# **UL TEST REPORT AND PROCEDURE**

Standard:	ANSI/AAMI ES60601-1 (2005 + C1:09 + A2:10)(Medical Electrical Equipment - Part 1: General Requirements for Basic Safety and Essential Performance) CAN/CSA-C22.2 No. 60601-1 (2008) (Medical Electrical Equipment - Part 1: General Requirements for Basic Safety and Essential Performance)			
Certification Type: CCN:	Component Recognition QQHM2, QQHM8 (Power Supplies, Medical and Dental)			
Product:	Switching Power Supply			
Model:	LPT100A-M, LPT100B-M, LPT100C-M, LPT101-M, LPT102-M, LPT103-M and LPT104-M			
Rating:	LPT100A-M and LPT101-M: Input: AC 100 - 250V; 50/60Hz; 1.9A DC 127Vmin 300Vmax Outputs: V1: +2.0 to +16.5V; 18.0A V2: +2.0 to +16.5V; 9.0A			
	V3: +7.2 to +16.5V; 2.3 A (at 130W with 200LFM forced-air cooling)			
	V1: +2.0 to +16.5V; 13.0A V2: +2.0 to +16.5V; 5.0A V3: +7.2 to +16.5V; 1.0A (80W at natural convection cooling)			
	Maximum continuous output power: - 130W with 200LFM Forced Air Cooling - 80W with Natural Convection Cooling			
	LPT100B-M, LPT102-M and LPT103-M: Input: AC 100 - 250V; 50/60Hz; 1.9A DC 127Vmin 300Vmax Outputs: V1: +2.0 to +16.5V; 18.0A V2: +2.0 to +16.5V; 9.0A V3: -7.2 to -16.5V; 2.3 A (at 130W with 200LFM forced-air cooling)			
	V1: +2.0 to +16.5V; 13.0A V2: +2.0 to +16.5V; 5.0A V3: -7.2 to -16.5V; 1.0A (80W at natural convection cooling)			
	Maximum continuous output power: - 130W with 200LFM Forced Air Cooling - 80W with Natural Convection Cooling			
	LPT100C-M and LPT104-M:			
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		Input: AC 100 -	250V; 50/60Hz; 1.9A	
		DC 127Vi	min 300Vmax	
		Outputs:		
		V1: +2.0 to +16.	5V; 18.0A	
		V2: +21.6 to +28	3.8V; 3.0A	
		V3: +7.2 to +16.		
		(at 130W with 2	00LFM forced-air cooling)	
		V1: +2.0 to +16.	5V; 13.0A	
		V2: +21.6 to +28	3.8V; 1.5A	
		V3: +7.2 to +16.	5V; 1.0A	
		(80W at natural	convection cooling)	
		Maximum contir	nuous output power:	
			LFM Forced Air Cooling	
		- 80W with Natu	ral Convection Cooling	
Applicant Name and Add	me and Address:	ASTEC INTERN 16TH FL	IATIONAL LTD - PHILIPPINE E	BRANCH
		2 WING YIP ST		
			KOWLOON HONG KONG	

Report Reference #

E182560-A61-UL

This is to certify that representative samples of the products covered by this Test Report have been investigated in accordance with the above referenced Standards. The products have been found to comply with the requirements covering the category and the products are judged to be eligible for Follow-Up Service under the indicated Test Procedure. The manufacturer is authorized to use the UL Mark on such products which comply with this Test Report and any other applicable requirements of UL LLC ('UL') in accordance with the Follow-Up Service Agreement. Only those products which properly bear the UL Mark are considered as being covered by UL's Follow-Up Service under the indicated Test Procedure.

The applicant is authorized to reproduce the referenced Test Report provided it is reproduced in its entirety.

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UL authorizes the applicant to reproduce the latest pages of the referenced Test Report consisting of the first page of the Specific Technical Criteria through to the end of the Conditions of Acceptability.

Any information and documentation involving UL Mark services are provided on behalf of UL LLC (UL) or any authorized licensee of UL.

Prepared by: Sammi Liang

Issue Date:

2014-02-28

Reviewed by: Calvin Tang

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#### **Supporting Documentation**

The following documents located at the beginning of this Procedure supplement the requirements of this Test Report:

- A. Authorization The Authorization page may include additional Factory Identification Code markings.
- B. Generic Inspection Instructions
  - i. Part AC details important information which may be applicable to products covered by this Procedure. Products described in this Test Report must comply with any applicable items listed unless otherwise stated in the body of this Test Report.
  - ii. Part AE details any requirements which may be applicable to all products covered by this Procedure. Products described in this Test Report must comply with any applicable items listed unless otherwise stated in the body of each Test Report.
  - iii. Part AF details the requirements for the UL Certification Mark which is not controlled by the technical standard used to investigate these products. Products are permitted to bear only the Certification Mark(s) corresponding to the countries for which it is certified, as indicated in each Test Report.

#### **Product Description**

1. The equipment is a Class I switching type power supply which electronic components mounted on PWB for building into medical electrical equipment.

2. Refer to Enclosure ID 7-02 for Maximum allowed rating for convection forced - Air Cooling.

#### Model Differences

See Enclosure 7-02 for details.

Model LPT100B-M is identical to model LPT100A-M except polarity of output V3 rating.

Model LPT100C-M is identical to model LPT100A-M except L8, L100, L101, L401, addition of Heatsink on V2 module and Output rating.

Model LPT101-M is identical to LPT100A-M excepted for the model designation.

Model LPT102-M, LPT103-M are identical to LPT100B-M excepted for the model designation.

Model LPT104-M is identical to LPT100C-M excepted for the model designation.

### **Technical Considerations**

- Classification of installation and use : Open frame Component to be evaluated in end product
- Device type (component/sub-assembly/ equipment/ system) : Component
- Intended use (Including type of patient, application location) : Component to be evaluated in end product
- Mode of operation : Continuous
- Supply connection : Primary connector to be evaluated in end product
- Accessories and detachable parts included : None
- Other options include : None
- The product was investigated to the following additional standards:: ANSI/AAMI ES60601-1 (2005 + C1:09 + A2:10) (Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance) Edition 1 Revision Date 2012/01/01;, CAN/CSA-C22.2 No. 60601-1 (2008) (Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance) Edition 2 Revision Date 2011/06/01;, IEC 60601-1: 2005 + CORR.1 (2006) + CORR.2 (2007) (Medical Electrical Equipment Part 1: General Requirements for Basic Safety and Essential Performance) Edition 3;
- The product was not investigated to the following standards or clauses:: Electromagnetic Compatibility (IEC 60601-1-2), Clause 14, Programmable Electronic Systems, Biocompatibility (ISO 10993-1)
- The degree of protection against harmful ingress of water is:: Ordinary
- The mode of operation is:: Continuous
- The product is suitable for use in the presence of a flammable anesthetics mixture with air or oxygen or with nitrous oxide:: No
- The product is Recognized only to the following hazards: Fire, Shock.

## Engineering Conditions of Acceptability

For use only in or with complete equipment where the acceptability of the combination is determined by UL LLC. When installed in an end-product, consideration must be given to the following:

- The power supplies have been judged on the basis of the required creepage and clearances in the Standard for Medical Electrical Equipment, ANSI/AAMI ES 60601-1, Sub clause 8.9.
- The power supplies have been evaluated as a Class I, continuous operation equipment and has not been evaluated for use in the presence of a flammable anesthetic mixture with air, oxygen, or nitrous

oxide. An additional evaluation shall be made if the power supply is intended for use in other than Class I equipment.

- The power supplies were tested on a 20 A branch circuit. If used on a branch circuit greater than this, additional testing may be necessary.
- The power supplies were evaluated as 2 MOPP provided between Primary and Secondary, and 1 MOPP provided between Primary and Earth; see insulation diagram for details.
- Consideration should be given to measuring the temperatures on power electronic components and transformer windings when the power supply is installed in the end-use equipment. The primary transformer (T1 / T2) incorporates a Class 155 (F) insulation system.
- Total continuous output power from all output rails shall not exceed 130W with 200LFM forced-air cooling. Total continuous output power at natural convection cooling is 80W. Output derates 2.5% per degree from 50degC to 70degC ambient temp. (half load at 70degC). The force air-cooling condition (Refer to Enclosure ID 7-03) shall be considered in end product installation.
- The output connectors are not acceptable for field connection and are only intended for connections to mating connectors of internal wiring inside the end use product. The acceptability of these and the mating connectors relative to secureness, insulating materials, and temperatures shall be considered in the end use product.
- A suitable Electrical, Mechanical and Fire enclosure shall be provided in the end use product.
- The end-product Electric Strength Test is to be based upon a maximum working voltage of T1: 279.5 Vrms, 460 Vpk; T2: 287.8 Vrms, 442 Vpk.
- Instructions and equipment marking shall be provided in a language, which is acceptable in the country in which the equipment is to be installed.
- The end product should ensure that the requirements related to accompanying documents, clause 7.9, are met.
- The component shall be installed in compliance with the enclosure, mounting, marking, spacing, and separation requirements of the end use application.
- Input terminal/connector shall be connected to the supply neutral in the end use for simultaneous disconnection of all supply poles.
- This secondary circuit of this power supply has not been evaluated for patient connected applications.
- Additional UL Recognized DC Fuse must be provided in end-system for DC input.
- The following tests shall be performed/further considered in the end-product evaluation: Impedance and Current Carrying Capability, Temperature Test, Dielectric Voltage Withstand Tests, Leakage Test and Interruption of Power Supply Test.
- End product Risk Management Process to include consideration of requirements specific to the Power Supply.
- End product Risk Management Process to consider the need for simultaneous fault condition testing.
- End product Risk Management Process to consider the need for different orientations of installation during testing.
- End product to determine the acceptability of risk in conjunction to the movement of components as part of the power supply.
- End product to determine the acceptability of risk in conjunction to the routing of wires away from moving parts and sharp edges as part of the power supply.
- Temperature Test was conducted without Test Corner. End product to determine the acceptability of risk in conjunction to temperature testing without test corner as part of the power supply.

- End product to determine the acceptability of risk in conjunction to the Cleaning and Disinfection Methods as part of the power supply.
- End product to determine the acceptability of risk in conjunction to the Leakage of Liquids as part of the power supply.
- End product to determine the acceptability of risk in conjunction to the selection of components as it
  pertains to the intended use, essential performance, transport, storage conditions as part of the
  power supply.
- The secondary output circuits of Transformer (T1 / T2) are SELV and are not at hazardous energy levels.
- The power supply shall be properly bonded to the main protective earthing termination in the End Product.
- Fuse of Littelfuse, type 392 does not have adequate breaking capacity 100A; Overcurrent releases of
  adequate breaking capacity must be employed in the end product.