



Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.
---	--

Produktname (deutsch): Thromborel S	Product name (English): Thromborel S
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Produkt-Nr. / Product No. (REF):	OUHP
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Packungsgröße(n) / Package Size(s) (REF):	OUHP 29, OUHP 49
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IVD-Kategorie / IVD Category:	Others
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Hersteller / Manufacturer:	Siemens Healthcare Diagnostics Products GmbH
----------------------------	--

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
---	--

Bestätigung / Authorization: Director Quality/Regulatory
Unterschrift / Signature
Dr. Jörg Amborn
Name /Name
2008-09-03
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.
---	--

Produktname (deutsch): Dade Actin FS Reagenz zur Bestimmung der APTT	Product name (English): Dade Actin FS Activated PTT Reagent
---	--

Produkt-Nr. / Product No. (REF): B4218-20, -100
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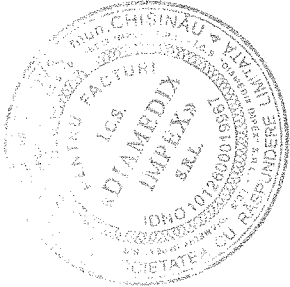
Packungsgröße(n) / Package Size(s) (REF): B4218-20, -100

IVD-Kategorie / IVD Category: Sonstige	Others
---	--------

Hersteller / Manufacturer: Siemens Healthcare Diagnostics Products GmbH
--

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
---	--

Bestätigung / Authorization: Director Quality/Regulatory	
J. AL	
Unterschrift / Signature	
Dr. Jörg Amborn	
Name /Name	
2008-09-03	
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:	





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.
---	--

Produktname (deutsch): Calciumchlorid-Lösung	Product name (English): Calcium Chloride Solution
---	--

Produkt-Nr. / Product No. (REF): ORHO
--

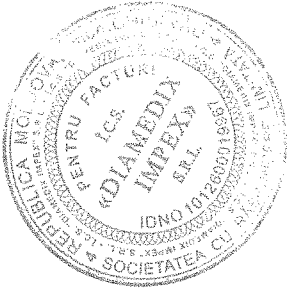
Packungsgröße(n) / Package Size(s) (REF): ORHO 37
--

IVD-Kategorie / IVD Category: Sonstige	Others
---	--------

Hersteller / Manufacturer: Siemens Healthcare Diagnostics Products GmbH
--

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
---	--

Bestätigung / Authorization: Director Quality/Regulatory
Unterschrift / Signature
Dr. Jörg Amborn
Name /Name
2009-11-05
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.
---	--

Produktname (deutsch): Dade Thrombin Reagenz	Product name (English): Dade Thrombin Reagent
---	--

Produkt-Nr. / Product No. (REF): B4233-25, -27

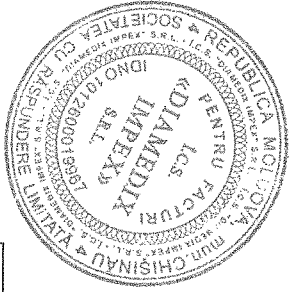
Packungsgröße(n) / Package Size(s) (REF): B4233-25, -27
--

IVD-Kategorie / IVD Category: Sonstige	Others
---	--------

Hersteller / Manufacturer: Siemens Healthcare Diagnostics Products GmbH
--

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
---	--

Bestätigung / Authorization:	
Director Quality/Regulatory	
J. A-L	
Unterschrift / Signature	
Dr. Jörg Amborn	
Name / Name	
2008-09-03	
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:	





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-Vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.

We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.

Produktname (deutsch):

Product name (English):

Dade Owren's Veronal-Puffer

Dade Owren's Veronal Buffer

Produkt-Nr. / Product No. (REF):

B4234-25

Packungsgröße(n) / Package Size(s) (REF):

B4234-25

IVD-Kategorie / IVD Category:

Sonstige

Others	
--------	--

Hersteller / Manufacturer:

Siemens Healthcare Diagnostics Products GmbH

Adresse (innerhalb Deutschland):

Siemens Healthcare Diagnostics Products GmbH
Emil-von-Behring-Str. 76
35041 Marburg

Address (international):

Siemens Healthcare Diagnostics Products GmbH
Emil-von-Behring-Str. 76
35041 Marburg
Germany

Bestätigung / Authorization:

Director Quality/Regulatory

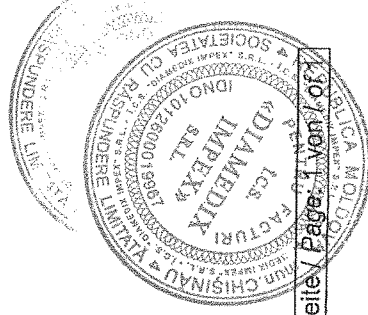
Unterschrift / Signature

Dr. Jörg Amborn

Name /Name

2008-09-03

Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	<i>We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance with Annex III assessment procedures of Annex III.</i>
---	--

Produktname (deutsch):	Produktname (English):
PT-Multi Calibrator	PT-Multi Calibrator

Produkt-Nr. / Product No. (REF):

OPAT

Packungsgröße(n) / Package Size(s) (REF):	OPAT 03

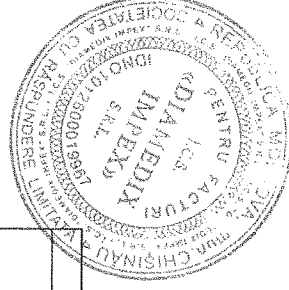
IVD-Kategorie / IVD Category:	
Sonstige	Others

Hersteller / Manufacturer: Siemens Healthcare Diagnostics Products GmbH

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
--	---

Bestätigung / Authorization:
Director Quality/Regulatory

Unterschrift / Signature
Dr. Jörg Amborn
Name /Name
2008-09-03
Datum [JJJJ-MM-IT] / Date [YYYY-MM-DD]:



Konformitätserklärung

Declaration of Conformity



Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.
---	--

Produktname (deutsch): Standard-Human-Plasma	Product name (English): Standard Human Plasma
---	--

Produkt-Nr. / Product No. (REF):	ORKL
----------------------------------	------

Packungsgröße(n) / Package Size(s) (REF):	ORKL 13, ORKL 17, ORKL 21
---	---------------------------

IVD-Kategorie / IVD Category:	
Sonstige	Others

Hersteller / Manufacturer:	Siemens Healthcare Diagnostics Products GmbH
----------------------------	--

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
---	--

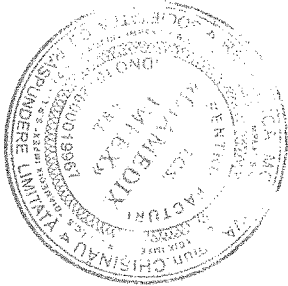
Bestätigung / Authorization:	
Director Quality/Regulatory	
J. A. C.	
Unterschrift / Signature	
Dr. Jörg Amborn	
Name /Name	
2011-04-05	
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:	





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.		We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.	
Produktname (deutsch): Kontroll-Plasma N		Product name (English): Control Plasma N	
Produkt-Nr. / Product No. (REF):		ORKE	
Packungsgröße(n) / Package Size(s) (REF):		ORKE 41	
IVD-Kategorie / IVD Category: Sonstige		Others	
Hersteller / Manufacturer:		Siemens Healthcare Diagnostics Products GmbH	
Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg		Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany	

Bestätigung / Authorization:	
Director Quality/Regulatory	
[Signature]	
Unterschrift / Signature	
Dr. Jörg Amborn	
Name /Name	
2008-09-03	
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:	





Wir erklären hiermit, dass die unten angegebenen In-vitro-Diagnostika-Produkte mit den Grundlegenden Anforderungen der Richtlinie 98/79/EG des Europäischen Parlaments und des Rates über In-vitro-Diagnostika übereinstimmen und die Anforderungen gemäß Annex III erfüllt werden.	We hereby declare that the in vitro diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on in vitro Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.
---	--

Produktname (deutsch): Kontroll-Plasma P	Product name (English): Control Plasma P
---	---

Produkt-Nr. / Product No. (REF):	OUPZ
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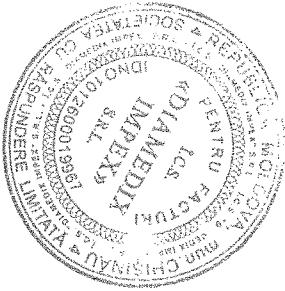
Packungsgröße(n) / Package Size(s) (REF):	OUPZ 17
---	---------

IVD-Kategorie / IVD Category:	Others
-------------------------------	--------

Hersteller / Manufacturer:	Siemens Healthcare Diagnostics Products GmbH
----------------------------	--

Adresse (innerhalb Deutschland): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg	Address (international): Siemens Healthcare Diagnostics Products GmbH Emil-von-Behring-Str. 76 35041 Marburg Germany
---	--

Bestätigung / Authorization:	
Director Quality/Regulatory	
Unterschrift / Signature	
Dr. Jörg Amborn	
Name /Name	
2008-09-03	
Datum [JJJJ-MM-TT] / Date [YYYY-MM-DD]:	



Konformitätserklärung



We hereby declare that the *in vitro* diagnostic devices described below conforms to all applicable Essential Requirements of Directive 98/79/EC on *in vitro* Diagnostic Medical Devices and accordance was shown by conformity assessment procedures of Annex III.

Product name (English):

Dade Ci-Trol 2

291071

291071

Others

Siemens Healthcare Diagnostics Products GmbH

Address (international):

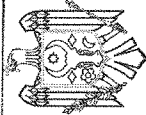
Healthcare Diagnostics Products GmbH
Emill-von-Behring-Str. 76
35041 Marburg
Germany

Director Quality/Regulatory

Unterschrift / Signature

Name / Name

Datum [JJJ-MM-TT] / Date [YYYY-MM-DD]:



Digitally signed by Marinescu Traian Alin
Date: 2019.11.04 08:57:48 EET
Reason: MoldSign Signature
Location: Moldova

Seite / Page: 1 von / of 1

EC Declaration of Conformity

Application of Council Directive:

98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

Means of conformity:

The following product is in conformity with Directive **98/79/EC** based on the test results using harmonised standards in accordance with Article 5 of the Directive.

Product identification:

Product name: CA CLEAN I

Manufacturer:

Name: SYSMEX CORPORATION

Address: 1-5-1 Wakoinohama-Kaigandori, Chuo-ku, Kobe 651-0073

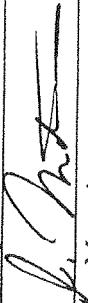
Country: Japan

Authorised representative:

Name: SYSMEX EUROPE GMBH

Address: Bornbarch 1, 22848 Norderstedt

Country: Germany

Authorised officer: 

Iwafne Matsui

Position: President

Date: 9TH JANUARY 2002

Place: NORDERSTEDT, GERMANY

This certificate was issued under sole responsibility of:

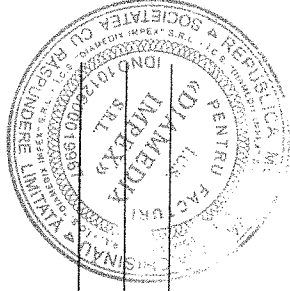
Authorised officer: 

Tokuhiko Okada

Position: Vice President, Technology Control

Date: November 7, 2001

Place: Japan



EC Declaration of Conformity

Application of Council Directive:

98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

Means of conformity:

The following product is in conformity with Directive 98/79/EC based on the test results using harmonised standards in accordance with Article 5 of the Directive.

Product identification:

Product name: CA CLEAN II

Manufacturer:

Name: SYSIMEX CORPORATION

Address: 1-5-1 Wakinohama-Kaigandori, Chuo-ku, Kobe 651-0073

Country: Japan

Authorised representative:

Name: SYSIMEX EUROPE GMBH

Address: Bornbarch 1, 22848 Norderstedt

Country: Germany

Authorised officer:


 Iwane Matsui

Position: President

Date: 9TH JANUARY 2002

Place: NORDERSTEDT, GERMANY

This certificate was issued under sole responsibility of:

Authorised officer:

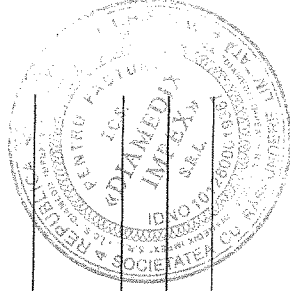


Tokuhiko Okada

Position: Vice President, Technology Control

Date: November 7, 2001

Place: Japan



EC Declaration of Conformity

Application of Council Directive:

98/79/EC of 27 October 1998 on In Vitro Diagnostic Medical Devices

Means of conformity:

The following product is in conformity with Directive **98/79/EC** based on the test results using harmonised standards in accordance with Article 5 of the Directive.

Product identification:

Product: REACTION TUBE

Model: SU-40

Manufacturer:

Name: SYSMEX CORPORATION

Address: 1-5-1 Wakoinohama-Kaigandori, Chuo-ku, Kobe 651-0073

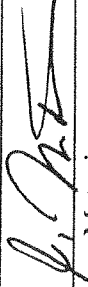
Country: Japan

Authorised representative:

Name: SYSMEX EUROPE GMBH

Address: Bornbarch 1, 22848 Norderstedt

Country: Germany

Authorised officer: 

Iwane Matsui

Position: President

Date: 10TH JANUARY 2002

Place: NORDERSTEDT, GERMANY

This certificate was issued under sole responsibility of:

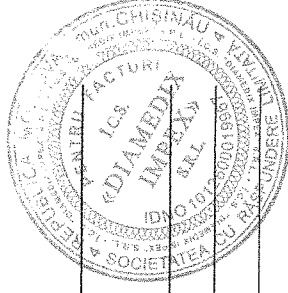
Authorised officer: 

Tokuhiro Okada

Position: Vice President, Technology Control

Date: November 16, 2001

Place: Japan



Siemens Healthcare Diagnostics Products GmbH, Marburg, Germany

Name Department	Tobias Thäns SHS LD LC ESB
Telephone Fax	+49 (6196) 7713-2414 +49 (6196) 7713-7007
E-Mail	tobias.thaens@siemens-healthineers.com
Our reference Date	19-0117 January 15, 2020

To whom it may concern
For presentation in Romania and Moldova

Letter of Authorization

Siemens Healthcare Diagnostics Products GmbH, Emil-von-Behring-Strasse 76, 35041 Marburg, Federal Republic of Germany (hereinafter referred to as the "Company") hereby confirms that pursuant to the Amended and Restated Global Supply, Distributorship, Sale and Service Agreement dated March 23, 2009 (the "Sysmex Agreement") between Siemens Healthineers and Sysmex Corporation, Japan ("Sysmex"), the Company is authorized to distribute, sell and service Sysmex hemostasis products ("Products") in the territory of Moldova ("Territory"), including the right to appoint sub-distributors.

In accordance with the rights conferred under the Sysmex Agreement,

ICS Diamedix Impex SRL

str. 31 August 1989, 108/2.
MD2004 Chisinau
Municipiul Chisinau
Moldova
("ICS Diamedix"),

has been appointed as a distributor of the Products in the Territory including the right, acting in its own name and on its own behalf,

1. to distribute and perform technical maintenance and service of the Products and
2. to participate in public tenders and enter into contracts relating to Products for the purpose of fulfilling its obligations under such tenders.

This letter of authorization shall terminate December 31, 2020 unless revoked sooner by the Company, save that in the event of a successful bid by ICS Diamedix during the intervening period, this letter shall remain valid until the date on which the contract entered into by ICS Diamedix as a result of such bid ends.

On behalf of Siemens Healthcare Diagnostics Products GmbH

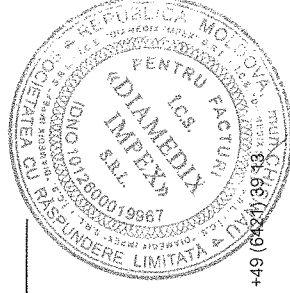

Electronically signed by: Claus
Prümper
Reason: I have reviewed this document
Date: 2020-01-16 11:22:16+01:00

Name: Dr. Claus Prümper
Title: Authorized Representative



Electronically signed by: Jürgen Vogt
Reason: I have reviewed this document
Date: 2020-01-15 15:56:49+01:00

Name: Jürgen Vogt
Title: Authorized Representative



Siemens Healthcare Diagnostics Products GmbH
Management: Michael Heindl, Joerg Berner, Tobias Thaens

Emil-von-Behring-Str. 76
35041 Marburg
Germany

Tel.: +49 (6421) 39-43

Chairman of the Supervisory Board: Deepak Nath; Registered Office: Marburg an der Lahn; Commercial registry: Marburg an der Lahn HRB 2054

Siemens Healthcare Diagnostics Products GmbH, Marburg, Germany

To whom it may concern
For presentation in
the Republic of Moldova

Name Department	Tobias Thäns SHS LC LD ESB
Telephone Fax	+49 (6196) 7713-2414 +49 (6196) 7713-7007
E-Mail	tobias.thaens@siemens-healthineers.com
Our reference Date	19-0116 January 15, 2020

MANUFACTURER'S AUTHORIZATION

SIEMENS HEALTHCARE DIAGNOSTICS PRODUCTS GMBH a company incorporated and existing under the laws of the Federal Republic of Germany, with offices located at Emil-von-Behring-Strasse 76, 35041 Marburg, Germany (the "Company"), hereby acknowledges that it has designated **ICS Diamedix Impex SRL**, a company incorporated in the Republic of Moldova under the fiscal code (IDNO) 1012600019967 ("Representative") as an **AUTHORIZED REPRESENTATIVE** for the Republic of Moldova ("Territory"). As such Representative is duly authorized to register and re-register the products of the Siemens Healthineers in-vitro diagnostic portfolio (the "Products") listed in Attachment A within the Territory. The manufacturing facilities of the Company and its affiliates are listed in Attachment B.

In accordance with an Authorized Representative Agreement, Siemens Healthineers will, among other things, provide to the Representative such product-related technical documentation relevant to market surveillance investigations being undertaken by the Medicines and Medical Devices Agency in the Territory as may reasonably be required. In connection with the requirements of Annex 3 to the Administrative Procedures for Registration of medical devices nr. A07PS-01Rg04-112 dated 03.07.2014 ("Procedure Document"), however, we point out that the following information relating to the Products, to the extent not disclosed in documentation accompanying the Products (such as Instructions for Use) concerns trade secrets and/or binding confidentiality obligations (numbering reflects that of the Procedure Document):

3. Information on design and manufacturing
4. General requirements on safety and performance
5. Risk-benefit analysis and risk management
6. Verification and validation of product.

This Manufacturer's **Authorization** does not confer any powers or authorizations beyond those contained herein. The present authorization should not be interpreted as an extension or renewal of any Authorized Representative Agreement or other contractual relationship. This letter of authorization shall remain valid until the process of registration ends or for **1 year** from the date of this letter, whichever comes later, unless revoked sooner by the Company.

On behalf of Siemens Healthcare Diagnostics Products GmbH


Electronically signed by: Claus Prümper
Reason: I have reviewed this document
Date: 2020-01-16 11:22:10+01:00

Name: Dr. Claus Prümper
Title: Authorized Representative


Electronically signed by: Jürgen Vogt
Reason: I have reviewed this document
Date: 2020-01-15 15:56:49+01:00

Name: Jürgen Vogt
Title: Authorized Representative

Siemens Healthcare Diagnostics Products GmbH
Management: Michael HeinoId, Joerg Berner, Tobias Thaens

Email-von-Behring-Str. 76
35041 Marburg
Germany

Tel.: +49 (6421) 35041

Chairman of the Supervisory Board: Deepak Nathi; Registered Office: Marburg an der Lahn; Commercial registry: Marburg an der Lahn HRB 20654

