



META BIOMED CO., LTD.

270, Osongsaengmyeong 1-ro, Osong-eup, Heungdeok-gu, Cheongju-si,
Chungcheongbuk-do, Korea

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EC Declaration of Conformity

**according to the Medical Devices Directive 93/42/EEC
as amended by Directive 2007/47/EC**

For the following medical device :

Product Name : Dental Composite Resin

Brand Name : Ezfil (detailed model names attached)

Classification : Class IIa by Rule 8 of Annex IX, Council Directive 93/42/EEC as amended by
Directive 2007/47/EC

Manufacturer's Name : Meta Biomed Co., Ltd.

Manufacturer's Address : 270, Osongsaengmyeong1-ro, Osong-eup, Heungdeok-gu, Cheongju-si,
Chungcheongbuk-do, Korea

is herewith the conformity with the requirements set out in the Council Directive 93/42/EEC of 14 June 1993 as amended by Directive 2007/47/EC of the European Parliament and of the Council of 5 September 2007 concerning Medical Devices declared, For the evaluation of the conformity with this Directive, the following standards have been applied:

Directive 93/42/EEC amended by Directive 2007/47/EC, MEDDEV 2.12-1 Rev.8, MEDDEV 2.7.1 Rev.4, MEDDEV 2.12.2 Rev.2, EN ISO 13485:2016, FDA 21 CFR 820, EN ISO 14971:2019, ISO/TR 24971:2020, EN 1041:2008, EN ISO 15223-1:2016, EN 62366-1:2015, IEC TR 62366-2:2016, ISO/TR 20416:2020, EN 1641:2009, EN ISO 4049:2019, EN ISO 7405:2018, EN ISO 10993-1:2009, EN ISO 10993-3:2014, EN ISO 10993-5:2009, EN ISO 10993-6:2009, EN ISO 10993-10:2023, EN ISO 10993-11:2018, EN ISO 10993-23:2021, EN ISO 14644-1:2015, EN ISO 14644-2:2015, EN ISO 14644-3:2019, EN ISO 14644-4:2022, EN ISO 14644-5:2004, EN ISO 14644-7:2004, EN ISO 14644-8:2022, EN ISO 14644-9:2022, EN ISO 14644-10:2022, ISO 14644-12:2018, EN ISO 14644-13:2017, EN ISO 14644-14:2016, EN ISO 14644-15:2017, EN ISO 14644-16:2019, EN ISO 14644-17:2021, EN ISO 14698-1:2003, EN ISO 14698-2:2003, BS EN 17141:2020, KS A 3011:2018, USP <1116>, ASTM F2847-17, ASTM G131-96(2016)e1, EN ISO 11737-1:2018/A1:2021, EN ISO/TS 17665-2:2009, Korean Pharmacopeia 12th, ASTM F1980-21, ASTM F88/F88M-21, KS Q ISO 2859-1:2014, ISTA 1A:2022

is subject to the procedure set out in Annex II(excluding section 4) of Directive 93/42/EEC under the supervision of Notified Body Number 1639, SGS Belgium NV, Noorderlaan 87, BE-2030 Antwerpen, Belgium.

GMDN Code: 35870(Dental composite resin)

EC Certificate Number : KR19/81826259

Responsible for making this declaration is the :

■ Manufacturer: META BIOMED CO., LTD.

European Representative is:

■ Meta Biomed Europe GmbH, Wiesenstr. 35, 45473 Mulheim an der Ruhr, Germany

■ Tel: +49-208-30991910, Fax: +49-208-30991999

We, the manufacturer's declaration, declare and ensure with sole responsibility, that the Medical Device meet the provisions of Council Directive 93/42/EEC (Medical Device Directive) which apply to them.

Person responsible for making this declaration:

Name, Surname : Suk Song Oh

Position/Title : PRESIDENT

Place: Cheongju-si

Date: November 08, 2023

Suk Song Oh
(Company stamp and legal signature)



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(Attachment: Model name list)

No	Brand Name	Model Name
1	Ezfil	Ezfil A1
2		Ezfil A2
3		Ezfil A3
4		Ezfil A3.5
5		Ezfil A4
6		Ezfil B1
7		Ezfil B2
8		Ezfil B3
9		Ezfil B4
10		Ezfil C1
11		Ezfil C2
12		Ezfil C3
13		Ezfil C4
14		Ezfil D2
15		Ezfil D3
16		Ezfil D4
17		Ezfil OP
18		Ezfil TL
19		Ezfil kit