#### **EU DECLARATION OF CONFORMITY**

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

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

## ActivArmr® General Purpose Hi-Viz 97-012

Products manufactured as of: [04/11/2016] and till: [12/09/2018]

PPE to be used against category II risks



is in conformity with the provisions of Regulation (EU) 2016/425 and with the European harmonized standards EN 388:2016, EN 420:2003 + A1:2009 and is identical to the PPE which is subject to the EU-Type examination (Module B, Annex V of the Regulation), under certificate number 032/2016/1068, issued by the Notified Body:

CENTEXBEL (0493) TECHNOLOGIEPARK 70 B-9052 ZWIJNAARDE BELGIUM

and is subject to the procedure set out in Annex VI (Module C) of the Regulation.

Guido Van Duren

Director - Regulatory affairs

Ansell

Place: Brussels Date: 04/11/2016

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Guido Van Duren

Director - Regulatory affairs

Ansell

Place: Brussels Date: 13/09/2018

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and is subject to the procedure set out in Annex VI (Module C) of the Regulation.

Guido Van Duren

Director - Regulatory affairs

Ansell

Place: Brussels Date: 20/01/2014