declaration of conformity



We, the manufacturer, declare in sole responsibility that the products mentioned below are in conformity with the respective regulations of the following directives.

Category	seca analytics
Product	115
Classification medical device	Class I
Conformity assessment procedure for medical devices	in accordance with Annex VII of the Medical Devices Directive 93/42/EEC

Directive:

93/42/EEC Directive concerning medical devices

Manufacturer: seca gmbh & co. kg

Hammer Steindamm 3-25 22089 Hamburg, Germany

Made in Germany

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This declaration of conformity is valid from the date of signature until a revised declaration of conformity is issued due to modification of the above-mentioned products.

Hamburg, 26 / 10 / 2016

Frederik Vogel

CEO Development & Manufacturing

declaration of conformity



Annex

Applied harmonised standards, national standards or other normative documents:

EN 62304

Medical device software - Software life-cycle processes