

3EC International a. s., Hraničná 18, 821 05 Bratislava, Slovak Republic Notified body No. 2265

### **EU QUALITY MANAGEMENT SYSTEM CERTIFICATE**

No. 2023-IVDR/QS-001

### TestLine Clinical Diagnostics s.r.o.

Registered place of business: Křižíkova 68, 612 00 Brno, Czech Republic Manufacturing sites:

1. Křižíkova 68, 612 00 Brno, Czech Republic

2. Karásek 1767/1, 621 00 Brno, Czech Republic

SRN No.: CZ-MF-000001803

This EU Quality Management System Certificate issued in accordance with the Regulation (EU) 2017/746 of the European Parliament and of the Council on in vitro diagnostic medical devices as amended confirms, that quality management system of in vitro diagnostic medical device:

INFECTIOUS DISEASES, In vitro diagnostic devices which require knowledge regarding immunoassays (EMDN W0105 + IVP 3007) (detailed list is stated in Annex I)

Intended purpose: Annex II

IVD MD class C

(detailed list is stated in the annex(es) if applicable)

meets the requirements on quality management system according to the Chapter I and III of Annex IX of the Regulation (EU) 2017/746 of the European Parliament and of the Council on in vitro diagnostic medical devices as amended.

Conditions for or limitations to the validity of the certificate: N/A

Validity of the certificate is conditional upon positive results of regular surveillance audits.

Notified body No. 2265 has performed assessment of the quality management system of the abovementioned in vitro diagnostic medical device and found that it meets the requirements stated above. The outcome of the assessment of the quality management system of the abovementioned in vitro diagnostic medical device is stated in the IVD MD Technical Documentation Assessment Report No. IVDR008\_2023 from 26.05.2023, IVD MD Performance Evaluation Assessment Report No. IVDR008\_2023 from 26.05.2023 and IVD MD Audit Report No. SK-0735-23 from 26.05.2023. Information on all examinations and tests performed is stated in the abovementioned reports and is available on request.

This **EU Quality Management System Certificate** applies only to the quality management system of the abovementioned in vitro diagnostic medical device. The certificate validity is conditional upon fulfilment of relevant legal requirements by the manufacturer.



Valid from: 01.10.2024 Valid until: 06.06.2028 First issue: 06.06.2023

Revision: 01
History: Annex III

NB 2265
NB 2265

3EC International a. s. Ing. Katarina Tomin Srdošová, PhD. Director of NB2265

In Bratislava, Slovakia, 01.10.2024



## ANNEX I TO EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2023-IVDR/QS-001

issued for the company

### TestLine Clinical Diagnostics s.r.o.

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List of in vitro diagnostic medical devices covered by the EU Quality Management System Certificate:

REF	Trade Name		
CMGMA48	Microblot-Array CMV IgG		
CMMMA48	Microblot-Array CMV IgM		
HSGMA48	Microblot-Array HSV 1+2 IgG		
HSMMA48	Microblot-Array HSV 1+2 IgM		
CL-PVG050	CLIA Parvovirus B19 IgG		
CL-PVM050	CLIA Parvovirus B19 IgM		
CL-HSVG100	CLIA HSV 1+2 IgG		
CL-HSVM100	CLIA HSV 1+2 IgM		
CL-VZVA100	CLIA VZV IgA		
CL-VZVG100	CLIA VZV IgG		
CL-VZVM100	CLIA VZV IgM		

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Katarina Tomin Srdošová, PhD. Director of NB 2265

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# ANNEX II TO EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2023-IVDR/QS-001

issued for the company

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Intended purpose of in vitro diagnostic medical devices covered by the EU Quality Management System Certificate:

**Microblot-Array CMV IgG**, **ref**. **CMGMA48**, Intended purpose: The Microblot-Array assay is intended for the diagnosis of CMV infection using IgG antibodies in human serum or plasma in the general population. The qualitative, semi-quantitative and quantitative automated assay is designed for professional use in a laboratory. The assay is not intended to assess the suitability for transfusion, transplantation or cell administration.

**Microblot-Array CMV IgM, ref. CMMMA48**, Intended purpose: The Microblot-Array assay is intended for the diagnosis of CMV infection using IgM antibodies in human serum or plasma in the general population. The qualitative, semi-quantitative and quantitative automated assay is designed for professional use in a laboratory. The assay is not intended to assess the suitability for transfusion, transplantation or cell administration.

**Microblot-Array HSV 1+2 IgG**, **ref**. **HSGMA48**, Intended purpose: The Microblot-Array assay is intended for the diagnosis of HSV infection using IgG antibodies in human serum or plasma in the general population. The qualitative, semi-quantitative and quantitative automated assay is designed for professional use in a laboratory.

**Microblot-Array HSV 1+2 IgM, ref. HSMMA48**, Intended purpose: The Microblot-Array assay is intended for the diagnosis of HSV infection using IgM antibodies in human serum or plasma in the general population. The qualitative, semi-quantitative and quantitative automated assay is designed for professional use in a laboratory.

**CLIA Parvovirus B19 IgG, ref. CL-PVG050**, Intended purpose: The chemiluminescence assay is intended for the diagnosis of parvovirus B19 infection using IgG antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.

**CLIA Parvovirus B19 IgM**, **ref**. **CL-PVM050**, Intended purpose: The chemiluminescence assay is intended for the diagnosis of parvovirus B19 infection using IgM antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.





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In Bratislava, Slovakia, 01.10.2024 Valid until 06.06.2028 Katarína Tomin Srdošová, PhD. Director of NB 2265



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Intended purpose of in vitro diagnostic medical devices covered by the EU Quality Management System Certificate:

**CLIA HSV 1+2 IgG**, **ref. CL-HSVG100**, Intended purpose: The chemiluminescence assay is intended for the diagnosis and screening of HSV infection using IgG antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.

**CLIA HSV 1+2 IgM, ref. CL-HSVM100**, Intended purpose: The chemiluminescence assay is intended for the diagnosis of HSV infection using IgM antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.

**CLIA VZV IgA**, **ref. CL-VZVA100**, Intended purpose: The chemiluminescence assay is intended for the diagnosis of VZV infection using IgA antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.

**CLIA VZV IgG, ref. CL-VZVG100**, Intended purpose: The chemiluminescence assay is intended for the diagnosis, monitoring and screening of VZV infection using IgG antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.

**CLIA VZV IgM, ref. CL-VZVM100,** Intended purpose: The chemiluminescence assay is intended for the diagnosis of VZV infection using IgM antibodies in human serum or plasma in the general population. The quantitative automated assay is designed for professional use in a laboratory.

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Katarína Tomin Srdošová, PhD.

Director of NB 2265

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## ANNEX III TO EU QUALITY MANAGEMENT SYSTEM CERTIFICATE No. 2023-IVDR/QS-001

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#### Certificate history:

Revision	EU QMS Certificate reference	Date of issue	Application for Conformity Assessment of IVD MD number	Description
00	2023-IVDR/QS-001	06.06.2023	IVDR008_2023 IVDR009_2023 IVDR010_2023	First issue
01	2023-IVDR/QS-001	01.10.2024	IVDR015_2024 IVDR016_2024	Scope extension

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Director of NB 2265

In Bratislava, Slovakia, 01.10.2024 Valid until 06.06.2028

International No 2206 SEC International NE 2266 SEC International NE 22