

# **DECLARATION OF CONFORMITY**

Replaces version dated:

22.06.2022

Valid until the issue of next version of this document.

We,

OneMed Group Oy,

SRN FI-MF-000000642,

Domicile Helsinki, Business ID 2039640-1.

Metsäläntie 20,

FI-00320 Helsinki, Finland,

declare under our sole responsibility that following products are in conformity with Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices as well as Regulation (EU) 2016/425 of the European Parliament and of the Council of 9 March 2016 on personal protective equipment.

#### Classification:

Class I according to Annex VIII of the Regulation (EU) 2017/745 on medical devices,
Category III according to the Regulation (EU) 2016/425 on personal protective equipment

### Intended Purpose (As per Regulation (EU) 2017/745 on medical devices):

Nitrile examination gloves are for medical purposes intended to be worn on care giver's hands to prevent cross-contamination between care giver and a patient. The gloves are non-sterile. The gloves are for single use. The device may be used by health care professional or lay person.

## List of devices under Basic UDI-DI 6438129B0001GU:

REF	Product name
210262 XS	SELEFA® Nitrile Medical Exam Gloves, SENSE LIGHT, white, size XS
210262 S	SELEFA® Nitrile Medical Exam Gloves, SENSE LIGHT, white, size S
210262 M	SELEFA® Nitrile Medical Exam Gloves, SENSE LIGHT, white, size M
210262 L	SELEFA® Nitrile Medical Exam Gloves, SENSE LIGHT, white, size L
210262 XL	SELEFA® Nitrile Medical Exam Gloves, SENSE LIGHT, white, size XL

#### These items are in conformity with following European standards and common specifications<sup>1</sup>:

Medical Devices – Quality management systems – Requirements for
regulatory purposes
Medical Devices – Application of risk management to medical devices
Information supplied by the manufacturer with medical devices
Medical Devices – Symbols to be used with medical device labels, labelling
and information to be supplied. Part 1: General requirements
Protective glove. General requirements and test method
Medical gloves for single use. Part 1 – Requirements and testing for
freedom from holes
Medical gloves for single use. Part 2 – Requirements and testing for
physical properties
Medical gloves for single use. Part 3 – Requirements and testing for
biological evaluation

T-084, Rev 5

DoC\_OneMed Group\_MDR class I\_PPER cat III\_Nitrile exam gloves\_SENSE LIGHT\_SELEFA







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EN 455-4	Medical gloves for single use. Part 4 – Requirements and testing for shelf
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life determination

EN ISO 374-1 Protective gloves against chemicals and micro-organism. Part 1 -

Terminology and performance requirements for chemical risks

EN ISO 374-2 Protective gloves against dangerous chemicals and micro-organism – Part

2: Determination of resistance to penetration

EN 374-4 Protective gloves against dangerous chemicals and micro-organism — Part

4: Determination of resistance to degradation by chemicals

EN ISO 374-5 Protective gloves against dangerous chemicals and micro-organisms. Part 5

- Terminology and performance requirements for micro-organisms risks

<sup>1</sup>Relevant standards, latest applied revisions and common specifications are listed in *T-079 Review of regulations and standards* 

The notified body 2777, SATRA Technology Europe Limited Bracetown, Business Park, Clonee, D15YN2P, Republic of Ireland performed the EU type-examination (Module B) and issued the EU type-examination certificate 2777/11581-02/E09-01. The product(s) is (are) subject to the conformity assessment procedure Module C2 under surveillance of the same notified body. Type C glove according to EN 374-1:2016.

Place and date of issue

Helsinki 13.06.2023

Name and signature of the authorized person

Hilleriikka Vaho

Regulatory Manager

OneMed Group Oy





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