PROCEDURE of 8.9.3.4 followed by humidity preconditioning per 5.7 (for 48 hours), followed by dielectric strength test (cl. 8.8.3 at 1,6 x test voltage):	See appended Table 8.9.3.2	N/A
	Cracks or voids in insulating compound affecting homogeneity of material didn't occur		N/A
3.9.3.3	Where insulating compound forms a cemented joint with other insulating parts, three samples tested for reliability of joint		N/A
	A winding of solvent-based enamelled wire replaced for the test by a metal foil or by a few turns of bare wire placed close to cemented joint, and three samples tested as follows:		N/A
	 One sample subjected to thermal cycling PROCEDURE of 8.9.3.4, and immediately after the last period at highest temperature during thermal cycling followed by dielectric strength test of cl. 8.8.3 at 1.6 x the test voltage 		N/A
	- The other two samples subjected to humidity preconditioning of 5.7, except for 48 hours only followed by a dielectric strength test of cl. 8.8.3 at 1.6 times the test voltage		N/A
3.9.4	Minimum spacing of grooves transvers to the CREEPAGE DISTANCES considered a MEANS OF OPERATOR PROTECTION adjusted based on pollution degree	Pollution degree: II	Р

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Clause	Requirement + Test	Result - Remark	Verdict
	Force was applied between bare conductors and outside metal enclosure when measuring CREEPAGE DISTANCES and AIR CLEARANCES	Refer to Insulation Diagram supplemental information for location and force used	Р
8.10	Components and wiring		Р
8.10.1	Components of ME EQUIPMENT likely to result in an unacceptable RISK by their movements mounted securely:	All components mounted securely.	N/A
	,	Unwanted movement of single component not result unacceptable risk.	
	RISK MANAGEMENT FILE includes an assessment of RISKS related to unwanted movement of components:	RMF Reference to specific RISKS: EL3	P
	(ISO 14791 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 Cl. 4.2-4.4, 5)	
8.10.2	Conductors and connectors of ME EQUIPMENT adequately secured or insulated to prevent	Accidental detachment prevented.	N/A
	accidental detachment:	Approved connectors used	
	Stranded conductors are not solder-coated when secured by clamping means to prevent HAZARDOUS SITUATIONS		N/A
8.10.3	Interconnecting flexible cords detachable without a TOOL used provided with means for connection to comply with requirements for metal ACCESSIBLE PARTS when a connection is loosened or broken:	No flexible cords provided.	N/A
8.10.4	Cord-connected HAND-HELD parts and cord-connected devices	ected foot-operated control	N/A
8.10.4.1	Control devices of ME EQUIPMENT and their connection cords contain only conductors and components operating at 42.4 V peak a.c., max, or 60 V d.c. in circuits isolated from MAINS PART by two MEANS OF PROTECTION	No cord-connected hand-held parts and cord-connected foot-operated control device provided.	N/A
8.10.4.2	Connection and anchorage of a flexible cord to a HAND-HELD or foot-operated control device of ME EQUIPMENT, at both ends of the cable to the control device, complies with the requirements for POWER SUPPLY CORDS in CI. 8.11.3		N/A
	Other HAND-HELD parts, if disturbance or breaking of one or more of the connections could result in a HAZARDOUS SITUATION, also comply with tests of Cl. 8.11.3		N/A
8.10.5	Mechanical protection of wiring		N/A
	a) Internal cables and wiring adequately protected against contact with a moving part or from friction at sharp corners and edges:	No moving parts provided.	N/A
	b) Wiring, cord forms, or components are not likely to be damaged during assembly or during opening or closing of ACCESS COVERS		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
8.10.6	Guiding rollers prevent bending of movable insulated conductors around a radius of less than five times the outer diameter of the lead		N/A
8.10.7	a) Insulating sleeve adequately secured:	See appended Table 8.10	N/A
		No insulation sleeve used.	
	b) Sheath of a flexible cord not used as a MEANS OF PROTECTION inside ME EQUIPMENT when it is subject to mechanical or thermal stresses beyond its RATED characteristics		N/A
	c) Insulated conductors of ME EQUIPMENT subject to temperatures exceeding 70 °C:	See appended Table 8.10	N/A
3.11	Mains parts, components and layout		Р
3.11.1	a) ME EQUIPMENT provided with means of	See appended Table 8.10	Р
	I MAINS SIMILITANGOLISIV ON ALL NOIGS	For means of disconnection considered appliance coupler.	
	PERMANENTLY INSTALLED ME EQUIPMENT connected to a poly-phase SUPPLY MAINS equipped with a device not interrupting neutral conductor, provided local installation conditions prevent voltage on neutral conductor from exceeding limits in 8.4.2 c)	EUT is not permanently installed poly-phase equipment.	N/A
	PERMANENTLY INSTALLED ME EQUIPMENT provided with means to isolate its circuits electrically from the SUPPLY MAINS are capable of being locked in the off position	EUT is not permanently installed equipment.	N/A
	- the isolation device specified in the ACCOMPANYING DOCUMENTS	Final determination in the end product.	N/A
	b) Means of isolation incorporated in ME EQUIPMENT, or if external, described in technical description:	See appended Table 8.10	N/A
	c) A SUPPLY MAINS switch used to comply with 8.11.1 a) complies with CREEPAGE / CLEARANCES for a MAINS TRANSIENT VOLTAGE of 4 kV:	No switch used as disconnect device.	N/A
	d) A SUPPLY MAINS switch not incorporated in a POWER SUPPLY CORD or external flexible lead		N/A
	e) Actuator of a SUPPLY MAINS switch used to comply with 8.11.1 a) complies with IEC 60447		N/A
	f) A suitable plug device used in non- PERMANENTLY INSTALLED ME EQUIPMENT with no SUPPLY MAINS SWITCH:	See appended Table 8.10	Р
	g) A fuse or a semiconductor device not used as an isolating means		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	h) ME EQUIPMENT not provided with a device causing disconnection of ME EQUIPMENT from SUPPLY MAINS by producing a short circuit resulting in operation of an overcurrent protection device		P
	i) Parts within ENCLOSURE of ME EQUIPMENT with a circuit > 42.4 V peak a.c. or 60 V d.c. that cannot be disconnected from its supply by an external switch or a plug device accessible at all times is protected against touch even after opening ENCLOSURE by an additional covering		N/A
	A clear warning notice is marked on outside of ME EQUIPMENT to indicate it exceeds allowable touch voltage		N/A
	For a part that could not be disconnected from supply by an external switch or a plug device accessible at all times, the required cover or warning notice complied with this clause		N/A
	Standard test finger applied		N/A
8.11.2	MULTIPLE SOCKET-OUTLETS integral with ME EQUIPMENT complied with 16.2 d), second dash; and 16.9.2	No MULTIPLE SOCKET-OUTLETS provided.	N/A
8.11.3	POWER SUPPLY CORDS		N/A
8.11.3.1	MAINS PLUG not fitted with more than one POWER SUPPLY CORD	EUT is power supply provided with appliance coupler. For connection to the mains power supply cord with appliance coupler shall be used. Power supply cord is not part of the investigation and is end product consideration.	N/A
8.11.3.2	Power Supply Cords are no less robust than ordinary tough rubber sheathed flexible cord (IEC 60245-1:2003, Annex A, designation 53) or ordinary polyvinyl chloride sheathed flexible cord (IEC 60227-1:1993, Annex A, design 53):	See appended Table 8.10	N/A
	Only polyvinyl chloride insulated POWER SUPPLY CORD with appropriate temperature rating used for ME EQUIPMENT having external metal parts with a temperature > 75 °C touchable by the cord in NORMAL USE:	See appended Table 8.10	N/A
8.11.3.3	NOMINAL cross-sectional area of conductors of POWER SUPPLY CORDS of ME EQUIPMENT is not less than in Table 17:		N/A
8.11.3.4	APPLIANCE COUPLERS complying with IEC 60320- 1 are considered to comply with 8.11.3.5 and 8.11.3.6		N/A
8.11.3.5	Cord anchorage		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	a) Conductors of POWER SUPPLY CORD provided with strain relief and insulation protected from abrasion at point of entry to ME EQUIPMENT or a MAINS CONNECTOR by a cord anchorage	No power supply cord provided.	N/A
	b) Cord anchorage of POWER SUPPLY CORD is an insulating material, or		N/A
	- metal, insulated from conductive ACCESSIBLE PARTS non-PROTECTIVELY EARTHED by a MEANS OF PROTECTION, or		N/A
	- metal provided with an insulating lining affixed to cord anchorage		N/A
	c) Cord anchorage prevents cord from being clamped by a screw bearing directly on cord insulation		N/A
	d) Screws to be operated when replacing POWER SUPPLY CORD do not serve to secure any components		N/A
	e) Conductors of POWER SUPPLY CORD arranged to prevent PROTECTIVE EARTH CONDUCTOR against strain as long as phase conductors are in contact with their terminals		N/A
	f) Cord anchorage prevents POWER SUPPLY CORD from being pushed into ME EQUIPMENT OR MAINS CONNECTOR		N/A
	Conductors of POWER SUPPLY CORD supplied by MANUFACTURER disconnected from terminals or from MAINS CONNECTOR and cord subjected 25 times to a pull applied with no jerks, each time for 1 s, on sheath of the value in Table 18:	See appended Table 8.11.3.5	N/A
	Cord subjected to a torque in Table 18 for one minute immediately after pull tests		N/A
	Cord anchorage did not allow cord sheath to be longitudinally displaced by more than 2 mm or conductor ends to move over a distance of more than 1 mm from their connected position		N/A
	CREEPAGE and CLEARANCES not reduced below limits in 8.9		N/A
	It was not possible to push the cord into ME EQUIPMENT OR MAINS CONNECTOR to an extent the cord or internal parts would be damaged		N/A
8.11.3.6	Power supply cords protected against excessive bending at inlet opening of equipment	Power supply cord not part of investigation (for C8 plug).	N/A
	Cord guard complied with test of IEC 60335-1:2001, Clause 25.14, or		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	ME EQUIPMENT placed such that axis of cord guard projected at an angle of 45° with cord free from stress, and a mass equal 10 x D ² gram attached to the free end of cord (g):	See appended Table 8.11.3.6	N/A
	Cord guard of temperature-sensitive material tested at 23 °C ± 2 °C, and flat cords bent in the plane of least resistance		N/A
	Curvature of the cord radius, immediately after mass attached, was not less than 1.5 x D:	See appended Table 8.11.3.6	N/A
8.11.4	MAINS TERMINAL DEVICES		N/A
8.11.4.1	PERMANENTLY INSTALLED and ME EQUIPMENT with non-DETACHABLE POWER SUPPLY CORD provided with MAINS TERMINAL DEVICES ensuring reliable connection	Approved appliance inlet used for connection to the mains.	N/A
	Terminals alone are not used to keep conductors in position		N/A
	Terminals of components other than terminal blocks complying with requirements of this Clause and marked accordingly used as terminals intended for external conductors		N/A
	Screws and nuts clamping external conductors do not serve to secure any other component		N/A
8.11.4.2	Arrangement of MAINS TERMINAL DEVICES		N/A
	a) Terminals provided for connection of external cords or POWER SUPPLY CORDS together with PROTECTIVE EARTH TERMINAL grouped to provide convenient means of connection		N/A
	d) Mains Terminal Devices not accessible without use of a TOOL		N/A
	e) A MEANS OF PROTECTION are not short circuited when one end of a flexible conductor with NOMINAL cross-sectional area is stripped 8 mm and a single free wire is bent in each possible direction		N/A
8.11.4.3	Internal wiring not subjected to stress and CREEPAGE and CLEARANCES not reduced after fastening and loosening a conductor of largest cross-sectional area 10 times		N/A
8.11.4.4	Terminals with clamping means for a rewireable flexible cord did not require special preparation of conductors and conductors were not damaged and did not slip out when clamping means tightened		N/A
8.11.4.5	Adequate space provided inside ME EQUIPMENT designed for FIXED wiring or a rewireable POWER SUPPLY CORD to allow for connection of conductors		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Correct connection and positioning of conductors before ACCESS COVER verified by an installation test		N/A
8.11.5	Mains fuses and OVER-CURRENT RELEASES		Р
	A fuse or OVER-CURRENT RELEASE provided in each supply lead for CLASS I and CLASS II ME EQUIPMENT with a functional earth connection:	See appended Table 8.10 Two primary fuses used (in line and neutral).	Р
		EUT is Class I equipment.	
	- in at least one supply lead for other single- phase CLASS II ME EQUIPMENT:	See appended table 8.10. Two primary fuses used (in line and neutral).	Р
		EUT is Class I equipment.	
		Short circuit tests performed and when components were failing, the test was repeated two times.	
	- neutral conductor not fused for PERMANENTLY INSTALLED ME EQUIPMENT	EUT is not permanently installed equipment.	N/A
	- fuses or OVER-CURRENT RELEASES omitted due to provision of two MEANS OF PROTECTION between all parts within MAINS PART		N/A
	Protective devices have adequate breaking capacity to interrupt the max. fault current:	See appended Table 8.10	Р
	A fuse or OVER-CURRENT RELEASE not provided in a PROTECTIVE EARTH CONDUCTOR	Not fused.	Р
	Justification for omission of fuses or OVER-CURRENT RELEASES documented:		N/A
3.11.6	Internal wiring of the MAINS PART		Р
	a) Cross-sectional area of internal wiring in a MAINS PART between MAINS TERMINAL DEVICE or APPLIANCE INLET and protective devices suitable		N/A
	b) Cross-sectional area of other wiring in MAINS PART and sizes of tracks on printed wiring circuits are sufficient:	See appended Table 8.10	Р

9	PROTECTION AGAINST MECHANICAL HAZARDS OF ME EQUIPMENT AND ME SYSTEMS		Р
9.2	HAZARDS associated with moving parts		N/A
9.2.1	When ME EQUIPMENT with moving parts PROPERLY INSTALLED, used per ACCOMPANYING DOCUMENTS or under foreseeable misuse, RISKS associated with moving parts reduced to an acceptable level:		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	RISK from contact with moving parts reduced to an acceptable level using protective measures, (access, function, shape of parts, energy, speed of motion, and benefits to PATIENT considered)		N/A
	RESIDUAL RISK associated with moving parts considered acceptable when exposure was needed for ME EQUIPMENT to perform its intended function, and		N/A
	RISK CONTROLS implemented:		N/A
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with moving parts:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
	All RISKS associated with moving parts have been reduced to an acceptable level		N/A
9.2.2	TRAPPING ZONE		N/A
9.2.2.1	ME EQUIPMENT with a TRAPPING ZONE complied with one or more of the following as feasible:	No trapping zones.	N/A
	- Gaps in Clause 9.2.2.2, or		N/A
	- Safe distances in Clause 9.2.2.3, or		N/A
	- GUARDS and other RISK CONTROL measures in 9.2.2.4, or		N/A
	- Continuous activation in Clause 9.2.2.5		N/A
	Control of relevant motion complied with 9.2.2.6 when implementation of above protective measures were inconsistent with INTENDED USE of ME EQUIPMENT OF ME SYSTEM		N/A
9.2.2.2	A TRAPPING ZONE considered not to present a MECHANICAL HAZARD when gaps of TRAPPING ZONE complied with dimensions per Table 20:	See appended Table 9.2.2.2	N/A
9.2.2.3	A TRAPPING ZONE considered not to present a MECHANICAL HAZARD when distances separating OPERATOR, PATIENT, and others from TRAPPING ZONES exceeded values in ISO 13857:2008:	See appended Table 9.2.2.2	N/A
9.2.2.4	GUARDS and other RISK CONTROL measures		N/A
9.2.2.4.1	A TRAPPING ZONE do not to present a MECHANICAL HAZARD when GUARDS or other RISK CONTROL measures are of robust construction, not easy to bypass or render non-operational, and did not introduce additional unacceptable RISK:	No trapping zones.	N/A
9.2.2.4.2	FIXED GUARDS held in place by systems that can only be dismantled with a TOOL		N/A
9.2.2.4.3	Movable GUARDS that can be opened without a TOOL remained attached when GUARD was open		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	- they are associated with an interlock preventing relevant moving parts from starting to move while TRAPPING ZONE is accessible, and stops movement when the GUARD is opened,		N/A
	- absence or failure of one of their components prevents starting, and stops moving parts		N/A
	Movable GUARDS complied with any applicable tests		N/A
9.2.2.4.4	Other RISK CONTROL designed and incorporated into to the control system stops movement and		N/A
	- SINGLE FAULT CONDITIONS have a second RISK CONTROL, or		N/A
	ME EQUIPMENT IS SINGLE FAULT SAFE		N/A
9.2.2.5	Continuous activation		N/A
	Continuous activation used as a RISK CONTROL, complies with the following		N/A
	a) movement was in OPERATOR'S field of view		N/A
	b) movement of ME EQUIPMENT or its parts was possible only by continuous activation of control by OPERATOR		N/A
	c) a second RISK CONTROL provided for SINGLE FAULT CONDITION of continuous activation system, or		N/A
	- the continuous activation system is SINGLE FAULT SAFE		N/A
9.2.2.6	Speed of movement(s) positioning parts of ME EQUIPMENT or PATIENT limited to allow OPERATOR control of the movement		N/A
	Over travel of such movement occurring after operation of a control to stop movement, did not result in an unacceptable RISK		N/A
9.2.3	Other MECHANICAL HAZARDS associated with moving	ng parts	N/A
9.2.3.1	Controls positioned, recessed, or protected by other means so that they cannot be accidentally actuated	No moving parts provided.	N/A
	- unless for the intended PATIENT, the USABILITY ENGINEERING PROCESS concludes otherwise (e.g. PATIENT with special needs), or		N/A
	- activation does not result in an unacceptable		N/A
9.2.3.2	Over travel past range limits of the ME EQUIPMENT prevented:		N/A
	Over travel means provided with mechanical strength to withstand loading in NORMAL CONDITION & reasonably foreseeable misuse:		N/A



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Clause	Requirement + Test	Result - Remark	Verdict

9.2.4	Emergency stopping devices		N/A
	Where necessary to have one or more emergency stopping device(s), emergency stopping device complied with all the following, except for actuating switch capable of interrupting all power:	No emergency stopping device provided.	N/A
	a) Emergency stopping device reduced RISK to an acceptable level		N/A
	RISK MANAGEMENT FILE indicates the use of an emergency stopping device reduces the RISK to an acceptable level: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.6)	RMF Reference to specific RISKS: (ISO 14971 CI) No emergency stopping device provided. No RM considered necessary.	N/A
	b) Proximity and response of OPERATOR to actuate emergency stopping device could be relied upon to prevent HARM		N/A
	c) Emergency stopping device actuator was readily accessible to OPERATOR		N/A
	d) Emergency stopping device(s) are not part of normal operation of ME EQUIPMENT		N/A
	e) Emergency switching operation or stopping means neither introduced further HAZARD nor interfered with operation necessary to remove original MECHANICAL HAZARD		N/A
	f) Emergency stopping device was able to break full load of relevant circuit, including possible stalled motor currents and the like		N/A
	g) Means for stopping of movements operate as a result of one single action		N/A
	h) Emergency stopping device provided with an actuator in red and easily distinguishable and identifiable from other controls		N/A
	i) An actuator interrupting/opening mechanical movements marked on or immediately adjacent to face of actuator with symbol 18 of Table D.1 or "STOP"		N/A
	j) Emergency stopping device, once actuated, maintained ME EQUIPMENT in disabled condition until a deliberate action, different from that used to actuate it, was performed		N/A
	k) Emergency stopping device is suitable for its application		N/A
9.2.5	Means provided to permit quick and safe release of PATIENT in event of breakdown of ME EQUIPMENT or failure of power supply, activation of a RISK CONTROL measure, or emergency stopping:	Equipment does not require fixation of patient.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	- and uncontrolled or unintended movement of ME EQUIPMENT that could result in an unacceptable RISK prevented		N/A
	- Situations where PATIENT is subjected to unacceptable RISKS due to proximity of moving parts, removal of normal exit routes, or other HAZARDS prevented		N/A
	- Measures provided to reduce RISK to an acceptable level when after removal of counterbalanced parts, other parts of ME EQUIPMENT can move in a hazardous way		N/A
	RISK MANAGEMENT FILE includes an assessment of RISKS to the PATIENT related to breakdown of the ME EQUIPMENT	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI) No moving parts provided. Equipment does not require fixation of patient. No RM considered necessary.	
9.3	Rough surfaces, sharp corners and edges of ME EQUIPMENT that could result in injury or damage avoided or covered:	No rough surfaces, no sharp corners and no sharp edges.	Р
9.4	Instability HAZARDS		Р
9.4.1	ME EQUIPMENT and its parts, other than FIXED, for placement on a surface did not overbalance (tip over) or move unexpectedly in NORMAL USE	Not overbalanced when tilted through an angle of 10°. Power supply unit is desk-top equipment.	Р
9.4.2	Instability – overbalance		P
9.4.2.1	ME EQUIPMENT or its parts did not overbalance	See appended Table 9.4.2.1	Р
	when prepared per ACCOMPANYING DOCUMENTS,	Not overbalanced when tilted through an angle of 10°.	
		Power supply unit is desk-top equipment.	
9.4.2.2	Instability excluding transport		N/A
	ME EQUIPMENT or its did not overbalance when placed in different positions of NORMAL USE,:	See appended Table 9.4.2.2	N/A
	A warning provided when overbalance occurred during 10° inclined plane test		N/A
9.4.2.3	Instability from horizontal and vertical forces		N/A
	a) ME EQUIPMENT or its parts with a mass of 25kg or more, intended to be used on the floor, didn't overbalance due to pushing, leaning against it		N/A
	Surfaces of ME EQUIPMENT or its parts where a RISK of overbalancing exists from pushing, etc., permanently marked with a warning of the RISK		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	ME EQUIPMENT did not overbalance when tested according to Cl. 9.4.2.3 a)	See appended Table 9.4.2.3	N/A
	b) ME EQUIPMENT, for use on the floor or on a table, did not overbalance due to sitting or stepping		N/A
	ME EQUIPMENT or its parts, for use on the floor or on a table, where RISK of overbalancing exists, permanently marked with the RISK warning:		N/A
	ME EQUIPMENT did not overbalance when tested according to Cl. 9.4.2.3b):	See appended Table 9.4.2.3	N/A
9.4.2.4	Castors and wheels		N/A
9.4.2.4.1	Means used for transportation of MOBILE ME EQUIPMENT did not result in an unacceptable RISK when MOBILE ME EQUIPMENT moved or parked in NORMAL USE	Not mobile equipment.	N/A
9.4.2.4.2	Force required to move MOBILE ME EQUIPMENT did not exceed 200 N:	Not mobile equipment.	N/A
9.4.2.4.3	MOBILE ME EQUIPMENT exceeding 45 kg able to pass over threshold:	Not mobile equipment.	N/A
9.4.3	Instability from unwanted lateral movement (incl	uding sliding)	N/A
9.4.3.1	a) Brakes of power-driven MOBILE ME EQUIPMENT normally activated and could only be released by continuous actuation of a control	Not mobile equipment.	N/A
	b) Mobile ME Equipment provided with locking means to prevent unwanted movements		N/A
	c) No unwanted lateral movement resulted when MOBILE ME EQUIPMENT placed in its transport position when test per 9.4.3.1		N/A
9.4.3.2	Instability excluding transport		N/A
	a) MOBILE ME EQUIPMENT provided with wheel locks or braking system compliant with 5° tilt test	Not mobile equipment.	N/A
	b) MOBILE ME EQUIPMENT provided with wheel locks or braking system compliant with lateral stability test		N/A
9.4.4	Grips and other handling devices		N/A
	a) ME EQUIPMENT with a mass of over 20 kg requiring lifting in NORMAL USE or transport provided with suitable handling means, or ACCOMPANYING DOCUMENTS specify safe lifting method		N/A
	Handles, suitably placed to enable ME EQUIPMENT or its part to be carried by two or more persons and by examination of EQUIPMENT, its part, or ACCOMPANYING DOCUMENTS		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	b) PORTABLE ME EQUIPMENT with a mass > 20 kg provided with one or more carrying-handles suitably placed to enable carrying by two or more persons as confirmed by actual carrying		N/A
	c) Carrying handles and grips and their means of attachment withstood loading test:		N/A
9.5	Expelled parts HAZARD		N/A
9.5.1	Suitability of means of protecting against expelled parts determined by assessment and examination of RISK MANAGEMENT FILE:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.3, 4.4, 5, 6.2-6.5)	(ISO 14971 CI) No risk of expelled parts. No RM considered necessary.	
	All identified RISKS associated with expelled parts mitigated to an acceptable level		N/A
9.5.2	Cathode Ray tube(s) complied with IEC 60065:2001, Clause 18, or IEC 61965:	No cathode ray tubes.	N/A
9.6	Acoustic energy (including infra- and ultrasound	l) and vibration	N/A
9.6.1	Human exposure to acoustic energy and vibration from ME EQUIPMENT doesn't result in unacceptable RISK and	Equipment does not produce significant acoustic energy or vibration.	N/A
	If necessary, confirmed in RISK MANAGEMENT FILE including audibility of auditory alarm signals, and PATIENT sensitivity:		N/A
	If necessary, confirmed in RISK MANAGEMENT FILE including audibility of auditory alarm signals, PATIENT sensitivity, and	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-44, 5, 6.2-6.5)	(ISO 14971 Cl)	
	(130 1437 1 31. 4.2-44, 3, 0.2-3.3)	Equipment does not produce significant acoustic energy or vibration. No RM considered necessary.	
	All identified RISKS mitigated to an acceptable level		N/A
9.6.2	Acoustic energy		N/A
9.6.2.1	PATIENT, OPERATOR, and other persons are not exposed to acoustic energy from ME EQUIPMENT in NORMAL USE	Equipment does not produce significant acoustic energy.	N/A
	- 80 dBA for a cumulative exposure of 24 h over a 24 h period (dBA):		_
	- 83 dBA (when halving the cumulative exposure time) (dBA):		_
	- 140 dBC (peak) sound pressure level for impulsive or impact acoustic energy (dB):		_



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Clause	Requirement + Test	Result - Remark	Verdict
9.6.2.2	RISK MANAGEMENT FILE examined: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	RMF Reference to specific RISKS: (ISO 14971 CI) Equipment does not produce infrasound and ultrasound.	N/A
9.6.3	Hand-transmitted vibration		N/A
	Means provided to protect PATIENT and OPERATOR when hand-transmitted frequency-weighted r.m.s. acceleration generated in NORMAL USE exceeds specified values	Equipment does not produce vibration.	N/A
	 2.5 m/s² for a cumulative time of 8 h during a 24 h period (m/s²) 		N/A
	 Accelerations for different times, inversely proportional to square root of time (m/s²): 		N/A
).7	Pressure vessels and parts subject to pneumation	c and hydraulic pressure	N/A
9.7.2	Pneumatic and hydraulic parts of ME EQUIPMENT or ACCESSORIES met requirements based on examination of RISK MANAGEMENT FILE	RMF Reference to specific RISKS: (ISO 14971 Cl) No pneumatic and hydraulic parts within the equipment.	N/A
	- No unacceptable RISK resulted from loss of pressure or loss of vacuum		N/A
	- No unacceptable RISK resulted from a fluid jet caused by leakage or a component failure		N/A
	- Elements of ME EQUIPMENT or an ACCESSORY, especially pipes and hoses leading to an unacceptable RISK protected against harmful external effects		N/A
	- Reservoirs and similar vessels leading to an unacceptable RISK are automatically depressurized when ME EQUIPMENT is isolated from its power supply		N/A
	Means provided for isolation, or local depressurizing reservoirs and similar vessels, and pressure indication when above not possible		N/A
	- All elements remaining under pressure after isolation of ME EQUIPMENT or an ACCESSORY from its power supply resulting in an unacceptable RISK provided with clearly identified exhaust devices, and a warning to depressurize these elements before setting or maintenance activity		N/A
9.7.3	Maximum pressure a part of ME EQUIPMENT can be subjected to in NORMAL and SINGLE FAULT CONDITIONS considered to be highest of following:		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	a) RATED maximum supply pressure from an external source		N/A
	b) Pressure setting of a pressure-relief device provided as part of assembly		N/A
	c) Max pressure that can develop by a source of pressure that is part of assembly, unless pressure limited by a pressure-relief device		N/A
9.7.4	Max pressure in NORMAL and SINGLE FAULT CONDITIONS did not exceed MAXIMUM PERMISSIBLE WORKING PRESSURE for EQUIPMENT part, except as allowed in 9.7.7, confirmed by inspection of THE MANUFACTURER'S data for the component, ME EQUIPMENT, and by functional tests::		N/A
9.7.5	A pressure vessel withstood a HYDRAULIC TEST PRESSURE when pressure was more than 50 kPa, and product of pressure and volume was more than 200 kPal:		N/A
9.7.6	Pressure-control device regulating pressure in ME EQUIPMENT with pressure-relief device completed 100,000 cycles of operation under RATED load and prevented pressure from exceeding 90 % of setting of pressure-relief device in different conditions of NORMAL USE .:		N/A
9.7.7	Pressure-relief device(s) used where MAXIMUM PERMISSIBLE WORKING PRESSURE could otherwise be exceeded met the following, as confirmed by MANUFACTURER'S data, ME EQUIPMENT, RISK MANAGEMENT FILE, and functional tests::		N/A
	a) Connected as close as possible to pressure vessel or parts of system it is to protect		N/A
	b) Installed to be readily accessible for inspection, maintenance, and repair		N/A
	c) Could be adjusted or rendered inoperative without a TOOL		N/A
	d) With discharge opening located and directed as to not to release material towards any person		N/A
	e) With discharge opening located and directed as to not to deposit material on parts that could result in an unacceptable RISK		N/A
	f) Adequate discharge capacity provided to ensure that pressure will not exceed MAXIMUM PERMISSIBLE WORKING PRESSURE of system it is connected to by more than 10 % when failure occurs in control of supply pressure		N/A
	g) No shut-off valve provided between a pressure-relief device and parts it is to protect		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	h) Min number of cycles of operation 100 000, except for one-time use devices (bursting disks)		N/A
	RISK MANAGEMENT FILE includes an assessment of the risks associated with the discharge opening of the pressure relief device: (ISO 14971 Cl. 4.3, 4.4, 5, 6.2-6.5)	RMF Reference to specific RISKS: (ISO 14971 CI) No pressure-relief devices provided. No RM considered necessary.	N/A
9.8	HAZARDS associated with support systems		N/A
9.8.1	ME EQUIPMENT parts designed to support loads or provide actuating forces when a mechanical fault could constitute an unacceptable RISK:	No support system provided.	N/A
	Construction of support, suspension, or actuation system complied with Table 21 and TOTAL LOAD		N/A
	 Means of attachment of ACCESSORIES prevent possibility of incorrect attachment that could result in an unacceptable RISK 		N/A
	- RISK ANALYSIS of support systems included MECHANICAL HAZARDS from static, dynamic, vibration, foundation and other movements, impact and pressure loading, temperature, environmental, manufacture and service conditions:	RMF Reference to specific RISKS: (ISO 14971 CI) No support system provided.	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5) RISK ANALYSIS included effects of failures such as excessive deflection, plastic deformation, ductile/brittle fracture, fatigue fracture, instability (buckling), stress-assisted corrosion cracking, wear, material creep and deterioration, and residual stresses from manufacturing PROCESSES		N/A
	- Instructions on attachment of structures to a floor, wall, ceiling, included in ACCOMPANYING DOCUMENTS making adequate allowances for quality of materials used to make the connection and list the required materials		N/A
	Additional instructions provided on checking adequacy of surface of structure parts will be attached to		N/A
9.8.2	Support systems maintain structural integrity during EXPECTED SERVICE LIFE, and TENSILE SAFETY FACTORS are not less than in Table 21, except when an alternative method used to demonstrate structural integrity throughout EXPECTED SERVICE LIFE, or for a foot rest	No support system provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Compliance with 9.8.1 and 9.8.2 confirmed by examination of ME EQUIPMENT, RISK MANAGEMENT FILE, specifications and material processing:		N/A
	RISK MANAGEMENT FILE includes an assessment of the structural integrity of support system:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.3-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		No support system provided.	
	All identified RISKS are mitigated to an acceptable level		N/A
	When test were conducted, testing consisted of application of a test load to support assembly equal to TOTAL LOAD times required TENSILE SAFETY FACTOR while support assembly under test was in equilibrium after 1 min, or not resulted in an unacceptable RISK::		N/A
	Where the equipment is not at equilibrium after 1 min, the RISK MANAGEMENT FILE includes an assessment of the test results: (ISO 14971 Cl. 4.3-4.4, 5, 6.2-6.5)	RMF Reference to specific RISK: (ISO 14971 CI) No support system provided.	N/A
9.8.3	Strength of PATIENT or OPERATOR support or susp		N/A
9.8.3.1	ME EQUIPMENT parts supporting or immobilizing PATIENTS presents no unacceptable RISK of physical injuries and accidental loosening of secured joints		N/A
	RISK MANAGEMENT FILE includes assessment of the RISKS associated with physical injuries and accidental loosening of fixings:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
	(130 1437 1 01. 4.2-4.4, 3, 0.2-0.3)	No support system provided.	
	SAFE WORKING LOAD of ME EQUIPMENT or its parts supporting or suspending PATIENTS or OPERATORS is sum of mass of PATIENTS or mass of OPERATORS plus mass of ACCESSORIES supported by ME EQUIPMENT or its parts		N/A
	Supporting and suspending parts for adult human PATIENTS or OPERATORS designed for a PATIENT or OPERATOR with a min mass of 135 kg and ACCESSORIES with a min mass of 15 kg, unless stated by MANUFACTURER		N/A
	Maximum mass of PATIENT included in SAFE WORKING LOAD of ME EQUIPMENT or its parts supporting or suspending PATIENTS adapted when MANUFACTURER specified applications		N/A
	Max allowable PATIENT mass < 135 kg marked on ME EQUIPMENT and stated in ACCOMPANYING DOCUMENTS		N/A



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Clause Re	equirement + Test	Result - Remark	Verdict

	Max allowable PATIENT mass over 135 kg stated in ACCOMPANYING DOCUMENTS	 N/A
	Examination of markings, ACCOMPANYING DOCUMENTS, and RISK MANAGEMENT FILE confirmed compliance:	 N/A
9.8.3.2	a) Entire mass of PATIENT OR OPERATOR distributed over an area of 0.1 m² on a foot rest temporarily supporting a standing PATIENT OR OPERATOR:	 N/A
	Compliance confirmed by examination of ME EQUIPMENT specifications of materials and their processing, and tests:	 N/A
	b) Deflection of a support surface from PATIENT or OPERATOR loading on an area of support/ suspension where a PATIENT or OPERATOR can sit did not result in an unacceptable RISK	 N/A
	Compliance confirmed by examination of ME EQUIPMENT, specifications of materials and their processing, and by a test:	 N/A
9.8.3.3	Dynamic forces that can be exerted on equipment parts supporting or suspending a PATIENT OF OPERATOR IN NORMAL USE maintained BASIC SAFETY and ESSENTIAL PERFORMANCE confirmed test	 N/A
9.8.4	Systems with MECHANICAL PROTECTIVE DEVICES	N/A
9.8.4.1	a) A MECHANICAL PROTECTIVE DEVICE provided for the support system	 N/A
	b) MECHANICAL PROTECTIVE complies with the requirements as follows:	 N/A
	- Designed based on TOTAL LOAD	 N/A
	 Has TENSILE SAFETY FACTORS for all parts not less than Table 21, row 7 	 N/A
	- Activated before travel produced an unacceptable RISK	 N/A
	- Takes into account Clauses 9.2.5 and 9.8.4.3	 N/A
	Compliance confirmed by examination of ME EQUIPMENT over travel calculations and evaluation plus functional tests:	 N/A
9.8.4.2	Activation of MECHANICAL PROTECTIVE DEVICE is made obvious to OPERATOR when ME EQUIPMENT can still be used after failure of suspension or actuation means and activation of a MECHANICAL PROTECTIVE DEVICE	 N/A
	MECHANICAL PROTECTIVE DEVICE requires use of a TOOL to be reset or replaced	 N/A



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Clause	Requirement + Test	Result - Remark	Verdict		
	-use of ME EQUIPMENT not possible until replacement of MECHANICAL PROTECTIVE DEVICE :		N/A		
	- ACCOMPANYING DOCUMENTS provided with required information on replacement by service personal		N/A		
	ME EQUIPMENT permanently marked with safety sign 2 of Table D.		N/A		
	- Marking is adjacent to MECHANICAL PROTECTIVE DEVICE		N/A		
	Compliance confirmed by examination and following test: :		N/A		
	A chain, cable, band, spring, belt, jack screw nut, pneumatic or hydraulic hose, structural part or the like, employed to support a load, defeated by a convenient means causing maximum normal load to fall from most adverse position permitted by construction of ME EQUIPMENT		N/A		
	Load included SAFE WORKING LOAD in 9.8.3.1 when system was capable of supporting a PATIENT OF OPERATOR		N/A		
	No evidence of damage to MECHANICAL PROTECTIVE DEVICE affecting its ability to perform its intended function		N/A		
).8.5	Systems without MECHANICAL PROTECTIVE DEVICES		N/A		
	Support Systems does not require MECHANICAL PROTECTIVE DEVICES:		N/A		
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with wear on the support system:	RMF Reference to specific RISKS:	N/A		
	(ISO 14971 Cl. 4.3,4.4,5,6.2-6.5)	(ISO 14971 Cl) No support system provided.			

10	PROTECTION AGAINST UNWANTED AND EXCESSIVE RADIATION HAZARDS		N/A
10.1	X-Radiation		N/A
10.1.1	The air kerma did not exceed 5 µGy/hat 5 cm from surface of ME EQUIPMENT	Equipment does not produce X-radiations.	N/A
	Annual exposure reduced taking into account the irradiated body part, national regulations, and/or international recommendations for ME EQUIPMENT that has permanent proximity to a PATIENT as part of the INTENDED USE		N/A
10.1.2	RISK from unintended X-radiation from ME EQUIPMENT producing X-radiation for diagnostic and therapeutic purposes addressed application of applicable particular and collateral standards, or		N/A



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Clause	Requirement + Test	Result - Remark	Verdict

	RISK MANAGEMENT PROCESS as indicated in RISK MANAGEMENT FILE	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		Equipment does not produce diagnostic or therapeutic X-radiation.	
10.2	RISK associated with alpha, beta, gamma, neutron, and other particle radiation,	RMF Reference to specific RISKS:	N/A
	addressed in RISK MANAGEMENT PROCESS as shown in RISK MANAGEMENT FILE:	(ISO 14971 CI)	
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	Equipment does not produce alpha, beta, gamma, neutron and other radiation.	
10.3	The power density of unintended microwave radiation at frequencies between 1 GHz and 100 GHz does not exceed 10 W/m2	Equipment does not produce microwave radiation.	N/A
	Microwave radiation is propagated intentionally		N/A
10.4	Relevant requirements of IEC 60825-1:2007 applied to lasers, laser light barriers or similar with a wavelength range of 180nm to 1 mm.	No such components incorporated within the equipment.	N/A
10.5	RISK associated with visible electromagnetic radiation other than emitted by lasers and LEDS,	RMF Reference to specific RISKS:	N/A
	when applicable, addressed in RISK MANAGEMENT PROCESS as indicated in RISK	(ISO 14971 CI)	
	MANAGEMENT FILE:: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	Equipment does not produce other visible electromagnetic radiation.	
10.6	RISK associated with infrared radiation other than emitted by lasers and LEDS addressed in	RMF Reference to specific RISKS:	N/A
	RISK MANAGEMENT PROCESS as indicated in RISK MANAGEMENT FILE:	(ISO 14971 CI)	
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	Equipment does not produce infrared radiation.	
10.7	RISK associated with ultraviolet radiation other than emitted by lasers and LEDS addressed in	RMF Reference to specific RISKS:	N/A
	RISK MANAGEMENT PROCESS as indicated in RISK MANAGEMENT FILE:	(ISO 14971 CI)	
	(ISO 14971 CI. 4.2-4.4, 5, 6.2-6.5)	Equipment does not produce ultraviolet radiation.	

11	PROTECTION AGAINST EXCESSIVE TEMPERATURES AND OTHER HAZARDS		Р
11.1	Excessive temperatures in ME EQUIPMENT		P
11.1.1	Temperatures on ME EQUIPMENT parts did not exceed values in Tables 22 and:	See appended Table 11.1.1	Р
	Surfaces of test corner did not exceed 90 °C	Temperature of test corner did not exceed 90°C in normal conditions.	Р



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Clause	Requirement + Test	Result - Remark	Verdict
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	THERMAL CUT-OUTS did not operate in NORMAL CONDITION	No thermal cut-outs provided.	N/A
	RISK MANAGEMENT FILE includes an assessment of the duration of contact for all APPLIED PARTS	RMF Reference to specific RISK: H2	Р
	and ACCESSIBLE PARTS: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	
11.1.2	Temperature of APPLIED PARTS		N/A
11.1.2.1	APPLIED PARTS (hot or cold intended to supply heat to a PATIENT comply:	No APPLIED PARTS provided.	N/A
	Clinical effects determined and documented in the RISK MANAGEMENT FILE	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
	Temperature (hot or cold) of APPLIED PARTS intended to supply heat to a PATIENT disclosed in the instructions for use		N/A
11.1.2.2	APPLIED PARTS not intended to supply heat to a PATIENT complies with the limits of Table 24 in NORMAL CONDITION and SINGLE FAULT CONDITION:	No APPLIED PARTS provided.	N/A
	APPLIED PARTS surface temperature exceeds 41°C disclosed in the instruction manual:		N/A
	Maximum Temperature:		_
	Conditions for safe contact, e.g. duration or condition of the PATIENT:		_
	Clinical effects with respect to characteristics taken or surface pressure documented in the	RMF Reference to specific RISKS:	N/A
	RISK MANAGEMENT FILE (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
	APPLIED PARTS surface temperature of equal to or less than 41°C		N/A
	Analysis documented in the RISK MANAGEMENT FILE show that APPLIED PART temperatures are not affected by operation of the ME EQUIPMENT including SINGLE FAULT CONDITIONS. Measurement of APPLIED PART temperature according to 11.1.3 is not conducted		N/A
	Surfaces of APPLIED PARTS that are cooled below ambient temperatures evaluated in the RISK MANAGEMENT PROCESS	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 Cl)	



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Clause	Requirement + Test	Result - Remark	Verdict
11.1.3	Measurements not made when engineering judgment and rationale by MANUFACTURER indicated temperature limits could not exceed, as documented in RISK MANAGEMENT FILE: (ISO 14971 CI. 4.2-4.4, 5, 6.2-6.5)	See appended Table 11.1.3d and RMF Reference to specific RISKS: (ISO 14971 Cl) Temperature measurement was performed. No heating effect of nearby surfaces. No RM considered necessary.	P
	Test corner not used where engineering judgment and rationale by MANUFACTURER indicated test corner will not impact measurements, as documented in RISK MANAGEMENT FILE: (ISO 14971 CI. 4.2-4.4, 5, 6.2-6.5)	No heating effect of nearby surfaces. No RM considered necessary.	N/A
	Probability of occurrence and duration of contact for parts likely to be touched and for APPLIED PARTS documented in RISK MANAGEMENT FILE	RMF Reference to specific RISKS: RISK H2 (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	P
	e) Where thermal regulatory devices make this method inappropriate, alternative methods for measurement are justified in the RISK MANAGEMENT FILE	Temperature measurement was performed.	N/A
11.1.4	GUARDS preventing contact with hot or cold accessible surfaces removable only with a TOOL		N/A
11.2	Fire prevention		Р
11.2.1	ENCLOSURE has strength and rigidity necessary to prevent a fire and met mechanical strength tests for ENCLOSURES in 15.3	See clause 15.3.	Р
11.2.2	Me equipment and me systems used in conjunction with OXYGEN RICH ENVIRONMENTS		N/A
11.2.2.1	RISK of fire in an OXYGEN RICH ENVIRONMENT reduced by means limiting spread of:	Component, not evaluated for use with Oxygen Rich Environment	N/A
	a) No sources of ignition discovered in an OXYGEN RICH ENVIRONMENT under any of the following conditions		N/A
	when temperature of material raised to its ignition temperature		N/A
	2) when temperatures affected solder or solder joints causing loosening, short circuiting, or other failures causing sparking or increasing material temperature to its ignition temperature		N/A
	3) when parts affecting safety cracked or changed outer shape exposing temperatures higher than 300°C or sparks due to overheating		N/A



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Clause	Requirement + Test	Result - Remark	Verdict		
	4) when temperatures of parts or components exceeded 300°C, atmosphere was 100 % oxygen, contact material solder, and fuel cotton		N/A		
	5) when sparks provided adequate energy for ignition by exceeding limits of Figs 35 to 37 (inclusive), atmosphere was 100 % oxygen, contact material solder, and fuel cotton		N/A		
	Deviations from worst case limits in 4) and 5) above based on lower oxygen concentrations or less flammable fuels justified and documented in RISK MANAGEMENT FILE	RMF Reference to specific RISKS: (ISO 14971 CI) ME Equipment is not intended to be used in an oxygen rich environment.	N/A		
	Alternative test in this clause did not identify existence of ignition sources at highest voltage or current, respectively:		N/A		
	A safe upper limit determined by dividing upper limit of voltage or current, respectively, with safety margin factor of three:		N/A		
	b) RESIDUAL RISK of fire in an OXYGEN RICH ENVIRONMENT as determined by application of RISK MANAGEMENT PROCESS is based on following configurations, or in combination	RMF Reference to specific RISKS: (ISO 14971 CI) ME Equipment is not intended to be used in an oxygen rich environment.	N/A		
	1) Electrical components in an OXYGEN RICH ENVIRONMENT provided with power supplies having limited energy levels lower than those considered sufficient for ignition in 11.2.2.1 a) as determined by examination, measurement or calculation of power, energy, and temperatures in NORMAL and SINGLE FAULT CONDITIONS identified in 11.2.3	See appended Tables 4.11, 11.1.1, 11.2.2.1 and 13.2	N/A		
	2) Max oxygen concentration measured until it did not exceed 25 % in ventilated compartments with parts that can be a source of ignition only in SINGLE FAULT CONDITION and can be penetrated by oxygen due to an undetected leak (%):		N/A		
	3) A compartment with parts or components that can be a source of ignition only under SINGLE FAULT CONDITION separated from another compartment containing an OXYGEN RICH ENVIRONMENT by sealing all joints and holes for cables, shafts, or other purposes		N/A		



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Clause	Requirement + Test	Result - Remark	Verdict	
	Effect of possible leaks and failures under SINGLE FAULT CONDITION that could cause ignition evaluated using a RISK ASSESSMENT to determine maintenance intervals by examination of documentation and RISK MANAGEMENT FILE	See Attachment No	N/A	
	4) Fire initiated in ENCLOSURE of electrical components in a compartment with OXYGEN RICH ENVIRONMENT that can become a source of ignition only under SINGLE FAULT CONDITIONS self-extinguished rapidly and no hazardous amount of toxic gases reached PATIENT as determined by analysis of gases:	See Attachment No	N/A	
1.2.2.2	RISK of ignition did not occur and oxygen concentration did not exceed 25% in immediate surroundings due to location of external exhaust outlets of an OXYGEN RICH ENVIRONMENT		N/A	
11.2.2.3	Electrical connections within a compartment containing an OXYGEN RICH ENVIRONMENT under NORMAL USE did not produce sparks		N/A	
	 Screw-attachments protected against loosening during use by varnishing, use of spring washers, or adequate torques 		N/A	
	 Soldered, crimped, and pin-and-socket connections of cables exiting ENCLOSURE include additional mechanical securing means 		N/A	
11.2.3	SINGLE FAULT CONDITIONS related to OXYGEN RICH ENVIRONMENTS ME EQUIPMENT and ME SYSTEMS considered		N/A	
	- Failure of a ventilation system constructed in accordance with 11.2.2.1 b) 2):	Component, not evaluated for use with Oxygen Rich Environment	N/A	
	- Failure of a barrier constructed in accordance with 11.2.2.1 b) 3):		N/A	
	- Failure of a component creating a source of ignition (as defined in 11.2.2.1 a):		N/A	
	- Failure of solid insulation or creepage and clearances providing equivalent of at least one MEANS OF PATIENT PROTECTION but less than two MEANS OF PATIENT PROTECTION that could create a source of ignition defined in 11.2.2.1 a):		N/A	
	- Failure of a pneumatic component resulting in leakage of oxygen-enriched gas:		N/A	
11.3	Constructional requirements for fire ENCLOSURES	S of ME EQUIPMENT	Р	



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Clause	Requirement + Test	Result - Remark	Verdict
	ME EQUIPMENT met this clause for alternate means of compliance with selected HAZARDOUS SITUATIONS and fault conditions in 13.1.2:	Power supply unit complies with the construction requirements for fire enclosure.	Р
		In addition, single fault as specified within the standard performed.	
	Constructional requirements were met, or	Power supply is provided with plastic enclosure without openings (Flammability UL94 V-1).	Р
		All internal components are mounted on the PCB rated UL94 V-0.	
		Output cable is classified as VW-1.	
		Short circuit on the cable output performed.	
		No deformation of the cable, no hazard.	
	- constructional requirements specifically analysed in RISK MANAGEMENT FILE:	RMF Reference to specific RISKS:	Р
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	Specific Requirements not met:	
		(ISO 14971 CI)	
		Power supply unit complies with the constructional requirements for fire enclosure.	
	Justification, when requirement not met:	EUT fulfils the constructional requirements of the standard.	Р
	a) Flammability classification of insulated wire	See appended Table 8.10	Р
	within fire ENCLOSURE is FV-1, or better, based on IEC 60695 series as determined by examination of data on materials	Output cable is classified as VW-1.	
		Short circuit on the cable performed.	
		No deformation of the cable, no fire, no emission of flames, no hazard.	
	Flammability classification of connectors,	See appended Table 8.10	Р
	printed circuit boards, and insulating material on which components are mounted is FV-2, or better, based on IEC 60695-11-10 as decided by examination of materials data	PCB is rated V-0.	
	If no FV Certification, FV tests based on IEC 60695-11-10 conducted on 3 samples of complete parts (or sections of it), including area with min. thickness, ventilation openings		N/A

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Clause	Requirement + Test	Result - Remark	Verdict
Olause	Requirement + rest	Result - Remark	verdict
	b) Fire ENCLOSURE met following:		Р
	1) No openings at bottom or, as specified in Fig 39, constructed with baffles as in Fig 38, or made of perforated metal as in Table 25, or a metal screen with a mesh \leq 2 × 2 mm centre to centre and wire diameter of at least 0.45 mm	Power supply unit is provided with plastic enclosure without openings.	P
	2) No openings on the sides within the area included within the inclined line C in Fig 39	Power supply unit is provided with plastic enclosure without openings.	Р
	3) ENCLOSURE, baffles, and flame barriers have adequate rigidity and are made of appropriate metal or of non-metallic materials:	See appended Table 8.10 Plastic enclosure is classified min. UL94 V-1.	P
11.4	ME EQUIPMENT and ME SYSTEMS intended for use w	vith flammable anaesthetics	N/A
	ME EQUIPMENT, ME SYSTEMS and parts described in ACCOMPANYING DOCUMENTS for use with flammable with Annex G	Not evaluated for use in the presence of flammable anaesthetics.	N/A
11.5	ME EQUIPMENT and ME SYSTEMS intended for use in conjunction with flammable agents		N/A
	MANUFACTURER'S RISK MANAGEMENT PROCESS addresses possibility of fire and associated mitigations as confirmed by examination of RISK MANAGEMENT FILE: (ISO 14971 CI. 4.2-4.4, 5, 6.2-6.5)	RMF Reference to specific RISKS: (ISO 14971 CI) Equipment is not intended to be used in conjunction with flammable agents.	N/A
11.6	Overflow, spillage, leakage, ingress of water or particulate matter, cleaning, disinfection, sterilization and compatibility with substances used with the ME EQUIPMENT		Р
11.6.1	Sufficient degree of protection provided against overflow, spillage, leakage, ingress of water or particulate matter, cleaning, disinfection and sterilization, and compatibility with substances used with ME EQUIPMENT:	See Appended Table 11.6.1	Р
11.6.2	Overflow in ME EQUIPMENT		N/A
	ME EQUIPMENT incorporates a reservoir or liquid storage that did not wet any MEANS OF PROTECTION, nor result in the loss of BASIC SAFETY OF ESSENTIAL PERFORMANCE:	Equipment does not contain a reservoir or liquid storage chamber.	N/A
	Maximum fill level is indicated by marking on the ME EQUIPMENT and a warning or safety notice is given, no HAZARDOUS SITUATION (as specified in 13.1) or unacceptable RISK due to overflow developed when the reservoir or liquid storage chamber is filled to its maximum capacity and the TRANSPORTABLE ME EQUIPMENT is tilted through an angle of 10°, or for MOBILE ME EQUIPMENT exceeding 45 kg, is moved over a threshold as described in 9.4.2.4.3.		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	No warning or safety notice provided regarding the maximum fill level, no HAZARDOUS SITUATION (as specified in 13.1) or unacceptable RISK due to overflow developed when the reservoir or liquid storage chamber was filled to 15 % above the maximum capacity and the TRANSPORTABLE ME EQUIPMENT was tilted through an angle of 10°, or in MOBILE ME EQUIPMENT exceeding 45 kg, was moved over a threshold as described in 9.4.2.4.3.		N/A
11.6.3	Spillage on ME EQUIPMENT and ME SYSTEM		N/A
	ME EQUIPMENT and ME SYSTEMS handling liquids constructed that spillage does not wet parts as determined by review of the RISK MANAGEMENT FILE and test	See appended Tables 11.6.1; 8.7, 8.8.3 and RMF Reference to specific RISK: (ISO 14971 CI) Equipment does not require handling of liquids in normal or foreseeable misuse.	N/A
	RISK ANALYSIS identifies the type of liquid, volume, duration and location of the spill:		N/A
11.6.5	Ingress of water or particulate matter into ME EQUIPMENT and ME SYSTEMS		Р
	ME EQUIPMENT with IP Code placed in least favourable position of NORMAL USE and subjected to tests of IEC 60529 (IP Code):	No protection against ingress of water provided.	N/A
	ME EQUIPMENT met dielectric strength and LEAKAGE CURRENT tests and there were no bridging of insulation or electrical components that could result in the loss of BASIC SAFETY or ESSENTIAL PERFORMANCE IN NORMAL CONDITION or in combination with a SINGLE FAULT CONDITION:	See appended Tables 8.7 8.8.3	P
11.6.6	Cleaning and disinfection of ME EQUIPMENT and ME SYSTEMS		N/A
	ME EQUIPMENT/ME SYSTEM and their parts and ACCESSORIES cleaned or disinfected using methods specified in instructions for use:	Cleaning or disinfection process not specified by the manufacturer.	N/A
	Effects of multiple cleanings/disinfections during EXPECTED SERVICE LIFE OF EQUIPMENT evaluated by MANUFACTURER:		N/A
11.6.7	Sterilization of ME EQUIPMENT and ME SYSTEMS		N/A
	ME EQUIPMENT, ME SYSTEMS and their parts or ACCESSORIES intended to be sterilized assessed and documented and compliant with tests:	Component, to be determined in end-product evaluation.	N/A
	RISK MANAGEMENT FILE includes an assessment of the RISKS associated with any deterioration following sterilization	RMF Reference to specific RISKS: (ISO 14971 CI) No parts subjected to sterilization.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
11.6.8	RISKS associated with compatibility of substances used with ME EQUIPMENT addressed in RISK MANAGEMENT PROCESS: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	RMF Reference to specific RISKS: (ISO 14971 CI) No special substances used in conjunction with the equipment.	N/A
11.7	ME EQUIPMENT, ME SYSTEM, and ACCESSORIES coming into direct or indirect contact with biological tissues, cells, or body fluids assessed and documented		N/A
11.8	Interruption and restoration of power supply did not result in a loss of BASIC SAFETY or ESSENTIAL PERFORMANCE	Shall be evaluated during end medical product approval.	N/A

12	ACCURACY OF CONTROLS AND INSTRUMENTS AND PROTECTION AGAINST HAZARDOUS OUTPUTS		N/A
12.1	RISKS associated with accuracy of controls and instruments stated:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		No controls and instruments provided.	
12.2	RISK of poor USABILITY, including identification, marking, and documents addressed in a USABILITY ENGINEERING:	USABILITY ENGINEERING PROCESS not applied for power supply.	N/A
12.3	MANUFACTURER implemented an ALARM SYSTEM compliant with IEC 60601-1-8:	No alarm system incorporated.	N/A
12.4	Protection against hazardous output		N/A
12.4.1	RISKS associated with hazardous output arising from intentional exceeding of safety limits	RMF Reference to specific RISKS:	N/A
	addressed in RISK MANAGEMENT PROCESS(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
12.4.2	- need for indication associated with hazardous output addressed in RISK	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
12.4.3	RISKS associated with accidental selection of excessive output values for ME EQUIPMENT with	RMF Reference to specific RISKS:	N/A
	a multi-purpose unit addressed in RISK MANAGEMENT PROCESS::	(ISO 14971 CI)	
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	No multi-purpose equipment.	
12.4.4	RISKS associated with incorrect output addressed in RISK MANAGEMENT PROCESS:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		No incorrect output possible.	
12.4.5	Diagnostic or therapeutic radiation		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
12.4.5.1	Adequate provisions to protect OPERATORS, PATIENTS, other persons and sensitive devices in vicinity of unwanted or excessive radiation		N/A
	Radiation safety ensured by compliance with requirements of appropriate standards		N/A
12.4.5.2	ME EQUIPMENT and ME SYSTEMS designed to produce X-radiation for diagnostic imaging purposes complied with IEC 60601-1-3:	Equipment does not produce diagnostic or therapeutic radiation.	N/A
12.4.5.3	RISKS associated with radiotherapy addressed in RISK MANAGEMENT PROCESS as(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	RMF Reference to specific RISKS:	N/A
	(130 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		Equipment does not produce diagnostic or therapeutic radiation.	
12.4.5.4	RISKS associated with ME EQUIPMENT producing diagnostic or therapeutic radiation other than	RMF Reference to specific RISKS:	N/A
	diagnostic X-rays and radiotherapy addressed in RISK MANAGEMENT PROCESS as	(ISO 14971 CI)	
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	Equipment does not produce diagnostic or therapeutic radiation.	
12.4.6	RISKS associated with diagnostic or therapeutic acoustic pressure addressed in RISK MANAGEMENT	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		Equipment does not produce diagnostic or therapeutic radiation.	

13	HAZARDOUS SITUATIONS AND FAULT CONDIT	IONS	Р
13.1	Specific HAZARDOUS SITUATIONS		Р
13.1.2	Emissions, deformation of ENCLOSURE or exceed	ing maximum temperature	Р
	- Emission of flames, molten metal, poisonous or ignitable substance in hazardous quantities did not occur	No fire, emission of molten metal or ignition of substances was noted during the tests.	Р
	- Deformation of ENCLOSURE impairing compliance with 15.3.1 did not occur	No deformation was noted during the tests.	Р
	- Temperatures of APPLIED PARTS did not exceed allowable values in Table 24:	No applied parts.	N/A
	- Temperatures of ME EQUIPMENT parts that are not APPLIED PARTS likely to be touched did not exceed values in Table 23:	See appended Table 11.1.1	Р
	-Allowable values for "other components and materials" in Table 22 times 1.5 minus 12.5 °C were not exceeded	Considered	Р



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Clause	Requirement + Test	Result - Remark	Verdict
	Limits for windings in Tables 26, 27, and 31 not exceeded	Temperature of transformer windings not exceeded.	Р
	Table 22 not exceeded in all other cases		Р
	After tests of this Clause, settings of THERMAL CUT-OUTS and OVER-CURRENT RELEASES did not change sufficiently to affect their safety function		N/A
13.1.3	- limits for LEAKAGE CURRENT IN SINGLE FAULT CONDITION did not exceed:	See appended Table 8.7	Р
	- voltage limits for ACCESSIBLE PARTS including	See appended Table 8.7	Р
	APPLIED PARTS did not exceed:	Output voltage of the medical power supply.	
13. 2	SINGLE FAULT CONDITIONS		Р
13.2.1	During the application of the SINGLE FAULT CONDITIONS listed in 13.2.2 to 13.2.13 (inclusive), the NORMAL CONDITIONS identified in 8.1 a) also applied in the least favourable combination	Considered.	Р
	ME EQUIPMENT complied with 13.2.2 -13.2.12:	See appended Table 13.2	Р
	RISK MANAGEMENT FILE includes and assessment of RISKS associated with leakage of liquid in a SINGLE FAULT CONDITION	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI) Equipment does not contain liquids.	
	RISK MANAGEMENT FILE defines the appropriate test conditions	Equipment does not contain liquids.	N/A
13.2.13	ME EQUIPMENT remained safe after tests of 13.2.13.2 to 13.2.13.4, and cooling down to	No heating elements or motors provided.	N/A
	within 3 °C of test environment temperature	EUT is intended for continuous operation.	
	ME EQUIPMENT examined for compliance or appropriate tests such as dielectric strength of motor insulation according to 8.8.3 conducted		N/A
	For insulation of thermoplastic materials relied upon as a MEANS OF PROTECTION, the ball-pressure test specified in 8.8.4.1 a) performed at a temperature 25 °C higher than temperature of insulation measured during tests of 13.2.13.2 to 13.2.13.4 (inclusive).		N/A
13.2.13.2	ME EQUIPMENT with heating elements		N/A
	a 1) thermostatically controlled ME EQUIPMENT with heating elements for building-in, r for unattended operation, or with a capacitor not protected by a fuse connected in parallel with THERMOSTAT contacts met tests	No heating elements provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	a 2) ME EQUIPMENT with heating elements RATED for non-CONTINUOUS OPERATION met tests		N/A
	a 3) other ME EQUIPMENT with heating elements met test		N/A
	When more than one test was applicable to same ME EQUIPMENT, tests performed consecutively		N/A
	Heating period stopped when a heating element or an intentionally weak part of a non-SELF-RESETTING THERMAL CUT-OUT ruptured, or current interrupted before THERMAL STABILITY without possibility of automatic restoration		N/A
	Test repeated on a second sample when interruption was due to rupture of a heating element or an intentionally weak part		N/A
	Both samples met 13.1.2, and open circuiting of a heating element or an intentionally weak part in second sample not considered a failure by itself		N/A
	b) ME EQUIPMENT with heating elements without adequate heat discharge, and supply voltage set at 90 or 110 % of RATED supply voltage, least favourable of the two (V):		N/A
	Operating period stopped when a non-SELF- RESETTING THERMAL CUT-OUT operated, or current interrupted without possibility of automatic restoration before THERMAL STABILITY		N/A
	ME EQUIPMENT switched off as soon as THERMAL STABILITY established and allowed to cool to room temperature when current not interrupted		N/A
	Test duration was equal to RATED operating time for non-CONTINUOUS OPERATION		N/A
	c) Heating parts of ME EQUIPMENT tested with ME EQUIPMENT operated in NORMAL CONDITION at 110 % of RATED supply voltage and as in 11.1, and		N/A
	1) Controls limiting temperature in NORMAL CONDITION disabled, except THERMAL CUT-OUTS		N/A
	2) When more than one control provided, they were disabled in turn		N/A
	3) ME EQUIPMENT operated at RATED DUTY CYCLE until THERMAL STABILITY achieved, regardless of RATED operating time		N/A
13.2.13.3	ME EQUIPMENT with motors		N/A
	a 1) For the motor part of the ME EQUIPMENT, compliance checked by tests of 13.2.8- 13.2.10, 13.2.13.3 b), 13.2.13.3 c), and 13.2.13.4, as applicable	No motor provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	To determine compliance with 13.2.9 and 13.2.10 motors in circuits running at 42.4 V peak a.c./ 60 V d.c. or less are covered with a single layer of cheesecloth which did not ignite during the test		N/A
	a 2) Tests on ME EQUIPMENT containing heating parts conducted at prescribed voltage with motor & heating parts operated simultaneously to produce the least favourable condition		N/A
	a 3) Tests performed consecutively when more tests were applicable to the same ME EQUIPMENT		N/A
	b) Motor met running overload protection test of this clause when:		N/A
	1) it is intended to be remotely or automatically controlled by a single control device with no redundant protection, or		N/A
	2) it is likely to be subjected to CONTINUOUS OPERATION while unattended		N/A
	Motor winding temperature determined during each steady period and maximum value did not exceed Table 27 (Insulation Class, Maximum temperature measured °C)::		N/A
	Motor removed from ME EQUIPMENT and tested separately when load could not be changed in appropriate steps		N/A
	Running overload test for motors operating at 42.4 V peak a.c./60 V d.c. or less performed only when examination and review of design indicated possibility of an overload		N/A
	Test not conducted where electronic drive circuits maintained a substantially constant drive current		N/A
	Test not conducted based on other justifications (justification):		N/A
	c) ME EQUIPMENT with 3-phase motors operated with normal load, connected to a 3-phase SUPPLY MAINS with one phase disconnected, and periods of operation per 13.2.10		N/A
13.2.13.4	ME EQUIPMENT RATED for NON-CONTINUOUS OPERATION	DN	N/A
	ME EQUIPMENT (other than HAND-HELD) operated under normal load and at RATED voltage or at upper limit of RATED voltage range until increase in temperature was \leq 5 °C in one hour, or a protective device operated	EUT is rated for continuous operation.	N/A
	When a load-reducing device operated in NORMAL USE, test continued with ME EQUIPMENT running idle		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Motor winding temperatures did not exceed values in 13.2.10:		N/A
	Insulation Class::		_
	Maximum temperature measured (°C):		-

14	PROGRAMMABLE ELECTRICAL MEDICAL SYSTEMS (PEMS)		N/A
14.1	Requirements in 14.2 to 14,12 not applied to PEMS when it provides no functionality necessary for BASIC SAFETY OF ESSENTIAL PERFORMANCE, or	No Such Parts / PESS not relied upon for BASIC SAFETY or ESSENTIAL PERFORMANCE.	N/A
	- when application of RISK MANAGEMENT showed that failure of PESS does not lead to unacceptable RISK:		N/A
	RISK MANAGEMENT FILE contains an assessment of RISKS associated with the failure of the PESS: (ISO 14971 Cl. 4.2-4.4, 5)	RMF Reference to specific RISKS: ISO 14971 CI)	N/A
	Requirements of 14.13 not applied to PEMS intended to be incorporated into an IT NETWORK	Equipment not connected to such network.	N/A
	When the requirements of 14.2 to 14.13 apply, the requirements of IEC 6204:2006 clause 4.3, 5, 7, 8 and 9 apply for the development or modification of software of each PESS	Software Class:	N/A
	Software development process for Software Classification applied in accordance with Clause 4.3 of IEC 62304:		N/A
	Software development process applied according to Clause 5 of IEC 62304:		N/A
	Software development process for Software risk management applied according to Clause 7 of IEC 62304:		N/A
	Software development process Configuration Management applied according to Clause 8 of IEC 62304:		N/A
	Software development process for Software Problem Resolution applied according to Clause 9 of IEC 62304:		N/A
14.2	Documents required by Clause 14 reviewed, approved, issued and revised according to a formal document control process:	The following documents were inspected:	N/A
14.3	RISK MANAGEMENT plan required by 4.2.2 includes reference to PEMS VALIDATION plan		N/A
14.4	A PEMS DEVELOPMENT LIFE-CYCLE including a set of defined milestones has been documented		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	At each milestone, activities to be completed, and VERIFICATION methods to be applied to activities have been defined		N/A
	Each activity including its inputs and outputs defined, and each milestone identifies RISK MANAGEMENT activities that must be completed before that milestone		N/A
	PEMS DEVELOPMENT LIFE-CYCLE tailored for a specific development by making plans detailing activities, milestones, and schedules		N/A
	PEMS DEVELOPMENT LIFE-CYCLE includes documentation requirements		N/A
14.5	A documented system for problem resolution within and between all phases and activities of PEMS DEVELOPMENT LIFE-CYCLE has been developed and maintained		N/A
14.6	RISK MANAGEMENT PROCESS		N/A
14.6.1	MANUFACTURER considered HAZARDS associated with software and hardware aspects of PEMS including those associated with the incorporating PEMS into an IT-NETWORK, components of third-party origin, legacy subsystems when compiling list of known or foreseeable HAZARDS		N/A
	RISK MANAGEMENT FILE includes known or foreseeable HAZARDS associated with software, hardware, incorporation of the PEMS into an IT-NETWORK, components of 3rd party origin and legacy subsystems	RMF Reference to specific HAZARDS: (ISO 14971 Cl)	N/A
14.6.2	Suitably validated tools and PROCEDURES assuring each RISK CONTROL measure reduces identified RISK(S) satisfactorily provided in addition to PEMS requirements in Clause 4.2.2.:		N/A
	RISK MANAGEMENT FILE documents the suitability of tools and procedures to validate each RISK CONTROL measure	RMF Reference to specific RISKS: (ISO 14971 Cl)	N/A
14.7	A documented requirement specification for PEMS and each of its subsystems (e.g. for a PESS) which includes ESSENTIAL PERFORMANCE and RISK CONTROL measures implemented by that system or subsystem	RMF Reference to specific RISK CONTROLS: (ISO 14971 Cl)	N/A
14.8	An architecture satisfying the requirement is specified for PEMS and each of subsystems: (ISO 14971 Cl. 6.3)	RMF Reference to specific RISK CONTROLS:	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
14.9	Design is broken up into sub systems and descriptive data on design environment documented:		N/A
14.10	A VERIFICATION plan containing the specified information used to verify and document functions implementing BASIC SAFETY, ESSENTIAL PERFORMANCE, or RISK CONTROL measures: (ISO 14971 CI. 6.3)	RMF Reference to specific RISK CONTROLS: (ISO 14971 Cl)	N/A
	 milestone(s) when VERIFICATION is to be performed for each function 		N/A
	- selection and documentation of VERIFICATION strategies, activities, techniques, and appropriate level of independence of the personnel performing the VERIFICATION		N/A
	- selection and utilization of VERIFICATION tools		N/A
	- coverage criteria for VERIFICATION		N/A
	The VERIFICATION performed according to the VERIFICATION plan and results of the VERIFICATION activities documented		N/A
14.11	A PEMS VALIDATION plan containing validation of BASIC SAFETY & ESSENTIAL PERFORMANCE:		N/A
	The PEMS VALIDATION performed according to the PEMS VALIDATION plan with results of PEMS VALIDATION activities and methods used for PEMS VALIDATION documented		N/A
	The person with overall responsibility for PEMS VALIDATION is independent		N/A
	All professional relationships of members of PEMS VALIDATION team with members of design team documented in RISK MANAGEMENT FILE (ISO 14971 CI. 6.3)	RMF Reference to specific RISK CONTROLS: (ISO 14971 Cl)	N/A
14.12	Continued validity of previous design documentation assessed under a documented modification/change PROCEDURE		N/A
	Software Classification for Software changes applied in accordance with Clause 4.3 of IEC 62304:	Software Class:	N/A
	Software Process for Software changes applied according to Clause 5 of IEC 62304:		N/A
	RISK MANAGEMENT for Software changes applied according to Clause 7 of IEC 62304:		N/A
	Configuration management of software changes applied per Clause 8 of IEC 62304:		N/A
	Problem resolution for Software changes applied according to Clause 9 of IEC 62304:		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
14.13	For PEMS incorporated into an IT-NETWORK not VALIDATED by the PEMS MANUFACTURER, instructions made available for implementing the connection include the following:	Equipment not connected to such network.	N/A
	a) Purpose of the PEMS connection to an IT- NETWORK		N/A
	b) required characteristics of the IT-NETWORK		N/A
	c) required configuration of the IT-NETWORK		N/A
	d) technical specifications of the network connection, including security specifications		N/A
	e) intended information flow between the PEMS, the IT-NETWORK and other devices on the IT-NETWORK, and the intended routing through the IT-NETWORK		N/A
	f) a list of HAZARDOUS SITUATIONS resulting from failure of the IT-NETWORK to provide the required	RMF Reference to specific hazardous situations:	N/A
	characteristics (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.3)	(ISO 14971 Cl)	
		Equipment not connected to such network. No RM considered necessary.	
	ACCOMPANYING DOCUMENTS for the RESPONSIBLE OR following:	GANIZATION include the	N/A
	- statement that connection to IT-NETWORKS including other equipment could result in previously unidentified RISKS TO PATIENTS, OPERATORS or third parties		N/A
	 Notification that the RESPONSIBLE ORGANIZATION should identify, analyse, evaluate and control these RISKS 		N/A
	- Notification that changes to the IT-NETWORK could introduce new RISKS that require additional analysis		N/A
	- Changes to the IT-NETWORK include: - changes in network configuration - connection of additional items - disconnection of items - update of equipment - upgrade of equipment		N/A

15	CONSTRUCTION OF ME EQUIPMENT		Р
15.1	and indicators of ME EQUIPMENT addressed	A green LED is illuminated when the unit is operating. No other controls or indicators provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
15.2	Parts of ME EQUIPMENT subject to mechanical wear, electrical, environmental degradation or ageing resulting in unacceptable RISK when unchecked for a long period, are accessible for inspection, replacement, and maintenance	No such parts provided.	N/A
	Inspection, servicing, replacement, and adjustment of parts of ME EQUIPMENT can easily be done without damage to or interference with adjacent parts or wiring		N/A
15.3	Mechanical strength		Р
15.3.1	Mould stress relief, push, impact, drop, and rough handling tests did not result in loss of BASIC SAFETY OR ESSENTIAL PERFORMANCE	See below	Р
15.3.2	Push test conducted:	See Appended Table 15.3	Р
	No damage resulting in an unacceptable RISK sustained	Equipment does not sustain any damage during testing.	Р
15.3.3	Impact test conducted:	See Appended Table 15.3	Р
	No damage resulting in an unacceptable RISK sustained	Equipment does not sustain any damage during testing.	Р
15.3.4	Drop test		Р
15.3.4.1	Sample of HAND-HELD ME EQUIPMENT, ACCESSORIES and HAND-HELD part with SAFE WORKING LOAD tested:	See Appended Table 15.3	N/A
	No unacceptable RISK resulted		N/A
15.3.4.2	Sample of PORTABLE ME EQUIPMENT, ACCESSORIES and PORTABLE part with SAFE WORKING LOAD withstood stress as demonstrated by test:	See Appended Table 15.3	Р
	No damage resulting in an unacceptable RISK sustained	EUT is provided with enclosure with adequate strength and rigidity.	Р
15.3.5	MOBILE ME EQUIPMENT and MOBILE part with SAFE WORKING LOAD and in most adverse condition in NORMAL USE passed Rough Handling tests:	Equipment is not mobile.	N/A
	No damage resulting in an unacceptable RISK sustained		N/A
15.3.6	Examination of ENCLOSURE made from moulded or formed thermoplastic material indicated that material distortion due to release of internal stresses by moulding or forming operations will not result in an unacceptable RISK	EUT is provided with enclosure with adequate strength and rigidity.	Р



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Clause	Requirement + Test	Result - Remark	Verdict
	Mould-stress relief test conducted by placing one sample of complete ME EQUIPMENT, ENCLOSURE or a portion of larger ENCLOSURE, for 7 hours in a circulating air oven at 10°C over the max temperature measured on ENCLOSURE in 11.1.3, but no less than 70 °C:	See Appended Table 15.3 Test was performed for 7 hours at 95°C ambient temperature. No damage of the enclosure. All plastic enclosures were tested.	P
	No damage resulting in an unacceptable RISK	No risk of shrinkage or distortion on enclosures due to release of internal stresses.	Р
15.3.7	INTENDED USE, EXPECTED SERVICE LIFE, and conditions for transport and storage were taken into consideration for selection and treatment of materials used in construction of ME EQUIPMENT		N/A
	Based on review of EQUIPMENT, ACCOMPANYING DOCUMENTS, specifications and processing of materials, and MANUFACTURER'S relevant tests or calculations, corrosion, ageing, mechanical wear, degradation of biological materials due to bacteria, plants, animals and the like, will not result in an unacceptable RISK		N/A
15.4	ME EQUIPMENT components and general assembl	у	Р
15.4.1	Incorrect connection of accessible connectors, removable without a TOOL, prevented where an unacceptable RISK exists,: (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	RMF Reference to specific RISKS: (ISO 14971 CI)	N/A
	a) Plugs for connection of PATIENT leads or PATIENT cables cannot be connected to outlets on same ME EQUIPMENT intended for other functions,:		N/A
	b) Medical gas connections on ME EQUIPMENT for different gases to be operated in NORMAL USE are not interchangeable inspection:	No medical gas connections provided.	N/A
15.4.2	Temperature and overload control devices		N/A
15.4.2.1	a) THERMAL CUT-OUTS and OVER-CURRENT RELEASES with automatic resetting not used in ME EQUIPMENT when their use could lead to a HAZARDOUS SITUATION: (ISO 14971 Cl. 4.2-4.4, 5)	RMF Reference to specific RISKS: (ISO 14971 CI) No automatic resetting thermal cut-out or over-current release provided.	N/A
	b) THERMAL CUT-OUTS with a safety function with reset by a soldering not fitted in ME EQUIPMENT	No thermal cut-out that is reset by soldering provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	c) An additional independent non-SELF- RESETTING THERMAL CUT-OUT is provided:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4)	(ISO 14971 CI)	
		No thermostat provided.	
	d) Operation of THERMAL CUT-OUT OR OVER CURRENT RELEASE doesn't result in a HAZARDOUS	RMF Reference to specific RISKS:	N/A
	SITUATION OF IOSS OF ESSENTIAL PERFORMANCE: (ISO 14971 CI. 4.2-4.4)	(ISO 14971 CI)	
	,	No thermal cut-out or over current release provided.	
	e) Capacitors or other spark-suppression devices not connected between contacts of THERMAL CUT-OUTS	No thermal cut-out provided.	N/A
	f) Use of THERMAL CUT-OUTS OR OVER-CURRENT RELEASES do not affect safety as verified by following tests	No thermal cut-out or over current release provided.	N/A
	- Positive temperature coefficient devices) complied with IEC 60730-1: 2010, Clauses 15, 17, J.15, and J.17		N/A
	- ME EQUIPMENT containing THERMAL CUT-OUTS and OVER-CURRENT RELEASES operated under the conditions of Clause 13:		N/A
	- SELF-RESETTING THERMAL CUT-OUTS and OVER-CURRENT RELEASES including circuits performing equivalent functions Certified according to appropriate standards		N/A
	- In the absence of Certification in accordance with IEC standards, SELF-RESETTING THERMAL CUT-OUTS and OVER-CURRENT RELEASES including circuits performing equivalent functions operated 200 times		N/A
	Manual reset THERMAL CUT-OUTS and OVER- CURRENT RELEASES Certified in accordance with appropriate IEC standards		N/A
	manual reset THERMAL CUT-OUTS and OVER-CURRENT RELEASES operated 10 times		N/A
	Thermal protective devices tested separately from ME EQUIPMENT when engineering judgment indicated test results would not be impacted		N/A
	g) Protective device incorporating a fluid filled container with heating means, operated when heater switched on with container empty and prevented an unacceptable RISK due to overheating	No fluid filled container with heating means provided.	N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	h) ME EQUIPMENT with tubular heating elements provided with protection against overheating:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4)	(ISO 14971 CI)	
		No tubular heating elements provided.	
15.4.2.2	Temperature settings clearly indicated when means provided to vary setting of THERMOSTATS	No thermostats incorporated.	N/A
15.4.3	Batteries		N/A
15.4.3.1	Battery housings provided with ventilation: (ISO 14971 Cl. 4.2-4.4)	RMF Reference to specific RISKS:	N/A
	(130 1437 1 Cl. 4.2-4.4)	(ISO 14971 CI)	
		Neither batteries nor battery compartments incorporated	
	Battery compartments designed to prevent accidental short circuiting		N/A
15.4.3.2	Means provided to prevent incorrect connection of polarity:		N/A
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with incorrect connection	RMF Reference to specific RISKS:	N/A
	or replacement of batteries:	(ISO 14971 CI)	
	(100 1407 1 011 412 414)	Neither batteries nor battery compartments incorporated.	
15.4.3.3	Overcharging of battery prevented by virtue of design:		N/A
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with overcharging of	RMF Reference to specific RISKS:	N/A
	batteries: (ISO 14971 Cl. 4.2-4.4)	(ISO 14971 CI)	
		Neither batteries nor battery compartments incorporated.	
15.4.3.4	Primary lithium batteries comply with IEC 80086-4	No primary lithium battery used.	N/A
	Secondary lithium batteries comply with IEC 62133	No secondary lithium battery used.	N/A
15.4.3.5	A properly RATED protective device provided within INTERNAL ELECTRICAL POWER SOURCE to protect against fire:		N/A
	Protective device has adequate breaking capacity		N/A
	Justification for OVER-CURRENT RELEASES or FUSE exclusion is documented		N/A
	Short circuit test between the positive and negative poles of an INTERNAL ELECTRICAL POWER SOURCE between the output and protective device(s) omitted where 2 MOOPs provided, or		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	Short circuit between the positive and negative poles of an INTERNAL ELECTRICAL POWER SOURCE between the output and protective device(s) does not result in any HAZARDOUS SITUATION		N/A
15.4.4	Indicator lights provided to indicate ME EQUIPMENT is ready for:	A green LED is illuminated when the unit is operating or ready for operation (delivering output voltage).	Р
	An additional indicator light provided on ME EQUIPMENT with a stand-by state or a warm-up state exceeding 15 s,		N/A
	Indicator lights provided on ME EQUIPMENT incorporating non-luminous heaters to indicate heaters are operational	No heaters provided.	N/A
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with the use of indicator lights for EQUIPMENT incorporating non-luminous heaters: (ISO 14971 CI. 4.2-4.4)	RMF Reference to specific RISKS: (ISO 14971 Cl) No heaters provided.	N/A
	Requirement not applied to heated stylus-pens for recording purposes		N/A
	Indicator lights provided on ME EQUIPMENT to indicate an output exists		N/A
	Colours of indicator lights complied with 7.8.1	Only green LED is illuminated when the unit is operating.	Р
	Charging mode visibly indicated	No charging mode incorporated.	N/A
15.4.5	RISKS associated with pre-set controls addressed in RISK MANAGEMENT PROCESS:	RMF Reference to specific RISKS:	N/A
	(ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	(ISO 14971 CI)	
		No pre-set controls provided.	
15.4.6	Actuating parts of controls of ME EQUIPMENT	No. and address of the second	N/A
15.4.6.1	a) Actuating parts cannot be pulled off or loosened during NORMAL USE	No actuating parts or controls provided.	N/A
	b) Controls secured so that the indication of any scale always corresponds to the position of the control		N/A
	c) Incorrect connection prevented by adequate construction when it could be separated without use of a TOOL		N/A
	When torque values per Table 30 applied knobs did not rotate:		N/A
	Tests conducted with no unacceptable RISK:		N/A
15.4.6.2	Stops on rotating/ movable parts of controls are of adequate mechanical strength:		N/A

N/A

N/A

Ρ

See appended Tables 15.5.1.2

and 15.5.1.3



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Clause	Requirement + Test	Result - Remark	Verdict
	Torque values in Table 30 applied:		N/A
	No unexpected change of the controlled parameter when tested:		N/A
15.4.7	Cord-connected HAND-HELD and foot-operated co	ontrol devices	N/A
15.4.7.1	a) HAND-HELD control devices of ME EQUIPMENT complied with 15.3.4.1	No hand held control device provided.	N/A
	b) Foot-operated control device supported an actuating force of 1350 N in its position of NORMAL USE with no damage:	No foot operated control device provided.	N/A
15.4.7.2	Control device of HAND-HELD and foot-operated control devices turned in all possible abnormal positions and placed on a flat surface:	No inadvertent change of control device possible.	N/A
	No unacceptable RISK caused by changing control setting when accidentally placed in an abnormal position		N/A
15.4.7.3	a) Foot-operated control device is at least rated IPX1		N/A
	b) ENCLOSURE of foot operated control devices containing electrical circuits is at least IPX6:		N/A
15.4.8	Aluminium wires less than 16 mm ² in cross- sectional area are not used	Aluminium wires not used.	Р
15.4.9	a) Oil container in PORTABLE ME EQUIPMENT allows for expansion of oil and is adequately sealed	No oil container provided.	N/A
	b) Oil containers in MOBILE ME EQUIPMENT sealed to prevent loss of oil during transport		N/A
	A pressure-release device operating during NORMAL USE is provided		N/A
			+

c) Partially sealed oil-filled ME EQUIPMENT and its

parts provided with means for checking the oil

MAINS SUPPLY TRANSFORMERS OF ME EQUIPMENT and transformers providing

ME EQUIPMENT and technical description

separation in accordance with 8.5

examined, and manual tests conducted to confirm compliance with above requirements

Transformers of ME EQUIPMENT are protected

During tests, windings did not open, no HAZARDOUS SITUATION occurred, and maximum temperatures of windings did not exceed

against overheating.....:

level to detect leakage

Overheating

values in Table 31

15.5

15.5.1

15.5.1.1



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Clause	Requirement + Test	Result - Remark	Verdict
	Dielectric strength test conducted after short circuit and overload tests:	See appended Table 15.5.2 Switch mode transformer incorporated within the equipment.	P
		Dielectric strength test according to Clause 8.8.3 performed.	
15.5.1.2	Transformer output winding short circuited, and test continued until protective device operated or THERMAL STABILITY achieved:	See appended Table 15.5.1.2	Р
	Short circuit applied directly across output windings	Short circuit performed on transformer output windings.	Р
15.5.1.3	Multiple overload tests conducted on windings	See appended Table 15.5.1.3	Р
15.5.2	Transformers operating at a frequency above 1kHz tested according to clause 8.8.3:	Switch mode transformer provided. Therefore this test is not applicable.	N/A
	Transformer windings provided with adequate insulation	Switch mode transformer provided. Therefore this test is not applicable.	N/A
	Dielectric strength tests were conducted:	Switch mode transformer provided. Therefore this test is not applicable.	N/A
15.5.3	Transformers forming MEANS OF PROTECTION as required by 8.5 comply with:	See appended Table 8.10	Р
		Transformer tested within the equipment.	
	- Means provided to prevent displacement of end turns		Р
	- protective earth screens with a single turn have insulated overlap		N/A
	- Exit of wires form internal windings of toroid transformers protected with double sleeving	No toroid transformer construction.	N/A
	- insulation between primary and secondary windings complies with 8.8.2		N/A
	- CREEPAGE DISTANCES and AIR CLEARANCE comply with 8.9.4		N/A

16	ME SYSTEMS		N/A
16.1	After installation or subsequent modification, ME SYSTEM didn't result in an unacceptable RISK		N/A



IEC 60601-1			
Clause	Requirement + Test	Result - Remark	Verdict
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with installation and modification of an ME SYSTEM	RMF Reference to specific RISKS: (ISO 14971 CI) No different configurations specified. ME Equipment is not intended to be combined with other equipment to create a system.	N/A
	Only HAZARDS arising from combining various equipment to form a ME SYSTEM considered		N/A
	- ME SYSTEM provides the level of safety within the PATIENT ENVIRONMENT equivalent to ME EQUIPMENT complying with this standard		N/A
	- ME SYSTEM provides the level of safety outside PATIENT ENVIRONMENT equivalent to equipment complying with their respective IEC or ISO safety standards		N/A
	- tests performed in NORMAL CONDITION, except as specified		N/A
	- tests performed under operating conditions specified by MANUFACTURER of ME SYSTEM		N/A
	Safety tests previously conducted on individual equipment of ME SYSTEM according to relevant standards not repeated		N/A
	RISK MANAGEMENT methods used by MANUFACTURER of an ME SYSTEM reconfigurable by RESPONSIBLE ORGANIZATION OF OPERATOR		N/A
	Non-ME EQUIPMENT used in ME SYSTEM complied with applicable IEC or ISO safety standards		N/A
	Equipment relying only on BASIC INSULATION for protection against electric shock not used in ME SYSTEM		N/A
16.2	ACCOMPANYING DOCUMENTS of an ME SYSTEM		N/A
	Documents containing all data necessary for ME SYSTEM to be used as intended by MANUFACTURER including a contact address accompany ME SYSTEM or modified ME SYSTEM		N/A
	ACCOMPANYING DOCUMENTS regarded as a part of ME SYSTEM		N/A
	a) ACCOMPANYING DOCUMENTS provided for each item of ME EQUIPMENT supplied by MANUFACTURER		N/A
	b) ACCOMPANYING DOCUMENTS provided for each item of non-ME EQUIPMENT supplied by MANUFACTURER		N/A
	c) the required information is provided:		N/A



IEC 60601-1			
Clause	Requirement + Test	Result - Remark	Verdict
	- specifications, instructions for use as intended by MANUFACTURER, and a list of all items forming the ME SYSTEM		N/A
	 instructions for installation, assembly, and modification of ME SYSTEM to ensure continued compliance with this standard 		N/A
	- instructions for cleaning and, when applicable, disinfecting and sterilizing each item of equipment or equipment part forming part of the ME SYSTEM		N/A
	- additional safety measures to be applied during installation of ME SYSTEM		N/A
	- identification of parts of ME SYSTEM suitable for use within the PATIENT ENVIRONMENT		N/A
	additional measures to be applied during preventive maintenance		N/A
	- a warning forbidding placement of MULTIPLE SOCKET-OUTLET, when provided and it is a separate item, on the floor		N/A
	- a warning indicating an additional MULTIPLE SOCKET-OUTLET or extension cord not to be connected to ME SYSTEM		N/A
	a warning to connect only items that have been specified as part of ME SYSTEM or specified as being compatible with ME SYSTEM		N/A
	- maximum permissible load for any MULTIPLE SOCKET-OUTLET(S) used with ME SYSTEM		N/A
	- instructions indicating MULTIPLE SOCKET- OUTLETS provided with the ME SYSTEM to be used only for supplying power to equipment intended to form part of ME SYSTEM		N/A
	- an explanation indicating RISKS of connecting non-ME EQUIPMENT supplied as a part of ME SYSTEM directly to wall outlet when non-ME EQUIPMENT is intended to be supplied via a MULTIPLE SOCKET-OUTLET with a separating transformer		N/A
	- an explanation indicating RISKS of connecting any equipment supplied as a part of ME SYSTEM to MULTIPLE SOCKET-OUTLET		N/A
	- permissible environmental conditions of use for ME SYSTEM including conditions for transport and storage		N/A
	- instructions to OPERATOR not to, simultaneously, touch parts referred to in 16.4 and PATIENT		N/A
	d) the following instructions provided for use by RESPONSIBLE ORGANIZATION:		N/A



	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict
	- adjustment, cleaning, sterilization, and disinfection PROCEDURES		N/A
	assembly of ME SYSTEMS and modifications during actual service life shall be evaluated based on the requirements of this standard		N/A
16.3	Instructions for use of ME EQUIPMENT intended to receive its power from other equipment in an ME SYSTEM, describe the other equipment to ensure compliance with these requirements		N/A
	Transient currents restricted to allowable levels for the specified IPS or UPS::		N/A
	Technical description and installation instructions specify the actual transient currents where an IPS or UPS is not specified		N/A
16.4	Parts of non-ME EQUIPMENT in PATIENT ENVIRONMENT subject to contact by OPERATOR during maintenance, calibration, after removal of covers, connectors operated at a voltage ≤ voltage in 8.4.2 c)		N/A
16.5	Safety measures incorporating a SEPARATION DEVICE applied when FUNCTIONAL CONNECTION between ME EQUIPMENT and other items of an ME SYSTEM or other systems can cause allowable values of LEAKAGE CURRENT to exceed		N/A
	SEPARATION DEVICE has dielectric strength, CREEPAGE and CLEARANCES required for one MEANS OF OPERATOR PROTECTION		N/A
	WORKING VOLTAGE was highest voltage across SEPARATION DEVICE during a fault condition, but not less than MAXIMUM MAINS VOLTAGE (V):		N/A
16.6	LEAKAGE CURRENTS		N/A
16.6.1	TOUCH CURRENT IN NORMAL CONDITION did not exceed 100 μA:		N/A
	TOUCH CURRENT did not exceed 500 μA in event of interruption of any non-PERMANENTLY INSTALLED PROTECTIVE EARTH CONDUCTOR:		N/A
16.6.2	Current in PROTECTIVE EARTH CONDUCTOR of MULTIPLE SOCKET-OUTLET didn't exceed 5 mA:		N/A
16.6.3	PATIENT LEAKAGE CURRENT and total PATIENT LEAKAGE CURRENT of ME SYSTEM in NORMAL CONDITION did not exceed values:		N/A
16.7	ME SYSTEM complied with applicable requirements of Clause 9:		N/A
16.8	Interruption and restoration power to the ME SYSTEM or any part of the ME SYSTEM did not result in a loss of BASIC SAFETY OR ESSENTIAL PERFORMANCE		N/A



	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict
16.9	ME SYSTEM connections and wiring		
16.9.1	Incorrect connection of accessible connectors, removable without a TOOL, prevented where unacceptable RISK can result:		N/A
	RISK MANAGEMENT FILE includes an assessment of RISKS associated with plugs for connection of PATIENT leads or cables likely to be located in the PATIENT ENVIRONMENT	RMF Reference to specific RISKS: (ISO 14971 CI)	N/A
	- Plugs for connection of PATIENT leads or PATIENT cables could not be connected to other outlets of the same ME SYSTEM likely to be located in PATIENT ENVIRONMENT, except when examination of connectors and interchanging them proved no unacceptable RISK results		N/A
	Medical gas connections on the ME SYSTEM for different gasses operated in NORMAL USE are not interchangeable	No medical gas connections provided.	N/A
16.9.2	MAINS PARTS, components and layout		N/A
16.9.2.1	a) – MULTIPLE SOCKET-OUTLET only allows connection using a TOOL, or		N/A
	- MULTIPLE SOCKET-OUTLET is of a type that cannot accept MAINS PLUGS of any of the kinds specified in IEC/TR 60083, or		N/A
	MULTIPLE SOCKET-OUTLET is supplied via a separating transformer		N/A
	b) – MULTIPLE SOCKET-OUTLET marked with safety sign 2 of Table D.2 visible in NORMAL USE, and		N/A
	- marked either individually or in combinations, with the maximum allowed continuous output in amperes or volt-amperes, or		N/A
	- marked to indicate the equipment or equipment parts it may safely be attached to		N/A
	- MULTIPLE SOCKET-OUTLET is a separate item or an integral part of ME EQUIPMENT or non-ME EQUIPMENT		N/A
	c) MULTIPLE SOCKET-OUTLET complied with IEC 60884-1 and the following requirements:		N/A
	- CREEPAGE and CLEARANCES complied with 8.9		N/A
	- It is CLASS I, and PROTECTIVE EARTH CONDUCTOR is connected to earthing contacts in socket-outlets		N/A
	- PROTECTIVE EARTH TERMINALS and PROTECTIVE EARTH CONNECTIONS comply with 8.6:		N/A
	- ENCLOSURE complied with 8.4.2 d)		N/A



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Clause	Requirement + Test	Result - Remark	Verdict	
	- MAINS TERMINAL DEVICES and wiring complied with 8.11.4, when applicable		N/A	
	- RATINGS of components are not in conflict with conditions of use:		N/A	
	Electrical terminals and connectors of MULTIPLE SOCKET-OUTLETS prevent incorrect connection of accessible connectors removable without a TOOL		N/A	
	- POWER SUPPLY CORD complied with 8.11.3		N/A	
	d) Additional requirements applied when MULTIPLE SOCKET-OUTLET combined with a separating transformer:		N/A	
	- Separating transformer complied with this standard or IEC 61558-2-1,:		N/A	
	- Separating transformer is CLASS I		N/A	
	Degree of protection against ingress of water specified as in IEC 60529		N/A	
	Separating transformer assembly marked according to 7.2 and 7.3		N/A	
	MULTIPLE SOCKET-OUTLET permanently connected to separating transformer, or socket-outlet of separating transformer assembly cannot accept MAINS PLUGS as identified in IEC/TR 60083		N/A	
16.9.2.2	The impedance between the protective earth pin in the MAINS PLUG and any part that is PROTECTIVELY EARTHED did not exceed 200 m Ω		N/A	
	Removal of any single item of equipment in ME SYSTEM will not interrupt the protective earthing of any other part without simultaneous disconnection of electrical supply to that part		N/A	
	Additional PROTECTIVE EARTH CONDUCTORS can be detachable only by use of a TOOL		N/A	
16.9.2.3	Conductors connecting different items within an ME SYSTEM protected against mechanical		N/A	

17	ELECTROMAGNETIC COMPATIBILITY OF ME EQUIPMENT AND ME SYSTEMS		Р
	RISKS associated confirmed by review:		Р
	- electromagnetic phenomena at locations where ME EQUIPMENT or ME SYSTEM is to be used as stated in ACCOMPANYING DOCUMENTS:	Power supply unit is not end medical product, shall be considered during end medical product approval.	N/A

damage



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Clause	Requirement + Test	Result - Remark	Verdict
	RISK MANAGEMENT FILE includes an assessment of risks associated with the introduction of electromagnetic phenomena into the environment by the EQUIPMENT or SYSTEM	RMF Reference to specific RISKS: M1 (ISO 14971 Cl. 4.2-4.4, 5, 6.2-6.5)	P
	- introduction of electromagnetic phenomena into environment by ME EQUIPMENT or ME SYSTEM that might degrade performance of other devices, electrical equipment, and systems	See IEC 60601-1-2 Report (WTX19X09063416E, performed by Shenzhen SEM Test Technology Co., Ltd.).	Р

ANNEX G	PROTECTION AGAINST HAZARDS OF IGNITION OF FLAMMABLE ANESTHETIC MIXTURES		
G.2	Locations and basic requirements		N/A
G.2.1	Parts of CATEGORY APG ME EQUIPMENT in which a FLAMMABLE ANAESTHETIC MIXTURE WITH AIR OCCURS are CATEGORY AP Or APG ME EQUIPMENT and complied with G.3, G.4, and G.5	Equipment is not intended to be used in conjunction with flammable anaesthetics mixtures.	N/A
G.2.2	FLAMMABLE AESTHETIC MIXTURE WITH		N/A
G.2.3	A FLAMMABLE AESTHETIC MIXTURE WITH OXYGEN OF NITROUS OXIDE		N/A
G.2.4	ME EQUIPMENT specified for use with FLAMMABLE AESTHETIC MIXTURE WITH AIR complied with G.4 and G.5		N/A
G.2.5	ME EQUIPMENT or parts thereof for use with FLAMMABLE AESTHETIC MIXTURE WITH OXYGEN OR NITROUS OXIDE comply with G.4 and G.6		N/A
	ME EQUIPMENT in G.2.4 to G.2.5 met appropriate tests of G.3-G.5 conducted after tests of 11.6.6 and 11.6.7		N/A
G.3	Marking, ACCOMPANYING DOCUMENTS		N/A
G.3.1	CATEGORY APG ME EQUIPMENT prominently marked "APG" (symbol 23 in Table D.1):		N/A
	Length of green-coloured band is ≥ 4 cm, and size of marking is as large as possible for particular case		N/A
	When above marking not possible, relevant information included in instructions for use:		N/A
	Marking complied with tests and criteria of 7.1.2 and 7.1.3		N/A
G.3.2	CATEGORY AP ME EQUIPMENT prominently marked, with a green-coloured circle "AP" (symbol 22 in Table D.1):		N/A
	Marking is as large as possible for the particular case		N/A
	When above marking not possible, the relevant information included in instructions for use:		N/A



	Marking complied with tests and criteria of		N/A
Clause	Requirement + Test	Result - Remark	Verdict
	IEC 60601-1		

	Marking complied with tests and criteria of 7.1.2 and 7.1.3		N/A
G.3.3	The marking placed on major part of ME EQUIPMENT for CATEGORY AP or APG parts		N/A
G.3.4	ACCOMPANYING DOCUMENTS contain an indication enabling the RESPONSIBLE ORGANIZATION to distinguish between CATEGORY AP and APG parts		N/A
G.3.5	Marking clearly indicates which parts are CATEGORY AP or APG when only certain ME EQUIPMENT parts are CATEGORY AP Or APG		N/A
G.4	Common requirements for CATEGORY AP and CATE	EGORY APG ME EQUIPMENT	N/A
G.4.1	a) CREEPAGE and CLEARANCES are according to Table 12 for one MEANS OF PATIENT PROTECTION		N/A
	b) Connections protected against accidental disconnection		N/A
	c) CATEGORY AP and APG not provided with a DETACHABLE POWER SUPPLY CORD,		N/A
G.4.2	Construction details		N/A
	a) Opening of an ENCLOSURE protecting against penetration of gases or vapours into ME EQUIPMENT or its parts possible only with a TOOL		N/A
	b) ENCLOSURE complies with:		N/A
	- no openings on top covers of ENCLOSURE,		N/A
	openings in side-covers prevented penetration of a solid cylindrical test rod		N/A
	openings in base plates prevented penetration of a solid cylindrical test		N/A
	c) Short circuiting conductor(s) to a conductive part (when no explosive gasses) did not result in loss of integrity of the part, an unacceptable temperature, or any HAZARDOUS SITUATION		N/A
G.4.3	a) Electrostatic charges prevented on CATEGORY AP and APG ME EQUIPMENT by a combination of appropriate measures		N/A
	Use of antistatic materials with a limited electrical resistance: :		N/A
	- Provision of electrically conductive paths from ME EQUIPMENT or its parts to a conductive floor, protective earth or potential equalization system, or via wheels to an antistatic floor		N/A
	b) Electrical resistance limits of aesthetic tubing, mattresses/ pads, castor tires & other antistatic material comply with ISO 2882:		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
Jiause	requirement + rest	Nesult - Nemark	Verdict
G.4.4	Corona cannot be produced by components or parts of ME EQUIPMENT operating at more than 2000 V a.c. or 2400 V d.c. and not included in ENCLOSURES complying with G.5.4 or G.5.5		N/A
3. 5	Requirements and tests for CATEGORY AP ME EQUI	PMENT, parts and components	N/A
G.5.1	ME EQUIPMENT, its parts or components do not ignite FLAMMABLE AESTHETIC MIXTURES WITH AIR under NORMAL USE and CONDITIONS based on compliance with G.5.2 to G.5.5		N/A
	Alternatively, ME EQUIPMENT, its parts, and components complied with requirements of IEC 60079-0 for pressurized ENCLOSURES (IEC 60079-2); for sand-filled ENCLOSURES, IEC 60079-5; or for oil immersed equipment, IEC 60079-6; and with this standard excluding G.5.2 to G.5.5:		N/A
3.5.2	Temperature limits:		N/A
G.5.3	ME EQUIPMENT, its parts, and components producing sparks in NORMAL USE and CONDITION complied with temperature requirements of G.5.2, and U_{max} and I_{max} occurring in their circuits, and complied as follows:		N/A
	Measured $U_{max} \le U_{zR}$ with I_{zR} as in Fig. G.1:		N/A
	Measured U _{max} ≤ U _c with C _{max} as in Fig. G.2:		N/A
	Measured $I_{max} \le I_{zR}$ with U_{zR} as in Fig G.1:		N/A
	Measured $I_{max} \le I_{zL}$ with L_{max} and a $U_{max} \le 24 \text{ V}$ as in Fig G.3:		N/A
	 Combinations of currents and corresponding voltages within the limitations IzR.UzR ≤ 50 W extrapolated from Fig G.1 		N/A
	No extrapolation made for voltages above 42 V		N/A
	 Combinations of capacitances and corresponding voltages within limitations of C/2U² ≤ 1.2 mJ extrapolated from Fig G.2 		N/A
	No extrapolation made for voltages above 242V		N/A
	U _{max} determined using actual resistance R		N/A
	- Combinations of currents and corresponding inductances within limitations L/2l² \leq 0.3 mJ extrapolated from Fig G.3		N/A
	No extrapolation made for inductances larger than 900 mH		N/A
	- U _{max} was the highest supply voltage occurring in circuit under investigation with sparking contact open		N/A
	 I_{max} was the highest current flowing in circuit under investigation with sparking contact closed 		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	- C _{max} and L _{max} taken as values occurring at the component under investigation producing sparks		N/A
	- Peak value considered when a.c. supplied		N/A
	– An equivalent circuit calculated to determine equivalent max capacitance, inductance, and equivalent U_{max} and I_{max} , either as d.c. or a.c. peak values in case of a complicated circuit:		N/A
	Temperature measurements made according to 11.1, and U_{max} , I_{max} , R , L_{max} , and C_{max} determined with application of Figs G.1-G.3:		N/A
	Alternatively, compliance was verified by examination of design data		N/A
3.5.4	External ventilation with internal overpressure		N/A
	ME EQUIPMENT, its parts, and components enclosed in an ENCLOSURE with external ventilation by means of internal overpressure complied with the following requirements:		N/A
	a) FLAMMABLE AESTHETIC MIXTURES WITH AIR t removed by ventilation before EQUIPMENT energized,		N/A
	b) Overpressure inside ENCLOSURE was 75 Pa, min., in NORMAL CONDITION (Pa):		N/A
	Overpressure maintained at the site of potential ignition		N/A
	ME EQUIPMENT could be energized only after the required minimum overpressure was present long enough to ventilate the ENCLOSURE		N/A
	ME EQUIPMENT energized at will or repeatedly when overpressure was continuously present		N/A
	c) Ignition sources de-energized automatically when during operation overpressure dropped below 50 Pa (Pa):		N/A
	d) External surface of ENCLOSURE did not exceed 150 °C in 25 °C:		N/A
3.5.5	ENCLOSURES with restricted breathing		N/A
	ME EQUIPMENT, its parts, and components enclosed in an ENCLOSURE with restricted breathing complied with the following:		N/A
	a) A FLAMMABLE AESTHETIC MIXTURE WITH AIR did not form inside ENCLOSURE with restricted breathing		N/A
	b) Gasket or sealing material used to maintain tightness complied with aging test B-b of IEC 60068-2-2, Clause 15, at 70 $^{\circ}$ C ± 2 $^{\circ}$ C and 96 h :		N/A



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Clause	Requirement + Test	Result - Remark	Verdict
	c) Gas-tightness of ENCLOSURE containing inlets for flexible cords maintained		N/A
	Cords are fitted with adequate anchorages to limit stresses as determined by test		N/A
	Overpressure not reduced below 200 Pa		N/A
	Tests waived when examination of ENCLOSURE indicated it is completely sealed or gas-tight without a doubt (100 % degree of certainty)		N/A
	Operating temperature of external surface of ENCLOSURE was ≤ 150 °C in 25 °C (°C):		N/A
	Steady state operating temperature of ENCLOSURE also measured (°C):		N/A
G.6	CATEGORY APG ME EQUIPMENT, parts and componer	nts thereof	N/A
G.6.1	ME EQUIPMENT, its parts, and components did not ignite FLAMMABLE AESTHETIC MIXTURE WITH OXYGEN OR NITROUS OXIDE under NORMAL USE and SINGLE FAULT CONDITION		N/A
	ME EQUIPMENT, its parts, and components not complying with G.6.3 subjected to a CONTINUOUS OPERATION test		N/A
G.6.2	Parts and components of CATEGORY APG ME EQUIPMENT operating in a FLAMMABLE AESTHETIC MIXTURE WITH OXYGEN OR NITROUS OXIDE supplied from a source isolated from earth by insulation equal to one MEANS OF PATIENT PROTECTION and from electrical parts by insulation twice the MEANS OF PATIENT PROTECTION:		N/A
G.6.3	Test of G.6.1 waived when the following requirements were met in NORMAL USE and under NORMAL and SINGLE FAULT CONDITIONS:		N/A
	a) no sparks produced and temperatures did not exceed 90 °C, or		N/A
	b) a temperature limit of 90 °C not exceeded, sparks produced in NORMAL USE, and SINGLE FAULT CONDITIONS, except U_{max} and I_{max} occurring in their circuits complied with requirements, taking C_{max} and L_{max} into consideration:		N/A
	Measured U _{max} ≤ U _{zR} with I _{zR} as in Fig. G.4:		N/A
	Measured U _{max} ≤ U _z c with C _{max} as in Fig. G.5:		N/A
	Measured I _{max} ≤ I _{zR} with U _{zR} as in Fig G.4:		N/A
	Measured $I_{max} \le I_{zL}$ with L_{max} and a $U_{max} \le 24$ V as in Fig G.6:		N/A
	- Extrapolation from Figs G.4, G.5, and G.6 was limited to areas indicated		N/A



	IEC 60601-1	·	
Clause	Requirement + Test	Result - Remark	Verdict
	- U _{max} was the highest no-load voltage occurring in the circuit under investigation, taking into consideration mains voltage variations as in 4.10		N/A
	- I _{max} was the highest current flowing in the circuit under investigation, taking into account MAINS VOLTAGE variations as in 4.10		N/A
	- C _{max} and L _{max} are values occurring in relevant circuit		N/A
	– U_{max} additionally determined with actual resistance R when equivalent resistance R in Fig G.5 was less than 8000 Ω		N/A
	- Peak value considered when a.c. supplied		N/A
	- An equivalent circuit calculated to determine max capacitance, inductance, and U _{max} and I _{max} , either as d.c. or a.c. peak values in case of a complicated circuit:		N/A
	- When energy produced in an inductance or capacitance in a circuit is limited by voltage or current-limiting devices, two independent components applied, to obtain the required limitation even when a first fault (short or open circuit) in one of these components		N/A
	- requirement not applied to transformers complying with this standard		N/A
	- requirement not applied to wire-wound current-limiting resistors provided with a protection against unwinding of the wire in case of rupture		N/A
	Compliance verified by examination of CATEGORY APG ME EQUIPMENT, parts, and components, or		N/A
	Temperature measurements made in accordance with 11.1:		N/A
	- or U _{max} , I _{max} , R, L _{max} and C _{max} determined together with application of Figs G.4-G.6:		N/A
	Alternatively, compliance verified by comparison with design data:		N/A
G.6.4	ME EQUIPMENT, its parts, and components heating a FLAMMABLE AESTHETIC MIXTURE WITH OXYGEN OR NITROUS OXIDE provided with a non-SELF-RESETTING THERMAL CUT-OUT and complied with 15.4.2.1		N/A
	Current-carrying part of heating element is not in direct contact with FLAMMABLE AESTHETIC MIXTURE WITH OXYGEN OR NITROUS OXIDE		N/A



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	IEC 60601	I-1	
Clause	Requirement + Test	Result - Remark	Verdict
G.7	Test apparatus for flammable mixtures according to this Clause and Fig G.7		N/A

ANNEX L	INSULATED WINDING WIRES FOR USE WITHOUNSULATION	N/A	
L.1	BASIC, SUPPLEMENTARY, DOUBLE, and REINFORCED INSULATION in wound components without interleaved insulation complied with this Annex		N/A
L.2	Wire construction		N/A
	Overlap of layers when wire is insulated with two or more spirally wrapped layers of tape is adequate to ensure continued overlap during manufacture of wound component		N/A
	Layers of spirally wrapped wire insulation are sufficiently secured to maintain the overlap		N/A
L.3	Type Test		N/A
	The wire subjected to tests of L.3.1 to L.3.4 at a temperature and a relative humidity specified		N/A
	Temperature (°C):		_
	Humidity (%):		_
L.3.1	Dielectric strength		N/A
	Dielectric strength test of Clause 8.8.3 for the appropriate type and number of MOP(s) conducted with no breakdown:		N/A
	- 3000 V for BASIC and SUPPLEMENTARY INSULATION (V):		N/A
	- 6000 V for REINFORCED INSULATION (V):		N/A
L.3.2	Flexibility and adherence		N/A
	Sample subjected to flexibility and adherence		N/A
	Sample examined per IEC 60851-3: 1997, cl. 5.1.1.4, followed by dielectric test of cl. 8.8.3, with no breakdown		N/A
	Test voltage was at least the voltage in Tables 6 and 7 but not less than the following:		N/A
	- 1500 V for BASIC and SUPPLEMENTARY INSULATION (V)		N/A
	- 3000 V for REINFORCED INSULATION (V):		N/A
	Tension applied to wire during winding on mandrel calculated from the wire diameter equivalent to 118 MPa ± 11.8 MPa		N/A
L.3.3	Heat Shock		N/A

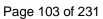


IEC 60601-1				
Clause	Requirement + Test	Result - Remark	Verdict	
	Sample subjected to heat shock test 9 of IEC 60851-6:1996, followed by dielectric strength test of clause 8.8.3		N/A	
	Test voltage was at least the voltage in Tables 6 and 7, but not less than the following:		N/A	
	- 1500 V for BASIC and SUPPLEMENTARY INSULATION (V):		N/A	
	- 3000 V for REINFORCED INSULATION (V):		N/A	
	Oven temperature based on Table L.2 (°C):		_	
	Mandrel diameter and tension applied as in clause L.3.2, (MPa; N/mm²):		N/A	
	Dielectric strength test conducted at room temperature after removal from the oven		N/A	
L.3.4	Retention of electric strength after bending		N/A	
	Five samples prepared as in L.3.2 subjected to dielectric strength and bending tests		N/A	
	Test voltage was at least the voltage in Tables 6 and 7, but not less than the following:		N/A	
	- 1500 V for BASIC and SUPPLEMENTARY INSULATION (V):		N/A	
	- 3000 V for REINFORCED INSULATION (V)::		N/A	
	Test voltage applied between the shot and conductor		N/A	
	Mandrel diameter and tension applied as in L.3.2, (MPa; N/mm²):		N/A	
L.4	Tests during manufacture		N/A	
L.4.1	Production line dielectric strength tests done by the manufacture per L.4.2 and L.4.3:		N/A	
L.4.2	Test voltage for routine testing (100 % testing) is at least the voltage in Tables 6 and 7 but not less than the following:		N/A	
	- 1500 V r.m.s. or 2100 V peak for BASIC and SUPPLEMENTARY INSULATION (V)		N/A	
	- 3000 V r.m.s. or 4200 V peak for REINFORCED INSULATION (V)		N/A	
L.4.3	Sampling tests conducted using twisted pair samples (IEC 60851-5:1996, clause 4.4.1):		N/A	
	Minimum breakdown test voltage at least twice the voltage in Tables 6 and 7 but not less than:		N/A	
	- 3000 V r.m.s. or 4200 V peak for BASIC and SUPPLEMENTARY INSULATION:		N/A	
	- 6000 V r.m.s. or 8400 V peak for REINFORCED INSULATION:		N/A	



		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

4.2.2	RM RESULTS TABLE: General requirements for RISK MANAGEMENT			Р
Clause of ISO	Document Ref. in RN paragraph/clause, ve		Result - Remarks	Verdict
14971	General process	Particular Medical Device		
3.1	GTQPR05000; 2017-1-13; cl. 6.1	_	Risk Management Process (excluding production and post-production)	Р
3.2	GTQPR05000; 2017-1-13; cl. 4	_	Adequate Resources —	
3.2	GTQPR05000; 2017-1-13; cl. 4	_	Assignment of qualified personnel	Р
3.2	GTQPR05000; 2017-1-13; cl. 5	_	Policy for determining criteria for risk acceptability	Р
3.3	_	GTQPR05000; 2017-1-13; cl. 4	Qualification of personnel provided.	
3.4a	_	GTQPR05000; Document name : Attachment B: Risk Management Plan for Device GT-500160-30 REV1		Р
3.4b	_	GTQPR05000; 2017-1-13; cl. 6.1	Document name : Attachment B: Risk Management Plan for Device GT-500160-30 REV1	Р
3.4c	_	GTQPR05000; 2017-1-13; cl. 6.1	Document name : Attachment B: Risk Management Plan for Device GT-500160-30 REV1	Р
3.4d	_	GTQPR05000; 2017-1-13; cl. 6.1	Document name : Attachment B: Risk Management Plan for Device GT-500160-30 REV1	P
3.4e	_	GTQPR05000; 2017-1-13; cl. 6.1	Document name : Attachment B: Risk Management Plan for Device GT-500160-30 REV1	Р
3.5	_		Document name :	Р
			Risk Management Report; GT- RM2019-001 contains all relevant attachments:	
		GTQPR05000; 2017-1-13;	Attachment A: ISO 14971 Gap Analysis Checklist	
		cl. 4	Attachment B: Risk Management Plan for Device GT-500160-30 REV1	
			Attachment C: Risk Management Procedure	





IEC 60601-1				
Clause	Requirement + Test		Result - Remark	Verdict

4.2.2	RM RESULTS TABL	E: General requirements	for RISK MANAGEMENT	Р
Clause of ISO	Document Ref. in RM paragraph/clause, ve		Result - Remarks	Verdict
14971	General process	Particular Medical Device		
4.1	_	GTQPR05000; 2017-1-13;	Risk analysis performed for particular medical device.	Р
		cl. 6	A description and identification that was analysed is stated.	
			Identification of the person(s) and organization that carried out the risk analysis is stated.	
			The scope and date of risk analysis is stated.	
			Document name : Risk Management Report; GT-RM2019-001	
4.2	1	GTQPR05000; Intended use and identification of characteristics related to safety stated.		Р
			Document name : Risk Management Report; GT-RM2019-001	
4.3			Identification of hazards conducted during risk analysis.	Р
		cl. 6	Document name : Risk Management Report; GT-RM2019-001	
4.4	_	GTQPR05000; 2017-1-13;	Estimation of risk conducted during risk analysis.	Р
		cl. 6	Document name : Risk Management Report; GT-RM2019-001	
5	-	GTQPR05000; 2017-1-13;	Risk evaluation conducted during risk analysis.	Р
		cl. 6	Document name : Risk Management Report; GT-RM2019-001	
6.2	_	GTQPR05000; 2017-1-13;	Risk reduction conducted during risk analysis.	Р
		cl. 6.2.3 - 6.2.5	Document name : Risk Management Report; GT-RM2019-001	
6.3	_	GTQPR05000; 2017-1-13; cl. 6.1	Implementation of risk control measure(s) conducted during risk analysis.	Р
			Document name : Risk Management Report; GT-RM2019-001	



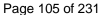
		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

4.2.2	RM RESULTS TABLE: General requirements for RISK MANAGEMENT			Р
Clause of ISO	Document Ref. in RI paragraph/clause, ve		Result - Remarks	Verdict
14971	General process	Particular Medical Device		
6.4	_	GTQPR05000; 2017-1-13;	Residual risk evaluation conducted during risk analysis.	Р
		cl. 6.2.6	Document name : Risk Management Report; GT-RM2019-001	
6.5	_	GTQPR05000; No Risk/benefit analysis requir		Р
		2017-1-13; The residual risk was acceptable.		
		CI. U.Z.3	Document name : Risk Management Report; GT-RM2019-001	
6.6a	_	GTQPR05000; 2017-1-13; cl. 6	There were no new risks identified after implementation of risk-reducing measures.	P
			Document name : Risk Management Report; GT-RM2019-001	
6.6b		GTQPR05000; 2017-1-13; cl. 6	Previously identified hazardous situations were not affected by the introduction of risk control measures.	Р
			Document name : Risk Management Report; GT-RM2019-001	
6.7	_	GTQPR05000; 2017-1-13;	Completeness of risk control conducted during risk analysis.	Р
		cl. 6	Document name : Risk Management Report; GT-RM2019-001	
7	-	GTQPR05000; 2017-1-13;	Evaluation of overall risk acceptability conducted.	Р
		cl. 6.2.6	Document name : Risk Management Report; GT-RM2019-001	
8	_	GTQPR05000; 2017-1-13;	A risk management report was prepared by the manufacturer.	Р
		cl. 6	Document name : Risk Management Report; GT-RM2019-001	

Supplementary Information:

Document Ref should be with regards to the policy/procedure documents and documents containing device specific output.

4.3	TABLE: ESSENTIAL PERFORMANCE			N/A
List of ESSENTIAL PERFORMANCE functions		MANUFACTURER'S document number reference or reference from this standard or collateral or particular standard(s)	Remarks	
		-		



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	IEC 60601-1	· · · · · · · · · · · · · · · · · · ·	
	IEC 00001-1		
Clause	Requirement + Test	Result - Remark	Verdict

4.3 TABLE: ESSENTIAL PERFORMANCE			N/A	
List of ESS PERFORMAN	ENTIAL NCE functions	MANUFACTURER'S document number reference or reference from this standard or collateral or particular standard(s)	Remarks	

Supplementary Information:

ESSENTIAL PERFORMANCE is performance, the absence or degradation of which, would result in an unacceptable risk.

Power supply unit is not end medical product; therefore no essential performance defined by the manufacturer.

4.11	TABLE: Power Input					Р
Operating	Conditions / Ratings	Voltage (V)	Frequency (Hz)	Current (mA)	Power (W)	Power factor (cos φ)
		5	0 Hz			
Rated outp	out load.	90	50	1,26	70,6	
Rated outp	out load.	100	50	1,16	69,7	
Rated outp	out load.	120	50	1,01	68,9	
Rated outp	out load.	132	50	0,94	68,6	
Rated outp	out load.	180	50	0,76	68,4	
Rated outp	out load.	200	50	0,70	68,2	
Rated outp	out load.	220	50	0,65	67,9	
Rated outp	out load.	230	50	0,63	67,6	
Rated outp	out load.	240	50	0,61	67,5	
Rated outp	out load.	264	50	0,57	67,8	
		6	0 Hz			
Rated outp	out load.	90	60	1,27	70,4	
Rated outp	out load.	100	60	1,17	69,7	
Rated outp	out load.	120	60	1,02	68,9	
Rated outp	out load.	132	60	0,95	68,6	
Rated outp	out load.	180	60	0,76	68,3	
Rated outp	out load.	200	60	0,71	68,1	
Rated outp	out load.	220	60	0,66	67,8	
Rated outp	out load.	230	60	0,63	67,5	
Rated outp	out load.	240	60	0,61	67,4	
Rated outp	out load.	264	60	0,58	67,7	

Supplementary Information:

Rated supply voltage: 100-240 Vac Rated supply frequency: 50-60 Hz



		1 agc 100 01 25	•	Report No. 12	20 000-/10
		IEC 60601-1			
Clause	Requirement + Test		Result - Remark		Verdict
4.11	TABLE: Power Input				Р
Rated inp	out current: 1,6-0,7 A				
See copy	of marking plate for details.				

5.9.2	TABLE: Determ	E: Determination of ACCESSIBLE parts						
Location		Determination method (NOTE1)	Comments					
EUT enclosure		Visual	Live parts not accessible.					
			Enclosure without opening to cover all live parts.	s provided				
			No doubts; therefore only vinspection performed.	risual				

Supplementary information:

¹⁾ NOTE: The determination methods are: visual; rigid test finger; jointed test finger; test hook.

EUT is provided approved appliance inlet.

During normal operation mains supply terminals are not accessible.

7.1.2	TABLE: Legibility of Marking							
Markings tested		Ambient Illuminance (lx)	Remarks					
Outside Ma	arkings (Clause 7.2):	970 lx	Marking plate					
Inside Mar	kings (Clause 7.3)		No markings to be read inside					
Controls &	Instruments (Clause 7.4):		No controls & instrument	S				
Safety Sign	ns (Clause 7.5):		No safety signs					
Symbols (0	Clause 7.6):	970 lx	Present on Marking plate	;				

Supplementary information:

Observer, with a visual acuity of 0 on the log Minimum Angle of Resolution (log MAR) scale or 6/6 (20/20) and is able to read N6 of the Jaeger test card in normal room lighting condition (~500lx), reads marking at ambient illuminance least favourable level in the range of 100 lx to 1,500 lx. The ME EQUIPMENT or its part was positioned so that the viewpoint was the intended position of the OPERATOR or if not defined at any point within the base of a cone subtended by an angle of 30° to the axis normal to the centre of the plane of the marking and at a distance of 1 m.

EUT is desktop power supply. Markings are provided on the enclosure surface and shall be visible from one of the normal positions.

In additional, all information provided within technical specifications.

7.1.3	TABLE: Durability of marking test						
Characteristics of the Marking Label tested:							
Material of	Marking Label:	In-print method	ı	Pass			
Ink/other pr	inting material or process:	Laser printing	ı	Pass			
Material (co	omposition) of Warning Label:			N/A			



	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict

7.1.3	TABLE: Durability of marking test				
Characteri		Remarks			
Ink/other pr	N/A				
Other	N/A				
		Remarks			
In-print met					

Supplementary information:

Marking rubbed by hand, first for 15 s with a cloth rag soaked with distilled water, then for 15 s with a cloth rag soaked with ethanol 96%, and then for 15 s with a cloth rag soaked with isopropyl alcohol.

Marking provided on the outer side of the power supply. See copy of marking plate and enclosed pictures of the unit for details.

8.4.2	TABLE: TABL		Р				
Test supply voltage/frequency (V/Hz) ¹⁾ :							/ 50 Hz
Location		I					
From/To	Vrms	Vpk or Vdc	Peak-to- peak ripple ²⁾	Power W/VA	Energy (J)	Remarks	
Measured between output minu and output plus		29,73 Vdc		62,30 VA		Maximum output current achieved: 2,2 A	
R13 short o	circuit						
Measured between output minu and output plus	IS	29,73 Vdc		177,2 VA		Maximum out achieved: 5,96 A	put current

Supplementary Information:

Output voltage during single fault conditions was recorded. Output voltage was below SELV limit. See table 13.2: TABLE: single fault conditions in accordance with 13.2.2 to 13.2.13, inclusive.

In additional, evaluation of voltage limiting components in SELV circuits performed. See table: evaluation of voltage limiting components in SELV circuits.

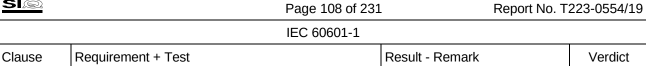
Output power less than 240 VA in normal and single fault conditions.

8.4.3	TABLE: ME EQUIPMENT for connection to a power source by a plug - measurement of voltage or calculation of stored charge 1 s after disconnection of plug from mains supply				
Maxin					

¹⁾The input supply voltage to the ME EQUIPMENT was the RATED voltage or the voltage within the RATED voltage range which results in the highest measured value. See clause 8.5.4.

^{2).} If the d.c peak-to-peak ripple >10%, waveform considered as a.c. See clause 8.4.2.2





·									•	
Voltage measured (V)										
Voltage Measured Between:	1	2	3	4	5	6	7	8	9	10
Plug pins 1 and 2	31	28								
Plug pin 1 and plug earth pin										
Plug pin 2 and plug earth pin										
Plug pin 1 and enclosure										
Plug pin 2 and enclosure										
Maximum allowable stored c	harge w	hen me	easured	voltage	e excee	ded 60 v	v (μc)	: 45		
		Calcula	ated sto	red cha	rge (μc))				
Voltage Measured Between:	1	2	3	4	5	6	7	8	9	10
Plug pins 1 and 2										
Plug pin 1 and plug earth pin										
Plug pin 2 and plug earth pin										
Plug pin 1 and enclosure										
Plug pin 2 and enclosure										

Supplementary information:

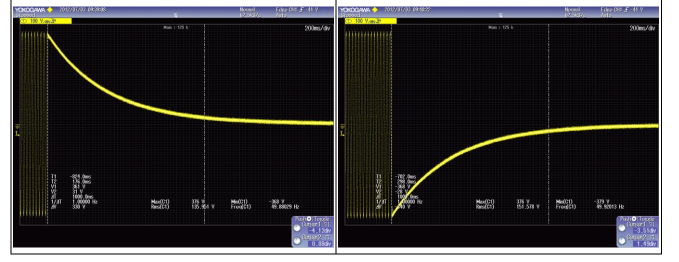
Supply voltage: 264 Vac.

SI®

Trigger starts at peak voltage, therefore no need to be done 10 measurements.

Voltage measured betweens Plug pins 1 and 2 - 1

Voltage measured betweens Plug pins 1 and 2 - 2



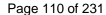


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		IEC 60601-	1				
Clause	Requirement + Test		Result - Remark		Verdict		
8.4.4	.4.4 TABLE: Internal capacitive circuits – measurement of residual voltage or calculation of the stored charge in capacitive circuits (i.e., accessible capacitors or circuit parts) after de-energizing ME EQUIPMENT						
Maximum a	illowable residual voltage (V	<u> </u>	······································	60 V			
Maximum a	Illowable stored charge whe	n residual voltage exc	eeded 60 V:	45 μC			
Description of the capacitive circuit (i.e., accessible capacitor or circuit parts) Measured residual voltage (V) charge (μC)					arks		
Suppleme	ntary information: \			•			

	5.1a TABLE: defibrillation-proof applied parts – measurement of hazardous N/A electrical energies						
Test Condition: Figs. 9 & 1		Applied part with test voltage	Test voltage polarity	Measured voltage between Y1 and Y2 (mV)	Remarks		
Supplemen	Supplementary information: \						

8.5.5.1b	TABLE: defibr	TABLE: defibrillation-proof applied parts – verification of recovery time N/A						
Applied part with test voltage Recovery time from polarity documents (s)		Measured Remar recovery time (s)		narks				
Suppleme	Supplementary information: \							

8.5.5.2 TABLE: DEFIBRILLATION-PROOF APPLIED PARTS or PATIENT CONNECTIONS of DEFIBRILLATION-PROOF APPLIED PARTS - Energy reduction test –measurement of Energy delivered to a 100 Ω load					
	Test Voltage applied to	Measured Energy E1 (mJ)	Measured Energy E2 (mJ)	Energy E1 as % of E2 (%)	
PATIENT CON	NNECTION 1 or APPLIED PART with INECTIONS 2, 3, and 4 of the same T connected to earth		-		
PATIENT CON	NNECTION 2 or APPLIED PART with INECTIONS 1, 3, and 4 of the same T connected to earth		-		
PATIENT CON	NNECTION 3 or APPLIED PART with INECTIONS 1, 2, and 4 of the same T connected to earth		-		
PATIENT CON	NNECTION 4 or APPLIED PART with INECTIONS 1, 2, and 3 of the same T connected to earth				





	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict

Supplementary information:

For compliance: E1 must at least 90% of E2

E1= Measured energy delivered to 100 Ω with ME Equipment connected; E2= Measured energy delivered to 100 Ω without ME equipment connected.

8.6.4 TABLE: Impedance and current-carrying capability of PROTECTIVE EARTH CONNECTIONS					Р
	of ME EQUIPMENT & impedance neasured between parts	Test current (A) /Duration (s)	Voltage drop measured between parts (V)	Maximum calculated impedance (mΩ)	Maximum allowable impedance (mΩ)
impedance l	LY INSTALLED ME EQUIPMENT, between PROTECTIVE EARTH d a PROTECTIVELY EARTHED part		1	1	100
	NT with an APPLIANCE INLET,	120 sec	0,143V	3,6	100
	between earth pin in the APPLIANCE PROTECTIVELY EARTHED part	40 A			
		60 sec	0,079V	3,1	100
		25 A			
SUPPLY CORI	NT with a non-DETACHABLE POWER D, impedance between the arth pin in the MAINS PLUG and a LY EARTHED part			-	200

Supplementary information:

PERMANENTLY INSTALLED ME EQUIPMENT, impedance between PROTECTIVE EARTH TERMINAL and a PROTECTIVELY EARTHED part - Limit 100 m Ω ME EQUIPMENT with an APPLIANCE INLET, impedance between earth pin in the APPLIANCE INLET and a PROTECTIVELY EARTHED part - Limit 100 m Ω

ME EQUIPMENT with an APPLIANCE INLET, impedance between earth pin in the protective earth pin on the DETACHABLE POWER SUPPLY CORD and a PROTECTIVELY EARTHED part - Limit 200 m Ω

ME EQUIPMENT with a non-DETACHABLE POWER SUPPLY CORD, impedance between the protective earth pin in the MAINS PLUG and a PROTECTIVELY EARTHED part - Limit 200 m Ω

8.7	3.7 TABLE: leakage current					Р	
Type of leakage current and test condition (including single faults)		Supply voltage (V)	Supply frequency (Hz)	Measured max. value (µA)	Remark	s	
Fig 12 Fo	urth Leakage (ER)				Maximum allowed val	ues:	
rig. 13 - Ea	iiiii Leakaye (EN)	age (ER) — — —		_	5 mA NC; 10 mA SFC		
	Before humidity treatment (Frequency weighted)						
	dition, normal polarity, idity treatment;	264	60	116,3			
	dition, reverse polarity, idity treatment;	264	60	116,6			
	condition (supply), normal polarity, before eatment;	264	60	229,8			



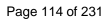
		Pa	ge 111 of 2	31	Repon	t No. T223-0554/19
		IEC	60601-1			
Clause	Requirement + Test			Result - Re	mark	Verdict
	t condition (supply n), reverse polarity, before eatment;	264	60	228,6		
		Rev	vision 1.0			
	Before hu	midity treat	ment (Freq	uency weigh	ted)	
	ndition, normal polarity, nidity treatment;	264	60	160,0		
	ndition, reverse polarity, nidity treatment;	264	60	172,0		
	t condition (supply n), normal polarity, before eatment;	264	60	298,0		
	t condition (supply n), reverse polarity, before eatment;	264	60	298,0		
	After hur	nidity treatn	nent (Frequ	uency weight	ed)	
	ndition, normal polarity, nidity treatment;	264	60	169,5		
	ndition, reverse polarity, nidity treatment;	264	60	167,3		
	t condition (supply n), normal polarity, before eatment;	264	60	298,3		
	t condition (supply n), reverse polarity, before eatment;	264	60	296,7		
	Before hum	idity treatmo	ent (Non-fr	equency weig	ghted)	
	ndition, normal polarity, nidity treatment;	264	60	162,0		
	ndition, reverse polarity, nidity treatment;	264	60	175,0		
	t condition (supply n), normal polarity, before eatment;	264	60	302,0		
	t condition (supply n), reverse polarity, before eatment;	264	60	301,0		
	After humi	dity treatme	nt (Non-fre	quency weig	hted)	
	ndition, normal polarity, nidity treatment;	264	60	185,7		
	ndition, reverse polarity, nidity treatment;	264	60	180,5		
	t condition (supply n), normal polarity, before eatment;	264	60	319,4		



	IEC	60601-1			
Clause Requirement + Test			Result - Re	emark	Verdict
Single fault condition (supply interruption), reverse polarity, before humidity treatment;	264	60	317,5		
Fig. 14 - Touch Current (TC)	_	_	_	Maximum allowed va 100 μA NC; 500 μA	
With frequency	weighted d	evice (before	e humidity tre	eatment)	
Normal condition, normal polarity, before humidity treatment	264	60	11,2	Measured on outp	ut plus
Normal condition, reverse polarity, before humidity treatment	264	60	11,1	Measured on outp	ut plus
Single fault condition (supply interruption), normal polarity, before humidity treatment	264	60	16,6	Measured on outp	ut plus
Single fault condition (supply interruption), reverse polarity, before humidity treatment	264	60	16,5	Measured on outp	ut plus
Single fault condition (PE interruption), normal polarity, before humidity treatment	264	60	39,1	Measured on outp	ut plus
Single fault condition (PE interruption), reverse polarity, before humidity treatment	264	60	39,6	Measured on outp	ut plus
With frequency	weighted d	evice (before	e humidity tre	eatment)	
Normal condition, normal polarity, before humidity treatment	264	60	11,14	Measured on outp	ut minus
Normal condition, reverse polarity, before humidity treatment	264	60	11,18	Measured on outp	ut minus
Single fault condition (supply interruption), normal polarity, before humidity treatment	264	60	16,64	Measured on outp	ut minus
Single fault condition (supply interruption), reverse polarity, before humidity treatment	264	60	16,56	Measured on outp	ut minus
Single fault condition (PE interruption), normal polarity, before humidity treatment	264	60	40,06	Measured on outp	ut minus
Single fault condition (PE interruption), reverse polarity, before humidity treatment	264	60	39,15	Measured on outp	ut minus
	Rev	ision 1.0			
With frequency	weighted d	evice (before	e humidity tre	eatment)	
Normal condition, normal polarity, before humidity treatment	264	60	7,8	Measured on outp	ut plus
Normal condition, reverse polarity, before humidity treatment	264	60	8,1	Measured on outp	ut plus



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Clause	Requirement + Test			Result - Re	emark	Verdict
0.000	1.040			1.1000		10.0.01
	condition (supply), normal polarity, before eatment	264	60	12,2	Measured on outp	ut plus
	condition (supply), reverse polarity, before eatment	264	60	12,2	Measured on outpo	ut plus
	condition (PE interruption), arity, before humidity	264	60	35,8	Measured on outp	ut plus
	condition (PE interruption), arity, before humidity	264	60	39,2	Measured on outpo	ut plus
		- I			<u> </u>	
	dition, normal polarity, idity treatment	264	60	7,7	Measured on outp	ut minus
	dition, reverse polarity, idity treatment	264	60	8,1	Measured on outpo	ut minus
	condition (supply), normal polarity, before eatment	264	60	12,3	Measured on outpo	ut minus
	condition (supply), reverse polarity, before eatment	264	60	12,2	Measured on outpo	ut minus
	condition (PE interruption), arity, before humidity	264	60	35,5	Measured on outpo	ut minus
	condition (PE interruption), arity, before humidity	264	60	37,3	Measured on outp	ut minus
	dition, normal polarity, idity treatment	264	60	5,7	Measured on plast (metal foil used)	ic enclosure
	dition, reverse polarity, idity treatment	264	60	5,9	Measured on plast (metal foil used)	ic enclosure
	condition (supply), normal polarity, before eatment	264	60	8,5	Measured on plast (metal foil used)	ic enclosure
	condition (supply), reverse polarity, before eatment	264	60	8,5	Measured on plast (metal foil used)	ic enclosure
	condition (PE interruption), arity, before humidity	264	60	8,0	Measured on plast (metal foil used)	ic enclosure
	condition (PE interruption), arity, before humidity	264	60	7,8	Measured on plast (metal foil used)	ic enclosure





	IEC 606	01-1	
Clause	Requirement + Test	Result - Remark	Verdict

With frequency wei	ghted devi	ce (after hur	nidity treatme	ent – 2 days)
Normal condition, normal polarity, after humidity treatment	264	60	8,6	Measured on output plus
Normal condition, reverse polarity, after humidity treatment	264	60	8,5	Measured on output plus
Single fault condition (supply interruption), normal polarity, after humidity treatment	264	60	12,9	Measured on output plus
Single fault condition (supply interruption), reverse polarity, after humidity treatment	264	60	12,8	Measured on output plus
Single fault condition (PE interruption), normal polarity, after humidity treatment	264	60	38,4	Measured on output plus
Single fault condition (PE interruption), reverse polarity, after humidity treatment	264	60	37,1	Measured on output plus
Normal condition, normal polarity, after humidity treatment	264	60	8,5	Measured on output minus
Normal condition, reverse polarity, after humidity treatment	264	60	8,4	Measured on output minus
Single fault condition (supply interruption), normal polarity, after humidity treatment	264	60	12,8	Measured on output minus
Single fault condition (supply interruption), reverse polarity, after humidity treatment	264	60	12,8	Measured on output minus
Single fault condition (PE interruption), normal polarity, after humidity treatment	264	60	38,1	Measured on output minus
Single fault condition (PE interruption), reverse polarity, after humidity treatment	264	60	37,6	Measured on output minus
Normal condition, normal polarity, after humidity treatment	264	60	6,3	Measured on plastic enclosure (metal foil used).
Normal condition, reverse polarity, after humidity treatment	264	60	6,3	Measured on plastic enclosure (metal foil used).
Single fault condition (supply interruption), normal polarity, after humidity treatment	264	60	8,9	Measured on plastic enclosure (metal foil used).
Single fault condition (supply interruption), reverse polarity, after humidity treatment	264	60	8,9	Measured on plastic enclosure (metal foil used).
Single fault condition (PE interruption), normal polarity, after humidity treatment	264	60	8,0	Measured on plastic enclosure (metal foil used).



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Clause Requirement + Test			Result - Re	emark	Verdict
Single fault condition (PE interruption), reverse polarity, after humidity treatment	264	60	8,4	Measured on plast (metal foil used).	ic enclosure
With non-frequer	ncy weighted	device (be	fore humidity	treatment)	
Normal condition, normal polarity, before humidity treatment	264	60	243,9	Measured on outp	ut plus
Normal condition, reverse polarity, before humidity treatment	264	60	244,0	Measured on outp	ut plus
Single fault condition (supply interruption), normal polarity, before humidity treatment	264	60	243,4	Measured on outpo	ut plus
Single fault condition (supply interruption), reverse polarity, before humidity treatment	264	60	13,9	Measured on outpo	ut plus
Single fault condition (PE interruption), normal polarity, before humidity treatment	264	60	160,5	Measured on outpo	ut plus
Single fault condition (PE interruption), reverse polarity, before humidity treatment	264	60	161,1	Measured on outpo	ut plus
Normal condition, normal polarity, before humidity treatment	264	60	244,6	Measured on outpo	ut minus
Normal condition, reverse polarity, before humidity treatment	264	60	244,7	Measured on outpo	ut minus
Single fault condition (supply interruption), normal polarity, before humidity treatment	264	60	243,8	Measured on outpo	ut minus
Single fault condition (supply interruption), reverse polarity, before humidity treatment	264	60	13,8	Measured on outpo	ut minus
Single fault condition (PE interruption), normal polarity, before humidity treatment	264	60	161,1	Measured on outpo	ut minus
Single fault condition (PE interruption), reverse polarity, before humidity treatment	264	60	161,7	Measured on outpo	ut minus
Manual and Britania	004		100-	1.4	
Normal condition, normal polarity, before humidity treatment	264	60	106,5	Measured on plast (metal foil used)	
Normal condition, reverse polarity, before humidity treatment	264	60	106,6	Measured on plast (metal foil used)	ic enclosure
Single fault condition (supply interruption), normal polarity, before humidity treatment	264	60	105,5	Measured on plast (metal foil used)	ic enclosure



		IEC	60601-1				
Clause	Requirement + Test			Result - Re	emark	Verdict	
	condition (supply), reverse polarity, before eatment	264	60	12,7	Measured on plast (metal foil used)	ic enclosure	
	condition (PE interruption), arity, before humidity	264	60	86,8	Measured on plastic enclosure (metal foil used)		
	condition (PE interruption), arity, before humidity	264	60	86,8	Measured on plast (metal foil used)	ic enclosure	
	With non-frequency w	veighted de	vice (after h	numidity treatr	nent – 2 days)		
Normal con humidity tre	dition, normal polarity, after eatment	264	60	246,6	Measured on outp	ut plus	
Normal cor humidity tre	dition, reverse polarity, after eatment	264	60	246,7	Measured on outp	ut plus	
	condition (supply), normal polarity, after eatment	264	60	245,9	Measured on outp	ut plus	
	condition (supply), reverse polarity, after eatment	264	60	15,3	Measured on outp	ut plus	
	condition (PE interruption), arity, after humidity treatment	264	60	181,5	Measured on output plus		
	condition (PE interruption), arity, after humidity	264	60	180,1	Measured on outp	ut plus	
Normal cor humidity tre	ndition, normal polarity, after eatment	264	60	247,4	Measured on outp	ut minus	
Normal con humidity tre	dition, reverse polarity, after eatment	264	60	247,6	Measured on outp	ut minus	
	condition (supply), normal polarity, after eatment	264	60	246,9	Measured on outp	ut minus	
	condition (supply), reverse polarity, after eatment	264	60	15,4	Measured on outp	ut minus	
	condition (PE interruption), arity, after humidity treatment	264	60	181,7	Measured on outp	ut minus	
	condition (PE interruption), arity, after humidity	264	60	181,0	Measured on outp	ut minus	
Normal cor	ndition, normal polarity, after eatment	264	60	90,7	Measured on plast (metal foil used).	ic enclosure	



	IEC 60601-1	· · · · · · · · · · · · · · · · · · ·	
	IEC 00001-1		
Clause	Requirement + Test	Result - Remark	Verdict

Normal condition, reverse polarity, after	264	60	90,7	Measured on plastic enclosure
humidity treatment				(metal foil used).
Single fault condition (supply interruption), normal polarity, after humidity treatment	264	60	89,9	Measured on plastic enclosure (metal foil used).
Single fault condition (supply interruption), reverse polarity, after humidity treatment	264	60	12,2	Measured on plastic enclosure (metal foil used).
Single fault condition (PE interruption), normal polarity, after humidity treatment	264	60	79,6	Measured on plastic enclosure (metal foil used).
Single fault condition (PE interruption), reverse polarity, after humidity treatment	264	60	80,0	Measured on plastic enclosure (metal foil used).
				Maximum allowed values:
Fig. 15 - Patient Leakage Current (P)	_	_	_	Type B or BF AP: 10 μA NC; 50 μA SFC (d.c. current); 100 μA NC; 500 μA SFC (a.c.) Type CF AP: 10 μA NC; 50 μA SFC (d.c. or a.c. current)
				Maximum allowed values:
Fig. 16 - Patient leakage current with				Type B: N/A
mains on the F-type applied parts (PM)	_	_	_	Type BF AP: 5000 μA
				Type CF AP: 50 μA
				Maximum allowed values:
Fig. 17 - Patient leakage current with external voltage on Signal Input/Output				Type B or BF AP: 10 μA NC; 50 μA SFC(d.c. current);
part (SIP/SOP)	_	_	_	100 μA NC; 500 μA SFC (a.c.) ;
				Type CF AP: 10 μA NC; 50 μA SFC (d.c. or a.c. current)
Fig. 18 - Patient leakage current with				Maximum allowed values:
external voltage on metal Accessible	_	_	_	Type B or BF AP: 500 μA
Part that is not Protectively Earthed				Type CF: N/A
_				Maximum allowed values:
Fig. 19 – Patient Auxiliary Current				Type B or BF AP: 10 μA NC; 50 μA SFC (d.c. current);
1 ig. 19 – Falletit Auxiliary Current		_	_	100 μA NC; 500 μA SFC (a.c.) ;
				Type CF AP: 10 μA NC;50 μA SFC (d.c. or a.c. current)





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Clause	Requirement + Test	Result - Remark	Verdict

	Maximum allowed values:
	Type B or BF AP: 50 µA NC; 100µA SFC (d.c. current);
_	500 μA NC; 1000 μA SFC (a.c.);
	Type CF AP: 50 μA NC; 100 μA SFC (d.c. or a.c. current)
	Maximum allowed values:
	Type B or BF AP: 50 µA NC; 100µA SFC (d.c. current);
_	500 μA NC;1000 μA SFC (a.c.);
	Type CF AP: 50 μA NC; 100 μA SFC (d.c. or a.c. current)
	Maximum allowed values:
_	Type B: NA
	Type BF: 5000 μA
	Type CF: 100 μA
	Maximum allowed values:
_	Type B & BF: 1000 μA
	Type CF: N/A
	<u></u>
	Maximum allowed values:
	5 mA NC; 10 mA SFC

Supplementary information:

- Note 1: For EARTH LEAKAGE CURRENT see 8.7.3 d) and 8.7.4.5;
- Note 2: For TOUCH CURRENT see 8.7.3 c) and 8.7.4.6;
- Note 3: For PATIENT LEAKAGE CURRENT SEE 8.7.3.b) and 8.7.4.7
- Note 4: Total PATIENT LEAKAGE CURRENT values are only relative to equipment with multiple APPLIED PARTS of the same type. See 8.7.4.7 h). The individual APPLIED PARTS complied with the PATIENT LEAKAGE CURRENT values.

Note 5: In addition to conditions indicated in the Table, tests conducted at operating temperature and after humidity preconditioning of 5.7, EQUIPMENT energized in stand-by condition and fully operating, max rated supply frequency, at 110 % of the max rated mains voltage, and after relevant tests of Clause 11.6 (i.e., overflow, spillage, leakage, ingress of water and particulate matter, cleaning & disinfection, & sterilization).

Touch current measured after 2 days and 5 days on humidity treatment.





	IEC 60601-1				
Clause	Requirement + Test		Result - Remark	Verdict	
TC – Touch of P - Patient le PA – Patient TP – Total Pa	akage current auxiliary current atient current leakage current with mains on the applied parts	B 1 0 N	- After humidity conditioning - Before humidity conditioning - Switch closed or set to normal pola - Switch open or set to reversed pola C - Normal condition FC - Single fault condition	,	

8.8.3		ABLE: Dielectric strength test of solid insulating materials with safety inction – MEANS OF OPERATOR PROTECTION (MOOP) / MEANS OF PATIENT PROTECTION MOPP)					
Inquiation	under teet	Reference Voltage Die					
(area from	under test insulation ram)	Insulation Type (1 or 2 MOOP/MOPP)	PEAK WORKING VOLTAGE (U) V peak	PEAK WORKING VOLTAGE (U) V d.c.	A.C. test voltages in V r.m.s ¹⁾	breakdown after 1 minute Yes/No ²⁾	
Basic insu	lation						
Primary to	PE	1 x MOOP	574 V peak		1864 Vac*	No	
Reinforced	l insulation						
Primary to	secondary	2 x MOOP	560 V peak		3000 Vac	No	
Primary to plastic enclused)		2 x MOOP	354 V peak		3000 Vac	No	
Two layers insulation to (provided by primary and secondary the transformation)	ape etween d windings of	2 x MOOP	354 Vpeak		3000 Vac	No	

Supplementary information:

¹ Alternatively, per the Table (i.e., __dc), a d.c. test voltage equal to the peak value of the a.c. test voltage used.

² A) Immediately after humidity treatment of 5.7, ME EQUIPMENT de-energized, B) after required sterilization PROCEDURE, ME EQUIPMENT de-energized, C) after reaching steady state operating temperature as during heating test of 11.1.1, and D) after relevant tests of 11.6 (i.e., overflow, spillage, leakage, ingress of water, cleaning, disinfection, and sterilization).

EUT was evaluated for 2 Means Of Operator Protection (2 x MOOP) between primary and secondary and between primary and accessible (plastic) enclosure and for 1 x MOOP between primary and protective earthing.

*) Based on measured working voltage between Transformer T2 and PE.

8.8.4.1	TABLE: Resistance to heat - Ball pressure test of thermoplastic parts			Р	
	Allowed impression diameter (mm):	≤ 2	≤ 2 mm		_
	Force (N):	20		_	
Part/mate	rial		Test temperature (°C)		ression eter (mm)



· · · · · · · · · · · · · · · · · · ·				
		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

Enclosure/External insulating parts				
Insulating material supporting un-insulated Mains Parts				

Supplementary information:

Manufacturer is using approved materials with adequate temperature characteristics. See Table 8.10. No additional test was considered required.

8.9.2 TABLE: Short circuiting of each single one of the CREEPAGE DISTANCES and AIR CLEARANCES for insulation in the MAINS PART between parts of opposite polarity in lieu of complying with the required measurements in 8.9.4					Α
	areas of circuits short- ed and test conditions	Test in lieu of CREEPAGE DISTANCE OF AIR CLEARANCE ¹⁾	HAZARDOUS SITUATION observed (i.e., fire hazard, shock hazard, explosion, discharge of parts, etc.)? Yes/No	Remarks	
					•

Supplementary information:

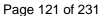
1) Note: AC - AIR CLEARANCE CD - CREEPAGE DISTANCE

Sufficient creepage and clearance distances between parts of opposite polarity provided before mains fuses. After mains fuses, distances verified by short circuit. See Table 13.2 for details.

8.9.3.2	Table: Thermal cycling tests on o solid insulation between conduct	ng N/A		
Part Test	8.9.3.4 - Test duration and temperature for 10 cycles after which the sample was subjected to Humidity Preconditioning per Cl. 5.7 Dielectric test voltage Dielectric strength test after humidity preconditioning per cl. 5.7 except for 48 h only, Breakdown: Yes/No			
	68 h at T1 ± 2 °C =°C 1)			
	1 h at 25 °C ± 2 °C			
	2 h at 0 °C ± 2 °C			
	1 or more h at 25 °C ± 2 °C			

Supplementary information:

¹⁾ T1 = 10 °C above the maximum temperature of relevant part determined per 11.1.1, or 85 °C, the higher of the two. 10 °C not added to T1 when temperature measured by an embedded thermocouple. Used gradual transition from one temperature to another.





IEC 60601-1				
Clause	Requirement + Test	Result - Remark	Verdict	

8.9.3.3		Table: Thermal cycling tests on one sample of cemented joint with other insulating parts (see 8.9.3.3)				
Part tested	Sample	pple Each test duration and temperature Dielectric test voltage Dielectric				
		10 Cycles conducted of the following:		-	_	
		1 - 68 h at T1 ± 2 °C =°C¹				
	1	2 - 1 h at 25 °C ± 2 °C				
		3 - 2 h at 0 °C ± 2 °C				
		4 - 1 or more h at 25 °C ± 2 °C				
	2	Humidity Conditioning per 5.7		-	-	
	3	Humidity Conditioning per 5.7		-	-	

Supplementary information:

¹⁾ T1 = 10 °C above the maximum temperature of relevant part determined per 11.1.1, or 85 °C, the higher of the two. 10 °C not added to T1 when temperature measured by an embedded thermocouple. Used gradual transition from one temperature to another.





IEC 60601-1				
Clause	Requirement + Test	Result - Remark	Verdict	

8.10 TA	BLE: List of critica	al components			Р
Component/ Part No.	Manufacturer/ Trademark	Type No./model No./	Technical data	Standard No./, Edition	Mark(s) & Certificates of conformity ¹⁾
		Unit from	n outside		
Enclosure (Electrical, Fire)	Min. thickness: 2,3	S x 62 mm x 35 mm s mm arts are fixed togethe	r with 2 screws.	IEC/EN 60601-1	Checked with appliance.
Enclosure Material	Teijin Chemicals Ltd.	LN-1250P LN-1250G	UL94-V0 at minimum thickness 1,5 mm 125°C	IEC/EN 60601-1 (QMFZ2)	Checked with appliance. UR E50075
Enclosure Material (alternative)	Sabic Innovative plastic	SE1X SE1 HF500R CX7211 EXCY0098	UL94-V1 at minimum thickness 1,5 mm 90°C	IEC/EN 60601-1 (QMFZ2)	Checked with appliance. UR E45329
Enclosure Material (alternative) Revision 1.0	Sabic Innovative plastic	945	UL94-V0 at minimum thickness 1,5 mm 90°C	IEC/EN 60601-1 (QMFZ2)	Accepted. UR E45329
Appliance inlet	Leci	DB-14	Min.10A, 250Vac, 70°C	IEC/EN 60320-1	VDE 40032137
Appliance inlet (alternative)	Rich Bay Co., Ltd.	R-301	Min.10A, 250Vac, 70°C	IEC/EN 60320-1	VDE 40029319
Appliance inlet (alternative)	Rong Feng Industrial	SS-120	Min.10A, 250Vac, 70°C	IEC/EN 60320-1	VDE 40029376
Appliance inlet (alternative)	Sun Fair	S-03	Min.10A, 250Vac, 70°C	IEC/EN 60320-1	VDE 40034447
Appliance inlet (alternative)	TECX-UNIONS Technology	TU-301-SP	Min.10A, 250Vac, 70°C	IEC/EN 60320-1	VDE 40025582
Appliance inlet (alternative)	Zhe Jiang Bei Er Jia	ST-A01 Series	Min.10A, 250Vac, 70°C	IEC/EN 60320-1 (AXUT2)	VDE 40013388 UR E225980



IEC 60601-1				
Clause	Requirement + Test	Result - Remark	Verdict	

Appliance	Steady	2107	Min.10A,	IEC/EN 60320-1	VDE 40011923
inlet			250Vac, 70°C	(AYVZ2)	UR E315002
(alternative)		 	m inside		
Fuse (F1, F2)	Bel Fuse Ltd.	MRT	T3.15A; 250V	IEC/EN 60127-3	VDE 139937
, ,		372	+		
Fuse (F1, F2) (alternative)	Littlefuse, Inc	372	T3.15A; 250V	IEC/EN 60127-3	VDE 097187
Fuse (F1, F2) (alternative)	Schurter	MST 250	T3.15A; 250V	IEC/EN 60127-3	VDE 40013529
Fuse (F1, F2) (alternative)	Cooper Bussmann	SR-5	T3.15A; 250V	IEC/EN 60127-3	VDE 122052
Fuse (F1, F2) (alternative)	Conquer Electronics Co., Ltd.	MET	T3.15A; 250V	IEC/EN 60127-3	VDE 40017157
Fuse (F1, F2)	Ever Island	2000	T3.15A,250V	IEC/EN 60127-3	VDE40018790
(alternative)	Electric Co., Ltd. & Walter Electric			(JDYX2)	UR E220181
Fuse (F1, F2) (alternative)	Enterprise Co.,	TSP, TMP	T3.15A, 250V	IEC/EN 60127-3	VDE 40027173 (TSP)
	Ltd.				VDE 40027218 (TMP)
				(JDYX2)	UR E133774
Capacitor CX1, CX2	+ Cheng Tung Industrial Co.,	СТХ	Min.250Vac, Max.0.33µF,	IEC/EN 60384- 14	VDE 40026382
	Ltd.		X1 (min. X2)	UL1414	
				(FOWX2)	UR E193049
Capacitor CX1, CX2	Dain Electronics Co., Ltd	MPX	Min.250Vac, Max.0.33µF,	IEC/EN 60384- 14	VDE 40018798
(alternative)			Min. X2	UL1414	
				(FOWX2)	UR E147776
Capacitor CX1, CX2	Ultra Tech Xiphi Enterprise Co.,	HQX	Min.250Vac, Max.0.33µF,	IEC/EN 60384- 14	VDE 40015608
(alternative)	Ltd.		Min. X2	UL1414	
				(FOWX2)	UR E183780
Y capacitor CY1	+ Success Electronics Co.,	SE, SB, SF	Max.1500pF, Min. 250Vac,	IEC/EN 60384- 14	VDE 40008996 (SE)
	Ltd.		Min.Y2		VDE 40016621 (SB)
					VDE 40016665 (SF)



		1 3.9 1 - 1 31 - 21		
		IEC 60601-1		
Clause	Requirement + Test	Result -	Remark Verdi	ct

Y capacitor CY1 (alternative)	TDK – EPC Corporation, Capacitors Group	CS, CD	Max.1500pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 122006 (CS) VDE 124321 (CD)
Y capacitor CY1 (alternative)	JYA-NAY Co., Ltd.	JY, JN	Max.1500pF, Min. 250VAC Min.Y2	IEC/EN 60384- 14	VDE 40001827 (JY) VDE 40001831 (JN)
Y capacitor CY2	+ Success Electronics Co., Ltd.	SE, SB, SF	Max.1000pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 40008996 (SE) VDE 40016621 (SB) VDE 40016665 (SF)
Y capacitor CY2 (alternative)	TDK – EPC Corporation, Capacitors Group	CS, CD	Max.1000pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 122006 (CS) VDE 124321 (CD)
Y capacitor CY2 (alternative)	JYA-NAY Co., Ltd.	JY, JN	Max.1000pF, Min. 250VAC Min.Y2	IEC/EN 60384- 14	VDE 40001827 (JY) VDE 40001831 (JN)
Y capacitor CY3	+ Success Electronics Co., Ltd.	SE, SB, SF	Max.470pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 40008996 (SE) VDE 40016621 (SB) VDE 40016665 (SF)
Y capacitor CY3 (alternative)	TDK – EPC Corporation, Capacitors Group	CS, CD	Max.470pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 122006 (CS) VDE 124321 (CD)
Y capacitor CY3 (alternative)	JYA-NAY Co., Ltd.	JY, JN	Max.470pF, Min. 250VAC Min.Y2	IEC/EN 60384- 14	VDE 40001827 (JY) VDE 40001831 (JN)
Y capacitor CY4	+ Success Electronics Co., Ltd.	SE, SB, SF	Max.220pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 40008996 (SE) VDE 40016621 (SB) VDE 40016665 (SF)



IEC 60601-1					
Clause	Requirement + Test	Result - Remark	Verdict		

Y capacitor CY4 (alternative)	TDK – EPC Corporation, Capacitors Group	CS, CD	Max.220pF, Min. 250Vac, Min.Y2	IEC/EN 60384- 14	VDE 122006 (CS) VDE 124321 (CD)
Y capacitor CY4 (alternative)	JYA-NAY Co., Ltd.	JY, JN	Max.220pF, Min. 250VAC Min.Y2	IEC/EN 60384- 14	VDE 40001827 (JY) VDE 40001831 (JN)
Optocoupler IC2, IC6	Lite-On	LTV-817	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40015248 UR E113898
Optocoupler IC2, IC6 (alternative)	Lite-On	LTV-357T	Isolation voltage: 3750 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 138213 UR E113898
Optocoupler IC2, IC6 (alternative)	Sharp	PC817	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40008087 UR E64380
Optocoupler IC2, IC6 (alternative)	Sharp	PC123	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40008087 UR E64380
Optocoupler IC2, IC6 (alternative)	Bright Led Electronics	BPC-817	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40007240 UR E236324
Optocoupler IC2, IC6 (alternative)	Bright Led Electronics	BPC-817M	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40007240 UR E236324
Optocoupler IC2, IC6 (alternative)	Bright Led Electronics	BPC-817S	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40007240 UR E236324
Optocoupler IC2, IC6 (alternative)	Cosmo	K1010	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 101347 UR E169586
Optocoupler IC2, IC6 (alternative)	Fairchild	H11A817B	Isolation voltage: 5000 Vac 100°C	VDE 0884 UL1577 (FPQU2)	VDE 40026857 UR E90700



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		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

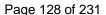
Optocoupler IC2, IC6 (alternative)	Fairchild	H11A817C	Isolation voltage: 5000 Vac	VDE 0884 UL1577	VDE 40026857
(diterriative)			100°C	(FPQU2)	UR E90700
Optocoupler	Fairchild	FOD817B	Isolation	VDE 0884	VDE 40026857
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577	
(======================================			100°C	(FPQU2)	UR E90700
Optocoupler	Everlight	EL817	Isolation	VDE 0884	VDE 132249
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577	
(anomany)			100°C	(FPQU2)	UR E214129
Bleeder Resistors	Various	SMD type	Max.330KΩ, 1/4W	IEC/EN 60601-1	Checked with appliance.
(R1, R2)					
Varistor MOV1	Panasonic	10K471U	300 Vrms; 385 Vdc;	IEC/EN 60601-1	VDE 40018677
(alternative)			diameter: 10	UL1449	
			mm	(VZCA2)	UR E321499
Varistor MOV1	Panasonic	14K471U	300 Vrms; 385 Vdc;	IEC/EN 60601-1	VDE 40018677
(alternative)			diameter: 14	UL1449	
			mm	(VZCA2)	UR E321499
Varistor	Thinking Electronic	TVR07471	300 Vrms;	IEC/EN 60601-1	VDE 005944
MOV1 (alternative)	Industrial		385 Vdc; diameter: 7 mm	UL1449	
, ,				(VZCA2)	UR E314979
Varistor	Thinking	TVR10471	300 Vrms;	IEC/EN 60601-1	VDE 005944
MOV1 (alternative)	Electronic Industrial		385 Vdc; diameter: 10	UL1449	
, ,			mm	(VZCA2)	UR E314979
Varistor	Thinking	TVR14471	300 Vrms;	IEC/EN 60601-1	VDE 005944
MOV1 (alternative)	Electronic Industrial		385 Vdc; diameter: 14	UL1449	
			mm	(VZCA2)	UR E314979
Varistor MOV1	Joyin	7N471K	300 Vrms;	IEC/EN 60601-1	VDE 005937
(alternative)			385 Vdc; diameter: 7 mm	UL1449	
				(VZCA2)	UR E325508
Varistor MOV1	Joyin	10N471K	300 Vrms; 385 Vdc;	IEC/EN 60601-1	VDE 005937
(alternative)			diameter: 10	UL1449	
,			mm	(VZCA2)	UR E325508
Varistor MOV1	Joyin	14N471K	300 Vrms;	IEC/EN 60601-1	VDE 005937
(alternative)			385 Vdc; diameter: 14	UL1449	
			mm	(VZCA2)	UR E325508





IEC 60601-1					
Clause	Requirement + Test	Result - Remark	Verdict		

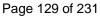
Varistor Centra CNR07D471K 300 Vrms; 385 Vdc;			IEC/EN 60601-1	VDE 40008220		
(alternative)			diameter: 7 mm	UL1449		
				(VZCA2)	UR E316325	
Varistor	Centra	CNR10D471K	300 Vrms;	IEC/EN 60601-1	VDE 40008220	
MOV1 (alternative)			385 Vdc; diameter: 10	UL1449		
			mm	(VZCA2)	UR E316325	
Varistor	Centra	CNR14D471K	300 Vrms;	IEC/EN 60601-1	VDE 40008220	
MOV1 (alternative)			385 Vdc; diameter: 14	UL1449		
(anomany)			mm	(VZCA2)	UR E316325	
Varistor	Success	SVR07D471K	300 Vrms;	IEC/EN 60601-1	VDE 123677	
MOV1 (alternative)	Electronics Co., Ltd.		385 Vdc; diameter: 7 mm	UL1449		
(anomative)	Ltd.			(VZCA2)	UR E330256	
Varistor	Success	SVR10D471K	300 Vrms;	IEC/EN 60601-1	VDE 123677	
MOV1 (alternative)	Electronics Co., Ltd.		385 Vdc; diameter: 10	UL1449		
(anomativo)			mm	(VZCA2)	UR E330256	
Varistor	Success	SVR14D471K	300 Vrms;	IEC/EN 60601-1	VDE 123677	
MOV1 (alternative)	Electronics Co., Ltd.		385 Vdc; diameter: 14	UL1449		
(anomany)			mm	(VZCA2)	UR E330256	
Thermistor	Various	Various	Min. 5A, 2,5Ω	IEC/EN 60601-1	Checked with	
TH1				UL 1434	appliance.	
				(XGPU2)	UR E138827	
Bridge Diode	+ Various	Various	Min. 4A;	IEC/EN 60601-1	Checked with	
(DB1)			Min. 800V		appliance.	
Electrolytic	+ Various	Various	Min.150µF	IEC/EN 60601-1	Checked with	
capacitor C9			Min. 400V		appliance.	
			105°C			
Choke L1	+ GlobTek	GS-150-L1	Min. 17mH	IEC/EN 60601-1	Checked with	
			125°C		appliance.	
Choke L2	+ GlobTek	GS-150-L2	145µH±10%	IEC/EN 60601-1	Checked with	
			130°C		appliance.	
Choke L5	+ GlobTek	GS-150-L5	Min. 380μH	IEC/EN 60601-1	Checked with	
			130°C		appliance.	





		IEC 60601-1		
Clause	Requirement + Test		Result - Remark	Verdict

Transformer	GlobTek or BOAM	1 or Haipuwei or HEJ	IIA	IEC/EN 60601-1	Checked with
T1	GS-150-T1 (HES4	19-xxxx)			appliance.
	Open type constru 33,5mm by 26mm	uction with overall dir by 16mm.			
	Primary and second bobbin.	ndary magnet wire w	ound around		
	3 layers of insulati and secondary wii	ion tape provided be ndings.	tween primary		
	Class B.				
	See enclosed tran Enclosure No. 3	sformer specification	ns for details in		
Bobbin	+ Chang Chun	T375J	UL94-V0	IEC/EN 60601-1	Checked with
			Min. 150°C	UL746	appliance. UR E59481
Bobbin	+ Sumitomo	PM9820	UL94-V0	IEC/EN 60601-1	Checked with
(alternative)		PM9630	Min. 150°C	UL746	appliance. UR E41429
Bobbin	Chang Chun	T375HF	Phenolic	IEC/EN 60601-1	Accepted.
(alternative) Revision 1.0	Plastics Co., Ltd.		V-0 at min. 0,43	UL746	UR E59481
			Min. 150°C		
Insulation tape	+ 3M	1350F-1 1350T-1	130°C	IEC/EN 60601-1	Checked with appliance.
		44		UL510	UR E17385
Internal PE	+ Various	Various	VW-1; 18AWG	IEC/EN 60601-1	Checked with
wire (green/yellow			105°C; 600V	UL758	appliance.
wire)				(AVLV2)	UR E108485
Output wire	+ Various	Various	VW-1; min.	IEC/EN 60601-1	Checked with
			22AWG; min. 80°C; min.	UL758	appliance.
			300V	(AVLV2)	UR E330069
PCB	Various	Various	Min. UL94V-0; 130°C	(ZPMV2)	UR
			Dimensions: approx. 115,5mm by 54,8mm		
			Min. thickness: 1,5mm		



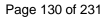


IEC 60601-1					
Clause	Requirement + Test	Result - Remark	Verdict		

Supplementary information:

- 1) Indicates a mark which assures the agreed level of surveillance. See Licenses and Certificates of Conformity for verification.
- 2) + means, that components from other vendor and other model number, but with the same characteristics and equivalent approvals are accepted.

8.10 b	TABLE: List of identified components with HIGH INTEGRITY CHARACTERISTICS P					Р
Componen Part No.	Manufacturer/ Trademark	Type No./model No./	Technical data	Standard No./, Edition	Се	Mark(s) & rtificates of onformity ¹⁾
Optocouple	Lite-On	LTV-817	Isolation	VDE 0884	VDE	40015248
IC2, IC6			voltage: 5000 Vac	UL1577		
			100°C	(FPQU2)	UR I	E113898
Optocouple	Lite-On	LTV-357T	Isolation	VDE 0884	VDE	138213
IC2, IC6 (alternative)			voltage: 3750 Vac	UL1577		
(4.1011141115)			100°C	(FPQU2)	UR I	E113898
Optocouple	Sharp	PC817	Isolation	VDE 0884	VDE	40008087
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577		
(anomalive)			100°C	(FPQU2)	UR I	E64380
Optocouple	Sharp	PC123	Isolation	VDE 0884	VDE	40008087
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577		
(ancimative)			100°C	(FPQU2)	UR I	E64380
Optocouple		BPC-817	Isolation	VDE 0884	VDE	40007240
IC2, IC6 (alternative)	Electronics		voltage: 5000 Vac	UL1577		
,			100°C	(FPQU2)	UR I	E236324
Optocouple		BPC-817M	Isolation	VDE 0884	VDE	40007240
IC2, IC6 (alternative)	Electronics		voltage: 5000 Vac	UL1577		
,			100°C	(FPQU2)	UR I	E236324
Optocouple		BPC-817S	Isolation	VDE 0884	VDE	40007240
IC2, IC6 (alternative)	Electronics		voltage: 5000 Vac	UL1577		
,			100°C	(FPQU2)	UR I	E236324
Optocouple	Cosmo	K1010	Isolation	VDE 0884	VDE	101347
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577		
,,			100°C	(FPQU2)	UR I	E169586





Report No. T223-0554/19 IEC 60601-1 Clause Requirement + Test Result - Remark Verdict

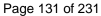
Optocoupler IC2, IC6 (alternative)	Fairchild	H11A817B	Isolation voltage: 5000 Vac	VDE 0884 UL1577	VDE 40026857
			100°C	(FPQU2)	UR E90700
Optocoupler	Fairchild	H11A817C	Isolation	VDE 0884	VDE 40026857
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577	
			100°C	(FPQU2)	UR E90700
Optocoupler	Fairchild	FOD817B	Isolation	VDE 0884	VDE 40026857
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577	
			100°C	(FPQU2)	UR E90700
Optocoupler	Everlight	EL817	Isolation	VDE 0884	VDE 132249
IC2, IC6 (alternative)			voltage: 5000 Vac	UL1577	
			100°C	(FPQU2)	UR E214129
Transformer	GlobTek or BOAM	1 or ZhongTong or H	EJIA	IEC/EN 60601-1	Checked with
T1	GS-150-T1 (HES4	19-xxxx)			appliance.
	Open type constru 33,5mm by 26mm	uction with overall dir by 16mm.	mension approx.		
	Primary and secondary magnet wire wound around bobbin.				
	3 layers of insulation tape provided between primary and secondary windings.				
	Class B.				
See enclosed transformer specifications for details Enclosure No. 3					

Supplementary information:

1) Indicates a mark which assures the agreed level of surveillance. See Licenses and Certificates of Conformity for verification.

8.11.3.5	TABLE: Cord anchorages					N/A
Cord under test		Mass of equipment (kg)	Pull (N)	Torque Nm)	Rem	narks
		-				-
Supplementary information: \						

8.11.3.6	TABLE: Cord guard						
Cord under test		Test mass	Measured curvature	Remarks			
Suppleme	Supplementary information: \						



Result - Remark

> 100

> 25 or < 4

IEC 60601-1

Requirement + Test

> 100

> 25 or < 8

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Verdict

9.2.2.2	TABLE:	Measurement of gap "a" according to Table 20 (ISO 13852: 1996)					
Part of body		Allowable adult gap ¹⁾ , mm	Measured adult gap, mm	Allowable children gap ¹⁾ , mm		ed children p, mm	
Body		> 500		> 500			
Head		> 300 or < 120		> 300 or < 60			
Leg		> 180		> 180			
Foot		> 120 or < 35		> 120 or < 25			
Toes		> 50		> 50			
Arm		> 120		> 120			

Supplementary information:

Hand, wrist, fist

Finger

Clause

¹⁾ In general, gaps for adults used, except when the device is specifically designed for use with children, values for children applied.

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9.2.3.2	TABLE: Over-trave	TABLE: Over-travel End Stop Test			
ME EQUIPMENT end stop Test Condition (Test Condition (cycles, load, speed)	Remarks		
		+			
Supplementary information: \					

9.4.2.1	9.4.2.1 TABLE: Instability—overbalance in transport position				
ME EQUIPMENT preparation		Test Condition (transport position)	Remarks		
Supplementary information: \					

9.4.2.2 TABLE: Instability—overbalance excluding transport position					
ME EQUIPMENT preparation		Test Condition (excluding transport position) Test either 5 ° incline and verify Warning marking or 10 ° incline)	Remarks		
Suppleme	Supplementary information: \				



IEC 60601-1				
Clause	Requirement + Test	Result - Remark	Verdict	

9.4.2.3 TABLE: Instability—overbalance from horizontal and vertical forces				
ME EQUIPMENT preparation		Test Condition (force used, direction of force, weight of equipment, location of force)		
Supplementary information: \				

9.4.2.4.2 TABLE: Castors and wheels – Force for propulsion					
ME EQUIPMENT preparation		Test Condition (force location and height)	t) Remarks		
		-			
Supplementary information: \					

9.4.2.4.3 TABLE: Castors and wheels – Movement over a threshold					
ME EQUIPMENT preparation		Test Condition (speed of movement)	Remarks		
		-			
Supplementary information: \					

9.4.3.1		TABLE: Instability from unwanted lateral movement (including sliding) in N/A ransport position				
ME EQUIPMENT Preparation		Test Condition (transport position, Remarks working load, locking device(s), caster position)		;		
Supplementary information: \						

9.4.3.2		TABLE: Instability from unwanted lateral movement (including sliding) N/A excluding transport position				
ME EQUIPMENT Preparation		Test Condition (working load, locking device(s), caster position, force, force location, force direction)				
Supplementary information: \						

9.4.4 TABLE: Grips and other handling devices					
Clause and Name of Test		Test Condition	Remarks		
Supplementary information: \					



IEC 60601-1				
	IEC 00001-1			
Clause	Requirement + Test	Result - Remark	Verdict	

9.7.5 TABLE: Pressure vessels					N/A	
Hydraulic, Pneumatic or Suitable Media and Test Pressure		Vessel Burst	Permanent Deformation	Leaks	Vessel fluid substance	Remarks
Supplemen	Supplementary Information: \					

9.8.3.2 TABLE: PATIENT support/suspension system - Static forces					N/A		
ME EQUIPMENT part or area		Position	Load	Area	Remar	·ks	
Suppleme	Supplementary Information: \						

9.8.3.3		TABLE: Support/Suspension System – Dynamic forces due to loading from PV/A persons						
ME EQUIPMENT part or area		Position	Safe Working Load	ng Area Remarks				
Supplementary Information: \								

10.1.1	N/A		
Maximum	allowable radiation pA/kg (μ Sv/h) (mR/h)	36 (5 μSv/h) (0.5 mR/h)	
	Surface area under test Surface no./ Description ¹⁾	Measured Radiation, pA/kg (μSv/h) (mR/h)	Remarks
1/ /			
2/ /			
3/ /			
4/ /			
5/ /			
6/ /			
7/ /			
8/ /			
9/ /			
10/ /			



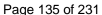
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Clause	Requirement + Test	Result - Remark	Verdict	

Supplementary information:

 $^{1)}$ Measurements made at a distance of 5 cm from any surface to which OPERATOR (other than SERVICE PERSONNEL) can gain access without a TOOL, is deliberately provided with means of access, or is instructed to enter regardless of whether or not a TOOL is needed to gain access



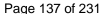


IEC 60601-1					
Clause	Requirement + Test	Result - Remark	Verdict		

11.1.1 TABLE: Excessive temperatures in ME EQUIPMENT						Р		
Model No			1)	2)	2) 3) 4)			
Test ambient (°C)		:	40	40	4	0	40	
Test supply	voltage/fre	quency (V/Hz)4:	90/60	100/60	120)/60	132/60	
Model No.	Thermo- couple No.	Thermocouple lo	cation ³	Max allowa temperature ¹ Table 22, 23 or RM file for AP	from r 24 or	temp	measured perature ² , (°C)	Remarks
1)	1.	Inductor L4		130			70,8	
1)	2.	Capacitor C9		105			95,7	
1)	3.	Transistor Q1		130		1	112,2	
1)	4.	Diode D1		130		1	106,3	
1)	5.	Bridge BR1		125		1	108,0	
1)	6.	Transformer T1 – wi	nding	120			91,7	
1)	7.	Transformer T1 – co	ore	120			87,0	
1)	8.	Capacitor C23		105			77,5	
1)	9.	Inductor L1		130		1	109,2	
1)	10.	Diode D15		130			80,2	
1)	11.	РСВ		130		88,8		
1)	12.	Optocoupler IC2		100		83,5		
1)	13.	Capacitor C9		105		76,1		
1)	14.	Inside enclosure		95		78,7		
1)	15.	Case Enclosure – T	OP	86	62,5		62,5	
1)	16.	Case Enclosure – B (test corner)	ОТТОМ	86			84,7	
0)		1. 1		400			70.5	
2)	1. 2.	Inductor L4 Capacitor C9		130 105			70,5 92,6	
2)								
2)	3.	Transistor Q1		130			108,3	
2)	4.	Diode D1		130			103,1	
2)	5.	Bridge BR1		125			103,4	
2)	6.	Transformer T1 – wi			120		90,8	
2)	7.	Transformer T1 – co	ore	120			86,2	
2)	8.	Capacitor C23		105			77,1	
2)	9.	Inductor L1		130			103,8	
2)	10.	Diode D15		130		79,6		



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Clause	Requirement + Test		Result - F	Remark	Verdict
2)	11.	PCB	130	87,8	
2)	12.	Optocoupler IC2	100	82,6	
2)	13.	Capacitor C9	105	75,2	
2)	14.	Inside enclosure	95	78,2	
2)	15.	Case Enclosure – TOP	86	62,4	
2)	16.	Case Enclosure – BOTTOM (test corner)	86	82,8	
3)	1.	Inductor L4	130	70,1	
3)	2.	Capacitor C9	105	88,3	
3)	3.	Transistor Q1	130	103,9	
3)	4.	Diode D1	130	99,4	
3)	5.	Bridge BR1	125	97,1	
3)	6.	Transformer T1 – winding	120	89,8	
3)	7.	Transformer T1 – core	120	85,4	
3)	8.	Capacitor C23	105	76,5	
3)	9.	Inductor L1	130	96,3	
3)	10.	Diode D15	130	79,0	
3)	11.	PCB	130	86,5	
3)	12.	Optocoupler IC2	100	81,5	
3)	13.	Capacitor C9	105	73,9	
3)	14.	Inside enclosure	95	77,6	
3)	15.	Case Enclosure – TOP	86	62,4	
3)	16.	Case Enclosure – BOTTOM (test corner)	86	79,6	
					,
4)	1.	Inductor L4	130	70,2	
4)	2.	Capacitor C9	105	86,9	
4)	3.	Transistor Q1	130	102,9	
4)	4.	Diode D1	130	98,5	
4)	5.	Bridge BR1	125	94,9	
4)	6.	Transformer T1 – winding	120	89,9	
4)	7.	Transformer T1 – core	120	85,5	
4)	8.	Capacitor C23	105	76,6	
4)	9.	Inductor L1	130	93,4	



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Clause	Requirer	ment + Test		Result - F	Verdict	
4)	10.	Diode D15	1	30	79,0	
4)	11.	PCB	1	30	86,5	
4)	12.	Optocoupler IC2	1	00	81,5	
4)	13.	Capacitor C9	1	05	73,7	
4)	14.	Inside enclosure	9	95	77,7	
4)	15.	Case Enclosure – TOP	8	36	62,4	
4)	16.	Case Enclosure – BOTTOM (test corner)	3	36	78,7	

Supplementary information:

- ¹ Maximum allowable temperature on surfaces of test corner is 90 °C
- ² Max temperature determined in accordance with 11.1.3e)
- ³ When thermocouples used to determine temperature of windings, limits of Table 22 reduced by 10 °C.
- ⁴ Supply voltage:
 - ME EQUIPMENT with heating elements 110 % of the maximum RATED voltage;
 - Motor operated ME EQUIPMENT least favourable voltage between 90 % of the minimum RATED and 110 % of the maximum RATED voltage. ME EQUIPMENT operated under normal load and normal DUTY CYCLE.
- Combined heating and motor operated and other ME EQUIPMENT tested both at 110 % of the maximum RATED voltage and at 90 % of the minimum RATED voltage.

Information from Risk Management, as applicable: Risk No.: H2

Model 1):

- Input current:1,30 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

Model 2):

- Input current: 1,19 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

Model 3):

- Input current: 1,04 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

Model 4):

- Input current: 0,97 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

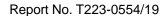
Temperature rises of the transformer was measured with thermocouples (Class B transformer provided inside the unit, 130°C - 10°C= 120°C).

The printed circuit board (PCB) is rated 130°C.

Desk-top power supply unit is not intended to be moved from one to another location during operation; therefore limit 86°C applied for external plastic surfaces.

⁵ APPLIED PARTS intended to supply heat to a PATIENT - See RISK MANAGEMENT FILE containing temperatures and clinical effects. Also, see instructions for use.





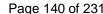


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Clause	Requirement + Test	Result - Remark	Verdict			

11.1.1 TABLE: Excessive temperatures in ME EQUIPMENT						Р		
Model No			1)	2)	2) 3) 4)			
Test ambie	nt (°C)	:	40	40	4	40		
Test supply	voltage/fre	quency (V/Hz)4:	200/50	230/50	240	/50	264/50	
Model No.	Thermo- couple No.	Thermocouple lo	cation ³	Max allowa temperature ¹ Table 22, 23 o RM file for AP	from r 24 or	temp	measured perature ² , (°C)	Remarks
1)	1.	Inductor L4		130		•	71,4	
1)	2.	Capacitor C9		105		;	83,7	
1)	3.	Transistor Q1		130		1	01,0	
1)	4.	Diode D1		130		!	97,6	
1)	5.	Bridge BR1		125		;	88,5	
1)	6.	Transformer T1 – wi	nding	120		,	92,6	
1)	7.	Transformer T1 – co	ore	120		;	88,0	
1)	8.	Capacitor C23		105		78,1		
1)	9.	Inductor L1		130		85,3		
1)	10.	Diode D15		130	130		80,7	
1)	11.	PCB		130		88,6		
1)	12.	Optocoupler IC2		100		83,0		
1)	13.	Capacitor C9		105		74,1		
1)	14.	Inside enclosure		95		79,4		
1)	15.	Case Enclosure – To		86			63,1	
1)	16.	Case Enclosure – B	ОТТОМ	86			76,1	
2)	1.	Inductor L4		130			71,9	
2)	2.	Capacitor C9		105			81,6	
2)	3.	Transistor Q1		130			96,2	
2)	4.	Diode D1					95,7	
-					130			
2)	5.	Bridge BR1		125			85,8	
2)	6.	Transformer T1 – wi		120			93,5	
2)	7.	Transformer T1 – co	ore		120 88,7			
2)	8.	Capacitor C23		105			78,6	
2)	9.	Inductor L1		130			82,5	
2)	10.	Diode D15		130		81,3		



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Clause	Requirer	ment + Test	Result - F	Verdict	
2)	11.	РСВ	130	89,4	
2)	12.	Optocoupler IC2	100	83,5	
2)	13.	Capacitor C9	105	74,0	
2)	14.	Inside enclosure	95	80,0	
2)	15.	Case Enclosure – TOP	86	63,5	
2)	16.	Case Enclosure – BOTTOM	86	74,6	
1					
3)	1.	Inductor L4	130	72,1	
3)	2.	Capacitor C9	105	81,2	
3)	3.	Transistor Q1	130	95,7	
3)	4.	Diode D1	130	95,5	
3)	5.	Bridge BR1	125	85,2	
3)	6.	Transformer T1 – winding	120	93,8	
3)	7.	Transformer T1 – core	120	89,0	
3)	8.	Capacitor C23	105	78,8	
3)	9.	Inductor L1	130	82,0	
3)	10.	Diode D15	130	81,4	
3)	11.	PCB	130	89,7	
3)	12.	Optocoupler IC2	100	83,7	
3)	13.	Capacitor C9	105	74,1	
3)	14.	Inside enclosure	95	80,2	
3)	15.	Case Enclosure – TOP	86	63,8	
3)	16.	Case Enclosure – BOTTOM	86	74,4	
4)	1	Inductor L4	420	72.0	
4)	1. 2.	Capacitor C9	130 105	73,0 83,2	
		Transistor Q1			
4)	3.		130	99,5	
4)	4.	Diode D1	130	98,1	
4)	5.	Bridge BR1	125	86,7	
4)	6.	Transformer T1 – winding	120	95,5	
4)	7.	Transformer T1 – core	120	90,6	
4)	8.	Capacitor C23	105	79,9	
4)	9.	Inductor L1	130	83,1	
4)	10.	Diode D15	130	82,7	
4)	11.	PCB	130	91,7	



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4)	12.	Optocoupler IC2	100	85,3	
4)	13.	Capacitor C9	105	75,4	
4)	14.	Inside enclosure	95	81,6	
4)	15.	Case Enclosure – TOP	86	64,4	
4)	16.	Case Enclosure – BOTTOM	86	76,1	

Supplementary information:

- ¹ Maximum allowable temperature on surfaces of test corner is 90 °C
- ² Max temperature determined in accordance with 11.1.3e)
- ³ When thermocouples used to determine temperature of windings, limits of Table 22 reduced by 10 °C.
- ⁴ Supply voltage:
 - ME EQUIPMENT with heating elements 110 % of the maximum RATED voltage;
 - Motor operated ME EQUIPMENT least favourable voltage between 90 % of the minimum RATED and 110 % of the maximum RATED voltage. ME EQUIPMENT operated under normal load and normal DUTY CYCLE.
- Combined heating and motor operated and other ME EQUIPMENT tested both at 110 % of the maximum RATED voltage and at 90 % of the minimum RATED voltage.

Information from Risk Management, as applicable: Risk No.: H2

Model 1):

- Input current:0,72 A - Output current: 2,0 A - Output voltage: 29,25 Vdc

Model 2):

- Input current: 0,66 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

Model 3):

- Input current: 0,63 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

Model 4):

- Input current: 0,60 A - Output current: 2,0 A - Output voltage: 29,17 Vdc

Temperature rises of the transformer was measured with thermocouples (Class B transformer provided inside the unit, 130°C - 10°C= 120°C).

The printed circuit board (PCB) is rated 130°C.

Desk-top power supply unit is not intended to be moved from one to another location during operation; therefore limit 86°C applied for external plastic surfaces.

⁵ APPLIED PARTS intended to supply heat to a PATIENT - See RISK MANAGEMENT FILE containing temperatures and clinical effects. Also, see instructions for use.





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Clause	Requirement + Test	Result - Remark	Verdict

11.1.3d	TABLE: Temperature of windings by change-of-resistance method							N/A
Temperature T of winding:		t₁ (°C)	R ₁ (Ω)	t ₂ (°C)	R ₂ (Ω)	T (°C)	Allowed T _{max} (°C)	Insulatio n class
Supplemen	Supplementary information: \							

11.2.2.1	TABLE: Alternative method tignition source	to 11.2.2.1 a) 5) to determine exis	tence of an	N/A
Areas whe	re sparking might cause igniti	on:	Remarks	
1.				
2.				
3.				
4.				
5.				
6.				
	f the parts between which spaignation, Manufacturer):	arks could occur (Composition,	Remark	S
1.				
2.				
3.				
4.				
5.				
6.				
Test param EQUIPMENT:	neters selected representing v	vorst case conditions for ME	Remark	S
Oxygen cor	ncentration (%):			
Fuel				
Current (A)	· · ·			
Voltage (V)	·			
Capacitanc	e (μF):			
Inductance	or resistance (h or Ω):			
No. of trials	(300 Min):			
Sparks resu	ulted in ignition (Yes/No):			



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Clause	Requirement + Test		Result - Remark	Verdict

Supplementary information:

Test procedure of 11.2.2.1 a) 5) & Figs 35-37 used for tests. For circuits not in Figs 35-37, test voltage or current set at 3 times the worst case values with other parameters set at worst case values to determine if ignition can occur.

Information from Risk Management, as applicable:

11.6.1	TABLE: overflow, spillage, leakage, ingress of water, cleaning, disinfection, sterilization, compatibility with substances						
Clause / Test Name		Test Condition	Part under test	Remarks			
Suppleme	Supplementary information: \						

13.1.2	TABLE: measurement of power or energy dissipation in parts & components to waive SINGLE FAULT CONDITIONS in 4.7, 8.1 b), 8.7.2, and 13.2.2 relative to emission of flames, molten metal, or ignitable substances					s to N/A
Power dissipated less than (W)						
Energy dissipated less than (J)						
Part or component dissipated (W) Calculated energy SINGLE FAULT CONDITIONS Restricted (W) Calculated energy waived (Yes/No)					Remarks	
Suppleme	ntary inform	ation: \	<u>'</u>		,	

13.2	TABLE: SINGLE FAULT CONDITIONS in accordance wi	th 13.2.2 to 13.2.13, inclusive	Р		
Clause No.	Description of SINGLE FAULT CONDITION	Results observed	HAZARDOUS SITUATION (Yes/No)		
13.2.2	Electrical SINGLE FAULT CONDITIONS per CI. 8.1:	_	_		
	SELV Reliability tests				
	D12 (Rated load)	Output immediately dropped	NO		
	Short	to 0V and 0A. Input current in hiccup mode (64-86mA). After			
	Input: 264Vac; 0,38A	removal of short output in			
		hiccup mode (0-8,4V; 0-0,23A). Input in hiccup mode			
		(57-69mA).			
		Unit damaged.			
		No hazard. No fire.			



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Clause	Requirement + Test	Result - Remark	Verdict	
	D15 (Rated load) Short Input: 264Vac; 0,38A	Output immediately dropped to 0V and 0A. Input current drops to 46mA. No peaks. After removal of short, no effect on function.	NO	
		No hazard. No fire.		
	IC2 (No load) Sec. pins short Input: 264Vac; 51mA	Output voltage goes up to 34,5 Vpk, then slowly drops to 0V. After removal of short no effect on function No damage. No hazard.	NO	
	IC2 (Pated load)		NO	
	IC2 (Rated load) Sec. pins short Input: 264Vac; 0,39A	Output voltage goes up from 28,5V to 29,7V. Output current goes up to 2,02A. input current rises to 0,4A. After removal of short output voltage first drops do 22.9Vpk, then unit operates normally.	NO	
		No damage. No hazard.		
	IC2 (No load) Prim. pins short Input: 264Vac; 51mA	Output voltage slowly dropped to 0V. After removal of short unit operates normally. No damage. No hazard.	NO	
	IC2 (Rated load) Prim. pins short Input: 264Vac; 0,39A	Output immediately dropped to 0V and 0A. Input current drops to 50mA. After removal of short unit operates normally.	NO	
		No damage. No hazard.		
	IC6 (No load) Sec. pins short Input: 264Vac; 50mA	No effect on function!	NO	
	IC6 (Rated load) Sec. pins short Input: 264Vac; 0,39A	No effect on function!	NO	
	IC6 (No load) Prim. pins short Input: 264Vac; 51mA	Output voltage slowly dropped to 0V. After removal of short no effect on function. No damage. No hazard.	NO	
	IC6 (Rated load) Prim. pins short Input: 264Vac; 0,39A	Output immediately dropped to 0V and 0A. Input current drops to 51mA. After removal of short no effect on function. No damage. No hazard.	NO	



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Clause	Requirement + Test	Result - Remark	Verdict
	C25 (Rated load) Short Input: 264Vac; 0,38A	Output voltage in hiccup mode (0,2-0,4V; 2-8mA). Input current in hiccup mode (0,064-0,088A). After remova of short unit operates normally.	
		No damage. No hazard.	
	C21 (Rated load) Short Input: 264Vac; 0,38A	Output voltage in hiccup mode (0-6,1V; 0-0,17A). Inpucurrent drops to 52mA. After removal of short unit operate normally.	
		No damage. No hazard.	
	C22 (Rated load) Short Input: 264Vac; 0,38A	Output voltage in hiccup mode (0-9V; 0-0,23A). Input current drops to 50mA. After removal of short unit operate normally.	
		No damage. No hazard.	
	C32 (Rated load) Short Input: 264Vac; 0,38A	Output voltage drops to 0V, 0A immediately. Input currendrops to 45mA. After removator of short unit operates normally.	
		No damage. No hazard.	
	C12 (Rated load) Short Input: 264Vac; 0,38A	Output voltage drops to 0V, 0A immediately. Input curren down to 45mA. After remova of short unit operates normally.	
		No damage. No hazard.	
	Q1 (Rated load) Short (D-S) Input: 264Vac; 0,38A	Output voltage drops to 0V, 0A immediately. Max input current is 0,41A. Unit damaged!	NO
		No fire. No hazard.	
	R12 (Rated load) Short Input: 264Vac; 0,39A	Input current first rises to 0,4A. Maximum input current is 1,26A, maximum output current is 5,96A, then unit goes in hiccup mode (output from 0 to 18,8V (0 – 0,37A); input from 66mA to 104mA). After removal of short unit stin hiccup mode. Unit damaged!	
		No fire. No hazard.	



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Clause	Requirement + Test	Result - Remark	Verdict
	C9 (Rated load) Short Input: 264Vac; 0,39A	Output voltage drops to 0A immediately. Primar blown! ** No fire. No hazard.	
	DB1 (Rated load) Short (~ o +) Input: 264Vac; 0,39A	Output voltage drops to 0A immediately. Primar blown! ** No fire. No hazard.	
	T1 (Rated load) Sec. pins short (pin 4S to pin 3F) Input: 264Vac; 0,38A	Output voltage first falls 30V to 8,1V for 0,5s the to 0V. Output current fall 0A. input current falls to 52mA. After removal of no effect on function.	en falls ills to o short
	T1 (Rated load) Sec. pins short (pin 4S to pin 4F) Input: 264Vac; 0,566A	Output voltage first falls 30V to 8,1V for 0,5s the to 0V. Output current falls to 52mA. After removal of no effect on function.	s from NO en falls ills to
	T1 (Rated load) Prim. pins short (pin 1F to pin 1S) Input: 264Vac; 0,39A	No damage. No hazard Output voltage immedia down to 0V. Input curre hiccup mode (65mA-10 After removal of short u operates normally. No damage. No hazard	ntely NO ent in (3mA). enit
	T1 (Rated load) Prim. pins short (pin 2F to pin 2S) Input: 264Vac; 0,39A	Output voltage in hiccup mode (0-14V). Output cin hiccup mode (0-0,39). Input current in hiccup (0,071-0,116A). After reof short unit operates normally.	p NO current A). mode emoval
	Output Short (+ to -) Input: 264Vac; 50mA	No damage. No hazard Output in hiccup mode max). Input current in h mode (0,054-0,074A). A removal of short unit op normally. No damage. No hazard	(2,38A NO iccup After perates



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Clause	Requirement + Test	Result - Remark	Verdict
	Output	Output voltage: 29,04Vdc	NO
	Overload	Output current: 2,24A	
	Input: 90Vac; 1,43A	Input current: 1,43A	
		Transformer temperatures:	
		T _{WINDING} = 96,8°C	
		T _{CORE} = 91,6°C	
		No excessive temperatures. No damage. No hazard.	
	Output	Output voltage: 29,11Vdc	NO
	Overload	Output current: 2,23A	
	Input: 264Vac; 0,66A	Input current: 0,66A	
		Transformer temperatures:	
		T _{WINDING} = 102,3°C	
		T _{CORE} = 96,9°C	
		No excessive temperatures. No damage. No hazard.	
13.2.3	Overheating of transformers per Clause 15.5:	-	_
	See table 15.5.1.2 and 15.5.1.3 for details.		No
13.2.4	Failure of THERMOSTATS according to 13.2.13 & 15.4.2, overloading - THERMOSTATS short circuited or interrupted, the less favourable of the two:	_	_
	No thermostats incorporated.		
13.2.5	Failure of temperature limiting devices according to 13.2.13 & 15.4.2, overloading, THERMOSTATS short circuited or interrupted, the less favourable of the two:	_	-
	No temperature limiting devices incorporated.		
13.2.6	Leakage of liquid - RISK MANAGEMENT FILE examined to determine the appropriate test conditions (sealed rechargeable batteries exempted)	-	_
	No liquid used.		
13.2.7	Impairment of cooling that could result in a HAZARD using test method of 11.1:	_	_
	Single ventilation fans locked consecutively	No fan provided.	
	Ventilation openings on top and sides impaired by covering openings on top of ENCLOSURE or positioning of ME EQUIPMENT against walls	No opening provided	
	Simulated blocking of filters	No such filter used.	
	Flow of a cooling agent interrupted	No cooling agent provided.	



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Clause	Requirement + Test	Result - Remark	Verdict

13.2.8	Locking of moving parts – Only one part locked at a time – Also see 13.2.10 below:	_	_
		-	
13.2.9	Interruption and short circuiting of motor capacitors – Motor capacitors short & open circuited ¹⁾ – Also see 13.10	_	_
		V measured =	
13.2.10	Additional test criteria for motor operated ME EQUIPMENT in 13.2.8 &13.2.9:	_	_
	For every test in SINGLE FAULT CONDITION of 13.2.8 and 13.2.9, motor-operated EQUIPMENT stared from COLD CONDITION at RATED voltage or upper limit of RATED voltage range for specified time:	No motors provided.	
	Temperatures of windings determined at the end of specified test periods or at the instant of operation of fuses, THERMAL CUT-OUTS, motor protective devices	No motors provided.	
	Temperatures measured as specified in 11.1.3 d)	No motors provided.	
	Temperatures did not exceed limits of Table 26	No motors provided.	
13.2.11	Failures of components in ME EQUIPMENT used in conjunction with OXYGEN RICH ENVIRONMENTS:	_	_
13.2.12	Failure of parts that might result in a MECHANICAL HAZARD (See 9 & 15.3):	_	_

Supplementary information:

There was no flame, extensive smoke or melted metal.

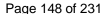
When components were failing, the test was repeated two times.

In case of primary fuse opened or fusible resistor, test repeated two times (three times in total) with all alternative fuses as specified within the Table 8.10 (List of critical components). There was no permanent arcing, no ignition and no bursting of the fuse. None of fuses exploded.

Information from Risk Management, as applicable:

15.3	TABLE: Mechanical Strength tests 1)			Р
Clause	Name of Test	Test conditions	Observed result	s/Remarks
15.3.2	Push Test	Force = 250 N ± 10 N for 5 s	No damage on the	enclosure
15.3.3	Impact Test	Steel ball (50 mm in dia., 500 g ± 25 g) falling from a 1.3 m	No damage on the enclosure	
15.3.4.2	Drop Test (hand-held and portable)	Drop height (cm) = 1 m	No damage on the	enclosure

¹⁾ Test with short-circuited capacitor not performed when motor provided with a capacitor complying with IEC 60252-1 and the ME EQUIPMENT not intended for unattended use including automatic or remote control. See Attachment # and appended Table 8.10.





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	IEC 60601-1		
Clause	Requirement + Test	Result - Remark	Verdict

15.3.6	Mould Stress Relief	7 h in oven at temperature (°C) = 95	No damage on the enclosure
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Supplementary information:

¹⁾ As applicable, Push, Impact, Drop, Mould Stress Relief and Rough Handling Tests (delete not applicable rows or state N/A in Remarks field).

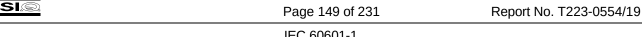
15.4.6	TABLE: ac	ABLE: actuating parts of controls of ME EQUIPMENT – torque & axial pull tests					
Rotating unde	£	Gripping diameter "d" of control knob (mm) ¹⁾	Torque from Table 30 (Nm)		Unacceptable RISK occurred Yes/No	Remarks	
-	-						

Supplementary information:

¹⁾ Gripping diameter (d) is the maximum width of a control knob regardless of its shape (e.g. control knob with pointer)

15.5.1.2	TABLE: transformer short circuit test short-circuit applied at end or at the first point that could be short circuited under SINGLE FAU		Р
Primary vol	Primary voltage (most adverse value from 90 % to 110 % of RATED voltage)(V) ¹⁾		_
RATED inpu	t frequency (Hz)	50 Hz	_

Winding tested	Class of insulation (A, B, E, F, or H)	Type of protective device (fuse, circuit breaker) /Ratings	Protective device operated Yes/No	Time to THERMAL STABILITY (when protective device did not operate)(Min)	Maximum allowed temp from Table 31 (°C)	Maximum winding temp measured (°C)	Ambient (°C)
Transf. T1 - Sec. pins short (pin 4S to pin 3F)	Class B	Fuse	No	10 min	165	*	25
Transf. T1 - Sec. pins short (pin 4S to pin 4F)	Class B	Fuse	No	10 min	165	*	25
Transf. T1 - Prim. pins short (pin 1F to pin 1S)	Class B	Fuse	No	10 min	165	*	25
Transf. T1 - Prim. pins short (pin 2F to pin 2S)	Class B	Fuse	No	10 min	165	*	25



	IEC 00001-1		
Clause	Requirement + Test	Result - Remark	Verdict

Supplementary information:

¹⁾Loads on other windings between no load and their NORMAL USE load. Short-circuit applied at end of windings or at the first point that could be short circuited under SINGLE FAULT CONDITION.

In both short tests the power supply reverts to hiccup mode. After removal of the short the power supply operated normally.

* See Table 13.2 – Fault condition tests. No excessive temperatures.

15.5.1.3 TABLE: transformer overload test – conducted only when protective device under short-circuit test operated		Р	
Primary vol	tage, most adverse value between 90 % to 110 % of RATED voltage (V)1):	90	Vac/264Vac
RATED input frequency (Hz):		50Hz	
Test current just below minimum current that would activate protective device and achieve THERMAL STABILITY under method a) (A)		Considered	
	Test current based on Table 32 when protective device that operated under method a) is external to transformer, and it was shunted (A)		N/A

Winding tested	Class of insulation (A, B, E, F, H)	Type of protective device used (fuse, circuit breaker)/Ratings	Maximum allowed temp from Table 31 (°C)	Maximum winding temp measured (°C)	Ambient (°C)
Output overload test also represent for T1 (secondary) overload test – Input voltage: 90Vac	Class B	Fuse	165	96,8	40
Output overload test also represent for T1 (secondary) overload test – Input voltage: 264Vac	Class B	Fuse	165	102,3	40

Supplementary information:

1) Loads on other windings between no load and their NORMAL USE load.

Time durations: - IEC 60127-1 fuse: 30 min at current from Table 32.

Non IEC 60127-1 fuse: 30 min at the current based on characteristics supplied by fuse manufacturer, specifically, 30 min clearing-time current. When no 30 min clearing-time current data available, test current from Table 32 used until THERMAL STABILITY achieved.

- Other types of protective devices: until THERMAL STABILITY achieved at a current just below minimum current operating the protective device in a). This portion concluded at specified time or when a second protective device opened.

15.5.2	TABLE	ABLE: Transformer dielectric strength after humidity preconditioning of 5.7			N/A	
Transfor Model/Type No		Test voltage applied between	Test voltage, (V)	Test frequency (Hz)	Breakdown Yes/No	Deterioration Yes/No
		Primary & secondary windings				

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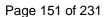
		i ago	100 01 20 1		Ropoli No.	1220 000 1/10
		IEC 6	0601-1			
Clause	Requir	ement + Test		Result - Remark	k	Verdict
	Primary winding & frame					
		Secondary winding & frame				

Supplementary information:

Tests conducted under the conditions of 11.1, in ME EQUIPMENT or under simulated conditions on the bench. See Clause 15.5.2 for test parameters & other details

16.6.1	TABLE: LEAKAGE CURRENTS IN ME SYSTEM _ TOUCH CURRENT MEASUREMENTS					N/A
Specific area where TOUCH CURRENT measured (i.e., from or between parts of ME SYSTEM within PATIENT ENVIRONMENT)		Allowable TOUCH CURRENT IN NORMAL CONDITION (μA) Measured TOUCH CURRENT IN Event of Interruption of PROTECTIVE EARTH CONDUCTOR, (μA)		Measured TOUCH CURRENT in event of interruption of PROTECTIVE EARTH CONDUCTOR, (μΑ)		
		100		500		
Supplemen	Supplementary information: \					

SP TABLE: Additional or special tests conducted				Р	
Clause and Name of Test		Test type and condition	Observed results		
Evaluation of voltage limiting components in SELV circuits		See table below	See table below		
Suppleme	Supplementary information: See Table below.				



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	9			, , , ,
	IEC	60601-1		
Clause	Requirement + Test	Resu	ult - Remark	Verdict

TABLE: evaluation of voltage li	imiting c	ompoi	nents in SELV	circuits	Р
Component (measured between)			oltage (V) operation)	Voltage Limiting Components	
	V p	eak	V d.c.		
T1: pin 3S to pin 3F	114	V _{pk}		D12	
	54,8	V_{rms}			
T1: pin 3S to after D12	_	-	30 V _{DC}	SELV	,
T1: pin 4S to pin 4F	114	V_{pk}		D15	
	54,8	V_{rms}			
T1: pin 4S to after D15	-	-	30 V _{DC}	SELV	,
Fault test performed on voltage limiting components			ge measured (ak or V d.c.)	V) in SELV circuits	3
D12 (Short)		curren short c	t in hiccup mod	ropped to 0V and 0A le (64-86mA). After i mode (0-8,4V; 0-0,2 nA).	emoval of
		Unit da	amaged.		
		No haz	zard. No fire.		
D15 (Short)		curren		ropped to 0V and 0A A. No peaks. After reaction.	
		No haz	zard. No fire.		
Supplementary information:	1				
Rated output load (30 Vdc / 2 A).					



Enclosure No. 1

National Differences to IEC 60601-1: 2005 + AM1: 2012

(25 pages including this cover page)



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National Differences to IEC 60601-1:2005 + A1:2012			
Clause	Requirement + Test	Result - Remark	Verdict

National	standard ANSI/AAMI ES60601-1: 2005	,	
4.8	Components of ME EQUIPMENT		Р
	When no relevant US ANSI standard existed, the requirements of this standard applied	Considered.	Р
4.10.2.	SUPPLY MAINS for ME EQUIPMENT and ME SYSTEMS		Р
	Replacement: Reference to "500 V" replaced with "600 V" in the second and third dashes to agree with the National Electrical Code (NEC) "and the NEC" added after the reference to "IEC 60364-4-41" in the text of the second-to-last dash of this sub-clause to agree with NEC	Considered.	P
8.2	Requirements related to power sources		Р
	Addition to agree with NEC: The requirement, "ALL FIXED ME EQUIPMENT and PERMANENTLY INSTALLED ME EQUIPMENT are CLASS I ME EQUIPMENT"	SMPS is Class I equipment.	Р
8.7.3	Allowable values:		Р
	Deleted the second sentence and note to sub- clause 8.7.3 d) to read as follows to agree with NFPA 99 which does not permit for allowances larger than the stated values:		Р
	d) The allowable values of the EARTH LEAKAGE CURRENT are 5 mA in NORMAL CONDITION and 10 mA in SINGLE FAULT CONDITION		Р
8.11.	MAINS PARTS, components and layout		N/A
	Addition to agree with NEC:		N/A
	The requirement, "Permanently connected ME EQUIPMENT shall have provision for the connection of one of the wiring systems that is in accordance with the NEC"	Not permanently connected equipment.	N/A
	Fixed and stationary X-ray ME EQUIPMENT supplied from a branch circuit rated at 30 A or less, and ME EQUIPMENT not strictly portable but obviously intended to be stationary, considered acceptable when supply connection provided with a length of attached Type S hard service flexible cord, or equivalent:		
	Installation of connecting cords between EQUIPMENT parts comply with NEC, as applicable	Connecting cord between equipment parts within end medical product not part of the investigation.	N/A



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	National Differences to IEC 60601-1	:2005 + A1:2012	
Clause	Requirement + Test	Result - Remark	Verdict
	Cable used as external interconnection between units was:		N/A
	1) Type SJT, SJTO, SJO, ST, SO, STO, or equivalent flexible cord when exposed to abuse, or similar multiple-conductor appliance-wiring material such as computer cable:		
	2) The cable was as in item 1) above when not exposed to abuse, or it was		
	i) Type SPT-2, SP-2, or SPE-2, or equivalent		
	ii) Type SVr, SVRO, SVE, or equivalent flexible cord or similar multiple-conductor appliance wiring material,		
	Or, ii) An assembly of insulated wires each with a nominal insulation thickness of 0.8 mm (1/32 inch) or more, enclosed in acceptable insulating tubing having a nominal wall thickness of 0.8 mm (1/32 inch) or more.		
	Receptacles provided as part of ME EQUIPMENT or ME SYSTEMS for use in the patient care areas of pediatric wards, rooms, or areas are listed (e.g., UL Certified) tamper resistant or employ a listed (e.g., UL Certified) tamper resistant cover in accordance with NEC		N/A
8.11.3.2.	Addition to agree with NEC:	Power supply cord not part of the investigation.	N/A
	The flexible cord is a type acceptable for the particular application, and it is acceptable for use at a voltage not less than the rated voltage of the appliance and has an ampacity as in NEC, not less than the current rating of the appliance.		N/A
	NATIONAL DIFFERENCES to IEC 60601-1 Third edit tandard CAN/CSA-C22.2 No. 60601-1:14	tion	
1.1	Scope		Р
	This Standard applies to the BASIC SAFETY and ESSENTIAL PERFORMANCE of MEDICAL ELECTRICAL	Considered.	Р
	EQUIPMENT and MEDICAL ELECTRICAL SYSTEMS designed to be used in accordance with CSA C22.1 (Canadian Electrical Code, Part I) and CSA Z32.		
	Note 1A: In the IEC 60601 Standards series adopted for use in Canada, the Canadian standards may modify, replace, or		
	delete requirements contained in the IEC		

and ME S

standard as appropriate to the ME EQUIPMENT



National Differences to IEC 60601-1:2005 + A1:2012				
Clause	Requirement + Test	Result - Remark	Verdict	

1.3	Collateral standards		Р
	Applicable Canadian 60601 collateral standards become normative at the date of their publication and apply together with this Standard.	Considered.	Р
1.4	Particular standards		Р
	Applicable Canadian 60601/80601 particular standards may modify, replace, or delete requirements contained in this Standard. The requirement of a Canadian 60601/80601 particular safety standard takes priority over this Standard	Considered.	Р
2	Normative references Where reference is made to CSA Group	Considered.	Р
	Standards, such reference shall be considered to refer to the latest edition and all amendments published to that edition. This Standard refers to the following Standards, and the years shown indicate the latest editions available at the time of printing:		
	CSA Group	Considered.	Р
	B51-09		
	Boiler, pressure vessel, and pressure piping code		
	C22.1-12		
	Canadian Electrical Code, Part I		
	CAN/CSA-C22.2 No. 0-10		
	General requirements — Canadian Electrical Code, Part II		
	C22.2 No. 21-95 (R2009)		
	Cord sets and power supply cords		
	C22.2 No. 42-10		
	General use receptacles, attachment plugs, and similar wiring devices		
	C22.2 No. 49-10		
	Flexible cords and cables		
	CAN/CSA-E61558-2-1-03 (R2012)		
	Safety of power transformers, power supply units and similar — Part 2: Particular requirements for separatingtransformers for general use		
	Z32-09		
	Electrical safety and essential electrical systems in health care facilities		



National Differences to IEC 60601-1:2005 + A1:2012			
Clause	Requirement + Test	Result - Remark	Verdict

CAN/CSA-Z305.8-03 (R2013)	Considered.	Р
Medical supply units		
Z305.12-06 (R2012)		
Safe storage, handling, and use of port oxygen systems in residential buildings health care facilities		
Z305.13-09		
Plume scavenging in surgical, diagnost therapeutic, and aesthetic settings	tic,	
CAN/CSA-Z5359-10		
Low-pressure hose assemblies for use medical gases	with	
CAN/CSA-Z9170-1-11		
Terminal units for medical gas pipeline — Part 1: Terminal units for use with compressed medical gases, vacuum, a anaesthetic gas scavenging systems		
CAN/CSA-Z10524-1:12		
Pressure regulators for use with medic — Part 1: Pressure regulators and pres regulators with flow-metering devices		
CAN/CSA-Z15002:12		
Flow-metering devices for connection terminal units of medical gas pipeline s		
ASME (American Society of Mechanica Engineers)	I	
PTC 25-2008		
Pressure Relief Devices		
CGA (Compressed Gas Association)		
V-1-2013		
Standard for Compressed Gas Cylinder Outlet and Inlet Connections	r Valve	
V-5-2008 (reaffirmed 2013)		
Diameter Index Safety System (Noninterchangeable Low Pressure Connections for Medical Gas Application	ons)	





National Differences to IEC 60601-1:2005 + A1:2012			
Clause	Requirement + Test	Result - Remark	Verdict

	ISO (International Organization for Standardization)		N/A
	32:1977		
	Gas cylinders for medical use — Marking for identification of content		
	407:2004		
	Small medical gas cylinders — Pin-index yoke- type valve connections		
	9170-2:2008		
	Terminal units for medical gas pipeline systems — Part 2: Terminal units for anaesthetic gas scavenging systems		
3.41	HIGH VOLTAGE		N/A
	voltage above 750 V, as defined in the Canadian Electrical Code, Part I	No such high voltage within the equipment.	N/A
4	GENERAL REQUIREMENTS		Р
	4.1A General	Considered.	Р
	General requirements applicable to ME EQUIPMENT and ME SYSTEMS are provided in CAN/CSA C22.2 No. 0.		
4.8	Components of ME EQUIPMENT		Р
	a) the applicable safety requirements of a relevant CSA Group, IEC, or ISO Standard; or	Considered.	Р
	b) where there is no relevant CSA Group, IEC, or ISO Standard, the requirements of this Standard shall be applied.	Considered.	Р
	Note 2: If there are neither requirements in this Standard nor in a CSA Group, IEC, or ISO Standard, any other applicable source (e.g., standards for other types of devices, national standards) could be used to demonstrate compliance with the RISK MANAGEMENT PROCESS.		
4.10.2	SUPPLY MAINS for ME EQUIPMENT and ME SYSTEMS		Р
	ME EQUIPMENT intended to be connected to SUPPLY MAINS shall be in accordance with the Canadian Electrical Code, Part I, and the following RATED voltages shall not be exceeded:		P
7.7	Colours of the insulation of conductors	Internal protective earth conductor.	N/A



National Differences to IEC 60601-1:2005 + A1:2012			
Clause	Requirement + Test	Result - Remark	Verdict

	Colours of the insulation of conductors shall be in accordance with the Canadian Electrical Code, Part I.		N/A
	A PROTECTIVE EARTH CONDUCTOR or a PROTECTIVE EARTH CONNECTION of any insulation on conductors shall be identified by either green or green and yellow colours at least at the termination of the conductors.		
	Identification by green or green and yellow insulation shall only be used for		
	- PROTECTIVE EARTH CONDUCTORS (see 8.6.2);		
	- conductors as specified in 7.7.2;		
	Note: In other safety standards such as IEC 60950-1, internal connections between conductive parts and the main protective earth are called "protective bonding conductors".		
	- POTENTIAL EQUALIZATION CONDUCTORS (see 8.6.7);		
	- FUNCTIONAL EARTH CONDUCTORS (see 8.6.9).		
	Colours of neutral conductors and POWER SUPPLY CORD conductors shall be in accordance with the Canadian Electrical Code, Part I, CSA C22.2 No. 21, and CSA C22.2 No. 49.		
	Compliance with the requirements of 7.7 is checked by inspection.		
8.7.3	Allowable values	Considered.	Р
	Allowable values shall also be in accordance with the Canadian Electrical Code, Part I.	Touch current did not exceed 100 µA in NC and 300 µA in SFC. Touch current measured on the output plus, output minus and plastic enclosure.	
8.11.3.2	Types		N/A



National Differences to IEC 60601-1:2005 + A1:2012				
Clause	Requirement + Test	Result - Remark	Verdict	

a) The MAINS PLUG of non-PERMANENTLY INSTALLED EQUIPMENT shall be	Power supply cord not part of the investigation.	N/A
i) if moulded-on type, a hospital-grade mains plug complying with CSA C22.2 No. 21;		
ii) a hospital-grade disassembly attachment plug type complying with CSA C22.2 No. 42; or		
ii) Class II equipment having fuses on the line side(s), and the neutral may use a non-polarized attachment plug or a polarized attachment plug. CSA configuration type 1-15P shall be required and shall meet all applicable requirements in CSA C22.2 No. 21 and CSA C22.2 No. 42. Where a polarized attachment plug is used, the POWER SUPPLY CORD shall be connected to the wiring of the EQUIPMENT on the ungrounded side of the line when any of the following devices are used in the primary circuit:		
the centre contact of an Edison base lampholder;		
2) a single pole switch;		
an automatic control with a marked off position;		
4) a solitary fuse/fuse holder; or		
any other single pole overcurrent protective device.		
b) A detachable POWER SUPPLY CORD for non- PERMANENTLY INSTALLED EQUIPMENT (cord- connected equipment) shall be of a type	Power supply cord not part of the investigation.	N/A
i) that can be shown to be unlikely to become detached accidentally, unless it can be shown that detachment will not constitute a safety HAZARD to a PATIENT or OPERATOR;		
ii) for which it can be shown that the impedance of the earth (ground) circuit contacts will not constitute a safety HAZARD to a PATIENT or OPERATOR; and		
iii) that has a terminal configuration or other constructional feature that will minimize the possibilityof its replacement by a detachable POWER SUPPLY CORD which could create a HAZARDOUS SITUATION.		



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	National Differer	nces to IEC 60601-1	:2005 + A1:2012		
Clause	Requirement + Test		Result - Remark	Verd	dict

	c) A detachable POWER SUPPLY CORD shall	Power supply cord not part of	N/A
	i) comply with the applicable requirements of CSA C22.2 No. 21; and	the investigation.	
	ii) not be smaller than No. 18 AWG, and the mechanical serviceability shall be not less than		
	Type SJ or equivalent for ME EQUIPMENT that is mobile or exposed to abuse; and		
	2) Type SV or equivalent for ME EQUIPMENT that is not exposed to abuse (or Type HPN if required because of temperature).		
	Note: See CSA C22.2 No. 49 for requirements for the cord types mentioned in Sub-item 2).		
	d) Installation of POWER SUPPLY CORDS shall meet the requirements of the Canadian Electrical Code, Part I, as applicable.	Power supply cord not part of the investigation.	N/A
8.11.5	Mains fuses and OVER-CURRENT RELEASES	See list of critical components.	Р
	Installation of overcurrent protective devices shall be in accordance with the Canadian Electrical Code, Part I.		
9.7.5	Pressure vessels	No pressure vessels provided.	N/A
	Pressure vessels shall comply with the requirements of CSA B51, as applicable		
9.7.7	Pressure-relief device	No pressure-relief devices	N/A
	A pressure-relief device shall comply, as applicable, with the requirements of ASME PTC 25 or equivalent Canadian requirements.	provided.	





National Differences to IEC 60601-1:2005 + A1:2012			
Clause	Requirement + Test	Result - Remark	Verdict

15.4.1	Construction of connectors	No gas cylinders provided.	N/A
	bA) The point of connection of gas cylinders to ME EQUIPMENT shall be gas-specific and clearly identified so that errors are avoided when a replacement is made. Medical gas inlet connectors on ME EQUIPMENT shall be		
	i) gas-specific, yoke type, or nut and nipple type valve connections complying with CGA V-1 for pressures over 1380 kPa (200 psi); or		
	ii) DISS type complying with CGA V-5 for pressures 1380 kPa (200 psi) or less and configured to permit the supply of medical gases from low-pressure connecting assemblies complying with CAN/CSA-Z5359.		
	Note: Users of this Standard should consult the CSA Z305 series of Standards, CAN/CSA-Z9170-1, ISO 9170-2, CAN/CSA-Z10524, and CAN/CSA-Z15002 for further information regarding inlet connectors; ISO 407 for requirements addressing yoke type valve connections; and ISO 32 for colour coding.		
15.4.8	Internal wiring of ME EQUIPMENT Flexible cords and equipment wire of ME EQUIPMENT shall be in accordance with the Canadian Electrical Code, Part I.		Р
16.1	General requirements for the ME SYSTEMS		N/A
	An ME SYSTEM shall provide		
	 within the PATIENT ENVIRONMENT, the level of safety equivalent to ME EQUIPMENT complying with this CSA Group Standard; and 		
	 outside the PATIENT ENVIRONMENT, the level of safety equivalent to equipment complying with their respective CSA Group, IEC, or ISO safety Standards. 		
	[Replace the third-last paragraph with the following]		
	Non-ME EQUIPMENT, when used in an ME SYSTEM, shall comply with the CSA Group, IEC, or ISO safety Standards that are relevant to that equipment.		



		- 3	-1	
National Differences to IEC 60601-1:2005 + A1:2012				
Clause	Requirement + Test		Result - Remark	Verdict

16.9.2.1	MULTIPLE SOCKET-OUTLET	No multiple socket outlets	N/A
	d) If the MULTIPLE SOCKET-OUTLET is combined with a separating transformer, the following additional requirements apply:	provided.	
	- The separating transformer shall comply with this Standard. Alternatively, the separating transformer may comply with the requirements of CAN/CSA-E61558-2-1, except that the requirements of maximum RATED output power of 1 kVA and degree of protection IPX4 do not apply.		
	Note 1: As a separating transformer is not a MAINS SUPPLY TRANSFORMER, it does not require more than BASIC INSULATION.		
	Note 2: Limitation of output power is not explained in CAN/CSA-E61558-2-1 and the RATED output power is defined by the fuse in the installation and by the allowable power supply cable used. However, the characteristics of the separating transformer need to be carefully selected, taking into account the variations in the load current of the ME SYSTEM to ensure that the voltage supplied to the various items of the ME SYSTEM remains within the limits specified for the equipment.		
	 The separating transformer assembly shall be of CLASS I construction. 		
	 The degree of protection against ingress of water as given in IEC 60529 shall be specified. 		
	 The separating transformer assembly shall be marked according to the requirements of 7.2 and 7.3. 		
	 The MULTIPLE SOCKET-OUTLET shall be permanently connected to the separating transformer or 		
	the socket-outlet of the separating transformer assembly shall be of a type that cannot accept MAINS		
	PLUGS of any of the kinds identified in Canadian Electrical Code, Part I (see Figure I.1 and Figure I.2 of this Standard).		
	Compliance is checked by inspection and as described in the relevant subclauses of this Standard.		
	[Add the following item]		
	dA) The MULTIPLE SOCKET-OUTLET shall comply with the requirements of CSA C22.2 No. 42, CSA C22.2 No. 49, and Item d) of this Standard, as applicable.		



National Differences to IEC 60601-1:2005 + A1:2012				
Clause	Requirement + Test	Result - Remark	Verdict	

JAPAN edition	NATIONAL DIFFERENCES to IEC 60601-1 Third	Not applicable for switch mode power supplies.	N/A
Nationa	al standard: JIS T0601-1:2012		
1.1	At the end, add the following:		N/A
	JIS T0601-1:1999 is applicable until 2017.05.31.		
1.3	In NOTE 3, add the following:		N/A
	In Japan, to check the concerned JIS standard is required.		
1.4	At the end of NOTE, add the following:		N/A
	In Japan, application of the concerned JIS standard(s) is required.		
2	Except the part of the first paragraph, Attention and NOTE, replace the existing part listing standards with the following, and apply these properly in the following clauses if any:		N/A
	JIS B7761-3, Hand-transmitted vibration-Part 3: General requirements for measurement and evaluation		
	NOTE: ISO 5349-1, Mechanical vibration - Measurement and evaluation of human exposure to hand-transmitted vibration - Part 1: General requirements (IDT)		
	JIS B9707, Safety of machinery-Safety distances to prevent danger zones being reached by the upper limbs		
	NOTE: ISO 13852, Safety of machinery - Safety distances to prevent danger zones being reached by the upper limbs (IDT)		
	JIS B9711, Safety of machinery-Minimum gaps to avoid crushing of parts of the human body		
	NOTE: ISO 13854, Safety of machinery - Minimum gaps to avoid crushing of parts of the human body (IDT)		
	JIS C0445, Identification of equipment terminals and of terminations of certain designated conductors, including general rules for an alphanumeric system		
	NOTE: IEC 60445, Basic and safety principles for man- machine interface, marking and identification - Identification of equipment terminals and of terminations of certain designated conductors, including general rules for an alphanumeric system (IDT)		
	JIS C0447, Man-machine interface (MMI) - Actuating principles		
	NOTE: IEC 60447, Basic and safety principles for man- machine interface, marking and identification - Actuating principles (IDT)		
	JIS C0920:2003, Degrees of protection provided by enclosures (IP Code)		



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Clause	Requirement + Test	Result - Remark	Verdict
	NOTE: IEC 60529:2001, Degrees of protection provided by		
	enclosures (IP Code) (IDT) JIS C1509-1, Electroacoustitcs - Sound level meters- Part 1: Specifications		
	NOTE: IEC 61672-1, Electroacoustics - Sound level meters - Part 1: Specifications (IDT)		
	JIS C1509-2, Electroacoustics -Sound level meters - Part 2: Pattern evaluation tests		
	NOTE: IEC 61672-2, Electroacoustics - Sound level meters - Part 2: Pattern evaluation tests (IDT)		
	JIS C2134, Method for the determination of the proof and the comparative tracking indices of solid insulating materials		
	NOTE: IEC 60112, Method for the determination of the proof and the comparative tracking indices of solid insulating materials (IDT)		
	JIS C3301:2000, Rubber insulated flexible cords		
	NOTE: IEC 60245-4:1994, Rubber insulated cables of rated voltages up to and including 450/750 V - Part 4: Cords and flexible cables, Amendment 1:1997 (NEQ)		
	JIS C3306:2000, Polyvinyl chloride insulated flexible cords		
	NOTE: IEC 60227-5:1997, Polyvinyl chloride insulated cables of rated voltages up to and including 450/750 V - Part 5: Flexible cables (cords) (NEQ)		
	JIS C4003, Electrical insulation-Thermal evaluation and designation		
	NITE: IEC 60085, Electrical insulation - Thermal evaluation and designation (MOD)		
	JIS C5101-14:2009, Fixed capacitors for use in		

electronic equipment - Part 14: Sectional specification: Fixed capacitors for electromagnetic interference suppression and connection to the supply mains

NOTE: IEC 60384-14:2005, Fixed capacitors for use in electronic equipment - Part 14: Sectional specification: Fixed capacitors for electromagnetic interference suppression and connection to the supply mains (IDT)

JIS C6065:2007, Audio, video and similar electronic apparatus-Safety requirements

NOTE: IEC 60065:2001, Audio, video and similar electronic apparatus - Safety requirements (MOD)

JIS C6802:2005, Safety of laser products

NOTE: IEC 60825-1:1993, Safety of laser products - Part 1: Equipment classification, requirements and user's guide, Amendment 1:1997 and Amendment 2:2001 (IDT)

JIS C6965, Mechanical safety of cathode ray tubes

NOTE: IEC 61965, Mechanical safety of cathode ray tubes (IDT)



National Differences to IEC 60601-1:2005 + A1:2012				
Clause	Requirement + Test	Result - Remark	Verdict	

JIS C8282-1, Plugs and socket-outlets for household and similar purposes - Part 1: General requirements

NOTE: IEC 60884-1, Plugs and socket-outlets for household and similar purposes - Part 1: General requirements (MOD)

JIS C8303, Plugs and receptacles for domestic and similar general use

NOTE: No corresponding JIS exists. This standard has been listed as normative reference corresponding to IEC60083, Plugs and socket-outlets for domestic and similar general use standardized in member countries of IEC, which has been listed in IEC 60601-1:2005. Refer to JIS T1021, too.

JIS C60068-2-2:1995, Environmental testing - Part 2-2:Tests -Test B: Dry heat

NOTE: IEC 60068-2-2:1974, Environmental testing - Part 2: Tests. Tests B: Dry heat, Amendment 1:1993 and Amendment 2:1994 (IDT)

JIS C60079-0, Explosive atmospheres-Part 0: Equipment-General requirements

NOTE: IEC 60079-0, Electrical apparatus for explosive gas atmospheres - Part 0: General requirements (IDT)

JIS C60079-2, Electrical apparatus for explosive gas atmospheres - Part 2: Pressurized enclosures "p"

NOTE: IEC 60079-2, Electrical apparatus for explosive gas atmospheres - Part 2: Pressurized enclosures "p" (IDT)

JIS C60079-6, Electrical apparatus for explosive gas atmospheres - Part 6:Oil immersion "o"

NOTE: IEC 60079-6, Electrical apparatus for explosive gas atmospheres - Part 6: Oil-immersion "o" (IDT)

JIS C60364-4-41, Low-voltage electrical installations-Part 4-41: Protection for safety - Protection against electric shock

NOTE: IEC 60364-4-41, Electrical installations of buildings -Part 4-41: Protection for safety - Protection against electric shock (IDT)

JIS C60664-1:2009, Insulation coordination for equipment within low-voltage systems - Part 1:Principles, requirements and tests

NOTE: IEC 60664-1:2007, Insulation coordination for equipment within low-voltage systems - Part 1: Principles, requirements and tests (IDT)

JIS C60695-11-10, Fire hazard testing-Part11-10:Test flames-50W horizontal and vertical flame test methods

NOTE: IEC 60695-11-10, Fire hazard testing - Part 11-10: Test flames - 50 W horizontal and vertical flame test methods (IDT)

JIS T0307, Medical devices-Symbols to be used with medical device labels, labelling and



		3	-1		
National Differences to IEC 60601-1:2005 + A1:2012					
Clause	Requirement + Test		Result - Remark	Verdict	t

information to be supplied

NOTE: ISO 15223, Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied (IDT)

JIS T0601-1-3, Medical electrical equipment-Part 1-3: General requirements for basic safety and essential performance-Collateral Standard: Radiation protection in diagnostic X-ray equipment

NOTE: IEC60601-1-3, Medical electrical equipment - Part 1: General requirements for safety - 3. Collateral standard: General requirements for radiation protection in diagnostic X-ray equipment (IDT)

JIS T14971:2003, Medical devices-Application of risk management to medical devices

NOTE: ISO 14971:2000, Medical devices - Application of risk management to medical devices (IDT)

JIS Z8202 (all parts), Quantities and units

NOTE: ISO 31 (all parts), Quantities and units (IDT)

JIS Z8203, SI units and recommendations for the use of their multiples and of certain other units

NOTE: ISO 1000, SI units and recommendations for the use of their multiples and of certain other units (IDT)

JIS Z8736-1, Acoustics - Determination of sound power levels of noise sources using sound intensity - Part 1 : Measurement at discrete points

NOTE: ISO 9614-1, Acoustics - Determination of sound power levels of noise sources using sound intensity - Part 1: Measurement at discrete points (IDT)

JIS Z9101:2005, Safety colours and safety signs-Design principles for safety signs in workplaces and public areas

NOTE: ISO 3864-1:2002, Graphical symbols - Safety colours and safety signs - Part 1: Design principles for safety signs in workplaces and public areas (IDT)

ISO 780, Packaging - Pictorial marking for handling of goods

NOTE: The corresponding JIS standard is JIS Z0150 Packaging-Pictorial marking for handling of goods (MOD)

ISO 1853, Conducting and dissipative rubbers, vulcanized or thermoplastic—Measurement of resistivity

NOTE: The corresponding JIS standard is JIS K6271 Rubber, vulcanized or thermoplastic-Determination of volume and surface resistivity (MOD)

ISO 2878, Rubber - Antistatic and conductive products - Determination of electrical resistance



National Differences to IEC 60601-1:2005 + A1:2012				
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ISO 2882, Rubber, vulcanized - Antistatic and conductive products for hospital use - Electrical resistance

Limits

ISO 3746, Acoustics - Determination of sound power levels of noise sources using sound pressure – Survey method using an enveloping measurement surface over a reflecting plane

ISO 7000-DB:2004, Graphical symbols for use on equipment - Index and synopsis

ISO 7010:2003, Graphical symbols - Safety colours and safety signs - Safety signs used in workplaces and public areas

ISO 10993 (all parts), Biological evaluation of medical devices

NOTE: The corresponding JIS standard is JIS T0993-1 Biological evaluation of medical devices-Part 1: Evaluation and testing within a risk management process (MOD). However, other Parts than Part 1 and Part 7 have still not been published as JIS.

ISO 11134, Sterilization of health care products
- Requirements for validation and routine
control -Industrial moist heat sterilization

NOTE: At present, as the corresponding JIS or international standards, the following exist:

JIS T0816-1:2010 Sterilization of health care products - Moist heat - Part 1: Requirements for the development, validation and routine control of a sterilization process for medical devices

ISO 17665-1:2006, Sterilization of health care products - Moist heat - Part 1: Requirements for the development, validation and routine control of a sterilization process for medical devices (IDT)

ISO 11135, Medical devices - Validation and routine control of ethylene oxide sterilization

NOTE: At present, as the corresponding JIS or international standards, the following exist:

JIS T0801-1:2010 Sterilization of health care products - Ethylene oxide - Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices

ISO 11135-1:2007, Sterilization of health care products -Ethylene oxide - Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices (IDT)

ISO 11137, Sterilization of health care products
- Requirements for validation and routine
control – Radiation Sterilization

NOTE: At present, as the corresponding JIS or international standards, the following exist:

JIS T0806-1:2010 Sterilization of health care products - Radiation - Part 1: Requirements for development, validation and routine control of a sterilization process for



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	National Differences to IEC 60601-1	1:2005 + A1:2012	
Clause	Requirement + Test	Result - Remark	Verdict
	medical devices	1	
	ISO 11137-1:2006, Sterilization of health care products - Radiation - Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices (IDT)		
	ISO 23529, Rubber - General procedures for preparing and conditioning test pieces for physical test methods		
	NOTE: The corresponding JIS standard is JIS K6250 Rubber-General procedures for preparing and conditioning test pieces for physical test methods (MOD)		
	IEC 60079-5, Explosive gas atmospheres — Part 5: Equipment protection by powder filling "q"		
	IEC/TR 60083, Plugs and socket-outlets for domestic and similar general use standardized in member countries of IEC		
	IEC 60086-4, Primary batteries - Part 4: Safety of lithium batteries		
	NOTE: The corresponding JIS standard is JIS C8513 Safety of primary lithium batteries (MOD)		
	IEC 60127-1, Miniature fuses - Part 1: Definitions for miniature fuses and general requirements for miniature fuse-links		
	NOTE: The corresponding JIS standard is JIS C6575-1 Miniature fuses-Part 1: Definitions of miniature fuses and general requirements for miniature fuse-links (MOD)		
	IEC 60227-1:1993, Polyvinyl chloride insulated cables of rated voltages up to and including 450/750 V – Part 1: General requirements, Amendment 1:1995 and Amendment 2:1998		
	NOTE: The corresponding JIS standard is JIS C3662-1:2009 Polyvinyl chloride insulated cables of rated voltages up to and including 450/750V - Part 1: General requirements (MOD)		
	IEC 60245-1:2003, Rubber insulated cables - Rated voltages up to and including 450/750 V - Part 1: General requirements		
	NOTE TI		

NOTE: The corresponding JIS standard is JIS C3663-1:2007 Rubber insulated cables-Rated voltages up to and including 450/750 V-Part 1: General requirements (MOD)

IEC 60252-1, AC motor capacitors - Part 1: General - Performance, testing and rating -Safety requirements -Guide for installation and operation

IEC 60320-1, Appliance couplers for household and similar general purposes - Part 1: General requirements

NOTE: The corresponding JIS standard is JIS C8283-1 Appliance couplers for household and similar general purposes-Part 1: General requirements (MOD)

IEC 60335-1:2001, Household and similar



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electrical appliances - Safety - Part 1: General requirements NOTE: The corresponding JIS standard is JIS C9335-1:2003 Household and similar electrical appliances - Safety - Part 1 **General requirements (MOD)** IEC 60417-DB:2002, Graphical symbols for use on equipment IEC 60601-1-2, Medical electrical equipment -Part 1 - 2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility -Requirements and tests NOTE: The current "JIS T0601-1-2:2012 Medical electrical equipment - Part 1-2: General requirements for safety -Electromagnetic compatibility - Requirements and tests" corresponds to IEC 60601-1-2:2001 and Amendment 1:2004. IEC 60601-1-6, Medical electrical equipment -Part 1 - 6: General requirements for basic safety and essential performance - Collateral standard: Usability NOTE: As the corresponding international standard, IEC 62336 is applicable. IEC 60601-1-8, Medical electrical equipment -Part 1 - 8: General requirements for basic safety and essential performance - Collateral standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems NOTE: The corresponding JIS standard is now under drafting. IEC 60730-1:1999, Automatic electrical controls for household and similar use - Part 1: General requirements, Amendment 1:2003 and Amendment 2:2007 NOTE: The corresponding JIS standard is JIS C9730-1:2010 Automatic electrical controls for household and similar use-Part 1:General requirements (MOD) IEC 60851-3:1996, Winding wires - Test methods - Part 3: Mechanical properties, Amendment 1:1997 and Amendment 2:2003 IEC 60851-5:1996, Winding wires - Test methods - Part 5: Electrical properties, Amendment 1:1997 and Amendment 2:2004 IEC 60851-6:1996, Winding wires - Test methods - Part 6: Thermal properties and Amendment 1:1997 IEC 60878:2003, Graphical symbols for electrical equipment in medical practice IEC 60884-1, Plugs and socket-outlets for household and similar purposes - Part 1:



National Differences to IEC 60601-1:2005 + A1:2012			
Clause	Requirement + Test	Result - Remark	Verdict
	General requirements		
	IEC 60950-1:2001, Information technology equipment – Safety - Part 1: General requirements		
	NOTE: The corresponding JIS standard is JIS C 6950-1:2009 Information technology equipment - Safety - Part 1: General requirements (MOD)		
	IEC 61058-1:2000, Switches for appliances - Part 1: General requirements and Amendment 1:2001		
	NOTE: The corresponding JIS standard is JIS C4526-1:2005 Switches for appliances - Part 1: General requirements (MOD)		
	IEC 61558-1:1997, Safety of power transformers, power supply units and similar products - Part 1: General requirements and tests and Amendment 1:1998		
	NOTE: No corresponding JIS exists. However, as the standard corresponding to IEC 61558-1:2005, the following exists:		
	JIS C 61558-1:2008 Safety of power transformers, power supplies, reactors and similar products - Part 1: General requirements and tests (MOD)		
	IEC 61558-2-1, Safety of power transformers, power supplies, reactors and similar products - Part 2-1: Particular requirements and tests for separating transformers and power supplies incorporating separating transformers for general applications		
	NOTE: The corresponding JIS standard is JIS C61558-2-1 Safety of power transformers, power supplies, reactors and similar products-Part 2-1: Particular requirements and tests for separating transformers and power supplies incorporating separating transformers for general applications (MOD)		
3.61	Add NOTE as follows:		N/A
	NOTE In this standard, MECHANICAL HAZARD is understandable suitably by replacing with mechanical HAZARD, mechanical HADARDOUS SITUATION, HARM or unacceptable RISK.		
3.70	Replace the existing text with:		N/A
	condition in which all means provided for protection against HAZARDOUS SITUATION or HAZARDS are intact		





National Differences to IEC 60601-1:2005 + A1:2012				
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4.2	Replace the existing NOTE 2 with the following:	N/A
	NOTE 2 Conditions or faults that can give rise to HAZARDOUS SITUATIONS are identified in the clauses of this standard. In these cases, it will often be necessary to carry out a RISK MANAGEMENT PROCESS to determine what the actual HAZARDOUS SITUATIONS are and the tests that need to be done to show that the identified HAZARDOUS SITUATIONS do not arise in the specified circumstances.	
4.10.1	In the existing text, replace "a separate power supply" with "a separate power supply (e.g., a power supply of other equipment)".	N/A
7.3.3	In the third paragraph, replace "could result in a HAZARD" with "could result in a HAZARDOUS SITUATION".	N/A
7.4.3	Replace the existing first paragraph with the following:	N/A
	Numeric indications of parameters on ME EQUIPMENT shall be expressed in SI units according to JIS Z8202 (ISO 31 (IDT)) except the base quantities listed in Table 1 may be expressed in the indicated units, which are used in conjunction with the SI units system or as the approved combination.	
	Replace the title of Table 1 with the following:	
	Units which are used in conjunction with the SI units system or as the approved combination	
7.7.4	Under the existing text, add the following:	N/A
	If polyvinyl chloride insulated flexible cord of JIS C3306 or rubber insulated flexible cord of JIS C3301 is used, the conductor may be coloured "white".	
7.7.5	Under the existing text, add the following:	N/A
	If polyvinyl chloride insulated flexible cord of JIS C3306 or rubber insulated flexible cord of JIS C3301 is used, conductors may be of the colour specified in the said standards.	
7.9.3.2	In the fourth dash, replace "the nature of the HAZARD" with "the HAZARDOUS SITUATION".	N/A



National Differences to IEC 60601-1:2005 + A1:2012			
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T	
	N/A
	N/A
	N/A
	N/A



National Differences to IEC 60601-1:2005 + A1:2012			
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8.9.1.12	At the end of the title of this sub-clause, add "(Apply to MOOP)".	 N/A
8.9.1.13	At the end of the title of this sub-clause, add "(Apply to MOOP)".	 N/A
8.9.1.14	At the end of the title of this sub-clause, add "(Apply to MOOP)".	 N/A
8.11.3.2	Add the following between the first paragraph and the second paragraph:	 N/A
	And, rubber insulated flexible cords of JIS C3301, polyvinyl chloride insulated flexible cords of JIS C3306 or cords of which the robustness is equal to or more than those are usable.	
	Add the following between the second paragraph and the last paragraph:	
	And, in the case of cords of JIS C3306, shall not use;	
	 for polyvinyl chloride insulated flexible cords, if the temperature of the above- mentioned external metal part exceeds 60 °C, and; 	
	 for grade heat-resistant polyvinyl chloride insulated flexible cords, if the temperature of the above-mentioned external metal part exceeds 75 °C. 	
9.2.2.2	In the bottom column of Table 20, replace the existing text with the following:	 N/A
	The values in this table are taken from JIS B9711 (ISO 13854 (IDT)).	
9.2.2.4.4	In the second dash, replace "no HAZARD or damage shall result" with "no HAZARDOUS SITUATION or unacceptable RISK shall result".	 N/A
9.2.4	In e), replace "no HAZARD or damage shall result" with "no HAZARDOUS SITUATION or unacceptable RISK shall result".	 N/A
9.4.4	In the first paragraph of a), replace "and no HAZARDS can develop" with "and no HAZARDOUS SITUATION can develop".	 N/A
9.7.5	In the last paragraph, delete "unmarked".	 N/A
9.8.4.1	Replace the existing NOTE with the following:	 N/A
	NOTE The upper carriage of the human body test mass apparatus is formed of wood or a similar material. The bottom portion is foam. The resiliency or spring factor of the foam (ILD or IFD ratings) has not been specified. The foam is cylindrical, rather than spherical.	



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10.1.1	In the paragraph, replace "0,5 mR/h" with "0,5 mR/h \approx 5 μ Gy/h"; and in NOTE 2, "0,1 mR/h" with "0,5 mR/h \approx 1 μ Gy/h".		N/A
11.1.1	To the existing text of a in the Table 22, add the following:		N/A
	(For example, the maximum temperature limit of a transformer with three insulating materials of Class A, Class B and Class E shall be 105 °C of Class A of the lowest limit.)		
13.2.7	In the title of this sub-clause, replace "in a HAZARD" with "in a HAZARDOUS SITUATION".		N/A
13.2.10	In Table 26, replace the existing NOTE with the following:		N/A
	NOTE The temperature limits in this table were derived from Table B.1 of IEC 60950-1:2001 (in the corresponding international standard, IEC 61010-1:2001 [22]).		
15.4.2.1	In c), replace "could constitute a HAZARD" with "could constitute a HAZARDOUS SITUATION".		N/A
15.4.3.4	In the first paragraph, replace "could become a HAZARD" with "could become a HAZARDOUS SITUATION".		N/A
16.1	Replace the last two paragraphs with the following:		N/A
	Otherwise, non-medical equipment shall be those which are in compliance with relevant JIS standards or the Technical Requirements of the Electrical Appliance and Material Safety Act or which ensure safety equivalent to the said standards/technical requirements.		
	Equipment in which protection against electric shock relies only on BASIC INSULATION shall not be used in an ME SYSTEM.		
	For the measures for ensuring safety, e.g., the case combined with a separating transformer with DOUBLE INSULATION or RAINFORCED INSULATION, equipment only with BASIC INSULATION may be used.		
	Compliance is checked by inspection of appropriate documents or certificates.		
16.6.4.1	In NOTE, replace "no possibility of any HAZARD" with "no possibility of any HAZARDOUS SITUATION".		N/A
16.9.2.1	In the text of c), replace "IEC 60884-1" with "IEC 60884-1 or JIS C8282-1".		N/A



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Annex D	In Table D.2, replace the sign of No. 10, which is shown as "IEC 60878 Safety 01 b", with the sign of "ISO 7010-M002 b".	 N/A
	In the bottom column if Table D.2, replace the existing a and b with the following:	
	^a The description of this commonly used safety sign appeared in Annex B of ISO 3864:1984.	
	b In accordance with the corrigendum of IEC 60601-1, Replaced "IEC 60878 Safety 01 " with "ISO 7010-M002	
Annex I	In 1.1.3, replace the first dash with the following:	 N/A
	- PATIENTS should only be connected to APPLIED PARTS of ME EQUIPMENT complying with this standard. Other equipment should comply with relevant IEC or ISO standards or comply with relevant JIS safety standards or the Technical Requirements of the Electrical Appliance and Material Safety Act, or ensure safety equivalent to the said standards/technical requirements.	
	Replace the existing NOTE 2 with the following:	
	NOTE 2 IEC 60601: MEDICAL ELECTRICAL EQUIPMENT in compliance with IEC 60601 (all parts) or JIS T0601 (all parts).	
	Replace the existing NOTE 3 with the following:	
	NOTE 3 IEC xxxxx: Non-medical equipment in compliance with relevant IEC safety standards. Include non-medical equipment in compliance with relevant JIS safety standards or the Technical Requirements of the Electrical Appliance and Material Safety Act, or non-medical equipment ensuring safety equivalent to the said standards/technical requirements.	
Annex L	In the first paragraph, replace "wound components" with "wound components (e.g., transformers, motors, etc.)"	 N/A
Bibliogra	Add the following at the end:	 N/A
phy	[55] JIS T1021, "Hospital grade" outlet-sockets and plugs	
	[56] JIS Q13485, Medical devices - Quality management systems - Requirements for regulatory purposes	

SWITZERLAND NATIONAL DIFFERENCES to IEC 60601-1 Third edition National standard SN EN 60601-1:06



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N/A N/A N/A
N/A
N/A
N/A
N/A
N/A
N/A



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EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
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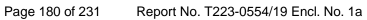
Annex ZA	Annex ZA		Р
	(normative)		
	Normative references to international publications with their corresponding European publications		
	The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.	Considered.	Р
	NOTE When an international publication has been modified by common modifications, indicated by (mod), the relevant EN/HD applies.	Considered.	Р



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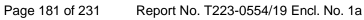
Publication	Voor	Titlo	EN/UD and IEC/IS/	Voor
Publication IEC 60065 (mod)	<u>Year</u> 2001	<u>Title</u> Audio, video and similar electronic	EN/HD and IEC/ISO EN 60065	2002
		apparatus - Safety requirements	+ corr. March	
A1 (mod)	2005		A1 + corr. August A11	2006 2007 2008
A2 (mod	2010		A2 A12	2010 2012
IEC 60068-2-2	2007	Environmental testing Part 2: Tests - Test B: Dry heat	EN 60068-2-2	2007
IEC 60079-0 (mod)	- 1)	Explosive atmospheres - Part 0: Equipment - General requirements	EN 60079-0	2012
IEC 60079-2	_ 1)	Explosive atmospheres - Part 2: Equipment protection by pressurized enclosure "p"	EN 60079-2	2007
IEC 60079-5	_ 1)	Explosive atmospheres - Part 5: Equipment protection by powder filling "q"	EN 60079-5	2007
IEC 60079-6	_ 1)	Explosive atmospheres - Part 6: Equipment protection by oil immersion "o"	EN 60079-6	2007
IEC 60083	_ 1)	Plugs and socket-outlets for domestic and similar general use standardized in member countries of IEC	IEC 60083	2009
IEC 60085	- 1)	Electrical insulation - Thermal evaluation and designation	EN 60085	2008
IEC 60086-4	- 1)	Primary batteries Part 4: Safety of lithium batteries	EN 60086-4	2007
IEC 60112	- 1)	Method for the determination of the proof and the comparative tracking indices of solid insulating materials	EN 60112	2003
IEC 60127-1	2006	Miniature fuses Part 1: Definitions for miniature fuses and general requirements for miniature fuse-links	EN 60127-1	2006





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	Publication IEC 60227-1 ²⁾ A1 A2	<u>Year</u> 1993 1995 1998	<u>Title</u> Polyvinyl chloride insulated cables of rated voltages up to and including 450/750 V Part 1: General requirements	EN/HD and IEC/ISC HD 21.1 S4	2002
	IEC 60245-1 3)	2003	Rubber insulated cables - Rated voltages up to and including 450/750 V Part 1: General requirements	IEC 60245-1	2003
	IEC 60252-1	- 1)	AC motor capacitors Part 1: General - Performance, testing and rating - Safety requirements - Guidance for installation and operation	EN 60252-1	2011
	IEC 60320-1	- 1)	Appliance couplers for household and similar general purposes Part 1: General requirements	rEN 60320-1	2001
	IEC 60335-1 (mod)	2010	Household and similar electrical appliances - Safety Part 1: General requirements	EN 60335-1	2012
	IEC 60364-4-41 (mod)	2005	Low-voltage electrical installations Part 4-41: Protection for safety - Protection against electric shock	HD 60364-4-41	2006
	IEC 60384-14	2005	Fixed capacitors for use in electronic equipment Part 14: Sectional specification - Fixed capacitors for electromagnetic interference suppression and connection to the supply mains	EN 60384-14	2005
	IEC 60417	Data base	Graphical symbols for use on equipment available from http://www.graphical-symbols.info/equipment	IEC 60417	2004
	IEC 60445	_ 1)	Basic and safety principles for man-machine interface, marking and identification - Identification of equipment terminals, conductor terminations and conductors	EN 60445	2010
	IEC 60447	- 1)	Basic and safety principles for man-machine interface, marking and identification - Actuating principles	EN 60447	2004
	IEC 60529	1989	Degrees of protection provided by enclosures (IP Code)	EN 60529 + corr. May	1991 1993
	A1	1999		A1	2000
	IEC 60601-1-2	- 1)	Medical electrical equipment Part 1-2: General requirements for basic safety and essential performance - Collateral standard: Electromagnetic compatibility - Requirements and tests	EN 60601-1-2	2007
	IEC 60601-1-3	_1)	Medical electrical equipment - Part 1-3: General requirements for basic safety and essential performance - Collatera	EN 60601-1-3 I+ corr. March	2008 2010
	A1	_1)	Standard: Radiation protection in diagnostic X-ray equipment	A1	2013
1					

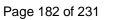




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 Clause
 Requirement + Test
 Result - Remark
 Verdict

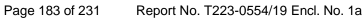
				1
Publication IEC 60601-1-6	Year -1)	Title Medical electrical equipment Part 1-6: General requirements for basic safety and essential performance - Collateral	EN/HD and IEC/ISC EN 60601-1-6	<u>2010</u>
IEC 60601-1-8	1)	standard: Usability	EN 60601-1-8	2007
IEC 60601-1-6		Medical electrical equipment Part 1-8: General requirements for basic safety and essential performance - Collateral Standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	+ corr. March	2010
A1	_1)		A1	2013
IEC 60664-1	2007	Insulation coordination for equipment within low-voltage systems Part 1: Principles, requirements and tests	EN 60664-1	2007
IEC 60695-11-10	- 1)	Fire hazard testing Part 11-10: Test flames - 50 W horizontal	EN 60695-11-10	1999
A1		and vertical flame test methods		2003
IEC 60730-1 (mod)	2010	Automatic electrical controls for household	EN 60730-1	2011
izo corco i (ilica)	2010	and similar use Part 1: General requirements	2.1 00100 1	2011
IEC 60825-1	2007	Safety of laser products Part 1: Equipment classification and requirements	EN 60825-1	2007
IEC 60851-3	2009	Winding wires - Test methods Part 3: Mechanical properties	EN 60851-3	2009
IEC 60851-5	2008	Winding wires - Test methods Part 5: Electrical properties	EN 60851-5	2008
IEC 60851-6 A1	1996 1997	Winding wires - Test methods Part 6: Thermal properties	EN 60851-6 A1	1996 1997
IEC/TR 60878	2003	Graphical symbols for electrical equipment in medical practice	IEC/TR 60878	2003
IEC 60884-1	- ¹⁾	Plugs and socket-outlets for household and similar purposes Part 1: General requirements	IEC 60884-1	2013
IEC 60950-1 (mod)	2001	Information technology equipment - Safety Part 1: General requirements	EN 60950-1 + corr. April A11	2001 2004 2004
IEC 61058-1 (mod) A1 A2	2000 2001 2007	Switches for appliances Part 1: General requirements	EN 61058-1 A2	2002 2008
IEC 61558-2-1	_ 1)	Safety of power transformers, power supplies, reactors and similar products - Part 2-1: Particular requirements and tests for separating transformers and power supplies incorporating separating transformers for general applications	EN 61558-2-1	2007
IEC 61672-1	- 1)	Electroacoustics - Sound level meters Part 1: Specifications	EN 61672-1	2003
IEC 61672-2	- 1)	Electroacoustics - Sound level meters Part 2: Pattern evaluation tests	EN 61672-2	2003





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Publication IEC 61965 IEC 62133	<u>Year</u> -1) -1)	<u>Title</u> Mechanical safety of cathode ray tubes Secondary cells and batteries containing alkaline or other non-acid electrolytes – Safety requirements for portable sealed secondary cells, and for batteries made from them, for use in portable applications	EN/HD and IEC/ISC EN 61965 EN 62133	<u>2003</u> 2013
IEC 62304	2006	Medical device software – Software lifecycle processes	EN 62304 + corr. November	2006 2008
ISO 780	- 1)	Packaging - Pictorial marking for handling of goods	EN ISO 780	1999
ISO 1853	- 1)	Conducting and dissipative rubbers, vulcanized or thermoplastic - Measurement of resistivity	ISO 1853	2011
ISO 2878	- ¹⁾	Rubber, vulcanized - Antistatic and conductive products - Determination of electrical resistance	ISO 2878	2011
ISO 2882	- ¹⁾	Rubber, vulcanized - Antistatic and conductive products for hospital use - Electrical resistance limits	ISO 2882	1997
ISO 3746	-1)	Acoustics - Determination of sound power levels of noise sources using sound pressure - Survey method using an enveloping measurement surface over a reflecting plane	EN ISO 3746	2010

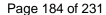




 EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)

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	Publication ISO 3864-1	<u>Year</u> 2002	<u>Title</u> Graphical symbols - Safety colours and	EN/HD and IEC/ISO ISO 3864-1	<u>OYear</u> 2011
			safety signs Part 1: Design principles for safety signs in workplaces and public areas		
	ISO 7000	2004	Graphical symbols for use on equipment – Collection of symbols	ISO 7000	2004
	ISO 7010	2011	Graphical symbols – Safety colours and safety signs – Registered safety signs	EN ISO 7010	2012
	ISO 9614-1	- 1)	Acoustics – Determination of sound power levels of noise sources using sound intensity – Measurement at discrete points	EN ISO 9614-1	2009
	ISO 10993 all parts	- 1)	Biological evaluation of medical devices	See list below	
	ISÓ 10993-1	2009	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process	EN ISO 10993-1	2009
	+ corr June	2010	-		
	ISO 10993-2	2006	Biological evaluation of medical devices - Part 2: Animal welfare requirements	EN ISO 10993-2	2006
	ISO 10993-3	2003	Biological evaluation of medical devices - Part 3: Tests for genotoxicity, carcinogenicity	EN ISO 10993-3 /	2003
	ISO 10994-4	2002	and reproductive toxicity Biological evaluation of medical devices - Part 4: Selection of tests for interactions with blood	1	
	A1	2006		EN ISO 10993-4	2009
	ISO 10993-5	2009	Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity	EN ISO 10993-5	2009
	ISO 10993-6	2007	Biological evaluation of medical devices - Part 6: Tests for local effects after implantation	EN ISO 10993-6	2009
	ISO 10993-7	2008	Biological evaluation of medical devices - Part 7: Ethylene oxide sterilization residuals	EN ISO 10993-7	2008
	+ corr November	2009		+ AC	2009
	ISO 10993-9	2009	Biological evaluation of medical devices - Part 9: Framework for identification and quantification of potential degradation products	EN ISO 10993-9	2009
	ISO 10993-10	2010	Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization	EN ISO 10993-10	2010
	ISO 10993-11	2006	Biological evaluation of medical devices - Part 11: Tests for systemic toxicity	EN ISO 10993-11	2009
	ISO 10993-12	2012	Biological evaluation of medical devices - Part 12: Sample preparation and reference materials	EN ISO 10993-12	2012
	ISO 10993-13	2010	Biological evaluation of medical devices - Part 13: Identification and quantification of degradation products from polymeric medica devices	EN ISO 10993-13	2010
	ISO 10993-14	2001	Biological evaluation of medical devices - Part 14: Identification and quantification of degradation products from ceramics	EN ISO 10993-14	2009
	ISO 10993-15	2000	Biological evaluation of medical devices - Part 15: Identification and quantification of degradation products from metals and alloys	EN ISO 10993-15	2009
	ISO 10993-16	2010	Biological evaluation of medical devices - Part 16: Toxicokinetic study design for degradation products and leachables	EN ISO 10993-16	2010
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EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)				
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	Publication ISO 10993-17	<u>Year</u> 2002	<u>Title</u> Biological evaluation of medical devices - Part 17: Establishment of allowable limits for	EN/HD and IEC/ISC EN ISO 10993-17	
	ISO 10993-18	2005	leachable substances Biological evaluation of medical devices - Part 18: Chemical characterization of materials	EN ISO 10993-18	2009
	ISO/TS 10993-19	2006	Biological evaluation of medical devices - Part 19: Physico-chemical, morphological and topographical characterization of materials	ISO/TS 10993-19	2006
	ISO/TS 10993-20	2006	Biological evaluation of medical devices - Part 20: Principles and methods for	ISO/TS 10993-20	2006
	ISO 11135-1	2007	immunotoxicology testing of medical devices Sterilization of health care products – Ethylene oxide – Part 1:Requirements for development, validation and routine control	EN ISO 11135-1	2007
	ISO 11137-1	2006	of a sterilization process formedical devices Sterilization of health care products – Radiation – Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices	EN ISO 11137-1	2006
	ISO 13857	2008	Safety of machinery – Safety distances to prevent hazard zones being reached by the upper and lower limbs	EN ISO 13857	2008
	ISO 14971	2007	Medical devices – Application of risk management to medical devices	EN ISO 14971	2012
	ISO 15223-1	2012	ISO 15223-1:2012, Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements	EN ISO 15223-1	2012
	ISO 17665-1	2006	Sterilization of health care products – Moist heat – Part 1: Requirements for the development, validation and routine control of a sterilization process for medical devices		2006
	ISO 23529	- 1)	Rubber – General process for medical devices and conditioning test pieces for physical test methods	ISO 23529	2010
	ISO 80000-1	2009	Quantities and units - Part 1: General	EN ISO 80000-1	2013

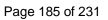
¹⁾ Undated reference, converted to dated reference in this European Standard.

Part 1: General requirements, which is related to, but not directly equivalent with, IEC 60227-1, applies instead.

3) HD 22.1 S4:2002, Cables of rated voltages up to and including 450/750 V and having cross-linked insulation – Part 1:General requirements, which is related to, but not directly equivalent with, IEC 60245-1, applies instead.

Annex	Annex ZZA	Power supply unit is not end	N/A
ZZA	(informative)	medical product.	
	Relationship between this European Standard and the Essential Requirements of EU Directive 93/42/EEC on Medical Devices		

²⁾ HD 21.1 S4:2002, Cables of rated voltages up to and including 450/750 V and having thermoplastic insulation –





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This European Standard has been prepared under a mandate given to CENELEC by the	 N/A
European Commission and the European Free Trade Association to provide a means of conforming to the Essential Requirements given in Annex I of the EC Directives 93/42/EEC as amended by 2007/47/EC.	
General Guidance:	 N/A
Once this standard will be cited in the Official Journal of the European Union under that Directive, compliance with the clauses of this standard given in Table ZZA.1 confers, within the limits of the scope of this standard, a presumption of conformity with the corresponding Essential Requirements (Ers) of that Directive and associated EFTA regulations.	 N/A
NOTE 1 This standard is intended to be applied in its entirety only. Selected clauses or subclauses may be not applicable due to the specific type of equipment under consideration. It is necessary to understand and apply Clauses 1 to 5. It is also recommended to understand and apply those clauses which contain general requirements related to a specific subclause. Elements of the standard that are not cited in Table ZZA.1 may be relevant for the appropriate fulfilment of certain essential requirements through indirect reference, and for safety and performance aspects of the device, that are not addressed through essential requirements.	 N/A
NOTE 2 Where a reference from a clause of this standard to the risk management process is made, the risk management process needs to be in compliance to the MDD (Directive 93/42/EEC amended by 2007/47/EC). This means that risks have to be reduced "as far as possible", "to a minimum", "to the lowest possible level", "minimized" or "removed", according to the wording of the corresponding essential requirement.	 N/A
NOTE 3 With respect to note 4 of clause 4.2.2 General requirement for risk management, the manufacturer's policy for determining acceptable risk must be in compliance with essential requirements 1, 2, 5, 6, 7, 8, 9, 11 and 12 of the directive.	 N/A



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	EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)				
Clause	Requirement + Test	Result - Remark	Verdict		
	NOTE 4 References in the clauses 3 to 17 or in the Annexes of this standard specify whether the normative references listed in Clause 2 as cited in Annex ZA are to be applied in whole or in part.		N/A		
	NOTE 5 This Annex ZZA is based on Normative References according to Annex ZA, replacing the references in the core text.		N/A		
	WARNING: Other requirements and other EU Directives and Regulations may be applicable to the product(s) falling within the scope of this standard.		N/A		

Table ZZA.1	Relationship between Essential Requirements of Directive 93/42/EEC amended by 2007/47/EC, and Clauses and Subclauses of this standard			
l.				
1.	General Guidance note 2 and 3 shall be observed	d		
1	The devices must be designed and manufactured in such a way that, when used under the conditions and for the purposes intended, they will not compromise the clinical condition or the safety of patients, or the safety and health of users or, where applicable, other persons, provided that any risks which may be associated with their intended use constitute acceptable risks when weighed against the benefits to the patient and are compatible with a high level of protection of health and safety. This shall include:	Not completely covered But If the manufacturer follows this standard in his design and manufacturing process, this European Standard gives a valuable set of technical requirements to assist in fulfilling this ER for equipment in the scope of this standard.		
	- reducing, as far as possible, the risk of use error due to the ergonomic features of the device and the environment in which the device is intended to be used (design for patient safety), and	Not covered See EN/IEC 60601-1-6, EN/IEC 62366, EN/IEC 60601-1-11 and EN/IEC 60601-1-12		
	- consideration of the technical knowledge, experience, education and training and where applicable the medical and physical conditions of intended users (design for lay, professional, disabled or other users).	Covered only for accompanying documents by: 7.9.1 Paragraphs 4 and 5, intended operator		
2.	General Guidance note 2 and 3 shall be observed			



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Clause	Requirement + Test	Result - Remark	Verdict
2	The solutions adopted by the manufacturer for the design and construction of the devices must conform to safety principles, taking account of the generally acknowledged state of the art.	1st paragraph: Covered only in respect of the funder the condition that 2nd par (including the following 3 bullets into account:	agraph
	principles in the following order: MI	8 Protection against electrical h ME equipment	nazards from
		9 Protection against mechanica ME equipment and ME systems	
		15 Construction of me equipme	ent
		2 nd paragraph (including the foll bullets)	lowing 3
		Not covered in the normative te	ext.
	- eliminate or reduce risks as far as possible (inherently safe design and construction),		
	- where appropriate take adequate protection measures including alarms if necessary, in relation to risks that cannot be eliminated,		
	- inform users of the residual risks due to any shortcomings of the protection measures adopted.		
3	The devices must achieve the performances intended by the manufacturer and be designed, manufactured and packaged in such a way that they are suitable for one or more of the functions referred to in Article 1 (2) (a), as specified by the manufacturer.	Not covered.	
4	The characteristics and performances referred to in Sections 1, 2 and 3 must not be adversely affected to such a degree that the clinical conditions and safety of the patients and, where applicable, of other persons are compromised during the lifetime of the device as indicated by the manufacturer, when the device is subjected to the stresses which can occur during normal conditions of use.	Not covered However, the standard provides procedure for the generation of that is necessary to document t device is in compliance with this	information that the
5.	General Guidance note 2 and 3 shall be observe	d	
5	The devices must be designed, manufactured and packed in such a way that their	Covered only in respect of the f	-
characteristics and performances during intended use will not be adversely affected during transport and storage taking acco	characteristics and performances during their intended use will not be adversely affected	Instructions and information promanufacturer	ovided by the
	during transport and storage taking account of the instructions and information provided by	7.2.17 Marking on protective pa	ackaging
	the manufacturer.	7.9.3.1 Technical description	
		15.3.7 Environmental influence	S
6.	General Guidance note 2 and 3 shall be observe	d	



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Clause	Requirement + Test		Result - Remark		Verdict

6	Any undesirable side-effect must constitute an acceptable risk when weighed against the performances intended.	Not covered.
6a	Demonstration of conformity with the essential requirements must include a clinical evaluation in accordance with Annex X.	Not covered.
II		
7.	Chemical, physical and biological properties	General Guidance note 2 and 3 shall be observed
7.1	The devices must be designed and manufactured in such a way as to guarantee the characteristics and performances referred to in Section I (3) on the 'General requirements'.	Not covered.
	Particular attention must be paid to:	
	regards toxicity and, where appropriate, flammability,	Partially covered in respect of the following:
		Toxicity:
		11.7 Biocompatibility, the manufacturer should apply the appropriate part of the EN ISO 10993 series
		13.1.2 Emissions, deformation of Enclosure or exceeding maximum temperature
		Flammability:
		11.2 Fire prevention
		11.3 Constructional requirements for fire enclosures
		11.4 ME equipment and ME systems intended for use with flammable anaesthetics
		Annex G Protection against hazards of ignition of flammable anaesthetic mixtures
	- the compatibility between the materials used	Not covered
	and biological tissues, cells and body fluids, taking account of the intended purpose of the device,	The manufacturer should apply the appropriate part of the EN ISO 10993 series
	 where appropriate, the results of biophysical or abelled research whose validity has been demonstrated beforehand. 	Not covered



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7.2	The devices must be designed, manufactured and packed in such a way as to minimize the risks posed by contaminants and residues to the persons involved in the transport, storage and use of the devices and to the patients, taking account of the intended purpose of the product. Particular attention must be paid to the tissues exposed and to the duration and frequency of exposure.	Not covered.
7.3	The devices must be designed and manufactured in such a way that they can be used safely with the materials, substances and gases with which they enter into contact during their normal use or during routine procedures;	Covered only for the physical properties dealt with in Subclauses: 11.2.2 ME equipment and ME systems used in conjunction with oxygen rich environments 11.2.3 Single fault conditions related to oxygen rich environments and 11.6.1, 11.6.2, 11.6.3, 11.6.4, 11.6.6, 11.6.7, 11.6.8 (Overflow, spillage, leakage, cleaning, disinfection, sterilization and compatibility with substances used)
	if the devices are intended to administer medicinal products they must be designed and manufactured in such a way as to be compatible with the medicinal products concerned according to the provisions and restrictions governing these products and that their performance is maintained in accordance with the intended use.	Not covered.
7.4	Where a device incorporates, as an integral part, a substance which, if used separately, may be considered to be a medicinal product as defined in Article 1 of Directive 2001/83/EC and which is liable to act upon the body with action ancillary to that of the device, the quality, safety and usefulness of the substance must be verified by analogy with the methods specified in Annex I to Directive 2001/83/EC.	Not covered.
	For the substances referred to in the first paragraph, the notified body shall, having verified the usefulness of the substance as part of the medical device and taking account of the intended purpose of the device, seek a scientific opinion from one of the competent authorities designated by the Member States or the European Medicines Agency (EMEA) acting particularly through its committee in accordance with Regulation (EC) No 726/2004 on the quality and safety of the substance including the clinical benefit/risk profile of the incorporation of the substance into the device. When issuing its opinion, the competent authority or the EMEA shall take into account	





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

the manufacturing process and the data related to the usefulness of incorporation of the substance into the device as determined by the notified body.

Where a device incorporates, as an integral part, a human blood derivative, the notified body shall, having verified the usefulness of the substance as part of the medical device and taking into account the intended purpose of the device, seek a scientific opinion from the EMEA, acting particularly through its committee, on the quality and safety of the substance including the clinical benefit/risk profile of the incorporation of the human blood derivative into the device. When issuing its opinion, the EMEA shall take into account the manufacturing process and the data related to the usefulness of incorporation of the substance into the device as determined by the notified body.

Where changes are made to an ancillary substance incorporated in a device, in particular related to its manufacturing process, the notified body shall be informed of the changes and shall consult the relevant medicines competent authority (i.e. the one involved in the initial consultation), in order to confirm that the quality and safety of the ancillary substance are maintained. The competent authority shall take into account the data related to the usefulness of incorporation of the substance into the device as determined by the notified body, in order to ensure that the changes have no negative impact on the established benefit/risk profile of the addition of the substance in the medical device.

When the relevant medicines competent authority (i.e. the one involved in the initial consultation) has obtained information on the ancillary substance, which could have an impact on the established benefit/risk profile of the addition of the substance in the medical device, it shall provide the notified body with advice, whether this information has an impact on the established benefit/risk profile of the addition of the substance in the medical device or not. The notified body shall take the updated scientific opinion into account in reconsidering its assessment of the conformity assessment procedure.



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7.5	The devices must be designed and manufactured in such a way as to reduce to a minimum the risks posed by substances leaking from the device.	Covered in respect of the following: 9.7 Pressure vessels and parts subject to pneumatic and hydraulic pressure,
	leaking from the device.	11.6.1 Protection against overflow, spillage, leakage, ingress of water or particulate matter, cleaning, disinfection and sterilization, compatibility with substances
		11.6.2 Overflow
		15.4.9 Oil containers
	Special attention shall be given to substances which are carcinogenic, mutagenic or toxic to reproduction, in accordance with Annex I to Council Directive 67/548/EEC of 27 June 1967 on the approximation of laws, regulations and administrative provisions relating to the classification, packaging and abelled of dangerous substances.	Not covered.
	If parts of a device (or a device itself) intended to administer and/or remove medicines, body liquids or other substances to or from the body, or devices intended for transport and storage of such body fluids or substances, contain phthalates which are classified as carcinogenic, mutagenic or toxic to reproduction, of category 1 or 2, in accordance with Annex I to Directive 67/548/EEC, these devices must be abelled on the device itself and/or on the packaging for each unit or, where appropriate, on the sales packaging as a device containing phthalates.	Not covered.
	If the intended use of such devices includes treatment of children or treatment of pregnant or nursing women, the manufacturer must provide a specific justification for the use of these substances with regard to compliance with the essential requirements, in particular of this paragraph, within the technical documentation and, within the instructions for use, information on residual risks for these patient groups and, if applicable, on appropriate precautionary measures.	Not covered.
7.6	Devices must be designed and manufactured in such a way as to reduce, as much as possible, risks posed by the unintentional ingress of substances into the device taking into account the device and the nature of the environment in which it is intended to be used.	Not covered.
8	Infection and microbial contamination	General Guidance note 2 and 3 shall be observed





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

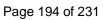
The devices and manufacturing processes must be designed in such a way as to eliminate or reduce as far as possible the risk of infection to the patient, user and third parties. The design must allow easy handling and, where necessary, minimize contamination of the device by the patient or vice versa during use.	Not covered.
Tissues of animal origin must originate from animals that have been subject to veterinary controls and surveillance adapted to the intended use of the tissues.	Not covered
Notified Bodies shall retain information on the geographical origin of the animals.	Not covered
Processing, preservation, testing and handling of tissues, cells and substances of animal origin must be carried out so as to provide optimal security. In particular safety with regard to viruses and other <i>transmissible</i> agents must be addressed by implementation of validated methods of elimination or viral inactivation in the course of the manufacturing process.	Not covered
Devices delivered in a sterile state must be designed, manufactured and packed in a non-reusable pack and/or according to appropriate procedures to ensure that they are sterile when placed on the market and remain sterile, under the storage and transport conditions laid down, until the protective packaging is damaged or opened.	Not covered
Devices delivered in a sterile state must have been manufactured and sterilized by an appropriate, validated method.	Not covered
Devices intended to be sterilized must be manufactured in appropriately controlled (e.g. environmental) conditions.	Not covered
Packaging system for non-sterile devices must	Covered in respect of
keep the product without deterioration at the level of cleanliness stipulated and, if the devices are to be sterilized prior to use,	7.2.17 Marking aspects of protective packaging
the packaging system must be suitable taking account of the method of sterilization indicated by the manufacturer.	Not covered
The packaging and/or label of the device must distinguish between identical or similar products sold in both sterile and non-sterile condition.	Not covered
	must be designed in such a way as to eliminate or reduce as far as possible the risk of infection to the patient, user and third parties. The design must allow easy handling and, where necessary, minimize contamination of the device by the patient or vice versa during use. Tissues of animal origin must originate from animals that have been subject to veterinary controls and surveillance adapted to the intended use of the tissues. Notified Bodies shall retain information on the geographical origin of the animals. Processing, preservation, testing and handling of tissues, cells and substances of animal origin must be carried out so as to provide optimal security. In particular safety with regard to viruses and other transmissible agents must be addressed by implementation of validated methods of elimination or viral inactivation in the course of the manufacturing process. Devices delivered in a sterile state must be designed, manufactured and packed in a nonreusable pack and/or according to appropriate procedures to ensure that they are sterile when placed on the market and remain sterile, under the storage and transport conditions laid down, until the protective packaging is damaged or opened. Devices delivered in a sterile state must have been manufactured and sterilized by an appropriate, validated method. Devices intended to be sterilized must be manufactured in appropriately controlled (e.g. environmental) conditions. Packaging system for non-sterile devices must keep the product without deterioration at the level of cleanliness stipulated and, if the devices are to be sterilized prior to use, minimize the risk of microbial contamination; the packaging system must be suitable taking account of the method of sterilization indicated by the manufacturer. The packaging and/or label of the device must distinguish between identical or similar products sold in both sterile and non-sterile



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9	Construction and environmental properties	General Guidance note 2 and 3 shall be observed
9.1	If the device is intended for use in combination	Covered in respect of the following:
	with other devices or equipment, the whole combination, including the connection system	9.1 Mechanical hazards
	must be safe and must not impair the specified	16.3 Power supply
	performances of the devices.	16.5 Separation devices
		16.6 Leakage currents
		16.8 Interruption of power supply
	Any restrictions on use must be indicated on the label or in the instructions for use.	Covered by 16.2 Accompanying documents of an ME system
9.2	Devices must be designed and manufactured in such a way as to remove or minimize as far as is possible:	
	- the risk of injury, in connection with their	Covered in respect of the following:
	physical features, including the volume/pressure ratio, dimensional and where	8.1 Electric shock
	appropriate ergonomic features;	9.1 Mechanical Hazards
		10 Radiation (all types)
		11.1 Excessive temperatures
		11.2 Fire prevention
	11	11.4 Flammable anaesthetics
	11.5 Flammable agent	
		11.6.3 Spillage
		11.8 Interruption of power supply
		12.4 Hazardous output
		13.1 Hazardous situations
		13.2 Single Fault condition
		15.3 Mechanical strength
		15.4 Components and general assembly
		15.5.3 Construction of transformers
		16.3 Power supply
		16.5 Separation devices
		16.6 Leakage currents
		16.8 Interruption of power supply





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

- risks connected with reasonably foreseeable environmental conditions, such as magnetic fields, external electrical influences, electrostatic discharge, pressure, temperature or variations in pressure and acceleration; - the risks of reciprocal interference with other devices normally used in the investigations or for the treatment given; - risks arising where maintenance or calibration are not possible (as with implants), from ageing of materials used or loss of accuracy of any measuring or control mechanism. 9.3 Devices must be designed and manufactured in such a way as to minimize the risks of fire or explosion during normal use and in single fault condition. Particular attention must be paid to devices whose intended use includes exposure to flammable substances or to substances which could cause combustion. Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered. See for EMC EN 60601-1-2 as re in Annex ZA Not covered in respect of the followin Mixtures Covered in respect of the followin Mixtures 11.4 Flammable anaesthetics Annex G ignition of flammable an Mixtures Annex G ignition of flammable an Mixtures	11 and Innex ZA g: according
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10 Devices with a measuring function	
10.1 Devices with a measuring function must be Not covered.	
designed and manufactured in such a way as to provide sufficient accuracy and stability See particular standards EN 6060	1-2-xx
within appropriate limits of accuracy and taking account of the intended purpose of the device. See 12.1 in respect of risks associated accuracy of controls and instruments and instruments accuracy.	
The limits of accuracy must be indicated by the manufacturer. Covered by 7.9.3.1 technical description.	iated with
10.2 The measurement, monitoring and display Not covered.	iated with nts
scale must be designed in line with ergonomic principles, taking account of the intended purpose of the device. See EN IEC 60601-1-6 and EN IEC principles in the intended purpose of the device.	iated with nts





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
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10.3	The managements made by decises with a	Covered in record of the fall available
measuring function must I	The measurements made by devices with a measuring function must be expressed in legal	Covered in respect of the following:
	units conforming to the provisions of Council	7.4.3 Units of measurement cmH2O is not included in 80/181/EEC
11	Protection against radiation	General Guidance note 2 and 3 shall be observed
11.1	General	
11.1.1	Devices shall be designed and manufactured in such a way that exposure of patients, users	For unintended radiation, covered in respect to the following:
	and other persons to radiation shall be reduced as far as possible compatible with the intended	10.1.1 (ionizing radiation),
	purpose, whilst not restricting the application	10.3 (microwave),
	of appropriate specified levels for therapeutic and diagnostic purposes.	10.4 (lasers).
		For intended radiation, covered in respect to the following:
		10.3 (microwave),
		10.4 (lasers).
		Other types of radiation of these devices and other devices not covered.
		For devices intended to produce radiation see EN 60601-1-3 for diagnostic x-radiation.
		For other radiation see particular standards EN 60601-2-xx.
11.2	Intended radiation	
11.2.1	Where devices are designed to emit hazardous levels of radiation necessary for a specific	1st and 2nd sentence covered in respect of the following:
	medical purpose the benefit of which is considered to outweigh the risks inherent in the emission, it must be possible for the user to control the emissions. Such devices shall be designed and manufactured to ensure reproducibility and tolerance of relevant variable parameters.	10.3, Microwave
		10.4 Lasers
		First sentence covered by subclauses 15.4.6, Actuating parts of controls and 15.4.7 hand or foot switches
		See particular standards EN 60601-2-xx
		See EN 60601-1-3 for diagnostic x-radiation
11.2.2	Where devices are intended to emit potentially hazardous, visible and/or invisible radiation, they must be fitted, where practicable, with visual displays and/or audible warnings of such emissions.	Not covered.
11.3	Unintended radiation	





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

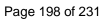
11.3.1	Devices shall be designed and manufactured in	Covered in respect to the following:
	such a way that exposure of patients, users and other persons to the emission of	10.1.1 (ionizing radiation),
	unintended, stray or scattered radiation is	10.3 (microwave),
	reduced as far as possible.	10.4 (lasers).
		Other types of radiation of these devices and other devices not covered.
11.4	Instructions	
11.4.1	The operating instructions for devices emitting radiation must give detailed information as to	Covered in respect of information relating to the nature of the emitted radiation:
	the nature of the emitted radiation, means of protecting the patient and the user and on ways of avoiding misuse and of eliminating the risks inherent in installation.	7.9.2.17 – ME equipment emitting radiation
11.5	Ionizing radiation	
11.5.1	Devices intended to emit ionizing radiation	Not covered.
	must be designed and manufactured in such a way as to ensure that, where practicable, the quantity, geometry and quality of radiation	For diagnostic x-radiation see EN 60601-1-3.
	emitted can be varied and controlled taking into account the intended use.	For other devices see particular standards EN 60601-2-xx
11.5.2	Devices emitting ionizing radiation intended for diagnostic radiology shall be designed and	Not covered For diagnostic x-radiation see EN 60601-1-3.
	manufactured in such a way as to achieve appropriate image and/or output quality for the intended medical purpose whilst minimizing radiation exposure of the patient and user.	For other devices see particular standards EN 60601-2-xx
11.5.3	Devices emitting ionizing radiation, intended for therapeutic radiology shall be designed and manufactured in such a way as to enable reliable monitoring and control of the delivered dose, the beam type and energy and where appropriate the quality of radiation.	Not covered.
12	Requirements for medical devices connected to or equipped with an energy source	General Guidance note 2 and 3 shall be observed
12.1	Devices incorporating electronic	Covered by
	programmable systems must be designed to ensure the repeatability, reliability and performance of these systems according to the intended use. In the event of a single fault condition (in the system) appropriate means should be adopted to eliminate or reduce as far as possible consequent risks.	14 Programmable electrical medical systems (PEMS)
12.1a	For devices which incorporate software or which are medical software in themselves, the software must be validated according to the state of the art taking into account the principles of development lifecycle, risk management, validation and verification.	Covered in respect of devices which incorporate SW by 14 Programmable electrical medical systems (PEMS)



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EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)				
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12.2	Devices where the safety of the patients depends on an internal power supply must be equipped with a means of determining the state of the power supply.	Not covered.
12.3	Devices where the safety of the patients depends on an external power supply must include an alarm system to signal any power failure.	Not covered.
12.4	Devices intended to monitor one or more clinical parameters of a patient must be equipped with appropriate alarm systems to alert the user of situations which could lead to death or severe deterioration of the patient's state of health.	Not covered
12.5	Devices must be designed and manufactured	Not covered
	in such a way as to minimize the risks of creating electromagnetic fields which could	EMC:
	impair the operation of other devices or equipment in the usual environment.	see EN 60601-1-2 as referenced in annex ZA
12.6	Protection against electrical risks	
12.6.1	Devices must be designed and manufactured	Covered in respect of the following:
	normal use and in single fault condition, provided the devices are installed correctly.	6.2 Protection against electric shock
		7.2.10 Applied parts
		7.9 Accompanying documents
		8 Protection against electrical hazard
		13.1 Specific hazardous situation
		13.2 Single fault conditions
		16.6 Leakage currents
12.7	Protection against mechanical and thermal risks	
12.7.1	Devices must be designed and manufactured	Covered in respect of the following:
	in such a way as to protect the patient and user against mechanical risks connected with, for	9.1 Mechanical Hazard
	example, resistance, stability and moving parts.	15.3 Mechanical strength
12.7.2	Devices must be designed and manufactured	Covered in respect of the following:
	in such a way as to reduce to the lowest possible level the risks arising from vibration	9.6 Acoustic energy and vibration
generated by the devices, taking account of technical progress and of the means availabl for limiting vibrations, particularly at source, unless the vibrations are part of the specified performance.	9.8.1 Support systems	





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
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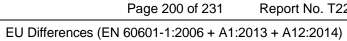
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12.7.3	Devices must be designed and manufactured in such a way as to reduce to the lowest	Covered in respect of
	possible level the risks arising from the noise emitted, taking account of technical progress and of the means available to reduce noise, particularly at source, unless the noise emitted is part of the specified performance.	9.6 Acoustic energy and vibration
12.7.4	gas or hydraulic and pneumatic energy	Covered in respect of the following:
		Electrical Risks:
	designed and constructed in such a way as to minimize all possible risks.	8.1 Fundamental rule of protection against electric shock
		8.2 Connection to power sources
		8.4 Limitation of voltage current or energy
		8,5 Separation of parts
		8.6 Functional earthing
		8.7 Leakage current
		8.11.3 Power supply cords
		Gas or Hydraulic and Pneumatic:
		9.7 Pressure vessels and parts
parts or areas intended to supply heat or re	Accessible parts of the devices (excluding the	Covered by
	given temperatures) and their surroundings must not attain potentially dangerous	11.1 Excessive temperatures
12.8	Protection against the risks posed to the patient by energy supplies or substances	
12.8.1	Devices for supplying the patient with energy	Covered in respect of the following:
	can be set and maintained accurately enough to guarantee the safety of the patient and of the user.	15.4.2 Temperature and overload control devices
		15.4.4 Indicators for standby and output
		15.4.6 Actuating parts of controls
		15.4.7 Cord-connected hand-held and foot- operated control devices
12.8.2	Devices must be fitted with the means of	Covered in respect of the following:
	preventing and/or indicating any inadequacies in the flow-rate which could pose a danger.	15.4.1 Construction of connectors
	and the state of t	15.4.2 Temperature and overload control devices
		15.4.4 indicators for standby and output
		15.4.5 Pre-set controls
		15.4.6 Actuating parts of controls
		15.4.7 Cord-connected hand-held and foot- operated control devices





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
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	prevent, as far as possible, the accidental release of dangerous levels of energy from an	Covered in respect of the following:
		Energy Source:
		12.4 Protection against hazardous output
		14 In respect of programmable electrical medical systems (PEMS)
		15.4.1 Construction of connectors
		15.4.2 Temperature and overload control devices
		15.4.4 Indicators for standby and output
		15.4.5 Pre-set controls
		15.4.6 Actuating parts of controls
		15.4.7 Cord-connected hand-held and foot- operated control devices
		Substance Source:
		9.7 Pressure vessels and parts subject to pneumatic and hydraulic pressure
		12.4 Protection against hazardous output
		14 In respect of programmable electrical medical systems (PEMS)
		15.4.4 Indicators for standby and output
		15.4.5 Pre-set controls
		15.4.6 Actuating parts of controls
12.9	The function of the controls and indicators	Covered in respect of the following:
	must be clearly specified on the devices	7.4 Marking of controls and instruments
	Where a device bears instructions required for	Covered in respect of the following:
	its operation or indicates operating or adjustment parameters by means of a visual	7.5 Safety signs
	system, such information must be understandable to the user and, as appropriate, the patient.	7.9.1 General requirements for accompanying documents
13	Information supplied by the manufacturer	
13.1	Each device must be accompanied by the	Covered in respect of the following:
	information needed to use it safely and properly, taking account of the training and knowledge of the potential users, and to	7.2.2 Identification
		7.2.4 Accessories
	identify the manufacturer.	7.2.5 Power from other equipment
	This information comprises the details on the label and the data in the instructions for use.	7.9 Accompanying documents



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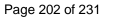
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Clause	Requirement + Test	Result - Remark	Verdict	
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	As far as practicable and appropriate, the information needed to use the device safely	Covered in respect of the following:		
	must be set out on the device itself and/or on	7.2.3 Consult accompanying do	cuments	
	the packaging for each unit or, where appropriate, on the sales packaging. If	7.9 Accompanying documents		
	individual packaging of each unit is not			
	practicable, the information must be set out in the leaflet supplied with one or more devices.			
	Instructions for use must be included in the	Covered in respect of the follow	/ina:	
	packaging for every device. By way of	Covered in respect of the following: 7.9.1 Accompanying documents, ge		
	exception, no such instructions for use are needed for devices in Class I or lia if they can	7.9.2 Instructions for use	s, general	
	be used safely without any such instructions.	7.9.2 Instructions for use		
13.2	Where appropriate, this information should	Covered in respect of the follow	/ing:	
	take the form of symbols. Any symbol or identification color used must conform to the	7.6 Symbols		
	harmonized standards.	Annex D Symbols on marking -	- informative	
	In areas for which no standards exist, the	annex for information only		
	symbols and colors must be described in the documentation supplied with the device.			
13.3	The label must bear the following particulars:	Covered in respect of the following:		
	the name or trade name and address of the	7.2.2 Identification (partially cov	· ·	
	manufacturer.	order to comply with this ER, no		
	view of their distribution in the Community, the label, or the outer packaging, or instructions for use, shall contain in addition the name and address of the authorized representative where the manufacturer does not have a registered place of business in the Community;	address must be used). Std. does not address the specifics of		
		imported devices (authorized		
		representative).		
		b) 7.2.2 Identification (limited to related to the identification of the		
			ie device)	
	(b) the details strictly necessary to identify the	c) 7.2.17 Protective packaging	occorios	
	device and the contents of the packaging especially for the users;	d) 7.2.2 Identification, 7.2.4 Accessories (the std. does not require to use the word		
	I where appropriate, the word 'STERILE';	LOT which has to be added)		
	(d) where appropriate, the batch code,	e) 7.2.2 Identification (std. does the format, however, the note d		
	preceded by the word 'LOT', or the serial number;	standard that specifies the form		
	(e) where appropriate, an indication of the date	f) 7.2.1 Marking (std. allows three	ee options,	
	by which the device should be used, in safety,	manufacturer needs to limit him one)	self on just	
	expressed as the year and month;	g) Not covered		
	(f) where appropriate, an indication that the device is for single use. A manufacturer's	h) Not covered		
	indication of single use must be consistent	i) 7.2.17 Protective packaging		
	across the Community;	i) Covered:		
	(g) if the device is custom-made, the words 'custom-made device';	7.2 Marking on the outside of e	auipment	
	(h) if the device is intended for clinical	and parts 7.3 Marking on the inside of equipmer		
	investigations, the words 'exclusively for			
	clinical investigations';			





	EU Differences (EN 60601-1:2006 + A1	I:2013 + A12:2014)	
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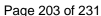
	(i) any special storage and/or handling	parts
	conditions;	7.5 Safety signs
	(j) any special operating instructions;	k) Covered:
	(k) any warnings and/or precautions to take;	7.2.2 Identification
	(I) year of manufacture for active devices other than those covered	7.2.20 Removable protective means
	by (e). This indication may be included in the batch or serial number;	7.3 Marking on the inside of equipment and parts
	(m) where applicable, method of sterilization;	I) 7.2.2 Identification
	(iii) where applicable, method of stermzation,	m) 7.2.17 Protective packaging
	(n) in the case of a device within the meaning of Article 1(4a), an indication that the device contains a human blood derivative.	n) Not covered.
13.4	If the intended purpose of the device is not obvious to the user, the manufacturer must clearly state it on the label and in the instructions for use.	Not covered.
13.5	Wherever reasonable and practicable, the devices and detachable components must be identified, where appropriate in terms of batches, to allow all appropriate action to detect any potential risk posed by the devices and detachable components.	Covered. 7.2.2 Identification
13.6	Where appropriate, the instructions for use must contain the following particulars:	a) Details referred to in Section 13.3, with the exception of (d) and (e):
	(a) the details referred to in Section 13.3, with the exception of (d) and (e);	13.3 a) Instructions for Use: authorized representative: not covered Instructions for Use: 7.9.2 Instructions for use
	(b) the performances referred to in Section 3 and any undesirable side-effects; I if the device must be installed with or connected to other medical devices or equipment in order to operate as required for	13.3 b) Instructions for Use: 7.9.1 General on accompanying documents (for electronic Instructions for Use adhere to EU legislation 2007/12)
	its intended purpose, sufficient details of its	13.3 c) Instructions for Use:
	characteristics to identify the correct devices or equipment to use in order to obtain a safe	7.9.2.18 Equipment and accessories supplied sterile (partly covered, the word "sterile" is not required by the standard)
	(d) all the information needed to verify whether	13.3 d) Exempted for Instructions for Use.
	the device is properly installed and can operate correctly and safely, plus details of the nature	13.3 e) Exempted for Instructions for Use
	and frequency of the maintenance and	13.3 f) Instructions for Use: not covered
	calibration needed to ensure that the devices operate properly and safely at all times;	13.3 g) Instructions for Use: not covered
	(e) where appropriate, information to avoid	13.3 h) Instructions for Use: not covered
	certain risks in connection with implantation of the device;	13.3 i) Instructions for Use:
	(f) information regarding the risks of reciprocal	Covered in respect of the following:
	interference posed by the presence of the device during specific investigations or	7.9.2.2 Warning and safety notices





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

treatment; (g) the necessary instructions in the event of damage to the sterile packaging and, where appropriate, details of appropriate methods of re-sterilization; 7.9.2.18 Equipment and accessories supplied sterile 7.9.3.1 General on Technical descriptions and other handling devices. Remark: handling is assumed to inclinate the following: 13.3 k) Instructions for Use: Covered in respect of the following: 7.9.2.2 Warning and safety notices, sentence 13.3 l) Instructions for Use: not cover 13.3 m) Instructions for Use: not cover 13.3 m) Instructions for Use: not cover 13.3 m) Instructions for Use: not cover 13.4 m) Instructions for Use: not cover 13.5 m) Instructions for Use: not cover 13.6 m) Instructions for Use: not cover 13.7 m) Instructions for Use: not cover 13.6 m) Instructions for Us	ption
(g) the necessary instructions in the event of damage to the sterile packaging and, where appropriate, details of appropriate methods of re-sterilization; 1. (a) 3.1 General on Technical description (a) 4.4.a Grips and other handling devices (a) 4.4.a Grips and other handling devic	ption
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connected to other medical devices Covered in respect of the following:	
	า or
7.9.1, General on accompanying documents	
7.9.2.1 General on instructions for us	se
7.9.2.14 Accessories, supplementary equipment, used material	y
7.9.3, Technical description	
14 Programmable electrical medical systems (PEMS)	
d) Covered in respect of the followin	g:
7.9.2.9 Operating instructions	
7.9.2.13 Maintenance	
e) Not covered	
f) Not covered	
g) Covered in respect of the followin	g:
7.2.17 Protective packaging	
7.9.2.18 ME equipment and accessor supplied sterile	ories
(h) if the device is reusable, information on the h) Covered in respect of	
appropriate processes to allow reuse, including cleaning, disinfection, packaging and, where appropriate, the method of	





EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

sterilization of the device to be re-sterilized, and any restriction on the number of reuses.

Where devices are supplied with the intention that they be sterilized before use, the instructions for cleaning and sterilization must be such that, if correctly followed, the device will still comply with the requirements in Section I:

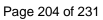
If the device bears an indication that the device is for single use, information on known characteristics and technical factors known to the manufacturer that could pose a risk if the device were to be re-used. If in accordance with Section 13.1 no instructions for use are needed, the information must be made available to the user upon request;

- (i) details of any further treatment or handling needed before the device can be used (for example sterilization, final assembly, etc.);
- (j) in the case of devices emitting radiation for medical purposes, details of the nature, type, intensity and distribution of this radiation.

The instructions for use must also include details allowing the medical staff to brief the patient on any contraindications and any precautions to be taken. These details should cover in particular:

- (k) precautions to be taken in the event of changes in the performance of the device;
- (I) precautions to be taken as regards exposure, in reasonably foreseeable environmental conditions, to magnetic fields, external electrical influences, electrostatic discharge, pressure or variations in pressure, acceleration, thermal ignition sources, etc.;
- (m) adequate information regarding the medicinal product or products which the device in question is designed to administer, including any limitations in the choice of substances to be delivered;
- (n) precautions to be taken against any special, unusual risks related to the disposal of the device:
- (o) medicinal substances, or human blood derivatives incorporated into the device as an integral part in accordance with Section 7.4;
- (p) degree of accuracy claimed for devices with a measuring function;
- (q) date of issue or the latest revision of the

- i) Covered in respect of
- 7.9 Accompanying documents
- i) Covered in respect of
- 7.9.2.17 ME equipment emitting radiation
- k) Not covered
- I) Not covered
- m) Not covered
- n) Not covered
- o) Not covered
- p) Not covered
- a) Not covered

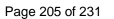




EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

instructions for use.

Annex	Annex ZZB	 N/A
ZZB	(informative)	
	Relationship between this European Standard and the Essential Requirements of EU Directive 90/385/EEC on Active Implantable Medical Devices	
	This European Standard has been prepared under a mandate given to CENELEC by the European Commission and the European Free Trade Association to provide a means of conforming to the Essential Requirements given in Annex I of the EC Directives 90/385/EEC as amended by 2007/47/EC.	 N/A
	General Guidance:	 N/A
	Once this standard will be cited in the Official Journal of the European Union under that Directive, compliance with the clauses of this standard given in Table ZZB.1 confers, within the limits of the scope of this standard, a presumption of conformity with the corresponding Essential Requirements (Ers) of that Directive and associated EFTA regulations.	 N/A
	NOTE 1 This standard is intended to be applied in its entirety only. Selected clauses or subclauses may be not applicable due to the specific type of equipment under consideration. It is necessary to understand and apply Clauses 1 to 16. It is also recommended to understand and apply those clauses which contain general requirements related to a specific subclause. Elements of the standard that are not cited in Table ZZB.1 may be relevant for the appropriate fulfilment of certain essential requirements through indirect reference, and for safety and performance aspects of the device, that are not addressed through essential requirements	N/A
	NOTE 2 Where a reference from a clause of this standard to the risk management process is made, the risk management process needs to be in compliance to the AIMD (Directive 90/385/EEC amended by 2007/47/EC). This means that risks have to be reduced "as far as possible", "to a minimum", "to the lowest possible level", "minimized" or "removed", according to the wording of the corresponding essential requirement.	 N/A





	EU Differences (EN 60601-1:2006 + A	1:2013 + A12:2014)	
Clause	Requirement + Test	Result - Remark	Verdict

NOTE 3 With respect to Note 4 of clause 4.2.2 General requirement for risk management, the manufacturer's policy for determining acceptable risk must be in compliance with essential requirements 1, 2, 5, 6, 7, 8, 9, 11 and 12 of the directive.	 N/A
NOTE 4 References in the Clauses 3 to 17 or in the Annexes of this standard specify whether the normative references listed in Clause 2 as cited in Annex ZA are to be applied in whole or in part.	 N/A
NOTE 5 This Annex ZZB is based on Normative References according to Annex ZA, replacing the references in the core text.	 N/A
WARNING: Other requirements and other EU Directives and Regulations may be applicable to the product(s) falling within the scope of this standard.	 N/A

Table ZZB.1	Relationship between Essential Requirements o 2007/47/EC, and Clauses and Subclauses of this	
No.	Essential Requirement	Coverage
l	GENERAL REQUIREMENTS	
1.	General Guidance notes 2 and 3 shall be observ	ed
1	The devices must be designed and manufactured in such a way that, when implanted under the conditions and for the purposes laid down, their use does not compromise the clinical condition or the safety of patients. They must not present any risk to the persons implanting them or, where applicable, to other persons.	Not covered This ER relates to the implanted part of the active implantable medical device.
2	The devices must achieve the performances intended by the manufacturer, viz. Be designed and manufactured in such a way that they are suitable for one or more of the functions referred to in Article 1 (2) (a) as specified by him.	Not covered.
3	The characteristics and performances referred to in sections 1 and 2 must not be adversely affected to such a degree that the clinical condition and safety of the patients or, as appropriate, of other persons are compromised during the lifetime of the device anticipated by the manufacturer, where the device is subjected to stresses which may occur during normal conditions of use.	Not covered. However, the standard provides a procedure for the generation of information that is necessary to document that the device is in compliance with this ER with regard to the external parts of an active implantable medical device.
4	General Guidance notes 2 and 3 shall be observ	ed



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	EU Differences (EN 60601-1:2006 + A	1:2013 + A12:2014)	
Clause	Requirement + Test	Result - Remark	Verdict
4	The devices must be designed, manufactured and packed in such a way that their characteristics and performances are not	Covered for the external part of implantable medical device only of the following:	
	adversely affected in the storage and transport conditions laid down by the manufacturer	7.2.17 Marking on protective pa	ackaging
	(temperature, humidity, etc.).	7.9.3.1 Technical description	
		15.3.7 Environmental influence	S
5	General Guidance notes 2 and 3 shall be observed	ed	
5	Any side effects or undesirable conditions must constitute acceptable risks when weighed against the performances intended.	Not covered.	
5a	Demonstration of conformity with the essential requirements must include a clinical evaluation in accordance with Annex 7.	Not covered.	
II	REQUIREMENTS REGARDING DESIGN AND CO	NSTRUCTION	
6	The solutions adopted by the manufacturer for the design and construction of the devices must comply with safety principles taking	Covered for the external part of implantable medical device only of the following:	
		8 Protection against electrical h ME equipment	azards from
		9 Protection against mechanica ME equipment and ME systems	
		15 Construction of ME equipme	ent
7	Implantable devices must be designed, manufactured and packed in a non-reusable pack according to appropriate procedures to ensure they are sterile when placed on the market and, in the storage and transport conditions stipulated by the manufacturer, remain so until the packaging is removed and	Not covered.	

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they are implanted.

as possible:

General Guidance notes 2 and 3 shall be observed

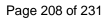
Devices must be designed and manufactured in such a way as to remove or minimize as far



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EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

- the risk of physical injury in connection with their physical, including dimensional, features,	Covered for the external part of an active implantable medical device only in respec of the following:
	8.1 Electric shock
	9.1 Mechanical Hazards
	10 Radiation (all types)
	11.1 Excessive temperatures
	11.2 Fire prevention
	11.4 Flammable anaesthetics
	11.5 Flammable agent
	11.6.3 Spillage
	11.8 Interruption of power supply
	12.4 Hazardous output
	13.1 Hazardous situations
	13.2 Single Fault condition
	15.3 Mechanical strength
	15.4 Components and general assembly
	15.5.3 Construction of transformers
	16.3 Power supply
	16.5 Separation devices
	16.6 Leakage currents
	16.8 Interruption of power supply
- risks connected with the use of energy sources with particular reference, where electricity is used, to insulation, leakage	Covered for the external part of an active implantable medical device only in respect of the following:
currents and overheating of the devices,	8.1 Electric shock
	13.2 Single Fault condition
	15.5.3 Construction of transformers
	16.3 Power supply
	16.5 Separation devices
	16.6 Leakage currents
	16.8 Interruption of power supply





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	EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict	
	environmental conditions such as magnetic fields, external electrical influences	Not covered.		
		See for EMC EN 60601-1-2 as in Annex ZA	referenced	
	in pressure and acceleration,	See for acceleration EN 60601 EN 60601-1-12 as referenced i		
	p	Covered in respect of the follow pressure, temperature: test in 5 to manufacturers' specification	.3 according	
	- risks connected with medical treatment, in particular those resulting from the use of defibrillators or high-frequency surgical	Covered for the external part of implantable medical device only of the following:		
	equipment,	For defibrillator protection		
		8.5.5 Defibrillation-proof applied	d parts	
	- risks connected with ionising radiation from radioactive substances included in the device, in compliance with the protection requirements laid down in Council Directive 96/29/Euratom of 13 May 1996 laying down basic safety standards for the protection of the health of workers and the general public against the dangers arising from ionising radiation (1) and Council Directive 97/43/Euratom of 30 June 1997 on health protection of individuals against the dangers of ionising radiation in relation to medical exposure (1),	Not covered.		
	- risks which may arise where maintenance and calibration are impossible, including:	Not covered.		
	- excessive increase of leakage currents,			
	- ageing of the materials used,			
	- excess heat generated by the device,			
	 decreased accuracy of any measuring or control mechanism. 			
9	The devices must be designed and manufactured in such a way as to guarantee the characteristics and performances referred to in I. 'General requirements', with particular attention being paid to:			
	- the choice of materials used, particularly as	Not covered.		
	regards toxicity aspects,	The manufacturer should apply appropriate part of EN ISO 109		
	- mutual compatibility between the materials used and biological tissues, cells and body fluids, account being taken of the anticipated use of the device,	Not covered.		
		The manufacturer should apply appropriate part of EN ISO 109		
	- compatibility of the devices with the substances they are intended to administer,	Not covered.		

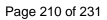


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EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark V	/erdict
	- the quality of the connections, particularly in respect of safety,	Covered for the external part of an a implantable medical device only in r	
		of the following: Covered in respect of the following:	
		15.4.1 Construction of connectors	
	- the reliability of the source of energy,	Not covered.	
	- if appropriate, that they are leakproof,	Covered for the external part of an a implantable medical device only in r of the following:	
		9.7 Pressure vessels and parts subj pneumatic and hydraulic pressure,	ect to
		11.6.1 Protection against overflow, s leakage, ingress of water or particul matter, cleaning, disinfection and sterilization, compatibility with subst	ate
		11.6.2 Overflow	
		15.4.9 Oil containers	
	- proper functioning of the programming and control systems, including software. For devices which incorporate software or which are medical software in themselves, the software must be validated according to the state of the art taking into account the principles of development lifecycle, risk management, validation and verification	Covered for the external part of an a implantable medical device which incorporates software by 14 Prograr electrical medical systems (PEMS)	
10	- Where a device incorporates, as an integral part, a substance which, if used separately, may be considered to be a medicinal product as defined in Article 1 of Directive 2001/83/EC, and which is liable to act upon the body with action ancillary to that of the device, the quality, safety and usefulness of the substance must be verified by analogy with the methods specified in Annex I to Directive 2001/83/EC. For the substances referred to in the first paragraph, the notified body shall, having verified the usefulness of the substance as part of the medical device and taking account of the intended purpose of the device, seek a scientific opinion from one of the competent authorities designated by the Member States or the European Medicines Agency (EMEA) acting particularly through its committee in accordance with Regulation (EC) No 726/2004	Not covered.	

(2) on the quality and safety of the substance including the clinical benefit/risk profile of the incorporation of the substance into the device. When issuing its opinion, the competent authority or the EMEA shall take into account the manufacturing process and the data related

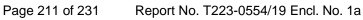




EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict

	to the usefulness of incorporation of the substance into the device as determined by the notified body. Where a device incorporates, as an integral part, a human blood derivative, the notified body shall, having verified the usefulness of the substance as part of the device and taking account of the intended purpose of the device, seek a scientific opinion from the EMEA, acting particularly through its committee, on the quality and safety of the substance, including the clinical benefit/risk profile of the incorporation of the human blood derivative into the device. When issuing its opinion, the EMEA shall take into account the manufacturing process and the data related to the usefulness of incorporation of the substance into the device as determined by the notified body. Where changes are made to an ancillary substance incorporated in a device, in particular related to its manufacturing process, the notified body shall be informed of the changes and shall consult the relevant medicines competent authority (i.e. the one involved in the initial consultation), in order to confirm that the quality and safety of the ancillary substance are maintained. The competent authority shall take into account the data related to the usefulness of the incorporation of the substance into the device as determined by the notified body, in order to ensure that the changes have no negative impact on the established benefit/risk profile of the addition of the substance in the device. When the relevant medicines competent authority (i.e. the one involved in the initial consultation) has obtained information on the ancillary substance, which could have an impact on the established benefit/risk profile of the addition of the substance to the device, it shall provide the notified body with advice, whether this information has an impact on the established benefit/risk profile of the addition of the substance to the device or not. The notified body shall take the updated scientific opinion into account in reconsidering its assessment of t	
11	The devices and, if appropriate, their component parts must be identified to allow any necessary measure to be taken following the discovery of a potential risk in connection with the devices and their component parts.	Not covered.





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	EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)			
Clause	Requirement + Test	Result - Remark	Verdict	

12	Devices must bear a code by which they and their manufacturer can be unequivocally identified (particularly with regard to the type of device and year of manufacture); it must be possible to read this code, if necessary, without the need for a surgical operation.	Not covered.
13	When a device or its accessories bear instructions required for the operation of the device or indicate operating or adjustment	Covered for the external part of an active implantable medical device only in respect of the following:
	parameters, by means of a visual system, such information must be understandable to the	7.5 Safety signs
	user and, as appropriate, the patient.	7.9.1 General requirements for accompanying documents
14	On the sterile pack:	Not covered.
	- the method of sterilization,	Not covered.
	- an indication permitting this packaging to be recognized as such,	Not covered.
	- the name and address of the manufacturer,	Not covered.
	- a description of the device,	Not covered.
	- if the device is intended for clinical investigations, the words: 'exclusively for clinical investigations	Not covered.
	- if the device is custom-made, the words 'custom-made device',	Not covered.
	- a declaration that the implantable device is in a sterile condition,	Not covered.
	- the month and year of manufacture,	Not covered.
	- an indication of the time limit for implanting a device safely.	Not covered.
14.2	On the sales packaging:	Not covered.
	- the name and address of the manufacturer and the name and address of the authorised representative, where the manufacturer does not have a registered place of business in the Community,	Not covered.
	- a description of the device,	Not covered.
	- the purpose of the device,	Not covered.
	- the relevant characteristics for its use,	Not covered.
	- if the device is intended for clinical investigations, the words: 'exclusively for clinical investigations',	Not covered.
	- if the device is custom-made, the words: 'custom-made device',	Not covered.





	EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)		
Clause	Requirement + Test	Result - Remark	Verdict
	- a declaration that the implantable device is in a sterile condition,	Not covered.	
	- the month and year of manufacture,	Not covered.	
	- an indication of the time limit for implanting a device safely,	Not covered.	
	- the conditions for transporting and staring the device,	Not covered.	
	- in the case of a device within the meaning of Article 1(4a), an indication that the device contains a human blood derivative.	Not covered.	
15	When placed on the market, each device must be accompanied by instructions for use giving the following particulars:	Not covered.	
	- the year of authorization to affix the CE mark,	Not covered.	
	- the details referred to in 14.1 and 14.2, with the exception of those referred to in the eighth and ninth indents,	Not covered.	
	- the performances referred to in section 2 and any undesirable side effects,	Not covered.	
	- information allowing the physician to select a suitable device and the corresponding software and accessories,	Not covered.	
	- information constituting the instructions for use allowing the physician and, where appropriate, the patient to use the device, its accessories and software correctly, as well as information on the nature, scope and times for operating controls and trials and, where appropriate, maintenance measures,	Not covered.	
	 information allowing, if appropriate, certain risks in connection with implantation of the device to be avoided, 	Not covered.	
	- information regarding the risks of reciprocal interference (1) in connection with the presence of the device during specific investigations or treatment,	Not covered.	
	- the necessary instructions in the event of the sterile pack being damaged and, where appropriate, details of appropriate methods of resterilization,	Not covered.	
	- an indication, if appropriate, that a device can be reused only if it is reconditioned under the responsibility of the manufacturer to comply with the essential requirements.	Not covered.	



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EU Differences (EN 60601-1:2006 + A1:2013 + A12:2014)				
Clause	Clause Requirement + Test Result - Remark Verdic			

	The instruction leaflet must also include details allowing the physician to brief the patient on the contra-indications and the precautions to be taken. These details should cover in particular:	Not covered.
	 information allowing the lifetime of the energy source to be established, 	Not covered.
	 precautions to be taken should changes occur in the device's performance, 	Not covered.
	- precautions to be taken as regards exposure, in reasonably foreseeable environmental conditions, to magnetic fields, external electrical influences, electrostatic discharge, pressure or variations in pressure, acceleration, etc.,	Not covered.
	 adequate information regarding the medicinal products which the device in question is designed to administer, 	Not covered.
	- date of issue or the latest revision of the instructions for use.	Not covered.
16	Confirmation that the device satisfies the requirements in respect of characteristics and performances, as referred to in I. 'General requirements', in normal conditions of use, and the evaluation of the side effects or undesirable effects must be based on clinical data established in accordance with Annex 7.	Not covered.



Enclosure No. 2

Photo documentation

(6 pages including this cover page)



Unit from outside



















Unit from inside

