



Verification of CE Registration

CIBG-20201610

This is to certify that during the examination of the Technical Documentation provided by the manufacturer:

Name: Soozly (Ningbo) Medical Instrument Co.,Ltd

Address: Zhoujiaduan, Zhangqi, Cixi, Zhejiang, China

On its product as follows:

Product Name: Disposable medical mask, Medical face shield, Medical protective clothing, Medical Safety Goggles

Classification: Class I.

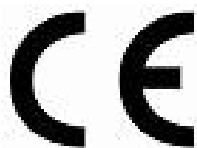
No Non-compliance according to the requirements of the Medical Device Regulation (EU) 2017/745 Annex II and Annex III was detected, and the aforementioned device complies with the Regulation including all General Safety and Performance Requirements defined in Annex I.

The manufacturer has provided the EU Declaration of Conformity according to the Medical Device Regulation (EU) 2017/745 - article 19 requirements, confirming that this medical device, as stipulated above, is fulfilling the applicable requirements of the Medical Device Regulation (EU) 2017/745.

The notification of aforementioned device has been completed by the European Representative in Netherlands. The Netherlands Competent Authority is notified of the manufacturer's medical devices and has allocated registration.

Memos:

This file is Not being issued by EU authorities or NB, all refer the CIBG registration certificate as the legal basis. We, Shanghai CV, as the 3rd party, produce this file, intended of facilitate client display & transmit information. The user can contact CIBG using the Eight-digit registration number to validate it, the CIBG web page: <https://hulpmiddelen.farmatec.nl>



Thomas E. Weast, P.E.
Executive Director

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Shanghai CV Certification Technology Co., Ltd.

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