

| 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.

1 ania (

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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| 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.

Danis Den

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



Lloyd's Register Group Limited, its affiliates and subsidiaries, including Lloyd's Register Quality Assurance Limited (LRQA), and their respective officers, employees or agents are, individually and collectively, referred to in this clause as 'Lloyd's Register'. Lloyd's Register assumes no responsibility and shall not be liable to any person for any loss, damage or expense caused by reliance on the information or advice in this document or howsoever provided, unless that person has signed a contract with the relevant Lloyd's Register entity for the provision of this information or advice and in that case any responsibility or liability is exclusively on the terms and conditions set out in that contract. Issued by: Lloyd's Register Quality Assurance Limited, 1 Trinity Park, Bickenhill Lane, Birmingham B37 7ES, United Kingdom



| 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



Lloyd's Register Group Limited, its affiliates and subsidiaries, including Lloyd's Register Quality Assurance Limited (LRQA), and their respective officers, employees or agents are, individually and collectively, referred to in this clause as 'Lloyd's Register'. Lloyd's Register assumes no responsibility and shall not be liable to any person for any loss, damage or expense caused by reliance on the information or advice in this document or howsoever provided, unless that person has signed a contract with the relevant Lloyd's Register entity for the provision of this information or advice and in that case any responsibility or liability is exclusively on the terms and conditions set out in that contract. Issued by: Lloyd's Register Quality Assurance Limited, 1 Trinity Park, Bickenhill Lane, Birmingham B37 7ES, United Kingdom



| Certificate Identification: | AIDD 3P36 |
|-------------------------------|-------------------------------------|
| Legal Manufacturer's Name: | Abbott Ireland Diagnostics Division |
| Legal Manufacturer's Address: | Finisklin Business Park |
| Bert 1. | Sligo |
| | Ireland |

| List Numbers and Size Code of Devices | GMDN Code | Names and Description of Devices | Classification |
|--|--|----------------------------------|------------------------------------|
| 3P36-20 3P36-25 3P36-30 3P36-35 | 17259 | ARCHITECT AFP Reagent | Self-declared |
| 3P36-01 | 38167 | ARCHITECT AFP Calibrators | Self-declared |
| 3P36-10 | 38166 | ARCHITECT AFP Controls | Self-declared |
| 1) | thorized European Representative Name and Address) | | |
| Storage site of technical documentation (Name and Address) | | Decision Address | ark, Sligo, County Sligo, 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.

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

| Signature: | - And | Signature: | homaine litistney |
|-------------------|-----------------|------------------------------------|-----------------------------------|
| Full Name: | Niall Plunkett | Full Name: | Lorraine Whitney |
| Position: | Quality Manager | Position: | Senior Manager Regulatory Affairs |
| Date of Approval: | 07 Jun 14 | Date of Approval: | 04 July 2014 |
| Date Issued: | 07 2014 | Place Issued: | AIDD Sligo |
| Supersedes: | 13 Jan 2013 | Effective (Date or Lot Number): | 07 Jul 14 |

1

Abbott

Declaration of Conformity

Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-7C18 (re-standardised Mag-Sep) -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 |
|---|-----------|----------------------------------|-----------------|
| 7C18-29 | 48316 | ARCHITECT Anti-HBs Reagent Kit | Annex II List A |
| 7C18-39 | 48316 | ARCHITECT Anti-HBs Reagent Kit | Annex II List A |
| 7C18-33 | 48316 | ARCHITECT Anti-HBs Reagent Kit | Annex II List A |
| 7C18-41 | 48316 | ARCHITECT Anti-HBs Reagent Kit | Annex II List A |
| 7C18-42 | 48316 | ARCHITECT Anti-HBs Reagent Kit | Annex II List A |
| 7C18-03 | 41997 | ARCHITECT Anti-HBs Calibrators | Annex II List A |
| 7C18-13 | 41998 | ARCHITECT Anti-HBs Controls | Annex II List A |

| Authorized European | N/A |
|-----------------------------------|--|
| Representative (name and address) | |
| Notified Body (name and address) | TÜV SÜD Product Service GmbH |
| u • | Ridlerstraße 65 |
| | 80339 Munich |
| | Germany |
| Notified Body number | 0123 |
| Approval Certificate No. | V7 001922 0012 |
| 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

Position:

Director Quality Assurance/Site Quality Head

18 OCT 2021

1200 2021

Date of Approval:

Date Issued:

Supersedes:

25 Nov 2019

Position:

Signature:

Full Name:

Noel Haren Manager Regulatory Affairs

Date of Approval:

18 OCT 2021

AIDD, Sligo

Effective (Date or Lot Number):

Place Issued:

18 OCT 2021



| Certificate Identification: | 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

Abbott

Declaration of Conformity

Certificate Identification:DoC-2K45-SD DELK TPMLegal Manufacturer's Name:Abbott GmbHLegal Manufacturer's Address:Max-Planck-Ring 2, 65205 Wiesbaden, Germany

| List Numbers and Size Code of Devices | GMDN Code | Names and Description of Devices | Classification |
|--|--------------|----------------------------------|----------------|
| 2K45-24 | 54588 | ARCHITECT CA 125 II Reagent Kit | Self-declared |
| 2K45-29 | 54588 | ARCHITECT CA 125 II Reagent Kit | Self-declared |
| 2K45-39 | 54588 | ARCHITECT CA 125 II Reagent Kit | Self-declared |
| 2K45-02 | 38231 | ARCHITECT CA 125 II Calibrators | Self-declared |
| 2K45-11 | 38230 | ARCHITECT CA 125 II Controls | Self-declared |

| Authorized European Representative (name and address) | N/A |
|--|--|
| Storage site of technical | Fujirebio Diagnostics, Inc., 201 Great Valley Parkway, Malvern, Pennsylvania |
| documentation (name and address) | 19355, 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.

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:

Dr. Jörg Amborn

Full Name:

Signature:

wanu

Susanne Ulrich

Position:

Director Quality Assurance

Position:

Senior Manager Regulatory Affairs

May-2020

65205 Wiesbaden, Germany

12-May-2017

Effective (Date or Lot Number):

14- May-2020

Date of Approval:

2020-05-14

Date of Approval:

.....

Date Issued:

Place Issued:

Supersedes:

Abbott

Declaration of Conformity

| Certificate Identification: | DoC-2K44-SD DELK TPM |
|-------------------------------|---|
| Legal Manufacturer's Name: | Abbott GmbH |
| Legal Manufacturer's Address: | Max-Planck-Ring 2, 65205 Wiesbaden, Germany |

| List Numbers and GMDN Size Code of Devices Code | | Names and Description of Devices | Classification | |
|--|-------|----------------------------------|----------------|--|
| 2K44-21 | 60975 | ARCHITECT CA 15-3 Reagent Kit | Self-declared | |
| 2K44-27 | 60975 | ARCHITECT CA 15-3 Reagent Kit | Self-declared | |
| 2K44-37 | 60975 | ARCHITECT CA 15-3 Reagent Kit | Self-declared | |
| 2K44-02 | 38223 | ARCHITECT CA 15-3 Calibrators | Self-declared | |
| 2K44-11 | 38222 | ARCHITECT CA 15-3 Controls | Self-declared | |

| Authorized European Representative (name and address) | N/A |
|--|--|
| Storage site of technical | Fujirebio Diagnostics, Inc., 201 Great Valley Parkway, Malvern, Pennsylvania |
| documentation (name and address) | 19355, 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.

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:

Claudia Becker

Director Quality Assurance

Full Name:

Signature:

lich wanne.

Susanne Ulrich

Assoc. Director Regulatory Affairs

Mav-

65205 Wiesbaden, Germany

2021

M94 2021

Date of Approval:

Date Issued:

Place Issued:

Supersedes:

12-May-2017 Effective (Date or Lot Number):

25- May-2021

Date of Approval:

Position:

- Abbott

Declaration of Conformity

Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-2K91-SD DLK TPM Abbott GmbH Max-Planck-Ring 2, 65205 Wiesbaden, Germany

| List Numbers and Size Code of Devices | GMDN Code | Names and Description of Devices | Classification |
|--|--------------|---|----------------|
| 2K91-24 | 60976 | ARCHITECT CA 19-9 _{XR} Reagent Kit | Self-declared |
| 2K91-32 | 60976 | ARCHITECT CA 19-9 _{XR} Reagent Kit | Self-declared |
| 2K91-39 | 60976 | ARCHITECT CA 19-9 XR Reagent Kit | Self-declared |
| 2K91-03 | 38225 | ARCHITECT CA 19-9 _{XR} Calibrators | Self-declared |
| 2K91-12 | 38224 | ARCHITECT CA 19-9 XR Controls | Self-declared |

| Authorized European Representative (name and address) | N/A |
|--|--|
| Storage site of technical | Fujirebio Diagnostics, Inc., 201 Great Valley Parkway, Malvern, Pennsylvania |
| documentation (name and address) | 19355, 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.

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

Signature:

C. Kecles

Full Name:

Claudia Becker

Position:

Date of Approval:

Director Quality Assurance

proanne lle

Susanne Ulrich

Assoc. Director Regulatory Affairs

Date Issued:

Date of Approval:

Signature:

Full Name:

Position:

Place Issued:

Supersedes:

19-June-2019

Effective (Date or Lot Number):

21- Dec- 2021

Associ Director Regula

211 Dec 2071

65205 Wiesbaden, Germany

1- Dec - 2021

- Abbott

Declaration of Conformity

Certificate Identification:DoC-7K68- AIDD SligoLegal Manufacturer's Name:Abbott Ireland Diagnostics DivisionLegal Manufacturer's Address:Finisklin Business Park, Sligo, Ireland

| List Numbers and Size Code of Devices | GMDN Code | Names and Description of Devices | Classification |
|---|-----------|----------------------------------|----------------|
| 7K68-22 | 54615 | ARCHITECT CEA Reagent Kit | Self-declared |
| 7K68-27 | 54615 | ARCHITECT CEA Reagent Kit | Self-declared |
| 7K68-32 | 54615 | ARCHITECT CEA Reagent Kit | Self-declared |
| 7K68-35 | 54615 | ARCHITECT CEA Reagent Kit | Self-declared |
| 7K68-02 | 38174 | ARCHITECT CEA Calibrators | Self-declared |
| 7K68-12 | 38173 | ARCHITECT CEA Controls | Self-declared |

| Authorized European | N/A |
|-----------------------------------|---|
| Representative (name and address) | |
| 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 III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

Signature: Full Name:

Position:

Date of Approval:

Joe Murray Quality Manager

OS Jan 17-

| Signature: | bomaine Culiiten |
|---------------------------------|-----------------------------------|
| Full Name: | Lorraine Whitney |
| Position: | Senior Manager Regulatory Affairs |
| Date of Approval: | 05 Jan 17 |
| Date Issued: | 05 Jan (7 |
| Place Issued: | AIDD Sligo |
| Supersedes: | 25 Sep 2014 |
| Effective (Date or Lot Number): | 05 Jan 17 |

ABBOTT

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

n

07K72 Abbott Ireland Diagnostics Division Lisnamuck, Longford, Co. Longford, Ireland.

| List Numbers and Size Code of Devices | GMDN Code | Names and Description of Devices | Classification |
|---|-----------|------------------------------------|----------------|
| 7K72-20 7K72-25 | 30321 | ARCHITECT Estradiol Reagent Kits | Self-declared |
| 7K72-01 | 38249 | ARCHITECT Estradiol Calibrators | Self-declared |
| 7K72-10 | 38248 | ARCHITECT Estradiol Controls | Self-declared |
| 7K72-50 | N/A | ARCHITECT Estradiol Manual Diluent | Self-declared |

| Authorized European Representative (name and address) | N/A |
|--|---|
| Storage of technical documentation (name and address) | Abbott Ireland Diagnostics Division, Lisnamuck, Longford, 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 and transposed Irish Regulation S.I. No 304 of 2001 and to the EC Directive 98/79/EC as it is 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 (printed): Position: | <u>Alschar</u> Wight Siobhán Wright Quality Manager | Signature: Full Name (printed): Position: | Lorraine Whitney Manager Regulatory Affairs |
|--|---|--|---|
| Date: | 05- NOV - 13 | Date: | 01 NOU 2013 |
| | | Date Issued: | 05 NOV 2013 |
| | | Place Issued: | Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, Ireland. |
| | | Supersedes: Effective (Lot number or date) | 30 MAR 2012 05 NOV 2013 |



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 NU1 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 ₃ Reagent Kit | Self-declared |
| 7K63-32 | | | |
| 7K63-37 | | | |
| 7K63-02 | 38261 | ARCHITECT Free T ₃ Calibrators | Self-declared |
| 7K63-12 | 54418 | ARCHITECT Free T ₃ 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



| Certificate Identification: | 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 (Name and Address) | | | N/A | |
| Storage of site technical documentation (Name and Address) | | | Abbott Ireland Diagnostics Division, Lisnamuck, Lo | 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.

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: | 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

2021 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:



Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: DoC-6C32/7P24-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 |
|---|-----------|---|-----------------|
| 6C32-20 | 48331 | ARCHITECT HBeAg Reagent Kit (4x100 Tests) | Annex II List A |
| 6C32-25 | 48331 | ARCHITECT HBeAg Reagent Kit (1x100 Tests) | Annex II List A |
| 6C32-27 | 48331 | ARCHITECT HBeAg Reagent Kit (1x100 Tests) | Annex II List A |
| 6C32-37 | 48331 | ARCHITECT HBeAg Reagent Kit (1x500 Tests) | Annex II List A |
| 6C32-01 | 42007 | ARCHITECT HBeAg Calibrators | Annex II List A |
| 6C32-10 | 42008 | ARCHITECT HBeAg Controls | Annex II List A |
| 7P24-01 | 42007 | ARCHITECT HBeAg Quantitative Calibrators | Annex II List A |
| 7P24-10 | 42008 | ARCHITECT HBeAg Quantitative 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 | TÜV SÜD: 0123 | |
| Approval Certificate No. | TÜV SÜD: V7 010051 0120 | |
| 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.

sang Signature: Signature: Full Name: Full Name: Susanne Ulrich Dr. Jörg Amborn Senior Manager Regulatory Affairs Position: **Director Quality Assurance** Position: 1020-07-12 Date of Approval: Date of Approval: 1070 12- Mar-2020 Date Issued: Place Issued: 65205 Wiesbaden, Germany 12- Mar- 2020 Supersedes: 19-Dec-2019 Effective (Date or Lot Number):







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,

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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

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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 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

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

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 P42 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 |

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: 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 Confirmatory | |
| 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: www.tuvsud.com/ps-cert?q=cert:V7_001922_0009_Rev.01

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 |
| | |

Page 3 of 11 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123







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 |
| | |

Page 4 of 11 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123







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 |
| | |

Page 5 of 11 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123







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)

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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)

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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 |
| | |

Page 7 of 11 TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123







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)

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| 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)

REF N°

6C19-09

6C19-19

6C19-74

6C19-77

6C20-09

6C20-19

6C20-74

6C20-77

6L37-74

REF N°

V1 010051 0103 Rev. 10

Annex II List B Products for ARCHITECT Platform Product Name Toxo IgG Calibrators Toxo IgG Controls Toxo IgG Reagent Kit Toxo IgG Reagent Kit Toxo IgM Calibrators Toxo IgM Controls Toxo IgM Reagent Kit Toxo IgM Reagent Kit Toxo IgM Reagent Kit

Annex II List B Products for Alinity Platform

Product Name

| Toxo IgG Calibrators | 07P4509 |
|------------------------------|---------|
| Toxo IgG Controls | 07P4519 |
| Toxo IgG Reagent Kit | 07P4574 |
| Toxo IgG Reagent Kit | 07P4577 |
| Toxo IgM Calibrators | 07P4709 |
| Toxo IgM Controls | 07P4719 |
| Toxo IgM Reagent Kit | 07P4774 |
| Toxo IgM Reagent Kit | 07P4777 |
| Toxo IgG Avidity Reagent Kit | 07P4674 |
| | |



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.

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

| <pre>Signature:</pre> | - fers | 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 | 16 DCT 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 European Representative (name and address) N/A | | |
| | Notified Body (name and address) TÜV SÜD Product Service GmbH Ridlerstraße 65 80339 Munich Germany Germany | | |
| Notified Body nu | Notified Body number 0123 | | |
| Approval Certif | al Certificate No. V1 0019220008 | | |
| Ų | rage site of technical Abbott Ireland Diagnostics Division, Finisklin Business Park, Sligo, Ireland | | n Business Park, Sligo, Ireland |
| documentation (| cumentation (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

Date of Approval:

05 OCT 2021

Oct 2021

Date Issued: Supersedes:

30 April 2021

05

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à

2021

Noel Haren

Manager Regulatory Affairs

Date of Approval:

Signature:

Full Name:

Position:

Place Issued: Effective (Date or

Lot Number):

AIDD Sligo

OS OL

OS OCT 2021

| Abbott | | | | | |
|---|-----------------------------------|---------------|--------------------------------------|--------|----------------|
| | | Declar | ation of Conformity | | |
| Certifica | ate Identification: | ARCH Sys | Acc LC | IRIS V | V4 |
| Legal Manu | ifacturer's Name: | Abbott La | | | |
| | | Diagnostic | | | |
| Legal Manuta | cturer's Address: | Abbott Par | k, IL 60064 USA | | |
| List Numbers | GMDN Code | Ν | ames and Description of Devices | | Classification |
| and Size Code of Devices | | | | | |
| 4D18-03 | 56701 | ARCHITECT | Septum | | Self-declared |
| 4D19-01 | 56701 | ARCHITECT | Replacement Caps | | Self-declared |
| 7C14-01 | 56676 | ARCHITECT | Sample Cups | | Self-declared |
| 7C15-02 | 56676 | ARCHITECT | Reaction Vessels | | Self-declared |
| 7C15-03 | 56676 | ARCHITECT | Reaction Vessels | | Self-declared |
| Autl | horized European | Abbott GmbH | Contract (1997-1998) (Network (1997) | | |
| | Representative | Max-Planck-F | | | |
| <u> </u> | ame and Address) | | den, Germany | | |
| Storage site of technical Abbott Labora | | | | | |
| (Ne | documentation ame and Address) | Diagnostics D | L 60064 USA | | |
| ······ | onized Standards | | 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: | 1AD/ | |
|-------------------|------------------|-------|
| Full Name: | Katerina Damia | moska |
| Position: | Site Quality Dir | ector |
| Date of Approval: | 5/29/2019 | |
| Date Issued: | 22 July 2019 | |
| Supersedes: | 02 June 2015 | |

Signature: Full Name: an redor Position:

Date of Approval:

Place Issued:

Abbott Laboratories, Diagnostics Division, Abbott Park, IL 60064 USA

Effective (Date or Lot Number):



Certificate Identification: Legal Manufacturer's Name: Legal Manufacturer's Address: 7K64 Abbott Ireland Diagnostics Division Lisnamuck, Longford, Co. Longford, Ireland.

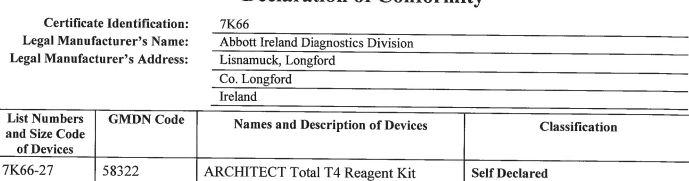
| List Numbers and Size Code of Devices | GMDN Code | Names and Description of Devices | Classification |
|---|-----------|---|----------------|
| 7K64-27 7K64-32 7K64-37 | 58330 | ARCHITECT Total T ₃ Reagent Kit | Self-declared |
| 7K64-02 | 58333 | ARCHITECT Total T ₃ Calibrators | Self-declared |
| 7K64-50 | 58208 | ARCHITECT Total T ₃ Manual Diluent | Self-declared |

| Authorized European Representative (name and address) | N/A |
|--|---|
| Storage of technical documentation (name and address) | Abbott Ireland Diagnostics Division, Lisnamuck, Longford, 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.

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

| Signature: | Asocher Wigh | Signature: | horraine Whitney |
|----------------------|--|-----------------------------------|---|
| Full Name: | Siobhan Wright | Full Name: | Lorraine Whitney |
| Position: | Director Quality Assurance/ Site Quality Head | Position: | Senior Manager Regulatory Affairs |
| Date of Approval: | 22 - MAY-10 | Date of Approval: | 22 MAY 2020 |
| Date Issued: | 22- MAY-20 | Place Issued: | Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, Ireland |
| Supersedes: | 24 April 2019 | Effective (Date or Lot Number) | 22 - MAY-20 |



ARCHITECT Total T4 Calibrators

ARCHITECT Total T4 Controls

N/A

Harmonized Standards | Listed in the Technical Documentation

Declaration of Conformity

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: | lidean Wigh |
|------------|--|
| Full Name: | Siobhan Wright |
| Position: | Director Quality Assurance/ Site Quality Head |
| Date of | 24-ADR-19 |

Approval:

Supersedes:

2

17-May-2016

| Signature: | foncia | Chihey | |
|------------|--------|--------|--|
| Juli Nomer | | | |

Self Declared

Self Declared

Full Name:

Lorraine Whitney Position: Senior Manager Regulatory Affairs

Date of Approval:

Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Ireland.

19 A1R 2019

Place Issued:

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

Date Issued:

24-APR-19

Effective (Date or Lot Number):

24- APR-19

Abbott

7K66-32 7K66-37 7K66-02

7K66-12

58324

58325

documentation (Name and

Authorized European

Representative (Name and Address) Storage of site technical

Address)



| 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 | ADDOUT IT CIAILU DIAGIIOSUCS DIVISIOII, LISIIAITIUCK, LOIIGIOIU, CO. LOIIGIOIU, ITCIAILU. |
| 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 III of the IVD Directive and is issued under the sole responsibility of the manufacturer.

| 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 |
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