

# EU-OVERENSSTEMMELSESERKLÆRING

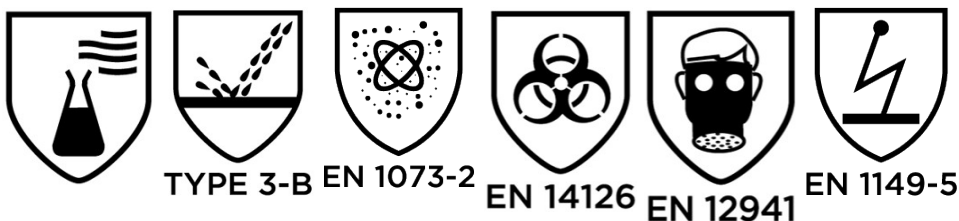
Producenten

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

erklærer under eneansvar, at PPE, der er beskrevet i det følgende:

## **AlphaTec 3000 PAPR Model 705**

**PPE, der skal anvendes mod kategori III-risici**



is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards , EN 14605:2005 + A1:2009, EN 1073-2:2002, EN 14126:2003, EN 12941:1998/A2:2008, EN 1149-5:2018 and is identical to the PPE which is subject to the EU Type Examination; under certificate number 0598/PPE/23/3439 issued by the Notified Body:

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

og er underlagt den procedure, der er fastlagt i Bilag VIII (Modul D) i forordningen under overvågning af testlaboratoriet:

**SGS FIMKO OY (0598)**  
**TAKOMOTIE 8,**  
**FI-00380 HELSINKI,**  
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**Ulf Nystrom**  
**Sr Manager, Regulatory Affairs PPE Products**

**Sted: Malmø**  
**Dato: 2024-10-11**