

EC CERTIFICATE AT SERTİFİKA

According to Annex V of the Directive 93/42/EEC on Medical Devices
93/42/AT Tıbbi Cihaz Yönetmeliği Ek V'e göre

Production Quality Assurance System Üretim Kalite Güvencesi

Certificate Number: 2195-MED-1816401
Sertifika Numarası

Manufacturer:
Üretici

R Vent Medikal Üretim A.Ş.
29 Ekim Mah. Balkan Cad. No:33 Torbalı, İzmir, TÜRKİYE

Product(s):
Ürün(ler)

- (1) Sterile and Non-Sterile Breathing Circuit Systems**
(1) Steril ve Steril Olmayan Solunum Devre Sistemleri
- (2) Sterile and Non-Sterile Breathing Filters**
(2) Steril ve Steril Olmayan Solunum Filtreleri
- (3) Sterile and Non-Sterile Catheter Mounts**
(3) Steril ve Steril Olmayan Katater Bağlantıları
- (4) Non-sterile Masks, BVM (Resuscitator), O₂ & Aerosol Therapy Set**
(4) Steril Olmayan Maskeler, BVM (Resusitatör), O₂ & Aeresol Terapi Seti
- (5) Sterile Closed Suction System**
(5) Steril Kapalı Emiş Sistemi

Reference Report No: MM0687-P004-R01, MM0687-P004-R02, MM0687-P005-R01
Referans Rapor No

Szutest, Notified Body 2195, declares that the aforementioned manufacturer has implemented a quality assurance system according to Annex V, Section 3 of the directive 93/42/EEC on medical devices. This quality assurance system covers those aspects of manufacturing concerned with securing and maintaining safe conditions of the respective product(s) and conforms to the provisions of this Directive. The approved quality system is subject to surveillance pursuant to Annex V, Section 4 of Directive 93/42/EEC and unannounced audits.

Szutest must be informed of any significant changes in the design and/or construction of the product(s). For class I devices with sterile conditions the quality management system evaluation is restricted to the aspects of manufacture concerned with securing and maintaining sterile conditions. For class I devices with measuring function the quality management system evaluation is restricted to the aspects of manufacture concerned with the conformity of the devices with metrological requirements.

2195 kimlik numaralı Onaylanmış Kuruluş Szutest, yukarıda belirtilen üreticinin 93/42/AT Tıbbi Cihaz Yönetmeliği EK V bölüm 3'üne göre bir kalite yönetim sistemi uyguladığını, bu yönetim sisteminin yönetmeliğin sadece bahsi geçen ürünün üretiminin güvenlik koşullarını sağlama ve devam ettirme ile ilgili gerekliliklerin karşıladığını beyan eder. Onaylanan bu kalite yönetim sistemi, 93/42/AT Tıbbi Cihaz Yönetmeliği EK V, bölüm 4'e göre periyodik olarak gözetime ve habersiz saha denetimlerine tabidir.

Üretici, ürünlerinin tasarımında ve yapısında gerçekleştirdiği önemli değişiklikleri Szutest'e bildirmek zorundadır. Steril kondisyondaki sınıf I ürünler için kalite yönetim sistemi değerlendirmesi üretimin steril kondisyonun sağlanması ve korunmasıyla limitlidir. Ölçüm fonksiyonlu sınıf I ürünler için Kalite yönetim sistemi değerlendirmesi üretimin cihazların metrolojik şartlara uyumunu sağlanmasıyla limitlidir.

This EC certificate is valid till 2024-05-26.
Bu AT Sertifikası 2024-05-26 tarihine kadar geçerlidir.

Issue Date/Yayın Tarihi: 2018-06-13
Revision No./ Revizyon No.: 02 Rev./Rev.
Revision Date/ Revizyon Tarihi: 2020-06-26



Rukiye BALKAN
Deputy General Manager
Genel Müdür Yardımcısı



STATEMENT: EXTENSION OF MDD CERTIFICATE

To whom it may concern,

The R Vent Medikal Üretim A.S. herewith declares that the conditions granted by Regulation (EU) 2023/607 are fulfilled for the extension of the validity of MDD certificate 2195-MED-1816401 until 2028-12-31.

In particular, R Vent Medikal Üretim A.S. declares that:

1. Before the date of expiry of the certificate, the R Vent Medikal Üretim A.S. signed a written agreement with SZUTEST Konformitätsbewertungsstelle GmbH (NB: 2975) in accordance with Section 4.3, second subparagraph, of Annex VII to MDR for the conformity assessment.
2. The devices covered by the MDD certificate 2195-MED-1816401 continue to comply with the MDD.
3. During the extension period, no significant changes will be applied to the design and intended purpose of the medical devices covered by the extension.
4. The devices do not present an unacceptable risk to the health or safety of patients, users or other persons, or to other aspects of the protection of public health.
5. R Vent Medikal Üretim A.S. Quality Management System complies with Annex IV of the MDR. For such Quality Management System, R Vent hold an ISO 13485:2016 certificate 31816401 issued by SZUTEST Konformitätsbewertungsstelle GmbH, valid until 10.06.2024.
6. The applications for certificate under MDR and the Technical Documentation have been acknowledged by SZUTEST Konformitätsbewertungsstelle GmbH in 2024.

A complete list of devices by the extension is provides in Table 1 below.

1. Sterile and Non-Sterile Breathing Circuit Systems
2. Sterile and Non-Sterile Breathing Filters
3. Sterile and Non-Sterile Catheter Mounts
4. Non-Sterile Masks, BVM (Resuscitator), O₂ & Aerosol Therapy Sets
5. Sterile Closed Suction Systems

Should you have any questions or comment, please do not hesitate to contact R Vent Medikal Üretim A.S.

Name of the Company: R Vent Medikal Üretim A.S.

Address: Yazıbaşı Mah. Balkan Cad. İztipsan Apt. No: 33/1 Torbalı, İzmir, Türkiye

E-mail: info@rventmedikal.com

Telephone: +90 232 853 95 00

Document No: SD.MDRT-CS-0905

Release Date: 09.05.2024

Rev. No: 00

Rev. Date: -

Signature:

For and on behalf of R Vent Medikal Üretim A.S.

Aybüke Elif US

Quality Management Representative

R VENT MEDİKAL ÜRETİM A.Ş.

Yazıbaşı Mah. Balkan Cd. İztipsan Apt. No:33

TORBALI/İZMİR

TORBALI V.D. 734 081 2763

R-Vent Medikal Üretim A.Ş.

A : Yazıbaşı Mah. Balkan Cad. İztipsan Apt. No:33/1 - 35860 Torbalı / İzmir / TURKEY

T: +90 232 853 95 00 · +90 232 853 94 95 · www.rventmedikal.com · info@rventmedikal.com

TORBALI V.D. 734 081 27 63 · Ticaret Sicil No: 5444 / TORBALI

Banka Hesap Numaraları / Bank Account No

Ziraat Bankası TL IBAN 73 0001 0022 9274 4921 1250 01

Ziraat Bankası EURO IBAN 46 0001 0022 9274 4921 1250 02

Ziraat Bankası USD IBAN 19 0001 0022 9274 4921 1250 03

Ziraat Bankası GBP IBAN 89 0001 0022 9274 4921 1250 04

NOTIFIED BODY CONFIRMATION LETTER
No: MD0045-CL-01

To whom it may concern,

Confirmation of the status of a formal application, written agreement, and appropriate surveillance in the framework of Regulation (EU) 2023/607 amending Regulation (EU) 2017/745 and implementing Regulation (EU) 2023/1194 amending implementing Regulation (EU) 2022/2346 as regards the transitional provisions for certain medical devices.

This letter confirms that **SZUTEST Konformitätsbewertungsstelle GmbH**, a Notified Body (NB) designated against Regulation (EU) 2017/745 (MDR) and identified by the number **2975** on NANDO, has received a formal application in accordance with Section 4.3, first subparagraph of Annex VII of MDR and has signed a written agreement in accordance with Section 4.3, second subparagraph of Annex VII of MDR with the following manufacturer:

Company Name	R Vent Medikal Üretim A.Ş.
Address	Yazıbaşı Mah. Balkan Cad. İztıpsan Apt. No:33/1 Torbalı, İzmir, TÜRKİYE
SRN Number (if available)	TR-MF-000028282

The devices covered by the formal application and the written agreement mentioned above are identified in the Tables below. Table 1 identifies the devices for which an MDR application has been received, written agreement concluded, and for which the SZUTEST Konformitätsbewertungsstelle GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive. Table 2 identifies the devices for which an MDR application has been received and a written agreement concluded, but SZUTEST Konformitätsbewertungsstelle GmbH has not yet taken responsibility for appropriate surveillance of the corresponding devices under the applicable Directive.

In the case of devices covered by certificates issued under Directive 93/42/EEC (MDD) that expired after 26 May 2021 and before 20 March 2023, without having been withdrawn, this letter also confirms that the manufacturer signed the written agreement under MDR by the date of MDD certificate expiry; or provided evidence that a competent authority of a Member State had granted a derogation or exemption from the applicable conformity assessment procedure in accordance with Article 59(1) of MDR or Article 97(1) of the MDR respectively, by the 20 Mar 2023 for the relevant devices.

The transition timelines that apply to the devices covered by this letter, subject to the manufacturer's continued compliance with the other conditions specified in Article 120.3c of MDR (as amended by (EU) 2023/607), are shown below:

- 31 December 2027 for Class III devices and Class IIb implantable devices excluding well-established technologies (WET - sutures, staples, dental fillings, dental braces, tooth crowns, screws, wedges, plates, wires, pins, clips, and connectors)
- 31 December 2028 for other Class IIb devices, Class IIa, Class I devices placed on the market in sterile condition or have a measuring function
- 31 December 2028 for devices not requiring the involvement of a notified body under MDD but requiring it under MDR
- 31 December 2028 for Annex XVI products which do not require a clinical investigation.
- 31 December 2029 for Annex XVI products which require a clinical investigation.

On behalf of SZUTEST Konformitätsbewertungsstelle GmbH,

MEHMET İŞIKLAR
General Manager

SZUTEST
Konformitätsbewertungsstelle GmbH
Friedrich-Ebert-Anlage 36
60325 Frankfurt am Main
VAT-IdNr. DE815819575
info@szutest-germany.de

SZUTEST Konformitätsbewertungsstelle GmbH-NB 2975
Friedrich-Ebert-Anlage 36 D-60325 Frankfurt am Main /GERMANY



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Table 1: Devices covered by this letter and for which SZUTEST Konformitätsbewertungsstelle GmbH is also responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (Under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD device	MDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
N/A	N/A	N/A	N/A



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Table 2: Devices covered by this letter and for which SZUTEST Konformitätsbewertungsstelle GmbH is NOT responsible for appropriate surveillance of the corresponding devices under the applicable Directive:

Device name or Basic UDI-DI (Under MDR application)	MDR Device classification (as proposed by the manufacturer and verified at the pre-application stage)	If the MDR device is a substitute device, identification of the corresponding MDD device	MDD Certificate Reference(s) of the devices under MDR application, and the NB Identification
Sterile and Non-Sterile Breathing Circuit Systems	Class IIa	Same	Certificate #1; 2195-MED-1816401 Revision No:2 Revision Date: 26.06.2020 Issue Date:13.06.2018 Expiry Date:26.05.2024 NB2195: Szutest Uygunluk Değerlendirme A.Ş.
Sterile and Non-Sterile Breathing Filters	Class IIa	Same	Certificate #1; 2195-MED-1816401 Revision No:2 Revision Date: 26.06.2020 Issue Date:13.06.2018 Expiry Date:26.05.2024 NB2195: Szutest Uygunluk Değerlendirme A.Ş.
Sterile and Non-Sterile Catheter Mounts	Class IIa	Same	Certificate #1; 2195-MED-1816401 Revision No:2 Revision Date: 26.06.2020 Issue Date:13.06.2018 Expiry Date:26.05.2024 NB2195: Szutest Uygunluk Değerlendirme A.Ş.
Non-sterile Masks, BVM (Resuscitator), O2 & Aerosol Therapy Set	Class IIa	Same	Certificate #1; 2195-MED-1816401 Revision No:2 Revision Date: 26.06.2020 Issue Date:13.06.2018 Expiry Date:26.05.2024 NB2195: Szutest Uygunluk Değerlendirme A.Ş.
Sterile Closed Suction System	Class IIa	Same	Certificate #1; 2195-MED-1816401 Revision No:2 Revision Date: 26.06.2020 Issue Date:13.06.2018 Expiry Date:26.05.2024 NB2195: Szutest Uygunluk Değerlendirme A.Ş.

Confirmation Letter Revision History

Date	Version of the letter	Action
2024/04/03	MD0045-CL-01	Initial issue



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