

RoHS EU Declaration of Conformity

No.: REG-001418

Product Name: Ambu® Neuroline Concentric

REF and Configuration
74025-30/NN
74025-45/NN
74030-35/NN
74038-45/NN
74050-45/NN
74075-65/NN

NN states number of electrodes in a box

Accessory: Ambu® Neuroline cable for Needle Electrodes: 1741, 1742, 1743

ROHS2 Category of the device
Category 8 – Medical devices

Identification: All products manufactured after issue date.

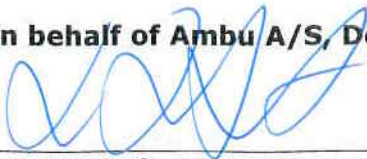
We the manufacturer hereby declare that this product is in conformity with the requirements in:

DIRECTIVE 2011/65/EU OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 8 June 2011
on the restriction of the use of certain hazardous substances in electrical and electronic equipment (RoHS recast) as it specified in Article 4.

This Declaration of Conformity is issued under the sole responsibility of Ambu A/S.

For and on behalf of Ambu A/S, Denmark

23 June 2014


Laila Strange Lundtoft, Senior Manager Corporate Regulatory Affairs

Current Issue: 1

First Issued: 23 June 2014