



EU DECLARATION OF CONFORMITY

Manufacturer	Etac A/S Parallelvej 3 DK-8751 Gedved Denmark
SRN	DK-MF-000017724
Statement	This EU declaration of conformity is issued under the sole responsibility of the manufacturer. The device(s) covered by present declaration is/are in conformity with EU Regulation 2017/745 on medical devices.
Basic UDI-DI	570799515TL
Device description	Patient slings
Intended purpose	The slings is an assistive device intended for alleviation of or compensation for a functional impairment due to an injury or disability. The device is designed for an individual lacking the ability to stand up and transfer oneself to/from a bed, the floor, a wheelchair, a chair, a toilet or similar, due to reduced mobility or physical strength.
Device name(s)	Molift RgoSling MediumBack Padded
	Molift RgoSling HighBack Padded
	Molift RgoSling MediumBack Net
	Molift RgoSling HighBack Net
	Molift RgoSling Toilet LowBack Padded
	Molift RgoSling Toilet HighBack Padded
	Molift RgoSling Comfort HighBack Net
	Molift RgoSling MediumBack Plus
	Molift EvoSling Hygiene
	Molift EvoSling LowBack Padded
	Molift EvoSling LowBack Net Padded
	Molift EvoSling Comfort MediumBack
	Molift EvoSling HighBack Net Padded
	Molift EvoSling HighBack Padded
	Molift RgoSling Shadow
	Molift EvoSling MediumBack Padded
,	Molift EvoSling MediumBack Net Padded
	Molift EvoSling Shadow
	Molift UnoSling HighBack
	Molift UnoSling Toilet
Brand	Molift
Risk class of the device	Class I, rule I
Place	Gedved, Denmark
Date of issue	24. August 2023
Name and function	Michael Bruun, Senior Vice President
	AM
	Signature, on behalf of Etac A/S