MOTOmed_®

EC-Declaration of Conformity on Medical Devices

According to annex II of Medical Device Directive 93/42/EEC (2007/47/EG)

Manufacturer:

RECK-Technik GmbH & Co. KG Engineering Division Sector Reckstraße 1–5, 88422 Betzenweiler, GERMANY phone +49 7374 18-85, fax +49 7374 18-480 email: vigilance@motomed.com

Name of Product:

Basic UDI-DI: 4260193710017M MOTOmed viva2 MOTOmed viva2 stativ MOTOmed viva2 light MOTOmed viva2 light stativ MOTOmed viva2 Parkinson	REF 200.003 200.012 200.004 200.021 200.008	SN xxx-VM2-003xxx xxx-VM2-012xxx xxx-VM2-004xxx xxx-VM2-021xxx xxx-VM2-008xxx
Basic UDI-DI: 4260193710047T MOTOmed letto2 MOTOmed letto2 MOTOmed letto2 MOTOmed letto2	REF 279.003 279.024 279.008 279.016	SN xxx-LET2-003xxx xxx-LET2-024xxx xxx-LET2-008xxx xxx-LET2-016xxx
Basic UDI-DI: 4260193710067X MOTOmed gracile12 MOTOmed gracile12 ATAP	REF 594.003 594.003 + 599.000	SN xxx-KIG12-003xxx xxx-KIG12-003xxx

Product options: all (according to current pricelist)

Product classification: Ila (rule 9, Medical Device Directive 93/42/EEC)



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We hereby declare under our sole responsibility that the above products comply with the relevant provisions of the Medical Devices Directive 93/42/EEC (2007), Annex II, Section 3 and that we are solely responsible for issuing this declaration of conformity.

This Declaration of Conformity is valid until 02.08.2023 (registration no. of certificate: 50293-16-05) or until a revised declaration is made due to product modifications.

Following party was involved in the evaluation of the conformity: DEKRA Certification GmbH, Handwerkstrasse 15, 70565 Stuttgart, GERMANY Notified Body No. 0124

Jus Beh

Betzenweiler, 03.08.2018

Andreas Reck Executive Director

