

EC Declaration of Conformity

Doc No : DC-114 Rev.No : 1

Manufacturer:

DiaDent Group International

16, Osongsaengmyeong 4-ro Osong-eup Heungdeok-gu,

Cheongju-si, Chungcheongbuk-do,

28161, Republic of Korea

We, the manufacturer, herewith declare that the products

Authorized Representative:

DiaDent Europe B.V.

Antennestraat 70, 1322AS Almere, the Netherlands

DIA-ROOT BIO Sealer

(including system components and accessories)

GMDN

36095

meet the provisions of The Medical Devices Directive 93/42/EEC amended by 2007/47/EC which apply to them.

The medical device has been assigned to Class IIa(Rule 8) according to Annex IX of the Medical Devices Directive 93/42/EEC amended by 2007/47/EC. It bears the mark

The product concerned has been designed and manufactured under a quality management system according to Annex II of the Medical Devices Directive 93/42/EEC amended by 2007/47/EC.

Compliance of the designated product with The Medical Devices Directive 93/42/EEC amended by 2007/47/EC has been assessed and certified by the Notified Body

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

Certificate No HD 60149568 0001

Issue date 2020-05-25 Expiry date 2024-05-26

following the procedure relating to the EC Declaration of Conformity set out in Annex II of the Medical Devices Directive 93/42/EEC amended by 2007/47/EC

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

DiaDent Group International

16, Osongsaengmyeong 4-ro Osong-eup Heungdeok-gu, Cheongju-si Chungcheongbuk-do, 28161, Republic of Korea

2020-06-01

Fax: 82-43-262-8658 Signature

Date

DiaDent Group International

16, Osongsaengmyeong 4-ro, Osong-eup, Heungdeok-gu, Cheongju-si, Chungcheongbuk-do, 28161, Korea

Tel: 82-43-266-2315

Fax: 82-43-262-8658

http://www.diadent.co.kr

E-mail: diadent@diadent.co.kr

