

# **EU Declaration of Conformity**

## with the Medical Device Regulation (MDR) 2017/745

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#### Wolturnus A/S, Skalhuse 31, 9240 Nibe, Denmark - DK-MF-000025274

Name and Address of Manufacturer – Single Registration Number (SRN)

hereby, under our sole responsibility declare, that the product			
Dalton			
Category Name			
with model names and reference			

Basic UDI-DI: 57138250097Y

Model	
Dalton	
Dalton F	
Dalton Light	
Dalton Low	
Dalton Tilt	
Dalton XXL	

is in conformity with the Medical Device Regulation (MDR) 2017/745 as class I medical devices based on Annex VIII and the following common specifications (CS) and standards

#### DS/EN ISO 14971:2019, ISO 13485:2016

Standards

The intended purpose: The Dalton wheelchair is intended to provide mobility to persons who are unable to walk or who have a mobility problem. It is designed for individual use, and it can be operated either by the patient or by another person. Dalton is a fixed-frame wheelchair with split, foldable footrests that are adjustable in both angle and height. Dalton can be used both indoors and outdoors.

This EU declaration of conformity was written according to Annex 4 in MDR, and all supporting documentation is retained at the premises of the manufacturer.

### Manufacturer

March 31, 2025 - Nibe

Peter Libak, CEO

Date and
Name and position of place of issue

Name and position of authorized person

Signature of authorized person

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