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## **EU DECLARATION OF CONFORMITY**

This EU 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

**SRN Number:** BE-MF-000000691

Risk Class: Class IIa

Intended Purpose: A sterile medical device intended as a surgical glove and

a protective barrier when worn on the hands of healthcare providers at the surgical site. It is used mainly as a two-way barrier to protect patient and staff from

microorganisms. This is a single-use device.

**EMDN Code and Description:** T01010102 - Non-Powdered Latex Surgical Gloves

**Basic UDI DI:** 5414566 GLUG33005 Z2

## **Product Name(s):**

Product Name	Product Code	Size	Region(s)
Gammex® Latex Underglove	330050055	5.5	EMEA
Gammex® Latex Underglove	330050060	6.0	EMEA
Gammex® Latex Underglove	330050065	6.5	EMEA
Gammex® Latex Underglove	330050070	7.0	EMEA
Gammex® Latex Underglove	330050075	7.5	EMEA
Gammex® Latex Underglove	330050080	8.0	EMEA
Gammex® Latex Underglove	330050085	8.5	EMEA
Gammex® Latex Underglove	330050090	9.0	EMEA

Conformity Assessment Procedure: Annex IX.

CE Certificate No: MDR 763361.

Certified through the British Standards Institution, Notified Body Number 2797.

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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

Name: Samantha Marshall

Position: Director Regulatory Affairs Medical EMEA & APAC

Date: 02 March 2023
Place of Issue: Nuneaton, England
Revision: MED\MDR\GLUG\001