

Declaration of Conformity

Regulation (EU) 2017/745

Manufacturer:	Ferno Canada Inc.
Manufacturer Address:	2460 Tedlo Street, Mississauga, Ontario, L5A 3V3, Canada
EU Representative:	FERNO S.r.l., Via B. Zallone n.26 – 40066 Pieve di Cento (BO) - Italy
Unique registration number: (Art.31 (2))	(available when the managing system will be implemented by the European Commission)

The manufacturer declares under its own responsibility that the medical device (s):

Name of device(s):	Patient Shield HD EMS
Class of device(s): (Annex VIII)	I
Annex applied for the CE marking:	Annex II and Annex III
Intended use:	Medical Equipment
Base UDI-DI: (Art. 29 (1))	(available when the managing system will be implemented by the European Commission)

complies with the essential requirements listed in Annex I of the European regulation 2017/745 concerning Medical Devices

Ferno Canada Inc. is a Health Canada registered company (Establishment #1447)

Dated: May 1, 2020

Sophia Hatsisavvas Director of Regulatory Affairs

This document is compiled in accordance with Annex IV - EU declaration of conformity

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