



To whom it may concern,

Budapest, June, 2020

LETTER OF AUTHORITY

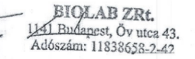
We **BIOLAB Inc** (Öv u. 43., H-1141 Budapest, Hungary) hereby certify that company **Mic-Tan** (C/f 1002600038282 Alexandru cel Bun 38 Chisinau Moldova) is authorized to register and distribute the products manufactured by **BIOLAB Inc** (Öv u. 43., H-1141 Budapest, Hungary) in **Moldova**.

In addition, Mic-Tan may present bids and make representations to official bodies in Moldova on behalf of BIOLAB Inc., in response to tenders.

This is valid until 31 December 2022.

For and on behalf of BIOLAB Inc.

Laszlo Ferenci General Manager





Certificate

Standard

ISO 9001:2015

Certificate Registr. No.

01 100 1824126

Certificate Holder:

BIOLAB Zrt. Öv u. 43. 1141 Budapest Hungary

Scope:

production and distribution of microbiological culture media and microbiological plastic disposables.

Proof has been furnished by means of an audit that the requirements of ISO 9001:2015 are met.

Validity:

The certificate is valid from 11-06-2018 until 10-06-2021.

11-06-2018

Negrid Sich

TÜV Rheinland Cert GmbH Am Grauen Stein 51105 Köln

Shedy !

«MIC-TAN»

TÜVRheinland Precisely Right.

DAKKS
Deutsche
Aktreditierungsstelle
D-2M-16031-01-00



1141 Budapest, Öv u. 43. Tel.: +36 1/221-96-14 Fax: +36 1/364-20-06

www.biolab.hu E-mail: biolab@biolab.hu

EC DECLARATION OF CONFORMITY

according to Annex III of the Directive 98/79/EC on "In Vitro Diagnostic Medical Devices"

Manufacturer:

BIOLAB Inc.

Address:

Öv u. 43., H-1141 Budapest

Phone:

+36 1 221 9614

Fax:

+36 1 364 2006

E-mail:

export@biolab.hu

Product identification: Microbiological culture media and supplements

Product classification: devices other than those mentioned in Annex II of

the Directive 98/79/EC

Hereby we declare

under our sole responsibility that the above mentioned devices meet the applicable provisions of the Directive 98/79/EC on "In Vitro Diagnostic Medical Devices".

All the supporting documents, as required by Annex III of the 98/79/EC Directive, in order to prove conformity to the Essential Requirements as listed in Annex I, are retained under premises of the Manufacturer.

Applicable standard: ISO 9001

Place and date: Budapest, 02. 03. 2019.

Signature:

László Ferenci Managing Director

BIOLAB ZRt.

1141 Budapest, Öv utca 43.

Adószám: 11838658-2-42







1141 Budapest, Öv u. 43. Tel.: +36 1/221-96-14 Fax: +36 1/364-20-06

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EC DECLARATION OF CONFORMITY

according to Annex III of the Directive 98/79/EC on "In Vitro Diagnostic Medical Devices"

Manufacturer:

BIOLAB Inc.

Address:

Öv u. 43., H-1141 Budapest

Phone:

+36 1 221 9614

Fax:

+36 1 364 2006

E-mail:

export@biolab.hu

Product identification: Antimicrobial susceptibility discs

Product classification: devices other than those mentioned in Annex II of

the Directive 98/79/EC

Hereby we declare

under our sole responsibility that the above mentioned devices meet the applicable provisions of the Directive 98/79/EC on "In Vitro Diagnostic Medical Devices".

All the supporting documents, as required by Annex III of the 98/79/EC Directive, in order to prove conformity to the Essential Requirements as listed in Annex I, are retained under premises of the Manufacturer.

Applicable standard: ISO 9001

Place and date: Budapest, 02.03. 2019.

Signature:

BIOLAB ZRt. 1141 Budapest, Öv utca 43. Adószám: 11838658-2-42





1141 Budapest, Öv u. 4 Tel.: +36 1/221-96-14 Fax: +36 1/364-20-0

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EC DECLARATION OF CONFORMITY

according to Annex III of the Directive 98/79/EC on "In Vitro Diagnostic Medical Devices"

Manufacturer: BIOLAB Inc.

Address:

Öv u. 43., H-1141 Budapest

Phone:

+36 1 221 9614

Fax:

+36 1 364 2006

E-mail:

export@biolab.hu

Product identification: Plastic disposables

Product classification: devices other than those mentioned in Annex II of

the Directive 98/79/EC

Hereby we declare

under our sole responsibility that the above mentioned devices meet the applicable provisions of the Directive 98/79/EC on "In Vitro Diagnostic Medical Devices".

All the supporting documents, as required by Annex III of the 98/79/EC Directive, in order to prove conformity to the Essential Requirements as listed in Annex I, are retained under premises of the Manufacturer.

Applicable standard: ISO 9001

Place and date: Budapest, 02.03. 2019.

Signature:

Adószám: 11838658-2-42

