

Test Report issued under the responsibility of:



TEST REPORT IEC 60601-1-11

Part 1-11: General requirements for basic safety and essential performance – Collateral Standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment

Report Number:	230500749SHA-003
Date of issue:	2023-06-07
Total number of pages:	32
Name of Testing Laboratory	Intertek Testing Services Shanghai
preparing the Report:	Building No. 86, 1198 Qinzhou Road (North) Shanghai 200233 China
Applicant's name:	GlobTek, Inc.
Address:	186 Veterans Drive Northvale NJ 07647, USA
Test specification:	
Standard:	IEC 60601-1-11:2015, IEC 60601-1-11:2015/AMD1:2020 for use in conjunction with IEC 60601-1:2005, IEC 60601- 1:2005/AMD1:2012, IEC 60601-1:2005/AMD2:2020
Test procedure:	CB Scheme
Non-standard test method:	N/A
TRF template used:	IECEE OD-2020-F1:2021, Ed.1.4
Test Report Form No	IEC60601_1_11G
Test Report Form(s) Originator :	UL(US)
Master TRF:	2021-09-16

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General disclaimer:

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Test item description	Medica	al Power Supply	
Trade Mark(s):	GlobTek, [°] Inc.		
Model/Type reference	186 Ve GT*46	ek, Inc. eterans Drive Northvale N 101-*05*-USB, GT*4610 to page 6 for details.)	
-	•	100-240V~, 50-60Hz, 0.3	А;
	Output		
		101-*05*-USB: 5Vdc,2A 101-*06*-USB: 5.1-5.5Vc	
Responsible Testing Laboratory (as ap	plicat	ole), testing procedure	and testing location(s):
CB Testing Laboratory:		Intertek Testing Services	s Shanghai
Testing location/ address	:	Building No. 86, 1198 Q 200233 China	linzhou Road (North) Shanghai
Tested by (name, function, signature).	:	Vivian Xu(Engineer)	Vi Vian - Xu.
Approved by (name, function, signatur	re):	Larry Zhong (Mandated reviewer)	Vi Vian . Xu. Lany Zhang
Testing procedure: CTF Stage 1:		N/A	
Testing location/ address	:		
Tested by (name, function, signature).	:		
Approved by (name, function, signatur	re):		
Testing procedure: CTF Stage 2:		N/A	
Testing location/ address	:		
Tested by (name + signature)	:		
Witnessed by (name, function, signatu	,		
Approved by (name, function, signatur	re):		
Testing procedure: CTF Stage 3:		N/A	
Testing procedure: CTF Stage 4:		N/A	
Testing location/ address	:		
Tested by (name, function, signature).	:		
Witnessed by (name, function, signatu	-		
Approved by (name, function, signatur			
Supervised by (name, function, signate	ure) :		

List of Attachments (including a total number of pages in each attachment):				
None				
Summary of testing:				
Tests performed (name of test and test	Testing location:			
clause):	Intertek Testing Services Shanghai			
4.2.2 Environmental condition test of transport and storage between uses	Building No. 86, 1198 Qinzhou Road (North) Shanghai 200233 China			
4.2.3.1 Environmental operating condition test				
10.1.2 a) Shock test				
10.1.2 b) Vibration test				
The sample tested complies with the requirements of IEC 60601-1-11:2020.				
Summary of compliance with National Difference	es (List of countries addressed):			
None.				
The product fulfils the requirements of IEC 60	0601-1-11:2015/AMD1:2020			
Use of uncertainty of measurement for decisions	s on conformity (decision rule) :			
⊠ No decision rule is specified by the IEC standard, when comparing the measurement result with the applicable limit according to the specification in that standard. The decisions on conformity are made without applying the measurement uncertainty ("simple acceptance" decision rule, previously known as "accuracy method").				
GMS-QC-12 Estimation of Measurement Uncerta	inty, 19-April-2018 Initial Release.			
Other: (to be specified, for example when required by the standard or client, or if national accreditation requirements apply)				
Information on uncertainty of measurement:				
The uncertainties of measurement are calculated by	y the laboratory based on application of criteria given of test methods, decision sheets and operational			
procedures of IECEE.				
the decision rule when reporting test results with	n of measurement uncertainty principles and applying in IECEE scheme, noting that the reporting of the t necessary unless required by the test standard or			
customer.	, , , , , , , , , , , , , , , , , , , ,			
Calculations leading to the reported values are on fi the testing.	le with the NCB and testing laboratory that conducted			

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. GlobTek, Inc. 186 Veterans Dr. Northvale, NJ 07647 USA ICT/ITE/MEDICAL POWER SUPPLY 电源供应器/電源供應器 圈 P/N/Número de pieza/номер/料号/料號:WR9QA2000USBNMNAR6B MODEL/Modelo/модель/型号/型號:GTM46101-1005-USB INPUT/Entrada/вход/输入/輸入:100-240V~,50-60Hz, 0.3A OUTPUT/Salida/выход/输出/輸出:5.0 V === 2.0A,10.0W Pin 1: (+) Pin 2&3: N/C Pin 4: Com T1.0A 250VAC CAN ICES-3 (B)/NMB-3(B) EFFICIENCY LEVEL RoHS LPS WWYY MADE IN CHINA (中国制造/中國製造)

Note:

The above markings are the minimum requirements required by the safety standard. For the final production samples, the additional markings which do not give rise to misunderstanding may be added. Other models are with similar label as corresponding above models except different model name and output ratings.

When the equipment is vended to EU, the name and address of the importer or authorized representative within the EEA shall be added on the equipment;

Test item particulars			
Classification of installation and use	Power adapter model.		
	Final evaluation in end product.		
Supply Connection	Power adapter model		
Possible test case verdicts:			
- test case does not apply to the test object::	N/A		
- test object does meet the requirement::	P (Pass)		
- test object does not meet the requirement::	F (Fail)		
Testing:			
Date of receipt of test item:	2023-05-13		
Date (s) of performance of tests:	2023-05-13 to 2023-06-01		
General remarks:			
"(See Enclosure #)" refers to additional information ap "(See appended table)" refers to a table appended to the			
Throughout this report a \square comma / $oxtimes$ point is u	sed as the decimal separator.		
☑ This Test Report Form contains requirements a 2015-12-25.	according to IEC 60601-1-11:2015 Standard dated		
Disclaimer:			
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Manufacturer's Declaration per sub-clause 4.2.5 of	IECEE 02:		
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 t	he General product information section.		

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Name and address of factory (ies): Factory 1

GlobTek, Inc.	
186 Veterans Drive Northvale NJ 07647, USA	
Factory 2	
GlobTek (Suzhou) Co., Ltd	
Building 4, No. 76, Jin Ling East Rd., Suzhou Industrial Park, Suzhou, JiangSu 215021, China	
Factory 3	
Shenzhen ENG Electronics Co., Ltd.	
Block B, Nuclear Group Industrial District, Baishix Fuyun Town, Bao'an, Shenzhen, China	

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General product information and other remarks:

Product covered by this report is medical power supply module, which can be used as a part of medical equipment. All the models have the same structure except with or without led.

Transformers used in all models are the same. All models have same PCB, but some non-critical components may be adjusted according to different output voltage. The parameters of these components depend on output voltage.

All the types are designed for continuous operation and no applied part is defined.

The insulation construction of EUT is evaluated as 2MOPP in this report as customer's request. Model Similarity:

GT*46101-*05*-USB, GT*46101-*06*-USB

The 1st "*" can be "M" or "-" or "H" for market identification and not related to safety.

The 2nd "*" can be "01" to "13", with interval of 1, denote the rated output wattage designation.

The 3rd "*" can be "-0.5" to "-0.9" with interval of 0.1, or blank indicate no voltage different, optional deviation, subtracted from standard output voltage.

The "05" or "06" and 3rd "*" together denote the output voltage, with a range of 5-5.5 volts.

Model list

Model	Rated output voltage range	Max. rated output current	Max. rated output power
GT*46101-*05*-USB	5Vdc	2A	10W
GT*46101-*06*-USB	5.1-5.5Vdc	2.54A	13W

Technical Considerations:

Models GTM46101-1005-USB, GTM46101-1306-0.9-USB and GTM46101-1306-0.5-USB are tested as typical models.

The products are not intended to use in environment which altitude exceed 5000m.

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Clause	Requirement + Test

Result - Remark

4	GENERAL REQUIREMENTS		Р	
4.1	Characteristics of SUPPLY MAINS specified in 4.10.2 of Part 1 applied, except ME EQUIPMENT or ME SYSTEMS intended for HOME HEALTHCARE ENVIRONMENT complied with the following:		Р	
	- SUPPLY MAINS in the HOME HEALTHCARE ENVIRONMENT did not exceed 110 % or was not below 85 % of NOMINAL voltage between any of the conductors of the system or between any of these conductors and earth (% V)	See the IEC 60601-1 report 2302500749SHA -001	_	
	 For ME EQUIPMENT OR ME SYSTEMS intended to actively keep alive or resuscitate a PATIENT, SUPPLY MAINS in the HOME HEALTHCARE ENVIRONMENT did not exceed 110 % or was not below 80 % of NOMINAL voltage between any of the conductors of the system or between any of these conductors and earth (% V	Not for life- supporting me equipment.	_	
	- RATED range of NOMINAL voltage did include at least 12.4 V to 15.1 V for operation from a 12 V dc supply mains	No such condition	N/A	
	- RATED range of NOMINAL voltage did include at least 24.8 V to 30.3 V for operation from a 12 V dc supply mains	No such condition	N/A	
	The equipment maintained BASIC SAFETY and ESSENTIAL PERFORMANCE during and following a 30 s dip to 10 V from a 12 V dc SUPPLY MAINS	No such condition	N/A	
	The equipment maintained BASIC SAFETY and ESSENTIAL PERFORMANCE during and following a 30 s dip to 20 V from a 24 V dc SUPPLY MAINS	No such condition	N/A	
4.2.2	Environmental conditions of transport and storage between uses, indicated in instructions for use			
	ME EQUIPMENT, except STATIONARY EQUIPMENT, after being removed from its protective packaging, and subsequently between uses, operated within its specified NORMAL USE after transport or storage in the specified environmental conditions		Ρ	
	temperature range:-25 °C to + 5 °C		Р	
	temperature range:+5 °C to +35 °C at a non- condensing relative humidity up to 90 %		Р	
	temperature range: >35 °C to 70 °C at a water vapour pressure up to 50 hPa		Р	
	For more restricted range of environmental transport and storage conditions between uses, the environmental conditions are specified	See RM table 4.2.2	Р	
	– Justified in the RISK MANAGEMENT FILE		Р	

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Clause	Requirement + Test	Result - Remark	Verdict
	- Marked on the ME EQUIPMENT		N/A
	When not practicable, the more restricted range is disclosed in the instructions for use		N/A
	 Marked on the carrying case when the instructions for use indicate the ME EQUIPMENT is intended to be transported or stored in a carrying case between uses 		N/A
	Symbol 5.3.5 (ISO 7000-0534), 5.3.6 (ISO 7000- 0533), or 5.3.7 (ISO 7000-0632) of ISO 15223- 1:2016 used to mark temperature range		N/A
	Symbol 5.3.8 (ISO 7000-2620) of ISO 15223-1:2016 used to mark humidity range		N/A
	Symbol 5.3.9 (ISO 7000-2621) of ISO 15223-1:2016 used to mark atmospheric pressure range		N/A
	Where ME EQUIPMENT used different marking for conditions of transport and storage between uses, continuous operating conditions and transient operating conditions, markings accompanied by supplementary markings except where the respective applicability was obvious		N/A
	Environmental transport and storage test		Р
	a) ME EQUIPMENT prepared for transportation or storage according to instructions for use		Р
	b) ME EQUIPMENT exposed to its lowest specified environmental transport and storage conditions (temperature -4 °C) (°C):	-40 °C	Р
	 For at least 16 h or, ensure ME EQUIPMENT reached THERMAL STABILITY for at least 2 h 	16h	Р
	c) Then ME EQUIPMENT exposed to 34 °C \pm 4 °C and 90 % - 0% + 6% relative humidity until the test chamber reached equilibrium and held for at least 2 hours. The transition from low to high temperature was made slowly enough to provide a non- condensing environment.	34°C and 90% relative humidity	Р
	d) ME EQUIPMENT exposed to its highest specified environmental transport and storage conditions, not requiring a water vapour pressure greater than 50	80 °C, 50hpa	Р
	hPa (temperature ⁺⁴ ⁰ °C); (°C, ± %):		
	 For at least 16 h or, ensured ME EQUIPMENT reached THERMAL STABILITY for at least 2 h 		Р
	e) At the end of this conditioning period, ME EQUIPMENT allowed to return and stabilize at the operating conditions of NORMAL USE		Р

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Clause	Requirement + Test	Result - Remark	Verdict	
	f) ME EQUIPMENT evaluated to its specifications and ensured it provides BASIC SAFETY and ESSENTIAL PERFORMANCE		Р	
4.2.3.1	Environmental operating conditions - Continuous	operating conditions	Р	
	Instructions for use indicated permissible environmental operating conditions of the ME EQUIPMENT		Р	
	ME EQUIPMENT complied with its specifications and all requirements of the standard when operated in NORMAL USE within temperature + 5 °C to +40 °C,		N/A	
	Relative humidity range of 15 % to 90%, non- condensing, but not requiring a water vapour partial pressure greater than 50 hPa; and		N/A	
	An atmospheric pressure range of 700 hPa to 1060 hPa		N/A	
	For more restricted range of environmental operating conditions	0 °C to +40 °C	Р	
	- justified in the risk management file;	See RISK MANAGEMENT Table 4.2.3.1	Р	
	-marked on the equipment; or were nor practical in the instructions for use	be disclosed in the instructions for use	Р	
	 Marked on the carrying case when the instructions for use indicate the ME EQUIPMENT is intended to be operated in a carrying case 	No carrying case	N/A	
	Symbol 5.3.5 (ISO 7000-0534), 5.3.6 (ISO 7000- 0533), or 5.3.7 (ISO 7000-0632) of ISO 15223- 1:2016 used to mark temperature range	No such symbol	N/A	
	Symbol 5.3.8 (ISO 7000-2620) of ISO 15223-1:2016 used to mark humidity range	No such symbol	N/A	
	Symbol 5.3.9 (ISO 7000-2621) of ISO 15223-1:2016 used to mark atmospheric pressure range	No such symbol	N/A	
	Where ME EQUIPMENT used different marking for conditions of continuous operating conditions and transient operating conditions, markings accompanied by supplementary markings		N/A	
	Environmental operating conditions test		Р	
	a) ME EQUIPMENT was set up for operation according to INTENDED USE		Р	
	b) ME EQUIPMENT exposed to 20 °C \pm 4 °C for at least 6 h or, ensured ME EQUIPMENT reached THERMAL STABILITY for at least 2 h, (h)	20°C, 6h	Р	
	c) ME EQUIPMENT evaluated to its specifications and ensured it continued to provide BASIC SAFETY and ESSENTIAL PERFORMANCE		Р	

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Clause	Requirement + Test	Result - Remark	Verdict			
	d) ME EQUIPMENT evaluated to its specifications and ensured it continued to provide BASIC SAFETY and ESSENTIAL PERFORMANCE while at the lowest specified atmospheric pressure.		P			
	e) ME EQUIPMENT evaluated to its specifications and ensured it continued to provide BASIC SAFETY and ESSENTIAL PERFORMANCE while at the highest specified atmospheric pressure.		P			
	f) Pressure in chamber relieved		Р			
	g) ME EQUIPMENT cooled to its lowest specified environmental operating conditions		Р			
	h) ME EQUIPMENT held at lowest specified environmental operating conditions for at least 6 h or, ensured the ME EQUIPMENT reached THERMAL STABILITY for at least 2 h	6h	P			
	i) ME EQUIPMENT met its specifications and BASIC SAFETY and ESSENTIAL PERFORMANCE		Р			
	j) ME EQUIPMENT warmed to its highest specified continuous environmental operating conditions		Р			
	k) ME EQUIPMENT held the conditions of j) for at least 6 h or, ensured the ME EQUIPMENT reached THERMAL STABILITY for at least 2 h	6h	Р			
	I) ME EQUIPMENT met its specifications and BASIC SAFETY and ESSENTIAL PERFORMANCE		Р			
4.2.3.2	Environmental shock to TRANSIT-OPERABLE EQUIPM	ENT	N/A			
	TRANSIT-OPERABLE EQUIPMENT with a stated wider range of continuous environmental operation conditions maintained BASIC SAFETY and ESSENTIAL PERFORMANCE in the presence of condensation and thermal shock from rapid changes in environmental temperature and humidity during INTENDED USE when test in accordance with 4.2.3.2 a)-j).	Not Transit-operable equipment	N/A			

5	GENERAL REQUIREMENTS FOR TESTING ME EQUIPMENT In addition to the requirements of 5.9.2.1 of with IEC 60601-1 standard, accessibility determined as indicated below:		Р
			Р
	ACCESSIBLE parts of ME EQUIPMENT identified by inspection and, when necessary, by testing		Р
	When in doubt, an ACCESSIBLE PART of ME EQUIPMENT determined by a test with the small finger probe of Fig 1, applied in a bent or straight position as follows:		Р
	 – for all positions of the ME EQUIPMENT operating in NORMAL USE 	No opening	N/A

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Clause	Requirement + Test	Result - Remark	Verdict		
	– after opening ACCESS COVERS and removal of parts, including lamps, fuses, and fuse holders when:		N/A		
	i) the ACCESS COVERS could be opened without the use of a TOOL, or		N/A		
	ii) the instructions for use instructed a LAY OPERATOR to open the relevant ACCESS COVER		N/A		

6	CLASSIFICATION OF ME EQUIPMENT AND ME SYSTEMS		Р
	ME EQUIPMENT intended for HOME HEALTHCARE ENVIRONMENT classified as follows, except for PERMANENTLY INSTALLED EQUIPMENT and as required by Part 1, Sub-clause 6.2:	Class II	Ρ
	- CLASS II OF INTERNALLY POWERED	class II	Р
	- Not provided with a FUNCTIONAL EARTH TERMINAL		Р
	– When equipped with APPLIED PARTS, they are TYPE BF or CF	No applied parts	N/A

7 ME EQUIPMENT IDENTIFICATION, MARKING AND DOCU		D DOCUMENTS	Р
7.1	USABILITY of identification, marking, and ACCOMPANYING DOCUMENTS intended for LAY OPERATOR or LAY RESPONSIBLE ORGANIZATION evaluated by an OPERATOR whose PROFILE included minimum eight years of education	230500749SHA -002 and usability engineering should be considered in end product again.	N/A
	ME EQUIPMENT and ME SYSTEMS intended for HOME HEALTHCARE ENVIRONMENT are simple to use and do not require referencing complex ACCOMPANYING DOCUMENTS		N/A
7.2	In addition to requirements of 7.2.9 of the general standard, the ME EQUIPMENT or its parts and, when appropriate, a carrying case are marked with the appropriate IP classification as tested in 8.3.1 :	IP20	N/A
	If the carrying case provide some or all of the ingress protection against water or particulate matter:	No carrying case	N/A
	a) The ENCLOSURE is marked with the safety sign ISO 7010-W001 and "keep dry" or symbol ISO 15223-1:2012, 5.3.4 (ISO 7000-0626)		N/A
	b) the carrying case marked with its degree of protection:		N/A
	Carrying case inspected, and tests and criteria of 7.1.2 and 7.1.3 of Part 1 applied:	No carrying case	N/A
7.3	ACCOMPANYING DOCUMENTS		Р

	1 age 12 01 02	Report No. 2000007	
	IEC 60601-1-11		
Clause	Requirement + Test	Result - Remark	Verdict
7.3.1	ACCOMPANYING DOCUMENTS indicate the LAY OPERATOR OF LAY RESPONSIBLE ORGANIZATION contact the MANUFACTURER OF MANUFACTURER'S representative on the following issues:	Accompany documents are provided for some critical issue like technical data, safety warnings, necessary information to set up	P
	– Assistance in setting up, using, or maintaining the ME EQUIPMENT or ME SYSTEM when needed, or		Р
	- To report unexpected operation or events		Р
	ACCOMPANYING DOCUMENTS include a postal address and either a phone number or web address for the LAY OPERATOR or LAY RESPONSIBLE ORGANIZATION to contact the MANUFACTURER or MANUFACTURER'S representative		P
7.3.2	ACCOMPANYING DOCUMENTS include necessary details for healthcare professional to brief the LAY OPERATOR or LAY RESPONSIBLE ORGANIZATION on any known contraindication(s) to the use of ME EQUIPMENT or ME SYSTEM and any precautions to be taken, including the following:	Accompany documents are provided for some critical issue like technical data, safety warnings, necessary information to set up	N/A
	 Precautions to be taken in the event of changes in the performance of ME EQUIPMENT or ME SYSTEM 		N/A
	 Precautions to be taken regarding the exposure of the ME EQUIPMENT or ME SYSTEM to reasonably foreseeable environmental conditions 		N/A
	 Adequate information regarding medicinal substances that ME EQUIPMENT is designed to administer, including any limitations in the choice of substances to be delivered as indicated below: 		N/A
	– Information on any medicinal substances or human blood derivatives incorporated into the ME EQUIPMENT or ACCESSORIES as an integral part; and		N/A
	 The degree of accuracy claimed for ME EQUIPMENT with a measuring FUNCTION 		N/A
7.4	Instructions for use		Р
7.4.1	Nature of the HAZARD, likely consequences that could occur if the advice is not followed, and precautions for reducing the RISK described in instructions for use corresponding to each warning and SAFETY SIGN	See RM table 7.4.1	P
	The instructions for use address the following issues,	, as applicable:	N/A
	 Strangulation due to cables and hoses, particularly due to excessive length 		N/A
	- Inhalation or swallowing of small parts		N/A
	 Potential allergic reactions to accessible materials used in the ME EQUIPMENT 		N/A
	– Contact injuries		N/A

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		10001110.200007		
IEC 60601-1-11				
Clause	Requirement + Test	Result - Remark	Verdict	
	The instructions for use include warnings to the effect that the following actions could be unsafe as applicable:		N/A	
	 Use of ACCESSORIES, detachable parts, and materials not described in the instructions for use (see 7.9.2.14 of Part 1) 		N/A	
	 Interconnection of this equipment to other equipment not described in the instructions for use (see 16.2 c) indent 9) of Part 1) 		N/A	
	- Modification of the equipment		N/A	
	- Use of the ME EQUIPMENT outside its carrying case when some part of the protection required by this standard is provided by that carrying case (see 8.3.1 and 10.1)		N/A	
7.4.2	When BASIC SAFETY OR ESSENTIAL PERFORMANCE dependents on the INTERNAL ELECTRICAL POWER SOURCE, the instructions for use describes the following:	No internal electrical power source	N/A	
	- Typical operation time or number of procedures :		N/A	
	– Typical service life of the INTERNAL ELECTRICAL POWER SOURCE; and		N/A	
	– Behaviour of ME EQUIPMENT while the rechargeable INTERNAL ELECTRICAL POWER SOURCE is charging:		N/A	
7.4.3	Instructions for use for ME EQUIPMENT intended for use by a LAY OPERATOR include easily understood diagrams, illustrations, or photographs of the fully assembled and ready-to-operate ME EQUIPMENT including all controls, visual INFORMATION SIGNALS, and indicators provided (see 7.1)	Necessary information to set up was provided in the instruction.	P	
7.4.4	Additional requirements for ME EQUIPMENT start-up	PROCEDURE:	N/A	
	- Easily understood diagrams, illustrations, or photographs showing proper connection of the PATIENT to the ME EQUIPMENT, ACCESSORIES and other equipment (see 7.1)	No connection to PATIENT	N/A	
	- the time from switching "ON" until the ME EQUIPMENT is ready for NORMAL USE, when it exceeds 15 s (see 15.4.4 of Part 1) (s):	No such feature.	N/A	
	-the time required for ME EQUIPMENT to warm from the minimum storage temperature between uses until it is ready for intended use; and	No such conditions	N/A	
	-the time required for ME EQUIPMENT to cool from the maximum storage temperature between uses until it is ready for intended use; and	No such conditions	N/A	
·		•		

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Clause	Requirement + Test	Result - Remark	Verdict
7.4.5	Instructions for use for ME EQUIPMENT intended for use by a LAY OPERATOR include a description of generally known conditions in the HOME HEALTHCARE ENVIRONMENT that can unacceptably affect the BASIC SAFETY and ESSENTIAL PERFORMANCE of the ME EQUIPMENT	See RM table 7.4.5	P
	The steps that can be taken by the LAY OPERATOR to identify and resolve the above conditions		N/A
	At least the following issues are also included as app	licable	N/A
	- The effects of lint, dust, light (including sunlight), etc.		N/A
	- A list of known devices or other sources that can potentially cause interference problems		N/A
	- The effects of degraded sensors and electrodes, or loosened electrodes, that can degrade performance or cause other problems		N/A
	- The effects caused by pets, pests or children		N/A
	The instructions for use explain the meaning of the IP classification marked on the ME EQUIPMENT, and on any carrying case provided with the ME EQUIPMENT as applicable		N/A
7.4.6	Instructions for use include a troubleshooting guide for use when there are indications of a ME EQUIPMENT malfunction during start-up or operation	No need of such guide for power supply. But final determination in the end product.	N/A
	Troubleshooting guide discloses the necessary steps in the event of an TECHNICAI ALARM CONDITION		N/A
7.4.7	Instructions for use for ME EQUIPMENT, ME SYSTEMS, parts, and ACCESSORIES for other than single use that can be contaminated by contact with PATIENT, body fluids, or expired gases, during INTENDED USE, indicate the following:	No cleaning, disinfection and sterilization required for power supply. But final determination in the end product.	N/A
	- Frequency of cleaning, cleaning and disinfection, or cleaning and sterilization, as appropriate, for ME EQUIPMENT, ME SYSTEMS, parts, and ACCESSORIES used on the same PATIENT including rinsing methods, drying, handling, and storage between uses (see 8.1 and 8.2); and		N/A
	- It is necessary to clean and disinfect, clean and sterilize the ME EQUIPMENT, ME SYSTEMS, parts, and ACCESSORIES for multiple PATIENT use between uses on different PATIENTS, including rinsing methods, drying, handling, and storage until re-use (see 8.1 and 8.2), or		N/A

			-	
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Clause	Requirement + Test	Result - Remark	Verdic	
	- ME EQUIPMENT, ME SYSTEMS and ACCESSORIES require professional hygienic maintenance prior to re-use and provide contact details for the source of these services (see 7.5.2)		N/A	
7.4.8	Instructions for use include:		Р	
	- EXPECTED SERVICE LIFE of the ME EQUIPMENT:	5 years	Р	
	– EXPECTED SERVICE LIFE of parts and ACCESSORIES shipped with the ME EQUIPMENT:	5 years	Р	
	- SHELF LIFE of parts and ACCESSORIES shipped with ME EQUIPMENT when SHELF LIFE is less than the EXPECTED SERVICE LIFE:	No such parts	N/A	
7.4.9	Instructions for use include:		N/A	
	- A statement indicating the LAY RESPONSIBLE ORGANIZATION must contact its local authorities to determine the proper method of disposal of potentially bio hazardous parts and ACCESSORIES, as applicable	Not applicable for power supply	N/A	
7.4.10	Instructions for use includes the recommended placement of the remote parts of the DISTRIBUTED ALARM SYSTEM, when applicable, to ensure the OPERATOR can be notified at all times by an appropriate element of DISTRIBUTED ALARM SYSTEM within its specified range	Not applicable for power supply	N/A	
7.5	Technical description	·	N/A	
7.5.1	The technical description for PERMANENTLY INSTALLED CLASS I ME EQUIPMENT includes:	Not permanently installed	N/A	
	– A warning indicating the ME EQUIPMENT installation, including a correct PROTECTIVE EARTH CONNECTION, must only be carried out by qualified SERVICE PERSONNEL		N/A	
	- Specifications of the PERMANENTLY INSTALLED PROTECTIVE EARTH CONDUCTOR		N/A	
	 A warning to verify the integrity of the external protective earthing system 		N/A	
	 A warning to connect and verify that the PROTECTIVE EARTH TERMINAL of the PERMANENTLY INSTALLED ME EQUIPMENT is connected to the external protective earthing system 		N/A	
7.5.2	Technical description includes methods for cleaning and disinfection or cleaning and sterilization for ME EQUIPMENT and ACCESSORIES requiring professional hygienic maintenance prior to reuse (see 7.4.7):	No cleaning, disinfection and sterilization required for power supply.	N/A	
	- Before and after any type of service PROCEDURE		N/A	
	– When the ME EQUIPMENT is transferred to another PATIENT		N/A	

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Clause	Requirement + Test	Result - Remark	Verdict
8	PROTECTION AGAINST EXCESSIVE TEMPERAT HAZARDS	URES AND OTHER	N/A
8.1	A LAY OPERATOR in the HOME HEALTHCARE ENVIRONMENT can perform the cleaning or cleaning and disinfection PROCESSES when intended (see 7.4.7)	No cleaning, disinfection and sterilization required for power supply	N/A
	USABILITY of each such PROCESS pertaining to a LAY OPERATOR was investigated by the USABILITY ENGINEERING PROCESS		N/A
8.2	A LAY OPERATOR in the HOME HEALTHCARE ENVIRONMENT can perform the cleaning and sterilization PROCESSES when intended (see 7.4.7)	No sterilization required for power supply.	N/A
	USABILITY of each such PROCESS pertaining to a LAY OPERATOR was investigated by the USABILITY ENGINEERING PROCESS		N/A
8.3	Additional requirements for ingress of water or p EQUIPMENT and ME SYSTEMS	particulate matter into ME	N/A
8.3.1	TRANSIT-OPERABLE, HANDHELD, and BODY-WORN ME EQUIPMENT maintained BASIC SAFETY and ESSENTIAL PERFORMANCE after undergoing the test of IEC 60529 for at least IP 22		N/A
	All other ME EQUIPMENT maintained BASIC SAFETY and ESSENTIAL PERFORMANCE after undergoing the test of IEC 60529 for at least IP21:		N/A
	For PORTABLE ME EQUIPMENT intended to be used only while in a carrying case, IP21 met with the ME EQUIPMENT in its the carrying case		N/A
	Maintenance of BASIC SAFETY and ESSENTIAL PERFORMANCE VERIFIED		N/A
8.3.2	ENCLOSURES of the non-ME EQUIPMENT parts of the ME SYSTEMS provide the degree of protection against harmful ingress of water or particulate matter equivalent to equipment complying with their respective IEC or ISO safety standards		N/A
	Tests of IEC 60529:1989 conducted with the equipment placed in the least favourable position of NORMAL USE and the ENCLOSURES inspected		N/A
8.4	Additional requirements for interruption of the period ME EQUIPMENT and ME SYSTEM	ower supply/SUPPLY MAINS to	N/A
	ME EQUIPMENT or ME SYSTEM with ESSENTIAL PERFORMANCE intended to actively keep alive or resuscitate a PATIENT maintained its ESSENTIAL PERFORMANCE for a sufficient time or for a sufficient number of PROCEDURES when loss or failure of SUPPLY MAINS or near depletion INTERNAL ELECTRICAL POWER SOURCE occurred		N/A

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Clause	Requirement + Test	Result - Remark	Verdict
	The time or number of PROCEDURES remaining allowed alternative life-supporting methods to be employed		N/A
	Optionally, an INTERNAL ELECTRICAL POWER SOURCE was used to maintain ESSENTIAL PERFORMANCE:		N/A
	Optionally, independent means were used to provide ESSENTIAL PERFORMANCE		N/A
	Instructions for use disclose the time or number of procedures available following a loss or failure of the SUPPLY MAINS or near depletion of the INTERNAL ELECTRICAL POWER SOURCE		N/A
	Instructions for use describes the alternative life- supporting methods to be employed		N/A
	The technical description describes methods that can be employed for longer periods		N/A
	ME EQUIPMENT OF ME SYSTEM with ESSENTIAL PERFORMANCE intended to actively keep alive or resuscitate a PATIENT with no INTERNAL ELECTRICAL POWER SOURCE is equipped with an ALARM SYSTEM that includes at least a MEDIUM PRIORITY ALARM CONDITION indicating power failure		N/A
	ME EQUIPMENT OF ME SYSTEM with ESSENTIAL PERFORMANCE intended to actively keep alive of resuscitate a PATIENT with an INTERNAL ELECTRICAL POWER SOURCE is equipped with an automatic switchover to INTERNAL ELECTRICAL POWER SOURCE		N/A
	ME EQUIPMENT OF ME SYSTEM with ESSENTIAL PERFORMANCE intended to actively keep alive or resuscitate a PATIENT with an INTERNAL ELECTRICAL POWER SOURCE is equipped with an ALARM SYSTEM that includes at least a MEDIUM PRIORITY TECHNICAL ALARM CONDITION indicating the INTERNAL ELECTRICAL POWER SOURCE is nearing insufficient remaining power for operation		N/A
	TECHNICAL ALARM CONDITION provides sufficient time or sufficient number of procedures for a LAY OPERATOR to act		N/A
	A TECHNICAL ALARM CONDITION of at least LOW PRIORITY remained active until the INTERNAL ELECTRICAL POWER SOURCE returned to a level above the ALARM LIMIT or until depleted		N/A
	It was not possible to inactivate the visual ALARM SIGNAL of this TECHNICAL ALARM CONDITION		N/A
	Functional tests conducted, and the RISK MANAGEMENT FILE inspected		N/A
8.5	Additional requirements for an INTERNAL ELECTRICA	AL POWER SOURCE	N/A

N/A

N/A

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Clause	Requirement + Test	Result - Remark	Verdict
8.5.1	ME EQUIPMENT provided with a means for the OPERATOR to determine state of the INTERNAL ELECTRICAL POWER SOURCE when the is essential for BASIC SAFETY OR ESSENTIAL PERFORMANCE OR to control risks associated with loss of ESSENTIAL PERFORMANCE	No internal electrical power source	N/A
	State of INTERNAL ELECTRICAL POWER SOURCE indicated by:		N/A
	- number of PROCEDURES remaining;		N/A
	-remaining operating time;		N/A
	-percentage of the remaining operating time or energy; or		N/A
	-"fuel" gauge		N/A
	Instructions described method to determine state of INTERNAL ELECTRICAL POWER SOURCE		N/A
8.5.2	Means, other than labelling, provided to prevent RISK of swallowing coin/button cells		N/A
	Replacement of button cell require use of TOOL		N/A
8.5.3	For ME EQUIPMENT or ME SYSTEM with an INTERNAL ELECTRICAL POWER SOURCE, if simultaneous connection of the ME EQUIPMENT to the PATIENT and the SUPPLY MAINS is possible, then APPLIED PARTS and parts that are likely to come into contact with the PATIENT have two MOPP from the SUPPLY MAINS	No internal power	N/A
	Parts which the PATIENT intentionally handles as the intended OPERATOR while the ME EQUIPMENT is not being used for its intended medical function are insulated with two MOOP or two MOPP from SUPPLY MAINS.		N/A
9	ACCURACY OF CONTROLS AND INSTRUMENTS AGAINST HAZARDOUS OUTPUTS	SAND PROTECTION	N/A
	The RISKS associated with USABILITY in the HOME HEA OPERATOR PROFILES including a LAY OPERATOR when ENGINEERING PROCESS include at least the following o	performing the USABILITY	N/A
	- changes of controls		N/A
	- unexpected movement		N/A
	 potential for misconnection 		N/A
	- potential for improper operation, or unsafe use		N/A

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mode

- potential for confusion as to current operational

- change in the transfer of energy or substance

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Clause	Requirement + Test	Result - Remark	Verdict
	- exposure to environmental conditions specified in this standard		N/A
	- exposure to biological materials, and		N/A
	- small parts being inhaled or swallowed		N/A
	Particular emphasis placed on the limited training of a LAY OPERATOR with respect to the ability to intervene and maintain BASIC SAFETY and ESSENTIAL PERFORMANCE.		N/A
	The MANUFACTURER'S USABILITY ENGINEERING PROCESS included the least capable intended LAY OPERATOR OF LAY RESPONSIBLE ORGANIZATION		N/A
	USABILITY ENGINEERING FILE inspected for compliance		N/A

10	CONSTRUCTION OF ME EQUIPMENT	Р
10.1	Additional requirements for mechanical strength	Р
10.1.1	Additions to Table 28 Mechanical strength test of the base standard, conducted as indicated in Table 1, Mechanical strength test applicability, non-TRANSIT- OPERABLE, and Table 2, Mechanical strength test applicability, TRANSIT-OPERABLENon-transit-operable	Р
10.1.2	ME EQUIPMENT, its parts, and mounting ACCESSORIES, intended for non-TRANSIT-OPERABLE use displayed adequate mechanical strength when subjected to mechanical stress caused by NORMAL USE, including pushing, impact, dropping and rough handling (not applicable to FIXED and STATIONARY ME EQUIPMENT)	P
	ME EQUIPMENT maintained BASIC SAFETY and ESSENTIAL PERFORMANCE after mechanical tests	Р
	OPERATOR-re-settable protective devices that can be reset without the use of a TOOL were, optionally, reset prior to the evaluation of BASIC SAFETY and ESSENTIAL PERFORMANCE	Р
	a) Shock tests conducted in accordance with IEC 60068-2-27:2008	Р
	b) Broad-band random vibration tests conducted in accordance with IEC 60068-2-64:2008, using the following conditions	Р
10.1.3	ME EQUIPMENT, parts, and mounting ACCESSORIES for TRANSIT-OPERABLE use displayed adequate mechanical strength when subjected to pushing, impact, dropping, rough handling, and rigorous conditions of PATIENT movement in NORMAL USE as well as transportation by trolleys, carts, road vehicles, trains, ships, and aircraft	N/A

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Clause	Requirement + Test	Result - Remark	Verdict
	ME EQUIPMENT maintained BASIC SAFETY and ESSENTIAL PERFORMANCE after the following tests:		N/A
	a) Shock tests conducted on other than HAND-HELD ME EQUIPMENT, parts, and mounting ACCESSORIES in accordance with IEC 60068-2-27:2008		N/A
	1) Test type: Type 1:		N/A
	2) Test type: Type 2:		N/A
	b) Shock tests conducted on HAND-HELD ME EQUIPMENT, parts, and mounting ACCESSORIES in accordance with IEC 60068-2-27:2008		N/A
	1) Test type: Type 1:		N/A
	2) Test type: Type 2:		N/A
	c) Broad-band random vibration test conducted on ME EQUIPMENT, parts, and mounting ACCESSORIES in accordance with IEC 60068-2-64:2008		N/A
	d) Free fall tests conducted on PORTABLE and MOBILE ME EQUIPMENT, parts, and mounting ACCESSORIES per IEC 60068-2-31:2008, using PROCEDURE 1		N/A
	BASIC SAFETY and ESSENTIAL PERFORMANCE were maintained		N/A
10.2	Controls of ME EQUIPMENT intended for use by a LAY OPERATORY that can affect BASIC SAFETY or ESSENTIAL PERFORMANCE protected from accidental or unauthorized changes or adjustments		N/A
	OPERATOR-adjustable controls used for calibration include a means to prevent unintentional changes from the intended position		N/A

11	PROTECTION AGAINST STRANGULATION OR ASPHYXIATION		Р
	Means provided to control the RISK of strangulation and asphyxiation of the PATIENT and others to an acceptable level		Р
	EQUIPMENT and RISK MANAGEMENT FILE inspected :	See RM table 11	Р

12	ADDITIONAL REQUIREMENTS FOR ELECTROMAGNETIC EMISSIONS OF ME EQUIPMENT AND ME SYSTEMS		
		Not applicable to component power supply system	N/A

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Clause	Requirement + Test	Result - Remark	Verdict
13	ADDITIONAL REQUIREMENTS FOR ALARM SYST AND ME SYSTEMS	TEMS OF ME EQUIPMENT	N/A
13.1	Each HIGH PRIORITY and MEDIUM PRIORITY ALARM CONDITION causes generation of auditory ALARM SIGNALS per IEC 60601-1-8:2006 and IEC 60601-1- 8:2006/AMD1:2012, except when equipment is connected to a DISTRIBUTED ALARM SYSTEM intended for confirmed deliver of ALARM CONDITIONS including the generation of auditory ALARM SIGNALS per IEC 60601-1-8:2006 and IEC 60601-1- 8:2006/AMD1:2012		N/A
13.2	For ME EQUIPMENT and ME SYSTEMS intended to actively keep alive or resuscitate a PATIENT, reducing the auditory ALARM SIGNAL volume T below audible levels resulted in the following was not possible, except when the ALARM SYSTEM was connected to a DISTRIBUTED ALARM SYSTEM that included generation of auditory ALARM SIGNALS per IEC 60601-1-8:2006 and IEC 60601-1-8:2006/AMD1:2012		N/A

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Clause Requirement + Test

Result - Remark

Verdict

4.2.2	RM RESULTS TABLE: Permissible environmental conditions of transport and storage, between uses, indicated in instructions for use		
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph)	Result - Remarks	Verdict
5.3	Risk Management Report GT-RM2015-001 6.1	Operational hazard is identified	Р
5.4	Risk Management Report GT-RM2015-001 6.2	The severity of the harm has been estimated as "3"	Р
		The probability of occurrence of the harm has been estimated in "1"	
5.5	Risk Management Report GT-RM2015-001 6.4	Intended use is identified	Р

4.2.3.1	RM RESULTS TABLE: Environmental operating conditions - Continuous operating conditions			
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph)		Verdict	
4.2	Risk management procedure GTQPR05000 A2 5.0	Intended use is identified	Р	
4.3	Risk management procedure GTQPR05000 A2 5.0	Hazardous situation is identified	Р	
4.4	Risk management procedure GTQPR05000 A2 5.0	Severity and probability is identified	Р	

7.4.1	RM RESULTS TABLE: Addition	RM RESULTS TABLE: Additional requirements for warning and safety notices		
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph)			
5.2		Intended use is identified	N/A	
5.3	Risk Management Report GT-RM2015-001 6.1	Operational hazard is identified	Р	
5.4	Risk Management Report GT-RM2015-001 6.2	The severity of the harm has been estimated as "2"	Р	
		The probability of occurrence of the harm has been estimated in "1"		
5.5	Risk Management Report GT-RM2015-001 6.4	The risk is evaluated as "ACC'	Р	
6	Risk Management Report GT-RM2015-001 7	Intended use is identified	Р	
7.1	Risk Management Report GT-RM2015-001 8.1	The risk is evaluated as "ACC'	Р	

7		E	
1.	4.	J.	

RM RESULTS TABLE: : Additional requirements for operating instructions

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Clause	Requirement + Test Result - Remark		Verdict
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph)	Result - Remarks	Verdict
5.4	Risk Management Report GT-RM2015-001 6.2	Intended use is identified	Р
5.5	Risk Management Report GT-RM2015-001 6.4	Operational hazard is identified	Р
6	Risk Management Report GT-RM2015-001 7	The severity of the harm has been estimated as "2" The probability of occurrence of the harm has been estimated in "1"	Ρ
7.1	Risk Management Report GT-RM2015-001 8.1	The risk is evaluated as "ACC'	Р

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Clause Requirement + Test

Result - Remark

8.4	RM RESULTS TABLE: Additional requirements for interruption of power supply / supply mains to ME Equipment and ME Systems		
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph) Result - Remarks		Verdict
5.2			
5.3			
5.4			
6			
7.1			
7.2			
7.3			
7.4			
7.5			
7.6			

10.1.2a	1.2a TABLE: Shock test (IEC 60068-2-27:2008), using the following conditions*:						
	Peak acceleration:				150 m/s2 (15 g)		
	Duration.		·····:	11 ms			
	Pulse sha	аре	:	half-sine			
Number of shocks 3 s				3 shocks p	3 shocks per direction per axis (18 total)		
Directio App	n Shock llied	Axis Shock Applied	Applied BASIC SAFETY and ESSENTIAL PERFORMANCE maintained? Yes/No				
Positive		X axis ²	Yes		The enclosure shows		
Negative		X axis ²	Yes		cracks and there is no damaged or loosing p		
Positive		Y axis ²	Yes		the product after test.		
Negative		Y axis ²	Yes		The EUT worked as n and passed the dielec		
Positive		Z axis ²	Yes		strength test.		
Negative	ative Z axis ² Yes		Yes				
	ntary inform This represe	ation: ents Class 7M1 as descril	bed in IEC TR 60	721-4-7:20	01 [6])		

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Clause Requirement + Test

Result - Remark

10.1.2b	TABLE: Broad-band random vibration test (IEC 60068-2-64:2008) using the following conditions*:				08) using the	Р
1	Acceleration ar	Acceleration amplitude:			Hz: 1,0 (m/s²)²/Hz	
2	Acceleration ar	nplitude	:	100 Hz to 20	0 Hz: – 3 db per octa	ve
3	Acceleration ar	nplitude	·······	200 Hz to 2 0	000 Hz: 0,5 (m/s²)²/H	Z
	Duration		·······	30 min per pe	erpendicular axis (3 t	otal)
Perpendicular axis subjected to broad-band random vibration test		Acceleration amplitude	ESSENTIAL F	C SAFETY and L PERFORMANCE Rema lined? Yes/No		
	1	1	Yes		The enclosure shows no	
	2	1	Y	′es	cracks and there is no damaged or loosing part	
	3	1	Yes		inside the product after te	
	1	2	Y	′es	The EUT worked as norr and passed the dielectric	
	2	2	2 Yes		strength test.	
	3 2 Yes		′es			
1 3 2 3 3 3		3	Yes		7	
		3	Y	′es	7	
		Y	′es			

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Clause Requirement + Test

Result - Remark

10.1.3a1		Shock test (IEC 600 Id mounting ACCESS				N/A
	Peak acc	eleration	:	150 m/s ² (15 g)		
	Duration.		:	11 ms		
	Pulse sha	аре	:	half-sine		
	Number of	of shocks	:	3 shocks per dir	rection per axis (18 t	otal)
Direction App		Axis Shock Applied	BASIC SAFETY a PERFORMANCE Yes/	maintained?	Remarks	
• •	ntary inform This repres	ation: ents Class 7M2 as de	escribed in IEC/TR	60721-4-7:2001	[6])	

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Clause Requirement + Test

Result - Remark

10.1.3a2		Shock test (IEC 6006 IT, parts, and mount be 2):				N/A
	Peak acc	eleration	:	300 m/s ² (15 g)		
	Duration.		:	6 ms		
	Pulse sha	ape	:	half-sine		
	Number o	of shocks	:	3 shocks per di	rection per axis (18 t	otal)
Directior Appl		Axis Shock Applied	BASIC SAFETY a PERFORMANCE Yes/	maintained?	Remarks	i
Supplemen	tary informa	ation:				

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Clause Requirement + Test

Result - Remark

10.1.3b1		Shock test (IEC 6006 g ACCESSORIES using				N/A
	Peak acc	eleration	:	300 m/s ² (30 g)		
	Duration.		:	11 ms		
	Pulse sha	аре	:	half-sine		
	Number of	of shocks	:	3 shocks per di	rection per axis (18 t	otal)
Direction Appli		Axis Shock Applied		and ESSENTIAL E maintained? Remarks s/No		
Supplement	ary inform	ation:				
	•	ents Class 7M3 as de	scribed in IEC/TR 6	60721-4-7:2001.	(Test Type 1)	

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Clause Requirement + Test

Result - Remark

10.1.3b2		Shock test (IEC 6006 ACCESSORIES using				N/A
	Peak acce	eleration	:	1000 m/s² (100	g)	
	Duration		:	6 ms		
	Pulse sha	pe	:	half-sine		
	Number o	f shocks	:	3 shocks per dir	ection per axis (18 t	otal)
Direction Appli		Axis Shock Applied	Basic safety a PERFORMANCE Yes/	maintained?	Remarks	
Supplement	ary informa	ation:				

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Clause Requirement + Test

Result - Remark

10.1.3c		ad-band random v arts, and mounting			4:2008) on ME llowing conditions*:	N/A
1	Acceleration amplitude:			10 Hz to	100 Hz: 1,0 (m/s²)²/Hz	
2	Acceleration a	amplitude	:	100 Hz t	o 200 Hz: - 3 db per octa	ve
3	Acceleration a	amplitude	:	200 Hz t	o 2 000 Hz: 0,5 (m/s²)²/H	Z
	Duration		:	30 min p	er perpendicular axis (3 t	total)
subjected	ndicular axis d to broad-band vibration test	Acceleration amplitude	BASIC SAFET ESSENTIAL PERFO maintained? Y	ORMANCE	Remarks	
	1	1				
	2	1				
	3	1				
	1	2				
	2	2				
	3	2				
1 3 2 3		3				
	3	3				

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Clause Requirement + Test

Result - Remark

Verdict

10.1.3d		UIPMENT, p			EDURE 1, ON PORTABLE NES (with carrying case if	N/A
1	Fall height for ma	ss ≤ 1 kg	0,25 m			
2	Fall height for ma	ss > 1 kg a	nd ≤ 10 Kg:	0,1 m		
3	Fall height for ma	ss > 10 kg	and ≤ 50 Kg:	0,05 m		
4	Fall height for ma	ss > 50 kg	:	0,01 m		
Specified altitude (m	Mass) (Kg)	Fall No.	BASIC SAFETY and ESS PERFORMANCE mainta Yes/No		Remarks	
0,25	≤ 1	1				
0,25	≤ 1	2				
0,1	> 1 & ≤ 10	1				
0,1	> 1 & ≤ 10	2				
0,05	> 10 & ≤ 50	1				
0,05	> 10 & ≤ 50	2				
0,01	> 50	1				
0,01	> 50	2				

(*NOTE 6 This represents Class 7M2 as described in IEC/TR 60721-4-7:2001)

11.0	RM RESULTS TABLE: PROTECTION AGAINST STRANGULATION AND ASPHYXIATION				
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph)	Result - Remarks	Verdict		
5.4	Risk Management Report	Intended use is identified			
	GT-RM2015-001 6.2				
5.5	Risk Management Report	Operational hazard is identified			
	GT-RM2015-001 6.4				
6	Risk Management Report	The severity of the harm has been estimated as			
	GT-RM2015-001 7	"2"			
		The probability of occurrence of the harm has been estimated in "1"			
7.1	Risk Management Report	The risk is evaluated as "ACC'			
	GT-RM2015-001 8.1				
7.2	Risk Management Report	specifications control is identified			
	GT-RM2015-001 8.1				
7.3	Risk Management Report	Product specification			
	GT-RM2015-001 8.2				

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Clause	Requirement + Test		Result - Remark	Verdict
11.0	RM RESULTS TABLE: PROTEC	CTION AGAINST ST	FRANGULATION AND	Р
Clause of ISO 14971	Document Ref. in RMF (Document No. & paragraph)	Result - Remarks		Verdict
7.4	Risk Management Report GT-RM2015-001 8.3	The residual risk	is evaluated as 'ACC'	
Suppleme	ntary information:	·		