



EU DECLARATION OF CONFORMITY

Manufacturer Etac Immedia A/S
Parallevej 3
DK-8751 Gedved
Denmark

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 57080123001LY

Intended purpose Support feet during seated transfer.

Product / device name Immedia PediTurn Soft
Immedia PediTurn Hard

Brand Immedia

Risk class of the device Class I

Place Gedved, Denmark

Date of issue 1 June 2022

Name and function Michael Bruun, Senior Vice President

Signature, on behalf of Etac Immedia A/S