

Current issue date:
Expiry date:
Certificate identity number

1 October 2021 30 September 2024 10393989 Original approval: ISO 13485 - 23 September 1994

# **Certificate of Approval**

This is to certify that the Management System of:

# Abbott GmbH & Abbott Diagnostics GmbH

Max-Planck-Ring 2, 65205 Wiesbaden, Germany

has been approved by Lloyd's Register to the following standards:

### EN ISO 13485:2016 | ISO 13485:2016

Approval number: ISO 13485 - 00004790

### The scope of this approval is applicable to:

Design, development, manufacture, control of contract manufacturers, registration, stockholding and distribution of in-vitro diagnostic devices.

David 1

David Derrick Area Operations Manager UK & Ireland

Issued by: Lloyd's Register Quality Assurance Limited



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Current issue date:
Expiry date:
Certificate identity number

1 October 2021 30 September 2024 10393990 Original approval: ISO 9001 - 23 September 1994

# **Certificate of Approval**

This is to certify that the Management System of:

# Abbott GmbH & Abbott Diagnostics GmbH

Max-Planck-Ring 2, 65205 Wiesbaden, Germany

has been approved by Lloyd's Register to the following standards:

### ISO 9001:2015

Approval number: ISO 9001 - 00004791

The scope of this approval is applicable to:

Design, development, manufacture, control of contract manufacturers, registration, stockholding and distribution of in-vitro diagnostic devices.

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David Derrick Area Operations Manager UK & Ireland Issued by: Lloyd's Register Quality Assurance Limited



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Certificate approval number:
Effective date:
Expiry date:
Certificate issue number:

LRQ0925480 2021 October 1 2024 September 30 10393988 Original approval: MDSAP/ISO 13485 - 2018 October 1

# **Certificate of Approval**

This is to certify that the Management System of:

# **Abbott GmbH**

Max-Planck-Ring 2, 65205 Wiesbaden, Germany

MDSAP Facility Identifier: F003705

has been audited by Lloyd's Register Quality Assurance and found to conform to the following audit criteria:

### ISO 13485:2016

#### Australia:

Therapeutic Goods (Medical Devices) Regulations, 2002, Schedule 3 Part 1 (Excluding Part 1.6) – Full Quality Assurance Procedure

#### Brazil:

RDC ANVISA n. 16/2013 RDC ANVISA n. 23/2012 RDC ANVISA n. 67/2009

#### Canada:

Medical Devices Regulations – Part 1- SOR 98/282 Japan:

MHLW Ministerial Ordinance 169, Article 4 to Article 68 PMD Act

### United States:

21 CFR 803 21 CFR 806 21 CFR 807 – Subparts A to D 21 CFR 820

Approval number: MDSAP - 0079011

#### The scope of this approval is applicable to:

Design, development, manufacture, control of contract manufacturers, registration, stockholding and distribution of in-vitro diagnostic devices.

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David Derrick Area Operations Manager UK & Ireland

Issued by: Lloyd's Register Quality Assurance Limited



MEDICAL DEVICE SINGLE AUDIT PROGRAM Lloyd's Register Quality Assurance Limited is an MDSAP authorised auditing organization.

To validate certificate authenticity visit: http://www.lrqausa.com/help-and-support/Request-for-certificate-verification

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<b>Certificate Identification:</b>	02K47 LC	IRIS V4	
Legal Manufacturer's Name:	Abbott Laboratories		
	Diagnostics Division		
Legal Manufacturer's Address:	Abbott Park, IL 60064 USA		

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
2K47-20	58729	ARCHITECT Anti-TPO Reagent Kit	Self-declared
2K47-25	58729	ARCHITECT Anti-TPO Reagent Kit	Self-declared
2K47-27	58729	ARCHITECT Anti-TPO Reagent Kit	Self-declared
2K47-01	55210	ARCHITECT Anti-TPO Calibrators	Self-declared
2K47-10	55211	ARCHITECT Anti-TPO Controls	Self-declared

Authorized European	Abbott GmbH
Representative	Max-Planck-Ring-2
(Name and Address)	65205 Wiesbaden, Germany
Storage Site of Technical	Fisher Diagnostics
Documentation	a division of Fisher Scientific Company LLC
(Name and Address)	a part of Thermo Fisher Scientific Inc.
	8365 Valley Pike, Middletown, VA 22645-1905
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states. This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the

## manufacturer.

Signature:

Full Name: Elizabeth Wernquist Position: Director QA, LC Site

Date of Approval:

8 November 2021 Date Issued: \_\_\_\_

Date of Approval:

8 November 2021

Associate Director Regulatory Affairs

Place Issued:

Signature:

Full Name:

Position:

Abbott Laboratories Diagnostics Division Abbott Park, IL 60064 U.S.A.

Jacek Gorzowski

Effective (Date or Lot Number):

8 November 2021

27 OCT 2021



**Certificate Identification:** Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-7K71- AIDD Sligo Abbott Ireland Diagnostics Division Finisklin Business Park, Sligo, Ireland

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7K71-20	54669	ARCHITECT Free PSA Reagent Kit	Annex II List B
7K71-25	54669	ARCHITECT Free PSA Reagent Kit	Annex II List B
7K71-01	38183	ARCHITECT Free PSA Calibrators	Annex II List B
7K71-10	38182	ARCHITECT Free PSA Controls	Annex II List B

Authorized European	N/A
Representative (name and address)	
Notified Body (name and address)	TÜV SÜD Product Service GmbH
	Ridlerstraße 65
	80339 Munich
	Germany
Notified Body number	0123
Approval Certificate No.	V1 0019220008
Storage site of technical	Abbott Ireland Diagnostics Division, Finisklin Business Park, Sligo, Ireland.
documentation (name and address)	Department: Regulatory Affairs.
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex IV of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature: Full Name:

Joe Murray Director Quality Assurance/Site

Position:

Quality Head 20 Nov 19 NUJ 2019

Date of Approval: Date Issued:

Supersedes:

14 October 2019

Position:

Signature:

Full Name:

Manager Regulatory Affairs

19 Nov 2019 Date of Approval:

Noel Haren

Place Issued:

AIDD, Sligo

Effective (Date or Lot Number):

20 NON 2019



Certificate Identification:	7K63	
Legal Manufacturer's Name:	Abbott Ireland Diagnostics Division	
Legal Manufacturer'sAddress:	Lisnamuck, Longford, Co. Longford, Ireland	

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7K63-27	54417	ARCHITECT Free T <sub>3</sub> Reagent Kit	Self-declared
7K63-32			
7K63-37			
7K63-02	38261	ARCHITECT Free T <sub>3</sub> Calibrators	Self-declared
7K63-12	54418	ARCHITECT Free T <sub>3</sub> Controls	Self-declared
Authorized Euro Representative (Name and Add		N/A	
Storage of technical documentation (Name and Address)		Abbott Ireland Diagnostics Division, Lisnamuck, Lo Ireland	ngford, Co. Longford,
Harmonized Standards		Listed in the Technical Documentation	

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:

Full Name: Siobhan Wright

Position: **Director Quality Assurance/** 

Sohan

Approval:

Supersedes:

01-MAY-2020

24-April-2019

01- MAY-2020

Date Issued:

Place Issued:

Date of Approval:

Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, Ireland.

Romaine Chihey

Effective (Date or Lot Number):

Signature:

Position:

Full Name: Lorraine Whitney

Senior Manager

**Regulatory Affairs** 

OI MAY 2020

01 - MAY - 2020

Site Quality Head

Date of



<b>Certificate Identification:</b>	7K65-22/-24/-27/-29/-32/-34/-35/-39, 7K65-02, 7K65-10
Legal Manufacturer's Name:	Abbott Ireland Diagnostics Division
Legal Manufacturer's Address:	Lisnamuck, Longford
	Co. Longford

Ireland

List Numbers and Size Code of Devices	GMDN Code		Names and Description of Devices	Classification
7K65-22 7K65-24 7K65-27 7K65-29 7K65-32 7K65-34 7K65-35 7K65-39	54413	ARG	CHITECT Free T4 Reagent Kit	Self-declared
7K65-02	38259	ARG	CHITECT Free T4 Calibrators	Self-declared
7K65-10	38258	ARG	CHITECT Free T4 Controls	Self-declared
Authorized European Representative N. (Name and Address)			N/A	
	Storage of site technical documentation (Name and Address)Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, I		ngford, Co. Longford, Ireland.	
Harmonized Standards Listed in the Technical Documentation				

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

Signature: Full Name:	Litchan Wight Siobhan Wright	Signature: Full Name:	<u>p.p. SANDRA GALLAGHER</u> Scalles les Lorraine Whitney
Position:	Director Quality Assurance/ Site Quality Head	Position:	Senior Manager Regulatory Affairs
Date of Approval:	29-1APR-19	Date of Approval:	25-19PR-2019.
Date Issued:	29-14/2-19	Place Issued:	Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, Ireland
Supersedes:	26-July-2017	Effective (Date or Lot Number):	29-AP.L-19



To Whom it may concern

I will be out of office Tues 23rd to Fri 26th April 19.

My signature during this time is delegated to Noel Haren and Sandra Gallagher.

poward Whitey 19 APR 2019

Lorraine Whitney Senior Manager Regulatory Affairs Site Operations Ireland



Abbott

**Certificate Identification:** Legal Manufacturer's Name:

Legal Manufacturer's Address:

DoC 8L44 AII DELK Abbott GmbH

Max-Planck-Ring 2, 65205 Wiesbaden, Germany

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
8L44-25	48304	ARCHITECT Anti-HBc II Reagent Kit (1x100 Tests)	Annex II List A
8L44-30	48304	ARCHITECT Anti-HBc II Reagent Kit (4x500 Tests)	Annex II List A
8L44-35	48304	ARCHITECT Anti-HBc II Reagent Kit (1x500 Tests)	Annex II List A
8L44-01	41983	ARCHITECT Anti-HBc II Calibrator	Annex II List A
8L44-10	41984	ARCHITECT Anti-HBc II Controls	Annex II List A

Authorized European	N/A
Representative (name and address)	
Notified Body (name and address)	TÜV SÜD Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany
Notified Body number	TÜV SÜD: 0123
Approval Certificate No.	TÜV SÜD: V7 010051 0130
Storage site of technical	Abbott GmbH, Max-Planck-Ring 2, 65205 Wiesbaden, Germany
documentation (name and address)	
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex IV of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:

Full Name:

**Claudia Becker** 

Position: **Director Quality Assurance** 

Date of Approval:

06 May 2021

Sula

Signature:

Full Name:

Manh

Susanne Ulrich

Assoc. Director Regulatory Affairs

1021 06- May-2021

65205 Wiesbaden, Germany

09-Mar-2020

Supersedes:

Place Issued:

Effective (Date or Lot Number):

06- May - 2021

Position:

Date of Approval:

Date Issued:







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 001922 0008 Rev. 03

Manufacturer:

## Abbott Ireland Diagnostics Division

Finisklin Business Park Sligo IRELAND

## Product Category(ies): Products for determination of infection markers and tumour markers

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. See also notes overleaf.

Report no.:

713158801-03

Valid from: Valid until: 2020-01-15 2024-05-26

Date,

œ

111 N 2020-01-15

**Christoph Dicks** Head of Certification/Notified Body

A4 / 07.17





# **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 001922 0008 Rev. 03

Model(s):

### Products for the determination of infection markers for Hepatitis B, cytomegalovirus, rubella and tumour marker PSA

REF N°

Facility(ies):

Abbott Ireland Diagnostics Division Finisklin Business Park, Sligo, IRELAND

The products detailed below are covered under the scope of this certificate:

### Annex II List A Products

Due duet Mana

Product Name	REFIN
ARCHITECT HBsAg Qualitative II Calibrators	2G22-01
ARCHITECT HBsAg Qualitative II Reagent Kit	2G22-25
ARCHITECT HBsAg Qualitative II Reagent Kit	2G22-30
ARCHITECT HBsAg Qualitative II Confirmatory Reagent Kit	2G23-25
ARCHITECT HBsAg Calibrators	3M61-01
ARCHITECT HBsAg Calibrators	3M61-02
ARCHITECT HBsAg Controls	6C36-10
ARCHITECT HBsAg Reagent Kit	6C36-22
ARCHITECT HBsAg Reagent Kit	6C36-27
ARCHITECT HBsAg Reagent Kit	6C36-32
ARCHITECT HBsAg Reagent Kit	6C36-29
ARCHITECT HBsAg Reagent Kit	6C36-34
ARCHITECT HBsAg Reagent Kit	6C36-35
ARCHITECT HBsAg Reagent Kit	6C36-43
ARCHITECT HBsAg Reagent Kit	6C36-44
ARCHITECT HBsAg Reagent Kit	6C36-41
ARCHITECT HBsAg Reagent Kit	6C36-42
ARCHITECT Anti-HBs Calibrators	7C18-01
ARCHITECT Anti-HBs Calibrators	7C18-03
ARCHITECT Anti-HBs Controls	7C18-10
ARCHITECT Anti-HBs Controls	7C18-13
ARCHITECT Anti-HBs Reagent Kit	7C18-20
ARCHITECT Anti-HBs Reagent Kit	7C18-25

Page 2 of 6 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

A4 / 07.17





# **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 001922 0008 Rev. 03

### Annex II List A Products

Product Name	REF N°
ARCHITECT Anti-HBs Reagent Kit	7C18-27
ARCHITECT Anti-HBs Reagent Kit	7C18-28
ARCHITECT Anti-HBs Reagent Kit	7C18-30
ARCHITECT Anti-HBs Reagent Kit	7C18-34
ARCHITECT Anti-HBs Reagent Kit	7C18-37
ARCHITECT Anti-HBs Reagent kit	7C18-38
ARCHITECT HBsAg Confirmatory V.1 Calibrators	9C94-01
ARCHITECT HBsAg Confirmatory V.1 Controls	9C94-10
ARCHITECT HBsAg Confirmatory V.1 Reagent Kit	9C94-25
ARCHITECT HBsAg Qualitative II Reagent Kit	2G22-35
ARCHITECT Anti-HBs Reagent Kit	7C18-29
ARCHITECT Anti-HBs Reagent Kit	7C18-41
ARCHITECT Anti-HBs Reagent Kit	7C18-39
ARCHITECT Anti-HBs Reagent Kit	7C18-42
ARCHITECT Anti-HBs Reagent Kit	7C18-33
Alinity i HBsAg Calibrators	08P0801
Alinity i HBsAg Controls	08P0810
Alinity i HBsAg Reagent Kit	08P0852
Alinity i HBsAg Confirmatory V.1 Calibrators	08P0901
Alinity i HBsAg Confirmatory V.1 Controls	08P0910
Alinity i HBsAg Confirmatory V.1 Reagent Kit	08P0922
Alinity i HBsAg Qualitative II Calibrators	08P1001
Alinity i HBsAg Qualitative II Controls	08P1010

A4 / 07.17





# **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 001922 0008 Rev. 03

### Annex II List A Products

Product Name	REF N°
Alinity i HBsAg Qualitative II Reagent Kit	08P1022
Alinity i HBsAg Qualitative II Confirmatory Reagent Kit	08P1122
Alinity i Anti-HBs Reagent Kit	07P8922
Alinity i Anti-HBs Controls	07P8910
Alinity i Anti-HBs Calibrators	07P8901
Alinity i Anti-HBs Reagent Kit	07P8952
Alinity s HBsAg Reagent Kit	06P0255
Alinity s HBsAg Reagent Kit	06P0260
Alinity s HBsAg Confirmatory Reagent Kit	06P0357
Alinity s HBsAg Confirmatory Reagent Kit	06P0359
Alinity s HBsAg Calibrator Kit	06P0202
Alinity s HBSAg Calibrator Kit	06P0204
Alinity s HBsAg Assay Control Kit	06P0210
Alinity s HBsAg Assay Control Kit	06P0213
Alinity s HBsAg Release Control Kit	06P0212
Alinity s HBsAg Release Control Kit	06P0215
ARCHITECT HBsAg Qualitative II Controls	2G22-10
Alinity i HBsAg Qualitative II Reagent Kit	08P1032
Alinity i HBsAg Reagent Kit	08P0832
Alinity i HBsAg Reagent Kit	08P0822
Alinity i HBsAg Reagent Kit	08P0857
Alinity i Anti-HBs Reagent Kit	07P8932
Alinity i Anti-HBs Reagent Kit	07P8957
Alinity i HBsAg Next Qualitative Calibrators	01R6401
Alinity i HBsAg Next Qualitative Controls	01R6410
Alinity i HBsAg Next Qualitative Reagent Kit	01R6422
Alinity i HBsAg Next Qualitative Reagent Kit	01R6432
Alinity i HBsAg Next Confirmatory Reagent Kit	01R6522
ARCHITECT HBsAg Next Qualitative Reagent Kit	4P76-25
ARCHITECT HBsAg Next Qualitative Reagent Kit	4P76-30
ARCHITECT HBsAg Next Qualitative Reagent Kit	4P76-35
ARCHITECT HBsAg Next Confirmatory Reagent Kit	4P77-25
ARCHITECT HBsAg Next Qualitative Calibrators	4P76-01
ARCHITECT HBsAg Next Qualitative Controls	4P76-10

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TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

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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 001922 0008 Rev. 03

### Annex II List B Products

REF N° Product Name 6C18-25 ARCHITECT Rubella IgM Reagent Kit 6C18-01 ARCHITECT Rubella IgM Calibrator 6C18-10 ARCHITECT Rubella IgM Controls 6C17-26/36 ARCHITECT Rubella IgG Reagent Kit 6C17-03 **ARCHITECT Rubella IgG Calibrators ARCHITECT Rubella IgG Controls** 6C17-13 7K71-20/25 **ARCHITECT Free PSA Reagent Kit** 7K71-01 **ARCHITECT Free PSA Calibrators** 7K71-10 **ARCHITECT Free PSA Controls** 7K70-20/25/30/35 **ARCHITECT Total PSA Reagent Kit** 7K70-01 **ARCHITECT Total PSA Calibrators ARCHITECT Total PSA Controls** 7K70-10 3L46-25 ARCHITECT CMV IgG Avidity Reagent Kit 3L46-11 ARCHITECT CMV IgG Avidity Calibrator and Controls 6C15-20/25/30 ARCHITECT CMV IgG Reagent Kit 6C15-01 **ARCHITECT CMV IgG Calibrators** 6C15-10 ARCHITECT CMV IgG Controls 6C16-20/25/30 ARCHITECT CMV IgM Reagent Kit 6C16-01 ARCHITECT CMV IgM Calibrator 6C16-10 ARCHITECT CMV IgM Controls 07P4222 / 07P4232 Alinity i CMV IgG Reagent Kit 07P4201 Alinity i CMV IgG Calibrators

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**REF N°** 

# **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)

www.złg.de

### No. V1 001922 0008 Rev. 03

### Annex II List B Products

Product Name

Alinity i CMV IgG Controls	07 <b>P42</b> 10
Alinity i CMV IgM Reagent Kit	07P4422 / 07P4432
Alinity i CMV IgM Calibrator	07P4401
Alinity i CMV IgM Controls	07P4410
Alinity i Rubella IgG Reagent Kit	08P4622 / 08P4632
Alinity i Rubella IgG Calibrators	08P4601
Alinity i Rubella IgG Controls	08P4610
Alinity i Rubella IgM Reagent Kit	08P4722 / 08P4732
Alinity i Rubella IgM Calibrator	08P4701
Alinity i Rubella IgM Controls	08P4710
Alinity i CMV IgG Avidity Reagent Kit	07P4322
Alinity i CMV IgG Avidity Controls	07P4310
Alinity s CMV IgG Qualitative Reagent Kit	06P1045
Alinity s CMV IgG Qualitative Calibrator Kit	06P1002
Alinity s CMV IgG Qualitative Assay Control Kit	06P1010
Alinity s CMV IgG Qualitative Release Control Kit	06P1012
Alinity i Free PSA Reagent Kit	07P9320 / 07P9330
Alinity i Free PSA Calibrators	07P9301
Alinity i Free PSA Controls	07P9310
Alinity i Total PSA Reagent Kit	07P9220 / 07P9230
Alinity i Total PSA Calibrators	07P9201
Alinity i Total PSA Controls	07P9210

Page 6 of 6 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

TÜV SÜD Product Service GmbH • Certification Body • Ridlerstraße 65 • 80339 Munich • Germany



Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-2G22-AIDD Sligo Abbott Ireland Diagnostics Division Finisklin Business Park, Sligo, Ireland

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
2G22-25	48321	ARCHITECT HBsAg Qualitative II Reagent Kit	Annex II List A
2G22-30	48321	ARCHITECT HBsAg Qualitative II Reagent Kit	Annex II List A
2G22-35	48321	ARCHITECT HBsAg Qualitative II Reagent Kit	Annex II List A
2G22-01	41999	ARCHITECT HBsAg Qualitative II Calibrators	Annex II List A
2G22-10	42000	ARCHITECT HBsAg Qualitative II Controls	Annex II List A

Authorized European	N/A
Representative (name and address)	
Notified Body (name and address)	TÜV SÜD Product Service GmbH
	Ridlerstraße 65
	80339 Munich
	Germany
Notified Body number	0123
Approval Certificate No.	V7 0019220009
Storage site of technical	Abbott Ireland Diagnostics Division, Finisklin Business Park, Sligo, Ireland.
documentation (name and address)	Department: Regulatory Affairs.
Harmonized Standards	Listed in the Technical Documentation

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This declaration is made in accordance with Annex IV of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature: Signature: Full Name: Full Name: Noel Haren Joe Murray Director Quality Assurance/Site Position: Position: Manager Regulatory Affairs Quality Head 27 Oct 2020 27 001 2020 Date of Approval: Date of Approval: 7 005 2020 Place Issued: AIDD, Sligo Date Issued: Effective (Date or 27 OCT 2020 Supersedes: 25 November 2019

Lot Number):







EC Design-Examination Certificate Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV (4) (List A)

## No. V7 001922 0009 Rev. 01

Manufacturer:	Abbott Ireland Diagnostics Division Finisklin Business Park Sligo IRELAND	
Product:	Screening and Confirmatory Test for Hepatitis B marker	
Model(s):	ARCHITECT HBsAg Qualitative II ARCHITECT HBsAg Qualitative II Confirmato	ry
Parameters:	Product Name	REF N°
	ARCHITECT HBsAg Qualitative II Reagent Kit	2G22-25
	ARCHITECT HBsAg Qualitative II Reagent Kit	2G22-30
	ARCHITECT HBsAg Qualitative II Reagent Kit	2G22-35
	ARCHITECT HBsAg Qualitative II Calibrators	2G22-01
	ARCHITECT HBsAg Qualitative II Controls	2G22-10
	ARCHITECT HBsAg Qualitative II Confirmatory Reagent Kit	2G23-25

The Certification Body of TÜV SÜD Product Service GmbH declares that a design examination has been carried out on the respective devices in accordance with IVDD Annex IV (4). The design of the devices conforms to the requirements of this Directive. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: <a href="https://www.tuvsud.com/ps-cert?q=cert:V7\_001922\_0009\_Rev.01">www.tuvsud.com/ps-cert?q=cert:V7\_001922\_0009\_Rev.01</a>

Report No.:

713190856-2

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

Date, 202

2021-05-25

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Christoph Dicks Head of Certification/Notified Body

Page 1 of 1 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

### Abbott

## **Declaration of Conformity**

Certificate Identification:

DOC-6C37-28/-33/-38-AII DLK

Legal Manufacturer's Name:

Abbott GmbH

Legal Manufacturer's Address: Max-

Max-Planck-Ring 2, 65205 Wiesbaden, Germany

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
6C37-28	48366	ARCHITECT Anti-HCV Reagent Kit (1 x 100 Tests)	Annex II List A
6C37-33	48366	ARCHITECT Anti-HCV Reagent Kit (4 x 500 Tests)	Annex II List A
6C37-38	48366	ARCHITECT Anti-HCV Reagent Kit (1 x 500 Tests)	Annex II List A
6C37-02	41972	ARCHITECT Anti-HCV Calibrator	Annex II List A
6C37-15	41973	ARCHITECT Anti-HCV Controls	Annex II List A

Authorized European Representative (name and address)	N/A
Notified Body (name and address)	TÜV SÜD Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany
Notified Body number	0123
Approval Certificate No.	V7 010051 0132
Storage site of technical documentation (name and address)	Abbott GmbH, Max-Planck-Ring 2, 65205 Wiesbaden, Germany
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex IV of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:

Signature:

Full Name: Herbert Hartmann

Full Name:

Position:

Date of Approval:

Manager Quality Systems

Position: Date of

Approval:

Date Issued: Place Issued:

Supersedes:

Effective (Date or Lot Number):

S. Ula

Stefan Veber

Manager Regulatory Affairs

2021-08-03

2021-08-03

65205 Wiesbaden, Germany 06-May-2021

2021-08-03









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)

### V1 010051 0103 Rev. 10

Manufacturer:

### Abbott GmbH

Max-Planck-Ring 2 65205 Wiesbaden GERMANY

### Product Category(ies): Products for determination of infection markers

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 010051 0103 Rev. 10 10

Report no.:	713215031

Valid from:	2021-08-20
Valid until:	2024-05-26

Date,

2021-08-20

Christoph Dicks Head of Certification/Notified Body







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)

## V1 010051 0103 Rev. 10

Model(s):

# Products for the determination of infection markers for HIV, Hepatitis B, Hepatitis C, HTLV, toxoplasmosis

Facility(ies):

Abbott GmbH Max-Planck-Ring 2, 65205 Wiesbaden, GERMANY







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)

## V1 010051 0103 Rev. 10

### The products detailed below are covered under the scope of this certificate:

Annex II List A Products	
Product Name	REF N°
ABBOTT PRISM HIV O Plus Assay Kit	3D34-48
ABBOTT PRISM HBsAg Assay Kit	3A47-48
ABBOTT PRISM HCV Assay Kit	6A52-48
ABBOTT PRISM HTLV-I/HTLV-II Assay Kit	6A53-48
ABBOTT PRISM Positive Run Control Kit	5E22-11
ABBOTT PRISM Run Control Kit	5E22-10
ABBOTT PRISM HBcore Assay Kit	1A77-48
ABBOTT PRISM HBsAg Confirmatory Assay Kit	6D16-48
ABBOTT PRISM HIV Ag/Ab Combo Assay Kit	7G46-48
ABBOTT PRISM Run Control Kit	2K24-10
ABBOTT PRISM Positive Run Control Kit	2K24-11
ARCHITECT Anti-HCV Reagent Kit	6C37-22
ARCHITECT Anti-HCV Reagent Kit	6C37-27
ARCHITECT Anti-HCV Reagent Kit	6C37-32
ARCHITECT Anti-HCV Reagent Kit	6C37-37
ARCHITECT Anti-HCV Calibrator	6C37-01
ARCHITECT Anti-HCV Controls	6C37-10
ARCHITECT Anti-HCV Reagent Kit	6C37-28
ARCHITECT Anti-HCV Reagent Kit	6C37-33
ARCHITECT Anti-HCV Reagent Kit	6C37-38
ARCHITECT Anti-HCV Calibrator	6C37-02
ARCHITECT Anti-HCV Controls	6C37-15
ARCHITECT Anti-HBc IgM Reagent Kit	6C33-22
ARCHITECT Anti-HBc IgM Reagent Kit	6C33-27
ARCHITECT Anti-HBc IgM Calibrators	6C33-02
ARCHITECT Anti-HBc IgM Controls	6C33-11
ARCHITECT Anti-HBe Reagent Kit	6C34-20
ARCHITECT Anti-HBe Reagent Kit	6C34-25
ARCHITECT Anti-HBe Reagent Kit	6C34-35
ARCHITECT Anti-HBe Calibrator	6C34-01
ARCHITECT Anti-HBe Controls	6C34-10

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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)

www.zlg.

## V1 010051 0103 Rev. 10

Annex II List A Products	
Product Name	REF N°
ARCHITECT HBeAg Reagent Kit	6C32-20
ARCHITECT HBeAg Reagent Kit	6C32-25
ARCHITECT HBeAg Reagent Kit	6C32-27
ARCHITECT HBeAg Reagent Kit	6C32-37
ARCHITECT HBeAg Calibrators	6C32-01
ARCHITECT HBeAg Quantitative Calibrators	7P24-01
ARCHITECT HBeAg Controls	6C32-10
ARCHITECT HBeAg Quantitative Controls	7P24-10
ARCHITECT HIV Ag/Ab Combo Reagent Kit	4J27-22
ARCHITECT HIV Ag/Ab Combo Reagent Kit	4J27-27
ARCHITECT HIV Ag/Ab Combo Reagent Kit	4J27-32
ARCHITECT HIV Ag/Ab Combo Reagent Kit	4J27-37
ARCHITECT HIV Ag/Ab Combo Calibrator	4J27-03
ARCHITECT HIV Ag/Ab Combo Controls	4J27-12
ARCHITECT rHTLV I/II Reagent Kit	6L61-25
ARCHITECT rHTLV I/II Reagent Kit	6L61-30
ARCHITECT rHTLV I/II Reagent Kit	6L61-35
ARCHITECT rHTLV I/II Calibrator	6L61-01
ARCHITECT rHTLV I/II Controls	6L61-10
ARCHITECT Anti-HBc II Reagent Kit	8L44-25
ARCHITECT Anti-HBc II Reagent Kit	8L44-30
ARCHITECT Anti-HBc II Reagent Kit	8L44-35
ARCHITECT Anti-HBc II Calibrator	8L44-01
ARCHITECT Anti-HBc II Controls	8L44-10
ARCHITECT HCV Ag Controls	6L47-11
ARCHITECT HCV Ag Controls	6L47-19
ARCHITECT HCV Ag Calibrators	6L47-02
ARCHITECT HCV Ag Calibrators	6L47-09
ARCHITECT HCV Ag Reagent Kit	6L47-29
ARCHITECT HCV Ag Reagent Kit	6L47-74

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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)

### V1 010051 0103 Rev. 10

Annex II List A Products	
Product Name	REF N°
Alinity i Anti-HBe Reagent Kit	07P6322
Alinity i Anti-HBe Reagent Kit	07P6332
Alinity i Anti-HBe Calibrator	07P6301
Alinity i Anti-HBe Controls	07P6310
Alinity i HIV Ag/Ab Combo Reagent Kit	08P0722
Alinity i HIV Ag/Ab Combo Reagent Kit	08P0732
Alinity i HIV Ag/Ab Combo Calibrator	08P0701
Alinity i HIV Ag/Ab Combo Controls	08P0710
Alinity i Anti-HCV Reagent Kit	08P0622
Alinity i Anti-HCV Reagent Kit	08P0632
Alinity i Anti-HCV Calibrator	08P0601
Alinity i Anti-HCV Controls	08P0610
Alinity i Anti-HCV Reagent Kit	08P0623
Alinity i Anti-HCV Reagent Kit	08P0633
Alinity i Anti-HCV Calibrator	08P0602
Alinity i Anti-HCV Controls	08P0611
Alinity i Anti-HBc IgM Reagent Kit	07P8622
Alinity i Anti-HBc IgM Calibrators	07P8601
Alinity i Anti-HBc IgM Controls	07P8610
Alinity i Anti-HBc II Reagent Kit	07P8722
Alinity i Anti-HBc II Reagent Kit	07P8732
Alinity i Anti-HBc II Calibrator	07P8701
Alinity i Anti-HBc II Controls	07P8710
Alinity i HCV Ag Reagent Kit	09P2322
Alinity i HCV Ag Controls	09P2310
Alinity i HCV Ag Calibrators	09P2301

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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)

www.zlg.

### V1 010051 0103 Rev. 10

Annex II List A Products	
Product Name	REF N°
	07P6122
Alinity i rHTLV-I/II Reagent Kit	0.1.0.122
Alinity i rHTLV-I/II Reagent Kit	07P6132
Alinity i rHTLV-I/II Calibrator	07P6101
Alinity i rHTLV-I/II Controls	07P6110
Alinity i HBeAg Reagent Kit	07P6422
Alinity i HBeAg Reagent Kit	07P6432
Alinity i HBeAg Calibrators	07P6401
Alinity i HBeAg Controls	07P6410
Alinity i HBeAg Quantitative Calibrators	09P1001
Alinity i HBeAg Quantitative Controls	09P1010
Alinity s Anti-HBc Reagent Kit	06P0655
Alinity s Anti-HBc Calibrator Kit	06P0602
Alinity s Anti-HBc Assay Control Kit	06P0610
Alinity s Anti-HBc Release Control Kit	06P0612
Alinity s HIV Ag/Ab Combo Reagent Kit	06P0155
Alinity s HIV Ag/Ab Combo Calibrator Kit	06P0102
Alinity s HIV Ag/Ab Combo Assay Control Kit	06P0110
Alinity s HIV Ag/Ab Combo Release Control Kit	06P0112
Alinity s HIV Ag/Ab Combo Reagent Kit	06P0160
Alinity s HIV Ag/Ab Combo Calibrator Kit	06P0103
Alinity s HIV Ag/Ab Combo Assay Control Kit	06P0120
Alinity s HIV Ag/Ab Combo Release Control Kit	06P0124





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)

www.zlg.

### V1 010051 0103 Rev. 10

Annex II List A Products	
Product Name	REF N°
Alinity s Anti-HCV Reagent Kit	06P0455
Alinity s Anti-HCV Calibrator Kit	06P0402
Alinity s Anti-HCV Assay Control Kit	06P0410
Alinity s Anti-HCV Release Control Kit	06P0412
Alinity s Anti-HCV Reagent Kit	06P0477
Alinity s Anti-HCV Calibrator Kit	06P0409
Alinity s Anti-HCV Assay Control Kit	06P0419
Alinity s Anti-HCV Release Control Kit	06P0418
Alinity s Anti-HCV Reagent Kit	06P0460
Alinity s Anti-HCV Calibrator Kit	06P0403
Alinity s Anti-HCV Assay Control Kit	06P0420
Alinity s Anti-HCV Release Control Kit	06P0424
Alinity s Anti-HCV II Reagent Kit	04W5655
Alinity s Anti-HCV II Calibrator Kit	04W5602
Alinity s Anti-HCV II Assay Control Kit	04W5610
Alinity s Anti-HCV II Release Control Kit	04W5612
Alinity s HTLV I/II Reagent Kit	06P0755
Alinity s HTLV I/II Calibrator Kit	06P0702
Alinity s HTLV I/II Assay Control Kit	06P0710
Alinity s HTLV I/II Release Control Kit	06P0712
Alinity s HIV Ag/Ab Combo Reagent Kit	06P0177
Alinity s HIV Ag/Ab Combo Calibrator Kit	06P0109
Alinity s HIV Ag/Ab Combo Assay Control Kit	06P0119
Alinity s HIV Ag/Ab Combo Release Control Kit	06P0118

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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)

**REF N°** 

### V1 010051 0103 Rev. 10

### Annex II List A Products for ARCHITECT Platform

Product Name

Anti-HCV Reagent Kit	6C37-74
Anti-HCV Reagent Kit	6C37-77
Anti-HCV Reagent Kit	6C37-78
Anti-HCV Calibrator	6C37-09
Anti-HCV Controls	6C37-19
Anti-HBc IgM Reagent Kit	6C33-74
Anti-HBc IgM Reagent Kit	6C33-75
Anti-HBc IgM Calibrators	6C33-09
Anti-HBc IgM Controls	6C33-19
Anti-HBe Reagent Kit	6C34-74
Anti-HBe Reagent Kit	6C34-77
Anti-HBe Calibrator	6C34-09
Anti-HBe Controls	6C34-19
HBeAg Reagent Kit	6C32-74
HBeAg Reagent Kit	6C32-77
HBeAg Calibrators	6C32-09
HBeAg Quantitative Calibrators	7P24-09
HBeAg Controls	6C32-19
HBeAg Quantitative Controls	7P24-19
HIV Ag/Ab Combo Reagent Kit	4J27-74
HIV Ag/Ab Combo Reagent Kit	4J27-77
HIV Ag/Ab Combo Reagent Kit	4J27-78
HIV Ag/Ab Combo Calibrator	4J27-09
HIV Ag/Ab Combo Controls	4J27-19
Anti-HBc II Reagent Kit	8L44-74
Anti-HBc II Reagent Kit	8L44-77
Anti-HBc II Reagent Kit	8L44-78
Anti-HBc II Calibrator	8L44-09
Anti-HBc II Controls	8L44-19







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)

### V1 010051 0103 Rev. 10

#### Annex II List A Products for Alinity i Platform Product Name REF N° Anti-HBe Reagent Kit 07P6374 Anti-HBe Reagent Kit 07P6377 Anti-HBe Calibrator 07P6309 Anti-HBe Controls 07P6319 HIV Ag/Ab Combo Reagent Kit 08P0774 HIV Ag/Ab Combo Reagent Kit 08P0777 HIV Ag/Ab Combo Calibrator 08P0709 HIV Ag/Ab Combo Controls 08P0719 Anti-HCV Reagent Kit 08P0674 Anti-HCV Reagent Kit 08P0677 Anti-HCV Calibrator 08P0609 Anti-HCV Controls 08P0619 Anti-HBc II Reagent Kit 07P8774 Anti-HBc II Reagent Kit 07P8777 Anti-HBc II Calibrator 07P8709 Anti-HBc II Controls 07P8719 Anti-HBc IgM Reagent Kit 07P8674 Anti-HBc IgM Calibrators 07P8609 Anti-HBc IgM Controls 07P8619 HBeAg Reagent Kit 07P6474 HBeAg Reagent Kit 07P6477 **HBeAg Calibrators** 07P6409 **HBeAg Controls** 07P6419 HBeAg Quantitative Calibrators 09P1009 HBeAg Quantitative Controls 09P1019

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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)

### V1 010051 0103 Rev. 10

Annex II List B Products	
Product Name	REF N°
ARCHITECT Toxo IgG Reagent Kit	6C19-25
ARCHITECT Toxo IgG Reagent Kit	6C19-35
ARCHITECT Toxo IgG Calibrators	6C19-01
ARCHITECT Toxo IgG Controls	6C19-10
ARCHITECT Toxo IgG Avidity Reagent Kit	6L37-25
ARCHITECT Toxo IgG Avidity Calibrator & Controls	6L37-11
ARCHITECT Toxo IgM Reagent Kit	6C20-25
ARCHITECT Toxo IgM Reagent Kit	6C20-35
ARCHITECT Toxo IgM Calibrator	6C20-01
ARCHITECT Toxo IgM Controls	6C20-10
Alinity i Toxo IgG Reagent Kit	07P4522
Alinity i Toxo IgG Reagent Kit	07P4532
Alinity i Toxo IgG Calibrators	07P4501
Alinity i Toxo IgG Controls	07P4510
Alinity i Toxo IgM Reagent Kit	07P4722
Alinity i Toxo IgM Reagent Kit	07P4732
Alinity i Toxo IgM Calibrator	07P4701
Alinity i Toxo IgM Controls	07P4710
Alinity i Toxo IgG Avidity Reagent Kit	07P4622
Alinity i Toxo IgG Avidity Controls	07P4610

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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)

07P4577

07P4709

07P4719

07P4774

07P4777

07P4674

### V1 010051 0103 Rev. 10

Toxo IgG Reagent Kit

Toxo IgM Calibrators

Toxo IgM Reagent Kit

Toxo IgM Reagent Kit

Toxo IgG Avidity Reagent Kit

Toxo IgM Controls

#### Annex II List B Products for ARCHITECT Platform Product Name REF N° Toxo IgG Calibrators 6C19-09 Toxo IgG Controls 6C19-19 Toxo IgG Reagent Kit 6C19-74 Toxo IgG Reagent Kit 6C19-77 **Toxo IgM Calibrators** 6C20-09 6C20-19 Toxo IgM Controls Toxo IgM Reagent Kit 6C20-74 Toxo IgM Reagent Kit 6C20-77 Toxo IgG Avidity Reagent Kit 6L37-74 Annex II List B Products for Alinity Platform Product Name REF N° Toxo IgG Calibrators 07P4509 Toxo IgG Controls 07P4519 Toxo IgG Reagent Kit 07P4574

Page 11 of 11 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123



TECHNOPATH CLINICAL DIAGNOSTICS

### **DECLARATION OF CONFORMITY**



### Manufacturer

Techno-path Manufacturing Ltd. Fort Henry Business Park, Ballina, Co. Tipperary, Ireland

### Product(s):

Product Name Multichem IA Plus Category Assayed/tri-level Catalogue Number 05P76-10

GMDN:	47869
Classification:	Annex II List B
Conformity Route:	Annex IV
Quality Management System:	EN ISO 13485:2016
QMS/CE Certification No.:	V11038520001
Issued By:	TÜV SÜD, Ridlerstraße 65, 80339 Munich,
	Germany
Expiry Date:	26 May 2024
Notified Body Number:	0123

Standards Applied: See attached list of standards for which documented evidence of compliance can be provided.

Techno-path Manufacturing Ltd. hereby declares that the product(s) specified above comply with the requirements listed in European Union In-vitro Diagnostic Medical Device Directive 98/79/EC.

I am fully responsible for all the information provided in this declaration. This declaration of conformity is valid from 3/(Day) 0/(Month) 20 (Year)

Signed for and on behalf of Techno-path Manufacturing Ltd.,

H1.

Bernd Hass, VP of Quality and Regulatory Affairs Techno-path Manufacturing Ltd. Ballina, Co.Tipperary <u>31-01-20</u>. Place and Date of Issue



TECHNOPATH CLINICAL DIAGNOSTICS

### STANDARDS USED IN FULL OR PART FOR CE MARKING AS PER IVDD 98/79/EC

Standard	Title	
EN ISO15223-1:2016	Symbols to be used with medical device labels, labelling	
	and information to be supplied.	
EN ISO13485:2016	Medical devices – Quality management systems –	
	Requirements for regulatory purposes	
EN 13612:2002 + AC:2002	Performance evaluation of in vitro diagnostic medical	
	devices	
EN 13641:2002	Elimination or reduction of risk of infection related to in	
*	vitro diagnostic reagents	
EN 13975:2003	Sampling procedures used for acceptance testing of in in	
	vitro diagnostic medical devices – statistical aspects	
EN ISO 14971:2012	Medical devices – Application of risk management to	
	medical devices	
EN ISO 18113-1:2011	In vitro diagnostic medical devices – Information supplied	
	by the manufacturer (labelling) – Part 1: Terms, definitions	
	and general requirements	
EN ISO 18113-2:2011	In vitro diagnostic medical devices – Information supplied	
	by the manufacturer (labelling) – Part 2: In vitro diagnostic	
	reagents for professional use	
EN 23640:2015	In vitro diagnostic medical devices - Evaluation of stability	
	of in vitro diagnostic reagents	
SOR/98-282, May 7, 1998	Canada Medical Device Regulations	

Abbott

Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address:		Declaration of Conformity   ARCHITECT Solutions   Abbott Ireland Diagnostics Division   Finisklin Business Park   Sligo   Ireland	1
List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
1L56-40	59058	ARCHITECT Probe Conditioning Solution	Self-declared
6C54-58	58236	ARCHITECT Concentrated Wash Buffer	Self-declared
6C54-82	58236	ARCHITECT Concentrated Wash Buffer	Self-declared
6C54-88	58236	ARCHITECT ARM Concentrated Wash Buffer	Self-declared
6C55-60	58793	ARCHITECT Trigger Solution	Self-declared
6C55-82	58793	ARCHITECT Trigger Solution	Self-declared
6E23-65	61163	ARCHITECT Pre-Trigger Solution	Self-declared
6E23-82	61163	ARCHITECT Pre-Trigger Solution	Self-declared
7D82-50	58208	ARCHITECT Multi-Assay Manual Diluent	Self-declared
	thorized European Representative Jame and Address)	N/A	
Storage site of technical documentation (Name and Address)		Abbott Ireland Diagnostics Division, Finisklin Business Park, Sligo, County Sligo, Ireland. Department: Regulatory Affairs.	
Harmonized Standards Listed in the Technical Documentation			

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

<pre>Signature:</pre>	- fens	Signature:	honore Weiter
Full Name:	Niall Plunkett	Full Name:	Lorraine Whitney
Position:	Quality Manager	Position:	Senior Manager Regulatory Affairs
Date of Approval:	16 Oct 14	Date of Approval:	14007 2014
Date Issued:	16 OCT 14	Place Issued:	AIDD Sligo
Supersedes:	07 July 2014	Effective (Date or Lot Number):	IDOCT IL



Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-7K70-AIDD Sligo Abbott Ireland Diagnostics Division Finisklin Business Park, Sligo, Ireland

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification	
7K70-20	54665	ARCHITECT Total PSA Reagent Kit	Annex II List B	
7K70-25	54665	ARCHITECT Total PSA Reagent Kit	Annex II List B	
7K70-30	54665	ARCHITECT Total PSA Reagent Kit	Annex II List B	
7K70-35	54665	ARCHITECT Total PSA Reagent Kit	Annex II List B	
7K70-01	38208	ARCHITECT Total PSA Calibrators	Annex II List B	
7K70-10	38207	ARCHITECT Total PSA Controls	Annex II List B	
Authorized Euro Representative (	opean name and address	) N/A		
	ame and address)			
Notified Body nu	umber	0123		
<b>Approval Certif</b>	icate No.	V1 0019220008	V1 0019220008	
Storage site of te		Abbott Ireland Diagnostics Division, Finisklin Business Park, Sligo, Ireland		
documentation (	name and address	Department: Regulatory Affairs		
Harmonized Sta	ndards	Listed in the Technical Documentation		

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex IV of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:	WALSH MURISM	Signature:	NO lon
Full Name:	VJoe Murray	Full Name:	Noel Haren
Position:	Director Quality Assurance/Site Quality Head	Position:	Manager Regulatory Affairs
Date of Approval:	25/0019	Date of Approval:	25 Nov 2019
Date Issued:	25 Nou 2019	Place Issued:	AIDD Sligo
Supersedes:	16 October 2019	Effective (Date or Lot Number):	25 NOU 2019

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Abbott

## **Declaration of Conformity**

Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-6S60-AIDD Sligo Abbott Ireland Diagnostics Division Finisklin Business Park, Sligo, Ireland

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
6\$60-22	64760	SARS-CoV-2 IgG II Quant Reagent Kit	Self Declared
6860-32	64760	SARS-CoV-2 IgG II Quant Reagent Kit	Self Declared
6S60-02	64854	SARS-CoV-2 IgG II Quant Calibrator Kit	Self Declared
6S60-12	64788	SARS-CoV-2 IgG II Quant Control Kit	Self Declared

Authorized European Representative (name and address) Storage site of technical documentation (name and address)	N/A Abbott Ireland Diagnostics Division, Finisklin Business Park, Sligo, Ireland. Department: Regulatory Affairs.
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:

Full Name: Position: Joe Murray Director Quality Assurance/Site Quality Head

Oct 2021

Date of Approval:

05 OCT 2021

Supersedes:

Date Issued:

30 April 2021

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2021

Noel Haren

Manager Regulatory Affairs

Date of Approval:

Signature:

Full Name:

Position:

Place Issued:

AIDD Sligo

OS OL

Effective (Date or Lot Number):

OS OCT 2021



Certificate Identification:	7K62
Legal Manufacturer's Name:	Abbott Ireland Diagnostics Division
Legal Manufacturer's Address: Lisnamuck, Longford	
	Co. Longford
	Ireland

List Numbers **GMDN** Code Names and Description of Devices Classification and Size Code of Devices 7K62-20 54386 ARCHITECT TSH Reagent Kit Self-declared 7K62-25 7K62-30 7K62-35 7K62-01 38272 **ARCHITECT TSH Calibrators** Self-declared 7K62-10 38271 **ARCHITECT TSH Controls** Self-declared

Authorized European Representative	N/A
(Name and Address)	
Storage of site technical documentation (Name and Address)	ADDOUL ITETATIO DIAGNOSTICS DIVISION, LISHAMUCK, LONGIDIU, CO. LONGIDIU, ITETATIO.
	Department: Regulatory Affairs
Harmonized Standards	Listed in the Technical Documentation

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

Signature:	lobled Wight	Signature:	ponaja aliphan
Full Name:	Siobhan Wright	Full Name:	Lorraine Whitney
Position:	Director Quality Assurance/ Site Quality Head	Position:	Senior Manager Regulatory Affairs
Date of Approval:	24-APR-19	Date of Approval:	19 APR 2019
Date Issued:	24- APR-19	Place Issued:	Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, Ireland
Supersedes:	25-May-2017	Effective (Date or Lot Number):	24-APR-19



08H59-01

08H59-02

55866

55866

### **Declaration of Conformity**

Certificate Identif	fication:	SC-08H59	
Legal Manufactur	rer's Name:	Abbott Laboratories Diagnostics Division	
Legal Manufactur	er's Address:	Abbott Park, IL 60064 USA	
List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification

CELL-DYN 26 Plus Control, Full Pack

CELL-DYN 26 Plus Control, Half Pack

Authorized European	ABBOTT	
Representative	Max-Planck-Ring-2	
(Name and Address)	65205 Wiesbaden, Germany	
Storage site of technical	Abbott Laboratories	
documentation	4551 Great America Parkway	
(Name and Address)	Santa Clara, CA 95054	
Harmonized Standards	Listed in the Technical Documentation	

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

## This declaration is made in accordance with Annex III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature:	Barny Com	Signature:	Juan Jogua
Full Name:	Barry Simpson	Full Name:	Marcy Jaqua
Position:	Site Quality Manager	Position:	Director, Regulatory Affairs
Date of Approval:	18. June , 2015	Date of Approval:	30 June 2015
Date Issued:	JUN 30 2015	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V5 February 26, 2015	Effective (Date or Lot Number):	JUL 06 2015

Self-declared

Self-declared



Certificate Identification:	SC-01H73	
	Abbott Laboratories	
Legal Manufacturer's Name:	Diagnostics Division	

Legal Manufacturer's Address:

Abbott Park, IL 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
01H73-01	58237	CELL-DYN Sapphire and CELL-DYN Ruby Systems DILUENT/SHEATH	Self-declared

Authorized European	ABBOTT	
Representative	Max-Planck-Ring-2	
(Name and Address)	65205 Wiesbaden, Germany	
Storage site of technical	Abbott Laboratories	
documentation	4551 Great America Parkway	
(Name and Address)	Santa Clara, CA 95054	
Harmonized Standards	Listed in the Technical Documentation	

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

Signature:	Barry Spor	Signature:	marcy Jogur
Full Name:	Barry Simpson	Full Name:	Marcy Jaqua
Position:	Site Quality Manager	Position:	Director, Regulatory Affairs
Date of Approval:	29. Jun. 2015	Date of Approval:	30 June 2015
Date Issued:	JUN 30 2015	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V2 January 10, 2014	Effective (Date or Lot Number):	JUL 06 2015

### Abbott

## **Declaration of Conformity**

SC-99644	
Abbott Laboratories	
Diagnostics Division	
Abbott Park, IL 60064 USA	_
	Abbott Laboratories Diagnostics Division

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
99644-01	59058	CELL-DYN ENZYMATIC CLEANER CONCENTRATE	Self-declared
93641-01	59058	CELL-DYN ENZYMATIC CLEANER CONCENTRATE	Self-declared

Authorized European	ABBOTT	
Representative (name and	Max-Planck-Ring 2	
address)	65205 Wiesbaden, Germany	
Storage site of technical	Abbott Laboratories	
documentation (name and	4551 Great America Parkway	
address)	Santa Clara, CA 95054 USA	
Harmonized Standards	Listed in the Technical Documentation	

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

Signature:	Barry Spa	Signature:	Juan Jua
Full Name:	Barry Simpson	Full Name:	Marcy Jaqua
Position:	Quality Manager	Position:	Regulatory Affairs, Director
Date of Approval:	04. Sept. 2015	Date of Approval:	04 Sep 2015
Date Issued:	SEP 0 4 2015	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V4, January 10, 2014	Effective (Date or Lot Number):	SEP 11 2015



Certificate Identification:	SC-03H80	
Legal Manufacturer's Name:	Abbott Laboratories Diagnostics Division	

Legal Manufacturer's Address:

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
03H80-02	61165	CELL-DYN Ruby, CELL-DYN 3200 Systems CN-FREE HGB/NOC LYSE	Self-declared

Abbott Park, IL 60064 USA

Authorized European	ABBOTT	
Representative	Max-Planck-Ring-2	
(Name and Address)	65205 Wiesbaden, Germany	
Storage site of technical	Abbott Laboratories	
documentation	4551 Great America Parkway	
(Name and Address)	Santa Clara, CA 95054	
Harmonized Standards	Listed in the Technical Documentation	

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

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Signature:	Barny Stores	Signature:	marco Squa
Full Name:	Barry Simpson	Full Name:	Marcy Jaqua
Position:	Site Quality Manager	Position:	Director, Regulatory Affairs
Date of Approval:	29. Jun. 2015	Date of Approval:	30 June 2015
Date Issued:	JUN <b>30</b> 2015	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V2 January 10, 2014	Effective (Date or Lot Number):	JUL 0 6 2015



Certificate Identification:	SC-08H52	
	Abbott Laboratories	
Legal Manufacturer's Name:	Diagnostics Division	

Legal Manufacturer's Address:

Abbott Park, IL 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification	
08H52-01 61165		CELL-DYN Ruby, CELL-DYN 3200 Systems WBC LYSE	Self-declared	

Authorized European	ABBOTT	
Representative	Max-Planck-Ring-2	
(Name and Address)	65205 Wiesbaden, Germany	
Storage site of technical	Abbott Laboratories	
documentation	4551 Great America Parkway	
(Name and Address)	Santa Clara, CA 95054	
Harmonized Standards Listed in the Technical Documentation		

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices as they are transposed into the laws of the member states.

Signature:	Barry Spin	Signature:	marco Jaguar
Full Name:	Barry Simpson	Full Name:	Marcy Jaqua
Position:	Site Quality Manager	Position:	Director, Regulatory Affairs
Date of Approval:	29. Jun, 2015	Date of Approval:	30 June 2015
Date Issued:	JUN 30 2015	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V2, January 10, 2014	Effective (Date or Lot Number):	JUL 0 6 2015



# **Certificate of Approval**

This is to certify that the Management System of:

# **Abbott Laboratories Diagnostics Division**

100 Abbott Park Road, Abbott Park, IL, 60064, United States

has been approved by LRQA to the following standards:

ISO 13485:2016



Cliff Muckleroy - Area Operations Manager Americas Issued by: Lloyd's Register Quality Assurance, Inc. for and on behalf of: Lloyd's Register Quality Assurance Limited

This certificate is valid only in association with the certificate schedule bearing the same number on which the locations applicable to this approval are listed.

Current issue date: 13 October 2018 Expiry date: 12 October 2021 Certificate identity number: 10155326 Original approval(s): ISO 13485 – 7 December 2017

Approval number(s): ISO 13485 - 0015680

The scope of this approval is applicable to:

Design, Manufacture, Development, Installation, Service and Support of In Vitro Diagnostic Products including Test Kits, Reagents, Accessories and Instruments.







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# **Certificate Schedule**

Certificate identity number: 10155326

Location	Activities	
100 Abbott Park Road, Abbott Park, IL, 60064,	ISO 13485:2016	
United States	Design, Manufacture, Development, Installation, Service and Support of In Vitro Diagnostic Products including Test Kits, Reagents, Accessories and Instruments.	
Conway Park, 675 North Field Drive, Lake Forest,	ISO 13485:2016	
IL, 60045, United States	Oversight of the Quality Management System for the Abbott Diagnostics Division Sites.	
K Complex - Distribution Center	ISO 13485:2016	
Route 41 & Martin Luther King Drive, North Chicago, IL, 60064, United States	Distribution of In Vitro Diagnostic Products including Test Kits, Reagents, Accessories and Instruments.	







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Certificate Identification:	SC-09H39	
	Abbott Laboratories	
Legal Manufacturer's Name:	Diagnostics Division	
Legal Manufacturer's Address:	Abbott Park, IL 60064 USA	

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
09H39-01 35476		CELL-DYN Emerald Instrument	Self-declared

Authorized European	Abbott GmbH		
Representative	Max-Planck-Ring-2		
(Name and Address)	65205 Wiesbaden, Germany		
Storage site of technical	Abbott Laboratories		
documentation	4551 Great America Parkway		
(Name and Address)	Santa Clara, CA 95054		
	BIT Group France,		
	Parc Euromedécine II,		
	Rue de la Valsière,		
	CEDEX 5, CS 14287		
	34 099 – Montpellier FRANCE		
Harmonized Standards	Listed in the Technical Documentation		

We, the undersigned, hereby declare that the in vitro diagnostic medical devices described above and bearing the CE marking, conform with the applicable provisions of the EC Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on In Vitro Diagnostic Medical Devices and Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011, as they are transposed into the laws of the member states.

Signature:	Church Nowlan	_ Signature:	alles
Full Name:	Cheryl Nowlan	Full Name:	Thao Phan
Position:	Director Quality Assurance	Position:	Associate Director Regulatory Affairs
Date of Approval:	14 JAN 2021	Date of Approval:	14 JAN 2021
Date Issued:	JAN 1 4 2021	Place Issued:	Abbott Santa Clara
Supersedes:	IRIS V7 (July 1, 2016)	Effective (Date or Lot Number):	JAN 1 4 2021