MOTOmed_®

Annex to EC-Declaration of Conformity MOTOmed gracile12 of 3 August 2018

Manufacturer:	RECK-Technik GmbH & Co. KG Medical Devices Reckstraße 1–5, 88422 Betzenweiler, GERMANY Email: vigilance@motomed.com
Name of Product:	MOTOmed gracile12 Basic UDI-DI: 4260193710067X REF 594.003, 594.003 + 599.000

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 gracile12 of 3 August 2018** 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. Fruch

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