



# CE Notification Confirmation

This is to confirm that, according to the council directive 93/42/EEC (MDD), SUNGO Europe B.V. performed the notification duties and responsibilities as the European authorized representative of:

**Xiantao Chenguang Protection Commodity Co., Ltd.**  
Gongqing Industry Garden, Yewang Road, Xiantao City,  
Hubei China

The Manufacturer has provided SUNGO Europe B.V. with the EC Declaration of Conformity confirming that the medical device, as stipulated here below, is fulfilling the applicable requirements of the European Council Directive 93/42/EEC.

According to 93/42/EEC (MDD), the European Databank on Medical Devices (EUDAMED) is established as of May 1 2011. The Netherlands Competent Authority is notified of the manufacturer's medical devices and has allocated registration number.

## Face Mask

Class I according to Annex IX of 93/42/EEC  
GMDN: 35177  
**CIBG Number: NL-CA002-2020-50268**

Where the manufacturer affix's the CE marking to the product listed they must ensure that all the requirements of the appropriate EU directive(s) have and continue to be met.

*This document should be used together with the competence authority notification letter and the Declaration of Conformity issued by the Manufacturer. This document will become to be invalid once the notification status is changed or the EAR agreement is terminated.*

Reference Number: EUCAN00135  
Issue date: Apr.23, 2020

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For and on behalf of  
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Authorized Signature  
Only used for the EU Representative Signature