

EU DECLARATION OF CONFORMITY

Fresenius Medical Care AG

Else-Kröner-Str. 1 61352 Bad Homburg – Germany SRN: DE-MF-000008193

declares under his sole responsibility that the product

Product Name: multiFiltratePRO SecuCas Ci-Ca HDF

Product Code: F00008117

GMN / Basic UDI-DI: 4039361-0000-0000-0117-NQ

Product Group: Acute dialysis tubing sets multifiltrate

EMDN Code: F020102

Intended Purpose: Channeling of blood and fluid in an extracorporeal

treatment

meets the applicable provisions as stated below and the relevant standards and common specifications as specified in the technical documentation.

Applicable regulation/s: European Medical Device Regulation 2017/745

Risk class: Class IIa according to Rule 2

(according to Annex VIII Medical Device

Regulation 2017/745)

Conformity assessment Annex IX (Chapters I and III) of the Regulation (EU)

procedure: 2017/745 (MDR)

Notified body: TÜV SÜD Product Service GmbH

Ridlerstraße 65

80339 München – Germany

Notified body no.: 0123

EU certificate: G10 077174 0005

Place, Date: Bad Homburg, 01-Dec-2023

Dr. Rüdiger Amberg

Product Center Responsible Person

Dr. Thomas Himstedt

Responsible person acc. MDR Art. 15

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