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## **EU DECLARATION OF CONFORMITY**

MANUFACTURER				
Name of Company and Address		EUDAMED SRN / Application ID		
www.ferno.it	FERNO S.r.I Via B. Zallone n.26 40066 Pieve di Cento (BO) Italy +39 (051) 6860028	IT-MF-000031330 / APP000027477		
SWISS AUTHORIZED REPRESENTATIVE AND IMPORTER				
Name of Company and Address		Swiss Single Registration Number (CHRN)		
CH REP	FERNO S.r.I Pieve di Cento, succursale di Savosa Via Tesserete, 67 6942 Savosa, Switzerland +41 (41) 259 60 00	CHRN-AR-20002332 - AUTHORIZED REPRESENTATIVE CHRN-IM-20002288 - IMPORTER		

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

PRODUCT IDENTIFICATION			
Product Brand Name		Photo	
FERNO, XT Serie			
EMDN			
V08050103 - EMERGENCY AND TRAUMATOLOGY STRETCHERS			
Intended Purpose			
KIT XT PLUS-B is a kit designed	d for the immobilization and maintenance of the		
	e rapid extrication of patients traumatized (and not)		
during the rescue procedures.			
REF (Item / Catalog)	Item Description	GTIN (UDI-DI)	GMN (Basic UDI-DI)
KIT XT PLUS-B	KIT XT PLUS-B, maximum load 160 kg	08051380870471	805138087V08050103XTRR
RISK CLASS FOR MEDICAL DEVICES			
Device Classification	Common Specifications		
Class I Rule 1	Not applicable		

## according to:

HARMONIZED AND NON-HARMONIZED STANDARDS		
Item	Description	
EN ISO 10993-1:2020	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process (ISO 10993-1:2018)	
EN ISO 14971:2019/A11:2021	Medical devices - Application of risk management to medical devices (ISO 14971:2019)	
EN ISO 15223-1:2021	Medical devices - Symbols to be used with information to be supplied by the manufacturer - Part 1: General requirements (ISO 15223-1:2021)	
EN ISO 13485:2016+A11:2021	Medical devices - Quality management systems - Requirements for regulatory purposes (ISO 13485:2016)	
EN ISO 9001:2015	Quality management systems - Requirements (ISO 9001:2015)	

complies with the essential requirements listed in Annex I of the Regulation (EU) 2017/745 concerning Medical Devices.

Pieve di Cento, December 15th 2022

Signature Enrico Carletti - Managing Director

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

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