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▼ Via B. Zallone 26 – 40066 Pieve di Cento (BO)

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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 REPR	RESENTATIVE 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 PRO Serie		R	0	
EMDN		OFFINISK OFF		
V08050103 - EMERGENCY AND TRAUMATOLOGY STRETCHERS				
NATO NUMBER (NSN)				
6515150288871				
Intended Purpose				
XT PRO MILITARY is medical device designed for spinal immobilization and		Special Forces Equipment NATO Suppliers List NCAGE No. AL707		
extrication during vertical rescue in isolated environments.				
		https://eportal.nspa.nato.int/Codification/CageTool/home		
REF (Item / Catalog)	Item Description	GTIN (UDI-DI)	GMN (Basic UDI-DI)	
XT PRO MILITARY	KIT XT PRO MILITARY WITH HUMAN LIFT BRIDLES	08051380870457	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 1498	Personal fall protection equipment - Rescue loops	
EASA CS-27.865(a) and CS-	European Union Aviation Safety Agency – "External loads" and "Helicopter External Loads Personnel	
29.865(a) EASA CM-CS-005	Carrying Device System" issued 08 December 2014	
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.







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LOOPS (TRUCK LOOP 120 cm BLACK - 2034120B) - Notified Body 0123 TÜV SÜD Product Service GmbH (module D)		
Regulation (EU) 2016/425	European Regulation on personal protective equipment (PPE)	
ANSI/ASSE Z359.1	The Fall Protection Code (USA) - Anchorage connector 1 person max 5000 lbs - 22.2 kN	
EN 354: 2010	Personal fall protection equipment – Lanyards - Breaking load 45 kN	
EN 795/B:2012	Personal fall protection equipment - Anchor devices	
	EU Type Examination (module B) APAVE SUDEUROPE SAS Notified Body 0082	
EAC TP TC 019/2011	Eurasian Conformity mark (EAC, Russia-Belarus-Kazakhstan-Armenia-Kyrgyzstan) is a certification mark to indicate	
	products that conform to all technical regulations of the Eurasian Customs Union - Breaking load <b>45 kN</b>	
CARABINER (3C4650A) – Notified Body 0333 AFNOR Certification (module D)		
Regulation (EU) 2016/425	European Regulation on personal protective equipment (PPE)	
EN 362:2004	Personal protective equipment against falls from a height – Connectors	
	EU Type Examination (module B) DOLOMITICERT s.c.ar.l. Notified Body 2008	

Pieve di Cento, December 15th 2022

Signature Enrico Carletti - Managing Director

Mies Colott!

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

DNV ISO 1948

