

EU - Declaration of Conformity



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 Treatment Systems

Basis UDI-DI: 426074548 01625019 A6

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-Extremities-Tourniquet P-E-T – fluorescent yellow | 138103908 | PAX-Extremities-Tourniquet P-E-T – blue | 138103907 |
| PAX-Extremities-Tourniquet P-E-T – olive | 138103921 | PAX-Extremities-Tourniquet P-E-T – black | 138103903 |

Intended use of the product group: Treatment of injuries and illnesses - to tie off the blood flow in cases of severe, life-threatening bleeding of the upper and lower extremities that cannot be controlled in any other way or that, depending on the situation, represent the best alternative therapy

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 – Medical devices – Quality management systems – Requirements for regulatory purposes

EN ISO 14971 – Medical devices – Application of risk management to medical devices

This EU Declaration of Conformity is
valid until **25.05.2026**

Nils-Lasse Schneider

Wardenburg, 25.05.2025

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

Manufacturers SRN: DE-MF-000009521

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|---|--------------------------------|-----------------------------|-----------------------------|-------------------------------|
| Version 1.1 | BasDok erstellt: TC-05.04.2021 | Freigabe QMB: TC-10.08.2023 | Dok erstellt: TC | Freigabe VP: NLS - 25.05.2025 |
| Datei: PAX CE KE-EN TQ 05-25.docx | | | Anlage: 28.04.2021 | Stand: 25.05.2025 |
| Firma X-CEN-TEK GmbH & Co.KG – 26203 Wardenburg | | | QM-System nach EN ISO 13485 | Seite 1 von 1 |