

## Declaration of Conformity

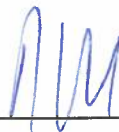
CE 0123

Manufacturer Address	Ivoclar Vivadent AG Bendererstrasse 2 9494 Schaan Liechtenstein
Product	<b>IPS e.max CAD</b>
Type of material	Ceramic System
Product category	Materials for inlays, onlays, crowns and bridges
Classification	Medical Device Class IIa

We hereby declare under our exclusive responsibility that the above mentioned products meet the provisions of the following EC Council Directives and its implementation in national law. All supporting documentation is retained on the premises of the manufacturer and the notified body.

Directives	Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)
Notified Body Address	TÜV SÜD Product Service GmbH Ridlerstrasse 65 80339 München Deutschland
Place, Valid from	Schaan, 2021-05-25 Replaces version of: 2020-02-20
Valid until	2024-05-26

Signature




Name  
Position

Dr. Thomas Hirt  
CTO

Dipl. Ing. Patrik Oehri  
Director CQM and Regulatory  
Sicherheitsbeauftragter  
(Product Safety Officer)

Date

2021-05-25

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