

## EC CERTIFICATE

According to Annex II of the Directive 93/42/EEC on Medical Devices

### Full Quality Assurance System

Certificate Number: 2195-MED-2030401

**Manufacturer:** Nucleoss Europe GMBH  
Graben17, 64646, Heppenheim / Germany

**Product(s):** 1. Sterile Dental Implant  
2. Non-Sterile Dental Abutments

**Model(s):** DC

**Reference Report No:** MM0827-P001-R02, MM0827-P001-R02

Szutest, Notified Body 2195, declares that the aforementioned manufacturer has implemented a quality assurance system according to Annex II (excluding section 4), 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 II, Section 5 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.

*This EC certificate is valid till 2024-05-26.*

**Issue Date:** 2020-10-30



Rukiye BALKAN  
Deputy General Manager

## CERTIFICATE INFO AMENDMENT

### SERTİFİKA BİLGİ DEĞİŞİKLİĞİ

According to Article 120(3) of the Regulation (EU) 2017/745 on Medical Devices

(AB) 2017/745 Tıbbi Cihazlar Yönetmeliği Madde 120(3)'ye göre

**Effected Certificate Number(s):** 2195-MED-2030401  
*Etkilenen Sertifika Numarası(ları):*

**Manufacturer:** Nucleoss Europe GMBH  
*Üretici* Tiergartenstr. 7b, 64646, Heppenheim (Bergstr.) GERMANY

**Product(s):** No change.  
*Ürün(ler)*

**Model(s):** No change.  
*Model(ler)*

**Reference Report No:** MM0827-P006-R01, MM0827-P006-R02  
*Referans Rapor No*

**Definition of the Change:** Address change.  
*Değişikliğin Tanımı*

SZUTEST, Notified Body 2195, declares and the above mentioned manufacturer has initiated an insignificant change according to Article 120(3) of (EU) 2017/745 and MDCG 2020-3 guidance and therefore the information on the effected 93/42/EEC certificate(s) has been changed as described above.

This document is a confirmation for authorities and cannot be used as other purposes.

2195 kimlik numaralı Onaylanmış Kuruluş SZUTEST, yukarıda belirtilen üreticinin (AB) 2017/745 Regülasyonu Madde 120(3)'e ve MDCG 2020-3 rehber dokümanına göre önemli olmayan bir değişiklik yürüttüğünü ve bu sebeple etkilenen 93/42/AT sertifika(lar)ındaki bilgilerin yukarıdaki gibi değiştiğini beyan eder.

Bu doküman yetkili otoriteler için bir onay niteliğinde olup farklı bir amaçla kullanılamaz.

**Issue Date/Yayın Tarihi:** 2024-03-06



Rukiye BALKAN  
Deputy General Manager  
Genel Müdür Yardımcısı