

DECLARATION OF CONFORMITY

This UK Declaration of Conformity is issued under the sole responsibility of Ansell Healthcare Europe NV.

Manufacturer Name/Address: Ansell Healthcare Europe NV
Boulevard International 55,
Brussels,
B-1070,
Belgium

UK Responsible Person: Ansell (U.K.) Limited
Building C,
Willerby Hill Business Park,
Willerby, Hull
HU10 6FE
United Kingdom

Risk Class: Class I

GMDN Code & Definition: 59821 - Ethylene vinyl acetate examination/treatment glove, non-powdered, non-sterile

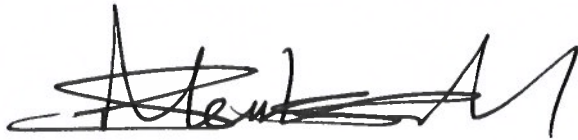
Product Name(s):

Product Name	Size	External Reference Code
ETHIPARAT	S	M2080
	M	M2079
	L	M2078
DISPOS-A-GLOVE® (pair)	S	P52370
DISPOS-A-GLOVE® (pair)	M	P52380
DISPOS-A-GLOVE® (pair)	L	P52390
DISPOS-A-GLOVE® (single)	S	S3027
DISPOS-A-GLOVE® (single)	M	S3047
DISPOS-A-GLOVE® (single)	L	S3067

Conformity Assessment Procedure: Part II – Annex VII

We hereby declare that the medical device(s) specified above meet the provision of The Medical Devices Regulations 2002 (UK MDR 2002).

Signed on behalf of the Manufacturer



Ansell Healthcare Europe NV
Riverside Business Park - Block J
Bid Internationalelaan 55
B-1070 Brussels
BELGIUM

Name: Samantha Marshall
Position: Director Regulatory Affairs Medical EMEA / APAC
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