

We, **X-CEN-TEK GmbH & Co. KG**, Westerburger Weg 30 - D-26203 Wardenburg, declare under our sole responsibility that the medical devices of the product group

PAX Rescue & Transport Systems Basic UDI-DI: 426074548 016018020019 MJ

complies with the relevant provisions of Regulation (EU) 2017/745 (MDR) on medical devices and, where applicable, other relevant Union legislation

The product group includes the following medical devices					
Commercial name	Article No.	Commercial name	Article No.		
PAX Rescue-BOA	118171908				

Intended use of the product group: Rescue and transport of injured and sick persons According to Annex VIII, Rule 1 MDR, all devices in the product group are class 1 medical devices and the applicable essential safety and performance requirements according to Annex I MDR are fulfilled. A conformity assessment procedure according to Annex II & III MDR has been carried out.

Applicable harmonized standards, national standards, or other regulatory documents:
EN ISO 13485:2016 – Medical devices – Quality management systems – Requirements for regulatory purposes
EN ISO 14971:2019 – Medical devices – Application of risk management to medical devices
EN 1865-1:2010 – Patient handling equipment used in road ambulances

This EU Declaration of Conformity is valid until **25th of May 2022**

Nils-Lasse Schneider

Wardenburg, the 25th of May 2021

Dr. Nils-Lasse Schneider PRRC according to Art.15 MDR

Manufacturers SRN: DE-MF-000009521

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Firma X-CEN-TEK GmbH & Co.KG – 26203 Wardenburg		QM-System nach EN 13485:2016		Seite 1 von 1	