

Test Report issued under the responsibility of:



## TEST REPORT IEC 60601-1-6 Medical electrical equipment - Part 1-6:

# General requirements for basic safety and essential performance -Collateral standard: Usability

Report Number:	220201761SHA-002
Date of issue:	2022-11-29
Total number of pages:	15
Name of Testing Laboratory preparing the Report:	Intertek Testing Services Shanghai Building No. 86, 1198 Qinzhou Road (North) Shanghai 200233 China
Applicant's name:	GlobTek, Inc.
Address:	186 Veterans Dr. Northvale, NJ 07647 USA
Test specification:	
Standard::	IEC 60601-1-6:2010, AMD1:2013, AMD2:2020 for use in conjunction with IEC 62366-1:2015, AMD1:2020, and IEC 60601-1:2005, AMD1:2012, AMD2:2020
Test procedure:	CB Scheme
Non-standard test method	N/A
TRF template used:	IECEE OD-2020-F1:2020, Ed.1.3
Test Report Form No	IEC60601_1_6K
Test Report Form(s) Originator :	TÜV Rheinland of North America
Master TRF:	Dated 2020-11-23
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Test	item description:	Medica	al Power Supply	
Trad	e Mark(s):	GlobTek, Inc.		
Manu	ıfacturer :	GlobTek, Inc. 186 Veterans Dr. Northvale, NJ 07647 USA		
Mode	el/Type reference:	GT*96180-******, GT*96300-******, GT*91120-*****, GTM91128LI*CEL**-****, GTM91128***-****, GTM91128LI1CEL, GTM91128LI2CEL, GTM91128LI3CEL		
Datir		(Refer	to IEC 60601-1 report 22	20201761SHA-001)
Ratir	igs	(Relei	10 IEC 60601-1 Tepolt 22	202017615HA-001)
Resp	oonsible Testing Laboratory (as a	pplicat	ble), testing procedure	and testing location(s):
$\square$	CB Testing Laboratory:		Intertek Testing Services	s Shanghai
Testi	ng location/ address	:	Building No. 86, 1198 Q 200233 China	linzhou Road (North) Shanghai
Teste	ed by (name, function, signature)	:	Vivian Xu (Engineer)	Vi Vian . Xu.
Approved by (name, function, signature):		Larry Zhong (Mandated reviewer)	Lany Zhang	
	Testing procedure: CTF Stage 1:		N/A	
Testi	ng location/ address	:		
Tested by (name, function, signature):				
Approved by (name, function, signature):				
	Testing procedure: CTF Stage 2:		N/A	
Testi	ng location/ address			
Teste	ed by (name + signature)	:		
Witn	essed by (name, function, signat	ure).:		
Appr	oved by (name, function, signatu	ire):		
	Testing procedure: CTF Stage 3:		N/A	
	resting procedure: CIF Stage 4:		N/A	
lesting location/ address:				
Nitropped by (name, function, signature)				
Approved by (name, function, signature) .:		ure)		
Approved by (name, function, signature):				
Supe	a vised by (name, function, signa	urej.		<u> </u>

List of Attachments (including a total number of pages in each attachment): ANNEX I – IEC 62366-1:2015, AMD1:2020 – Usability engineering process checklist (Pages: 9)			
Summary of testing:			
Tests performed (name of test and test clause):	Testing location: N/A		
Process standard only, no testing			
Summary of compliance with National Difference None	es (List of countries addressed):		
The product fulfils the requirements of IEC 60601-1-6:2010, AMD1:2013, AMD2:2020			
Statement concerning the uncertainty of the mea	surement systems used for the tests		
$\boxtimes$ Internal procedure used for type testing through which traceability of the measuring uncertainty has been established:			
Procedure number, issue date and title: GMS-QC-12 Estimation of Measurement Uncertainty, 1-July-2012 Initial Release.			
Calculations leading to the reported values are on file with the NCB and testing laboratory that conducted the testing.			
Statement not required by the standard used	for type testing		

#### 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.

See IEC 60601-1 report 220201761SHA-001

Test item particulars:		
Classification of installation and use	Direct plug-in for power adapter model.	
	Appliance coupler for one type of open frame model series.	
	Final evaluation in end product.	
Supply Connection	Direct plug-in for 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:	No test need	
Date of receipt of test item:	N/A	
Date (s) of performance of tests:	N/A	
General remarks:		
"(See Enclosure #)" refers to additional information ap "(See appended table)" refers to a table appended to th	pended to the report. ne report.	
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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 the state of the s	he General product information section.	
Name and address of factory (ies):	Factory 1 GlobTek, Inc. 186 Veterans Dr. 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	

General product information and other remarks: See IEC 60601-1 report 220201761SHA-001

### IEC 60601-1-6:2010, AMD1:2013, AMD2:2020

Clause	Requirement + Test

Result - Remark

Verdict

4.0	GENERAL REQUIREMENTS		Р
4.2	USABILITY ENGINEERING PROCESS complies with IEC 62366-1 including amended definitions. Excludes production and post-production monitoring, and maintenance of the USABILITY ENGINEERING PROCESS	See attached IEC 62366-1 ANNEX I	Ρ
	Inspection of the USABILITY ENGINEERING FILE verified	that the MANUFACTURER	Р
	- established a USABILITY ENGINEERING PROCESS	See QF-GT-DJD-7.3.2-13 Usability Engineering File P2/1.2	Р
	- established acceptance criteria for USABILITY; and	See QF-GT-DJD-7.3.2-13 Usability Engineering File P5/1.15	Ρ
	<ul> <li>demonstrated that the acceptance criteria for USABILITY have been met.</li> </ul>	See QF-GT-DJD-7.3.2-13 Usability Engineering File P5/1.15	Р

5	ME EQUIPMENT ACCOMPANYING DOCUMENTS	;	Р
	The instructions for use shall contain a summary of the USE SPECIFICATION as specified in IEC 62366-1:2015, AMD1:2020, Clause 5.1	Refer to "POWER SUPPLY INFORMATION" and "ELECTRICAL SPECIFICATIONS" of SPEC	Ρ
	The same information is also included in the technical description, if this is provided as a separate document from instructions for use		Р

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	ANNEX I - IEC 62366-1:2015, AMD1:2020 - Usability	engineering process checklist	
Clause	Requirement + Test	Result - Remark	Verdict

4	PRINCIPLES		Р
4.1.1	The MANUFACTURER has established, documented and maintains a USABILITY ENGINEERING PROCESS addressing USER interactions with the MEDICAL DEVICE according to the ACCOMPANYING DOCUMENT	QF-GT-DJD-7.3.2-13 Usability Engineering File	Ρ
4.1.2	The USABILITY ENGINEERING PROCESS complies with this standard and the acceptance criteria in the USABILITY VALIDATION plan have been met	QQF-GT-DJD-7.3.2-13 Usability Engineering File	Р
4.1.3	Information for SAFETY used as a RISK CONTROL measure has been evaluated according to the USABILITY ENGINEERING PROCESS	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 5 section1.15	Ρ
4.2	The results of the USABILITY ENGINEERING PROCESS are recorded in the USABILITY ENGINEERING FILE	QQF-GT-DJD-7.3.2-13 Usability Engineering File	Р
4.3	The USABILITY ENGINEERING PROCESS is scaled-up or scaled-down based on the significance of the modification as determined by the results of the RISK ANALYSIS:	Document Reference No. in usability engineering file: QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 6	Ρ

5	USABILITY ENGINEERING PROCESS		Р
5.1	The MANUFACTURER shall prepare a USE SPECIFCATION. The USE SPECIFICATION shall include the following	Document Reference No. in usability engineering file: QQF-GT-DJD-7.3.2-13 Usability Engineering File	Ρ
	- intended medical indication	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.4	Р
	- intended PATIENT population	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.4	Р
	<ul> <li>intended part of the body or type of tissue applied to or interacted with</li> </ul>	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.4	Р
	- intended USER PROFILE	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.5	Р
	- intended USE ENVIRONMENT	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.6	Р
	- operating principle	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.7	Р

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	ANNEX I - IEC 62366-1:2015, AMD1:2020 – Usability engineering process checklist		
Clause	Requirement + Test	Result - Remark	Verdict

5.2	The MANUFACTURER shall identify USER INTERFACE characteristics that could be related to SAFETY as part of a RISK ANALYSIS performed according to ISO 14971:2019, Clause 5.3	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 6, section 2	Р
5.3	As part of this RISK ANALYSIS, the MANUFACTURER shall identify known or foreseeable HAZARDS and HAZARDOUS SITUATIONS which could affect PATIENTS, USERS or others, related to the use of the MEDICAL DEVICE.	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 6, section 2	Ρ
5.4	The RISK ANALYSIS includes a description of all the reasonably foreseeable HAZARD-RELATED USE SCENARIOS associated with the identified HAZARD and HAZARDOUS SITUATIONS.	Document Reference No. in usability engineering file: QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.7	Ρ
	The description of each identified HAZARD-RELATED USE SCENARIO includes all TASKS and their sequences	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.7	Р
	The SEVERITY of the possible resulting associated HARM was determined	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.7	Р
5.5	The MANUFACTURER shall select the HAZARD- RELATED USE SENARIOS to be included in a SUMMATIVE EVALUATION as part of the USABILITY FILE. This SUMMATIVE EVALUATION shall include:	Document Reference No. in usability engineering file:	Р
	- all HAZARD-RELATED USE SCENARIOS;		Р
	- a subset of the HAZARD-RELATED USE SCENARIOS based on the SEVERITY of the potential HARM that could be caused by USE ERROR (e.g. for which medical intervention would be needed); or		Р
	- a subset of the HAZARD-RELATED USE SCENARIOS based on the SEVERITY of the potential HARM and based on other circumstances specific to the MEDICAL DEVICE and the MANUFACTURER		N/A
	A summary of any selection scheme, the rationale for its use and the results of applying it shall be stored in the USABILITY ENGINEERING FILE	:	N/A
5.6	The MANUFACTURER shall establish and maintain a USER INTERFACE SPECIFICATION	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Р
5.7	The MANUFACTURER shall establish and maintain a USER INTERFACE EVALUATION plan for the USER INTERFACE	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Р

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	ANNEX I - IEC 62366-1:2015, AMD1:2020 - Usability	engineering process checklist	
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5.8	The MANUFACTURER shall design and implement the USER INTERFACE, including the ACCOMPANYING DOCUMENTATION if needed, and training capability, if needed, as described in the USER INTERFACE SPECIFICATION	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Ρ
	Where new USE ERRORS, HAZARDS, HAZARDOUS SITUATIONS or HAZARD-RELATED USE SCENARIOS are discovered during this step the MANUFACTURER shall repeat the steps of Clause 5 as appropriate	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 6	Р
	If training on the specific MEDICAL DEVICE is required for the safe us of the MEDICAL DEVICE by the intended USER, the MANUFACTURER shall design and implement a training capability for the EXPECTED SERVICE LIFE of the MEDICAL DEVICE by doing at least one of the following:	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 9, section 7.2	Ρ
	- provide the materials necessary for training;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 9, section 7.2	Ρ
	<ul> <li>ensure the materials necessary for training are available;</li> </ul>	QF-GT-DJD-7.3.2-12Usability Engineering File Page 9, section 7.2	Р
	- make the training available; or	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 9, section 7.2	Ρ
	- make training available to the RESPONSIBLE ORGANIZATION that enables it to train its USERS	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 9, section 7.2	Р
5.9	The MANUFACTURER shall perform a SUMMATIVE EVALUATION of each HAZARD-RELATED USE SCENARIO selected in Clause 5.5	Document Reference No. in usability engineering file: QQF-GT-DJD-7.3.2-13 Usability Engineering File Page89, section 6	Ρ
	All USE ERRORS and use difficulties that occurred shall be identified	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page89, section 6	Р
	Where USE ERROR or use difficulty can lead to a HAZARDOUS SITUATION the root causes should be determined	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page89, section 6	Ρ
	If new USE ERRORS, HAZARDS, HAZARDOUS SITUATIONS or HAZARD-RELATED USE SCENARIOS are discovered during this data analysis:		—
	- if yes, then the MANUFACTURER shall repeat the activities of Clause 5 as appropriate;		Р
	- if not, then the MANUFACTURER determine whether further improvement of the USER INTERFACE design as it relates to SAFETY is necessary and practicable		Р

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	ANNEX I - IEC 62366-1:2015, AMD1:2020 – Usability	engineering process checklist	
Clause	Requirement + Test	Result - Remark	Verdict

	1)	If yes, then the MANUFACTURER shall re- enter the USABILITY ENGINEERING PROCESS at Clause 5.6		Р
	2)	If not then the MANUFACTURER shall:		Р
	i)	Document why improvement is not necessary or not practicable;		Р
	ii)	Identify the data from the USABILITY ENGINEERING PROCESS needed to determine the RESIDUAL RISK related to use; and		Р
	iii)	Evaluate the RESIDUAL RISK according to ISO 14971:2019, Clause 7.3		Р
5.10	USER IN was ev the req	TERFACE OF UNKNOWN PROVENANCE (UOUP) valuated according to Annex C rather than uirements of 5.1 through 5.9.	See Annex C below	Р

Annex C	Evaluation of a USER INTERFACE OF UNKNOW	VN PROVENANCE (UOUP)	Р
C.2.1	The MANUFACTURER shall establish a USE SPECIFICATION as required in 5.1.	Document Reference No. in usability engineering file:	Р
		QQF-GT-DJD-7.3.2-13 Usability Engineering File	
C.2.2	The MANUFACTURER of a device with UOUP shall review POST-PRODUCTION information including	Document Reference No. in usability engineering file:	Р
	complaints and field reports for incidents and near incidents. All identified cases of USE ERROR shall be stored in the USABILITY ENGINEERING FILE and addressed in C.2.3 and C.2.4	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.8	
C.2.3	The MANUFACTURER shall review the RISK ANALYSIS of the MEDICAL DEVICE with UOUP and ensure that all	Document Reference No. in usability engineering file:	Р
	HAZARDS and HAZARDOUS SITUATIONS associated with USABILITY have been identified and documented	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 6	
C.2.4	The MANUFACTURER shall verify and document that adequate RISK CONTROL measures have been	Document Reference No. in usability engineering file:	Р
	implemented for all identified HAZARDS and HAZARDOUS SITUATIONS identified in C.2.3 and that all RISKS are reduced to an acceptable level as indicated by the RISK ASSESSMENT	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 6	
C.2.5	Based on any new information identified in performing steps C.2.3 and C.2.4 the MANUFACTURER re-evaluated the overall RESIDUAL RISK according to ISO 14971:2019, Clause 7.3 and documented the results in either the USABILITY ENGINEERING FILE OR RISK MANAGEMENT FILE	Document Reference No. in usability engineering file or Risk Management File: GT-RM2019-002	P

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ANNEX I - IEC 62366-1:2015, AMD1:2020 – Usability engineering process checklist

Clause	Requirement + Test	Result - Remark	Verdict
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Table 5.3	USABILITY ENGINEERING	FILE RESULTS TABLE: RISK A	ANALYSIS	Р
		Document Ref. in USABILITY ENGINEERING FILE	Result - Remarks	Verdict
An identification of known or foreseeable HAZARDS and HAZARDOUS SITUATIONS which could affect PATIENTS, USERS or others, related to the use of the MEDICAL DEVICE. was performed according to ISO 14971:2019, Clause 5.3		QQF-GT-DJD-7.3.2-13 Usability Engineering File	Acceptable according to IEC 62366-1	Ρ
During the	identification of HAZARD	DS and HAZARDOUS SITUATIONS, the following was considered:		_
– USE SPEC USER PROF	SIFICATION, including ILE(S) (See 5.1)	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.5	Acceptable according to IEC 62366-1	Р
<ul> <li>information</li> <li>HAZARDOUS</li> <li>for existing</li> <li>MEDICAL DE</li> <li>type, if available</li> </ul>	on on HAZARDS and S SITUATIONS known USER INTERFACES of VICES of a similar ilable; and	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.5	Acceptable according to IEC 62366-1	Ρ
<ul> <li>identified</li> <li>5.2).</li> </ul>	USE ERRORS (See	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 4, section 1.5	Acceptable according to IEC 62366-1	Р

Table 5.6	USABILITY ENGINEERING FILE	RESULTS TABLE: USER INTE	RFACE SPECIFICATION	Р
		Document Ref. in USABILITY ENGINEERING FILE	Result - Remarks	Verdict
USER INTERFACE SPECIFICATION		QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Р
The USER I	The USER INTERFACE SPECIFICATION shall consider:			
– the USE S	PECIFICATION (See 5.1)	QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Р
– the know ERRORS as device (Se	n or foreseeable USE sociated with the medical e 5.2); and	QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Р
– the наzаl (See 5.4)	RD-RELATED USE SCENARIOS	QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Р
Inputs to th	USER INTERFACE SPECIFICA	TION shall include the following	:	_

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Clause Requirement + Test

Result - Remark

Verdict

Table 5.6	USABILITY ENGINEERING FILE RESULTS TABLE: USER INTERFACE SPECIFICATION			
		Document Ref. in USABILITY ENGINEERING FILE	Result - Remarks	Verdict
<ul> <li>testable to relevant to including the parts of the associated CONTROL m</li> </ul>	echnical requirements the USER INTERFACE, ne requirements for those USER INTERFACE with the selected RISK neasures;	QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Ρ
– an indica ACCOMPAN <sup>®</sup> required; a	tion as to whether YING DOCUMENTATION is nd	QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Р
– an indica DEVICE spe	tion as to whether MEDICAL cific training is required	QF-GT-DJD-7.3.2-13 Usability Engineering File Page 7, section 3	Acceptable according to IEC 62366-1	Р

Table 5.7	USABILITY ENGINEER	RING FILE RESULTS TABLE: USER	INTERFACE EVALUATION plan	Р
		Document Ref. in USABILITY ENGINEERING FILE	Result - Remarks	Verdict
The manufacturer shall establish and maintain a USER INTERFACE EVALUATION plan for the USER INTERFACE		QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ
The USER I	NTERFACE EVALUATIO	N plan shall document:		_
a) the object the method FORMATIVE SUMMATIVE	ctive and identify I of any planned EVALUATIONS and EVALUATIONS	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
<ul> <li>b) if USABILITY TESTS are employed,</li> <li>document which USER</li> <li>GROUPS are intended to be included in the test;</li> </ul>		QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ
<ul> <li>document</li> <li>environment</li> <li>conditions</li> <li>the USE SPR</li> </ul>	nt the test nt and other of use, based on ECIFICATION;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ
<ul> <li>– specify w ACCOMPANY DOCUMENTA during the state</li> </ul>	vhether YING ATION is provided test; and	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
- specify w DEVICE-spe provided pi the minimu between th beginning of	thether MEDICAL cific training is rior to the test and m elapsed time e training and the of the test.	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ

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ANNEX I - IEC 62366-1:2015, AMD1:2020 – Usability engineering process checklist

Clause Requirement + Test

Result - Remark

Verdict

Table 5.7	USABILITY ENGINEER	ING FILE RESULTS TABLE: USER	INTERFACE EVALUATION plan	Р
		Document Ref. in USABILITY ENGINEERING FILE	Result - Remarks	Verdict
The USER I	NTERFACE evaluation	plan for FORMATIVE EVALUATION sl	nall address:	—
a) the evalution being used	uation methods ;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
b) which pa INTERFACE and	art of the USER is being evaluated;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
c) when in ENGINEERIN perform ea INTERFACE	the USABILITY IG PROCESS to ch of the USER EVALUATIONS.	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
For each s	elected HAZARD-RELA	TED USE SCENARIO (see 5.5), the use shall specify:	JSER INTERFACE EVALUATION	_
a) the evaluation of the evalu	uation method and a rationale athod produces EVIDENCE;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	
b) which pa INTERFACE	art of the USER is being evaluated;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
c) where a criteria for whether the SAFETY is p understand CORRECT U DEVICE (4.1	oplicable, the determining e information for erceivable, lable and supports SE of the MEDICAL .3);	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
d) the avail ACCOMPAN DOCUMENT/ provision o the SUMMA and	ability of the YING ATION and f training during TIVE EVALUATION;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ
e) for a US/ – how the o the test pair representa USER PROFI	ABILITY TEST, characteristics of rticipants are tive of the intended LES;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ
<ul> <li>justifying participants distinct USE purpose of number of</li> </ul>	how the test s are grouped into ER GROUPS for the determining the test participants;	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ

ANNEX I - IEC 62366-1:2015, AMD1:2020 – Usability engineering process checklist

Clause	Requirement + Test

Result - Remark

Verdict

Table 5.7	USABILITY ENGINEERING FILE RESULTS TABLE: USER INTERFACE EVALUATION plan			Р
		Document Ref. in USABILITY ENGINEERING FILE	Result - Remarks	Verdict
- the test e conditions rationale fo adequately	environment and of use and a or how they are	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
USE ENVIRONMENT;				
- the defin USE for eac USE SCENA	ition of CORRECT Ch HAZARD-RELATED RIO; and	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Р
- the meth data during TEST for the analysis of ERRORS an	od of collecting the USABILITY subsequent observed USE d use difficulties.	QQF-GT-DJD-7.3.2-13 Usability Engineering File Page 8, section 4	Acceptable according to IEC 62366-1	Ρ

(ANNEX1)