

EU-DECLARATION OF CONFORMITY

Manufacturer	Wernli AG Eggasse 4 4852 Rothrist Switzerland
Single Registration Number CH:	CHRN-MF-20000003
EC Representative:	Wero Swiss Med Kft. Ipartelep utca 6 4220 Hajdúböszörmény Hungary
Single Registration Number HU:	HU-AR-000049642

We declare under our sole responsibility that

the medical device with reference no with Basic-UDI-DI (product code)	Wero Swiss Arm Sling see page 2 76300161A999812NF
Risk Class	class 1, unsterile according to Annex VIII, Rule 1

meets all the provisions of the Regulation (EU) 2017 /745 which apply to it.

Applied harmonised standards, national standards or other normative documents	EN ISO 10993-1:2021 EN ISO 15223-1:2021 EN ISO 14971:2019 / A11:2021
Conformity assessment procedure:	Regulation (EU) 2017/745 according to MDR Article 10 and MDR Article 52 (7) Annex I and Annex II
Place, date	Rothrist, 09.03.2026

Name and function

A handwritten signature in blue ink, appearing to read "K. Schönlé".

Katja Schönlé, Head of Sales

Reference Numbers:

Adults

A9998120	Wero Swiss Arm Sling Adult red
A9998121	Wero Swiss Arm Sling Adult blue
A9998122	Wero Swiss Arm Sling Adult green
A9998123	Wero Swiss Arm Sling Adult yellow
A9998124	Wero Swiss Arm Sling Adult black

Kids

A9998130	Wero Swiss Arm Sling Kids red
A9998131	Wero Swiss Arm Sling Kids blue
A9998132	Wero Swiss Arm Sling Kids green
A9998133	Wero Swiss Arm Sling Kids yellow
A9998134	Wero Swiss Arm Sling Kids black