



Declaration of Conformity

Manufacturer:

ResMed Ltd 1 Elizabeth Macarthur Drive

Bella Vista NSW 2153 Australia **EU Representative:**

ResMed SAS
Parc Technologique de Lyon
292 Allée Jacques Monod
69791 Saint Priest Cedex
France

Notified Body:

TÜV SÜD Product Service GmbH Ridlerstraße 65 80339 München Germany

Product: Reusable Breathing Tube

Intended Use: The Reusable Air Tube is a non-invasive accessory used for conveying the air-flow

(with or without supplemental oxygen) generated by flow generators to a face mask or

nasal pillow-system for the treatment of CPAP or Bi-level therapy.

Classification: IIa according to Rule 2

GMDN: 37705 Breathing circuit, ventilator, reusable

Conformity Assessment Route: Annex II (excluding Section 4), 93/42/EEC

We herewith declare that the above mentioned products meet the transposition into national law of the provisions of Council Directive 93/42/EEC including the MDD amendment 2007/47/EC, for medical devices. Compliance to the MDD is applicable from the date listed below. All supporting documentation is retained at the premises of the manufacturer.

This declaration is issued under the sole responsibility of ResMed Ltd.

EC Certificate Number: G1 17 08 49861 149

Signed at Sydney, Australia on: 26-Jun-18

Johanna Wright

Director of Regulatory Affairs

ResMed Ltd

First issued: 06-Mar-09