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EU DECLARATION OF CONFORMITY

Manufacturer Name/Address: Ansell Healthcare Europe NV/SA

Riverside Business Park,

Block J. Boulevard International 55.

1070 Brussels,

Belgium

SRN Number: BE-MF-000000691

Risk Class: Class I

Intended Purpose: A non-sterile medical device intended as an

examination/treatment glove and a protective barrier when worn on the hands of healthcare providers, during patient examination, or for other sanitary purposes. The device is used mainly as a two-way barrier to protect both the patient and wearer against contaminants. This

is a single-use device.

EMDN Code and Description: T01020204 - Nitrile Examination/Treatment Glove

Basic UDI-DI: 5414566 MF93244 E2

Product Name(s):

Product Name	Size	Product Code	Market Regions
Microflex® 93-244	XS (5.5-6)	93244060	EMEA/NA/APAC
Microflex® 93-244	S (6.5-7)	93244070	EMEA/NA/APAC
Microflex® 93-244	M (7.5-8)	93244080	EMEA/NA/APAC
Microflex® 93-244	L (8.5-9)	93244090	EMEA/NA/APAC
Microflex® 93-244	XL (9.5-10)	93244100	EMEA/NA/APAC
Microflex® 93-244	XXI (10.5-11)	93244110	EMEA/NA/APAC

Conformity Assessment Procedure: Annex I & Annex II + Annex III

This EU Declaration of Conformity is issued under the sole responsibility of Ansell Healthcare Europe NV/SA.

We hereby declare that the medical device(s) specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices.

Ansell Healthcare Europe NV/SA

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Signed on behalf of Ansell Healthcare Europe NV

Name: Samantha Marshall

Position: Director Regulatory Affairs Medical EMEA / APAC

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