



America

CERTIFICATE

No. QS6 001922 0023 Rev. 03

Certificate Holder:

Abbott Ireland Diagnostics Division
Finisklin Business Park
Sligo
IRELAND

Certification Mark:**Scope of Certificate:**

Design, Develop and Manufacture of In-Vitro Diagnostic Test Kits, Reagents and Common Liquid Accessories for Donor Screening and / or the Detection and / or Monitoring of Hepatitis, Cancers, Cardiac Markers, Congenital Transmitted Diseases, Determination of Congenital Disorders of the Foetus, Endocrine Disorders and Haematological Disorders and Therapeutic Drug Monitoring and Infectious Viral Diseases and Traumatic Brain Injury

Standard(s):**ISO 13485:2016****Regulatory Authority(ies):**

Australia TGA, Brazil ANVISA, Health Canada, USA FDA, MHLW / PMDA. See attached for listing of specific regulatory requirements.

The Certification Body of TÜV SÜD America Inc. certifies that the quality management system of the manufacturer listed above has been audited against the stated criteria and found to conform to those criteria for the scope of certification listed. Validity of this certificate can be obtained by visiting the website www.tuvsud.com/ps-cert
TÜV SÜD America Inc. is an MDSAP Recognized Auditing Organization.

REPs Facility ID:**F004677****Effective Date:****2023-03-02****Expiry Date:****2026-03-01**

Page 1 of 2

Date of Issue: 2023-02-16

(Renee Walker)
Director, US Certification Body, MHS

CERTIFICATE

No. QS6 001922 0023 Rev. 03

Regulatory Requirements: Audit/Certification Criteria

Australia

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

Brazil

- RDC ANVISA n. 665/2022 - Good Manufacturing Practices
- RDC ANVISA n. 551/2021
- RDC ANVISA n. 67/2009 - Vigilance

Canada

- Medical Device Regulations – Part 1- SOR 98/282

Japan

- MHLW Ministerial Ordinance No. 169 (2004), as amended by MHLW Ministerial Ordinance No.60 (2021)
- Japan PMD Act (as applicable)

United States

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

Facility(ies):

Abbott Ireland Diagnostics Division
Finisklin Business Park, Sligo, IRELAND

Facility Scopes:

Design, Develop and Manufacture of In-Vitro Diagnostic Test Kits, Reagents and Common Liquid Accessories for Donor Screening and / or the Detection and / or Monitoring of Hepatitis, Cancers, Cardiac Markers, Congenital Transmitted Diseases, Determination of Congenital Disorders of the Foetus, Endocrine Disorders and Haematological Disorders and Therapeutic Drug Monitoring and Infectious Viral Diseases and Traumatic Brain Injury
REPs Facility ID: F004677

Page 2 of 2

Date of Issue: 2023-02-16



(Renee Walker)
Director, US Certification Body, MHS



Certificate

No. Q5 001922 0022 Rev. 04

Holder of Certificate: **Abbott Ireland Diagnostics Division**
Finisklin Business Park
Sligo
IRELAND

Facility(ies): **Abbott Ireland Diagnostics Division**
Finisklin Business Park, Sligo, IRELAND

See Scope of Certificate

Certification Mark:



Scope of Certificate: **Design and Development, Production and Distribution of In Vitro Diagnostic Reagents for Immunochemistry, Immunology, Hematology and Infectious Diseases**

Applied Standard(s): EN ISO 13485:2016
Medical devices - Quality management systems -
Requirements for regulatory purposes
(ISO 13485:2016)
DIN EN ISO 13485:2016

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). 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:Q5 001922 0022 Rev. 04

Report No.: 713293822-01

Valid from: 2023-06-12

Valid until: 2026-03-24

Date, 2023-06-12

Christoph Dicks
Head of Certification/Notified Body



Management Service

CERTIFICATE

The Certification Body
of TÜV SÜD Management Service GmbH

certifies that



Abbott Ireland Diagnostics Division
Finisklin Business Park
Sligo
Ireland

has established and applies
a Quality Management System for

**Design and Development, Production and Distribution of
In Vitro Diagnostic Reagents for Immunochemistry,
Immunology and Infectious Diseases.**

An audit was performed, Order No. **707114974**.

Proof has been furnished that the requirements
according to

DIN EN ISO 9001:2015

are fulfilled.

The certificate is valid from **2023-04-01** until **2026-03-31**.

Certificate Registration No.: **12 100 59742 TMS**.

Head of Certification Body
Munich, 2023-01-03



ZERTIFIKAT ◆ CERTIFICATE ◆ CERTIFICADO ◆ CERTIFICAT ◆
CERTIFIKAT ◆ CERTIFICATE ◆ CERTIFICADO ◆ CERTIFICAT ◆
認證證書 ◆



Certificate

No. Q5 010051 0139 Rev. 01

Holder of Certificate: **Abbott GmbH**
Max-Planck-Ring 2
65205 Wiesbaden
GERMANY

Certification Mark:



Scope of Certificate: **Design and Development, Manufacture and Warehousing of In-vitro Diagnostic Reagents for Clinical Chemistry, Immunochemistry, Hematology and Infectious Immunology. The provision of Warehousing and Distribution services of In-vitro Diagnostic medical devices and medical devices.**

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the Testing, Certification, Validation and Verification Regulations TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:Q5 010051 0139 Rev. 01

Report No.: 713332354_IVDR

Valid from: 2024-10-01
Valid until: 2027-09-30

Date, 2024-07-11

Christoph Dicks
Head of Certification/Notified Body

Certificate

No. Q5 010051 0139 Rev. 01

Applied Standard(s): ISO 13485:2016
(EN ISO 13485:2016/AC:2018, EN ISO 13485:2016/A11:2021)
Medical devices - Quality management systems -
Requirements for regulatory purposes

Facility(ies): **Abbott GmbH**
Max-Planck-Ring 2, 65205 Wiesbaden, GERMANY

Design and Development, Manufacture and Warehousing of
In-vitro Diagnostic Reagents for Clinical Chemistry,
Immunochemistry, Hematology and Infectious Immunology.

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

The provision of Warehousing and Distribution services of
In-vitro Diagnostic medical devices and medical devices.

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Certificate

No. Q5 054869 0011 Rev. 02

Holder of Certificate: **Abbott Ireland Diagnostics Division**
Lisnamuck
Longford
Co. Longford
IRELAND

Certification Mark:



Scope of Certificate: **Design and Development, Manufacture and Distribution of In-Vitro Diagnostic Reagents for Clinical Chemistry and Immunochemistry.**

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). 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:Q5 054869 0011 Rev. 02](http://www.tuvsud.com/ps-cert?q=cert:Q5_054869_0011_Rev_02)

Report No.: 713280794

Valid from: 2023-09-01

Valid until: 2026-08-31

Date, 2023-07-14

Christoph Dicks
Head of Certification/Notified Body

Certificate

No. Q5 054869 0011 Rev. 02

Applied Standard(s): EN ISO 13485:2016
Medical devices - Quality management systems -
Requirements for regulatory purposes
(ISO 13485:2016)
DIN EN ISO 13485:2016

Facility(ies): **Abbott Ireland Diagnostics Division**
Lisnamuck, Longford, Co. Longford, IRELAND

See Scope of Certificate

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EU Quality Management System Certificate (IVDR)

Pursuant to Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices,
Annex IX Chapters I and III (Class C and B Devices excluding self-/near-patient-testing and
Companion Diagnostics)

No. V12 054869 0013 Rev. 01

Manufacturer: **Abbott Ireland Diagnostics Division**
Lisnamuck
Longford
Co. Longford
IRELAND

SRN Manufacturer: IE-MF-000010070

The Certification Body of TÜV SÜD Product Service GmbH certifies that the manufacturer has established, documented and implemented a quality management system as described in Article 10 (8) of the Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices. Details on devices covered by the quality management system are described on the following page(s).

The Report referenced below summarizes the result of the assessment and includes reference to relevant CS, harmonized standards, audit and test reports. The conformity assessment has been carried out according to Annex IX Chapter I and III of this regulation with a positive result.

The quality management system assessment was accompanied by the assessment of technical documentation for devices selected on a representative basis.

The certified quality management system is subject to periodical surveillance by TÜV SÜD Product Service GmbH. The surveillance assessment includes an assessment of the technical documentation for the device or devices concerned on the basis of further representative samples.

For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:V12 054869 0013 Rev. 01

Report No.: 713235160-02
Preceding Certificate No.: V12 054869 0013 Rev. 00
Valid from: 2021-12-03
Valid until: 2026-11-25
Date of Initial Issuance: 2021-11-26

Christoph Dicks
Head of Certification/Notified Body

Issue date: 2021-12-03



EU Quality Management System Certificate (IVDR)

Pursuant to Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices,
Annex IX Chapters I and III (Class C and B Devices excluding self-/near-patient-testing and
Companion Diagnostics)

No. V12 054869 0013 Rev. 01

Classification:	B
Device Group:	W0102 - IMMUNOCHEMISTRY (IMMUNOLOGY)
Intended Purpose:	IVR 0504 - Devices intended to be used to determine the infectious load, to determine infective disease status or immune status and devices used for infectious disease staging
Classification:	B
Device Group:	W0102 - IMMUNOCHEMISTRY (IMMUNOLOGY)
Intended Purpose:	IVR 0602 - Devices intended to be used for screening, determination or monitoring of physiological markers for a specific disease
Classification:	B
Device Group:	W0102 - IMMUNOCHEMISTRY (IMMUNOLOGY)
Intended Purpose:	IVR 0607 - Devices intended to be used for detection of pregnancy or fertility testing
Classification:	B
Device Group:	W0101 - CLINICAL CHEMISTRY
Intended Purpose:	IVR 0608 - Devices intended to be used for screening, determination or monitoring of physiological markers
Classification:	B
Device Group:	W0102 - IMMUNOCHEMISTRY (IMMUNOLOGY)
Intended Purpose:	IVR 0608 - Devices intended to be used for screening, determination or monitoring of physiological markers
Classification:	C
Device Group:	W0101 - CLINICAL CHEMISTRY
IVP Code:	IVP 3002 - In vitro diagnostic devices which require knowledge regarding biochemistry
Intended Purpose:	IVR 0301 - Devices intended to be used in screening, diagnosis, staging or monitoring of cancer
Classification:	C
Device Group:	W0101 - CLINICAL CHEMISTRY
IVP Code:	IVP 3002 - In vitro diagnostic devices which require knowledge regarding biochemistry
Intended Purpose:	IVR 0602 - Devices intended to be used for screening, determination or monitoring of physiological markers for a specific disease



EU Quality Management System Certificate (IVDR)

Pursuant to Regulation (EU) 2017/746 on in Vitro Diagnostic Medical Devices,
Annex IX Chapters I and III (Class C and B Devices excluding self-/near-patient-testing and
Companion Diagnostics)

No. V12 054869 0013 Rev. 01

Classification: C
Device Group: W0102 - IMMUNOCHEMISTRY (IMMUNOLOGY)
IVP Code: IVP 3007 - In vitro diagnostic devices which require knowledge
regarding immunoassays
Intended Purpose: IVR 0602 - Devices intended to be used for screening,
determination or monitoring of physiological markers for a specific
disease

Classification: C
Device Group: W0102 - IMMUNOCHEMISTRY (IMMUNOLOGY)
IVP Code: IVP 3002 - In vitro diagnostic devices which require knowledge
regarding biochemistry
Intended Purpose: IVR 0504 - Devices intended to be used to determine the
infectious load, to determine infective disease status or immune
status and devices used for infectious disease staging

**The validity of this certificate
depends on conditions and/or
is limited to the following:** -none-

Revision History:	Rev.	Dated	Report
	00	2021-11-26	713198595



America

CERTIFICATE

No. QS6 054869 0012 Rev. 04

Certificate Holder:

Abbott Ireland Diagnostics Division

Lisnamuck
Longford
Co. Longford
IRELAND

Certification Mark:



Scope of Certificate:

Design, Development and Manufacture of In-Vitro Diagnostic Test Kits and Reagents used in the Diagnosis of Prenatal Screening, Disease Status, Cardiac Markers, Protein Metabolism, Endocrine Disorders, Renal Dysfunction, Fertility Testing, Pregnancy Testing and for Therapeutic Drug Monitoring

Standard(s):

ISO 13485:2016

Regulatory Authority(ies):

Australia TGA, Brazil ANVISA, Health Canada, Japan MHLW / PMDA, USA FDA. See attached for listing of specific regulatory requirements.

The Certification Body of TÜV SÜD America Inc. certifies that the quality management system of the manufacturer listed above has been audited against the stated criteria and found to conform to those criteria for the scope of certification listed. For details and certificate validity see:

[www.tuvsud.com/ps-cert?q=cert:QS6 054869 0012 Rev. 04](http://www.tuvsud.com/ps-cert?q=cert:QS6_054869_0012_Rev._04)

TÜV SÜD America Inc. is an MDSAP Recognized Auditing Organization.

REPs Facility ID:

F005102

Report No.:

713319707

Effective Date:

2024-05-28

Expiry Date:

2026-05-30

Page 1 of 2

Date of Issue: 2024-06-05

(Renee Walker)
Director, US Certification Body, MHS

CERTIFICATE

No. QS6 054869 0012 Rev. 04

Regulatory Requirements: Audit/Certification Criteria

Australia

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

Brazil

- RDC ANVISA n. 665/2022 - Good Manufacturing Practices
- RDC ANVISA n. 551/2021
- RDC ANVISA n. 67/2009 - Vigilance

Canada

- Medical Device Regulations – Part 1- SOR 98/282

Japan

- MHLW Ministerial Ordinance No. 169 (2004), as amended by MHLW Ministerial Ordinance No.60 (2021)
- Japan PMD Act (as applicable)

United States

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

Facility(ies):

Abbott Ireland Diagnostics Division

Lisnamuck, Longford, Co. Longford, IRELAND

Facility Scopes:

Design, Development and Manufacture of In-Vitro Diagnostic Test Kits and Reagents used in the Diagnosis of Prenatal Screening, Disease Status, Cardiac Markers, Protein Metabolism, Endocrine Disorders, Renal Dysfunction, Fertility Testing, Pregnancy Testing and for Therapeutic Drug Monitoring
REPs Facility ID: F005102

Page 2 of 2

Date of Issue: 2024-06-05



(Renee Walker)
Director, US Certification Body, MHS



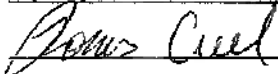
EU Declaration of Conformity

Basic UDI-DI: 038074DAL0002FQ
Basic UDI-DI Name: Alinity c-series Acid Probe Wash
Risk Class: Class A

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
01R6070	Alinity c-series Acid Probe Wash	58236	W0201010185
Manufacturer (Name and Address)		Abbott Laboratories 1915 Hurd Drive Irving, TX 75038 USA	
Manufacturer SRN		US-MF-00001777	
Authorized Representative (Name and Address)		Abbott GmbH Max-Planck-Ring 2 65205 Wiesbaden, Germany	
Authorized Representative SRN		DE-AR-000009457	
Produced by (Site of Manufacture) (Name and Address)		Fisher Diagnostics 8365 Valley Pike Middletown VA 22645 USA	
Conformity Assessment Procedure		Annex II and III	

We, the undersigned, hereby declare that the in vitro diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on In Vitro Diagnostic Medical Devices; and additionally conforms applicable provisions of Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, and to applicable provisions of Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery, and amending Directive 95/16/EC as transposed into the laws of the member states.

This declaration is made in accordance with Annex IV of the IVD Regulation, Annex VI of the ROHS Directive, and Annex II of the Machinery Directive and is issued under the sole responsibility of the manufacturer.

Full Name: Thomas Creel
Sr. Director, Instrument and Automation
Function: Quality
Signature: 
Date of Approval: 23-May-2022
Signed for, and on behalf of: Abbott Laboratories, 1915 Hurd Drive, Irving, TX 75038

Full Name: Michele Smith-Waheed
Function: Associate Director, Regulatory Affairs
Signature: 
Date of Approval: 23-May-2022

Date Issued: 23-May-2022
Supersedes: N/A

Place Issued: Irving, Texas
Effective (Date or Lot Number): 23-May-2022

EN	EU Declaration of Conformity	Basic UDI-DI	Basic UDI-DI Name
BG	ЕС ДЕКЛАРАЦИЯ ЗА СЪОТВЕТСТВИЕ	Базов UDI-DI	Наименование на базов UDI-DI
CS	EU PROHLAŠENÍ O SHODĚ	Základní UDI-DI	Název základního UDI-DI
DA	EU-OVERENSSTEMMELSESESKLÆRING	Grundlæggende UDI-DI	Grundlæggende UDI-DI-navn
DE	EU-KONFORMITÄTSESKLÄRUNG	Basis-UDI-DI	Basis-UDI-DI Name
EL	ΔΗΛΩΣΗ ΣΥΜΜΟΡΦΩΣΗΣ ΕΕ	Βασικό UDI-DI	Ονομασία βασικού UDI-DI
ES	DECLARACIÓN UE DE CONFORMIDAD	UDI-DI Básico	Nombre UDI-DI Básico
ET	ELi vastavusdeklaratsioon	Põhi-UDI-DI	Põhi-UDI-DI nimi
FR	Déclaration de conformité UE	IUD-ID de base	Nom IUD-ID de base
HR	EU IZJAVA O SUKLADNOSTI	Osnovni UDI-DI	Naziv osnovnog UDI-DI
HU	EU-MEGFELELŐSÉGI NYILATKOZAT	Alapvető UDI-DI	Alapvető UDI-DI neve
IT	Dichiarazione di conformità UE	UDI-DI di base	Nome UDI-DI di base
LV	ES atbilstības deklarācija	Pamata UDI-DI	Pamata UDI-DI nosaukums
LT	ES ATITIKTIES DEKLARACIJA	Bazinis UDI-DI	Bazinio UDI-DI pavadinimas
NO	EU-samsvarserklæring	Grunnleggende UDI-DI	Grunnleggende UDI-DI-navn
PL	DEKLARACJA ZGODNOŚCI UE	Kod Basic UDI-DI	Nazwa kodu Basic UDI-DI
PT	DECLARAÇÃO UE DE CONFORMIDADE	UDI-DI básico	Nome UDI-DI Básico
RO	Declarația de Conformitate UE	UDI-DI de bază	Nume UDI-DI de bază
SK	EÚ VYHLÁSENIE O ZHODE	Základný UDI-DI	Názov základného UDI-DI
SV	EU-FÖRSÄKRAN OM ÖVERENSSTÄMMELSE	Grundläggande UDI-DI	Namn på grundläggande UDI-DI
TR	AB Uygunluk Beyanı	Temel UDI-DI	Temel UDI-DI İsmi

EN	Risk Class	List Number and Size Code	Product and Trade Name
BG	Клас според риска	Каталожен номер и код на размера	Име на продукта и търговско наименование
CS	Riziková třída	Katalogové číslo a koncové dvojčíslí určující velikost soupravy	Název produktu a obchodní název
DA	Risikoklasse	Bestillingsnummer og størrelseskode	Produkt- og varemærkenavn
DE	Risikoklasse	Bestellnummer und Größencode	Produkt- und Handelsname
EL	Κατηγορία κινδύνου	Κωδικός Προϊόντος και Κωδικός Συσκευασίας	Προϊόν και Εμπορική Ονομασία
ES	Clase de riesgo	Número de referencia y código de tamaño	Producto y marca comercial
ET	Riskiklass	Katalooginumbr ja suurusekood	Toote nimetus ja kaubanimi
FR	Classe de risque	Référence	Nom de produit et de marque
HR	Klasa rizika	Kataloški broj i oznaka pakiranja	Naziv proizvoda i zaštiteni naziv
HU	Kockázati osztály	Listaszám és készletkiszerezés-kód	Termék- és kereskedelmi név
IT	Classe di rischio	Numero di listino e codice formato	Prodotto e nome commerciale
LV	Riska klase	Kataloga numurs un izmēra kods	Produkta nosaukums un tirdzniecības nosaukums
LT	Rizikos klasė	Katalogo numeris ir dydžio kodas	Gaminio ir prekybos pavadinimai
NO	Risikoklasse	Bestillingsnummer og størrelseskode	Produkt- og handelsnavn
PL	Klasa ryzyka	Numer katalogowy	Nazwa produktu i nazwa handlowa
PT	Classe de risco	Número de lista e código de apresentação	Produto e nome comercial
RO	Clasă de risc	Număr de listă și cod dimensiune	Denumirea produsului și denumirea comercială
SK	Riziková trieda	Katalógové číslo	Názov produktu a obchodný názov
SV	Riskklass	Listnummer och storlekskod	Produkt och firmanamn
TR	Risk Sınıfı	Liste Numarası ve Boyut Kodu	Ürün ve Ticari İsmi

EN	GMDN Code	EMDN Code	Manufacturer (Name and Address)	Manufacturer SRN
BG	Код GMDN	Код EMDN	Производител (име и адрес)	EPH на производителя
CS	Kód GMDN	Kód EMDN	Výrobce (název a adresa)	Jediné registrační číslo výrobce
DA	GMDN-kode	EMDN-kode	Fabrikant (navn og adresse)	Fabrikants SRN
DE	GMDN-Code	EMDN-Code	Hersteller (Name und Adresse)	Hersteller-SRN
EL	Κωδικός GMDN (Ονοματολογία ιατροτεχνολογικών προϊόντων)	Κωδικός EMDN (Ονοματολογία ιατροτεχνολογικών προϊόντων)	Κατασκευαστής (Όνομα και Διεύθυνση)	SRN (Μοναδικός Αριθμός Μητρώου) Κατασκευαστή
ES	Código GMDN	Código EMDN	Fabricante (nombre y dirección)	SRN (número de registro único) del fabricante
ET	GMDN-kood	EMDN-kood	Tootja (nimi ja aadress)	Tootja unikaalne registreerimisnumber
FR	Code GMDN	Code EMDN	Fabricant (nom et adresse)	Numéro d'enregistrement unique du fabricant
HR	GMDN kod	EMDN kod	Proizvođač (naziv i adresa)	SRN (jedinstveni registracijski broj) proizvođača
HU	GMDN-kód	EMDN-kód	Gyártó (név és cím)	Gyártó egyedi regisztrációs száma (SRN)
IT	Codice GMDN	Codice EMDN	Fabbricante (nome e indirizzo)	SRN (numero di registrazione unico) del fabbricante
LV	GMDN kods	EMDN kods	Ražotājs (nosaukums un adrese)	Ražotāja vienotais reģistrācijas numurs (VRN)
LT	Visuotinės medicinos priemonių nomenklatūros kodas	Europos medicinos priemonių nomenklatūros kodas	Gamintojas (pavadinimas ir adresas)	Gamintojo unikalusi registracijos numeris
NO	GMDN-kode	EMDN-kode	Produsent (navn og adresse)	Produsentens SRN
PL	Kod GMDN	Kod Europejskiej Nomenklatury Wyrobów Medycznych	Producent (nazwa i adres)	Niepowtarzalny numer rejestracyjny producenta
PT	Código GMDN	Código EMDN	Fabricante (Nome e Morada)	Número único de registo do fabricante
RO	Cod GMDN	Cod EMDN	Producător (nume și adresă)	SRN producător
SK	Kód GMDN	Kód EMDN	Výrobca (Názov a adresa)	Jediné registračné číslo (SRN) výrobcu
SV	GMDN-kod	EMDN-kod	Tillverkare (namn och adress)	Tillverkarens SRN
TR	GMDN Kodu	EMDN Kodu	Üretici (İsim ve Adres)	Üretici SRN'si

EN	Authorized Representative (Name and Address)	Authorized Representative SRN	Produced by (Site of Manufacture) (Name and Address)
BG	Упълномощен представител (име и адрес)	ЕРН на упълномощения представител	Произведено от (място на производство) (име и адрес)
CS	Zplnomocněný zástupce (název a adresa)	Jediné registrační číslo zplnomocněného zástupce	Vyrobeno (místo výroby) (název a adresa)
DA	Autoriseret repræsentant (navn og adresse)	Autoriseret repræsentants SRN	Produceret af (fabrikationssted) (Navn og adresse)
DE	Bevollmächtigter (Name und Adresse)	SRN des Bevollmächtigten	Hergestellt von (Herstellungsstandort) (Name und Adresse)
EL	Εξουσιοδοτημένος Αντιπρόσωπος (Όνομα και Διεύθυνση)	SRN Εξουσιοδοτημένου Αντιπροσώπου	Κατασκευάζεται από (Εργοστάσιο παραγωγής) (Όνομασία και Διεύθυνση)
ES	Representante autorizado (nombre y dirección)	SRN (número de registro único) del representante autorizado	Producido por (Lugar de fabricación) (Nombre y dirección)
ET	Volitatud esindaja (nimi ja aadress)	Volitatud esindaja unikaalne registreerimisnumber	Tootnud (tootmiskoht) (nimi ja aadress)
FR	Mandataire (nom et adresse)	Numéro d'enregistrement unique du mandataire	Produit par (site de fabrication) (nom et adresse)
HR	Ovlašteni zastupnik (naziv i adresa)	SRN (jedinstveni registracijski broj) ovlaštenog zastupnika	Proizvodi (Mjesto proizvodnje) (Naziv i adresa)
HU	Meghatalmazott képviselő (név és cím)	Meghatalmazott képviselő egyedi regisztrációs száma (SRN)	Gyártó (gyártás helye) (név és cím)
IT	Mandatario (nome e indirizzo)	SRN (numero di registrazione unico) del mandatario	Prodotto da (sito di fabbricazione) (nome e indirizzo)
LV	Pilnvarotais pārstāvis (nosaukums un adrese)	Pilnvarotā pārstāvja vienotais reģistrācijas numurs (VRN)	Ražots (ražošanas vieta) (nosaukums un adrese)
LT	Igaliotasis atstovas (pavadinimas ir adresas)	Igaliojojo atstovo unikalusi registracijos numeris	Pagaminta (gamybos vieta) (pavadinimas ir adresas)
NO	Autorisert representant (navn og adresse)	Den autoriserte representantens SRN	Produsert av (produksjonssted) (navn og adresse)
PL	Upoważniony przedstawiciel (nazwa i adres)	Niepowtarzalny numer rejestracyjny upoważnionego przedstawiciela	Wyprodukowano przez (miejsce produkcji) (nazwa i adres)
PT	Mandatário (Nome e Morada)	Número único de registo do mandatário	Produzido por (Local de fabrico) (Nome e Morada)
RO	Reprezentant autorizat (nume și adresă)	SRN reprezentant autorizat	Produs de către (locuție producție) (nume și adresă)
SK	Autorizovaný zástupca (názov a adresa)	Jediné registračné číslo (SRN) autorizovaného zástupcu	Výrobené (miesto výroby) (názov a adresa)
SV	Auktoriserad representant (namn och adress)	Auktoriserad representants SRN	Tillverkas av (tillverkningsort) (namn och adress)
TR	Yetkili Temsilci (İsim ve Adres)	Yetkili Temsilci SRN'si	Üretici (Üretim Tesisi) (İsim ve Adres)

EN	Conformity Assessment Procedure	Annex II and III	Full Name
BG	Процедура за оценка на съответствието	Приложение II и III	Пълно наименование
CS	Postup posuzování shody	Příloha II a III	Celý název
DA	Overensstemmelsesvurderingsprocedure	Bilag II og III	Fulde navn
DE	Konformitätsbewertungsverfahren	Anhang II und III	Vollständiger Name
EL	Διαδικασία αξιολόγησης συμμόρφωσης	Παράρτημα II και III	Πλήρης ονομασία
ES	Procedimiento de evaluación de la conformidad	Anexos II y III	Nombre completo
ET	Vastavushindamismenetlus	II ja III lisa	Täisnimi
FR	Procédure d'évaluation de la conformité	Annexes II et III	Nom complet
HR	Postupak ocjenjivanja sukladnosti	Prilog II. i III.	Puni naziv
HU	Megfelelőségértékelési eljárás	II. és III. melléklet	Teljes név
IT	Procedura di valutazione della conformità	Allegati II e III	Nome completo
LV	Atbilstības novērtēšanas procedūra	II un III pielikums	Pilns nosaukums
LT	Atitikties vertinimo procedūra	II ir III priedai	Vardas ir pavardė
NO	Framgangsmåte for samsvarsvurdering	Vedlegg II og III	Fullt navn
PL	Procedura oceny zgodności	Załącznik II oraz III	Imię i nazwisko
PT	Procedimento de avaliação da conformidade	Anexo II e III	Nome completo
RO	Procedură de evaluare a conformităţii	Anexa II şi III	Numele complet
SK	Postup posudzovania zhody	Príloha II a III	Celý názov
SV	Förfarande för bedömning av överensstämmelse	Bilaga II och III	Fullständigt namn
TR	Uygunluk Değerlendirme Prosedürü	Ek II ve III	Adı Soyadı

EN	Function	Signed for, and on behalf of	Date Issued
BG	Длъжност	Подписано за и от името на	Дата на издаване
CS	Funkce	Podepsáno za a jménem	Datum vydání
DA	Funktion	Underskrevet for og på vegne af	Udstedelsesdato
DE	Funktion	Unterzeichnet für und im Auftrag von	Datum
EL	Λειτουργία	Υπογράφεται για και εκ μέρους του/της	Ημερομηνία έκδοσης
ES	Función	Firmada por, y en nombre de	Fecha
ET	Funktsioon	Alla kirjutanud (kelle poolt ja nimel)	Väljaandmise kuupäev
FR	Fonction	Signé par et au nom de	Date d'établissement
HR	Funkcija	Potpisano za i u ime	Datum izdavanja
HU	Beosztás	Aláíró a következő képviselőtől és nevétől	Kiadás dátuma
IT	Funzione	Firmato a nome e per conto di	Data di rilascio
LV	Amats	Parakstīts šādas personas vārdā	Izdošanas datums
LT	Pareigos	Subjekto, kurio vardu pasirašoma, pavadinimas	Išdavimo data
NO	Funksjon	Signert for, og på vegne av	Utstedelsesdato
PL	Funkcja	Podpisano w imieniu	Data wydania
PT	Função	Assinado e em nome de	Data de emissão
RO	Funcția	Semnat pentru și în numele	Data eliberării
SK	Funkcia	Podpísané za a v mene	Dátum vydania
SV	Funktion	Undertecknat för och på uppdrag av	Datum för utfärdande
TR	Görevi	Namına ve temsilen imza	Düzenlenme Tarihi

EN	Supersedes	Signature	Date of Approval
BG	Замества	Подпис	Дата на одобрение
CS	Nahrazuje	Podpis	Datum schválení
DA	Erstatter	Underskrift	Godkendelsesdato
DE	Ersetzt	Unterschrift	Datum der Genehmigung
EL	Αντικαθιστά	Υπογραφή	Ημερομηνία έγκρισης
ES	Sustituye	Firma	Fecha de aprobación
ET	Asendab	Allkiri	Heakskiitmise kuupäev
FR	Annule et remplace	Signature	Date de l'autorisation
HR	Zamjenjuje	Potpis	Datum odobrenja
HU	Hatálytalanítja a következő dokumentumot:	Aláírás	Jóváhagyás dátuma
IT	Sostituisce	Firma	Data di approvazione
LV	Aizstāj	Paraksts	Apstiprināšanas datums
LT	Pakeičia	Parašas	Patvirtinimo data
NO	Erstatter	Signatur	Godkjenningsdato
PL	Zastępuje	Podpis	Data zatwierdzenia
PT	Substitui	Assinatura	Data de aprovação
RO	Înlocuitor	Semnătură	Data aprobării
SK	Nahrádza	Podpis	Dátum schválenia
SV	Ersätter	Namnteckning	Datum för godkännande
TR	Yerini aldığı belge	İmza	Onay Tarihi

EN	Place Issued	Effective (Date or Lot Number)
BG	Място на издаване	В сила от/за (дата или номер на партида)
CS	Místo vydání	Účinné od (datum nebo číslo šarže)
DA	Udstedelsessted	Ikrafttrædelse (dato eller lotnummer)
DE	Ort	Gültig ab (Datum oder Chargenbezeichnung)
EL	Τόπος έκδοσης	Σε ισχύ από (Ημερομηνία ή αρ. παρτίδας)
ES	Expedida en	Efectiva (fecha o número de lote)
ET	Väljaandmise koht	Jõustumine (kuupäev või partiinumber)
FR	Lieu d'établissement	Entrée en vigueur (date ou numéro de lot)
HR	Mjesto izdavanja	Stupa na snagu (datum ili broj serije)
HU	Kiadás helye	Hatálybalépés (dátum vagy tételszám)
IT	Luogo di rilascio	Valido da (data o numero di lotto)
LV	Izdošanas vieta	Spēkā no (datums vai partijas numurs)
LT	Išdavimo vieta	Isigalioja (data arba partijos numeris)
NO	Ustedelsessted	Gjelder fra (dato eller lotnummer)
PL	Miejsce wydania	Obowiązuje od (data lub numer partii)
PT	Local de emissão	Efetividade (Data ou número de lote)
RO	Locul eliberării	Valabilitate (data sau numărul lotului)
SK	Miesto vydania	Účinnosť od (dátum alebo číslo šarže)
SV	Plats för utfärdande	Verkställtigt (datum eller lotnummer)
TR	Düzenlendiği Yer	Yürürlük (Tarih veya Lot Numarası)

EN	We, the undersigned, hereby declare that the in vitro diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on In Vitro Diagnostic Medical Devices; and additionally conforms applicable provisions of Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, and to applicable provisions of Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery, and amending Directive 95/16/EC as transposed into the laws of the member states. This declaration is made in accordance with Annex IV of the IVD Regulation, Annex VI of the ROHS Directive, and Annex II of the Machinery Directive and is issued under the sole responsibility of the manufacturer.
BG	Ние, долуподписаните, с настоящото декларираме, че гореописаното(ите) медицинско(и) изделие(я) за инвитро диагностика отговаря(т) на приложимите разпоредби на Регламент (ЕС) 2017/746 на Европейския парламент и на Съвета от 5 април 2017 г. относно медицинските изделия за инвитро диагностика; освен това отговаря(т) на приложимите разпоредби на Директива 2011/65/ЕС на Европейския парламент и на Съвета от 8 юни 2011 г. относно ограничението на употребата на определени опасни вещества в електрическото и електронното оборудване и на приложимите разпоредби на Директива 2006/42/ЕО на Европейския парламент и на Съвета от 17 май 2006 г. относно машините, и за изменение на Директива 95/16/ЕО, както е транспонирана в националното законодателство на държавите членки. Тази декларация се прави в съответствие с Приложение IV на Регламента за IVD, Приложение VI на Директивата за ограничаване на опасните вещества (ROHS) и Приложение II на Директивата относно машините и за нейното издаване отговорност носи единствено производителят.
CS	My, níže podepsaní, tímto prohlašujeme, že diagnostický(-ě) zdravotnický(-ě) prostředek (prostředky) in vitro uvedený(-é) výše je (jsou) ve shodě s příslušnými ustanoveními nařízení Evropského parlamentu a Rady (EU) 2017/746 ze dne 5. dubna 2017 o diagnostických zdravotnických prostředcích in vitro; a že je (jsou) dále ve shodě s příslušnými ustanoveními směrnice Evropského parlamentu a Rady 2011/65/EU ze dne 8. června 2011 o omezení používání některých nebezpečných látek v elektrických a elektronických zařízeních a s příslušnými ustanoveními směrnice Evropského parlamentu a Rady 2006/42/ES ze dne 17. května 2006 o strojních zařízeních a o změně směrnice 95/16/ES, jak byla provedena ve vnitrostátním právu členských států. Toto prohlášení je v souladu s Přílohou IV nařízení IVD, Přílohou VI směrnice ROHS a Přílohou II směrnice o strojních zařízeních a je vydáno na výhradní odpovědnost výrobce.
DA	Vi, undertegnede, erklærer herved, at det in vitro-diagnostiske medicinske udstyr, der er beskrevet ovenfor, er i overensstemmelse med de gældende bestemmelser i Europa-Parlamentets og Rådets forordning (EU) 2017/746 af 5. april 2017 om in vitro-diagnostisk medicinsk udstyr, ligesom det overholder gældende bestemmelser i Europa-Parlamentets og Rådets direktiv 2011/65/EU af 8. juni 2011 om begrænsning af brugen af visse farlige stoffer i elektrisk og elektronisk udstyr samt overholder gældende bestemmelser i Europa-Parlamentets og Rådets direktiv 2006/42/EF af 17. maj 2006 om maskiner og ændring af direktiv 95/16/EF, som det er transponeret i medlemsstaternes lovgivning. Denne erklæring afgives i overensstemmelse med IVD-forordningens bilag IV, ROHS-direktivets bilag VI samt maskindirektivets bilag II og udstedes under fabrikantens eneansvar.
DE	Wir, die Unterzeichner, erklären hiermit, dass das oben beschriebene In-vitro-Diagnostikum/die oben beschriebenen In-vitro-Diagnostika die entsprechenden Bestimmungen der Verordnung (EU) 2017/746 des Europäischen Parlaments und des Rates vom 5. April 2017 über In-vitro-Diagnostika erfüllen und zusätzlich die entsprechenden Bestimmungen der Richtlinie 2011/65/EU des Europäischen Parlaments und des Rates vom 8. Juni 2011 zur Beschränkung der Verwendung bestimmter gefährlicher Stoffe in Elektro- und Elektronikgeräten sowie der Richtlinie 2006/42/EG des Europäischen Parlaments und des Rates vom 17. Mai 2006 über Maschinen und zur Änderung der Richtlinie 95/16/EG gemäß Umsetzung in den Gesetzen der Mitgliedsstaaten. Diese Erklärung erfolgt gemäß Anhang IV der IVD-Verordnung, Anhang VI der RoHS-Richtlinie und Anhang II der Maschinen-Richtlinie und wird unter alleiniger Verantwortung des Herstellers ausgestellt.
EL	Εμείς, οι υπογράφωντες, δηλώνουμε με το παρόν ότι τα προαναφερόμενα διαγνωστικά ιατροτεχνολογικά προϊόντα συμμορφώνονται με τις ισχύουσες διατάξεις του Κανονισμού (ΕΕ) 2017/746 του Ευρωπαϊκού Κοινοβουλίου και του Συμβουλίου της 5 ^{ης} Απριλίου 2017 σχετικά με τα in vitro διαγνωστικά ιατροτεχνολογικά προϊόντα και επίσης συμμορφώνονται με τις ισχύουσες διατάξεις της Οδηγίας 2011/65/ΕΕ του Ευρωπαϊκού Κοινοβουλίου και του Συμβουλίου της 8 ^{ης} Ιουνίου 2011 σχετικά με τους περιορισμούς στη χρήση συγκεκριμένων επικίνδυνων ουσιών στον ηλεκτρικό και ηλεκτρονικό εξοπλισμό, καθώς και με τις ισχύουσες διατάξεις της Οδηγίας 2006/42/ΕΚ του Ευρωπαϊκού Κοινοβουλίου και του Συμβουλίου της 17 ^{ης} Μαΐου 2006 σχετικά με τον μηχανικό εξοπλισμό και την τροποποιητική Οδηγία 95/16/ΕΚ όπως αυτή μεταφέρθηκε στη νομοθεσία των κρατών μελών. Η δήλωση αυτή γίνεται σύμφωνα με το Παράρτημα IV του Κανονισμού IVD, το Παράρτημα VI της Οδηγίας ROHS και το Παράρτημα II της Οδηγίας για τον μηχανικό εξοπλισμό και εκδίδεται με αποκλειστική ευθύνη του κατασκευαστή.
ES	Nosotros, los abajo firmantes, por la presente declaramos que el(los) producto(s) sanitario(s) para diagnóstico <i>in vitro</i> descrito(s) anteriormente cumple(n) las disposiciones aplicables del reglamento (UE) 2017/746 del Parlamento Europeo y del Consejo del 5 de abril de 2017 sobre productos sanitarios para diagnóstico <i>in vitro</i> ; y además cumple(n) las disposiciones aplicables de la Directiva 2011/65/UE del Parlamento Europeo y del Consejo del 8 de junio de 2011 sobre restricciones a la utilización de determinadas sustancias peligrosas en aparatos eléctricos y electrónicos, y las disposiciones aplicables de la Directiva 2006/42/EC del Parlamento Europeo y del Consejo del 17 de mayo de 2006 sobre maquinaria, y la Directiva de enmienda 95/16/EC tal y como se ha incorporado en las leyes de los Estados Miembros. Esta declaración se realiza en conformidad con el Anexo IV del Reglamento IVD, Anexo VI de la Directiva ROHS y Anexo II de la Directiva de máquinas y es emitida bajo la exclusiva responsabilidad del fabricante.
ET	Meie, allakirjutanud, kinnitame, et eespool kirjeldatud <i>in vitro</i> diagnostikameditsiiniseadmed vastavad Euroopa Parlamendi ja nõukogu 5. aprilli 2017. aasta määruse (EL) 2017/746 (<i>in vitro</i> diagnostikameditsiiniseadmete kohta) kohaldatavatele sätetele ning lisaks vastab see kohaldatavatele sätetele Euroopa Parlamendi ja nõukogu 8. juuni 2011. aasta direktiivis 2011/65/EL (teatavate ohtlike ainete kasutamise piiramise kohta elektri- ja elektroonikaseadmetes) ja Euroopa Parlamendi ja nõukogu direktiivis 2006/42/EÜ, 17. mai 2006, mis käsitleb masinaid ja millega muudetakse direktiivi 95/16/EÜ, nagu see on üle võetud liikmesriikide seadustes. See deklaratsioon on koostatud vastavalt IVD määruse IV lisale, ROHS direktiivi VI lisale ja masinadirektiivi II lisale ning see on välja antud tootja vastutusel.
FR	Nous soussigné(e)s, déclarons par la présente que le(s) dispositif(s) médical(aux) de diagnostic <i>in vitro</i> indiqué(s) ci-dessus est/sont conforme(s) aux dispositions applicables du Règlement (UE) 2017/746 du Parlement européen et du Conseil du 5 avril 2017 relatif aux dispositifs médicaux de diagnostic <i>in vitro</i> , aux dispositions applicables de la Directive 2011/65/UE du Parlement européen et du Conseil du 8 juin 2011 relative à la limitation de l'utilisation de certaines substances dangereuses dans les équipements électriques et électroniques, aux dispositions applicables de la Directive 2006/42/CE du Parlement européen et du Conseil du 17 mai 2006 relative aux machines et modifiant la Directive 95/16/CE, telles que transposées dans le droit national des États membres. Cette déclaration est établie conformément à l'Annexe IV du Règlement DIV, à l'Annexe VI de la Directive ROHS ainsi qu'à l'Annexe II de la Directive Machines sous la seule responsabilité du fabricant.
HR	Mi, niže potpisani, ovim putem izjavljujemo da su gore navedeni <i>in vitro</i> dijagnostički medicinski proizvod(i) sukladni primjenjivim odredbama Uredbe (EU) 2017/746 Europskog parlamenta i Vijeća od 5. travnja 2017. o <i>in vitro</i> dijagnostičkim medicinskim proizvodima; i dodatno primjenjivim odredbama Direktive 2011/65/EU Europskog parlamenta i Vijeća od 8. lipnja 2011. o ograničenju uporabe određenih opasnih tvari u električnoj i elektroničkoj opremi, te primjenjivim odredbama Direktive 2006/42/EZ Europskog parlamenta i Vijeća od 17. svibnja 2006. o strojevima zamjenjujući Direktivu 95/16/EZ kako je pretočeno u zakone država članica. Ova je izjava sastavljena u skladu s Prilogom IV. Uredbe IVD, Prilogom VI. Direktive ROHS i Prilogom II. Direktive o strojevima i izdaje se pod isključivom odgovornošću proizvođača.
HU	Alulírottak ezennel kijelentjük, hogy a fent leírt in vitro orvostechnikai eszköz(ök) megfelel(nek) az Európai Parlament és a Tanács in vitro diagnosztikai orvostechnikai eszközökről szóló (EU) 2017/746 (2017. április 5.) rendelete vonatkozó rendelkezéseit; továbbá az Európai Parlament és a Tanács egyes veszélyes anyagok elektromos és elektronikus berendezésekben való alkalmazásának korlátozásáról szóló 2011/65/EU (2011. június 8.) irányelve (RoHS irányelv) vonatkozó rendelkezéseit; valamint az Európai Parlament és a Tanács a gépekről és a 95/16/EK irányelv módosításáról szóló 2006/42/EK (2006. május 17.) irányelve vonatkozó rendelkezéseit a tagállamok jogrendjébe áttettető rendelkezéseknek. A jelen nyilatkozat megfelel az IVD rendelet IV. mellékletében, a RoHS irányelv VI. mellékletében és a gépekről szóló irányelv II. mellékletében foglalt előírásoknak, és a gyártó kizárólagos felelőssége alapján került kiadásra.

EN	We, the undersigned, hereby declare that the <i>in vitro</i> diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on <i>In Vitro</i> Diagnostic Medical Devices; and additionally conforms applicable provisions of Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, and to applicable provisions of Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery, and amending Directive 95/16/EC as transposed into the laws of the member states. This declaration is made in accordance with Annex IV of the IVD Regulation, Annex VI of the ROHS Directive, and Annex II of the Machinery Directive and is issued under the sole responsibility of the manufacturer.
IT	Noi, i sottoscritti, con la presente dichiariamo che il(i) dispositivo(i) medico-diagnostico(i) <i>in vitro</i> sopra descritto(i) è(sono) conforme(i) alle disposizioni applicabili del regolamento (UE) 2017/746 del Parlamento europeo e del Consiglio del 5 aprile 2017 relativo ai dispositivi medico-diagnostici <i>in vitro</i> ; è(sono) inoltre conforme(i) alle disposizioni applicabili della direttiva 2011/65/UE del Parlamento europeo e del Consiglio dell'8 giugno 2011 sulla restrizione dell'uso di determinate sostanze pericolose nelle apparecchiature elettriche ed elettroniche, e alle disposizioni applicabili della direttiva 2006/42/CE del Parlamento europeo e del Consiglio del 17 maggio 2006 relativa alle macchine e che modifica la direttiva 95/16/CE come recepite nelle legislazioni degli Stati membri. Questa dichiarazione è redatta in conformità all'allegato IV del regolamento IVD, all'allegato VI della direttiva ROHS e all'allegato II della direttiva macchine ed è rilasciata sotto la responsabilità esclusiva del fabbricante.
LV	Mēs, apakšā parakstījušies, ar šo paziņojam, ka iepriekš aprakstītā(-s) <i>in vitro</i> diagnostikas medicīniskā(-s) ierīce(-es) atbilst Eiropas Parlamenta un Padomes Regulas (ES) 2017/746 (2017. gada 5. aprīlis) piemērojamajām prasībām par <i>in vitro</i> diagnostikas medicīniskām ierīcēm un papildus prasībām, kas noteiktas Eiropas Parlamenta un Padomes Direktīvā 2011/65/ES (2011. gada 8. jūnijs) par dažu bīstamu vielu izmantošanas ierobežošanu elektriskās un elektroniskās iekārtās un Eiropas Parlamenta un Padomes Direktīvā 2006/42/EK (2006. gada 17. maijs) par mašīnām, un ar kuru groza Direktīvu 95/16/EK, kā tā ieviesta dalībvalstu tiesību aktos. Šī deklarācija ir sagatavota saskaņā ar IVD regulas IV pielikumu, ROHS direktīvas VI pielikumu un Direktīvas par mašīnām II pielikumu un par izdošanu atbild vienīgi ražotājs.
LT	Mes, toliau pasirašiusieji (-iusiosios), pareiškiame, kad anksčiau minėta (-os) <i>in vitro</i> diagnostikos medicinos priemonė (-ės) atitinka 2017 m. balandžio 5 d. Europos Parlamento ir Tarybos reglamento (ES) 2017/746 dėl <i>in vitro</i> diagnostikos medicinos priemonių taikytinas nuostatas; taip pat ji (jos) atitinka 2011 m. birželio 8 d. Europos Parlamento ir Tarybos direktyvos 2011/65/ES dėl tam tikrų pavojingų medžiagų naudojimo elektros ir elektroninėje įrangoje apribojimo taikomas nuostatas ir 2006 m. gegužės 17 d. Europos Parlamento ir Tarybos direktyvos 2006/42/EB dėl mašinų, iš dalies keičiančios Direktyvą 95/16/EB, taikomas nuostatas, perkeltas į valstybių narių teisės aktus. Ši deklaracija yra parengta vadovaujantis IVD reglamento IV priedu, ROHS direktyvos VI priedu ir Mašinų direktyvos II priedu ir yra išduodama tik gamintojo atsakomybe.
NO	Vi, undertegnede, erklærer herved at utstyret til <i>in vitro</i> -diagnostikk som er anført ovenfor, er i samsvar med gjeldende bestemmelser i Europaparlaments- og rådsforordning (EU) 2017/746 av 5. april 2017 om medisinsk utstyr til <i>in vitro</i> -diagnostikk, og ytterligere overholder gjeldende bestemmelser i Europaparlaments- og rådsdirektiv 2011/65/EU av 8. juni 2011 om bruksbegrensninger av visse farlige stoffer i elektrisk og elektronisk utstyr, og til gjeldende bestemmelser i Europaparlaments- og rådsdirektiv 2006/42/EF av 17. mai 2006 om maskiner, og endring av direktiv 95/16/EF som innarbeidet i medlemsstatenes lovgivning. Denne erklæringen er utarbeidet i overensstemmelse med vedlegg IV i IVD-forordningen, vedlegg VI i ROHS-direktivet og vedlegg II i maskindirektivet og er utstedt under produsentens eneansvar.
PL	My, niżej podpisani, niniejszym oświadczamy, że wymieniony(-e) powyżej wyrób(wyroby) medyczny(-e) do diagnostyki <i>in vitro</i> spełnia(-ja) odpowiednie wymagania Rozporządzenia (UE) 2017/746 Parlamentu Europejskiego i Rady z dnia 5 kwietnia 2017 r. w sprawie wyrobów medycznych do diagnostyki <i>in vitro</i> , a ponadto wymagania Dyrektywy 2011/65/UE Parlamentu Europejskiego i Rady z dnia 8 czerwca 2011 r. w sprawie ograniczenia stosowania niektórych niebezpiecznych substancji w sprzęcie elektrycznym i elektronicznym, Dyrektywy 2006/42/WE Parlamentu Europejskiego i Rady z dnia 17 maja 2006 r. w sprawie maszyn, zmieniającej Dyrektywę 95/16/WE, w sposób, w jaki zostały one wdrożone do ustawodawstwa państw członkowskich. Niniejsza deklaracja została sporządzona zgodnie z Załącznikiem IV Rozporządzenia IVDR, Załącznikiem VI Dyrektywy ROHS oraz Załącznikiem II Dyrektywy Maszynowej i wydana na wyłączną odpowiedzialność producenta.
PT	Nós, abaixo assinados, declaramos que os dispositivos médicos para diagnóstico <i>in vitro</i> descritos acima estão em conformidade com as disposições aplicáveis do Regulamento (UE) 2017/746 do Parlamento Europeu e do Conselho, de 5 de abril de 2017, relativo aos dispositivos médicos para diagnóstico <i>in vitro</i> ; e adicionalmente, em conformidade com as disposições aplicáveis da Diretiva 2011/65/UE do Parlamento Europeu e do Conselho, de 8 de junho de 2011, relativa à restrição do uso de determinadas substâncias perigosas em equipamentos elétricos e eletrônicos, e com as disposições aplicáveis da Diretiva 2006/42/CE do Parlamento Europeu e do Conselho, de 17 de maio de 2006, sobre máquinas e que altera a Diretiva 95/16/CE, conforme transposta nas leis dos Estados membros. Esta declaração é feita de acordo com o Anexo IV do Regulamento IVD, o Anexo VI da Diretiva ROHS e o Anexo II da Diretiva relativa às Máquinas e é emitida sob a exclusiva responsabilidade do fabricante.
RO	Subsemnatii, declaram că dispozitivul (dispozitivele) medical(e) pentru diagnostic <i>in vitro</i> descrie mai sus sunt conforme cu dispozițiile aplicabile din Regulamentul (UE) 2017/746 al Parlamentului European și al Consiliului din 5 aprilie 2017 privind Dispozitivele medicale pentru diagnosticul <i>in vitro</i> ; și, în plus, respectă dispozițiile aplicabile din Directiva 2011/65/UE a Parlamentului European și a Consiliului din 8 iunie 2011 privind restricția utilizării anumitor substanțe periculoase în echipamentele electrice și electronice și cu dispozițiile aplicabile din Directiva 2006/42/CE a Parlamentului European și a Consiliului din 17 mai 2006 privind utilajele și modificarea Directivei 95/16/CE, transpusă în legile statelor membre. Prezenta declarație este emisă în conformitate cu anexa IV la Regulamentul IVD, anexa VI la Directiva ROHS și anexa II la Directiva utilajelor și este emisă sub responsabilitatea exclusivă a producătorului.
SK	My, dolupodpisani, týmto vyhlasujeme, že diagnostická(-é) zdravotnícka(-e) pomôcka(-y) <i>in vitro</i> uvedená(-é) vyššie je (sú) v zhode s príslušnými ustanoveniami Nariadenia Európskeho parlamentu a Rady (EÚ) 2017/746 z 5. apríla 2017 o diagnostických zdravotníckych pomôckach <i>in vitro</i> ; a že je (sú) ďalej v zhode s príslušnými ustanoveniami Smernice Európskeho parlamentu a Rady 2011/65/EÚ z 8. júna 2011 o obmedzení používania určitých nebezpečných látok v elektrických a elektronických zariadeniach a s príslušnými ustanoveniami Smernice Európskeho parlamentu a Rady 2006/42/ES zo 17. mája 2006 o strojových zariadeniach a o zmene a doplnení Smernice 95/16/ES tak, ako boli transponované do zákonov členských štátov. Toto vyhlásenie je v súlade s Prílohou IV k Nariadeniu IVD, Prílohou VI k Smernici ROHS a Prílohou II k Smernici o strojových zariadeniach a vydáva sa na výhradnú zodpovednosť výrobcu.
SV	Vi, undertecknade, försäkrar härmed att den eller de medicintekniska produkter för <i>in vitro</i> -diagnostik som beskrivs ovan överensstämmer med de tillämpliga bestämmelserna i Europaparlamentets och rådets förordning (EU) 2017/746 av den 5 april 2017 om medicintekniska produkter för <i>in vitro</i> -diagnostik samt även överensstämmer med de tillämpliga bestämmelserna i Europaparlamentets och rådets direktiv 2011/65/EU av den 8 juni 2011 om begränsning av användning av vissa farliga ämnen i elektrisk och elektronisk utrustning samt med de tillämpliga bestämmelserna i Europaparlamentets och rådets direktiv 2006/42/EG av den 17 maj 2006 om maskiner och om ändring av direktiv 95/16/EG (omarbetning) som införlivats i medlemsstaternas lagstiftning. Denna försäkran görs i enlighet med bilaga IV till IVD-förordningen, bilaga VI till ROHS-direktivet samt bilaga II till maskindirektivet och utfärdas under tillverkarens enskilda ansvar.
TR	Biz, aşağıda imzaları bulunan, yukarıda belirtilen <i>in vitro</i> diagnostik tıbbi cihazların, 2017/746 sayılı Avrupa Parlamentosu (AB) Yönetmeliği ile 5 Nisan 2017 tarihli <i>In Vitro</i> Diagnostik Tıbbi Cihazlar Konseyinin ilgili hükümlerine uygun olduğunu ve ayrıca elektrikli ve elektronik cihazlarda belirli tehlikeli maddelerin kullanımının sınırlandırılmasına ilişkin 8 Haziran 2011 tarihli Konseyin ve 2011/65/EU sayılı Avrupa Parlamentosu Direktifinin ilgili hükümlerine, makinelere ilişkin 17 Mayıs 2006 tarihli Konseyin ve 2006/42/EC sayılı Avrupa Parlamentosu Direktifinin ilgili hükümlerine ve üye devlet yasalarına aktarılan 95/16/EC sayılı ek Direktife uygun olduğunu beyan ederiz. Bu beyan IVD Yönetmeliği Ek IV, ROHS Direktifi Ek VI ve Makineler Direktifi Ek II uyarınca yapılmıştır ve üreticinin münhasır sorumluluğu altında yayınlanmıştır.

End of form



EU Declaration of Conformity

Basic UDI-DI: 038074ACU0430JT
Basic UDI-DI Name: Albumin BCG2
Risk Class: Class B

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
04U3020	Albumin BCG2	59071	W01010201
04U3030	Albumin BCG2	59071	W01010201
Manufacturer (Name and Address)	Abbott Ireland Diagnostics Division Lisnamuck, Longford Co. Longford Ireland		
Manufacturer SRN	IE-MF-000010070		
Authorized Representative (Name and Address)	N/A		
Authorized Representative SRN	N/A		
Produced by (Site of Manufacture) (Name and Address)	Abbott Ireland Diagnostics Division Lisnamuck, Longford Co. Longford Ireland		
Notified Body (Name and Identification Number)	TÜV SÜD Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany Notified Body Number 0123		
Conformity Assessment Procedure	Quality Management System Annex IX Chapters I and III, Including an assessment of the technical documentation for devices concerned on the basis of representative samples	EU Certificate No. No. V12 054869 0013	
Common Specifications (CS)	N/A		

We, the undersigned, hereby declare that the in vitro diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on In Vitro Diagnostic Medical Devices. This declaration is made in accordance with Annex IV of the IVD Regulation and is issued under the sole responsibility of the manufacturer.

Full Name: <u>David Spellman</u>	Full Name: <u>Sandra Gallagher</u>
Function: <u>Director Quality Assurance/ Site Quality</u>	Function: <u>Manager Regulatory Affairs</u>
Signature: <u></u>	Signature: <u></u>
Date of Approval: <u>10 SEP 2024</u>	Date of Approval: <u>09-SEP-2024</u>
Signed for, and on behalf of: <u>Abbott Ireland Diagnostics Division Lisnamuck, Longford Co. Longford Ireland</u>	
Date Issued: <u>10 SEP 2024</u>	Place Issued: <u>Lisnamuck, Longford Co. Longford Ireland</u>
Supersedes: <u>13-Mar 2023</u>	Effective (Date or Lot Number): <u>10 SEP 2024</u>



EU Declaration of Conformity

Basic UDI-DI: 038074ACT0483K5
Basic UDI-DI Name: Alkaline Phosphatase2
Risk Class: Class B

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
04T8320	Alkaline Phosphatase2	52929	W01010105
04T8330	Alkaline Phosphatase2	52929	W01010105

Manufacturer (Name and Address)	Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford Ireland		
Manufacturer SRN	IE-MF-000010070		
Authorized Representative (Name and Address)	N/A		
Authorized Representative SRN	N/A		
Produced by (Site of Manufacture) (Name and Address)	Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford Ireland		
Notified Body (Name and Identification Number)	TÜV Süd Product Service GmbH Zertifizierstellen, Ridlerstraße 65 • 80339 Munich Germany Notified Body Number 0123		
Conformity Assessment Procedure	Quality Management System Annex IX Chapters I and III, Including an assessment of the technical documentation for devices concerned on the basis of representative samples	EU Certificate No. No. V12 054869 0013	
Common Specifications (CS)	N/A		

We, the undersigned, hereby declare that the in vitro diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on In Vitro Diagnostic Medical Devices. This declaration is made in accordance with Annex IV of the IVD Regulation and is issued under the sole responsibility of the manufacturer.

Full Name: <u>Siobhan Wright</u>	Full Name: <u>Sandra Gallagher</u>
Function: <u>Director Quality Assurance/Site Quality</u>	Function: <u>Manager Regulatory Affairs</u>
Signature: <u><i>Siobhan Wright</i></u>	Signature: <u><i>S. Gallagher</i></u>
Date of Approval: <u>16-DEC-2021</u>	Date of Approval: <u>16-DEC-2021</u>

Signed for, and on behalf of: Abbott Ireland Diagnostics Division Lisnamuck, Longford, Co. Longford Ireland

Date Issued: <u>16-DEC-2021</u>	Place Issued: <u>Lisnamuck, Longford, Co. Longford, Ireland</u>
Supersedes: <u>N/A</u>	Effective (Date or Lot Number): <u>16-DEC-2021</u>



EU Declaration of Conformity

Basic UDI-DI: 0380741DA1.00021 Q
Basic UDI-DI Name: Alinity c-series Alkaline Wash
Risk Class: Class A

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
08P7840	Alinity c-series Alkaline Wash	58236	W0201010185
Manufacturer (Name and Address)	Abbott Laboratories 1915 Hurd Drive Irving, TX 75038 USA		
Manufacturer SRN	US-MF-000017777		
Authorized Representative (Name and Address)	Abbott GmbH Max-Planck-Ring 2 65205 Wiesbaden, Germany		
Authorized Representative SRN	DE-AR-000009457		
Produced by (Site of Manufacture) (Name and Address)	Fisher Diagnostics 8365 Valley Pike Middletown, VA 22645 USA		
Conformity Assessment Procedure	Annex II and III		

We, the undersigned, hereby declare that the in vitro diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on In Vitro Diagnostic Medical Devices; and additionally conforms applicable provisions of Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, and to applicable provisions of Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery, and amending Directive 95/16/EC as transposed into the laws of the member states.

This declaration is made in accordance with Annex IV of the IVD Regulation, Annex VI of the ROHS Directive, and Annex II of the Machinery Directive and is issued under the sole responsibility of the manufacturer.

Full Name: Kevin Richardson

Function: Director, Instrument Quality

Signature: 

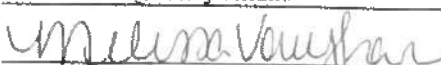
Date of Approval: 20-July-2023
Signed for, and on behalf of: Abbott Laboratories, 1915 Hurd Drive, Irving, TX 75038 USA

Date Issued: 20-JULY-2023

Supersedes: 23-May-2022

Full Name: Melissa Vaughan

Function: Director, Regulatory Affairs

Signature: 

Date of Approval: July 20, 2023

Place Issued: Irving, Texas

Effective (Date or Lot Number):

20-JULY-2023

EN	EU Declaration of Conformity	Basic UDI-DI	Basic UDI-DI Name
BG	ЕС ДЕКЛАРАЦИЯ ЗА СЪОТВЕТСТВИЕ	Базов UDI-DI	Наименование на базов UDI-DI
CS	EU PROHLÁŠENÍ O SHODĚ	Základní UDI-DI	Název základního UDI-DI
DA	EU-OVERENSSTEMMELSESESKLÆRING	Grundlæggende UDI-DI	Grundlæggende UDI-DI-navn
DE	EU-KONFORMITÄTSESKLÄRUNG	Basis-UDI-DI	Basis-UDI-DI Name
EL	ΔΗΛΩΣΗ ΣΥΜΜΟΡΦΩΣΗΣ ΕΕ	Βασικό UDI-DI	Όνομασία βασικού UDI-DI
ES	DECLARACIÓN UE DE CONFORMIDAD	UDI-DI Básico	Nombre UDI-DI Básico
ET	ELi vastavusdeklaratsioon	Põhi-UDI-DI	Põhi-UDI-DI nimi
FR	Déclaration de conformité UE	IUD-ID de base	Nom IUD-ID de base
HR	EU IZJAVA O SUKLADNOSTI	Osnovni UDI-DI	Naziv osnovnog UDI-DI
HU	EU-MEGFELELŐSÉGI NYILATKOZAT	Alapvető UDI-DI	Alapvető UDI-DI neve
IT	Dichiarazione di conformità UE	UDI-DI di base	Nome UDI-DI di base
LV	ES atbilstības deklarācija	Pamata UDI-DI	Pamata UDI-DI nosaukums
LT	ES ATITIKTIES DEKLARACIJA	Bazinis UDI-DI	Bazinio UDI-DI pavadinimas
NO	EU-samsvarserklæring	Grunnleggende UDI-DI	Grunnleggende UDI-DI-navn
PL	DEKLARACJA ZGODNOŚCI UE	Kod Basic UDI-DI	Nazwa kodu Basic UDI-DI
PT	DECLARAÇÃO UE DE CONFORMIDADE	UDI-DI básico	Nome UDI-DI Básico
RO	Declarația de Conformitate UE	UDI-DI de bază	Nume UDI-DI de bază
SK	EÚ VYHLÁSENIE O ZHODE	Základný UDI-DI	Názov základného UDI-DI
SV	EU-FÖRSÄKRAN OM ÖVERENSSTÄMMELSE	Grundläggande UDI-DI	Namn på grundläggande UDI-DI
TR	AB Uygunluk Beyanı	Temel UDI-DI	Temel UDI-DI İsmi

EN	Risk Class	List Number and Size Code	Product and Trade Name
BG	Клас според риска	Каталожен номер и код на размера	Име на продукта и търговско наименование
CS	Riziková třída	Katalogové číslo a koncové dvojčíslí určující velikost soupravy	Název produktu a obchodní název
DA	Risikoklasse	Bestillingsnummer og størrelseskode	Produkt- og varemærkenavn
DE	Risikoklasse	Bestellnummer und Größencode	Produkt- und Handelsname
EL	Κατηγορία κινδύνου	Κωδικός Προϊόντος και Κωδικός Συσκευασίας	Προϊόν και Εμπορική Ονομασία
ES	Clase de riesgo	Número de referencia y código de tamaño	Producto y marca comercial
ET	Riskiklass	Katalooginumber ja suurusekood	Toote nimetus ja kaubanimi
FR	Classe de risque	Référence	Nom de produit et de marque
HR	Klasa rizika	Kataloški broj i oznaka pakiranja	Naziv proizvoda i zaštićeni naziv
HU	Kockázati osztály	Listaszám és készletkészlet-kód	Termék- és kereskedelmi név
IT	Classe di rischio	Numero di listino e codice formato	Prodotto e nome commerciale
LV	Riska klase	Kataloga numurs un izmēra kods	Produkta nosaukums un tirdzniecības nosaukums
LT	Rizikos klasė	Katalogo numeris ir dydžio kodas	Gaminio ir prekės pavadinimai
NO	Risikoklasse	Bestillingsnummer og størrelseskode	Produkt- og handelsnavn
PL	Klasa ryzyka	Numer katalogowy	Nazwa produktu i nazwa handlowa
PT	Classe de risco	Número de lista e código de apresentação	Produto e nome comercial
RO	Clasă de risc	Număr de listă și cod dimensiune	Denumirea produsului și denumirea comercială
SK	Riziková trieda	Katalógové číslo	Názov produktu a obchodný názov
SV	Riskklass	Listnummer och storlekskod	Produkt och firmanamn
TR	Risk Sınıfı	Liste Numarası ve Boyut Kodu	Ürün ve Ticari İsmi

EN	GMDN Code	EMDN Code	Manufacturer (Name and Address)	Manufacturer SRN
BG	Код GMDN	Код EMDN	Производител (име и адрес)	EPH на производителя
CS	Kód GMDN	Kód EMDN	Výrobce (název a adresa)	Jediné registrační číslo výrobce
DA	GMDN-kode	EMDN-kode	Fabrikant (navn og adresse)	Fabrikants SRN
DE	GMDN-Code	EMDN-Code	Hersteller (Name und Adresse)	Hersteller-SRN
EL	Κωδικός GMDN (Ονοματολογία ιατροτεχνολογικών προϊόντων)	Κωδικός EMDN (Ονοματολογία ιατροτεχνολογικών προϊόντων)	Κατασκευαστής (Όνομα και Διεύθυνση)	SRN (Μοναδικός Αριθμός Μητρώου) Κατασκευαστή
ES	Código GMDN	Código EMDN	Fabricante (nombre y dirección)	SRN (número de registro único) del fabricante
ET	GMDN-kood	EMDN-kood	Tootja (nimi ja aadress)	Tootja unikaalne registreerimisnumber
FR	Code GMDN	Code EMDN	Fabricant (nom et adresse)	Numéro d'enregistrement unique du fabricant
HR	GMDN kod	EMDN kod	Proizvođač (naziv i adresa)	SRN (jedinstveni registracijski broj) proizvođača
HU	GMDN-kód	EMDN-kód	Gyártó (név és cím)	Gyártó egyedi regisztrációs száma (SRN)
IT	Codice GMDN	Codice EMDN	Fabbricante (nome e indirizzo)	SRN (numero di registrazione unico) del fabbricante
LV	GMDN kods	EMDN kods	Ražotājs (nosaukums un adrese)	Ražotāja vienotais reģistrācijas numurs (VRN)
LT	Visuotinės medicinos priemonių nomenklatūros kodas	Europos medicinos priemonių nomenklatūros kodas	Gamintojas (pavadinimas ir adresas)	Gamintojo unikalusi registracijos numeris
NO	GMDN-kode	EMDN-kode	Produsent (navn og adresse)	Produsentens SRN
PL	Kod GMDN	Kod Europejskiej Nomenklatury Wyrobów Medycznych	Producent (nazwa i adres)	Niepowtarzalny numer rejestracyjny producenta
PT	Código GMDN	Código EMDN	Fabricante (Nome e Morada)	Número único de registo do fabricante
RO	Cod GMDN	Cod EMDN	Producător (nume și adresă)	SRN producător
SK	Kód GMDN	Kód EMDN	Výrobca (Názov a adresa)	Jediné registračné číslo (SRN) výrobcu
SV	GMDN-kod	EMDN-kod	Tillverkare (namn och adress)	Tillverkarens SRN
TR	GMDN Kodu	EMDN Kodu	Üretici (İsim ve Adres)	Üretici SRN'si

EN	Authorized Representative (Name and Address)	Authorized Representative SRN	Produced by (Site of Manufacture) (Name and Address)
BG	Упълномощен представител (име и адрес)	ЕРН на упълномощения представител	Произведено от (място на производство) (име и адрес)
CS	Zplnomocněný zástupce (název a adresa)	Jediné registrační číslo zplnomocněného zástupce	Vyrobeno (místo výroby) (název a adresa)
DA	Autoriseret repræsentant (navn og adresse)	Autoriseret repræsentants SRN	Produceret af (fabrikationssted) (Navn og adresse)
DE	Bevollmächtigter (Name und Adresse)	SRN des Bevollmächtigten	Hergestellt von (Herstellungsstandort) (Name und Adresse)
EL	Εξουσιοδοτημένος Αντιπρόσωπος (Όνομα και Διεύθυνση)	SRN Εξουσιοδοτημένου Αντιπροσώπου	Κατασκευάζεται από (Εργοστάσιο παραγωγής) (Όνομασία και Διεύθυνση)
ES	Representante autorizado (nombre y dirección)	SRN (número de registro único) del representante autorizado	Producido por (Lugar de fabricación) (Nombre y dirección)
ET	Volitatud esindaja (nimi ja aadress)	Volitatud esindaja unikaalne registreerimisnumber	Tootnud (tootmiskoht) (nimi ja aadress)
FR	Mandataire (nom et adresse)	Numéro d'enregistrement unique du mandataire	Produit par (site de fabrication) (nom et adresse)
HR	Ovlašteni zastupnik (naziv i adresa)	SRN (jedinstveni registracijski broj) ovlaštenog zastupnika	Proizvodi (Mjesto proizvodnje) (Naziv i adresa)
HU	Meghatalmazott képviselő (név és cím)	Meghatalmazott képviselő egyedi regisztrációs száma (SRN)	Gyártó (gyártás helye) (név és cím)
IT	Mandatario (nome e indirizzo)	SRN (numero di registrazione unico) del mandatario	Prodotto da (sito di fabbricazione) (nome e indirizzo)
LV	Pilnvarotais pārstāvis (nosaukums un adrese)	Pilnvarotā pārstāvja vienotais reģistrācijas numurs (VRN)	Ražots (ražošanas vieta) (nosaukums un adrese)
LT	Igaliojasis atstovas (pavadinimas ir adresas)	Igaliojojo atstovo unikalusi registracijos numeris	Pagaminta (gamybos vieta) (pavadinimas ir adresas)
NO	Autorisert representant (navn og adresse)	Den autoriserte representantens SRN	Produsert av (produksjonssted) (navn og adresse)
PL	Upoważniony przedstawiciel (nazwa i adres)	Niepowtarzalny numer rejestracyjny upoważnionego przedstawiciela	Wyprodukowano przez (miejsce produkcji) (nazwa i adres)
PT	Mandatário (Nome e Morada)	Número único de registo do mandatário	Produzido por (Local de fabrico) (Nome e Morada)
RO	Reprezentant autorizat (nume și adresă)	SRN reprezentant autorizat	Produs de către (locuție producție) (nume și adresă)
SK	Autorizovaný zástupca (názov a adresa)	Jediné registračné číslo (SRN) autorizovaného zástupcu	Výrobené (miesto výroby) (názov a adresa)
SV	Auktoriserad representant (namn och adress)	Auktoriserad representants SRN	Tillverkas av (tillverkningsort) (namn och adress)
TR	Yetkili Temsilci (İsim ve Adres)	Yetkili Temsilci SRN'si	Üretici (Üretim Tesisi) (İsim ve Adres)

EN	Conformity Assessment Procedure	Annex II and III	Full Name
BG	Процедура за оценка на съответствието	Приложение II и III	Пълно наименование
CS	Postup posuzování shody	Příloha II a III	Celý název
DA	Overensstemmelsesvurderingsprocedure	Bilag II og III	Fulde navn
DE	Konformitätsbewertungsverfahren	Anhang II und III	Vollständiger Name
EL	Διαδικασία αξιολόγησης συμμόρφωσης	Παράρτημα II και III	Πλήρης ονομασία
ES	Procedimiento de evaluación de la conformidad	Anexos II y III	Nombre completo
ET	Vastavushindamismenetlus	II ja III lisa	Täisnimi
FR	Procédure d'évaluation de la conformité	Annexes II et III	Nom complet
HR	Postupak ocjenjivanja sukladnosti	Prilog II. i III.	Puni naziv
HU	Megfelelőségértékelési eljárás	II. és III. melléklet	Teljes név
IT	Procedura di valutazione della conformità	Allegati II e III	Nome completo
LV	Atbilstības novērtēšanas procedūra	II un III pielikums	Pilns nosaukums
LT	Atitikties vertinimo procedūra	II ir III priedai	Vardas ir pavardė
NO	Framgangsmåte for samsvarsvurdering	Vedlegg II og III	Fullt navn
PL	Procedura oceny zgodności	Załącznik II oraz III	Imię i nazwisko
PT	Procedimento de avaliação da conformidade	Anexo II e III	Nome completo
RO	Procedură de evaluare a conformității	Anexa II și III	Numele complet
SK	Postup posudzovania zhody	Príloha II a III	Celý názov
SV	Förfarande för bedömning av överensstämmelse	Bilaga II och III	Fullständigt namn
TR	Uygunluk Değerlendirme Prosedürü	Ek II ve III	Adı Soyadı

EN	Function	Signed for, and on behalf of	Date Issued
BG	Длъжност	Подписано за и от името на	Дата на издаване
CS	Funkce	Podepsáno za a jménem	Datum vydání
DA	Funktion	Underskrevet for og på vegne af	Udstedelsesdato
DE	Funktion	Unterzeichnet für und im Auftrag von	Datum
EL	Λειτουργία	Υπογράφεται για και εκ μέρους του/της	Ημερομηνία έκδοσης
ES	Función	Firmada por, y en nombre de	Fecha
ET	Funktsioon	Alla kirjutanud (kelle poolt ja nimel)	Väljaandmise kuupäev
FR	Fonction	Signé par et au nom de	Date d'établissement
HR	Funkcija	Potpisano za i u ime	Datum izdavanja
HU	Beosztás	Aláíró a következő képviselőtől és nevében	Kiadás dátuma
IT	Funzione	Firmato a nome e per conto di	Data di rilascio
LV	Amats	Parakstīts šādas personas vārdā	Izdošanas datums
LT	Pareigos	Subjekto, kurio vardu pasirašoma, pavadinimas	Išdavimo data
NO	Funksjon	Signert for, og på vegne av	Ustedelsesdato
PL	Funkcja	Podpisano w imieniu	Data wydania
PT	Função	Assinado e em nome de	Data de emissão
RO	Funcția	Semnăt pentru și în numele	Data eliberării
SK	Funkcia	Podpísané za a v mene	Dátum vydania
SV	Funktion	Undertecknat för och på uppdrag av	Datum för utfärdande
TR	Görevi	Namına ve temsilen imza	Düzenlenme Tarihi

EN	Supersedes	Signature	Date of Approval
BG	Замества	Подпис	Дата на одобрение
CS	Nahrazuje	Podpis	Datum schválení
DA	Erstatter	Underskrift	Godkendelsesdato
DE	Ersetzt	Unterschrift	Datum der Genehmigung
EL	Αντικαθιστά	Υπογραφή	Ημερομηνία έγκρισης
ES	Sustituye	Firma	Fecha de aprobación
ET	Asendab	Allkiri	Heakskiitmise kuupäev
FR	Annule et remplace	Signature	Date de l'autorisation
HR	Zamjenjuje	Potpis	Datum odobrenja
HU	Hatálytalanítja a következő dokumentumot:	Aláírás	Jóváhagyás dátuma
IT	Sostituisce	Firma	Data di approvazione
LV	Aizstāj	Paraksts	Apstiprināšanas datums
LT	Pakeičia	Parašas	Patvirtinimo data
NO	Erstatter	Signatur	Godkjenningsdato
PL	Zastępuje	Podpis	Data zatwierdzenia
PT	Substitui	Assinatura	Data de aprovação
RO	Înlocuitor	Semnătură	Data aprobării
SK	Nahrádza	Podpis	Dátum schválenia
SV	Ersätter	Namnteckning	Datum för godkännande
TR	Yerini aldiği belge	İmza	Onay Tarihi

EN	Place Issued	Effective (Date or Lot Number)
BG	Място на издаване	В сила от/за (дата или номер на партида)
CS	Místo vydání	Účinné od (datum nebo číslo šarže)
DA	Udstedelsessted	Ikrafttrædelse (dato eller lotnummer)
DE	Ort	Gültig ab (Datum oder Chargenbezeichnung)
EL	Τόπος έκδοσης	Σε ισχύ από (Ημερομηνία ή αρ. партидаς)
ES	Expedida en	Efectiva (fecha o número de lote)
ET	Väljaandmise koht	Jõustumine (kuupäev või partiinumber)
FR	Lieu d'établissement	Entrée en vigueur (date ou numéro de lot)
HR	Mjesto izdavanja	Stupa na snagu (datum ili broj serije)
HU	Kiadás helye	Hatálybalépés (dátum vagy tételszám)
IT	Luogo di rilascio	Valido da (data o numero di lotto)
LV	Izdošanas vieta	Spēkā no (datums vai partijas numurs)
LT	Išdavimo vieta	Įsigalioja (data arba partijos numeris)
NO	Ustedelsessted	Gjelder fra (dato eller lotnummer)
PL	Miejsce wydania	Obowiązuje od (data lub numer partii)
PT	Local de emissão	Efetividade (Data ou número de lote)
RO	Locul eliberării	Valabilitate (data sau numărul lotului)
SK	Miesto vydania	Účinnosť od (dátum alebo číslo šarže)
SV	Plats för utfärdande	Verkställtigt (datum eller lotnummer)
TR	Düzenlendiği Yer	Yürürlük (Tarih veya Lot Numarası)

EN	We, the undersigned, hereby declare that the <i>in vitro</i> diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on <i>In Vitro</i> Diagnostic Medical Devices; and additionally conforms applicable provisions of Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, and to applicable provisions of Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery, and amending Directive 95/16/EC as transposed into the laws of the member states. This declaration is made in accordance with Annex IV of the IVD Regulation, Annex VI of the ROHS Directive, and Annex II of the Machinery Directive and is issued under the sole responsibility of the manufacturer.
BG	Ние, долуподписаните, с настоящото декларираме, че гореписаното(ите) медицинско(и) изделие(я) за <i>in vitro</i> диагностика отговаря(т) на приложимите разпоредби на Регламент (ЕС) 2017/746 на Европейския парламент и на Съвета от 5 април 2017 г. относно медицинските изделия за <i>in vitro</i> диагностика; освен това отговаря(т) на приложимите разпоредби на Директива 2011/65/ЕС на Европейския парламент и на Съвета от 8 юни 2011 г. относно ограничаването на употребата на определени опасни вещества в електрическото и електронното оборудване и на приложимите разпоредби на Директива 2006/42/ЕО на Европейския парламент и на Съвета от 17 май 2006 г. относно машините, и за изменение на Директива 95/16/ЕО, както е транспонирана в националното законодателство на държавите членки. Тази декларация се прави в съответствие с Приложение IV на Регламента за IVD, Приложение VI на Директивата за ограничаване на опасните вещества (ROHS) и Приложение II на Директивата относно машините и за нейното издаване отговорност носи единствено производителят.
CS	My, níže podepsaní, tímto prohlašujeme, že diagnostický(-é) zdravotnický(-é) prostředek (prostředky) <i>in vitro</i> uvedený(-é) výše je (jsou) ve shodě s příslušnými ustanoveními nařízení Evropského parlamentu a Rady (EU) 2017/746 ze dne 5. dubna 2017 o diagnostických zdravotnických prostředcích <i>in vitro</i>; a že je (jsou) dále ve shodě s příslušnými ustanoveními směrnice Evropského parlamentu a Rady 2011/65/EU ze dne 8. června 2011 o omezení používání některých nebezpečných látek v elektrických a elektronických zařízeních a s příslušnými ustanoveními směrnice Evropského parlamentu a Rady 2006/42/ES ze dne 17. května 2006 o strojních zařízeních a o změně směrnice 95/16/ES, jak byla provedena ve vnitrostátním právu členských států. Toto prohlášení je v souladu s Přílohou IV nařízení IVD, Přílohou VI směrnice ROHS a Přílohou II směrnice o strojních zařízeních a je vydáno na výhradní odpovědnost výrobce.
DA	Vi, undertegnede, erklærer herved, at det <i>in vitro</i>-diagnostiske medicinske udstyr, der er beskrevet ovenfor, er i overensstemmelse med de gældende bestemmelser i Europa-Parlamentets og Rådets forordning (EU) 2017/746 af 5. april 2017 om <i>in vitro</i>-diagnostisk medicinsk udstyr, ligesom det overholder gældende bestemmelser i Europa-Parlamentets og Rådets direktiv 2011/65/EU af 8. juni 2011 om begrænsning af brugen af visse farlige stoffer i elektrisk og elektronisk udstyr samt overholder gældende bestemmelser i Europa-Parlamentets og Rådets direktiv 2006/42/EF af 17. maj 2006 om maskiner og ændring af direktiv 95/16/EF, som det er transponeret i medlemsstaternes lovgivning. Denne erklæring afgives i overensstemmelse med IVD-forordningens bilag IV, ROHS-direktivets bilag VI samt maskindirektivets bilag II og udstedes under fabrikantens eneansvar.
DE	Wir, die Unterzeichner, erklären hiermit, dass das oben beschriebene <i>In-vitro</i>-Diagnostikum/die oben beschriebenen <i>In-vitro</i>-Diagnostika die entsprechenden Bestimmungen der Verordnung (EU) 2017/746 des Europäischen Parlaments und des Rates vom 5. April 2017 über <i>In-vitro</i>-Diagnostika erfüllen und zusätzlich die entsprechenden Bestimmungen der Richtlinie 2011/65/EU des Europäischen Parlaments und des Rates vom 8. Juni 2011 zur Beschränkung der Verwendung bestimmter gefährlicher Stoffe in Elektro- und Elektronikgeräten sowie der Richtlinie 2006/42/EG des Europäischen Parlaments und des Rates vom 17. Mai 2006 über Maschinen und zur Änderung der Richtlinie 95/16/EG gemäß Umsetzung in den Gesetzen der Mitgliedsstaaten. Diese Erklärung erfolgt gemäß Anhang IV der IVD-Verordnung, Anhang VI der RoHS-Richtlinie und Anhang II der Maschinen-Richtlinie und wird unter alleiniger Verantwortung des Herstellers ausgestellt.
EL	Εμείς, οι υπογράφωντες, δηλώνουμε με το παρόν ότι τα προαναφερόμενα διαγνωστικά ιατροτεχνολογικά προϊόντα συμμορφώνονται με τις ισχύουσες διατάξεις του Κανονισμού (ΕΕ) 2017/746 του Ευρωπαϊκού Κοινοβουλίου και του Συμβουλίου της 5^{ης} Απριλίου 2017 σχετικά με τα <i>in vitro</i> διαγνωστικά ιατροτεχνολογικά προϊόντα και επίσης συμμορφώνονται με τις ισχύουσες διατάξεις της Οδηγίας 2011/65/ΕΕ του Ευρωπαϊκού Κοινοβουλίου και του Συμβουλίου της 8^{ης} Ιουνίου 2011 σχετικά με τους περιορισμούς στη χρήση συγκεκριμένων επικινδυνών ουσιών στον ηλεκτρικό και ηλεκτρονικό εξοπλισμό, καθώς και με τις ισχύουσες διατάξεις της Οδηγίας 2006/42/ΕΚ του Ευρωπαϊκού Κοινοβουλίου και του Συμβουλίου της 17^{ης} Μαΐου 2006 σχετικά με τον μηχανικό εξοπλισμό και την τροποποιητική Οδηγία 95/16/ΕΚ όπως αυτή μεταφέρθηκε στη νομοθεσία των κρατών μελών. Η δήλωση αυτή γίνεται σύμφωνα με το Παράρτημα IV του Κανονισμού IVD, το Παράρτημα VI της Οδηγίας ROHS και το Παράρτημα II της Οδηγίας για τον μηχανικό εξοπλισμό και εκδίδεται με αποκλειστική ευθύνη του κατασκευαστή.
ES	Nosotros, los abajo firmantes, por la presente declaramos que el(los) producto(s) sanitario(s) para diagnóstico <i>in vitro</i> descrito(s) anteriormente cumple(n) las disposiciones aplicables del reglamento (UE) 2017/746 del Parlamento Europeo y del Consejo del 5 de abril de 2017 sobre productos sanitarios para diagnóstico <i>in vitro</i>; y además cumple(n) las disposiciones aplicables de la Directiva 2011/65/UE del Parlamento Europeo y del Consejo del 8 de junio de 2011 sobre restricciones a la utilización de determinadas sustancias peligrosas en aparatos eléctricos y electrónicos, y las disposiciones aplicables de la Directiva 2006/42/EC del Parlamento Europeo y del Consejo del 17 de mayo de 2006 sobre maquinaria, y la Directiva de enmienda 95/16/EC tal y como se ha incorporado en las leyes de los Estados Miembros. Esta declaración se realiza en conformidad con el Anexo IV del Reglamento IVD, Anexo VI de la Directiva ROHS y Anexo II de la Directiva de máquinas y es emitida bajo la exclusiva responsabilidad del fabricante.
ET	Meie, allakirjutanud, kinnitame, et eespool kirjeldatud <i>in vitro</i> diagnostikameditsiiniseadmed vastavad Euroopa Parlamendi ja nõukogu 5. aprilli 2017. aasta määruse (EL) 2017/746 (<i>in vitro</i> diagnostikameditsiiniseadmete kohta) kohaldatavatele sätetele ning lisaks vastab see kohaldatavatele sätetele Euroopa Parlamendi ja nõukogu 8. juuni 2011. aasta direktiivis 2011/65/EL (teatavate ohtlike ainete kasutamise piiramise kohta elektri- ja elektroonikaseadmetes) ja Euroopa Parlamendi ja nõukogu direktiivis 2006/42/EÜ, 17. mai 2006, mis käsitleb masinaid ja millega muudetakse direktiivi 95/16/EÜ, nagu see on üle võetud liikmesriikide seadustes. See deklaratsioon on koostatud vastavalt IVD määruse IV lisale, ROHS direktiivi VI lisale ja masinadirektiivi II lisale ning see on välja antud tootja vastutusel.
FR	Nous soussigné(s), déclarons par la présente que le(s) dispositif(s) médical(aux) de diagnostic <i>in vitro</i> indiqué(s) ci-dessus est/sont conforme(s) aux dispositions applicables du Règlement (UE) 2017/746 du Parlement européen et du Conseil du 5 avril 2017 relatif aux dispositifs médicaux de diagnostic <i>in vitro</i>, aux dispositions applicables de la Directive 2011/65/UE du Parlement européen et du Conseil du 8 juin 2011 relative à la limitation de l'utilisation de certaines substances dangereuses dans les équipements électriques et électroniques, aux dispositions applicables de la Directive 2006/42/CE du Parlement européen et du Conseil du 17 mai 2006 relative aux machines et modifiant la Directive 95/16/CE, telles que transposées dans le droit national des États membres. Cette déclaration est établie conformément à l'Annexe IV du Règlement DIV, à l'Annexe VI de la Directive ROHS ainsi qu'à l'Annexe II de la Directive Machines sous la seule responsabilité du fabricant.
HR	Mi, niže potpisani, ovim putem izjavljujemo da su gore navedeni <i>in vitro</i> dijagnostički medicinski proizvod(i) sukladni primjenjivim odredbama Uredbe (EU) 2017/746 Europskog parlamenta i Vijeća od 5. travnja 2017. o <i>in vitro</i> dijagnostičkim medicinskim proizvodima; i dodatno primjenjivim odredbama Direktive 2011/65/EU Europskog parlamenta i Vijeća od 8. lipnja 2011. o ograničenju uporabe određenih opasnih tvari u električnoj i elektroničkoj opremi, te primjenjivim odredbama Direktive 2006/42/EZ Europskog parlamenta i Vijeća od 17. svibnja 2006. o strojevima zamjenjujući Direktivu 95/16/EZ kako je preočeno u zakone država članica. Ova je izjava sastavljena u skladu s Prilogom IV. Uredbe IVD, Prilogom VI. Direktive ROHS i Prilogom II. Direktive o strojevima i izdaje se pod isključivom odgovornošću proizvođača.
HU	Alulírottak ezennel kijelentjük, hogy a fent leírt <i>in vitro</i> orvostechnikai eszköz(ök) megfelel(nek) az Európai Parlament és a Tanács <i>in vitro</i> diagnosztikai orvostechnikai eszközökről szóló (EU) 2017/746 (2017. április 5.) rendelete vonatkozó rendelkezésének; továbbá az Európai Parlament és a Tanács egyes veszélyes anyagok elektromos és elektronikus berendezéseiben való alkalmazásának korlátozásáról szóló 2011/65/EU (2011. június 8.) irányelve (RoHS irányelv) vonatkozó rendelkezéseit; valamint az Európai Parlament és a Tanács a gépekről és a 95/16/EK irányelv módosításáról szóló 2006/42/EK (2006. május 17.) irányelve vonatkozó rendelkezéseit a tagállamok jogrendjébe áttetté rendelkezéseknek. A jelen nyilatkozat megfelel az IVD rendelet IV. mellékletében, a RoHS irányelv VI. mellékletében és a gépekről szóló irányelv II. mellékletében foglalt előírásoknak, és a gyártó kizárólagos felelőssége alapján került kiadásra.

EN	We, the undersigned, hereby declare that the <i>in vitro</i> diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on <i>In Vitro</i> Diagnostic Medical Devices; and additionally conforms applicable provisions of Directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment, and to applicable provisions of Directive 2006/42/EC of the European Parliament and of the Council of 17 May 2006 on machinery, and amending Directive 95/16/EC as transposed into the laws of the member states. This declaration is made in accordance with Annex IV of the IVD Regulation, Annex VI of the ROHS Directive, and Annex II of the Machinery Directive and is issued under the sole responsibility of the manufacturer.
IT	Noi, i sottoscritti, con la presente dichiariamo che il(i) dispositivo(i) medico-diagnostico(i) <i>in vitro</i> sopra descritto(i) è(sono) conforme(i) alle disposizioni applicabili del regolamento (UE) 2017/746 del Parlamento europeo e del Consiglio del 5 aprile 2017 relativo ai dispositivi medico-diagnostici <i>in vitro</i>; è(sono) inoltre conforme(i) alle disposizioni applicabili della direttiva 2011/65/UE del Parlamento europeo e del Consiglio dell'8 giugno 2011 sulla restrizione dell'uso di determinate sostanze pericolose nelle apparecchiature elettriche ed elettroniche, e alle disposizioni applicabili della direttiva 2006/42/CE del Parlamento europeo e del Consiglio del 17 maggio 2006 relativa alle macchine e che modifica la direttiva 95/16/CE come recepite nelle legislazioni degli Stati membri. Questa dichiarazione è redatta in conformità all'allegato IV del regolamento IVD, all'allegato VI della direttiva ROHS e all'allegato II della direttiva macchine ed è rilasciata sotto la responsabilità esclusiva del fabbricante.
LV	Mēs, apakšā parakstījušies, ar šo paziņojam, ka iepriekš aprakstītā(-s) <i>in vitro</i> diagnostikas medicīniskā(-s) ierīce(-es) atbilst Eiropas Parlamenta un Padomes Regulas (ES) 2017/746 (2017. gada 5. aprīlis) piemērojamajām prasībām par <i>in vitro</i> diagnostikas medicīniskām ierīcēm un papildus prasībām, kas noteiktas Eiropas Parlamenta un Padomes Direktīvā 2011/65/ES (2011. gada 8. jūnijs) par dažu bīstamu vielu izmantošanas ierobežošanu elektriskās un elektroniskās iekārtās un Eiropas Parlamenta un Padomes Direktīvā 2006/42/EK (2006. gada 17. maijs) par mašīnām, un ar kuru groza Direktīvu 95/16/EK, kā tā ieviesta dalībvalstu tiesību aktos. Šī deklarācija ir sagatavota saskaņā ar IVD regulas IV pielikumu, ROHS direktīvas VI pielikumu un Direktīvas par mašīnām II pielikumu un par izdošanu atbild vienīgi ražotājs.
LT	Mes, toliau pasirašiusieji (-iusiosios), pareiškiame, kad anksčiau minėta (-os) <i>in vitro</i> diagnostikos medicinos priemonė (-ės) atitinka 2017 m. balandžio 5 d. Europos Parlamento ir Tarybos reglamento (ES) 2017/746 dėl <i>in vitro</i> diagnostikos medicinos priemonių taikytinas nuostatas; taip pat ji (jos) atitinka 2011 m. birželio 8 d. Europos Parlamento ir Tarybos direktyvos 2011/65/ES dėl tam tikrų pavojingų medžiagų naudojimo elektros ir elektroninėje įrangoje apribojimo taikomas nuostatas ir 2006 m. gegužės 17 d. Europos Parlamento ir Tarybos direktyvos 2006/42/EB dėl mašinų, iš dalies keičiančios Direktyvą 95/16/EB, taikomas nuostatas, perkeltas į valstybių narių teisės aktus. Ši deklaracija yra parengta vadovaujantis IVD reglamento IV priedu, ROHS direktyvos VI priedu ir Mašinų direktyvos II priedu ir yra išduodama tik gamintojo atsakomybe.
NO	Vi, undertegnede, erklærer herved at utstyret til <i>in vitro</i>-diagnostikk som er anført ovenfor, er i samsvar med gjeldende bestemmelser i Europaparlaments- og rådsforordning (EU) 2017/746 av 5. april 2017 om medisinsk utstyr til <i>in vitro</i>-diagnostikk, og ytterligere overholder gjeldende bestemmelser i Europaparlaments- og rådsdirektiv 2011/65/EU av 8. juni 2011 om bruksbegrensninger av visse farlige stoffer i elektrisk og elektronisk utstyr, og til gjeldende bestemmelser i Europaparlaments- og rådsdirektiv 2006/42/EF av 17. mai 2006 om maskiner, og endring av direktiv 95/16/EF som innarbeidet i medlemsstatenes lovgivning. Denne erklæringen er utarbeidet i overensstemmelse med vedlegg IV i IVD-forordningen, vedlegg VI i ROHS-direktivet og vedlegg II i maskindirektivet og er utstedt under produsentens eneansvar.
PL	My, niżej podpisani, niniejszym oświadczamy, że wymieniony(-e) powyżej wyrób(wyroby) medyczny(-e) do diagnostyki <i>in vitro</i> spełnia(-ją) odpowiednie wymagania Rozporządzenia (UE) 2017/746 Parlamentu Europejskiego i Rady z dnia 5 kwietnia 2017 r. w sprawie wyrobów medycznych do diagnostyki <i>in vitro</i>, a ponadto wymagania Dyrektywy 2011/65/UE Parlamentu Europejskiego i Rady z dnia 8 czerwca 2011 r. w sprawie ograniczenia stosowania niektórych niebezpiecznych substancji w sprzęcie elektrycznym i elektronicznym, Dyrektywy 2006/42/WE Parlamentu Europejskiego i Rady z dnia 17 maja 2006 r. w sprawie maszyn, zmieniającej Dyrektywę 95/16/WE, w sposób, w jaki zostały one wdrożone do ustawodawstwa państw członkowskich. Niniejsza deklaracja została sporządzona zgodnie z Załącznikiem IV Rozporządzenia IVDR, Załącznikiem VI Dyrektywy ROHS oraz Załącznikiem II Dyrektywy Maszynowej i wydana na wyłączną odpowiedzialność producenta.
PT	Nós, abaixo assinados, declaramos que os dispositivos médicos para diagnóstico <i>in vitro</i> descritos acima estão em conformidade com as disposições aplicáveis do Regulamento (UE) 2017/746 do Parlamento Europeu e do Conselho, de 5 de abril de 2017, relativo aos dispositivos médicos para diagnóstico <i>in vitro</i>; e adicionalmente, em conformidade com as disposições aplicáveis da Diretiva 2011/65/UE do Parlamento Europeu e do Conselho, de 8 de junho de 2011, relativa à restrição do uso de determinadas substâncias perigosas em equipamentos elétricos e eletrônicos, e com as disposições aplicáveis da Diretiva 2006/42/CE do Parlamento Europeu e do Conselho, de 17 de maio de 2006, sobre máquinas e que altera a Diretiva 95/16/CE, conforme transposta nas leis dos Estados membros. Esta declaração é feita de acordo com o Anexo IV do Regulamento IVD, o Anexo VI da Diretiva ROHS e o Anexo II da Diretiva relativa às Máquinas e é emitida sob a exclusiva responsabilidade do fabricante.
RO	Subsemnatii, declarăm că dispozitivul (dispozitivele) medical(e) pentru diagnostic <i>in vitro</i> descrie mai sus sunt conforme cu dispozițiile aplicabile din Regulamentul (UE) 2017/746 al Parlamentului European și al Consiliului din 5 aprilie 2017 privind Dispozitivele medicale pentru diagnosticul <i>in vitro</i>; și, în plus, respectă dispozițiile aplicabile din Directiva 2011/65/UE a Parlamentului European și a Consiliului din 8 iunie 2011 privind restricția utilizării anumitor substanțe periculoase în echipamentele electrice și electronice și cu dispozițiile aplicabile din Directiva 2006/42/CE a Parlamentului European și a Consiliului din 17 mai 2006 privind utilajele și modificarea Directivei 95/16/CE, transpusă în legile statelor membre. Prezenta declarație este emisă în conformitate cu anexa IV la Regulamentul IVD, anexa VI la Directiva ROHS și anexa II la Directiva utilajelor și este emisă sub responsabilitatea exclusivă a producătorului.
SK	My, dolupodpisani, týmto vyhlasujeme, že diagnostická(-é) zdravotnícka(-e) pomôcka(-y) <i>in vitro</i> uvedená(-é) vyššie je (sú) v zhode s príslušnými ustanoveniami Nariadenia Európskeho parlamentu a Rady (EÚ) 2017/746 z 5. apríla 2017 o diagnostických zdravotníckych pomôckach <i>in vitro</i>; a že je (sú) ďalej v zhode s príslušnými ustanoveniami Smernice Európskeho parlamentu a Rady 2011/65/EÚ z 8. júna 2011 o obmedzení používania určitých nebezpečných látok v elektrických a elektronických zariadeniach a s príslušnými ustanoveniami Smernice Európskeho parlamentu a Rady 2006/42/ES zo 17. mája 2006 o strojových zariadeniach a o zmene a doplnení Smernice 95/16/ES tak, ako boli transponované do zákonov členských štátov. Toto vyhlásenie je v súlade s Prílohou IV k Nariadeniu IVD, Prílohou VI k Smernici ROHS a Prílohou II k Smernici o strojových zariadeniach a vydáva sa na výhradnú zodpovednosť výrobcu.
SV	Vi, undertecknade, försäkrar härmed att den eller de medicintekniska produkter för <i>in vitro</i>-diagnostik som beskrivs ovan överensstämmer med de tillämpliga bestämmelserna i Europaparlamentets och rådets förordning (EU) 2017/746 av den 5 april 2017 om medicintekniska produkter för <i>in vitro</i>-diagnostik samt även överensstämmer med de tillämpliga bestämmelserna i Europaparlamentets och rådets direktiv 2011/65/EU av den 8 juni 2011 om begränsning av användning av vissa farliga ämnen i elektrisk och elektronisk utrustning samt med de tillämpliga bestämmelserna i Europaparlamentets och rådets direktiv 2006/42/EG av den 17 maj 2006 om maskiner och om ändring av direktiv 95/16/EG (omarbeting) som införlivats i medlemsstaternas lagstiftning. Denna försäkran görs i enlighet med bilaga IV till IVD-förordningen, bilaga VI till ROHS-direktivet samt bilaga II till maskindirektivet och utfärdas under tillverkarens enskilda ansvar.
TR	Biz, aşağıda imzaları bulunan, yukarıda belirtilen <i>in vitro</i> diagnostik tıbbi cihazların, 2017/746 sayılı Avrupa Parlamentosu (AB) Yönetmeliği ile 5 Nisan 2017 tarihli <i>In Vitro</i> Diagnostik Tıbbi Cihazlar Konseyinin ilgili hükümlerine uygun olduğunu ve ayrıca elektrikli ve elektronik cihazlarda belirli tehlikeli maddelerin kullanımının sınırlandırılmasına ilişkin 8 Haziran 2011 tarihli Konseyin ve 2011/65/EU sayılı Avrupa Parlamentosu Direktifinin ilgili hükümlerine, makinelere ilişkin 17 Mayıs 2006 tarihli Konseyin ve 2006/42/EC sayılı Avrupa Parlamentosu Direktifinin ilgili hükümlerine ve üye devlet yasalarına aktarılan 95/16/EC sayılı Ek Direktife uygun olduğunu beyan ederiz. Bu beyan IVD Yönetmeliği Ek IV, ROHS Direktifi Ek VI ve Makineler Direktifi Ek II uyarınca yapılmıştır ve üreticinin münhasır sorumluluğu altında yayımlanmıştır.

End of form

Declaration of Conformity

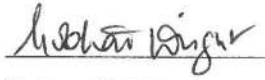
Certificate Identification: 04T84
Legal Manufacturer's Name: Abbott Ireland Diagnostics Division
Legal Manufacturer's Address: Lisnamuck, Longford, Co. Longford, Ireland.

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
04T8420	52925	Alanine Aminotransferase2	Self-declared
04T8430	52925	Alanine Aminotransferase2	Self-declared

Authorized European Representative (name and address)	Not Applicable
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: 
Full Name (printed): **Siobhan Wright**
Position: **Director Quality Assurance/
Site Quality Head**

Signature: 
Full Name (printed): **Thomas Breslin**
Position: **Manager Regulatory Affairs**

Date of Approval: 17-SEP-2021

Date of Approval: 17-SEP-2021

Date Issued: 17-SEP-2021

Place Issued: Abbott Ireland Diagnostics Division,
Lisnamuck, Longford, Co. Longford, Ireland.

Supersedes: Not Applicable

Effective Date: 17-SEP-2021

Declaration of Conformity

Certificate Identification: 04T85
 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
04T8520	52941	Amylase2	Self-declared

Authorized European Representative (name and address)	Not Applicable
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: *Siobhan Wright*
 Full Name (printed): **Siobhan Wright**
 Position: **Director Quality Assurance/
Site Quality Head**

Signature: *Lorraine Whitney*
 Full Name (printed): **Lorraine Whitney**
 Position: **Director Regulatory Affairs**

Date of Approval: 25-05-20

Date of Approval: 25 OCT 2020

Date Issued: 25-05-20

Place Issued: Abbott Ireland Diagnostics Division,
Lisnamuck, Longford, Co. Longford, Ireland.

Supersedes: Not Applicable

Effective Date: 25-05-20

 Biokit A Werfen Company	CE DECLARATION OF CONFORMITY	DRC-726
		Edition 5
	P-172	Page 1 of 2

CE DECLARATION OF CONFORMITY

Manufacturer: Hersteller Fabricante Fabricant Produttore	Fabricante Producent Tilvirkare Κατασκευαστής	BIOKIT, S.A. Av. Can Montcau, 7 08186 Lliçà d'Amunt Barcelona Spain
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Biokit hereby declares that the product(s) listed below conform to the European Union directive and standards identified in this declaration.

Biokit erkl rt, dass die aufgef hrten Produkt(e) mit den Bestimmungen der angegebenen EU-Richtlinien und mit den aufgef hrten normativen Dokumenten in  bereinstimmung sind.

Biokit declara por la presente que los producto(s) abajo mencionados, est n conformes con las directivas y normas Europeas identificadas en esta declaraci n.

Biokit d clare par la pr sente, que le(s) produit(s) sous-mentionn (s), est (sont) conforme(s) aux directives et normes Europ ennes identifi es dans cette d claration.

Biokit dichiara con la presente che il(l) prodotto(i) sottomenzionato(i)  (sono) conformi alla direttiva e agli standard specificati in questa dichiarazione.

Biokit declara pelo presente que o(s) produto(s) abaixo mencionado(s) est /est o conforme a Directiva e normas da Comiss o Europeia especificadas nesta declara  o.

Biokit erkl rer herved, at det (de) nedenfor anf rte produkt(er) er i overensstemmelse med de EU-direktiver og standarder, der er anf rt i denne erkl ring.

Biokit bekr ftar h rmed att nedan uppr knade produkt(er)  r f renlig(a) med de EU-direktiv och standarder som identifieras i denna deklaration

Η Biokit με το παρόν δηλώνει ότι το προϊόν(-τα) που αναφέρονται κατωτέρω συμμορφώνονται με την οδηγία της Ευρωπαϊκής Ένωσης και τα πρότυπα που παρατίθενται στην παρούσα δήλωση.

EU Directive:

EU-Richtlinie Directiva UE Directive Europ enne Direttiva Europea Directiva UE EU-Direktiv EU Direktiv Οδηγία ΕΕ

IVD - 98/79/EC (27/10/1998)

Standard(s):

Normen und Richtlinien Est andar(es) Norme(s) Norma(e) Padr o/Padr es Standard(er) Standard(er) Πρότυπο(-α)

ISO 13485

 Biokit A Werfen Company	CE DECLARATION OF CONFORMITY	DRC-726
		Edition 5
	P-172	Page 2 of 2

Notified Body:

Benannte Stelle Organismo Notificado Organisme Notifié Organismo Notificato Organismo Notificado Teknisk Kontrollorgan
Anmält Organ Κοινοποιημένος Οργανισμός

Name: Other Devices	Code: N/A
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• Certificate N°: N/A

Annex III

Product(s):

Produkt(e) Producto(s) Produkt(s) Prodotto(i) Produto(s) Produkt(er) Produkt(er) Προϊόν(-τα)

Product(s)	
Produkt(e)	Produto(s)
Producto(s)	Produkt(er)
Produkt(s)	Produkt(er)
Prodotto(i)	Προϊόν(-τα)
P/N	
01R0620	Alinity c ASO Reagent (300 test)
01R0630	Alinity c ASO Reagent (780 test)
01R0601	Alinity c ASO Standard

Signature
Pau Planas
CEO
Biokit, S.A

Date

August 28th, 2018

Declaration of Conformity

Certificate Identification:	<u>04T86</u>
Legal Manufacturer's Name:	<u>Abbott Ireland Diagnostics Division</u>
Legal Manufacturer's Address:	<u>Lisnamuck, Longford, Co. Longford, Ireland.</u>

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
04T8620	52954	Aspartate Aminotransferase2	Self-declared
04T8630	52954	Aspartate Aminotransferase2	Self-declared

Authorized European Representative (name and address)	Not Applicable
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: 
Full Name (printed): **Siobhan Wright**
Position: **Director Quality Assurance/
Site Quality Head**

Signature: 
Full Name (printed): **Thomas Breslin**
Position: **Manager Regulatory Affairs**

Date of Approval: 17-SEP-2021

Date of Approval: 17-SEP-2021

Date Issued: 17-SEP-2021

Place Issued: **Abbott Ireland Diagnostics Division,
Lisnamuck, Longford, Co. Longford, Ireland.**

Supersedes: **Not Applicable**

Effective Date: 17-SEP-2021

Declaration of Conformity

Certificate Identification: DOC-08P6101-SD DLK TPM
Legal Manufacturer's Name: Abbott GmbH
Legal 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
08P6101	41830	Alinity c Bilirubin Calibrator Kit	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038, 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. Becker

Full Name: **Claudia Becker**

Position: **Director Quality Assurance**

Date of Approval: 02 Feb 2022

Signature: Tiffini Jenkins

Full Name: **Tiffini Jenkins**

Position: **Manager Regulatory Affairs**

Date of Approval: 1-Feb-2022

Date Issued: 02 Feb 2022

Place Issued: **65205 Wiesbaden, Germany**

Supersedes: **05-April-2017**

Effective (Date or Lot Number): 02 Feb 2022

Declaration of Conformity

Certificate Identification: DOC-07P9720-SD DELK TPM
Legal Manufacturer's Name: Abbott GmbH
Legal 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
07P9720	53236	Alinity c Direct Bilirubin Reagent Kit	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038, 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. Becker

Full Name:

Claudia Becker

Position:

Director Quality Assurance

Date of Approval:

22 Jul 2021

Signature:

Tiffini Jenkins

Full Name:

Tiffini Jenkins

Position:

Manager Regulatory Affairs

Date of Approval:

11-Jul-2021

Date Issued:

22-Jul-2021

Place Issued:

65205 Wiesbaden, Germany

Supersedes:

19-Feb-2019

Effective (Date or Lot Number):

22-Jul-2021

Declaration of Conformity

Certificate Identification: DoC-04V5121, 04V5131-SD DELK
Legal Manufacturer's Name: Abbott GmbH
Legal 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
04V5121	53229	Alinity c Total Bilirubin Reagent Kit	Self-declared
04V5131	53229	Alinity c Total Bilirubin Reagent Kit	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038, 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:

Joerg Amborn

Position:

Director, Quality Assurance

Date of Approval:

2020-06-09

Signature:



Full Name:

Noah Lermer

Position:

Director Regulatory Affairs

Date of Approval:

12-Jun-20

Date Issued:

12-Jun-20

Place Issued:

65205 Wiesbaden, Germany

Supersedes:

27-Feb-2019

Effective (Date or Lot Number):

12-Jun-20

Declaration of Conformity

Certificate Identification: DOC-07P5720, 07P5730-SD DELK TPM
Legal Manufacturer's Name: Abbott GmbH
Legal 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
07P5720	45789	Alinity c Calcium Reagent Kit	Self-declared
07P5730	45789	Alinity c Calcium Reagent Kit	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038, 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. Becker

Full Name: **Claudia Becker**

Position: **Director Quality Assurance**

Date of Approval: 22 Jul 2021

Signature: Tiffini Jenkins

Full Name: **Tiffini Jenkins**

Position: **Manager Regulatory Affairs**

Date of Approval: 11-Jul-2021

Date Issued: 22-Jul-2021

Place Issued: 65205 Wiesbaden, Germany

Supersedes: 31-Dec-2016

Effective (Date or Lot Number): 22-Jul-2021



EU Declaration of Conformity

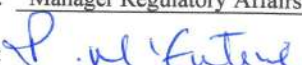
Basic UDI-DI: 038074ACT0488KF
Basic UDI-DI Name: Cholesterol2
Risk Class: Class B

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
04T8820	Cholesterol2	53359	W01010205
04T8830	Cholesterol2	53359	W01010205

Manufacturer (Name and Address)	Abbott Ireland Diagnostics Division Lisnamuck, Longford Co. Longford Ireland		
Manufacturer SRN	IE-MF-000010070		
Authorized Representative (Name and Address)	N/A		
Authorized Representative SRN	N/A		
Produced by (Site of Manufacture) (Name and Address)	Abbott Ireland Diagnostics Division Lisnamuck, Longford Co. Longford Ireland		
Notified Body (Name and Identification Number)	TÜV SÜD Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany Notified Body Number 0123		
Conformity Assessment Procedure	Quality Management System Annex IX Chapters I and III, Including an assessment of the technical documentation for devices concerned on the basis of representative samples	EU Certificate No. No. V12 054869 0013	
Common Specifications (CS)	N/A		

We, the undersigned, hereby declare that the in vitro diagnostic medical device(s) described above conform with the applicable provisions of the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on In Vitro Diagnostic Medical Devices. This declaration is made in accordance with Annex IV of the IVD Regulation and is issued under the sole responsibility of the manufacturer.

Full Name: David Spellman
Function: Director Quality Assurance/ Site Quality Head
Signature: 

Full Name: Rosemary McEntire
Function: Manager Regulatory Affairs
Signature: 

Date of Approval: 31 OCT 2024

Date of Approval: 31 Oct 2024

Signed for, and on behalf of: Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford Ireland

Date Issued: 31 OCT 2024

Place Issued: Lisnamuck, Longford Co. Longford Ireland
Effective (Date or Lot Number): 31 OCT 2024

Supersedes: 25-Sep-2023

Declaration of Conformity

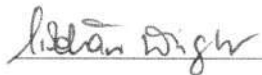
Certificate Identification: 04V62
Legal Manufacturer's Name: Abbott Ireland Diagnostics Division
Legal Manufacturer's Address: Lisnamuck, Longford, Co. Longford, Ireland.

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
04V6201	47868	Consolidated Chemistry Calibrator	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 and Randox Laboratories Ltd, 30 Randalstown Road, Antrim, Co. Antrim, BT41 4FL, UK
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 (printed): **Siobhan Wright**
Position: **Director Quality Assurance/
Site Quality Head**

Signature: 
Full Name (printed): **Thomas Breslin**
Position: **Manager Regulatory Affairs**

Date of Approval: 14-DEC-2021

Date of Approval: 15-DEC-2021

Date Issued: 15-DEC-2021

Place Issued: Abbott Ireland Diagnostics Division,
Lisnamuck, Longford, Co. Longford, Ireland.

Supersedes: 21 October 2021

Effective (Date or Lot Number): 15-DEC-2021