

## **DECLARATION OF CONFORMITY**

Manufacturer	ABIGO Medical AB	
	Vapenvägen 1, SE-696 33 Askersund, Sweden	
Device classification and rule (Regulation EU 2017/745	IIb, Rule 4	
Annex VIII)		
SRN of the Manufacturer:	SE-MF-000000736	

Basic UDI-DI: 07392130Sorb	act2DU	
EMDN: M040416		

**Intended Purpose:** Cutimed Sorbact Ribbon® is intended for use in management of clean, contaminated, colonized or infected exuding wounds, such as surgical wounds, traumatic wounds, pressure ulcers, diabetic foot ulcers and leg ulcers.

Cutimed Sorbact Ribbon® can be used on superficial and deep wounds such as cavity wounds and fistulas. Cutimed Sorbact Ribbon® is also intended to treat fungal infections in skin folds (intertrigo).

Trade and Product Name	Catalogue number (REF)	
Cutimed Sorbact Ribbon®	Healthcare: 72166-16, 72167-10, 72167-11,	

Conformity assessment based on a quality management system and on assessment of technical documentation per **Annex IX Chapters I & III of Regulation (EU) 2017/745** has been performed by the following Notified Body:

Name and address	Notified Body id no	EC Certificate no and validity
Intertek Medical Notified Body AB, Torshamnsgatan 43,	2862	28620115063-02, 28 June 2026
Box 1103, SE-164 22 Kista, Sweden		

This declaration of conformity is issued under the sole responsibility of ABIGO Medical AB as the manufacturer. I hereby declare that the above-mentioned devices comply with **Regulation (EU) 2017/745** concerning medical devices.

Mölndal, date of issue

Marie Skoglund, Regulatory Affairs Director On behalf of Anna Arvidsson, Managing Director

Document ID: QAD-8452-v.1.0 Based on BLA-4260-v.11.0