## EU DECLARATION OF CONFORMITY

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

SRN Number:
BE-MF-000000691

Risk Class:
Intended Purpose:

Class I
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: T010201 - Latex Examination / Treatment Gloves
Basic UDI-DI:
Product Name(s):

| Code | Product Description | Size | Region |
| :---: | :---: | :---: | :---: |
| 553301 | Micro-Touch $®$ Coated | XS | EMEA |
| 553302 | Micro-Touch $®$ Coated | S | EMEA |
| 553303 | Micro-Touch $®$ Coated | M | EMEA |
| 553304 | Micro-Touch $®$ Coated | L | EMEA |
| 553305 | Micro-Touch® Coated | XL | EMEA |

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.

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

Signed on behalf of the Manufacturer


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

## Name:

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Date of issue: 02 March 2023
Place of issue: Nuneaton, England
Version No: MED\MTCOAT\003

