

## 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:	<u>export@biolab.hu</u>

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





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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.

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 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

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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

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Applicable standard: ISO 9001

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