## **DECLARATION OF CONFORMITY EU - MD**

MANUFACTURER:	REFLEXX S.p.A.
	Via Passeri 2 - 46019 Viadana (MN) Italy e-mail: <u>info@reflexx.com</u> website: <u>www.reflexx.com</u>
Unique manufacturer registration number:	IT-MF-000021631

The undersigned REFLEXX S.p.A. with registered office in Via Passeri 2-46019 Viadana (MN) Italy, Share Capital € 1.200.000 (i.v.) VAT 02085450209 R.E.A. 223166, on their own and sole responsibility, as a manufacturer of the subject devices

## DECLARES

that the group of Medical Devices described below complies with the instructions of EU REGULATION 2017/745 (MDR) and complies with the general safety and performance requirements (Annex I) and with the applicable technical standards, reported in the technical file (EN 455 1,2,3 & 4).

The Technical File containing the relevant documentation is prepared in accordance with Annex II and is kept at the Manufacturer and made available to the Competent Authority. The Manufacturer has implemented and maintains a procedure for post-sales surveillance in accordance with Annex III.

Medical device (MD):	Family:DISPOSABLE EXAMINATION NON-SURGICAL GLOVESSub-family : non-sterile nitrile gloves CND T01020204Progressive number Attributed to the DM:
	art. N73: S/ 2543185 M/ 2543188 L/ 2543190 XL/ 2543191
	Code: reflexx N73 art. N73/S – art. N73/M – art. N73/L– art. N73/XL
Basic UDI-DI:	803289163GNPFEQ
Classification:	Class I not sterile - Rule 5 of Annex VIII of MDR

The company has certified its Quality Management System in compliance with EN ISO 9001: 2015 and EN ISO 13485:2016 (Certificate No. 1427.2023 and No. 0580.2024, issued by IMQ on 28.05.2024).

Place, Date

Signature Legal Representative

Viadana, 04/09/2024

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