

# DECLARATION OF CONFORMITY

MEDICAL DEVICE REGULATION (EU) 2017/745  
PERSONAL PROTECTIVE EQUIPMENT REGULATION (EU) 2016/425

Legal Manufacturer

**Semperit Investments Asia Pte. Ltd.**  
8 Jurong Town Hall Road,  
#29-03 to 06 The JTC Summit,  
Singapore 609434, Singapore  
sempermed@semperitgroup.com  
SRN: SG-MF-000001645

Authorized representative in the EU

**Semperit Technische Produkte Gesellschaft m.b.H.**  
Triester Bundesstraße 26, 2632 Wimpassing, Austria  
sempermed@semperitgroup.com  
SRN: AT-AR-000000735

This certificate is valid for the following product:

## Non-sterile examination and protective glove for single use

Classification: Class I according to MD Regulation (EU) 2017/745  
Category III according to PPE Regulation (EU) 2016/425

Basic UDI-DI: 9001570N\*F-035VB-N-3TY

## semperguard nitrile Xtra Lite

Sizes	X-Small	Small	Medium	Large	X-Large
Article codes	-	816780233	816780235	816780237	816780239
	-	3000001617	3000001618	3000001619	3000001620
	3000012020	3000012021	3000012022	3000012023	3000012024

**We hereby declare under sole responsibility that the CE marked product described above conforms to the requirements of the regulation for medical devices (EU) 2017/745.**

Declaration based on Annex IV. Classification according to rule 5, Annex VIII. The conformity assessment is based on Annex II.

Applied standards: ASTM D3578-19, ASTM D6319-19 or ASTM D5250-19, ASTM F1671/F1671M-13, EN ISO 20417:2021, EN 455-1:2020, EN 455-2:2015, EN 455-3:2015, EN 455-4:2009, EN 62366-1:2015 + AC:2016 + A1:2020, EN ISO 10993-1:2020, EN ISO 13485:2016 + AC:2018 + A11:2021, EN ISO 14971:2019 + A11:2021, EN ISO 15223-1:2021, ISO 10282:2014, ISO 2230:2002, MEDDEV 2.7/1 revision 4, Regulation (EC) 1907/2006

**We hereby declare under sole responsibility that the CE marked product described above conforms with the applicable provisions of Regulation (EU) 2016/425 on personal protective equipment and is identical to the personal protective equipment which is subject to EU Type Examination Certificate No. 2777/11464-01/E09-02 issued by:**

**SATRA Technology Europe Ltd, ID No. 2777**

**Bracetown Business Park, Clonee, Dublin D15 YN2P Ireland**

The products are subject to the procedure set out in Annex VII (Module C2) of Regulation (EU) 2016/425 under the supervision of

**SATRA Technology Europe Ltd, ID No. 2777**

**Bracetown Business Park, Clonee, Dublin D15 YN2P Ireland**

Applied standards: EN 420:2003+A1:2009, ISO 374-1:2016/AMD 1:2018, EN 374-2:2014, EN 16523-1:2015+A1:2018, EN 374-4:2013, EN ISO 374-5:2016, ISO 2859-1:1999 AMD 1:2011



Andreas Wöss  
Director



Released by: Christian Rohrbach

**This signed document is valid for all translations attached.**

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## IZJAVA O SKLADNOSTI

UREDBA O MEDICINSKIH PRIPOMOČKIH (EU) 2017/745/EGS  
UREDBA (EU) 2016/425 ZA OSEBNO VAROVALNO OPREMO

Proizvajalec

**Semperit Investments Asia Pte. Ltd.**  
8 Jurong Town Hall Road,  
#29-03 to 06 The JTC Summit,  
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SRN: SG-MF-000001645

Pooblaščen zastopnik EU

**Semperit Technische Produkte Gesellschaft m.b.H.**  
Triester Bundesstraße 26, 2632 Wimpassing, Austria  
sempermed@semperitgroup.com  
SRN: AT-AR-000000735

To potrдіlo velja za naslednje izdelke:

### Nesterilne zaščitne rokavice in rokavice za preglede za enkratno uporabo

Klasifikacija: Razred I v skladu z Uredbo o medicinskih pripomočkih (EU) 2017/745/EGS  
Kategorija III v skladu z Uredbo OVO (EU) 2016/425

Basic UDI-DI: 9001570N\*F-035VB-N-3TY

## semperguard nitrile Xtra Lite

Velikosti	X-Small	Small	Medium	Large	X-Large
Številke izdelkov	-	816780233	816780235	816780237	816780239
	-	3000001617	3000001618	3000001619	3000001620
	3000012020	3000012021	3000012022	3000012023	3000012024

**S to izključno odgovornostjo izjavljamo, da so izdelki z oznako CE v skladu z zahtevami Uredbe za medicinske pripomočke (EU) 2017/745.**

Izjava na podlagi Priloge IV. Razvrstitev v skladu s Prilogo VIII k Pravilniku 5. Ocenjevanje skladnosti temelji na Prilogi II.

Uporabljeni standardi: ASTM D3578-19, ASTM D6319-19 or ASTM D5250-19, ASTM F1671/F1671M-13, EN ISO 20417:2021, EN 455-1:2020, EN 455-2:2015, EN 455-3:2015, EN 455-4:2009, EN 62366-1:2015 + AC:2016 + A1:2020, EN ISO 10993-1:2020, EN ISO 13485:2016 + AC:2018 + A11:2021, EN ISO 14971:2019 + A11:2021, EN ISO 15223-1:2021, ISO 10282:2014, ISO 2230:2002, MEDDEV 2.7/1 revision 4, Regulation (EC) 1907/2006

**S to izključno odgovornostjo izjavljamo, da so zgoraj navedeni izdelki z oznako CE v skladu z bistvenimi zahtevami Uredbe (EU) 2016/425 za osebno varovalno opremo in so predmet certifikata ES o pregledu tipa št. 2777/11464-01/E09-02 izdano :**

**SATRA Technology Europe Ltd, ID No. 2777**

**Bracetown Business Park, Clonee, Dublin D15 YN2P Ireland**

Izdelki so predmet postopka v skladu s Prilogo VII (modul C2) uredbe pod nadzorom

**SATRA Technology Europe Ltd, ID No. 2777**

**Bracetown Business Park, Clonee, Dublin D15 YN2P Ireland**

Uporabljeni standardi: EN 420:2003+A1:2009, ISO 374-1:2016/AMD 1:2018, EN 374-2:2014, EN 16523-1:2015+A1:2018, EN 374-4:2013, EN ISO 374-5:2016, ISO 2859-1:1999 AMD 1:2011

Izdano dne:

Singapore, 2022-01-10

Veljavno do: 2024-01-09