



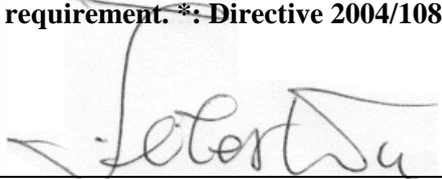
## VERIFICATION OF COMPLIANCE

**This Verification of Compliance is hereby issued to the product designated below.**

Product	<b>DC to DC Converter</b>
Model	<b>MSHU100 Series, MSDEU02-HI Series, MDHU100 Series, MDEU02-HI Series</b>
Brand	
Applicant	<b>MINMAX TECHNOLOGY CO., LTD.</b> No. 18, Sin-Sin Road, An-Ping Industrial District, Tainan, 702, Taiwan, R.O.C
Manufacturer	<b>MINMAX TECHNOLOGY CO., LTD.</b> No. 18, Sin-Sin Road, An-Ping Industrial District, Tainan, 702, Taiwan, R.O.C
Applicable Standard(s)	<b>EN 55011: 2009+A1: 2010</b> <b>EN 60601-1-2: 2007+AC: 2010</b> IEC 61000-4-2: 2008 IEC 61000-4-3: 2010 IEC 61000-4-4: 2012 IEC 61000-4-5: 2014 IEC 61000-4-6: 2013 IEC 61000-4-8: 2009
Reference No.	<b>T150817N12-E1</b>
Test Laboratory	<b>Compliance Certification Services Inc. (Tainan Lab.)</b> No.8,Jiucengling, Xinhua Dist., Tainan City 712, Taiwan (R.O.C.)

**This device has been tested and found to comply with the stated standard(s), which is (are) required by the Council Directive of 93/42/EEC, 2004/108/EC\* and 2014/30/EU, Amended by 92/31/EEC, 2006/95/EC & 98/13/EC. The test results are indicated in the test report and are applicable only to the tested sample identified in the report.**

**Note: The above EN basic standards are applied with latest version if customer has no special requirement. \*: Directive 2004/108/EC is repealed with effect from 20 April 2016.**

  
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*Jeter Wu / Assistant Manager*

*Tainan Lab.*

*Date: September 21, 2015*



# EC-Declaration of Conformity

The following equipment:

**DC to DC Converter**

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( Product Name )

**MSHU100 Series, MSDEU02-HI Series, MDHU100 Series, MDEU02-HI Series /**



( Model Designation / Brand Name )

**MINMAX TECHNOLOGY CO., LTD.**

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( Manufacturer Name )

**No. 18, Sin-Sin Road, An-Ping Industrial District, Tainan, 702, Taiwan, R.O.C**

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( Manufacturer Address )

is hereby confirmed to comply with the requirements set out in the Council Directive on the Approximation of the Laws of the Member States relating to Electromagnetic Compatibility Directive & Medical Devices (93/42/EEC, 2004/108/EC\* and 2014/30/EU, Amended by 92/31/EEC, 2006/95/EC & 98/13/EC (\*: Directive 2004/108/EC is repealed with effect from 20 April 2016)). For the evaluation regarding the Electromagnetic Compatibility & Medical (93/42/EEC, 2004/108/EC and 2014/30/EU, Amended by 92/31/EEC, 2006/95/EC & 98/13/EC) the following standards are applied:

**EN 55011: 2009+A1: 2010**

**EN 60601-1-2: 2007+AC: 2010**

IEC 61000-4-2: 2008

IEC 61000-4-3: 2010

IEC 61000-4-4: 2012

IEC 61000-4-5: 2014

IEC 61000-4-6: 2013

IEC 61000-4-8: 2009

The following manufacturer / importer or authorized representative established within the EUT is responsible for this declaration:

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( Company Name )

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( Company Address )

**Person responsible for making this declaration:**

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( Name, Surname )

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( Position / Title )

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( Place )

( Date )

( Legal Signature )