

## EU Declaration of Conformity according to the Medical Devices Regulation (EU) 2017/745 and the PPE Regulation (EU) 2016/425

The manufacturer:  
Ampri Handelsgesellschaft mbH  
Benzstr. 16  
21423 Winsen (Luhe)  
Germany  
SRN: DE-MF-000007622  
declares under its own responsibility that  
art. no.

**01251 S-XL MED COMFORT VITRIL (Vinyl-Nitril-Gemisch)**  
**Vinyl examination gloves**  
Basic-UDI-DI: 4044941001003RE

1) Complies with the requirements of regulation (EU) 2017/745 and the standards:

EN 455-1:2020	EN 455-2:2015	EN 455-3:2015	EN 455-4:2009
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This product is a Class 1 medical device according to the classification in Annex VIII.

and

2) complies with the requirements of regulation (EU) 2016/425 and the harmonized standards of

EN ISO 374-1:2016+A1:2018			
EN ISO 374-5:2016	EN 420:2003+A1:2009		

and the standards

EN ISO 374-4:2019	ISO 16604:2004		
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This product is a PPE of category III in accordance with attachment I of the regulation and is identical with the PPE which was subject to the EU type examination certificate no. 2777/15012-01/E05-01

issued by Satra, identification number 2777 and that is subject to the procedure according to Modul C2 of the regulation (EU) 2016/425 under the control of the notified body Satra (2777 identification number)

Technical documentation is available to prove this is accordance with the requirements.

Winsen, 06.10.2021

  
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ppa. Stephan Welzin  
Head of Quality Management & Operational Purchasing

This Declaration of Conformity is valid until 08.09.2025