

To whom it may concern

## Manufacturer's Authorization

Date: April 22, 2020

We Boditech Med Inc., who are official manufacturers of the ichroma and the AFIAS products, having factories at 43, Geodudanji 1-gil, Dongnae-myeon, Chuncheon-si Gang-won-do, Korea 200-883, do hereby declare that

ECHIPAMED PLUS SRL str. Valea Trandafirilor 24 "B", of. 80 MD-2001, Chisinau Republic of Moldova

is our official distributor and local representative for the ichroma and the AFIAS products of Boditech Med Inc., in the territory of the Republic of Moldova.

We declare that above mentioned company is authorized to quote, sell, subsequently negotiate and sign contracts, as well as to perform installation and after sales service of the ichroma and the AFIAS products, manufactured by us.

We hereby extend our full warranty with respect to the Goods offered by the above company.

This authorization letter will remain valid until 31.12.2020.

Boditech Med Inc.

Hye-sung Kim Sales manger

43, Geodudanji 1-gil, Dongnae-myeon, Chuncheon-si, Gangwon-do, South Korea

Boditech Med Inc.

PRESIDENT EUI YUL CHOI









## Certificate

No. Q5 053112 0026 Rev. 00

Holder of Certificate: Boditech Med Inc.

> 43, Geodudanji 1-gil, Dongnae-myeon Chuncheon-si, Gang-won-do 24398

REPUBLIC OF KOREA

Boditech Med Inc. Facility(ies):

43, Geodudanji 1-gil, Dongnae-myeon, Chuncheon-si, Gang-won-

do 24398, REPUBLIC OF KOREA

Boditech Med Inc.

14, Geodudanji 1-gil, Dongnae-myeon, Chuncheon-si, Gang-won-

do 24398, REPUBLIC OF KOREA

**Certification Mark:** 



Scope of Certificate: Design, Development, Production and Distribution

> of In Vitro Diagnostic Medical Devices - Reagents and Instruments for Point of Care Testing(POCT)

Applied Standard(s):

EN ISO 13485:2016

Medical devices - Quality management systems -

Requirements for regulatory purposes

(ISO 13485:2016) DIN EN ISO 13485:2016

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). See also notes overleaf.

Report No.:

74954566

Valid from:

2019-11-01

Valid until:

2022-10-31

Date,

2019-10-15

Stefan Preiß

1. Punil

Head of Certification/Notified Body

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TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany

TÜV®

## **DECLARATION OF CONFORMITY**

Manufacturer:

Boditech Med Incorporated

43, Geodudanji 1-gil, Dongnae-myeon,

Chuncheon-si, Gang-won-do, 24398

REPUBLIC OF KOREA

European Representative:

**OBELIS S.A** 

Bd. Général Wahis 53,

1030 Brussels,

Belgium

Product:

AFIAS COVID-19 Aq

Cat. No.: SMFP-71

Classification:

Others (Neither listed in the annex II of the IVDD, Non-self-

testing device)

Conformity Assessment Route:

Self-Declaration Route According to the Annex III of the IVDD

We herewith declare that the above mentioned products meet the provisions of the Council Directive 98/79/EC for In Vitro Diagnostic medical devices. All supporting documentation is retained under the premises of the manufacturer and the manufacturer is exclusively responsible for the declaration of conformity.

Standards applied:

EN ISO 15223-1:2016, EN ISO 13485:2016, EN 13612:2002,

EN ISO 23640:2015, EN 13641:2002, EN ISO 14971:2012,

EN 13975:2003, EN ISO 17511:2003, EN ISO 18113-1:2011,

EN ISO 18113-2:2011

Place, Date of Issue:

Chuncheon, Korea, June 1, 2020

Signature:

Dr. Eui Yul Choi / CEO

Boditech Med Inc. www.boditech.co.kr

43, Geodudanji 1-gil, Dongnae-myeon, Chuncheon-si, Gang-won-do, 24398, Republic of Korea 바디텍메드(주) 강원도 춘천시 동내면 거두단지 1길 43 Tel +82-33-243-1400 Fax +82-33-243-9373 RA-DOC-II-165 (Rev. 00)

## **DECLARATION OF CONFORMITY**

Manufacturer:

Boditech Med Incorporated

43, Geodudanji 1-gil, Dongnae-myeon,

Chuncheon-si, Gang-won-do, 24398

REPUBLIC OF KOREA

European Representative:

**OBELIS S.A** 

Bd. Général Wahis 53,

1030 Brussels,

Belgium

Product:

Boditech COVID-19 Ag Control

Cat. No.: CFPO-293

Classification:

Others (Neither listed in the annex II of the IVDD, Non-self-testing

device)

Conformity Assessment Route:

Self-Declaration Route According to the Annex III of the IVDD

We herewith declare that the above mentioned products meet the provisions of the Council Directive 98/79/EC for In Vitro Diagnostic medical devices. All supporting documentation is retained under the premises of the manufacturer and the manufacturer is exclusively responsible for the declaration of conformity.

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Place, Date of Issue:

Chuncheon, Korea, June 1, 2020

Signature:

Dr. Eui Yul Choi / CEO

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