

## DECLARATION OF CONFORMITY (DOC)

Manufacturer/EU Representative:

Ferno-Washington, Inc. 70 Weil Way Wilmington, Ohio 45177-9371 U.S.A. 1.937.382.1451

EC **REP** 

FERNO S.r.I. via 8. Zallone, n. 26, 40066 Pieve di Cento, Bologna, Italy +39.051.6860028

Trade Name:

**FERNO®** 

SRN: N/A

Item/Catalogue #	Item Description	UDI-DI Number (GTIN)	Risk Class
0313065	78 PEDI PAC IMOBIL DEVC	Not implemented to date	I

Intended Use of Medical Device:

a non-inflatable extremity splint intended to immobilize limbs or an extremity

Conformity Assessment: Class I medical device, self-certification by manufacturer; no requirement for NB

In accordance with Council Directive 93/42/EEC (MDD), Ferno-Washington, Inc. ("Ferno") declares the above named product(s) comply with the applicable provisions of the Medical Device Directive (MDD).

Ferno maintains an ISO 13485:2016 certification for its Quality Management System ensuring all medical devices are manufactured and distributed using consistent quality standards and post market surveillance and vigilance is maintained.

Compliance to additional standards/directives is noted as applicable:

day of July, 2020 in Wilmington, Ohio, USA, under the sole This Declaration of Conformity is issued on this responsibility of the manufacturer.

FERNO-WASHINGTON, INC.

By:

**Dorothy Ramsey** 

Title:

VP, Global Legal & Regulatory