



# EU-TYPE EXAMINATION CERTIFICATE



The following model of Personal Protective Equipment has been subjected to an EU-type examination in accordance with the module B of the PPE regulation (2016/425) and has been shown to satisfy to essential health and safety requirements.

**Certificate N°**            **0075/4324/162/05/21/0869**

Issued by CTC, Notified Body N°0075, to the following model of personal protective equipment :

**Manufacturer :**            **SINOAOO MEDICAL CORP**  
Room 1310, building 1, window of Green Space Business Plaza,  
Jinyuan Road 2, Dongshan street, Jiangning District, Nanjing  
China

### Description

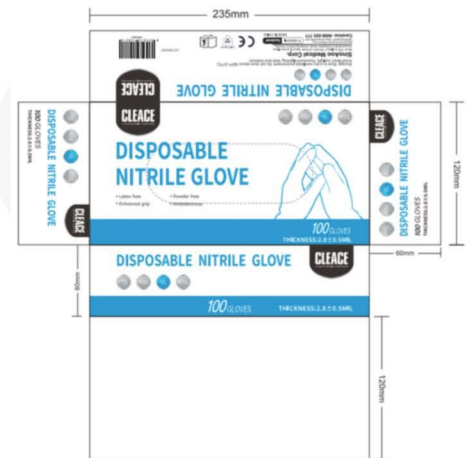
**PPE Type :**                    **protective glove against microorganisms risks**

**Product reference :**        **SANG002**

**Glove description :**        **Disposable nitrile gloves**

**Available sizes :**            **7/S 8/M 9/L 10/XL**

**Pictures :**



**Reference standard :**

**Levels of performance / class of protection**

« X » indicates that the glove has not been submitted to the test or the test method appears not to be suitable for the glove design or material.  
« 0 » indicates that the glove falls below the minimum performance level for the individual hazard.

**EN ISO 21420:2020**

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**EN ISO 374-5:2016**

**MICRO-ORGANISMS**

**This glove is intended to activity that does not involve a deliberate intention to work with or use a biological agent but may result in the workers' being exposed to a biological agent.**

At the date of the certificate, the product is in compliance with Annex XVII of REACH regulation (n° 1907/2006 and revisions)

Full description of the PPE, reference rules verified in the context of the EU-type examination and information given on the product are detailed in the manufacturer's technical file and the Instruction for Use index 01 dated from MAY 2021

NOTA : Any modification to new items of the personal protective equipment object of this EU type approval certificate or any modification of the information contained in the manufacturer technical file which served for the deliverance of the EU type approval certificate (change of address, change of company status) should be brought to the attention of the notified body in accordance with Annex V §7.2 of Regulation 2016/425. Any marking on the PPE which is not concerned by the Regulation (UE) 2016/425, is not covered by this certificate.

**Issued in Lyon by**  
**Didier GUISSADO**  
Certification and Quality Manager

**Date of first issue :** 18 May 2021  
**End of validity date :** 18 May 2026



In application of the Regulation 2016/425 of the European parliament and the Council of 9th March 2016 related to Personal Protective Equipment and repealing the Directive 89/686/EEC.



Accreditation n° 5-0594  
Scope available on:  
www.cofrac.fr

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Loi 78-654 du 22.06.1978 - Siret 77564972600160 - Code NAF 9412Z - TVA FR 88775649726