

欧盟符合性声明

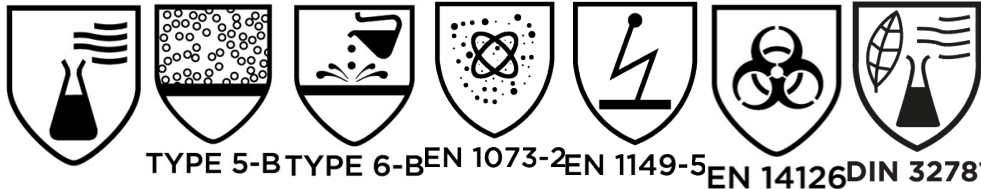
制造商

ANSELL HEALTHCARE EUROPE N.V.
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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:

AlphaTec® 2000 STANDARD Model 122

用于防护category III风险的PPE



is in conformity with the provisions of Regulation (EU) 2016/425 and with the standards , EN ISO 13982-1:2004 + A1:2010, EN 13034:2005 + A1:2009, EN 1073-2:2002, EN 1149-5:2018, EN 14126:2003, EN ISO 13688:2013, DIN 32781:2010 (with exceptions for puncture resistance being class 1 and resistance to ignition not tested) and is identical to the PPE which is subject to the EU-Type examination (Module B, Annex V of the Regulation), under certificate number 0598/PPE/23/3077, issued by the Notified Body:

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

并须遵守该规例附件VI (模块D) 所载的程序:

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

Ulf Nystrom
Sr Manager, Regulatory Affairs PPE Products

地点: Malmö
日期: 2024/09/26