

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: | proHD Set S |
|---------------------|---|
| Product Code: | F00001621 |
| GMN / Basic UDI-DI: | 4039361-0000-0000-0084-PB |
| Product Group: | Connection and disconnection set for dialysis |
| EMDN Code: | V0599 |
| Intended Purpose: | Connection and disconnection for 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: (according to Annex VIII Medical Device Regulation 2017/745) | Class Is according to Rule 4 |
| Conformity assessment procedure: | Annex IX (Chapters I and III) of the Regulation (EU) 2017/745 (MDR) |
| Notified body: | TÜV SÜD Product Service GmbH Ridlerstraße 65 80339 München – Germany |
| Notified body no.: | 0123 |
| EU certificate: | G11 077174 0007 |
| Place, Date: | Bad Homburg, 01-Dec-2023 |

i.V.

Dr. Rüdiger Amberg Product Center Responsible Person

ppa

Dr. Thomas Himstedt Responsible person acc. MDR Art. 15

This Declaration of Conformity is valid until withdrawn or reissued due to significant product change, new product code or new/changed regulatory/legal requirement.