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Revised: 2014-08-20

UL TEST REPORT AND PROCEDURE

Standard: ANSI/AAMI ES 60601-1:2005 (Medical electrical equipment – Part 1:

General requirements for basic safety and essential performance) CSA C22.2 No. 60601-1:08 (Medical Electrical Equipment – Part 1: General Requirements for Basic Safety and Essential Performance)

Certification Type: Component Recognition

CCN: QQHM2, QQHM8 (Power Supplies, Medical and Dental)

Product: Switching Power Supply

Model: AHM250PSXX-ZZ (where XX is any number between 12-48

designating output voltage, ZZ can be blank, "A", "6", or "6A")

AHM250PSXXT-ZZ

Rating: Input Rated: 100-240 Vac, 50/60 Hz, 3 A

Output Rated: See Model Differences for details

Applicant Name and Address: XP POWER LLC

SUITE 150

1241 E DYER RD SANTA ANA CA 92705 UNITED STATES

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 Underwriters Laboratories Inc. ('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.

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 Underwriters Laboratories Inc. (UL) or any authorized licensee of UL.

Prepared by: Melissa DeGuia

ed by. UL LLC

Reviewed by: Timothy L. Gambrell

UL LLC

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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

Products covered are external power supplies intended to be used with Medical Electrical Equipment. Units are Class I. Earthing symbol may only be provided for Class I power supplies.

Model Differences

All models in the Model AHM250PSXX-ZZ series are identical with exception to the Mains Transformer, T2, and minor secondary components that allow for different output voltage ratings per the output voltage range noted below. See Table below for Model Ratings at 40°C:

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Model AHM250PS12: 10.1-13.5 Vdc, 17.5 A max. (250W max.) Model AHM250PS15: 13.6-17.0 Vdc, 16.67 A max. (250W max.) Model AHM250PS19: 17.1-21.0 Vdc, 13.16 A max. (250W max.) Model AHM250PS24: 21.1-26.0 Vdc, 10.41 A max. (250W max.) Model AHM250PS28: 26.1-31 Vdc, 8.93 A max. (250W max.) Model AHM250PS33: 31.1-33 Vdc, 7.58 A max. (250W max.) Model AHM250PS36: 33.1-42 Vdc, 6.94 A max. (250W max.) Model AHM250PS48: 42.1-54 Vdc, 5.21 A max. (250W max.)
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Model AHM250PSXXT-ZZ is identical to AHM250PSXX-ZZ except for model designation for marketing purposes.

See Enclosure - Miscellaneous for de-rated output values for higher ambient.

Suffix -ZZ when provided denotes the following:

- A Optional Retention Clamp provided
- 6 Optional C6 Type appliance inlet provided
- 6A Both Optional Retention Clamp and C6 Type appliance inlet provided.

Technical Considerations

Classification of installation and use: External Transportable

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Supply connection : Appliance coupler

- Accessories and detachable parts included in the evaluation: None
- Options included: None
- The product was investigated to the following additional standards:: ANSI/AAMI ES60601-1:2005/C1:2009 +AM1 (2012) (includes National Differences for USA); CAN/CSA-C22.2 No. 60601-1:08 +AM1 (2014) (includes National Differences for Canada), EN 60601-1:2006 + AM1 (2013)
- Scope of Power Supply evaluation defers the following clauses to the be determined as part of the end product: Clause 7.5 (Safety Signs), Clause 7.9 (Accompanying Documents), Clause 9 (ME Hazard), Clause 10 (Radiation), Clause 14 (PEMS), Clause 16 (ME Systems)
- Supply connection: OVC II
- The product is Classified only to the following hazards: Casualty, Fire, Shock
- The degree of protection against harmful ingress of water is: Ordinary
- The mode of operation is: Continuous
- Software is relied upon for meeting safety requirements related to mechanical, fire and shock: No
- The product is suitable for use in the presence of a flammable anesthetics mixture with air or oxygen or with nitrous oxide: No
- Unit also complied with spacing requirements of UL60601-1 (1st), CSA C22.2 No. 60601-1 (2nd), and IEC 60601-1 (2nd) for Basic for 250 Vac from Primary to Ground, Double/Reinforced for 250 Vac from Primary to Secondary, and Supplementary for 250 Vac from Secondary to Earth.

Risk Controls/Engineering Conditions of Acceptability

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

The component shall be installed in compliance with the Marking (clause 7) and Separation (clause 8) requirements of the end use application.

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■ The power supply was evaluated for use in 40°C ambient at Full Rated Output and 60% of the Rated Output in 60°C ambient. (See De-rating Curve, Enclosure 7-01 for details)

- Repeating leakage current testing should be considered in the end product application.
- This power supply was evaluated with Two MOPP between Primary and Secondary; One MOPP primary and Earth; and One MOPP between Secondary and Earth.
- This power supply has been evaluated as a continuous operation, ordinary equipment and has not been evaluated for use in the presence of a flammable anesthetic mixture with air, oxygen, or nitrous oxide. The output circuits have not been evaluated for direct patient connection (Type B, BF or CF).
- The end product should ensure that the requirements related to accompanying documents, clause 7.9. are met.
- The available voltage for the secondary outputs does not exceed 25 Vac or 60 Vdc, under normal and single fault conditions.
- The output connectors are not acceptable for field connections; they are only intended for connection to mating connectors of the end-use machine.
- The Electric Strength Test conducted on this power supply was based upon a maximum working voltage of: Primary-Earthed Dead Metal (Class I units): 430 Vpk, 240 Vrms; Primary-SEC: 591 Vpk, 279 Vrms.
- The following magnetic devices (e.g. transformers or inductor) are provided with an OBJY2 insulation system with the indicated rating greater than Class A (105°C): L1-L4 and T1 are Class B (130°C).
- Cleaning test to be considered as part of end product evaluation.
- The need for Marking Legibility Testing and repeating the Marking Durability to be considered as part
 of the end product installation.
- Power cord suitable for the application to be provided as part of the end product evaluation.

Additional Information

Marking label is representative of all models. The nameplate labels included in this report depict the draft artwork for the marking plate pending approval by National Certification Bodies and it will not be affixed to products prior to such approval.

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Earthing	<u></u>			
Serial Number or lot or batch identifier	Eight alpha numeric characters (A BB CC DDD where A = factory code; BB = year; CC=week; DDD = serial number)			
Date of Manufacturer	Provided as part of the serial number			
Special Instructions to UL Representative				
N/A				

Production-Line Testing Requirements					
Test Exemptions - The following models are exempt from the indicated test					
Model	Grounding Continuity	Dielectric Voltage Withstand	Patient Circuit Dielectric Voltage Withstand		
All Models	Test	Test	Exempt		
Solid-State Component Test Exemptions - The following solid-state components may be disconnected from the remainder of the circuitry during either Dielectric Voltage Withstand Test: N/A					
Sample and Test Specifics for Follow-Up Tests at UL					
The following tests shall be conducted in accordance with the Generic Inspection Instructions					
Model	Samples	Test	Test Details		
N/A					