

Formularul ofertei (F3.1)

Data depunerii ofertei: "20" noiembrie 2019

Licitația Nr.: 2101397

Către: **IMSP AMT Buiucani.**

ICS Diamedix Impex SRL declară că:

- a) Au fost examinate și nu există rezervări față de documentele de atribuire.
- b) **ICS Diamedix Impex SRL** se angajează să furnizeze/presteze, în conformitate cu documentele de atribuire și condițiile stipulate în specificațiile tehnice și preț, următoarele bunuri și/sau servicii: reagenti si consumabile de laborator.
- c) Suma totală a ofertei fără TVA constituie:
329064,00 lei (trei sute douazeci si noua mii saizeci si patru lei 00 bani).
- d) Suma totală a ofertei cu TVA constituie:
371 010. 24 lei (trei sute saptezeci si unu mii zece lei 24 bani).
- e) Prezenta ofertă va rămâne valabilă pentru perioada de timp specificată în **FDA4.8.**, începînd cu data-limită pentru depunerea ofertei, în conformitate cu **FDA5.2.**, va rămîne obligatorie și va putea fi acceptată în orice moment pînă la expirarea acestei perioade;
- f) În cazul acceptării prezentei oferte, **ICS Diamedix Impex SRL** se angajează să obțină o Garanție de bună execuție în conformitate cu **FDA7**, pentru executarea corespunzătoare a contractului de achiziție publică.
- g) Nu sîntem în nici un conflict de interese, în conformitate cu punctul **IPO5.4**.
- h) Compania semnatară, afiliații sau sucursalele sale, inclusiv fiecare partener sau subcontractor ce fac parte din contract, nu au fost declarate neeligibile în baza prevederilor legislației în vigoare sau a regulamentelor cu incidență în domeniul achizițiilor publice, în conformitate cu punctul **IPO5.5**.

Semnat: _____

Turcanu

L.Ș.

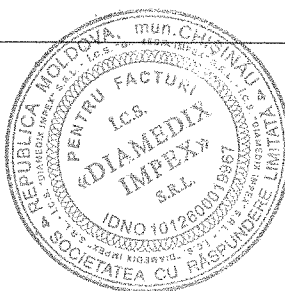
Nume: **Dorel Turcanu**

În calitate de: **Reprezentant vanzari**

Ofertantul: **I.C.S. "Diamedix Impex" S.R.L.**

Adresa: **mun. Chisinau, str. 31 august 1989 108/2**

Data: **"19" noiembrie 2019**

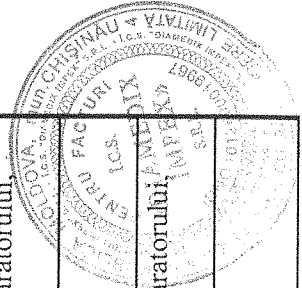


Specificații de preț (F4.2)

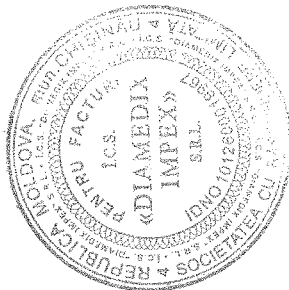
[Acest tabel va fi completat de către ofertant în coloanele 5,6,7,8, iar de către autoritatea contractantă — în coloanele 1,2,3,4,9]

Numărul licitației:	21013977			Data: „20” noiembrie 2019		Alternativa nr.: _____
Denumirea licitației:	Achiziționarea reactivelor de laborator pentru 2020			Lot: _____		Pagina: 1 din 1

Cod CPV	Denumirea bunurilor și/sau a serviciilor	Unitatea de măsură	Canti-tatea	Preț unitar (fără TVA)	Preț unitar (cu TVA)	Suma fara TVA	Suma cu TVA	Termenul de livrare/prestare
1	2	3	4	5	6	7	8	9
2	Analizator hematologic automat ADVIA 2120							
33696500-0	CBC Timepac	set	9	7800,00	8424,00	70200,00	75816,00	La comanda Cumparatorului, timp de 10 zile
3	Analizator hematologic automat ADVIA 2120							
33696500-0	Diff Timepac	set	9	8880,00	9590,40	79920,00	86313,60	La comanda Cumparatorului, timp de 10 zile
4	Analizator hematologic automat ADVIA 2120							
33696500-0	Sheath / Rinse	buc	16	4464,00	5356,80	71424,00	85708,80	La comanda Cumparatorului, timp de 10 zile
5	Analizator hematologic automat ADVIA 2120							
33696500-0	EZ Wash	set	16	3672,00	4406,40	58752,00	70502,40	La comanda Cumparatorului, timp de 10 zile
6	Analizator hematologic automat ADVIA 2120							
33696500-0	TestPoint 3in1 Hematology Control Abnormal1	set	3	4800,00	5184,00	14400,00	15552,00	La comanda Cumparatorului, timp de 10 zile
7	Analizator hematologic automat ADVIA 2120							



Cod CPV	Denumirea bunurilor și/sau a serviciilor	Unitatea de măsură	Cantitatea	Preț unitar (fără TVA)	Preț unitar (cu TVA)	Suma fără TVA	Suma cu TVA	Termenul de livrare/prestare
1	2	3	4	5	6	7	8	9
33696500-0	TestPoint 3in1 Hematology Control Abnormal2	set	3	4800,00	5184,00	14400,00	15552,00	La comanda Cumparatorului, timp de 10 zile
8	Analizator hematologic automat ADVIA 2120							
33696500-0	TestPoint 3in1 Hematology Control Normal	set	3	4800,00	5184,00	14400,00	15552,00	La comanda Cumparatorului, timp de 10 zile
9	Analizator hematologic automat ADVIA 2120							
33696500-0	OptiPoint	set	1	2976,00	3214,08	2976,00	3214,08	La comanda Cumparatorului, timp de 10 zile
10	Analizator hematologic automat ADVIA 2120							
33696500-0	Set Point Calibrator	set	1	2592,00	2799,36	2592,00	2799,36	La comanda Cumparatorului, timp de 10 zile
	TOTAL					329064,00	371010,24	



Semnat: Турчану Дорел Numele, Prenumele: Turcanu Dorel În calitate de: Reprezentant vanzari

Ofertantul: L.C.S. "DIAMEDIX IMPEX" S.R.L. Adresa: mun. Chisinau, str 31 august 1989, 108/2

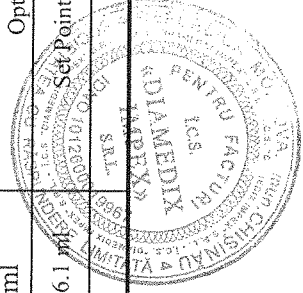
Specificații tehnice (F4.1)

[Acest tabel va fi completat de către ofertant în coloanele 3, 4, 5, 7, iar de către autoritatea contractantă – în coloanele 1, 2, 6, 8]

Numărul licitației:	21013977		Data: „20” noiembrie 2019		Alternativa nr.:
Denumirea licitației:	Achiziționarea reactivilor de laborator pentru 2020		Lot: _____		Pagina: 1 din 1

Semnat: Turcanu Numele, Prenumele: Turcanu Dorel În calitate de: Reprezentant vânzări

Ofertantul: I.C.S. "D IAMEDIX IMPEX" S.R.L. Adresa: mun. Chisinau, str. 31 august 1989, 108/2



SIEMENS

EC Declaration of Conformity



We hereby declare that the product described below conforms to all applicable requirements of Council Directive 98/79/EC for in vitro diagnostic medical devices.

Legal Manufacturer: Siemens Healthcare Diagnostics Inc.
511 Benedict Avenue
Tarrytown, NY, 10591-5097, USA

Place of Manufacture: ThermoFisher Scientific
8365 Valley Pike
Middletown, VA, 22645-0307, USA

EC Authorized Representative: Siemens Healthcare Diagnostics Ltd.
Sir William Siemens Square
Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i CBC TIMEPAC

Catalogue Number (REF): 09826813

Siemens Material Number (SMN): 10312269

Legacy Product Code: T01-3620-52

Classification: General IVD

Conformity Assessment Route: ANNEX III

Document Control Number: DoC_ADVIA 120/2120/2120i CBC TIMEPAC

Version: 1.0

*This declaration of conformity is issued under the sole responsibility of Siemens Healthcare Diagnostics Inc.
This declaration supersedes any declaration issued previously for the same product.*

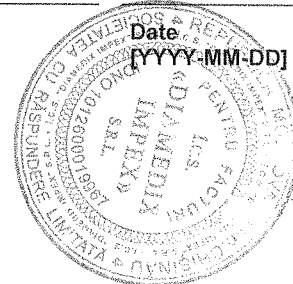
Signature:

Digitally signed by Gee Matthew
Date: 2015.11.12 15:54:48 -05'00'

Matthew Gee
Sr. Manager, Regulatory Affairs
Siemens Healthcare Diagnostics Inc.
Tarrytown, NY, USA

2015-11-12

Date
[YYYY-MM-DD]



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511 Benedict Avenue
Tarrytown, NY, 10591-5097, USA

Place of Manufacture: ThermoFisher Scientific
8365 Valley Pike
Middletown, VA, 22645-0307, USA

EC Authorized Representative: Siemens Healthcare Diagnostics Ltd.
Sir William Siemens Square
Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i DIFF TIMEPAC

Catalogue Number (REF): 00739500

Siemens Material Number (SMN): 10312270

Legacy Product Code: T01-3621-52

Classification: General IVD

Conformity Assessment Route: ANNEX III

Document Control Number: DoC_ADVIA 120/2120/2120i DIFF TIMEPAC

Version: 1.0

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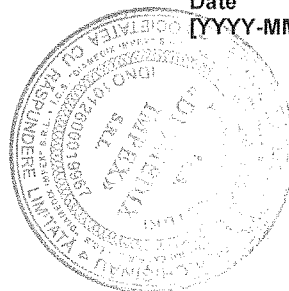
Signature:

Digitally signed by Gee Matthew
Date: 2015.11.12 15:59:38 -05'00'

2015-11-12

Matthew Gee
Sr. Manager, Regulatory Affairs
Siemens Healthcare Diagnostics Inc.
Tarrytown, NY, USA

Date
[YYYY-MM-DD]



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511 Benedict Avenue
Tarrytown, NY, 10591-5097, USA

Place of Manufacture: ThermoFisher Scientific
8365 Valley Pike
Middletown, VA, 22645-0307, USA

EC Authorized Representative: Siemens Healthcare Diagnostics Ltd.
Sir William Siemens Square
Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i EZ WASH

Catalogue Number (REF): 04871500

Siemens Material Number (SMN): 10285021

Legacy Product Code: N/A

Classification: General IVD

Conformity Assessment Route: ANNEX III

Document Control Number: DoC_ADVIA 120/2120/2120i EZ WASH

Version: 1.0

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This declaration supersedes any declaration issued previously for the same product.

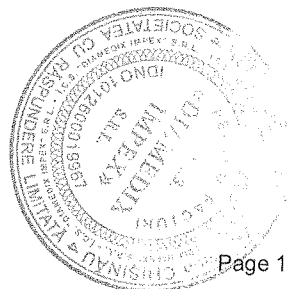
Signature:

Digitally signed by Gee Matthew
Date: 2015.11.12 16:00:43 -05'00'

Matthew Gee
Sr. Manager, Regulatory Affairs
Siemens Healthcare Diagnostics Inc.
Tarrytown, NY, USA

2015-11-12

Date
[YYYY-MM-DD]



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Legal Manufacturer: Siemens Healthcare Diagnostics Inc.
511 Benedict Avenue
Tarrytown, NY, 10591-5097, USA

Place of Manufacture: ThermoFisher Scientific
8365 Valley Pike
Middletown, VA, 22645-0307, USA

EC Authorized Representative: Siemens Healthcare Diagnostics Ltd.
Sir William Siemens Square
Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i OPTIpoint

Catalogue Number (REF): 08100525

Siemens Material Number (SMN): 10312283

Legacy Product Code: T03-3682-54

Classification: General IVD

Conformity Assessment Route: ANNEX III

Document Control Number: DoC_ADVIA 120/2120/2120i OPTIpoint

Version: 1.0

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This declaration supersedes any declaration issued previously for the same product.*

Signature:

Digitally signed by Gee Matthew
Date: 2015.11.12 16:01:44 -05'00'

Matthew Gee
Sr. Manager, Regulatory Affairs
Siemens Healthcare Diagnostics Inc.
Tarrytown, NY, USA

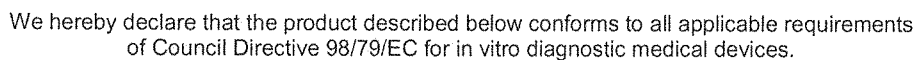
2015-11-12

Date
[YYYY-MM-DD]



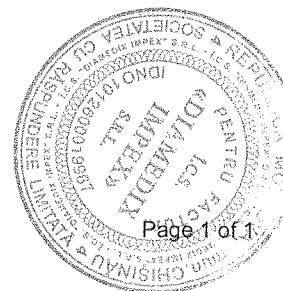
EC DECLARATION OF CONFORMITY

DECLARATION OF CONFORMITY



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Date
[YYYY-MM-DD]



SIEMENS

EC Declaration of Conformity



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Legal Manufacturer: Siemens Healthcare Diagnostics Inc.
511 Benedict Avenue
Tarrytown, NY, 10591-5097, USA

Place of Manufacture: ThermoFisher Scientific
8365 Valley Pike
Middletown, VA, 22645-0307, USA

EC Authorized Representative: Siemens Healthcare Diagnostics Ltd.
Sir William Siemens Square
Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i Sheath/Rinse

Catalogue Number (REF): 02337140 (10L)
01554628 (20L)

Siemens Material Number (SMN): 10316869 (10L)
10312272 (20L)

Legacy Product Code: T01-3664-01 (10L)
T01-3623-01 (20L)

Classification: General IVD

Conformity Assessment Route: ANNEX III

Document Control Number: DoC_ADVIA 120/2120/2120i Sheath/Rinse

Version: 1.0

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Signature:

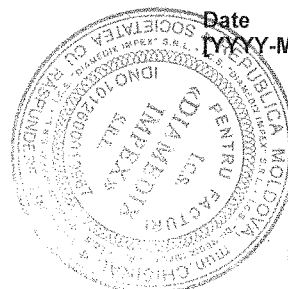
Digitally signed by Gee Matthew
Date: 2015.11.12 16:04:01 -05'00'

2015-11-12

Matthew Gee
Sr. Manager, Regulatory Affairs
Siemens Healthcare Diagnostics Inc.
Tarrytown, NY, USA

Date
[YY-MM-DD]

EC DECLARATION OF CONFORMITY



SIEMENS

EC Declaration of Conformity



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Legal Manufacturer: Siemens Healthcare Diagnostics Inc.
511 Benedict Avenue
Tarrytown, NY, 10591-5097, USA

Place of Manufacture: Streck
7002 South 109th Street
La Vista, NE, 68128, USA

EC Authorized Representative: Siemens Healthcare Diagnostics Ltd.
Sir William Siemens Square
Frimley, Camberley, GU16 8QD, UK

Product Name: ADVIA 120/2120/2120i TESTpoint Hematology Controls

Catalogue Number (REF): 00848547 (Low Control)
05147873 (Norm Control)
08822644 (High Control)

Siemens Material Number (SMN): 10312287 (Low Control)
10312289 (Norm Control)
10312291 (High Control)

Legacy Product Code: T03-3686-54 (Low Control)
T03-3687-54 (Norm Control)
T03-3688-54 (High Control)

Classification: General IVD

Conformity Assessment Route: ANNEX III

Document Control Number: DoC_ADVIA 120/2120/2120i TESTpoint Controls

Version: 1.0

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This declaration supersedes any declaration issued previously for the same product.*



EC DECLARATION OF CONFORMITY

SIEMENS

EC Declaration of Conformity

Signature:

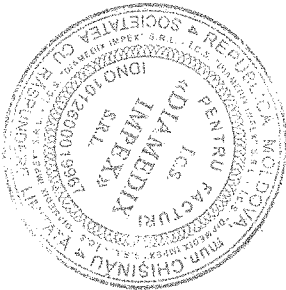


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Date: 2015.11.12 16:04:42 -05'00'

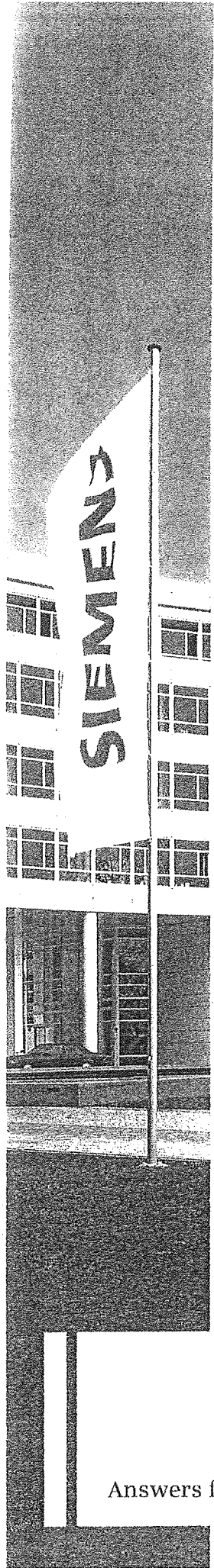
2015-11-12

Matthew Gee
Sr. Manager, Regulatory Affairs
Siemens Healthcare Diagnostics Inc.
Tarrytown, NY, USA

Date
[YYYY-MM-DD]



EC DECLARATION OF CONFORMITY



Daniel Tudor

Siemens Healthcare Diagnostics confirm
that the participant
has passed successfully the training

**ADVIA 2120/120 Field Service Training
(EMEA)**

from

10.09.2012 to 28.09.2012

Eschborn, 28.09.2012

A handwritten signature in black ink, appearing to read 'Philippe Renard'.

Philippe Renard
Trainer



Answers for life.

SIEMENS