

## **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: PatientOnLine

Product Code: F989901

GMN / Basic UDI-DI: 4039361-0000-0000-0010-MU

Product Group: PatientOnLine (POL)

EMDN Code: *Z12099092* 

Intended Purpose: Acquire, calculate, model, save, display and transfer

parameters of dialysis treatments

Software-Version: 6.4

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 11

(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

Daniel Pelzl

Product Center Responsible Person

Dr. Thomas Himstedt

Responsible person acc. MDR Art. 15

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This Declaration of Conformity is valid until withdrawn or reissued due to significant product change, new product code or new/changed regulatory/legal requirement.

Property of Fresenius Medical Care



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## 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: PatientOnLine Multi-User

Product Code: F989951

GMN / Basic UDI-DI: 4039361-0000-0000-0010-MU

Product Group: PatientOnLine (POL)

EMDN Code: *Z12099092* 

Intended Purpose: Acquire, calculate, model, save, display and transfer

parameters of dialysis treatments

Software-Version: 6.4

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 11

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