## **EC DECLARATION OF CONFORMITY**

Name and address of the manufacturer: Foshan SOCO Precision Instrument Co.,Ltd.

2FL.Bldg 3,District A Guangdong New Light Source Industrial Base,Luocun Shishan Town,Nanhai District Foshan City

528226 Guangdong China

EC Representative: Lotus NL B.V.

Koningin Julianaplein 10,1e Verd,2595AA,The Hague,

Netherlands.

We declare under our sole responsibility that

the medical device: Product Name: Dental Root Canal Instruments

Model: SC, SC-PRO, SC plus, S-one plus, S-one pro,

SX-F3, SC-GOLD

of class: Ila, rule 6

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 2056447-1

Notified Body: TÜV Rheinland LGA Products GmbH

Tillystraße 2 90431 Nürnberg Deutschland CE 0197

(FoShan), PR China 2021-05-25

Place, date

Name: (Mr) Zeng, You Miles Signature:

Name and function

Title: General Manage