

## EC Declaration of Conformity

**Manufacturer:**

Shenzhen Comen Medical Instruments Co.,Ltd

**Address:**

Floor 10, Floor 11 and Section C of Floor 12 of  
Building 1A & Floor 1 to Floor 5 of Building 2,  
FIYTA Timepiece Building, Nanhuan Avenue,  
Matian Sub-district, Guangming District,  
Shenzhen, Guangdong, 518106, P.R. China.

**Whose Single Authorized Representative:**

Lotus NL B.V.

**Address:**

Koningin Julianaplein 10, 1e Verd, 2595AA,  
The Hague, Netherlands.

We, the manufacturer, declare at our sole responsibility that following products

| Product name          | Model          |
|-----------------------|----------------|
| Defibrillator Monitor | S8, S6, S5, S3 |

meet the provisions of Directive 93/42/EEC.

The medical device has been assigned to class IIb according to rule 10 in Annex IX of the Directive 93/42/EEC. It bears the mark

CE 1639

The product concerned has been designed and manufactured under a quality management system according to Annex II (excluding Section 4) of Directive 93/42/EEC.

The product meet the following standard: (See Chapter 4 of Document No. 0039-63)

Compliance of the designated product with the Annex II (excluding Section 4) of Directive 93/42/EEC has been assessed and certified by the Notified Body

**SGS Belgium NV**  
**SGS House Noorderlaan**  
**87 2030 Antwerp Belgium**

CertificateNo.: CN19/41057

Issuedate: 2021.03.22

Expirydate: 2023.02.05

The above mentioned declaration of conformity is exclusively under the responsibility of

Company: Shenzhen Comen Medical Instruments Co.,Ltd

Address: Floor 10, Floor 11 and Section C of Floor 12 of Building 1A & Floor 1  
to Floor 5 of Building 2, FIYTA Timepiece Building, Nanhuan Avenue,  
Matian Sub-district, Guangming District, Shenzhen, Guangdong,  
518106, P.R. China.

Shenzhen, 2021.05.08  
Place, date

Gary Dunn Management Representative  
Legally binding signature, Function

