

## EC Certificate

### Production Quality Assurance System according to Medical Devices Directive 93/42/EEC Annex-V

Certificate Number: 1984-MDD-17-433

We hereby declare that an examination has been carried out following the requirements of the national legislation to which the undersigned is subject, transposing Annex-V of the Directive 93/42/EEC on medical devices. We certify that the production quality system conforms with the relevant provisions of the aforementioned legislation.

**Organization:**

### MEDİMPORT SAĞLIK ÜRÜNLERİ SANAYİ VE TİCARET LİMİTED ŞİRKETİ

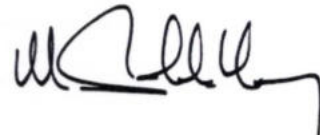
İkitelli OSB Mah. Aykosan 4'lü A Blok Sk. Aykosan Sitesi 4'lü A Blok No:4 İç Kapı  
No:245 Başakşehir/İstanbul, Turkey

Products	Types
Portable suction unit	AL-01, MSC-101, AL-03
Compressor nebuliser device	AL-20, NB-01, AL-50

The certificate is valid till expiration date, subject to successful completion of periodical surveillance audits. Please contact Kiwa for details.

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Kiwa Belgelendirme Hizmetleri A.Ş. is Notified Body under Council Directive 93/42/EEC concerning medical devices with identification number: 1984



Muhteşem Gökhan Yücel  
Head of Notified Body

10 May 2021, Istanbul, Turkey