

# EU DECLARATION OF CONFORMITY

The Manufacturer  
**ANSELL HEALTHCARE EUROPE N.V.**  
**RIVERSIDE BUSINESS PARK, BLOCK J**  
**BOULEVARD INTERNATIONAL 55**  
**B-1070 BRUSSELS**  
**BELGIUM**  
**WWW.ANSELL.COM**

declares under his sole responsibility, that the PPE described hereafter:

## **BioClean™ Non-Sterile Nitrile RABS/Isolator Glove CGL20NIT59**

*Products manufactured as of: [2024/03/19]*

**PPE to be used against category III risks**

EN388: 2016



**4102X**

EN ISO 374-1:2016  
Type A



**AJKNOTS**

EN ISO 374-5



**VIRUS**

EN 421



is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 388:2016 +A1:2018, EN ISO 374-1:2016, EN ISO 374-5:2016, EN 421:2010 and is identical to the PPE which is subject to the EU-Type examination (Module B, Annex V of the Regulation), under certificate number 032/2024/0182, issued by the Notified Body:

**CENTEXBEL (0493)**  
**TECHNOLOGIEPARK 70**  
**B-9052 ZWIJNAARDE**  
**BELGIUM**

and is subject to the procedure set out in Annex VIII (Module D) of the Regulation under the supervision of the Notified Body:

**CENTEXBEL (0493)**  
**TECHNOLOGIEPARK 70**  
**B-9052 ZWIJNAARDE**  
**BELGIUM**



**Guido Van Duren**  
**Director - Regulatory affairs**  
**Ansell**

**Place: Brussels**  
**Date: 2024/03/19**

# EU DECLARATION OF CONFORMITY

The Manufacturer  
**ANSELL HEALTHCARE EUROPE N.V.**  
**RIVERSIDE BUSINESS PARK, BLOCK J**  
**BOULEVARD INTERNATIONAL 55**  
**B-1070 BRUSSELS**  
**BELGIUM**  
**WWW.ANSELL.COM**

declares under his sole responsibility, that the PPE described hereafter:

## **BioClean™ Non-Sterile Nitrile RABS/Isolator Glove CGL20NIT59**

*Products manufactured as of: [2019/06/04] and till: [2024/03/18]*

**PPE to be used against category III risks**

EN388: 2016



**4101X**

EN 421



EN ISO 374-1:2016  
Type A



**AJKNOTS**

EN ISO 374-5



**VIRUS**

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 388:2016 +A1:2018, EN 421:2010, EN ISO 21420:2020, EN ISO 374-1:2016, EN ISO 374-5:2016 and is identical to the PPE which is subject to the EU-Type examination (Module B, Annex V of the Regulation), under certificate number 032/2023/0462, issued by the Notified Body:

**CENTEXBEL (0493)**  
**TECHNOLOGIEPARK 70**  
**B-9052 ZWIJNAARDE**  
**BELGIUM**

and is subject to the procedure set out in Annex VIII (Module D) of the Regulation under the supervision of the Notified Body:

**CENTEXBEL (0493)**  
**TECHNOLOGIEPARK 70**  
**B-9052 ZWIJNAARDE**  
**BELGIUM**



**Guido Van Duren**  
**Director - Regulatory affairs**  
**Ansell**

**Place: Brussels**  
**Date: 2023/07/25**

# EU DECLARATION OF CONFORMITY

The Manufacturer  
**NITRITEK (M) SDN BHD,**  
**NO.2, JALAN JURUNILAI U1/20,**  
**SEKSYEN U1, HICOM GLENMARIE**  
**INDUSTRIAL PARK,**  
**40150 SHAH ALAM,**  
**SELANGOR, MALAYSIA**

and authorized representative:  
**NITRITEK LTD**  
**UNIT 4, MINTON ENTERPRISE PARK**  
**OAKS DRIVE, NEWMARKET**  
**SUFFOLK, CB8 7YY, UK**

declare under their sole responsibility, that the PPE described hereafter:

## **BioClean Non-Sterile, Clean Nitrile 5 finger glove CGLxxNIT59**

*Products manufactured till: [2019/06/03]*

**PPE to be used against category III risks**

**EN 421**



**EN ISO 374-5**



**VIRUS**

**2016**

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 421:2010, EN ISO 374-5:2016 and is identical to the PPE which is subject to the EU-Type examination (Module B, Annex V of the Regulation), under certificate number 060/2019/1025, issued by the Notified Body:

**CENTEXBEL (0493)**  
**TECHNOLOGIEPARK 70**  
**B-9052 ZWIJNAARDE**  
**BELGIUM**

and is subject to the procedure set out in Annex VIII (Module D) of the Regulation under the supervision of the Notified Body:

**SGS FIMKO OY (0598)**  
**TAKOMOTIE 8,**  
**FI-00380 HELSINKI,**  
**FINLAND**

**Guido Van Duren**  
**Director - Regulatory affairs**  
**Ansell**

**Place: Brussels**  
**Date: 2019/06/03**