

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



Hereby declare under our sole responsibility as a legal manufacture 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 %echnical 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.

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

PRODUCT LIST

Telescopic ramp

TM-055; TM-150; TM-210; TM-300

Harmonized norms used during conformity estimation:

EN 12182:1999, PN-EN ISO 14971:2004, EN 1041:2001

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Skanderborg, 2020-03-11, Thomas N. Christensen, Managing Director

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