

EU Declaration of Conformity

This declaration of conformity is issued under the sole responsibility of the manufacturer in accordance with Regulation 2017/745/EU Article 120 in its currently valid version. Article 120 (3a) case b) applies to the products. The requirements from Article 120 (3c) are met.

Product:

Oxygen Sensors

Basic UDI-DI:

4036616OOMCK

Type:

Honeywell	Viamed	Honeywell	Viamed	Honeywell	Viamed
OOM101	R15V	OOM105	T-7V	OOM201	R-23V
OOM102	R-22Vi	OOM106	R-45V	OOM202	R-49V
OOM102-A	R-22VA	OOM110	R-30V	OOM202-1	R-41V
OOM102-1	R-17Vi	OOM111	R-45V	OOM202-2	R-44V
OOM103	R-22MEDV	OOM112	R-43V	OOM202-2S	R-42V
OOM103-1	R-17V	OOM113	R-48V	OOM204	R-47V
OOM104	R-23Vi				

Classification:
(Annex VIII)

Class IIa

CE Mark:



Notified Body:

TÜV SÜD Product Service GmbH, Ridlerstr. 65, 80339 München, Germany

Conformity
Assessment Process:

Annex II, section 3 of Directive 93/42/EEC

Particular product
standard applied:

ISO 80601-2-55

Issued by:

Honeywell Healthcare Solutions GmbH
Eudamed-SRN: DE-MF-000004924
Alter Holzhafen 18
D-23966 Wismar
Germany

Valid from:

04.03.2024

Valid until:

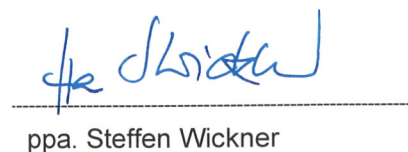
31.12.2028

Place, Date:

Wismar, 04.03.2024

Authorized
Signature:


ppa. Marko Sprössel


ppa. Steffen Wickner