Issued to:



SATRA customer number: P21130 Notified Body: 2777

Anhui Intco Medical Products Co., Ltd No. 6, Haitang South Road Suixi Wuhu Modern Industrial Park Suixi County Huaibei City Anhui Province China

EU Type-Examination Certificate

Certificate number: 2777/22223-01/E00-00

This EU Type-Examination Certificate covers the following product group(s) supported by testing to the relevant standards/technical specifications and examination of the technical file documentation. It has been issued Under Module B of Regulation 2016/425 on personal protective equipment. This product group has been shown to satisfy the applicable essential health and safety requirements as a Category III product.

Description: Product reference:

NGX/S/TEXXXXX Disposable Nitrile Gloves, Non-sterile

Blue	Green
NGX/S/TE10014-NGX/S/TE10018	NGX/S/TE10084- NGX/S/TE10088
NGX/S/TE20014-NGX/S/TE20018	NGX/S/TE20084- NGX/S/TE20088
NGX/S/TE10F14-NGX/S/TE10F18	NGX/S/TE10F84- NGX/S/TE10F88
NGX/S/TE20F14-NGX/S/TE20F18	NGX/S/TE20F84- NGX/S/TE20F88
NGX/S/TE10P14-NGX/S/TE10P18	NGX/S/TE10P84- NGX/S/TE10P88
NGX/S/TE20P14-NGX/S/TE20P18	NGX/S/TE20P84- NGX/S/TE20P88
Violet	Black
Violet NGX/S/TE10034- NGX/S/TE10038	Black NGX/S/TE10044- NGX/S/TE10048
	= 10.011
NGX/S/TE10034- NGX/S/TE10038	NGX/S/TE10044- NGX/S/TE10048
NGX/S/TE10034- NGX/S/TE10038 NGX/S/TE20034- NGX/S/TE20038	NGX/S/TE10044- NGX/S/TE10048 NGX/S/TE20044- NGX/S/TE20048
NGX/S/TE10034- NGX/S/TE10038 NGX/S/TE20034- NGX/S/TE20038 NGX/S/TE10F34- NGX/S/TE10F38	NGX/S/TE10044- NGX/S/TE10048 NGX/S/TE20044- NGX/S/TE20048 NGX/S/TE10F44- NGX/S/TE10F48

Sizes:

Classification:

7-11(S-XXL)	EN 150 374-1:2016+A1:2016/19pe B		Levei
	(K) Sodium hydroxide 40%	6	6

		Degradation %
K) Sodium hydroxide 40%	6	-30.1
P) Hydrogen peroxide 30%	3	43.1
T) Formaldehyde 37%	6	-17.5
O) Ammonium hydroxide 25%	2	-3.58

EN ISO 374-5:2016	Level
Protection against Bacterial and Fungi	Pass
Protection against Viruses	Pass

Standards/Technical specifications applied:

EN ISO 21420:2020; EN ISO 374-1:2016+A1:2018; EN ISO 374-5:2016

Technical reports/Approval documents:

SATRA: CHT0316811/2129, CHM0317791/2132/EN/A, CHM0317783/2132/LC, CHT0329910/2216, CHT0335653/2232,

CHT0335234/2231, CHT03333975/2228

SGS: QDHL2206007432HM, QDHL2206008535HM

TUV: 7191266610-EEC21-WBH-CR2

Signed on behalf of SATRA:

Geoff Graham

Date first issued: 29/08/2022 Date of issue: 29/08/2022 Expiry date: 29/08/2027

EN ISO 374-4:2019

TERMS AND CONDITIONS

The following conditions apply in addition to SATRA's standard terms and conditions of business and those given in the current certification agreement. This certificate has been issued in accordance with Annex V (Module B) of the applicable legislation (see note 11).

Please note:

- 1. Where the product is classified as category III then CE or UKCA Marking of production is reliant on current compliance with module C2 or Module D of the applicable legislation (See note 11). (Except that specifically produced to fit an individual user).
- 2. Full details of the scope of the certification and product(s) certified are contained within the manufacturer's technical documentation.
- 3. Where a translation of this certificate exists, the English language version shall be considered as the authoritative text.
- 4. Certification is limited to production undertaken at the sites listed in the manufacturer's technical documentation.
- 5. Ongoing manufactured product shall be consistent with the product(s) certified and listed on this certificate and an EU declaration of product conformity shall be made available in accordance with the applicable legislation (See note 11)
- 6. The Manufacturer shall inform SATRA of any changes to the certified product or technical documentation.
- 7. Where results obtained during type testing are within the budget of uncertainty when compared to the pass requirement, classification or performance level, then it is the responsibility of the manufacturer to ensure that the factory production control and manufacturing tolerances are such that the product placed on the market meets with the stated requirements, classifications or performance levels.
- 8. This certificate shall be kept together with the relevant technical documentation in a safe place by the client named on this certificate. Production of this certificate and other documentation may be required by a representative of the EC member state, or UK government.
- 9. This certificate relates only to the condition of the testable items at the time of the certification procedure and is subject to the expiry date shown.
- 10. SATRA reserves the right to withdraw this certificate if it is found that a condition of manufacture, design, materials or packaging have been changed and therefore no longer comply with the requirements of the applicable legislation (See note 11).
- 11. These terms and conditions shall apply to the requirements set out in Regulation (EU) 2016/425 of the European Parliament and of the council of 9th March 2016 on personal protective equipment or to UK legislation relating to UKCA Marking as defined within the issued certificate.



Document Number: INTCO-CE-DC-NBR-001

Version: A/2

EU DECLARATION OF CONFORMITY

Manufacturer

Authorized Representative

Name: Anhui Intco Medical Products

Name: Lotus NL B.V.

Co.,Ltd.

Address: No. 6, Haitang South Road, Suixi Wuhu Modern Industrial Park, Suixi County, Huaibei City, Anhui Province

Address: Koningin Julianaplein 10, le Verd, 2595AA, The Hague, Netherlands

Declares that the MDR described hereafter

Product name and model:

Disposable Nitrile Exam Gloves

EMDN code: T01020204

Model:XS/S/M/L/XL/XXL

Product Code: See the following annex I

Basic UDI-DI: 697306977NitrileFR

SRN: CN-MF-000002356

This declaration of conformity is issued under the sole responsibility of the manufacturer:

Anhui Intco Medical Products Co., Ltd.

Conformity Assessment Route: Annex II and Annex III according to EU 2017/745.

Applicable Standard:

EN ISO 13485:2016; EN 14971:2019; EN 1041:2008; EN 15223-1:2016

EN 455-1:2020; EN 455-2:2015; EN 455-3:2015; EN 455-4:2009;

ISO 10993-1:2018; ISO 10993-10:2010. ISO 10993-11:2017

Meet the provisions of the Council Regulation EU 2017/745 and Annex I which apply to them,

The medical device has been assigned to Class I, based on rule 1 & rule 5 of Annex VIII

Chapter III of the Regulation EU 2017/745 MDR. It bears the mark



We agree to develop, implement and maintain a documented post-production monitoring process



Document Number : INTCO-CE-DC-NBR-001

Version: A/2

Annex 1 product code

Color	Product Code
Blue	NGV/B/H/PEM10013-NGV/B/H/PEM10018
	NGV/B/H/PEM20013-NGV/B/H/PEM20018
	SNV/B/H/PE10013-SNV/B/H/PE10018
	SNV/B/H/PE20013-SNV/B/H/PE20018
White	NGV/B/H/PEM10023-NGV/B/H/PEM10028
	NGV/B/H/PEM20023-NGV/B/H/PEM20028
	SNV/B/H/PE10023-SNV/B/H/PE10028
	SNV/B/H/PE20023-SNV/B/H/PE20028
	NGV/B/H/PEM10033-NGV/B/H/PEM10038
Voilet	NGV/B/H/PEM20033-NGV/B/H/PEM20038
	SNV/B/H/PE10033-SNV/B/H/PE10038
	SNV/B/H/PE20033-SNV/B/H/PE20038
Black	NGV/B/H/PEM10043-NGV/B/H/PEM10048
	NGV/B/H/PEM20043-NGV/B/H/PEM20048
	SNV/B/H/PE10043-SNV/B/H/PE10048
	SNV/B/H/PE20043-SNV/B/H/PE20048
Pink	NGV/B/H/PEM10053-NGV/B/H/PEM10058
	NGV/B/H/PEM20053-NGV/B/H/PEM20058

Anhui 2021-10-30

Place , date

Quality Manager

Legally binding signature, Function