Declaration of Conformity



Dr. Müller Gerätebau GmbH Burgker Strasse 133 DE-01705 Freital

Herewith we declare on our own responsibility that the medical device meets all applicable requirements of the Directive 98/79/EC and applied standards and guidelines.

We do not guarantee the fulfilment of these norms and guidelines after unauthorized modification of the device.

Name of the product:

SUPER GL compact (incl. accessories)

Applied norms / guidelines

DIN EN 13485 Medical devices – Quality management systems – Requirements for

regulatory purposes

98/79/EG In vitro diagnostic medical devices

DIN EN 61010-1 Safety requirements for electrical equipment for measurement, control and

laboratory use - Part 1: General requirements

DIN EN 61010-2-081 Safety requirements for electrical equipment for measurement, control and

laboratory use - Part 2-081: Particular requirements for automatic and

semi-automatic laboratory equipment and other purposes

DIN EN 61010-2-101 Safety requirements for electrical equipment for measurement, control and

laboratory use - Part 2-101: Particular requirements for in vitro diagnostic

(IVD) medical equipment

DIN EN 61326-1 Electrical equipment for measurement, control and laboratory use – EMC-

requirements - Part 1: General requirements

DIN EN 61326-2-6 Electrical equipment for measurement, control and laboratory use – EMC-

requirements - Part 2-6: Particular requirements - In-vitro-diagnostic (IVD)

medical equipment

The CE mark was fixed to the device.

Freital, 20.03.2013

Ralf Günther General Manager

Stand: 03/2013 Rev. 01