





Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 9001:2015

This is to certify that:

Abbott Informatics Corporation

4000 Hollywood Blvd

Suite 333 - S South Hollywood

Florida 33021 USA

Holds Certificate No: FM 636368

and operates a Quality Management System which complies with the requirements of ISO 9001:2015 for the following scope:

Design, manufacture, distribution, installation and servicing of Laboratory Information Management Systems software for various industries.

For and on behalf of BSI:

Original Registration Date: 2016-05-20 Latest Revision Date: 2021-04-22





Carlos Pitanga, Chief Operating Officer Assurance – Americas

Effective Date: 2021-06-26 Expiry Date: 2024-06-25

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Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

Abbott Informatics Corporation

4000 Hollywood Blvd

Suite 333 - S South Hollywood

Florida 33021 USA

Holds Certificate No: FM 636367

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 for the following scope:

Design, manufacture, distribution, installation and servicing of Laboratory Information Management Systems software for the medical device industry.

For and on behalf of BSI:

Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2016-05-20 Effective Date: 2021-06-26 Latest Revision Date: 2021-04-22 Expiry Date: 2024-06-25

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EC Certificate

Full Quality Assurance System Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6) (List A and B and devices for self-testing)

No. V1 103108 0003 Rev. 02

Manufacturer: Abbott Diagnostics Medical Co., Ltd.

> 357 Matsuhidai Matsudo-shi, Chiba 270-2214 JAPAN

Product Category(ies): Products for determination of infection

markers

HIV-1/-2 marker and Hepatitis B marker

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device families in accordance with IVDD Annex IV. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of List A devices an additional Annex IV (4) certificate is mandatory. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:V1 103108 0003 Rev. 02

Report no.: JN1648745

Valid from: 2021-03-18 Valid until: 2024-05-26

Date, 2021-03-03

> Christoph Dicks Head of Certification/Notified Body



EC Certificate

Full Quality Assurance System Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6) (List A and B and devices for self-testing)

No. V1 103108 0003 Rev. 02

Model(s): **Not Applicable**

Abbott Diagnostics Medical Co., Ltd. Chiba Plant Facility(ies): 357 Matsuhidai, Matsudo-shi, Chiba, 270-2214 JAPAN

Abbott Diagnostics Medical Co., Ltd. Chiba Logistic Center

483-2 Matsuhidai, Matsudo-shi, Chiba, 270-2214 JAPAN

-/-