

EC DECLARATION OF CONFORMITY

Name and address of the manufacturer: Foshan COXO Medical Instrument Co., Ltd.
Site 1: BLDG 4, District A Guangdong New Light Source Industrial Base, South of Luocun Avenue Nanhai District Foshan 528226 Guangdong China
Site 2: No. 17, Guangming Ave., New Light Source Industrial Base, Nanhai National High-tech Zone, Foshan 528226, Guangdong P.R. China

EC Representative: Lotus NL B.V.
Koningin Julianaplein 10, 1e Verd, 2595AA, The Hague, Netherlands.
E-mail: peter@lotusnl.com

Trademarks: **COXO, YUSENDENT, YSDENT, CODENTAL**

We declare under our sole responsibility that

the medical device:

Product Name: Root Apex Locators

Model: C-Root I, C-Root I(III), C-Root I(V), C-Root I(VI),
C-Root i+

of class: IIa, rule 10
according to annex IX of directive 93/42/EEC

meets the provisions of the directive 93/42/EEC as amended by Directive 2007/47/EC and its transpositions in national laws which apply to it. The declaration is valid in connection with the "final inspection report" of the device.

Conformity assessment procedure: **Directive 93/42/EEC Annex V**

Registration No.: **DD 60151346 0001**

Notified Body:
TÜV Rheinland LGA Products GmbH
Tillystraße 2
90431 Nürnberg
Deutschland
CE 0197

(FoShan), PR China 2020-12-08

Place, date

Title: General Manager
Name: (Mr) Zheng Yongliang
Signature:

