



Product Service

EU Quality Management System Certificate (MDR)

Pursuant to Regulation (EU) 2017/745 on Medical Devices, Annex IX Chapters I and III (Class IIa and Class IIb Devices)

No. G10 083904 0009 Rev. 01

Manufacturer: ResMed Corp.

> 9001 Spectrum Center Blvd. San Diego CA 92123

USA

SRN Manufacturer - US-MF-000011805

Authorized ResMed SAS

Parc Technologique de Lyon, 292 Allée Jacques Monod, 69791 Representative:

Saint-Priest Cedex, FRANCE

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has established, documented and implemented a quality management system as described in Article 10 (9) of the Regulation (EU) 2017/745 on medical devices. Details on device categories covered by the quality management system are described on the following page(s). The Report referenced below summarises the result of the assessment and includes reference to relevant CS, harmonized standards and test reports. The conformity assessment has been carried out according to Annex IX Chapter I and III of this regulation with a positive result.

The quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis.

The certified quality management system is subject to periodical surveillance by TÜV SÜD Product Service GmbH. The surveillance assessment shall also include an assessment of the technical documentation for the device or devices concerned on the basis of further representative samples. All applicable requirements of the Testing, Certification, Validation and Verification Regulations TÜV SÜD Group have to be complied with.

For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:G10 083904 0009 Rev. 01

Report No.: 1482186793

G10 083904 0009 Rev. 00 **Preceding Certificate No.:**

Valid from: 2024-05-29 Valid until: 2027-09-12

Date of Initial Issuance: 2022-09-13

Christoph Dicks

Issue date: 2024-05-29 Head of Certification/Notified Body





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No. G10 083904 0009 Rev. 01

Classification: Class IIa

Device Group: Z1203019092 - VARIOUS INSTRUMENTS FOR ANAESTHESIA

AND PULMONARY VENTILATION SUPPORT - MEDICAL

DEVICE SOFTWARE

Z12040192 - GENERAL MEDICINE DIAGNOSIS AND

MONITORING INSTRUMENTS - MEDICAL DEVICE SOFTWARE

Intended Purpose:

The validity of this certificate depends on conditions and/or is limited to the following:

None

Revision History:

Rev. Dated Report 00 2022-09-13 72174274 01 2024-05-29 1482186793 Description

Supplemented: Device(s)/group of

device(s) added