

## **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:** 56286 - Nitrile examination/treatment glove, non-powdered, non-antimicrobial

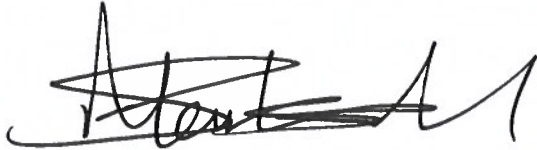
**Product Name(s):**

Product Name	Size	External Reference Code
MICROFLEX® 93-853	XS	93853060
	S	93853070
	M	93853080
	L	93853090
	XL	93853100
	XXL	93853110
	XXXL	93853120

**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



Name: Samantha Marshall  
Position: Director Regulatory Affairs Medical EMEA / APAC  
Date: 25 May 2023  
Revision: MED\UKMDR\MFX93853\001

**Ansell Healthcare Europe NV**  
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