

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.			
medical device	n of conformity is issued under the sole responsibility of SAM [®] Medical Products. We hereby declare that the s specified above meet the applicable provisions of the Medical Devices Regulation (EU) MDR 2017/745. documentation is retained at the premises of the manufacturer.			
Person authorized to sign on behalf of SAM [®] Medical Products:	Signature & date: Name: Jeff Lipps Position: Director RA/QA, SAM® Medical Products Place of Issue: 12200 SW Tualatin Road, Suite 200, Tualatin, OR 97062, USA			

SAM	SAM [®] XT Extremity Tourniquet Declaration of Conformity		
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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

Standard #	Title	Year / Version		
	Applied Standards			
EN ISO 10993-1	Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk	2020		
	management process			
EN ISO 10993-18	Biological evaluation of medical devices — Part 18: Chemical characterization of medical	2020		
	device materials within a risk management process			
EN ISO 13485	Medical devices - Quality management systems - Requirements for regulatory purposes	2016+A11:2021		
EN ISO 14971	Medical devices - Application of Risk Management to Medical Devices	2019+A11:2021		
EN ISO 15223-1	Medical devices - Symbols to be used with medical device labels, labelling and information to	2021		
	be supplied - Part 1: General requirements			
EN ISO 16061	Instrumentation for use in association with non-active surgical implants - General	2021		
	requirements	See Footnote ¹		
EN ISO 20417	Medical Devices - Information to be supplied by the manufacturer	2021		
EN 62366-1	Medical devices – Part 1: Application of usability engineering to medical devices	2015+A1:2020		
	Other relevant standards			
ISTA 3A	Packaged Products for Parcel Delivery System Shipment 70 kg (150 lb) or Less	2018		
MIL-STD-810G	Environmental Engineering Considerations and Laboratory Tests	G		
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.			
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¹Annex A was utilized for biocompatibility considerations.