

# Xianning Full Guard Medical Products Co.,Ltd

ADD: Yong'an East Avenue, Xian'an Distrct, Xianning City, P.R.C

TEL: +86-715-8210013 www.fullguardmedical.com

# **EC** Declaration of Conformity

### Manufacturer:

Name: Xianning Full Guard Medical Products Co., Ltd

Address: Yongan East Avenue, Xian'an Economic Development Zone, Xianning City, Hubei

Province, China

**Tel/Fax:** 0715-8200113

**SRN:** CN-MF-000013685

## Whose single Authorized Representative:

Name: ZOUSTECH S.L

Address: Pso.Castellana, 141- planta 19, 28046-Madrid, Spain

Tel/Fax: +34694426446

SRN: ES-AR-000002008

## **Disposable Surgical Gown**

**UMDNS CODE: 11901** 

**Product Code: FGSG** 

Product Size: XS, S, M, L, XL, XXL, XXXL, XXXXL

Classification According To MDD, Annex VII: Class I Sterile, Rule 1

**Applied Common Specification/Standard:** 

EN 13795-1:2019 Surgical clothing and drapes-Requirements and test methods part1:Surgical drapes and gown

EN ISO 11607-1:2020 Packaging for terminally sterilized medical devices - Part 1: Requirements for materials, sterile barrier systems and packaging systems

EN ISO 11607-2:2020 Packaging for terminally sterilized medical devices - Part 2: Validation requirements for forming, sealing and assembly processes

EN ISO 11135:2014 Sterilization of health-care products-Ethylene oxide - Requirements for the development, validation and routine control of a sterilization process for medical devices

EN ISO 14971:2019 Medical devices - Application of risk management to medical devices (ISO 14971:2019)

ISO 10993-1:2018 Biological evaluation of medical devices - Part 1: Evaluation and testing within a risk management process

EN ISO 10993-5:2009 Biological evaluation of medical devices - Part 5: Tests for in vitro cytotoxicity (ISO 10993-5:2009)

EN ISO 10993-10:2013 Biological evaluation of medical devices - Part 10: Tests for irritation and skin sensitization (ISO 10993-10:2010)



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EN ISO 1041:2008 Information supplied by the manufacturer of medical devices

EN ISO 15223-1:2016 Medical devices – Symbols to used with medical devices labels, labeling and information to be supplied.

We, the manufacturer, herewith declare under our sole responsibility that the above-mentioned product, meets the provision of the following EC Council Directives and All applicable harmonized Standards. All supporting documentations are retained under the premises of the manufacturer.

#### **DIRECTIVES**

Medical Device Directive:

COUNCIL DIRECTIVE 93/42/EEC of 14 June 1993 concerning medical devices (MDD/93/42/EEC) Amended by DIRECTIVE 2007/47/EC of 5 Sep 2007.



Notified Body: TÜV SÜD Product Service GmbH, Ridlerstr. 65, 80339, MÜnchen, Germany

NB Identification number: 0123

Certificate No: G2S 003747 0002 Rev.00

(EC) Certificate(s): YES

Expire date of Certificate: Nov 04, 2023

Start of CE Marking: Nov 05, 2018

Place of Issue: Xianng, Hubei

Date of Issue: 2021.5.26

Signature:

Name: 姜迪 Rosen Jiang

Position: Managing Director

Stamp:

Issued By: Molly Zheng Approved By: Rosen Jiang Revision: FD01 Effective Date: 2021.5.26