

欧盟符合性声明

制造商

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

声明以下所述的个人防护设备由其全权负责：

ActivArmr® 80-409

适用产品起始日期 [2022/03/10]

用于防护category II风险的PPE



020

EN388: 2016



4121C

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 511:2006, EN 388:2016 +A1:2018, EN ISO 21420:2020 and is identical to the PPE which is subject to the EU-Type examination (Module B, Annex V of the Regulation), under certificate number 032/2022/0364, issued by the Notified Body:

CENTEXBEL (0493)
TECHNOLOGIEPARK 70
B-9052 ZWIJNAARDE
BELGIUM

符合法规（欧盟）2016/425的规定和欧洲统一标准EN 511:2006, EN 388:2016 +A1:2018, EN ISO 21420:2020，并与进行EC型式检验的个人防护设备（PPE）相同；根据公告机构颁发的证书编号

Guido Van Duren
Director - Regulatory affairs
Ansell

地点： Brussels
日期： 2022/03/10

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

声明以下所述的个人防护设备由其全权负责：

ActivArmr[®] 80-409

适用产品起始日期 [2019/01/07] and till: [2022/03/09]

用于防护category II风险的PPE



020



4121C

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 511:2006, 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 032/2019/0010, issued by the Notified Body:

CENTEXBEL (0493)
TECHNOLOGIEPARK 70
B-9052 ZWIJNAARDE
BELGIUM

符合法规（欧盟）2016/425的规定和欧洲统一标准EN 511:2006, EN 388:2016, EN 420:2003 + A1:2009，并与进行EC型式检验的个人防护设备（PPE）相同；根据公告机构颁发的证书编号



Guido Van Duren
Director - Regulatory affairs
Ansell

地点： Brussels
日期： 2019/01/07

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

声明以下所述的个人防护设备由其全权负责：

Powerflex 80-409

适用产品起始日期 [2016/12/07] and till: [2019/01/06]

用于防护category II风险的PPE



020



4121C

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 511:2006, 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 032/2016/1235, issued by the Notified Body:

CENTEXBEL (0493)
TECHNOLOGIEPARK 70
B-9052 ZWIJNAARDE
BELGIUM

符合法规（欧盟）2016/425的规定和欧洲统一标准EN 511:2006, EN 388:2016, EN 420:2003 + A1:2009，并与进行EC型式检验的个人防护设备（PPE）相同；根据公告机构颁发的证书编号

Guido Van Duren
Director - Regulatory affairs
Ansell

地点： Brussels
日期： 2016/12/07

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制造商

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

声明以下所述的个人防护设备由其全权负责：

Powerflex 80-409

适用期至： [2016/12/06]

用于防护category II风险的PPE



020



4121

is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards EN 511:2006, EN 388:2003, EN 420:2003 + A1:2009 and is identical to the PPE which is subject to the EC Type examination; under certificate number 032/2014/1604 issued by the Notified Body:

CENTEXBEL (0493)
TECHNOLOGIEPARK 70
B-9052 ZWIJNAARDE
BELGIUM

符合法规（欧盟）2016/425的规定和欧洲统一标准EN 511:2006, EN 388:2003, EN 420:2003 + A1:2009，并与进行EC型式检验的个人防护设备（PPE）相同；根据公告机构颁发的证书编号

Guido Van Duren
Director - Regulatory affairs
Ansell

地点： Brussels
日期： 2014/11/27