CERTIFICATE OF TRAINING

THIS CERTIFIES THAT

Sergiu Sorocovici

HAS SUCCESSFULLY COMPLETED THE TRAINING COURSE

ARCHITECT c8000 & RSH Service

March 6th – 14th, 2018

Ali Güntekin

TRAINER NAME

ABBOTT DIAGNOSTICS

DATE DD.MM.YYYY

TRAINER SIGNATURE

Germany - Delkenheim

14.03.2018

CERTIFICATE OF TRAINING

THIS CERTIFIES THAT

Alexei Legun

HAS SUCCESSFULLY COMPLETED THE TRAINING COURSE

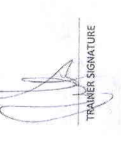
ARCHITECT c8000 Service & c8000 RSH

November 27th- December 5th, 2018

Vlassis Tsompanidis

TRAINER NAME

ABBOTT DIAGNOSTICS



05.12.2018

DATE DD.MM.YYYYY



Certificate Identification:

List Numbers and

Storage site of technical

documentation (name and address)

7D56

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038

Legal Manufacturer's Address:

GMDN

Abbott Park, Illinois 60064 USA

Devices	Code	Names and Description of Devices	Classification
7D56-21	52925	Alanine Aminotransferase	Self-declared
Authorized Euro	pean	Abbott GmbH & Co. KG Max-Planck-Ring 2	41-
Representative (name and address)		65205 Wincharden, Germany	

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:	Energy.	Signature:	mark Little
Full Name:	Erik Muegge	Full Name:	Mark Littlefield
Position:	QA Manager Ops	Position:	Assoc Director Dogulatory Ass.

Date of Approval: 8-SEP-2017 Date of Approval: 8-SEP-2017

Date Issued: 8-SEP-2017

Abbott Laboratories
1921 Hurd Drive

Place Issued: 1921 Hurd Drive Irving, TX 75038

Supersedes: _September 3, 2015____

Effective (Date or Lot Number): 8-SEP-2017



Certificate Identification:

7D81

Legal Manufacturer's Name:

Abbott Laboratories Diagnostic Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D81-21	52954	Aspartate Aminotransferase	Self-declared

Authorized European Representative (name and address)	Abbott GmbH & Co. KG Max-Planck-Ring 2 65205 Wiesbaden, Germany
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038
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:

Mark Littlefield

Full Name:

Thomas Creel

Director, Site QA

Position:

Assoc. Director Regulatory Affairs

Date of Approval:

15 13. L = 2015

Date of Approval:

15-0xT-2018

Date Issued:

15-CCT-2018

Abbott Laboratories

1921 Hurd Drive

Place Issued:

Irving, TX 75038

Supersedes:

08-SEP-2017

Effective (Date or

Lot Number):

15-0× T-2018



Certificate Identification:

DoC-7D55-SD DELK

Legal Manufacturer's Name:

Abbott GmbH & Co. KG

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
7D55-22	52929	Alkaline Phosphatase	Self-declared
7D55-32	52929	Alkaline Phosphatase	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical documentation (name and address)	Abbott Laboratories 1021 Hurd Drive Lating Town 75020 Mg.
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:

Olana Pomeco

Signature:

Full Name:

Diana Romero

Full Name:

Mark Littlefield

Position:

Director Quality Assurance

Position:

Assoc. Director Regulatory Affairs

Date of Approval:

22-MAY.2017

Date of Approval:

22 10177 2017

Date Issued:

22-MAY-2017

Place Issued:

65205 Wiesbaden, Germany

Supersedes:

Not applicable

Effective (Date or

Lot Number):

22-MAY-2017

□ ABBOTT

Declaration of Conformity

Certificate Identification: Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D53-23	53599	Albumin BCG	Self-declared
	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	ge site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038	
		Department - Regulatory Affairs	
Harm	onized 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

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: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: 9-3-2015

> Date Issued: 9-3-2015

Supersedes: November 5, 2014

Signature:

Full Name: Mark Littlefield

Associate Director, Regulatory Affairs Position:

9-3-2015 Date of Approval:

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

9-3-2015 Lot Number):

Certificate Identification: Legal Manufacturer's Name: 7D58

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D58-21	52941	Amylase	Self-declared
	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	ge site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038	
		Department - Regulatory Affairs	
Harm	onized 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

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: Diana Romero

Position: Site Director, Quality Assurance

9-3-2015 Date of Approval:

> 9-3-2015 Date Issued:

Supersedes: November 5, 2014

Signature:

Full Name: Mark Littlefield

Position: Associate Director, Regulatory Affairs

9-3-2015 Date of Approval:

Abbott Laboratories

Place Issued: 1921 Hurd Drive Irving, TX 75038

Effective (Date or

9-3-2015 Lot Number):



Certificate Identification:

7D81

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
6L45-21	53229	Total Bilirubin	Self-declared
6L45-41	53229	Total Bilirubin	Self-declared

Authorized European Representative (name and address)	Abbott GmbH & Co. KG Max-Planck-Ring 2 65205 Wiesbaden, Germany
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038
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:

Thomas Creel

Signature:

Mark Littlefield

Full Name:
Position:

Director, Site QA

Full Name:
Position:

Assoc. Director Regulatory Affairs

Date of Approval:

12-Oct - 2018

Date of Approval:

12-007-2018 12-007-2018

Date Issued:

101-10-1 De 10

Abbott Laboratories

Place Issued:

1921 Hurd Drive Irving, TX 75038

Supersedes:

September 8,2017

Effective (Date or

Lot Number):

12-007-2018



Certificate Identification:

8G63

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
8G63-21	53236	Direct Bilirubin	Self-declared

Authorized European	Abbott GmbH & Co. KG	
Representative (name and address)	Max-Planck-Ring 2	
(name and address)	65205 Wiesbaden, Germany	
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038	
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:

Ellen

Signature:

Mark Littlefield

Full Name:

Position:

Erik Muegge

QA Manager Ops

Full Name:
Position:

Assoc. Director Regulatory Affairs

Date of Approval-

8-550- 2017

Date of Approval:

8-5EP-2017

8-SEP-2017

Date Issued:

Abbott Laboratories

1921 Hurd Drive

Place Issued:

Irving, TX 75038

Supersedes:

_September 3, 2015

Effective (Date or

Lot Number):

8-5EP-2017

Certificate Identification:

1E66

Legal Manufacturer's Name:

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers nd Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
1E66-04	41830	Bilirubin Calibrator	Self-declared
	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
Storage site of technical documentation (Name and Address)		Abbott 1921 Hurd Drive Irving, TX 75038 Department - Regulatory Affairs	
Harm	олized 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

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:

Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: November 5, 2014

November 5, 2014

Date Issued:

Supersedes:

September 28, 2006

Signature:

Full Name: Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval: November 5, 2014

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

November 17, 2014 Lot Number):

Certificate Identification: Legal Manufacturer's Name: 3L79

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
3L79-21;3L79-31; 3L79-41	45789	Calcium	Self-declared
	orized European Representative me and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	e site of technical documentation me and Address)	Abbott 1921 Hurd Drive Irving, TX 75038	
Harmi	onized Standards	Department - Regulatory Affairs 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:

Dlana 50

New at Jones

Position: Site Director, Quality Assurance

Date of Approval:

November 5, 2014

Diana Romero

Date Issued: //-5-2014

Supersedes: December 31, 2012

Signature:

Full Name: Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval:

November 5, 2014

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

Lot Number): November 17, 2014



Certificate Identification:

7D62

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D62-21	53362	Cholesterol	Self-declared

Authorized European	Abbott GmbH & Co. KG	
Representative (name and address)	Max-Planck-Ring 2	
	65205 Wiesbaden, Germany	
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038	
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:

QA Manager Ops

Signature:

Mark Littlefield

Full Name: Position:

Erik Muegge

Full Name: Position:

Assoc. Director Regulatory Affairs

Date of Approval:

8-SEP-2017

8-SEP-2017

Date Issued:

Abbott Laboratories 1921 Hurd Drive

Place Issued:

Irving, TX 75038

Supersedes:

9-3-2015

Effective (Date or

Lot Number):

8-SEP-2017



EC DECLARATION OF CONFORMITY

For in vitro diagnostic medical devices (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 and immunochemistry, coagulation and rapid tests for immunology and serology" declares, under its own responsibility that the below listed devices comply with all 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. 322/2000).

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

- 1. comply with the applicable provisions of the Directive
- 2. are not included in the list A and B of Annex II of the Directive
- 3. are designed, manufactured and placed on the market according to the company certified quality system, in compliance with ISO 9001 and ISO 13485 as per indication expressed in Annex III of the Directive.

DICHIARAZIONE DI CONFORMITÀ CE

per dispositivi medico diagnostici in vitro IVD) - Direttiva 98/79/CE

La scrivente Sentinel CH. SpA con sede in Milano, Italia – Via Robert Koch 2, fabbricante dei dispositivi appartenente alla famiglia denominata "kit per chimica clinica, immunochimica, coagulazione, e test rapidi per immunologia e serologia" dichiara sotto la propria responsabilità che i dispositivi sotto elencati soddisfano 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 i dispositivi:

- 1. soddisfano le disposizioni applicabili della Direttiva
- 2. non sono inclusi nell'elenco A e B dell'Allegato III della Direttiva
- sono progettati, fabbricati ed immessi in commercio nell'ambito dell'applicazione di un sistema di qualità aziendale dichiarato conforme e certificato secondo le norme ISO 9001 e ISO 13485 come descritto dall'Allegato III della Direttiva.

Code/Codice	Product Description/Nome prodotto
6K26-30	CRP Vario
6K26-41	CRP Vario
6K26-10	CRP Calibrator Set
6K26-12	CRP Calibrator WR
6K26-14	CRP Calibrator HS
6K26-21	CRP Control HS
6K89-30	Ammonia Ultra
6K91-30	Ceruloplasmin
4P79-30	UIBC Liquid
8L24-31	Creatinine (Enzymatic)
8L24-41	Creatinine (Enzymatic)
8L25-30	Lithium
6K89-20	Ammonia Controls
6K30-10	Clin Chem Cal
6K31-10	Plasmaproteins Cal 3x
1P93-30	Cystatin C
1P93-10	Cystatin C Calibrator





Code/Codice	Product Description/Nome prodotto
1P93-20	Cystatin C Control Set
6K25-10	CK-MB Calibrator
6K25-20	CK-MB Control
6K30-20	Clin Chem Control 1
6K30-21	Clin Chem Control 2
6K32-20	Immuno Control 1
6K32-21	Immuno Control 2
6K32-22	Immuno Control Set
6K90-20	Bile Acids Controls
6K98-10	Fructosamine Control 1
6K98-20	Fructosamine Control 2
4P80-30	Lambda Light Chains
6K24-30	Cholinesterase
6K25-30	CK-MB
6K22-30	Pancreatic Amylase
6K96-30	Kappa Light Chains
6K23-30	HBDH
6K90-30	Bile Acids
6K92-30	Dibucaine CHE
6K93-30	Copper
6K94-30	Fructosamine
6K95-30	Iron
6K95-41	Iron

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 (10) 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.

Il fabbricante dichiara inoltre di:

- conservare e tenere a disposizione delle 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 almeno di dieci anni dalla data di produzione dell'ultimo lotto
- 2. avere istituito e di mantenere un'idonea procedura per garantire la sorveglianza postvendita richiesta dalla Direttiva.

Sentinel Ch. SpA

A Legal Representative
Un Legale Rappresentante
Dr. Filippo De Luca

Date/Data 소의/06/20년5

Certificate Identification: Legal Manufacturer's Name:

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
3L81-22; 3L81-32; 3L81-41	53251	Creatinine	Self-declared
	norized European Representative ime and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	Proceedings of the second
Storage site of technical documentation (Name and Address)		Abbott 1921 Hurd Drive Irving, TX 75038	
Harme	onized Standards	Department - Regulatory Affairs 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

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:

Diana Romero

Full Name: Position:

Mark Littlefield Associate Director, Regulatory Affairs

Position:

Site Director, Quality Assurance

November 5, 2014

Date of Approval:

November 5, 2014

Date of Approval:

Abbott Laboratories

Date Issued:

11-5-2014

Place Issued:

1921 Hurd Drive

Irving, TX 75038

Supersedes: July 16, 2013

Effective (Date or

Lot Number):

November 17, 2014



CE DECLARATION OF CONFORMITY

DRC-726

Edition 3

P-172

Page 1 of 3

CE

DECLARATION OF CONFORMITY

Manufacturer:

Hersteller Fabricante Fabricant Produttore Fabricante Producent Tillverkare Καταοκευαστής BIOKIT, S.A. Can Malé s/n. 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.

Bìokit 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.

Biokít 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 específicadas 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 nedan uppräknade produkt(er) är förenlig(a) med de EU-direktiv och standarder som identifieras i denna deklaration

Η Biokil 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

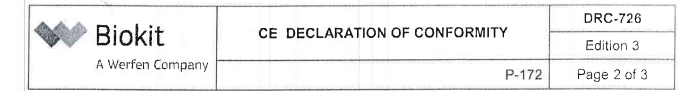
IVD - 98/79/EC (27/10/1998)

Standard(s):

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

ISO 9001

ISO 13485



Notified Body:
Benannte Stelle Organismo Notificado Organisme Notifié Organismo Notificato Organismo Notificado Teknisk Kontrollorgon Anmält Organ Κοινοποιημένος Οργανισμός

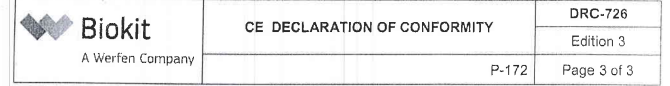
Name: Other Devices Code:N/A

Certificate Nº: N/A

Annex III

Product(s):
Produkt(e) Producto(s) Produit(s) Produit(s) Produto(s) Produkt(er) Produkt(er) (Προϊόν(-τα)

Product(s) Produkt(e) Producto(s) Produit(s Prodolto(i)	Produto(s) Produkt(er) Produkt(er) Проїо́ν(-τα)	
P/N	复源、斯勒斯科· 通信和限制。	
6L34-42	Quantia A-1-AGP	
6K38-01	Quantia ASO	
6K39-01	Quantia β2-Microglobulin	
6K40-01	Quantia Digitoxin	
6K41-01	Quantia Ferritin	
6K42-01	Quantia IgE	
6L32-42	Quantia Myoglobin	
6K44-01	Quantia RF	
6K99-01	Quantia A1-Antitrypsin	
7K02-01	Quantia D-Dimer	
7K00-01	Quantia Lp (a)	
6K45-01	Quantia PROTEINS Standard	
6K46-01	Quantia ASO Standard	
6K47-01	Quantia β2-Microglobulin Standard	
6K48-01	Quantia Digitoxin Standard	
6K49-01	Quantia Ferritin Standard	
6K50-01	Quantia IgE Standard	
6L33-04	Quantia Myoglobin Standard	
6K52-01	Quantia RF Standard	
7K02-10	Quantia D-Dimer Standard	
7K00-10	Quantia Lp (a) Standard	
5P83-01	Lp (a) Calibrators	
6K53-01	Quantia PROTEINS Control	
6K54-01	Quantia ASO-RF Control I	
6K55-01	Quantia ASO-RF Control II	



Product(s) Produkt(e) Producto(s) Produit(s Prodotto(i)	Produto(s) Produkt(er) Produkt(er) Προϊόν(-τα)
P/N	STANDARD SERVICE SERVI
6K56-01	Quantia Ferritin/Myoglobin/lgE Control
6K57-01	Quantia Digitoxin Control
7K02-20	Quantia D-Dimer Control
7K00-20	Quantia Lp (a) Control
5P84-10	Lp (a) Control

Signature

20/3/2015



Certificate Identification:

3L82

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
3L82-21, 3L82-41	53301	Glucose	Self-declared

Authorized European	Abbott GmbH & Co. KG	
Representative (name and address)	Max-Planck-Ring 2	
	65205 Wiesbaden, Germany	
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038	
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:

frage of the same of the same

Signature:

Mark Littlefield

Full Name:
Position:

Erik Muegge

Full Name: Position:

Assoc. Director Regulatory Affairs

QA Manager Ops

Date of Approval:

8-5EP-2017

Date of Approval:

0-36-6011

Date Issued:

8-SEP-2017

Abbott Laboratories

1921 Hurd Drive

Place Issued:

Irving, TX 75038

Supersedes:

_November 17, 2014

Effective (Date or

Lot Number):

8-SEP-2017

Certificate Identification: Legal Manufacturer's Name: 7D65

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D65-21 7D65-41	53030	Gamma-Glutamyl Transferase	Self-declared

Authorized European	Abbott
Representative	Max-Planck-Ring 2
(Name and Address)	65205 Wiesbaden, Germany
Storage site of technical	Abbott
documentation	1921 Hurd Drive
(Name and Address)	Irving, TX 75038
	Department - Regulatory Affairs
Harmonized Standards	Listed in the Technical Documentation

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

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

Signature:

Jana Romero

Full Name: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: 9-3-2015

> Date Issued: 9-3-2015

Supersedes: November 5, 2014 Signature:

Full Name: Mark Littlefield

Associate Director, Regulatory Affairs Position:

Date of Approval: 9-3-2015

Abbott Laboratories 1921 Hurd Drive Place Issued:

Irving, TX 75038

Effective (Date or

9-3-2015 Lot Number):

Certificate Identification: Legal Manufacturer's Name:

3K33

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
3K33-21	30169	Ultra HDL	Self-declared
	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	ge site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038	
		Department - Regulatory Affairs	0
Harm	onized 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: Dana Bom

Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: November 5, 2014

Full Name:

Date Issued: November 5, 2014

Supersedes: April 4, 2013

Signature:

Full Name: Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval: November 5, 2014

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving TV 75020

Irving, TX 75038

Effective (Date or

Lot Number): November 17, 2014



DECLARATION OF CONFORMITY

Manufacturer:

Sekisui Diagnostics P.E.I. Inc

70 Watts Avenue Charlottetown

Prince Edward Island

C1E 2B9 Canada

European Representative:

Sekisui Diagnostics (UK) Ltd

Liphook Way Allington Maidstone

Kent ME16 0LQ

Product:

Direct LDL

Catalogue Number: 1E31-20; 1E31-02

GMDN Code:

53395; 41728

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:

Allington, UK

Signature:

David Torrens

20-NOV-2018

Date

Senior Manager Regulatory Affairs Sekisui Diagnostics (UK) Ltd

Sekisul Diagnostics (UK) Ltd Liphook Way Allington, Kent, ME16 0LQ Tel: 01622 607800 Fax: 01622 607801 Info@sekisui-dx.com www.sekisuidlagnostics.com

Certificate Identification:

5P56

Legal Manufacturer's Name:

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers nd Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
5P56-01	53356	Lipid Multiconstituent Calibrator	Self-declared

Authorized European	Abbott
Representative	Max-Planck-Ring 2
(Name and Address)	65205 Wiesbaden, Germany
Storage site of technical	Abbott
documentation	1921 Hurd Drive
(Name and Address)	Irving, TX 75038
	Department - Regulatory Affairs
Harmonized Standards	Listed in the Technical Documentation

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

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

Signature:

Full Name: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: November 5, 2014

Date Issued: //-5-7014

Supersedes: January 30, 2014

Signature:

Full Name: Marl

Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval: November 5, 2014

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

Lot Number):

November 17, 2014



Certificate Identification:

DoC-4P5220, 4P5201, 4P5211-SD DELK

Legal Manufacturer's Name:

Abbott GmbH & Co. KG

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
4P52-20	59090	Hemoglobin A1c Reagent	Self-declared
4P52-01	53315	Hemoglobin A1c Calibrator	Self-declared
4P52-11	44435	Hemoglobin Alc Controls	Self-declared

Authorized European Representative (name and address)	N/A
Storage site of technical	
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:

Diana Romero

Signature:

Full Name:

Mark Littlefield

Full Name:
Position:

Director, Site QA

Position:

Assoc. Director, Regulatory Affairs

Date of Approval:

17-NOV-2017

Date of Approval:

17-200-2017

Date Issued:

17-100-2017

Place Issued:

65205 Wiesbaden, Germany

Supersedes:

N/A

Effective (Date or

Lot Number):

17-Nov-2017



Certificate Identification:

7D80

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D80-31	53114	Lipase	Self-declared

Authorized European	Abbott GmbH & Co. KG
Representative (name and address)	Max-Planck-Ring 2
	65205 Wiesbaden, Germany
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038
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:

Erik Muegge

Signature:

Mark Littlefield

Full Name: Position:

QA Manager Ops

Full Name: Position:

Assoc. Director Regulatory Affairs

Date of Approval:

Date of Approval:

8-SEP-2017

8-SEV-2017

Date Issued:

Abbott Laboratories

1921 Hurd Drive

Place Issued: Irving, TX 75038

Supersedes:

_November 17, 2014

Effective (Date or

Lot Number):

8-SEP-2017

□ ABBOTT

Declaration of Conformity

Certificate Identification: Legal Manufacturer's Name: 3E16

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

nd Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
3E16-02	53109	Lipase Calibrator	Self-declared
	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	ge site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038 Department - Regulatory Affairs	
Harm	onized 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:

Diana Rom

Chara Homera

Full Name: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: 9-3-2015

Date Issued: 9-3-2015

Supersedes: November 5, 2014

Signature:

Full Name: Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval: 9-3-2015

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

Lot Number): 9-3-2015

□ ABBOTT

Declaration of Conformity

Certificate Identification: Legal Manufacturer's Name: 7D73

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D73-21	53989	Total Protein	Self-declared
	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	ge site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038 Department - Regulatory Affairs	
Harm	onized 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:

ignature: (C)(U) (C) 7)(U)

Full Name: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: 9-3-2015

Date Issued: 9-3-2015

Supersedes: November 5, 2014

Signature:

Full Name: Mark

Mark Littlefield

Position:

Associate Director, Regulatory Affairs

Date of Approval:

9-3-2015

Abbott Laboratories

1921 Hurd Drive

Irving, TX 75038

or ,

Effective (Date or Lot Number):

Place Issued:

9-3-2015



Certificate Identification:

7D74

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Legal Manufacturer's Address:

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
7D74-21	53462	Triglyceride	Self-declared

Authorized European Representative (name and address)	Abbott GmbH & Co. KG	
	Max-Planck-Ring 2	
	65205 Wiesbaden, Germany	
Storage site of technical documentation (name and address)	Abbott Laboratories, 1921 Hurd Drive, Irving, Texas 75038	
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:

Erik Muegge

Signature:

Mark Littlefield

Full Name: Position:

QA Manager Ops

Full Name: Position:

Assoc. Director Regulatory Affairs

Date of Approval:

Date of Approval:

8-SEP-2017

8-SEP-2017

Date Issued:

Abbott Laboratories

1921 Hurd Drive

Place Issued:

Irving, TX 75038

Supersedes:

9-3-2015

Effective (Date or

Lot Number):

8-SEP-2017

List Numbers

Declaration of Conformity

Certificate Identification: Legal Manufacturer's Name:

GMDN Code

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
3P39-21; 3P39-41	53583	Uric Acid	Self-declared
(Na Storag	norized European Representative me and Address) e site of technical documentation me and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany Abbott 1921 Hurd Drive Irving, TX 75038	
Harmo	onized Standards	Department - Regulatory Affairs 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

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:

Diana Romero

Position: Site Director, Quality Assurance

Date of Approval:

November 5, 2014

Date Issued:

11-5-2014

Supersedes:

December 31, 2012

Signature:

Full Name:

Mark Littlefield

Position:

Associate Director, Regulatory Affairs

Date of Approval:

Lot Number):

November 5, 2014

Place Issued:

Abbott Laboratories 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

November 17, 2014

List Numbers

Declaration of Conformity

Names and Description of Devices

Certificate Identification: Legal Manufacturer's Name:

GMDN Code

1E65

Abbott Laboratories

Diagnostics Division

Abbott Park, Illinois 60064 USA

and Size Code of Devices		Names and Description of Devices	Classification
1E65-04	30216	Multiconstituent Calibrator	Self-declared
1E65-05	30216	Multiconstituent Calibrator	Self-declared
(Na	norized European Representative nme and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	e site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038	

Department - Regulatory Affairs Listed in the Technical Documentation Harmonized Standards

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

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

Signature:

Supersedes:

Full Name: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: November 5, 2014

> November 5, 2014 Date Issued:

March 6, 2014

Date of Approval:

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

November 5, 2014

Mark Littlefield

Associate Director, Regulatory Affairs

Classification

Effective (Date or

Signature:

Full Name:

Position:

November 17, 2014 Lot Number):



DECLARATION OF CONFORMITY



Manufacturer

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

Product(s):

Product Name	Catalogue Numbe
Multichem S Plus (Unassayed)	05P79-10
Multichem S Plus (Unassayed)	05P79-11
Multichem S Plus (Unassayed)	05P79-12
Multichem S Plus	CH100CRP
Multichem S Plus	CH101CRP
Multichem S Plus	CH102CRP
Multichem S Plus	CH103CRP
Multichem S Plus (Assayed)	05P78-10
Multichem S Plus (Assayed)	05P78-11
Multichem S Plus (Assayed)	05P78-12
Multichem S Plus (Unassayed)	CH110CRP.05
Multichem S Plus (Unassayed)	CH111CRP.05
Multichem S Plus (Unassayed)	CH112CRP.05
Multichem S Plus (Unassayed)	CH113CRP.05
Multichem S Plus	CH100PLA
Multichem S Plus	CH101PLA
Multichem S Plus	CH102PLA
Multichem S Plus	CH103PLA
Multichem S Plus (Assayed)	CH110PLA.05
Multichem S Plus (Assayed)	CH111PLA.05
Multichem S Plus (Assayed)	CH112PLA.05
Multichem S Plus (Assayed)	CH113PLA.05



GMDN:

47869

Conformity Route:

Annex III Self-Declared

Quality Management System:

EN ISO 13485:2012/ ISO 13485:2003

QMS Certification No.:

LRQ 4008261/A

Issued By:

Lloyds Register LRQA, 71 Fenchurch Street,

London EC3M 4BS United Kingdom.

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

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

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

Bernd Hass , Head of Quality and Regulatory Affairs

24-Jan-2014.

Techno-path Manufacturing Ltd.

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

Standard	Title	
EN ISO 15223-1:2012	Symbols for use in the labelling of medical devices	
ISO 13485:2012 + AC:2012	Medical devices – Quality management systems – Requirements for regulatory purposes	
EN 13612:2002 + AC:2002	Performance evaluation of in vitro diagnostic medical devices	
EN 13641:2002	Elimination or reduction of risk of infection related to in vitro diagnostic reagents	
EN 13975:2003	Sampling procedures used for acceptance testing of in in vitro diagnostic medical devices – statistical aspects	
EN ISO 14971:2012	Medical devices – Application of risk management to medical devices	
EN ISO 18113-1:2011	In vitro diagnostic medical devices – Information suppli by the manufacturer (labelling) – Part 1: Terms, definiti and general requirements	
EN ISO 18113-2:2011	In vitro diagnostic medical devices – Information supplied by the manufacturer (labelling) – Part 2: In vitro diagnostic reagents for professional use	
EN 13640:2002	Stability Testing of In vitro diagnostic reagents	

List Numbers

Declaration of Conformity

Certificate Identification:

Legal Manufacturer's Name: Abbott Laboratories

GMDN Code

Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers nd Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
6K01-20	56676	Acid Wash	Self-declared
(N:	horized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
Storage site of technical documentation (Name and Address)		Abbott 1921 Hurd Drive Irving, TX 75038 Department - Regulatory Affairs	
Harmonized Standards		Listed in the Technical Documentation	

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

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: Diana Romero

Position: Site Director, Quality Assurance

Date of Approval: November 5, 2014

> 11-5-2014 Date Issued:

Supersedes: December 11, 2006

Full Name: Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval: November 5, 2014

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

Lot Number):

November 17, 2014

Certificate Identification:

9D31

Legal Manufacturer's Name:

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
9D31-20	58236	Alkaline Wash	Sclf-declared
(N	horized European Representative ame and Address) ge site of technical	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany Abbott	
(N	documentation ame and Address)	1921 Hurd Drive Irving, TX 75038 Department - Regulatory Affairs	
Harmonized Standards		Listed in the Technical Documentation	

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

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: Diana Romero

Position: Site Director, Quality Assurance

5-28-2015 Date of Approval:

> Date Issued: 5-28-2015

Supersedes: March 28, 2013

Signature:

Full Name: Mark Littlefield

Associate Director, Regulatory Affairs Position:

Date of Approval: 5-28-2015

Abbott Laboratories

1921 Hurd Drive Place Issued: Irving, TX 75038

Effective (Date or Lot Number):

5-28-2015

Certificate Identification: Legal Manufacturer's Name: 1J72

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
1J72-20	59058	Detergent A	Self-declared
(Na Storag	horized European Representative ame and Address) ge site of technical documentation ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany Abbott 1921 Hurd Drive Irving, TX 75038 Department - Regulatory Affairs	
Harm	onized 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

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: Diana Romero

Site Director, Quality Assurance Position:

5-28-2015 Date of Approval:

> 5-28-2015 Date Issued:

Supersedes: March 28, 2013

Full Name: Mark Littlefield

Associate Director, Regulatory Affairs Position:

Date of Approval: 5-28-2015

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

5-28-2015 Lot Number):

Certificate Identification:

Legal Manufacturer's Name: **Abbott Laboratories**

Diagnostics Division

Abbott Park, Illinois 60064 USA

and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
2J94-21	59058	Detergent B	Self-declared
	norized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
	e site of technical documentation ame and Address)	Abbott 1921 Hurd Drive Irving, TX 75038	
		Department - Regulatory Affairs	

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

Listed in the Technical Documentation

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: Diana Romero

Position:

Site Director, Quality Assurance

Harmonized Standards

Date of Approval: December 4, 2014

> December 4, 2014 Date Issued:

Supersedes: New

Signature:

Full Name:

Mark Littlefield

Position: Associate Director, Regulatory Affairs

Date of Approval:

December 4, 2014

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

Lot Number):

December 4, 2014

Certificate Identification: Legal Manufacturer's Name: 9D29

Abbott Laboratories Diagnostics Division

Abbott Park, Illinois 60064 USA

and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
9D29-20	56676	Water Bath Additive	Self-declared
9D29-21	56676	Water Bath Additive	Self-declared
	norized European Representative ame and Address)	Abbott Max-Planck-Ring 2 65205 Wiesbaden, Germany	
Storage site of technical documentation (Name and Address)		Abbott 1921 Hurd Drive Irving, TX 75038	

Department - Regulatory Affairs

Listed in the Technical Documentation Harmonized Standards

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

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:

Diana Romero

Position: Site Director, Quality Assurance

Date of Approval:

(0-11-2015 Date Issued:

6-11-2015

Supersedes: March 28,2013

Signature:

Full Name: Mark Littlefield

Associate Director, Regulatory Affairs Position:

6-11-2015 Date of Approval:

Abbott Laboratories

Place Issued: 1921 Hurd Drive

Irving, TX 75038

Effective (Date or

Lot Number):

6-11-2015



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

ARCH Sys Acc LC	IRIS V3
Abbott Laboratories	
Diagnostics Division	
Abbott Park, IL 60064 USA	

List Numbers and Size Code of Devices	GMDN Code	Names and Description of Devices	Classification
4D18-03	NA	ARCHITECT Septum	Self-declared
4D19-01	NA	ARCHITECT Replacement Caps	Self-declared
7C14-01	NA	ARCHITECT Sample Cups	Self-declared
7C15-02	NA	ARCHITECT Reaction Vessels	Self-declared
7C15-03	NA	ARCHITECT Reaction Vessels	Self-declared

Authorized European Representative (name and address)	Abbott GmbH & Co. KG Max-Planck-Ring 2 65205 Wiesbaden, Germany
Storage site of technical documentation (name and address)	Abbott Laboratories Diágnostics División Abbott Park, 1L 60064 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

Deborah Hinkley

Full Name:

Lauren Sieber `

Full Name:

.

Position:

Product Quality Assurance

Position:

Regulatory Affairs

Manager

Director

Date of Approval

5 28 2015

Date of Approval

Abbott Laboratories

Dare Issued:

ni las lane

Place Issued:

Diagnostics Division Abbott Park, IL 60064 USA

Supersedes:

June 13, 2013

Effective (Date or Lot Number):

24/02/235