



EC DECLARATION OF CONFORMITY

(Following the provisions of the medical devices directive 93/42/EEC, Annex II and of the directive 2011/65/EU)

We

Manufacturer Datex-Ohmeda, Inc. 3030 Ohmeda Drive PO Box 7550 Madison, WI 53707-7550 USA EU Authorized Representative GE Healthcare Finland Ov Kuortaneenkatu 2 Helsinki, Finland FI-00510

Declare under our sole responsibility that the class IIb device:

Avance Or s/5 Avance

Ref: 1009-9002-000

GMDN Code: 37710, UMDNS Code: 10-134

Classification rule (93/42/EC Annex IX): Class IIb. Rule 11

To which this declaration relates is in conformity with the requirements of the medical devices directive 93/42/EEC which apply to it. In addition, the product is in conformity with the requirements of the directive 2011/65/EU on the restriction of the use of certain hazardous substances in electrical and electronic equipment (as assessed by the manufacturer).

This conformity is based on the following elements:

- Information included in the documents: Technical Documentation of the product to which this declaration relates is maintained at Datex-Ohmeda, Inc.
- EC certificate: approval of full quality assurance system (Annex II of the medical devices directive 93/42 EEC) delivered by TÜV Rheinland LGA Products GmbH, Tillystraße 2, 90431 Nuremberg, Germany, Notified Body # 0197, Certificate N° HD 60098566 0001 valid until 9 June 2016.

List of harmonized standards applied for CE marking is located in the Technical Documentation file for this product.

Monica Merrison

Regulatory Affairs Manager

12 JAN 2015 Madison, USA, Day Month - Year

This EC declaration of conformity supersedes the previous declaration dated 1 July 2014.

Reference of the Declaration: DOC0929222