

## EU Declaration of Conformity

|                                          |                                                            |                                                                 |
|------------------------------------------|------------------------------------------------------------|-----------------------------------------------------------------|
| <b>Manufacturer:</b>                     | <b>Granberg AS</b><br>Bjoavegen 1442, NO-5584 Bjoa, Norway | <b>E-mail:</b> post@granberg.no<br><b>Phone:</b> +47 53 775 300 |
| <b>Single Registration number (SRN):</b> | NO-MF-000000207                                            | <b>Doc Nr:</b> F/MD/008                                         |

**REF. 114.621**

Disposable Examination and Protective Gloves Granberg®

The item is in conformity with (EU) 2017/745 Medical Device Regulation (MDR) as:

|                                    |                                                                                                                                                                                                                                                                                                                        |
|------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <b>Name</b>                        | Disposable Examination and Protective Nitrile Gloves Granberg®                                                                                                                                                                                                                                                         |
| <b>Description</b>                 | non-sterile, powder-free, accelerators free                                                                                                                                                                                                                                                                            |
| <b>Risk Classification</b>         | Class I according to Rule 1 and 5 of Annex VIII of Regulation MDR (EU) 2017/745                                                                                                                                                                                                                                        |
| <b>Applied normative standards</b> | EN 455-1:2020+A2:2024, EN 455-2:2024, EN 455-3:2023, EN 455-4:2009, ISO 10993-1:2018, ISO 10993-5:2009, ISO 10993-10:2021, ISO 10993-23:2021, EN ISO 14971:2019+A11:2021, ISO 15223-1:2021, EN ISO 20417:2021                                                                                                          |
| <b>Intended Use</b>                | Powder-free examination and protective disposable nitrile gloves intended for use in the medical field to protect patients and users from cross-contamination. These gloves are also intended to protect against certain chemicals and microorganisms, and radioactive contamination, where hand protection is needed. |
| <b>Basic UDI-DI</b>                | 702377MT114NNSUV                                                                                                                                                                                                                                                                                                       |
| <b>Available sizes</b>             | XS - XL                                                                                                                                                                                                                                                                                                                |
| <b>EMDN Code</b>                   | T01020201                                                                                                                                                                                                                                                                                                              |

The technical documentation for assessing the conformity of the medical devices with MDR has been developed in accordance with Annexes II and III of MDR.

The product described above is also meeting requirements of **Category III** Personal Protective Equipment (PPE) and complies with the Regulation (EU) 2016/425 of the European Parliament and of the Council of 9 March 2016 on Personal Protective Equipment and European harmonized Standards EN ISO 21420:2020, EN ISO 374-1:2016+A1:2018, as **Type B**, EN ISO 374-5:2016 including **virus** protection and EN 421:2010 protection against radioactive contamination.

The product is identical to the PPE, which is the subject of the EU Type Examination (Module B) certificate of conformity no. **2777/10013-06/E01-01** issued by Notified Body:

**Satra Technology Europe Limited, (NB No. 2777)**, Bracetown Business Park, Clonee, Dublin D15YN2P, Republic of Ireland.

Furthermore, the product is subject to the Conformity to type, based on quality assurance of the production process (Module D) of the Regulation 2016/425 under the surveillance of the Notified Body:

**BSI Group The Netherlands B.V. (NB No. 2797)**, Say Building, John M. Keynesplein 9, 1066 EP Amsterdam, Netherlands.

This Declaration of Conformity is issued under the sole responsibility of the manufacturer – Granberg AS.

Signed for Granberg AS:



Ole Marthon Granberg

Managing Director

Place and date of issue:

Bjoa, 06.02.2025