



REHASENSE

EU DECLARATION OF CONFORMITY

**REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE
COUNCIL OF 05 APRIL 2017 ON MEDICAL DEVICES (MDR)**

Manufacturer:

Rehasense Sp. z o.o.
Sulejowska 45G
97-300 Piotrków Trybunalski, Poland
SRN: PL-MF-000004772

Declare with sole responsibility that product (an ultra-light four-wheeled support for the disabled)

Product name: **ATHLON SL**

Catalog number: CRaab700cc, CRaab600cc, CRaab500cc, CRaab550cc,
CRaab60Ncc, CRaab70cc, CRaab60cc, CRaab55cc

(aa – colour, b- size, cc- accesories)

Intended use: The rollator has been designed as a tool to assist walking for people who have mobility problems.

Basic UDI-DI: 59074678ROL6U

meet requirements of the Regulation 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices and applicable international standards: ISO14971:2019; ISO 20417:2021; ISO 11199-2:2005;

Class of the medical device 1, in accordance with rule 1 (technical aid for disabled person). The product classification was carried out in accordance with the rules at Annex VIII of the Regulation 2017/745.

Manufacturer declares that follows conformity assessments procedure described in art. 52 para. 7 of the Regulation 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices after drawing up the technical documentation set out at Annexes II and III of the Regulation 2017/745.



02-08-2021/ Piotrków Trybunalski/ CEO Roger Spencer Dutton

Rehasense Sp. z o.o.
Prezes Zarządu

Roger Spencer Dutton