

SAM® XT Extremity Tourniquet Declaration of Conformity

EUDOC-0008-C

Valid through: 2026-05-18

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EU DECLARATION OF CONFORMITY AS PER ANNEX IV OF THE REGULATION (EU) MDR 2017/745

Manufacturer:



SAM® Medical Products

12200 SW Tualatin Road, Suite 200, Tualatin, OR 97062, USA

Tel: + 1 (503) 639-5474 | Fax: +1 (503) 639-5425

quality@sammedical.com

Single Registration Number (SRN):

US-MF-000002589

EU Authorized Representative:



Emergo Europe

Westervoortsedijk 60, 6827 AT Arnhem, The Netherlands

Tel: +31 (0)70 345 8570 emergoeurope@ul.com

Single Registration Number (SRN):

NL-AR-000000116

Product Family Name

SAM® XT Extremity Tourniquet (SAM® XT)

Basic UDI-DI:

0822045XT01WH (see details in Table 1 attached)

Device(s) concerned:

This Declaration applies to all devices and variants included within the SAM® XT Extremity Tourniquet Product Family (see details in Table 1 attached).

Intended Purpose

The SAM® XT Extremity Tourniquet is intended to be applied around a limb to occlude arterial blood flow.

Risk Class per Annex VIII:

Class I (non-sterile) as per Rule 1

GMDN Code

58128 (Limb tourniquet, manual, single-use)

EMDN Code

V9003 (Tourniquets)

Notified Body:

Not applicable. Class I (non-sterile, non-measuring, non-reusable) devices are not reviewed by a Notified body.

Conformity Assessment Route: SAM Medical® Products utilizes Annex II and Annex III Technical Documentation (including PMS) for Class I EU medical devices and issues a Declaration of Conformity (self-certification).

Applicable CE Certificate(s):

Not applicable - Class I (non-sterile, non-measuring, non-reusable) devices are self-certified.

Standards and Common Specifications (CS):

This certificate further declares that the products covered herein also comply with the applicable requirements of relevant standards and Common Specifications specified in Table 2.

This declaration of conformity is issued under the sole responsibility of SAM® Medical Products. We hereby declare that the medical devices specified above meet the applicable provisions of the Medical Devices Regulation (EU) MDR 2017/745.

All supporting documentation is retained at the premises of the manufacturer.

Person authorized to sign on behalf of SAM®

Medical

Signature & date:

Name: Jeff Lipps

Position: Director RA/QA, SAM® Medical Products

Products: Place of Issue: 12200 SW Tualatin Road, Suite 200, Tualatin, OR 97062, USA

2023-06-26



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Table 1: Medical devices and variants included in the SAM® XT Extremity Tourniquet Product Family

Basic UDI-DI	GTIN	Product	Packaging Level	SKU
0822045XT01WH	10822045000206	SAM® XT Extremity Tourniquet – Tactical Black	Case	XT600-BK-EN
	00822045000209		Each	
	10822045000213	SAM® XT Extremity Tourniquet – Hi-Viz Orange	Case	XT600-OR-EN
	00822045000216		Each	
	10822045000220	SAM® XT Extremity Tourniquet – Hi-Viz Blue	Case	XT600-BL-EN
	00822045000223		Each	

Table 2: Standards and Common Specifications (CS) applied

Title	Year / Version				
Standard # Title Applied Standards					
Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk	2020				
management process					
18 Biological evaluation of medical devices — Part 18: Chemical characterization of medical					
device materials within a risk management process					
Medical devices - Quality management systems - Requirements for regulatory purposes	2016+A11:2021				
Medical devices - Application of Risk Management to Medical Devices	2019+A11:2021				
Medical devices - Symbols to be used with medical device labels, labelling and information to	2021				
be supplied - Part 1: General requirements					
Instrumentation for use in association with non-active surgical implants - General	2021				
requirements	See Footnote ¹				
Medical Devices - Information to be supplied by the manufacturer	2021				
Medical devices – Part 1: Application of usability engineering to medical devices	2015+A1:2020				
Other relevant standards					
Packaged Products for Parcel Delivery System Shipment 70 kg (150 lb) or Less	2018				
Environmental Engineering Considerations and Laboratory Tests	G				
Translation services — Requirements for translation services	2015+A1:2017				
EN ISO 17100 Translation services — Requirements for translation services — 2015+A1:2017 Common Specifications					
No common specifications relevant to the device family have been published in OJ at this time.					
	Applied Standards Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process Biological evaluation of medical devices — Part 18: Chemical characterization of medical device materials within a risk management process Medical devices - Quality management systems - Requirements for regulatory purposes Medical devices - Application of Risk Management to Medical Devices Medical devices - Symbols to be used with medical device labels, labelling and information to be supplied - Part 1: General requirements Instrumentation for use in association with non-active surgical implants - General requirements Medical Devices - Information to be supplied by the manufacturer Medical Devices - Part 1: Application of usability engineering to medical devices Other relevant standards Packaged Products for Parcel Delivery System Shipment 70 kg (150 lb) or Less Environmental Engineering Considerations and Laboratory Tests Translation services — Requirements for translation services Common Specifications				

¹Annex A was utilized for biocompatibility considerations.