



## **UK Declaration of Conformity**

For the following equipment :

Product Name: Medical Type Switching Power Supply

Model Designation:RPS-120S-x (x=12,15,24,27,48)

The designated product(s) is(are) in conformity with the relevant legislation:

The Restriction of the Use of Certain Hazardous Substances in Electrical and Electronic Equipment Regulations 2012: SI 2012 No. 3032

Medical Devices Regulations 2002 (SI 2002 No 618, as amended) (UK MDR 2002)

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3S EN 60601-1:2006+A1+A12+A2		TUV certificate No: TA50433767		
BS EN60601-1-2:2015+A1	:2021			
EMI (Electro-Magnetic Int				
Conducted emission	BS EN 55011:2016+A2:2021	Class B		
Radiated emission	BS EN 55011:2016+A2:2021Class A(for Class II ) ; Class B(for Class I )			
Harmonic current	BS ENIEC 61000-3-2:2019+A1:202	21		
Voltage flicker	BS EN61000-3-3:2013+A1:2019+A2:2021			
EMS (Electro-Magnetic S	usceptibility)			
ESD air	BS EN 61000-4-2:2009	Level 4	15KV	
ESD contact	BS EN 61000-4-2:2009	Level 4	8KV	
RF field susceptibility	BS EN IEC 61000-4-3:2020	Level 3	10V/m(80MHz-2.7GHz)	
RF field susceptibility	BS EN IEC 61000-4-3:2020	Table 9	9~28V/m (385MHz~5.78GHz	
EFT bursts	BS EN 61000-4-4:2012	Level 3	2KV/100KHz	
Surge susceptibility	BS EN 61000-4-5:2014+A1:2017	Level 4	2KV/Line-Line	
Surge susceptibility	BS EN 61000-4-5:2014+A1:2017	Level 4	4KV/Line-Earth	
Conducted susceptibility	BS EN 61000-4-6:2014	Level 3	10V	
Magnetic field immunity	BS EN 61000-4-8:2010	Level 4	30A/m	
Voltage dip, interruption	BS EN IEC 61000-4-11:2020 0% residu 70% residual v		ycles,0% residual voltage for 1 cycle es, 0% residual voltage for 250 cycle	

## Note:

The power supply is considered as a component that will be operated in combination with final equipment. Since EMC performance will be affected by the complete system, the final equipment manufacturers must re-qualify EMC Directive on the complete system again.

For guidance on how to perform these EMC tests, please refer to TDF (Technical Documentation File). The product under declaration is just a unit without medical function. Complete MDR should only be verified when it is used together with particular medical device(s).

This Declaration is effective from serial number SC3xxxxxx

## Person responsible for marking this declaration :

MEAN WELL Enterprises Co	o., Ltd.				
(Manufacturer Name)					
No.28, Wuquan 3rd Rd., Wugu Dist., New Taipei City 24891, Taiwan					
(Manufacturer Address)	$\wedge$ -				
Aries Jian/ Director, Group R&D :	Tries	Alex Tsai/ Director, Product Strategy Center :	( ) S		
(Name / Position)	(Signature)	(Name / Position)	(Signature)		
Taiwan	Dec. 27th, 2023	_			
(Place)	(Date)				

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