

Basic UDI-DI: Basic UDI-DI Name:

**Conformity Assessment Procedure** 

Common Specifications (CS)

038074ACT0491K4

Notified Body Number 0123

Quality Management System

Annex IX Chapters I and III,

representative samples

Including an assessment of the technical

DI Name: Creatinine2

Risk Class: Class B

List Number and Size Code	Product and Trade Name		GMDN Code	EMDN Code	
04T9120	Creatinine2		53251	W01010207	
(1)	Manufacturer Name and Address)	Abbott Ireland Diagnostics Division Lisnan	nuck, Longford Co. Longford I	reland	
Manufacturer SRN		IE-MF-000010070			
Authorized Representative		N/A			
(1	Name and Address)	0.000,930,000			
Authorized Representative SRN		N/A			
Produced by (Site of Manufacture)		Abbott Ireland Diagnostics Division Lisnamuck, Longford Co. Longford Ireland			
(Name and Address)			,		
Notified Body		TÜV SÜD Product Service GmbH,			
(Name and Identification Number)		Ridlerstraße 65, 80339 Munich, Germany			

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.

documentation for devices concerned on the basis of

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

EU Certificate No.

No. V12 054869 0013



for in vitro diagnostic medical device (IVD) - Directive 98/79/EC

The undersigned Sentinel CH. SpA, having its premises in Milan, Italy - Via Robert Koch 2, manufacturer of the family of devices named as "kits for clinical chemistry, immunochemistry, coagulation, molecular biology and rapid tests for immunology and serology" declares, under its own responsibility that the device

REF: 07P5620

Description: Alinity c CRP Vario Reagent Kit

EDMA: 12.11.01.09

complies with all the essential requirements listed in Annex I of the 98/79/EC Directive, as prescribed in Annex III of such Directive and its Italian transposition (legislative decree nr. 332/2000).

It therefore declares and assures, under its own responsibility, that the device:

- 1. complies with the applicable provisions of the Directive
- 2. is not included in the list A and B of Annex II of the Directive
- 3. is designed, manufactured and placed on the market according to the company certified quality system, in compliance with the current quality regulations as expressed in Annex III of the Directive.

Furthermore, the manufacturer declares to:

- 1. keep and make available for the Competent Authority the product technical file, as specified in Annex III of the 98/79/CE Directive, as well as to retain the batch records for a period of at least ten years after the production date of the last lot
- 2. have instituted and keep up to date an adequate procedure to guarantee the market surveillance requested by the Directive.

### DICHIARAZIONE DI CONFORMITA' CE

per dispositivo medico diagnostico in vitro (IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia - Via Robert Koch 2, fabbricante del dispositivo appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coagulazione, biologia molecolare e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che il dispositivo

REF:

07P5620

Descrizione: Alinity c CRP Vario Reagent Kit

EDMA: 12.11.01.09

soddisfa i requisiti essenziali applicabili richiesti dall'Allegato I della Direttiva 98/79/CE, come prescritto dall'Allegato III della medesima Direttiva e suo recepimento italiano (Decreto Legislativo 332/2000).

A tale scopo garantisce e dichiara sotto la propria responsabilità che il dispositivo:

- 1. soddisfa le disposizioni applicabili della Direttiva
- 2. non è incluso nell'elenco A e B dell'Allegato II della Direttiva
- 3. è progettato, fabbricato e immesso in commercio nell'ambito dell'applicazione di un sistema qualità aziendale dichiarato conforme e certificato secondo le norme vigenti così come descritto dall'Allegato III della Direttiva.

Il fabbricante dichiara inoltre di:

- 1. conservare e tenere a disposizione dell'Autorità Competente il fascicolo tecnico di prodotto, specificato nell'Allegato III della Direttiva 98/79/CE, nonché le registrazioni di produzione e controllo per un periodo di almeno dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza post-vendita richiesta dalla Direttiva.

Sentinel CH. SpA

A Legal Representative

Un Legale Rappresentante

Filippo De Luca

Date / Data

30/06/2017



for in vitro diagnostic medical device (IVD) - Directive 98/79/EC

The undersigned Sentinel CH. SpA, having its premises in Milan, Italy - Via Robert Koch 2, manufacturer of the family of devices named as "kits for clinical chemistry, immunochemistry, coagulation, molecular biology and rapid tests for immunology and serology" declares, under its own responsibility that the device

REF: 07P5601

Description: Alinity c CRP Vario Wide Range Calibrator Kit

EDMA: 12.50.03.13

complies with all the essential requirements listed in Annex I of the 98/79/EC Directive, as prescribed in Annex III of such Directive and its Italian transposition (legislative decree nr. 332/2000).

It therefore declares and assures, under its own responsibility, that the device:

- 1. complies with the applicable provisions of the Directive
- 2. is not included in the list A and B of Annex II of the Directive
- 3. is designed, manufactured and placed on the market according to the company certified quality system, in compliance with the current quality regulations as expressed in Annex III of the Directive.

### Furthermore, the manufacturer declares to:

- 1. keep and make available for the Competent Authority the product technical file, as specified in Annex III of the 98/79/CE Directive, as well as to retain the batch records for a period of at least ten years after the production date of the last lot
- 2. have instituted and keep up to date an adequate procedure to guarantee the market surveillance requested by the Directive.

### DICHIARAZIONE DI CONFORMITA' CE

per dispositivo medico diagnostico in vitro (IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia - Via Robert Koch 2, fabbricante del dispositivo appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coagulazione, biologia molecolare e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che il dispositivo

REF:

07P5601

Descrizione: Alinity c CRP Vario Wide Range Calibrator Kit

EDMA: 12.50.03.13

soddisfa i requisiti essenziali applicabili richiesti dall'Allegato I della Direttiva 98/79/CE, come prescritto dall'Allegato III della medesima Direttiva e suo recepimento italiano (Decreto Legislativo 332/2000).

A tale scopo garantisce e dichiara sotto la propria responsabilità che il dispositivo:

- 1. soddisfa le disposizioni applicabili della Direttiva
- 2. non è incluso nell'elenco A e B dell'Allegato II della Direttiva
- 3. è progettato, fabbricato e immesso in commercio nell'ambito dell'applicazione di un sistema qualità aziendale dichiarato conforme e certificato secondo le norme vigenti così come descritto dall'Allegato III della Direttiva.

### Il fabbricante dichiara inoltre di:

- 1. conservare e tenere a disposizione dell'Autorità Competente il fascicolo tecnico di prodotto, specificato nell'Allegato III della Direttiva 98/79/CE, nonché le registrazioni di produzione e controllo per un periodo di almeno dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza post-vendita richiesta dalla Direttiva.

Sentinel CH. SpA

A Legal Representative Un Legale Rappresentante

Filippo De Luca

Date / Data

30/06/2017



for in vitro diagnostic medical device (IVD) - Directive 98/79/EC

The undersigned Sentinel CH. SpA, having its premises in Milan, Italy - Via Robert Koch 2, manufacturer of the family of devices named as "kits for clinical chemistry, immunochemistry, coagulation, molecular biology and rapid tests for immunology and serology" declares, under its own responsibility that the device

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07P5602

Description: Alinity c CRP Vario High Sensitivity Calibrator Kit

EDMA: 12.50.03.13

complies with all the essential requirements listed in Annex I of the 98/79/EC Directive, as prescribed in Annex III of such Directive and its Italian transposition (legislative decree nr. 332/2000).

It therefore declares and assures, under its own responsibility, that the device:

- 1. complies with the applicable provisions of the Directive
- 2. is not included in the list A and B of Annex II of the Directive
- 3. is designed, manufactured and placed on the market according to the company certified quality system, in compliance with the current quality regulations as expressed in Annex III of the Directive.

### Furthermore, the manufacturer declares to:

- 1. keep and make available for the Competent Authority the product technical file, as specified in Annex III of the 98/79/CE Directive, as well as to retain the batch records for a period of at least ten years after the production date of the last lot
- 2. have instituted and keep up to date an adequate procedure to guarantee the market surveillance requested by the Directive.

### DICHIARAZIONE DI CONFORMITA' CE

per dispositivo medico diagnostico in vitro (IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia - Via Robert Koch 2, fabbricante del dispositivo appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coaqulazione, biologia molecolare e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che il dispositivo

REF:

07P5602

Descrizione: Alinity c CRP Vario High Sensitivity Calibrator Kit

EDMA: 12.50.03.13

soddisfa i requisiti essenziali applicabili richiesti dall'Allegato I della Direttiva 98/79/CE, come prescritto dall'Allegato III della medesima Direttiva e suo recepimento italiano (Decreto Legislativo 332/2000).

A tale scopo garantisce e dichiara sotto la propria responsabilità che il dispositivo:

- 1. soddisfa le disposizioni applicabili della Direttiva
- 2. non è incluso nell'elenco A e B dell'Allegato II della Direttiva
- è progettato, fabbricato e immesso in commercio nell'ambito dell'applicazione di un sistema qualità aziendale dichiarato conforme e certificato secondo le norme vigenti così come descritto dall'Allegato III della Direttiva.

#### Il fabbricante dichiara inoltre di:

- 1. conservare e tenere a disposizione dell'Autorità Competente il fascicolo tecnico di prodotto, specificato nell'Allegato III della Direttiva 98/79/CE, nonché le registrazioni di produzione e controllo per un periodo di almeno dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza post-vendita richiesta dalla Direttiva.

Sentine CH. SpA

A Legal Representative Un Legale Rappresentante

Filippo De Luca

Date / Data

30/06/2017



Mod. 98 - Rev.4 - Data:03/09/2013

## **EC DECLARATION OF CONFORMITY**

for in vitro diagnostic medical device (IVD) - Directive 98/79/EC

The undersigned Sentinel CH. SpA, having its premises in Milan, Italy - Via Robert Koch 2, manufacturer of the family of devices named as "kits for clinical chemistry, immunochemistry, coagulation, molecular biology and rapid tests for immunology and serology" declares, under its own responsibility that the device

REF:

07P5621

Description: Alinity c CRP Vario Reagent Kit

EDMA: 12.11.01.09

complies with all the essential requirements listed in Annex I of the 98/79/EC Directive, as prescribed in Annex III of such Directive and its Italian transposition (legislative decree nr. 332/2000).

It therefore declares and assures, under its own responsibility, that the device:

- 1. complies with the applicable provisions of the Directive
- 2. is not included in the list A and B of Annex II of the Directive
- 3. is designed, manufactured and placed on the market according to the company certified quality system, in compliance with the current quality regulations as expressed in Annex III of the Directive.

### Furthermore, the manufacturer declares to:

- 1. keep and make available for the Competent Authority the product technical file, as specified in Annex III of the 98/79/CE Directive, as well as to retain the batch records for a period of at least ten years after the production date of the last lot
- 2. have instituted and keep up to date an adequate procedure to guarantee the market surveillance requested by the Directive.

### DICHIARAZIONE DI CONFORMITA' CE

per dispositivo medico diagnostico in vitro (IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia - Via Robert Koch 2, fabbricante del dispositivo appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coagulazione, biologia molecolare e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che il dispositivo

REF: 07P5621

1 Descrizion

Descrizione: Alinity c CRP Vario Reagent Kit

EDMA: 12.11.01.09

soddisfa i requisiti essenziali applicabili richiesti dall'Allegato I della Direttiva 98/79/CE, come prescritto dall'Allegato III della medesima Direttiva e suo recepimento italiano (Decreto Legislativo 332/2000).

A tale scopo garantisce e dichiara sotto la propria responsabilità che il dispositivo:

- 1. soddisfa le disposizioni applicabili della Direttiva
- 2. non è incluso nell'elenco A e B dell'Allegato II della Direttiva
- 3. è progettato, fabbricato e immesso in commercio nell'ambito dell'applicazione di un sistema qualità aziendale dichiarato conforme e certificato secondo le norme vigenti così come descritto dall'Allegato III della Direttiva.

### Il fabbricante dichiara inoltre di:

- 1. conservare e tenere a disposizione dell'Autorità Competente il fascicolo tecnico di prodotto, specificato nell'Allegato III della Direttiva 98/79/CE, nonché le registrazioni di produzione e controllo per un periodo di almeno dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza post-vendita richiesta dalla Direttiva.

Sentinel CH. SpA

A Legal Representative Un Legale Rappresentante

Ugo De Luca / hun

Date / Data

12/12/2013



Mod. 98 - Rev.4 - Data:03/09/2013

### EC DECLARATION OF CONFORMITY

for in vitro diagnostic medical device (IVD) - Directive 98/79/EC

The undersigned Sentinel CH. SpA, having its premises in Milan, Italy - Via Robert Koch 2, manufacturer of the family of devices named as "kits for clinical chemistry, immunochemistry, coagulation, molecular biology and rapid tests for immunology and serology" declares, under its own responsibility that the device

REF:

07P5624

Description: Alinity c CRP Vario Reagent Kit

EDMA: **12.11.01.09** 

complies with all the essential requirements listed in Annex I of the 98/79/EC Directive, as prescribed in Annex III of such Directive and its Italian transposition (legislative decree nr. 332/2000).

It therefore declares and assures, under its own responsibility, that the device:

- 1. complies with the applicable provisions of the Directive
- 2. is not included in the list A and B of Annex II of the Directive
- 3. is designed, manufactured and placed on the market according to the company certified quality system, in compliance with the current quality regulations as expressed in Annex III of the Directive.

#### Furthermore, the manufacturer declares to:

- 1. keep and make available for the Competent Authority the product technical file, as specified in Annex III of the 98/79/CE Directive, as well as to retain the batch records for a period of at least ten years after the production date of the last lot
- 2. have instituted and keep up to date an adequate procedure to guarantee the market surveillance requested by the Directive.

### DICHIARAZIONE DI CONFORMITA' CE

per dispositivo medico diagnostico in vitro (IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia - Via Robert Koch 2, fabbricante del dispositivo appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coagulazione, biologia molecolare e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che il dispositivo

REF:

07P5624

Descrizione: Alinity c CRP Vario Reagent Kit

EDMA: 12.11.01.09

soddisfa i requisiti essenziali applicabili richiesti dall'Allegato I della Direttiva 98/79/CE, come prescritto dall'Allegato III della medesima Direttiva e suo recepimento italiano (Decreto Legislativo 332/2000).

A tale scopo garantisce e dichiara sotto la propria responsabilità che il dispositivo:

- 1. soddisfa le disposizioni applicabili della Direttiva
- 2. non è incluso nell'elenco A e B dell'Allegato II della Direttiva
- 3. è progettato, fabbricato e immesso in commercio nell'ambito dell'applicazione di un sistema qualità aziendale dichiarato conforme e certificato secondo le norme vigenti così come descritto dall'Allegato III della Direttiva.

#### Il fabbricante dichiara inoltre di:

- 1. conservare e tenere a disposizione dell'Autorità Competente il fascicolo tecnico di prodotto, specificato nell'Allegato III della Direttiva 98/79/CE, nonché le registrazioni di produzione e controllo per un periodo di almeno dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza post-vendita richiesta dalla Direttiva.

Sentinel CH. SpA

A Legal Representative Un Legale Rappresentante

Ugo De Luca

Date / Data

12/12/2018



as per Annex IV of the Reg. EU 2017/746 on in vitro diagnostic medical devices

#### SENTINEL CH. S.p.A.

Via Robert Koch, 2 - Milano (MI) - 20152 - Italy REGISTERED TRADEMARK: SENTINEL DIAGNOSTICS

SRN NUMBER: IT-MF-000012556

The present EU Declaration of Conformity is released under the sole responsibility of the SENTINEL CH. S.p.A., manufacturer of the devices listed in table 1.

REF	Device Commercial Name	Basic UDI-DI
06T3220	Cystatin C	805805668CICYSC02ZZ231N42
06T3230	Cystatin C	805805668CICYSC02ZZ231N42
06T3201	Cystatin C Calibrators	805805668CICYSC02ZZ231N42
06T3210	Cystatin C Controls	805805668CICYSC02ZZ231N42

Table 1 - Product list

REF	Intended purpose
06T3220	The Cystatin C assay is an in vitro diagnostic test used in the quantitative immunoturbidimetric determination
	of cystatin C in human serum and plasma on the Alinity c system.
06T3230	Measurement of cystatin C aids in the diagnosis and treatment of renal diseases.
	For laboratory professional use only.
06T3201	The Cystatin C Calibrators are for the calibration of the Alinity c system when used in the quantitative
	immunoturbidimetric determination of cystatin C in human serum and plasma.
	For additional information, refer to the Cystatin C reagent package insert and the Alinity ci-series Operations
	Manual. For laboratory professional use only.
06T3210	The Cystatin C Controls are for the estimation of test precision and the detection of systematic analytical
	deviations of the Alinity c system when used in the quantitative immunoturbidimetric determination of cystatin
	C in human serum and plasma.
	For additional information, refer to the Cystatin C reagent package insert and the Alinity ci-series Operations
	Manual. For laboratory professional use only.

Table 2 - Intended purposes

The above listed devices are classified in Class B according to the rules set out in Annex VIII of the Reg. EU 2017/746.

The above listed devices covered by the present declaration are in conformity with Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on in vitro diagnostic medical devices.

Common specifications: not applicable

### **Notified Body identification**

Name and identification number of the notified body: BSI Group The Netherlands B.V., 2797.

Description of the conformity assessment procedure performed as per Article 48:

- Quality Management System Assurance: Annex IX;
- Assessment of Technical Documentation: Annex IX.

Identification of the certificate: IVDR 752785.

Name:

Paolo Bosio

Place and Date: Milowe, 13 12 2024

Function:

A legal representative

(place, dd/mm/yyyy)

Signature:

(on behalf of the company)

Version: 2

mod. 468 - rev. 3 - 07/02/2023

Page 1/1



DRC-1269 Edition 8

File name: DRC-1269\_D-Dimer\_R02

Page 1 of 2

P-172-08

### **EU DECLARATION OF CONFORMITY**

	REF	Basic UDI-DI	EMDN Code
D-Dimer	01R1022	8436003070726XV	W0103 (Level 3) W010302 (Level 4)
D-Dimer Calibrator	01R1001	8436003070726XV	W0103 (Level 3) W010302 (Level 4)
D-Dimer Controls	01R1010	8436003070726XV	W0103 (Level 3) W010302 (Level 4)

#### INTENDED PURPOSE

**D-Dimer:** The D-Dimer is an automated clinical chemistry assay used for the quantitative determination of D-Dimer in human citrated plasma on the Alinity c system. The D-Dimer assay is to be used as an aid for diagnosing thrombosis and monitoring thrombolytic therapy in patients suspected to suffer a thrombotic event. Elevated levels of D-Dimer are found in clinical conditions such as deep vein thrombosis (DVT), pulmonary embolism (PE), and disseminated intravascular coagulation (DIC). D-Dimer levels also rise during the normal pregnancy, but very high levels are associated with complications. For laboratory professional use only.

**D-Dimer Calibrator:** The D-Dimer Calibrator is for the calibration of the Alinity c system when used for the quantitative determination of D-Dimer in human citrated plasma. For laboratory professional use only.

**D-Dimer Controls:** The D-Dimer Controls are for the estimation of test precision and the detection of systematic analytical deviations of the Alinity c system when used for the quantitative determination of D-Dimer in human citrated plasma. For laboratory professional use only.

	BIOKIT, S.A.
	Av. Can Montcau 7
	Lliçà d'Amunt (Barcelona) 08186 SPAIN
	SRN: ES-MF-000002406

We, as the manufacturer of the device(s), take sole responsibility for and hereby declare that the above-mentioned product(s) meet(s) the provisions of the following Regulation(s)/Directive(s):



DRC-1269 Edition 8

File name: DRC-1269\_D-Dimer\_R02

Page 2 of 2

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Regulation EU 2017/746 on In vitro Diagnostic Medical Devices					
RISK CLASS					
□A □B ☑C □D					

CONFORMITY ROUTE				
	EU CERTIFICATE #: V12 111106 0002 Rev.03 Name of Notified Body: TÜV SÜD Product Service GmbH, Ridlerstraße 65, 80339 Munich, Germany. Notified Body Identification.: 0123			

Androde .	Biokit, S.A., Lliçà d'Amunt	2025-01-15
Signature	Issued in	Date
José Luis Zarroca	CEO	
Name	Function	



Manufacturer:

Hersteller Fabricante Fabricant Produttore Fabricante Producent Tillverkare Κατασκευαστής BIOKIT, S.A. Av. Can Montcau, 7. 08186 Llicà d'Amunt

Barcelona - Spain

Biokit hereby declares that the product(s) listed below conform to the European Union directive and standards identified in this declaration.

Biokit erklärt, dass die aufgeführten Produkt(e) mit den Bestimmungen der angegebenen EU-Richtlinien und mit den aufgeführten normativen Dokumenten in Übereinstimmung sind.

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

Biokit déclare par la présente, que le(s) produit(s) sous-mentionné(s), est (sont) conforme(s) aux directives et normes Européennes identifiées dans cette déclaration.

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

Biokit declara pelo presente que o(s) produto(s) abaixo mencionado(s) está/estão conforme a Directiva e normas da Comissão Europeia especificadas nesta declaração.

Biokit erklærer herved, at det (de) nedenfor anførte produkt(er) er i overensstemmelse med de EU-direktiver og standarder, der er anført i denne erklæring.

Biokit bekräftar härmed att produkt(er) listade nedan, vara förenlig(a) med Europeiska Union-ens direktiv och standarder identifierade i denna deklaration.

Η Biokit με το παρόν δηλώνει ότι τα προϊόντα που αναφέρονται κατωτέρω συμμορφούνται με την οδηγία της Ευρωπαϊκής Ενωσης και τα πρότυπα που προσδιορίζονται στην παρούσα δήλωση.

#### **EU Directive:**

EU-Richtlinie Directiva UE Directive Européenne Direttiva Europea Directiva UE EU-direktiv EU Direktiv Οδηγία ΕΕ

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

### Standard(s):

Normen und Richtlinien Estándar(es) Norme(s) Norma(e) Padrāo/Padrões Standard(er) Standard(er) Πρότυπα

ISO 13485

Name:

José Luis Zarroca

CEO

Biokit, S.A.

Lliçà d'Amunt, 23<sup>rd</sup> November 2020 R01

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Product(s)		
Produkt(e)	Produto(s)	
Producto(s)	Produkt(er)	GMDN code
Produit(s Prodotto(i)	Produkt(er)	The American
P/N	Προϊόντα	
6K38-02	Quantia ASO	59055
6K39-02	Quantia 62-Microglobulin	53927
6K40-02	Quantia Digitoxin	59084
6K41-02	Quantia Ferritin	53718
6K42-02	Quantia IgE	61274
6K44-02	Quantia RF	55111
6K45-03	Quantia PROTEINS Standard	30505
6K46-03	Quantia ASO Standard	51744
6K47-03	Quantia ß2-Microglobulin Standard	38215
6K48-02	Quantia Digitoxin Standard	55330
6K49-03	Quantia Ferritin Standard	41927
6K50-03	Quantia IgE Standard	53777
6K52-03	Quantia RF Standard	42230
6K53-02	Quantia PROTEINS Control	30506
6K57-02	Quantia Digitoxin Control	38533
6K54-02	Quantia ASO RF Control I	30506
6K55-02	Quantia ASO RF Control II	30506
6K56-02	Quantia Ferritin/Myoglobin/IgE Control	30506
6K99-02	Quantia A1-Antitrypsin	53602
6L32-43	Quantia Myoglobin	59042
6L33-05	Quantia Myoglobin Standard	41733
6L34-43	Quantia A-1-AGP	53606
7K00-03	Quantia Lp(a)	53438
5P83-02	Lp(a) Calibrators	41417
5P84-11	Lp(a) Control	41418
7K02-02	Quantia D-Dimer	47346
7K02-21	Quantia D-Dimer Control	47347
7K02-11	Quantia D-Dimer Standard	47348

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Annex E	
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Spenned	ABBOTT

Certificate Identification:

04T96

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
04T9620	53030	Gamma-Glutamyl Transferase2	Self-declared
04T9630	53030	Gamma-Glutamyl Transferase2	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:

Signature:

Thomas Breslin

Full Name (printed):

Siobhan Wright

Full Name (printed):

Position:

Director Quality Assurance/

Position:

Manager Regulatory Affairs

Site Quality Head

Date of

09-SEP-2021

Date of

09 - Sep - 2021

Approval:

Approval:

Date Issued:

09-564-2021

Place Issued:

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

Supersedes:

Not Applicable

Effective Date:

09 - Sep - 2021



Certificate Identification:

DOC-07P5520, 07P5530-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
07P5520	53301	Alinity c Glucose Reagent Kit	Self-declared
07P5530	53301	Alinity c Glucose Reagent Kit	

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: Signature: Signature: Signature: Tiffini Jenkins

Position: Director Quality Assurance Position: Manager Regulatory Affairs

Date of Approval: 11-Jul-2021

Date of Approval: 11-Jul-2021

Date Issued: 22- Jul- 2021

Place Issued: 65205 Wiesbaden, Germany

Supersedes: 13-Oct-2017

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



Basic UDI-DI:	038074ACT0498KJ	
Basic UDI-DI Name:	Iron2	
Risk Class:	Class B	

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
04T9820	Iron2	54758	W01010216

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	
<b>Authorized Representative SRN</b>	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, Certification Body, Ridlerstraße 65, 80339 Munich Germany Notified Body Number 0123	
Conformity Assessment Procedure	Annex IX Chapters I and III,  Including an assessment of the technical documentation for devices concerned on the basis of	EU Certificate No. No. V12 054869 0013
Common Specifications (CS)	representative samples 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	Full Name:	Rosemary McEntire
Function:	Director Quality Assurance/Site Quality Head	Function:	Manager Regulatory Affairs
Signature:	Sell -	Signature:	L'M'Entrie
Date of Approval:	21 Nov 2023	Date of Approval:	21 NOV 2023
Signed for, and on behalf of:	Abbott Ireland Diagnostics Division Lisnamu	ck, Longford, Co. Lon	gford Ireland
Date Issued:	21 Nov 2023	Place Issued:	Lisnamuck, Longford, Co. Longford, Ireland
Supersedes:	09 December 2021	Effective (Date or Lot Number):	21 Nov 2023



Basic UDI-DI:

038074ACT0499KL

Basic UDI-DI Name:

Lactate Dehydrogenase2

Risk Class:

Class C

List Number and Size Code		Product and Trade Name	GMDN Code	EMDN Code
04T9920		Lactate Dehydrogenase2	53072	W01010119
04T9930		Lactate Dehydrogenase2	53072	W01010119
	Manufacturer Name and Address)	Abbott Ireland Diagnostics Division, Lisna	amuck, Longford, Co. Longford	Ireland
	Manufacturer SRN	IE-MF-000010070		
Authorized Representative N/A				

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	N/A	
<b>Authorized Representative SRN</b>	N/A		
Produced by (Site of Manufacture) (Name and Address)	Abbott Ireland Diagnostics Division, Lisnamuck, Lon	gford, 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:	Siobhan Wright Director Quality Assurance/Site Quality	Full Name:	Sandra Gallagher
Function:		Function:	Manager Regulatory Affairs
Signature:	libla Drigh	Signature:	S. Gallafor
Date of Approval:	14-062-2021		13-DEC-2021
Signed for, and on behalf of:	Abbott Ireland Diagnostics Division, Lisnamuc	k, Longford, Co. Lo	ngford Ireland
Date Issued:	14-0EC-2021	Place Issued:	Lisnamuck, Longford, Co. Longford, Ireland
Supersedes:	N/A	Effective (Date or Lot Number):	14-DEC-2021



B #			-		
IV /I	an	ıufa	30	rıır	or.
IVI	all	uli	20	LUI	CI.

Sekisui Diagnostics P.E.I. Inc

70 Watts Avenue Charlottetown

Prince Edward Island

C1E 2B9 Canada

European Representative:

MDSS GmbH

Schiffgraben 41 30175 Hannover

Germany

Product:

**Product Code** 

Name

**GMDN** Code

07P7120

Alinity c Direct LDL Reagent Kit

53395

Classification:

General IVD

Conformity Assessment Route: Annex III, self-certified

We hereby declare that the above-mentioned products meet the provisions of the Council Directive 98/79EC for in vitro diagnostic medical devices. All supporting documents are held by the manufacturer.

Place of Issue:

Prince Edward Island, Canada

Signature:

Penny White

Senior Manager Regulatory Affairs

Sekisui Diagnostics PEI Inc.

29-Jun-2021 Date



for in vitro diagnostic medical device (IVD) - Directive 98/79/EC

The undersigned Sentinel CH. SpA, having its premises in Milan, Italy - Via Robert Koch 2, manufacturer of the family of devices named as "kits for clinical chemistry, immunochemistry, coagulation, molecular biology and rapid tests for immunology and serology" declares, under its own responsibility that the device

REF: 04Y85-20

Description: Lipase NG OC Reagent Kit

EDMA: 11.01.01.23

complies with all the essential requirements listed in Annex I of the 98/79/EC Directive, as prescribed in Annex III of such Directive and its Italian transposition (legislative decree nr. 332/2000).

It therefore declares and assures, under its own responsibility, that the device:

- 1. complies with the applicable provisions of the Directive
- 2. is not included in the list A and B of Annex II of the Directive
- 3. is designed, manufactured and placed on the market according to the company certified quality system, in compliance with the current quality regulations as expressed in Annex III of the Directive.

### Furthermore, the manufacturer declares to:

- 1. keep and make available for the Competent Authority the product technical file, as specified in Annex III of the 98/79/CE Directive, as well as to retain the batch records for a period of at least ten years after the production date of the last lot
- 2. have instituted and keep up to date an adequate procedure to guarantee the market surveillance requested by the Directive.

### DICHIARAZIONE DI CONFORMITA' CE

per dispositivo medico diagnostico in vitro (IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia - Via Robert Koch 2, fabbricante del dispositivo appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coagulazione, biologia molecolare e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che il dispositivo

REF:

04Y85-20

Descrizione: Lipase NG OC Reagent Kit

EDMA: 11.01.01.23

soddisfa i requisiti essenziali applicabili richiesti dall'Allegato I della Direttiva 98/79/CE, come prescritto dall'Allegato III della medesima Direttiva e suo recepimento italiano (Decreto Legislativo 332/2000).

A tale scopo garantisce e dichiara sotto la propria responsabilità che il dispositivo:

- 1. soddisfa le disposizioni applicabili della Direttiva
- 2. non è incluso nell'elenco A e B dell'Allegato II della Direttiva
- 3. è progettato, fabbricato e immesso in commercio nell'ambito dell'applicazione di un sistema qualità aziendale dichiarato conforme e certificato secondo le norme vigenti così come descritto dall'Allegato III della Direttiva.

#### Il fabbricante dichiara inoltre di:

- 1. conservare e tenere a disposizione dell'Autorità Competente il fascicolo tecnico di prodotto, specificato nell'Allegato III della Direttiva 98/79/CE, nonché le registrazioni di produzione e controllo per un periodo di almeno dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza post-vendita richiesta dalla Direttiva.

Sentinel CH. SpA

A Legal Representative Un Legale Rappresentante

Ugo De Luça/4

Date / Data

28/02/2019



Certificate Identification:

DOC-08P19-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
08P1920	46795	Alinity c Magnesium Reagent Kit	Self-declared
08P1930	46795	Alinity c Magnesium Reagent Kit	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Fisher Diagnostics, a division of Fisher Scientific Company, LLC, a part of Thermo Fisher Scientific Inc., 8365 Valley Pike Middletown, VA 22645, 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:

a. .. . .

Signature:

Seffer genters

Full Name:

Claudia Becker

Full Name:

Tiffini Jenkins

Position:

Director Quality Assurance

Position:

Assoc. Director Regulatory Affairs

Date of Approval:

24 May 2022

Date of Approval:

24 - May - 2022

Date Issued:

Place Issued: 65

65205 Wiesbaden, Germany

Supersedes:

11-Jan-2019

Effective (Date or

Lot Number):

24 May 2022



04U0330

# **EU Declaration of Conformity**

Basic UDI-DI: Basic UDI-DI Name: 038074ACU0403JQ

Phosphorus2

Risk Class:

Phosphorus2 Class B

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
04U0320	Phosphorus2	59123	W01010307
0400320	Phoenhous?	59123	W01010307

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	C. J. C. I am afound Implement
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 Certification Body, 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 we, the undersigned, hereby declare that the in vide diagnostic frequent device(s) described above combinitivitin 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	Full Name:	Sandra Gallagher
Function:	Director Quality Assurance/ Site Quality Head	Function:	Manager Regulatory Affairs
Signature:	Buch	Signature:	S. Callafor
Date of Approval:	17 Apr 2023	Date of Approval:	17-APR-2023
Signed for, and on behalf of:	Abbott Ireland Diagnostics Division Lisnamuc	ck, Longford, Co. Lon	gford Ireland
Date Issued:	17 APR 2023	Place Issued:	Lisnamuck, Longford, Co. Longford, Ireland
Supersedes:	N/A	Effective (Date or Lot Number):	17 APR 2023

7	ADDOTT
	ABBOTT

Certificate Identification:

04T81

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
04T8120	53989	Total Protein2	Self-declared
04T8130	53989	Total Protein2	Self-declared

Authorized European	Not ∧pplicable
Representative (name and address)	
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:

Signature:

dorraine Whitney

Full Name (printed):

Siobhan Wright

Full Name (printed):

Lorraine Whitney

Position:

Director Quality Assurance/

Position:

**Director Regulatory Affairs** 

Site Quality Head

Date of Approval:

22-OLT-20

Date of Approval: 22 OLT 2020

Date Issued:

22.00 -20

Place Issued:

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

Supersedes:

Not Applicable

Effective Date:

22-005-20



**DRC-726** 

Edition 5

P-172

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# CE DECLARATION OF CONFORMITY

Manufacturer:

Hersteller Fabricante Fabricant Produttore Fabricante Producent Tillverkare Κατασκευαστής BIOKIT, S.A. Av. Can Montcau, 7 08186 Lliçà d'Amunt Barcelona Spain

Biokit hereby declares that the product(s) listed below conform to the European Union directive and standards identified in this declaration.

Biokit erklärt, dass die aufgeführten Produkt(e) mit den Bestimmungen der angegebenen EU-Richtlinien und mit den aufgeführten normativen Dokumenten in Übereinstimmung sind.

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

Biokit déclare par la présente, que le(s) produit(s) sous-mentionné(s), est (sont) conforme(s) aux directives et normes Européennes identifiées dans cette déclaration.

Biokit dichiara con la presente che ii(i) prodotto(i) sottomenzionato(i) è(sono) conformi alla direttiva e agli standard specificati in questa dichiarazione,

Biokit declara pelo presente que o(s) produto(s) abaixo mencionado(s) está/estão conforme a Directiva e normas da Comissão Europeia especificadas nesta declaração.

Biokit erklærer herved, at det (de) nedenfor anførte produkt(er) er i overensstemmelse med de EU-direktiver og standarder, der er anført i denne erklæring.

Biokit bekräfter härmed att nedan uppräknade produkt(er) är förenlig(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 Européenne Direttiva Europea Directiva UE EU-Direktiv EU Direktiv Οδηγία ΕΕ

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

### Standard(s):

Normen und Richtlinien Estándar(es) Norme(s) Norma(e) Padrão/Padrões Standard(er) Standard(er) Πρότυπο(-α)

ISO 13485



**DRC-726** 

Edition 5

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**Notified Body:** 

Benannte Stelle Organismo Notificado Organismo Notificado Organismo Notificado Organismo Notificado Teknisk Kontrollorgon Anmält Organ Κοινοποιημένος Οργανισμός

Name: Other Devices

Code: N/A

Mast 28th, 2018

Certificate Nº: N/A

Annex III

Product(s):

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

Product(s)	
Produkt(e)	Produto(s)
Producto(s)	Produkt(er)
Produit(s	Produkt(er)
Produit(s Prodotto(i)	Προϊόν(-τα)
P/N	
01R1622	Alinity c RF Reagent Kit (400 T)
01R1632	Alinity c RF Reagent Kit (920 T)
01R1601	Alinity c RF Standard

Signature Pau Planas CEO

Biokit, S.A

Date



Basic UDI-DI:

038074LFD0018KQ

Basic UDI-DI Name:

Transferrin Class B

Risk Class:

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
08P3824	Transferrin	59041	W0102010307

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, Certification Body, Ridlerstraße 65, 80339 Munich, Germany Notified Body Number 0123	
	Quality Management System	EU Certificate No.
Conformity Assessment Procedure	Annex IX Chapters I and III,  Including an assessment of the technical documentation for devices concerned on the basis of representative samples	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	Full Name:	Sandra Gallagher
Function:	Director Quality Assurance/ Site Quality Head	Function:	Manager Regulatory Affairs
Signature:	Belle	Signature:	S. Calleyler
Date of Approval:	15 DGC 2023	Date of Approval:	15-DEC-2023
Signed for, and on behalf of:	Abbott Ireland Diagnostics Division, Lisnamu	ick, Longford, Co. Lor	ngford Ireland
Date Issued:	15 DEC ZOZZ	Place Issued:	Lisnamuck, Longford, Co. Longford, Ireland
Supersedes:	30 June 2022	Effective (Date or Lot Number):	15 DEC 2023

SCHOOL SECTION	
grante S	
	ARROTT
Street, or other Designation of the last o	ADDUII

Certificate Identification:

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
04U0620	53462	Triglyceride2	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:

Signature:

Full Name (printed):

Siobhan Wright

Full Name (printed):

Thomas Breslin

Position:

Director Quality Assurance/

Position:

**Manager Regulatory Affairs** 

Site Quality Head

Date of Approval: 24-JUN-7021

25-JUNE -2021

Date of Approval:

Date Issued:

24- JUN- 2021

Place Issued:

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

Supersedes:

Not Applicable

Effective Date:

25-JUNE - 2021



**Basic UDI-DI:** 

038074ACP0775J9

**Basic UDI-DI Name:** 

Alinity c Ultra HDL

**Risk Class:** 

Class B

List Number and Size Code	Product and Trade Name	GMDN Code	EMDN Code
07P7520	Alinity c Ultra HDLReagent Kit	53391	W01010215
07P7530	Alinity c Ultra HDLReagent Kit	53391	W01010215

Manufacturer (Name and Address)	Abbott GmbH, Max-Planck-Ring 2, 65205 Wiesbaden, Germany		
Manufacturer SRN	DE-MF-000009455		
Authorized Representative (Name and Address)	N/A		
Authorized Representative SRN	N/A		
Produced by (Site of manufacture) (Name and Address)	Sekisui Diagnostics P.E.I. Inc. 70 Watts Avenue Charlottetown Prince Edward Island C1E 2B9 Canada		
Notified Body (Name and Identification Number)	TÜV SÜD Product Service GmbH Zertifizierstellen Ridlerstraße 65, 80339 München, 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 010051 02		
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:	Claudia Becker	Full Name:	Susanne Ulrich
Function:	Director Quality Assurance		Assoc. Director Regulatory Affairs
Signature:	C. Teclas	Signature:	moanne 116-la 12/08/2023
Date of Approval:	12 Oct 2023	_ Date of Approval:	12/00/12073
Signed for, and on behalf of:	Abbott GmbH, Wiesbaden, Germany		
Date Issued:	12 Oct 2023	_ Place Issued:	65205 Wiesbaden, Germany
Supersedes:	08-Jul-2022	Effective (Date or Lot Number):	12-0ct-2023



Certificate Identification:

DOC-08P1620, 08P1630-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
08P1620	53590	Alinity c Urea Nitrogen Reagent Kit	Self-declared
08P1630	53590	Alinity c Urea Nitrogen 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. Fiches	Signature:	ayfur Jenkins
Full Name:	Claudia Becker	Full Name:	Tiffini Jenkins
Position:	Director Quality Assurance	Position:	Manager Regulatory Affairs
Date of Approval:	22 Jul 2021	Date of Approval:	_11-JU1-2021
	U	Date Issued:	22-Jul- 2021
		Place Issued:	65205 Wiesbaden, Germany

Effective (Date or

05-Jan-2018

22-Jul- 2021

Supersedes:

Lot Number):

 4 80 Wh 40 000000
ABBOTT

Certificate Identification: 04U09

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
04U0920	53583	Uric Acid2	Self-declared
04U0930	53583	Uric Acid2	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): Position:	Siobhan Wright  Director Quality Assurance/ Site Quality Head	Signature: Full Name (printed): Position:	Lorraine Whitney Director Regulatory Affairs
Date of Approval:	[8- NOV-70	Date of Approval:	18 NOU 2020
Date Issued:	18-NOV-70	Place Issued:	Abbott Ireland Diagnostics Division, Lisnamuck, Longford, Co. Longford, Ireland.
Supersedes:	Not Applicable	Effective Date:	18-NOV-20



Certificate Identification:

DOC-07P5920, 07P5930-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
07P5920	53480	Alinity c Urine/CSF Protein Reagent Kit	Self-declared
07P5930	53480	Alinity c Urine/CSF Protein 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:

Position:

Claudia Becker

**Director Quality Assurance** 

C. Precus

Date of Approval:

Signature:

Full Name:

Tiffini Jenkins

11-1111-2021

Position:

**Manager Regulatory Affairs** 

Super Jentine

Date of Approval:

Date Issued:

Place Issued:

22- Jul- 2021 65205 Wiesbaden, Germany

Supersedes:

05-Feb-2019

Effective (Date or

Lot Number):

21- Jul- 2021