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EC Declaration of Conformity

*Manufacturer:**whose single Authorized Representative:***Guilin Woodpecker Medical Instrument Co., Ltd.****MedNet EC-REP GmbH • Borkstrasse 10
• 48163 Muenster • Germany**

We, the manufacturer, herewith declare that the products
Gutta Percha Obturation Device, GMDN-Code: 44859
MODEL: Fi-P, Fi-G, Fi-E

meet the provisions of Directive 93/42/EEC which apply to them.

The medical device has been assigned to class IIa according to Annex IX of the Directive 93/42/EEC. It bears the mark

CE 0197

The product concerned has been manufactured under a quality management system according to Annex II of Directive 93/42/EEC.

Compliance of the designated product with the Directive 93/42/EEC has been assessed and certified by the Notified Body

TÜV Rheinland LGA Products GmbH
Tillystraße 2, 90431, Nürnberg, Germany

Certificate No.: HD 2158053-1

Issue date: 2021-05-19

Expiry date: 2024-05-26

following the procedure relating to the EC Declaration of Conformity set out in Annex II of Directive 93/42/EEC.

This Declaration of Conformity covers all medical devices as specified in the product list belonging to this declaration and is only valid in connection with a batch specific Certificate of Compliance for all products concerned bearing the CE mark

The above mentioned declaration of conformity is exclusively under the responsibility of

Company: **Guilin Woodpecker Medical Instrument Co., Ltd.**

Address: Information Industrial Park, Guilin National High Tech Zone, Guilin, GuangXi,
541004, P.R.China

陈燕 2022.3.9
Preparation, Date

王波 2022.3.9
Review, Date

丁加 2022.3.9
Legally binding signature, Location, Function

