

# EU Declaration of Conformity



## We, the responsible manufacturer;

Company Name:	Mascot Electronics AS		
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## declare that this Declaration is issued under our sole responsibility and belongs to the following product(s):

Product and intended purpose:	Battery Charger for Li-Ion-, LiFePO <sub>4</sub> -, Li-Titanate, Lead-Acid or NiMH/NiCd Batteries
Brand(s):	and/or <b>MASCOT</b> (may also carry additional customer name, logo or trade mark)
Type(s)/Model(s)/UDI-DI:	<b>3546</b> (may also carry additional customer model name or part number)
Batch / Serial No./UDI-PI:	<b>all CE-marked products produced from the date indicated below</b> (for production date: see marking on the product)
Description:	<b>Input: max. 0.35 A 100-240 VAC 50-60 Hz, Class II</b> <b>Output: max. 28 W</b> (see product specific technical information) <b>1- to 16-cell for Lithium-Ion Batteries or</b> <b>1- to 16-cell for LiFePO<sub>4</sub> Batteries or</b> <b>1-to 20-cell for Li-Titanate Batteries or</b> <b>12V, 24V, 36V or 48V for Lead Acid Batteries or</b> <b>2- to 20-cell for NiMH/NiCd Batteries.</b>

## The product(s) described above are in conformity with the relevant European Union harmonisation legislation:

2014/35/EU	EU Directive - Safety of electrical equipment ("Low-Voltage Directive") (LVD) recast, repealing Directives 2006/95/EC & 73/23/EEC
2014/30/EU	EU Directive - Electromagnetic Compatibility (EMC) recast, repealing Directives 2004/108/EC & 89/336/EEC
(EU) 2017/745	EU Regulation - Medical Devices Regulation (MDR), Risk Class I Device Note : The product complies with the necessary EMC and safety standards but is not registered as a medical device according to the procedure in the MDR. amending Directive 2001/83/EC, Regulations (EC) 178/2002 & (EC) 1223/2009 and repealing Directives 90/385/EEC & 93/42/EEC
2009/125/EC	EU Directive - Energy Related Products, Ecodesign (ERP) recast, repealing Directive 2005/32/EC (EUP)
2015/863/EU Also 2011/65/EU	EU Directive - Restriction on use of Hazardous Substances in EEE ("RoHS3") recast, repealing Directives 2002/95/EC, 2008/35/EC & 2011/65/EU

## The following harmonised standards and technical specifications have been applied:

(International editions and comments indicated in brackets)

### Electrical Safety (to LVD- Directive):

EN 60335-1	EN 60335-1:2012/AC:2014/A11:2014/A13:2017 /A1:2019/A14:2019/A2:2019/A15:2021 (IEC 60335-1:2010 modified + /A1:2013 + /A2:2016, Edition 5.2)	Household and similar appliances-General req, Edition 5.2
EN 60335-2-29	EN 60335-2-29:2021 +/A1:2021 + /A11:2024 (IEC 60335-2-29:2016/A1:2019 , Edition 5.1)	Requirements for battery chargers, Edition 5.1

### Electrical Safety and Electromagnetic Compatibility (to MDR-Directive):

EN 60601-1	EN 60601-1:2006 + /AC:2010 +/A1:2013+/A2:2021 (IEC 60601-1:2005 + /A1:2012+/A2:2020)	Medical electrical equipment, Edition 3.2
EN 60601-1-2	EN 60601-1-2:2015 + A1:2021 (IEC 60601-1-2:2014 +A1:2020, Edition 4.1)	Medical equipment, EMC - Requirements and tests, Edition 4.1

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## Electromagnetic Compatibility (to EMC- Directive):

EN 61000-6-1	EN 61000-6-1:2019 (IEC 61000-6-1:2016, Edition 3.0)	Immunity-residential, comm. & light-industrial environment, Edition 3.0
EN 61000-6-3	EN 61000-6-3:2021 (IEC 61000-6-3:2020, Edition 3.0)	Emission-residential, comm. & light-industrial environment, Edition 3.0
EN 55014-1	EN IEC 55014-1:2021 (CISPR 14-1:2020, Edition 7.0)	Emission-household appliances, Edition 7.0
EN 55014-2	EN IEC 55014-2:2021 (CISPR 14-2:2020, Edition 3.0)	Immunity-household appliances, Edition 3.0
EN 55024	EN 55024:2010 (CISPR 24:2010, Edition 2.0) (also CISPR 24:2010 + /Corr.1:2011 + /A1:2015, Edition 2.1, but not yet an EN-norm)	Immunity-IT-Equipment, Edition 2.0
EN 55032	EN 55032:2015 + /A11:2020 (CISPR 32:2015, Edition 2.0)	Emission-Multimedia Equipment, Edition 2.0

## Additional Information:

Compliance with harmonised standards and technical specifications may have been verified by the manufacturer, by third party testing or by a Certification Body (NCB).

The products are considered Risk Class I devices according to EU Medical Devices Regulation and the U.K. The Medical Devices Regulations 2002.

The product(s) may be produced at production sites (for specific product: see "Made in"-marking on the product):

- Mascot Baltic OÜ, Taevakivi 15, EE-13619 Tallinn, ESTONIA
- MASCOT POWER SUPPLIES NINGBO CO LTD Building 9, No. 1188, Zhongguan Road, Zhenhai District NINGBO Zhejiang 315200 CN

The production sites are certified to standard EN 29001:2015 (ISO 9001:2015):

- Mascot Baltic OÜ: Metrosert, certificate ref. K-144
- Mascot Power Supplies (Ningbo) Co.,Ltd: DNV-GL, certificate ref. 179027-2015

The most recent issue of this Declaration is available at [www.mascot.com](http://www.mascot.com).

Fredrikstad, Norway

Place of issue

2025-11-24

Date of issue

Signed on behalf of Mascot Electronics AS

Fredrik Johansen, Compliance Manager

Name, function, signature