

## 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](http://WWW.ANSELL.COM)

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

### **BioClean-C™ Chemotherapy Protective Apron BCDA**

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



PARTIAL BODY  
PROTECTION ONLY



TYPE PB [4]



TYPE PB [6]

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN ISO 13688:2013+A1:2021, Partial Body Protection Only, EN 14605:2005 + A1:2009, EN 13034:2005 + A1:2009 and is identical to the PPE which is subject to the EU Type Examination; under certificate number 032/2023/0236 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**

Ulf Nystrom  
Sr Manager, Regulatory Affairs PPE Products

Place: Malmö  
Date: 2023/04/19