



EU Declaration of Conformity

Manufacturer

Etac Immedia A/S

Parallelvej 3

DK-8751 Gedved

Denmark

www.etac.com

SRN

DK-MF-000019241

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

57080125001ME

Device description

Support

EMDN

Y1299

Intended purpose

The assistive device is 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 transfer or position themselves due to reduced mobility or physical strength.

Device name(s)

Sling

Risk class of the device

Class I, rule I

Harmonized/Established Standards

Separate list available upon request

Place

Gedved, Denmark

Date of issue

15. January 2024

Name and function

Michael Bruun, Senior Vice President

Signature, on behalf of Etac Immedia A/S