

Declaration of Conformity

For the following equipment :

Product Name: Medical Type Switching Power Supply Model Designation: RPxy-160z, RPTy-160a-C (x = S, D or T; y = G or blank; z = -5, -12, -15, -24, -48, A, B, C or D; a = A, B, C or D)

is herewith confirmed to comply with the requirements set out in the Council Directive, the following standards were applied :

RoHS Directive (2011/65/EU), (EU)2015/863

MDR Directive (EU) 2017/745

EN 60601-1:2006+A1+A12+A2

TUV certificate No : TA50147896

EN 60601-1-2:2015+A1:2021

EMI (Electro-Magnetic Interference)

Conducted emission / Radiated emission

		EN 55011:2016+A2:2021	Class B
Harmonic current		EN IEC 61000-3-2:2019	
Volta	ge flicker	EN 61000-3-3:2013+A1:2019	

EMS ((Electro-Magnetic Susce	ptibility)
	LICOLO Magnetio Ousee	publicy

ESD air	EN 61000-4-2:2009	Level 4	15KV				
ESD contact	EN 61000-4-2:2009	Level 4	8KV				
RF field susceptibility	EN IEC 61000-4-3:2020	Level 3	10V/m(80MHz-2.7GHz)				
RF field susceptibility	EN IEC 61000-4-3:2020	Table 9	9~28V/m (385MHz~5.78GHz)				
EFT bursts	EN 61000-4-4:2012	Level 3	2KV/100KHz				
Surge susceptibility	EN 61000-4-5:2014+A1:2017	Level 3	1KV/Line-Line				
Surge susceptibility	EN 61000-4-5:2014+A1:2017	Level 3	2KV/Line-Earth				
Conducted susceptibility	EN 61000-4-6:2014	Level 3	10V				
Magnetic field immunity	EN 61000-4-8:2010	Level 4	30A/m				
Voltage dip, interruption		oltage for 1 cycles, 709 ge for 250 cycles	% residual voltage for 25 cycles,				

Note:

A component power supply with load will be installed into final equipment which consists of an electronically shielded metal enclosure. Since EMC performance will be affected by the complete installation, the final equipment manufacturers must re-qualify EMC Directive on the complete installation again.

The EMC tests mentioned above are performed using a well defined metal plate to simulate said metal enclosure.

For guidance on how to perform these EMC tests, please refer to "EMI testing of component power supplies".(as available on http://www.meanwell.com)".

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 Enterprise	es Co., Ltd.		
(Manufacturer Name)			
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(Manufacturer Address)	\wedge -		(DE
Aries Jian/ Director, Group R	&D: MYLES	Alex Tsai/Director, Product Strategy Center	er:
(Name / Position)	(Signature)	(Name / Position)	(Signature)
Taiwan	Oct. 19th, 2023		
(Place)	(Date)		