



EC Declaration of Conformity

for

Class 1 Medical Devices MDR 2017/745

Alu Rehab AS
Bedriftsveien 23
4353 Klepp Stasjon, Norway

SRN (single registration number) APP000023082

Netti Dynamic AdaptPro and accessories

GMDN 41620 Wheelchair, attendant / occupant driven, rear wheels, non-collapsible
Basic UDI-DI 704764NettiDynAdaptProET
Conforms to the Medical Device Regulation MDR 2017/745 EU

Intended purpose: Wheelchair for severely disabled people. The wheelchair enables function and mobility, varying seating positions and comfort during long time seating, allowing for more independence and improvements of quality of life.

Alu Rehab AS uses the procedures for the CE-labelling of Netti products according to Regulation MDR 2017/745 Annex IV

The following standards and the standards they request are used:

EN 12182:2012 Technical aids for disabled persons. General requirements and test methods.
EN 12183:2014 Manual wheelchairs – Requirements and test methods.
EN ISO 14971:2019 Medical devices – Application of risk management to medical devices.
EN 1021-2:2014 Assessment of the ignitability of upholstered furniture.

This declaration of conformity is issued under the sole responsibility of Alu Rehab AS.

We hereby declare that the medical device(s) specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices. This declaration is supported by the Quality System approval according to ISO 13485:2016 issued by TÜV SÜD.

All supporting documentation is retained at the premises of the manufacturer.

Klepp Stasjon 2022-02-24



Björn Carlzon
Managing Director



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