

EC DECLARATION OF CONFORMITY

Manufacturer: Accriva Diagnostics, Inc.
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San Diego, CA 92121 USA
Tel: (858) 263-2300 Fax: (858) 875-0603

European Authorized Representative: MDSS GmbH
Schiffgraben 41
Hannover, Germany

Product Group or Family: AVOXimeter Whole Blood Oximeter and CO-Oximeter
and Accessories
Product Name(s): See Attachment 1
Device Nomenclature: See Attachment 1
Classification: "Other" in vitro diagnostic medical device

We, Accriva Diagnostics, Inc., hereby declare that the products listed in Attachment 1 are in conformity with the In Vitro Diagnostic Medical Device Directive 98/79/EC (IVDD).

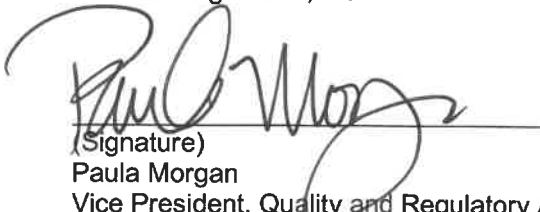
Conformity Assessment Procedure: IVDD, Annex III, excluding section 6

Notified Body Name: Not applicable, self-declared
Notified Body Identification Number: N/A
EC Certificate Number: N/A
EC Certificate Expiry: N/A

Additional Information: Applied Standards: See Attachment 2

Quality Management System: BSI, UK
EN ISO 13485:2016
Certificate Number: MD632084
Expiry Date: 2021-07-23

This declaration of conformity is issued under the sole responsibility of the manufacturer, Accriva Diagnostics, Inc.


(Signature)
Paula Morgan
Vice President, Quality and Regulatory Affairs
Accriva Diagnostics, Inc.

Place: San Diego, CA

Date: 5/7/2019

Attachment 1

Product Name	Catalogue No. / Ref No.	Device Nomenclature	Device Nomenclature Code	Device Nomenclature Term
Instruments, Disposables & Controls				
AVOXimeter 1000 E Cuvettes	C100B	EDMA	21 07 11 09	Other POC Hardware + acc + cons + software
AVOXimeter 4000 Cuvettes	QV8	EDMA	21 07 11 09	Other POC Hardware + acc + cons + software
AVOXimeter 1000E QC Optical Filters	E-QCYO	EDMA	21 07 11 09	Other POC Hardware + acc + cons + software
AVOXimeter 4000 QC Optical Filters	4-QCYO	EDMA	21 07 11 09	Other POC Hardware + acc + cons + software
Accessories				
Temperature Probe	4-TMPB	EDMA	21 07 11 09	Other POC Hardware + acc + cons + software

Attachment 2

Standard No.	Title
EN 13612:2002	Performance evaluation of in vitro diagnostic medical devices
EN 23640:2015	In Vitro Diagnostic Medical Devices - Evaluation Of Stability Of In Vitro Diagnostic Reagents
EN 13641:2002	Elimination or reduction of risk of infection related to in vitro diagnostic reagents
EN 13975:2003	Sampling procedures used for acceptance testing of in vitro diagnostic medical devices – Statistical aspects
EN ISO 13485: 2016	Medical Devices - Quality Management Systems - Requirements for Regulatory Purposes
EN 61010-1:2010	Safety requirements for electrical equipment for measurement, control , and laboratory use - Part 1: General requirements
EN 61010-2-101:2015	Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 2-101 :Particular requirements for in vitro diagnostic (IVD) medical equipment
EN 61010-2-010:2014	Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 2-010: Particular requirements for laboratory equipment for the heating of materials
EN ISO 14971:2012	Medical Devices - Application of Risk Management to Medical Devices
EN ISO 15193:2009	In Vitro Diagnostic Medical Devices - Measurement of Quantities in Samples of Biological Origin - Requirements for Content and Presentation of Reference Measurement Procedures
EN ISO 15194:2009	In Vitro Diagnostic Medical Devices - Measurement of Quantities in Samples of Biological Origin - Requirements for Certified Reference Materials and the Content of Supporting Documentation
EN ISO 15223-1:2016	Medical Devices - Symbols to be used with Medical Device Labels. Labelling and Information to be Supplied - Part 1: General Requirements
EN ISO 18113-1:2011	In Vitro diagnostic medical devices -Information supplied by the manufacturer (labelling)- Part 1: Terms, definitions and general requirements
EN ISO 18113-2:2011	In Vitro diagnostic medical devices -Information supplied by the manufacturer (labelling) –Part 2: In vitro diagnostic reagents for professional use
EN ISO 18113-3:2011	In Vitro diagnostic medical devices -Information supplied by the manufacturer(labelling)-Part 3: In vitro diagnostic instruments for professional use
EN 61326-1:2006	Electrical equipment for measurement, control and laboratory use -EMC requirements - Part 1: General requirements
EN 61326-2-6:2006	Electrical equipment for measurement, control, and laboratory use - EMC requirements - Part 2-6: Particular requirements - In vitro diagnostic (IVD) medical equipment