

Warsaw, 25.05.2021

EU DECLARATION OF CONFORMITY MEDICAL DEVICE

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acting as Manufacturer registered in Eudamed, SRN number: PL-MF-000001583, according to Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017, herein declare that medical devices:

ROLLATOR: AT51013, AT51014, AT51015

The Basic UDI-DI: 59015714AT510133N, 59015714AT510143Q, 59015714AT510153S

have been classified as medical device class I, rule 1.

Intended purpose: rollators are designed for disable people to enable them to move. User rests on the rollator, makes a step forward and then pushes the rollator

We herein declare that the medical devices are in conformity with Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017. The assessment of conformity has been proceeded in compliance with the requirements of the above Regulation.

Applied standards:

PN-EN ISO 9001 (current edition)
PN-EN ISO 13485 (current edition)
PN-EN ISO 15223 (current edition)
PN-EN ISO 14971 (current edition)
PN-EN ISO 10993 (current edition)

The EU declaration of conformity is issued under the sole responsibility of the Manufacturer.

Andrzej Tarnkowski

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co-owner

Independent representation of the company
based on the Company Register

