



## Mobile lifters

**Manufacturer**  
 V. Guldmann A/S  
 Graham Bells Vej 21-23A  
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 Phone +45 8741 3151  
 SRN: DK-MF-000003602

**Distributor/  
 Subsidiary**

Company

Address

Country

Phone

Hereby declare that the requirements specified in EU Regulation 2017/745 (MDR) regarding medical devices have been fulfilled in relation to the below listed device groups. The devices are also in conformity EU Directive RoHS 2011/65/EU and EU Directive REACH - 1907/2006/EC.

The declared medical devices comply where appropriate, with the following European standards  
 ISO 10535 Hoist for transfer of disabled persons – Requirements and test method. EN 12182 Assistive products for persons with disability - General requirements and test methods. IEC 60601-1 Medical electric equipment – Part 1 General requirements for basic safety and essential performance. EN 62304 Medical devices software – Software life-cycle process.

Guldmann A/S is certified as meeting quality management systems according with standards ISO 9001 and ISO 14001. Where appropriate, we comply with ISO 13485 Medical Devices – Quality Management Systems – Requirements for regulatory purposes and FDA 21 CFR part 820.

Device Group

**Mobile lifters**

GL5.1, GLS5.1  
 Class I, Rule 13

Basic UDI-DI

15707287mobileliftKV

Intended purpose

The Guldmann Mobile lifters are intended for lifting and transferring a person with disabilities.

**On behalf of V. Guldmann A/S**

Skejby, 2021.07.07

Place and date of issue

Ulrik Møller, Technical Manager