

Annex to EC-Declaration of Conformity MOTOmed layson edition of 20 July 2020

Manufacturer: RECK-Technik GmbH & Co. KG Medical Devices

Reckstraße 1–5, 88422 Betzenweiler, GERMANY

Email: vigilance@motomed.com

Name of Product: MOTOmed layson edition

Basic UDI-DI: 4260193710037R

REF 261.xxx

We, as the manufacturer declare under our sole responsibility in reference to Regulation (EU) 2017/745 in conjunction with Regulation (EU) 2023/607 of 15 March 2023 that our Declaration of Conformity according to Annex II of Council Directive 93/42/EEC (2007/47/EC) for our medical devices MOTOmed layson edition of 20 July 2020 is valid until 31 December 2028 due to the extended transition period for class IIa medical devices.

In this context, we reference our manufacturer's declaration according to Regulation (EU) 2023/607 of 24 July 2023 (P03D56rev02).

RECK-Technik GmbH & Co. KG Betzenweiler, 25 July 2023

i.A. A. Fruil

i. A. Andreas Brauch, PRRC ra@motomed.com

