EC DECLARATION OF CONFORMITY



Name of basic product:

MTA+ (MINI, STANDARD, MAXI, PRO)

Variant:

BIO MTA+ (MINI, STANDARD, MAXI, PRO)

Manufacturer:

Przedsiębiorstwo Produkcyjno-Handlowe CERKAMED Wojciech Pawłowski 37-450 Stalowa Wola, ul. Kwiatkowskiego 1, POLAND

Purpose and range of use:

The product is intended for dental treatment as a material for filling and rebuilding of root canals.

Medical device of class IIa, according to the rule 8 of Annex IX MDD 93/42/ EEC. Evaluation of conformity was conducted following the procedure relating to Annex II MDD 93/42 / EEC excluding section 4.

Reference documents:

- Council Directive 93/42 / EEC of 14 June 1993 concerning medical devices with a changes 2007/47/ EC
- The Act of May 20, 2010. about medical devices with a changes
- Regulation of the Minister of Health of February 17, 2016 on essential requirements and conformity assessment procedures for medical devices

Evaluation of conformity was conducted with participation of the notified body: CE Certiso Ltd., Organisation for Certification and Testing on the Field of Medical and Hospital Engineering, nr ID 2409 Erdő utca 101, 2092 Budakeszi, Hungary.

Conformity of the product confirmed with certificate No.144731-18-02-18 valid until 2023-02-17.

We declare with full responsibility that the manufactured product with variant, which this statement refers to, complies with the reference documents mentioned above.

PRZEDSIEBIORSTWO PRODUKCYJNO-HANDLOWE

Honorata Sołowiej, Person responsible for regulatory compliance,

On behalf of Wojciech Pawłowski

Stalowa Wola

WOJCIECH PAWŁOWSKI
ul. Kwiatkowskiego 1
37-450 STALOWA WOLA
tel./fax 15 842 35 85
www.cerkamed.pt NIP 865-204-87-70

010me 29.11.202

signature, company stamp, date

updated: 06.04.2021/10