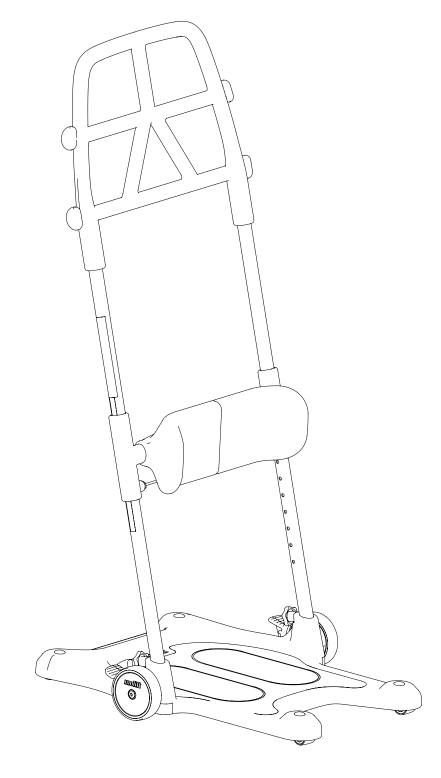
Molift Raiser Pro

- EN User manual
- SV Bruksanvisning
- NO Brukermanual
- DK Brugsvejledning
- FI Käyttöohje
- DE Gebrauchsanweisun
- NL Handleiding
- FR Manual d`utilisati
- IT Manual utente
- ES Manual de usuario

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General

CE declaration

The product, and its related accessories, described in this instruction for use (IFU), is in compliance with the regulation (EU) 2017/745 of 5. April 2017 – as a medical device, risk class I. The product is manufactured and tested according to EN 12182:2012 and selected parts of EN/ ISO 10535:2006. Any serious incident that occurs in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

Conditions for Use:

Transferring a person may pose an element of risk. Only trained personnel should use the equipment covered by this user manual.

Warranty notice:

Two-year warranty against defects in workmanship and materials of our products. Please refer to www.etac.com for terms and conditions.

Visit www.etac.com to download the latest version of our doccumentation.

Important

Read User Manual before use!

It is important to fully understand the content of the user manual before attempting to use the equipment. This User Manual contains important safety instructions and information regarding the use of the Molift Raiser Pro. For instructions/information regarding the accessories, see separate user manual.

In this manual the user is the person being transferred. The assistant is the person operating the Molift Raiser Pro.

Modifications and use of components made by other manufacturers:

We recommend only using Etac components and spare parts. Declaration of conformity is not valid and Etac is not responsible for warranty if any modifications are made to the Product. Etac shall not be liable for faults or accidents that can occur when using components made by other manufacturers.

Risk assessment

It is important that the Molift Raiser Pro has been tested with the individual user and for the intended transfer. Assess the risks and take notes. Evaluate the user's ability to use the Product and if needed use any suitable Molift Raiser Pro accessories. You as a carer are responsible for the safety of the user.



Only technical personnel are allowed to perform repairs.

Introduction

The person referred to as "the user" is the person who is standing on the product. "The carer" is the person who manoeuvres the product.

Intended Purpose

The sit-to-stand aid 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 themselves over shorter distances to another sitting position to/from a bed, a wheelchair, a chair, a toilet or similar due to reduced mobility or physical strength.

Intended User

The device can be used for children, adults and elderly people with a need for support to stand up and be transferred over shorter distances between sitting positions.

Intended Environment

For indoor use on horizontal surfaces in acute care, long-term care and home care.

Indications

For users who are in need of support to stand up and be transferred over shorter distances between sitting positions; e.g. people suffering from pain, reduced range of motion in their joints, stiffness and muscle weakness. Not an exhaustive list.

The user group for the device is based on individual health and mobility function and not on a specific diagnosis or age.

Contraindications

There are no known contraindications.

Precaution

The user should be able to follow instructions, grip and pull up their own body weight, maintain a good core stability and load at least one leg.

Training Requirements

Only carers who have read and understood the user manual are allowed to use the device.