

EU Declaration of Conformity

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| Manufacturer SRN (Single Registration Number) | GX Corporation Sdn Bhd (Specialty Plant) MY-MF-000035320 |
| Address | Plot 6497-A, Lorong Haji Abdul Manan, Batu 5 ¾ Sementa Jalan Kapar, 42100 Klang, Selangor, Malaysia |
| Authorized Representative Name SRN (Single Registration Number) | Emergo Europe B.V NL-AR-000000116 |
| Authorized Representative Address | Westervoortsedijk 60, 6827 AT Arnhem, The Netherlands |
| Notify Body Name | SATRA Technology Europe Ltd (Number: 2777) |
| Notify Body Address | Bracetown Business Park Clone, D15 YN2P, Ireland |
| PPE Certification Number | 2777/12322-02/E00-00 |
| Name of the Device | Powder Free Nitrile Non-Sterile Examination Glove |
| Trade Name | Velo® Soft |
| Reference Number | 5044001, 5044002, 5044003, 5044004, 5044005 |
| Glove Sizes Available | XS, S, M, L and XL |
| Classification | Class I, Rule 5 for MDR Regulation (EU) 2017/745 Category III for Annex II of the PPE Regulation (EU) 2016/425 |
| Intended Use | Intended to wear on the hands for medical examination to provide a barrier against cross contamination and other contaminants. |
| Conformity assessment route | GX Corporation Sdn Bhd (Specialty Plant) uses the following procedures for the CE-labelling of our products: Class I: EC conformity declaration according to Annex IV + Annex VIII of MDR Regulation (EU) 2017/745 Category III: The PPE is subject to the conformity assessment procedure based on internal procedure control plus supervised product checks at random intervals (Module C2) under the surveillance of the notified body. |

This declaration of conformity is issued under the sole responsibility of GX Corporation Sdn Bhd (Specialty Plant). We hereby declare that the device specified above meets the provision of the MDR Regulation (EU) 2017/745 for medical devices, PPE Regulation (EU) 2016/425 for personal protective equipment and relevant harmonized standard as follows:



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| EN ISO 13485:2016 | Medical Devices – Quality Management Systems – Requirements for Regulatory Purposes |
| EN ISO 13485:2016 MDSAP | Medical Devices – Quality Management Systems – Requirements for Regulatory Purposes |
| ISO 9001:2015 | Quality Management Systems Requirements |
| EN 455-1:2020 | Medical Gloves for Single Use – Part 1: Requirements and Testing for Freedom from Holes |
| EN455-2:2015 | Medical Gloves for Single Use – Part 2: Requirements and Testing for Physical Properties |
| EN 455-3:2015 | Medical Gloves for Single Use – Part 3: Requirements and Testing for Biological Evaluation |
| EN 455-4:2009 | Medical Gloves for Single Use – Part 4: Requirements and Testing for Shelf-Life Determination |
| ASTM D6319-19 | Standard Specification for Nitrile Examination Gloves for Medical Application |
| EN 420:2003+A1:2009 | Protective gloves – General Requirements and Test Methods |
| EN 374-1 | Protective Gloves against Dangerous Chemicals and Micro-organisms – Part 1: Terminology and Performance Requirements |
| EN 374-2 | Protective Gloves against Dangerous Chemicals and Micro-organisms – Part 2: Determination of Resistance to Penetration |
| EN 374-4 | Protective Gloves against Dangerous Chemicals and Micro-organisms – Part 4: Determination of Resistance to Degradation by Chemicals |
| EN 374-5 | Protective Gloves against Dangerous Chemicals and Micro-organisms – Part 5: Terminology and Performance Requirements for Micro-organisms Risks |
| EN 16523-1 | Determination of Material Resistance to Permeation by Chemicals – Part 1: Permeation by Liquid Chemical under Conditions of Continuous Contact |

This declaration is supported by the Quality System approval to ISO 13485 issued by TUV SUD Product Service GmbH. All supporting documentation is retained at the premises of the manufacturer.

7th July 2023

Date of Issue

Klang, Selangor

Place of Issue

Mr. Hew Yew Fook

Printed Name

Signature

Chief Operating Officer

Title

GX Corporation Sdn Bhd (Specialty Plant)

Company

