

EC DECLARATION of CONFORMITY

Regulation (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL on medical devices

We, MOBILEX A/S Registered place of business Grønlandsvej 5 8660 Skanderborg Denmark



SRN: DK-MF-000021885

Hereby declare under our sole responsibility as a legal manufacturer that the product specified on the product list below, meet the essential health and safety requirements and is in conformance with the provisions of the Regulation (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL on medical devices. The product specified on the product list below is "technical aid for the disabled", classified as Class I, medical device. The classification is based on the requirements of Rule 1 of annex VIII, of the Regulation (EU) 2017/745.

Intended purpose: Designed for individuals who require support standing to keep the balance.

The CE marking has been affixed on the product according to Annex V of the Regulation (EU) 2017/745.

PRODUCT LIS	T			
REF / item no.	311316	311325	311330	311340
UDI-DI	5740001403572	5740001403589	5740001403596	5740001403602
BASIC-UDI-DI	57400014GRABBAR9C			
REF / item no.	311350	311360	311375	311390
UDI-DI	5740001403619	5740001403626	5740001403633	5740001403640
BASIC-UDI-DI	57400014GRABBAR9C			

ACCESSORIES LIST

Item nr.	Accessories item nr.	
311316+25+30+40+50+60+75+90	NO	

Harmonized norms used during conformity estimation: PN-EN ISO 14971:2012, PN-EN 12182:2012, PN-EN 1041:2009;

Skanderborg, 2022-04-25, Thomas N. Christensen, Managing Director

