



EU Quality Management System Certificate

Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices, Annex IX Chapter I

Certificate No. V13 114209 0003 Rev. 00

Manufacturer:

VITRO, S.A.

Calle Luis Fuentes Bejarano, 60, Edificio Nudo Norte,
Local 3,
41020 Sevilla
SPAIN

SRN Manufacturer - ES-MF-000001498

The quality management system has been evaluated in accordance with Regulation (EU) 2017/746, Annex IX Chapter I with a positive result.

Details on devices covered by the quality management system are described on the following page(s). The report referenced below summarises the results of the assessment and includes reference to relevant CS, harmonised standards and test reports.

The certified quality management system is subject to periodical surveillance.

If class A devices in sterile conditions are covered by this certificate, the audit was limited to the aspects relating to establishing, securing, and maintaining sterile conditions.

If class B or C excluding self-/near-patient-testing, or class C companion diagnostics devices are covered by this certificate, the quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis. The periodical surveillance includes further assessment of the technical documentation on the basis of representative samples.

If class D devices, class B or C self-/near-patient testing, or class C companion diagnostics devices are covered by this certificate, an EU Technical Documentation Assessment Certificate in accordance with Annex IX Chapter II is required before placing them on the market.

All applicable requirements of the Testing, Certification, Validation and Verification Regulations TÜV SÜD Group have to be complied with.

For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:V13 114209 0003 Rev. 00

Report No.: ITA200220013641

Preceding Certificate No.: V12 114209 0001 Rev. 01

Valid from: 2026-01-26

Valid until: 2028-10-22

Marta Carnielli
Head of Certification IVD

Issue date: 2026-01-26



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Certificate No. V13 114209 0003 Rev. 00

Classification: Class C
Device Group: W0103 + IVP 3010 - Haematology / Haemostasis / Immunohaematology / Histology / Cytology
Intended Purpose: IVD Reagents for Haematology, Haemostasis, Immunohaematology, Histology and Cytology

Classification: Class C
Device Group: W0105 + IVP 3011 - Infectious diseases
Intended Purpose: IVD Reagents for Infectious diseases

Classification: Class C
Device Group: W0103 + IVP 3004 - Haematology / Haemostasis / Immunohaematology / Histology / Cytology
Intended Purpose: IVD Reagents for Haematology, Haemostasis, Immunohaematology, Histology and Cytology

Classification: Class C
Device Group: W0103 + IVP 3011 - Haematology / Haemostasis / Immunohaematology / Histology / Cytology
Intended Purpose: IVD Reagents for Haematology, Haemostasis, Immunohaematology, Histology and Cytology

The validity of this certificate
depends on conditions and/or
is limited to the following:

Revision History:

Rev.	Dated	Report	Description
00	2026-01-26	ITA200220013641	Supplemented: Device(s)/group of device(s) added Administrative merge / transfer to new Certificate Type