

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 Forte see page 2 76300161A17PB
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 and applicable parts of the EN ISO 10993 Series EN ISO 15223-1:2021 EN ISO 14971:2019 / A11:2021
Conformity assessment procedures	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



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Felix Schoenle, CEO

Reference Numbers:

Length: 7m stretched

A1720060.070	Wero Swiss Forte 6cm x 7m brown
A1720080.070	Wero Swiss Forte 8cm x 7m brown
A1720100.070	Wero Swiss Forte 10cm x 7m brown
A1720120.070	Wero Swiss Forte 12cm x 7m brown
A1720150.070	Wero Swiss Forte 15cm x 7m brown