

Ansell

Ansell Healthcare Europe N.V.

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EC DECLARATION OF PRODUCT CONFORMITY

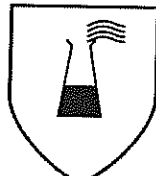
Category III

The manufacturer, established in the European Economic Community:

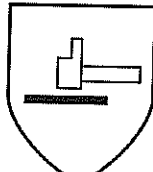
**ANSELL HEALTHCARE EUROPE N.V.
RIVERSIDE BUSINESS PARK, BLOCK J
BOULEVARD INTERNATIONAL 55
B-1070 BRUSSELS**

declares that the PPE described hereafter:

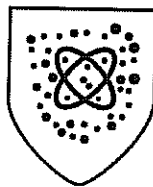
Bi-Colour™ 87-900



AKL



X120



is in conformity with the provisions of the Council Directive 89/686/EEC and with the European harmonised standards EN420:2003+A1:2009, EN374: 2003, EN388: 2003, EN421: 2010, and is identical to the PPE which is subject to the EC Type Examination certificate number 032/2015/0223 issued by the Notified Body:

**CENTEXBEL (0493)
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B-9052 ZWIJNAARDE**

is subject to the procedure set out in Article 11 point A of Directive 89/686/EEC under the supervision of the Notified Body

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**Monday, October 24, 2016
Alison Arnot-Bradshaw
Senior Director – EMEA/APAC Regulatory Affairs
Ansell**