

Warsaw, 25.05.2021

EU DECLARATION OF CONFORMITY MEDICAL DEVICE

*Irena Groniecka - Tarnkowska, Andrzej Tarnkowski „ANTAR” Spółka Jawna
ul. Zawiślańska 43
03-068 Warszawa*

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: AT51111, AT51112, AT51113

The Basic UDI-DI: not issued

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

Intended purpose: rollators are designed for disable people, to enable them to move

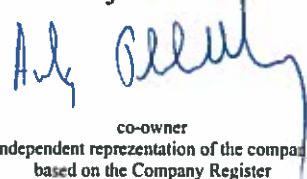
We herein declare that the medical devices are in conformity with Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017. The assessment of conformity has been proceeded in compliance with the requirements of the above Regulation.

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



co-owner
independent representation of the company
based on the Company Register

„ANTAR” Sp. Jawna
Irena Groniecka-Tarnkowska
Andrzej Tarnkowski
ul. Zawiślańska 43, 03-068 Warszawa
NIP: 524-21-23-915, REGON 012853911
Fax: 22 518 36 30, 22 518 36 31
Tel.: 22 518 36 00

