



## Declaration of conformity Medical Devices, class I

Legal Manufacturer:	Abena A/S including Abena International A/S Egelund 35 DK - 6200 Aabenraa
Conformity assessment procedure	Annex VII of the Medical Devices Directive 93/42/EEC as amended by the Council Directive 2007/47/EEC. Module B and Module C2 of PPE Regulation 2016/425 for category III.
Classification and harmonized standards	<b>MDD Class I non sterile</b> EN 455 part 1,2,3,4 <b>PPE CAT III Type B,</b> EN 374-1:2016 EN 374-2:2015 EN 16523-1:2015 EN 374-4:2013 EN 374-5:2016 VIRUS EN 420: 2003+A1: 2009
Product	Classic Examination Latex Gloves, Powder-Free, Natural colour. Article no. 8817,4387,4388,4389,4381
<p>We, the legal manufacturer hereby declare that the above-mentioned product complies with the European Medical Device Directive 93/42/EEC as amended by the Council Directive 2007/47/EEC and its relevant transposition into all national laws of the member states into which we place the devices.</p> <p>EU Type Examination Module B and On-going conformity Module C2 performed by notified body 2777 SATRA Technology Europe Limited, Bracetown Business Park, Clonee, D15YN2P Ireland. Issued the EU Type Examination certificate 2777/10906-02/E01-01</p>	
Signed in Aabenraa	23.04.2019
Name and authority	Khalid Elamri Global Category Manager
Signature	

Date: 03-04-2019  
Version: 07  
Updated by: KHTA

Document responsible: Khaled Taj  
Approved by: Khaled Taj  
Doc. no. and name: 2.0.2S ENG Declaration of Conformity, Class I

Page 1 of 1