EU DECLARATION OF CONFORMITY



Name of product: RUBBER-DAM TEMPLATE

Manufacturer:

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

SRN (Single Registration Number):

PL-MF-000003211

Purpose and range of use:

RUBBER-DAM TEMPLATE enables selecting particular points on Rubber-Dam, in which the hole should be made.

Medical device of class I, according to the rule 1 of Annex VIII MDR (EU) 2017/745. Evaluation of conformity was conducted following the procedure relating to Annex II and III Regulation (EU) 2017/745.

BASIC UDI-DI:

590755302SZABLONX9

Common specifications:

EN ISO 14971:2012 Medical devices — Application of risk management to medical

devices

EN 1041:2008+A1:2013 Information provided by the manufacturer of medical devices.

EN ISO 15223-1:2016 Medical devices — Symbols to be used with medical device labels, labelling and information to be supplied — Part 1:

General requirements

EN 1639:2009 Dentistry - Medical devices for dentistry - Instruments

Ph. Eur. 10 2020 European Pharmacopoeia

Reference documents:

- Regulation (EU) 2017/745 of 5 April 2017 on medical devices, amending Directive 2001/83/EC, Regulation (EC) No 178/2002 and Regulation (EC) No 1223/2009 and repealing Council Directives 90/385/EEC and 93/42/EEC
- The Act of May 20, 2010 about medical devices with a changes

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

This declaration of conformity is issued under the sole responsibility of manufacturer.

Honorata Sołowiej,

Person responsible for regulatory compliance,

On behalf of Wojciech Pawłowski

Stalowa Wola

WOJCIECH PAWLOWSKI
ul. Kwiatkowskiego 1
37-450 STALOWA WOLA
tel./fax 18 842 35 85
(www.cerkamed.pt NIP 865,204-87-70

PRZEDSIĘBIORSTWO PRODUKCYJNO-HANDLOWE

signature, company stamp, date