

## **STATEMENT**

We, ACON Laboratories, Inc., having a registered office at 5850 Oberlin Drive #340, San Diego, CA 92121 authorize SRL Sanmedico having a registered office at A. Corobceanu street 7A, apt. 9, Chisinău, MD-2012, Moldova

to register, notify, renew or modify the registration of medical devices on the territory of the Republic of Moldova.

Date: March 18, 2024

Signature:

Qiyi Xie, Md, MPH

V.P. of Regulatory & Clinical Affairs

ACON Laboratories, Inc.







## **Product Service**

# **Certificate**

No. Q5 104507 0001 Rev. 03

**Holder of Certificate: ACON Laboratories, Inc.** 

5850 Oberlin Drive, #340 San Diego CA 92121 **USA** 

**Certification Mark:** 



Design and Development, Manufacture and distribution Scope of Certificate: of In Vitro Diagnostic Test Kits and Reagents for the

**Determination of Infectious Diseases, Clinical** Chemistry, Drugs of Abuse, Tumor/Cardiac Marker, Fertility/Pregnancy and Blood Glucose Monitoring

System, Lancing Devices and Lancets

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:Q5 104507 0001 Rev. 03

SH22743A01 Report No.:

Valid from: 2022-09-15 Valid until: 2025-09-06

Christoph Dicks

Head of Certification/Notified Body



# **Certificate**

No. Q5 104507 0001 Rev. 03

Applied Standard(s): EN ISO 13485:2016

Medical devices - Quality management systems -

Requirements for regulatory purposes

(ISO 13485:2016) DIN EN ISO 13485:2016

Facility(ies): ACON Laboratories, Inc.

5850 Oberlin Drive, #340, San Diego CA 92121, USA

Address holder for registration only

**ACON Laboratories, Inc.** 

10125 Mesa Rim Road, San Diego CA 92121, USA

Manufacture and distribution of

In Vitro Diagnostic Test Kits and Reagents for the Determination of Infectious Diseases, Clinical Chemistry, Drugs of Abuse, Tumor/Cardiac Marker, Fertility/Pregnancy and Blood Glucose Monitoring System, Lancing Devices and Lancets

3 7 7

ACON Laboratories, Inc. 6865 Flanders Dr., Suite B, San Diego CA 92121, USA

Storage of

In Vitro Diagnostic Test Kits and Reagents for the Determination of Infectious Diseases, Clinical Chemistry, Drugs of Abuse, Tumor/Cardiac Marker, Fertility/Pregnancy and Blood Glucose Monitoring System, Lancing Devices and Lancets

AZURE Institute, Inc.

10125 Mesa Rim Road, San Diego CA 92121, USA

Design and Development of

In Vitro Diagnostic Test Kits and Reagents for the Determination of Infectious Diseases, Clinical Chemistry, Drugs of Abuse, Tumor/Cardiac Marker, Fertility/Pregnancy and Blood Glucose Monitoring System, Lancing Devices and Lancets

Acon Laboratories Inc.

Guerrero Negro 9942 Parque Industrial Pacifico IV, 22644 Tijuana B.C. CP, MEXICO

Manufacture of

blood glucose test strips, antigen rapid test and IgG/IgM antibody rapid test for infectious disease.







# **EC** Certificate

Full Quality Assurance System Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6) (List A and B and devices for self-testing)

No. V1 104507 0003 Rev. 06

Manufacturer: ACON Laboratories, Inc.

> 5850 Oberlin Drive, #340 San Diego CA 92121

USA

Product Category(ies): Blood glucose measuring systems for self testing

and self-testing devices for clinical chemistry, hematology and pregnancy and ovulation

The Certification Body of TÜV SÜD Product Service GmbH declares that the aforementioned manufacturer has implemented a quality assurance system for design, manufacture and final inspection of the respective devices / device families in accordance with IVDD Annex IV. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of List A devices an additional Annex IV (4) certificate is mandatory. All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:V1 104507 0003 Rev. 06

SH22743EXT01 Report no.:

Valid from: 2022-05-04 Valid until: 2025-05-26

2022-05-04 Date,

> Christoph Dicks Head of Certification/Notified Body



## **EC** Certificate

Full Quality Assurance System Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6) (List A and B and devices for self-testing)

## No. V1 104507 0003 Rev. 06

On Call Plus Blood Glucose Monitoring System, Model(s):

On Call Plus Blood Glucose Test Strips,

On Call EZ II Blood Glucose Monitoring System.

On Call Advanced Blood Glucose Monitoring System,

On Call Advanced Blood Glucose Test Strips. On Call Chosen Blood Glucose Test Strips,

On Call Vivid Blood Glucose Monitoring System (OGM-101), On Call Vivid Blood Glucose Test Strips (OGS-101),

On Call Sharp Blood Glucose Monitoring System (OGM-

121),

On Call Sharp Blood Glucose Test Strips (OGS-121)

On Call Plus II Blood Glucose Monitoring System (OGM-

On Call Plus II Blood Glucose Test Strips (OGS-171),

On Call Extra Blood Glucose Monitoring System (OGM-191).

On Call Extra Blood Glucose Test Strips (OGS-191),

On Call GK Dual Blood Glucose & Ketone Monitoring

System (OGM-161),

On Call Blood Ketone Test Strips (OGS-161),

Urinalysis Reagent Strips (Urine),

UTI Urinary Tract Infection Test Strips.

Cholesterol Monitoring System (CCM-111),

CHOL Total Cholesterol Test Devices (CCS-111).

TRIG Triglycerides Test Devices (CCS-112),

HDL High Density Lipoprotein Test Devices (CCS-113),

3-1 Lipid Panel Test Devices (CCS-114),

Cholesterol CTRL Control Devices,

Cholesterol Monitoring System (CCM-101),

CHOL Total Cholesterol Test Strips (CCS-101).

PT/INR Monitoring System (CCM-151),

PT/INR Test Strips (CCS-151),

Hemoglobin Testing System (CCM-141),

Hemoglobin Test Strips (CCS-141),

hCG Pregnancy Rapid Test Cassette (Urine),

Pregnancy Rapid Test Midstream,

On Call Extra Mobile Blood Glucose Monitoring System

(OGM-281),

On Call Sure Blood Glucose Monitoring System (OGM-211), On Call Sure Sync Blood Glucose Monitoring System (OGM-

212),

On Call Sure Blood Glucose Test Strips (OGS-211),

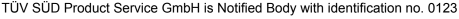
GIMA Blood Glucose Monitoring System,

GIMA Bluetooth Blood Glucose Monitoring System,

GIMA Blood Glucose Test Strips,

On Call GU Dual Blood Glucose & Uric Acid Monitoring









## **EC** Certificate

Full Quality Assurance System Directive 98/79/EC on In Vitro Diagnostic Medical Devices (IVDD), Annex IV excluding (4, 6) (List A and B and devices for self-testing)

## No. V1 104507 0003 Rev. 06

System (OGM-201),

On Call Blood Uric Acid Test Strips (OGS-201),

LH Ovulation Rapid Test Cassette (Urine).

Ovulation Rapid Test Midstream,

Ovulation & Pregnancy Test Combo Pack,

On Call Extra Voice Blood Glucose Monitoring System (OGM-291),

Early Detection Pregnancy Test,

Digital Pregnancy Test.

Go-Keto Blood Glucose & Ketone Monitoring System (OGM-

Go-Keto Blood Ketone Test Strips (OGS-161),

Go-Keto Blood Glucose Test Strips,

On Call Extra GM Blood Glucose Monitoring System(OGM-

On Call Extra GM Blood Glucose Test Strips (OGS-191),

On Call Plus GM Blood Glucose Monitoring System,

On Call Plus GM Blood Glucose Test Strips,

Go-Keto Urinalysis Reagent Strips

ACON Laboratories, Inc. Facility(ies):

5850 Oberlin Drive, #340, San Diego CA 92121, USA

ACON Laboratories, Inc.

10125 Mesa Rim Road, San Diego CA 92121, USA

AZURE Institute, Inc.

10125 Mesa Rim Road, San Diego CA 92121, USA

Acon Laboratories Inc.

Guerrero Negro 9942 Parque Industrial Pacifico IV, 22644 Tijuana

B.C. CP, MEXICO

TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

# **Declaration of Conformity**

# ACON Laboratories, Incorporated 5850 Oberlin Drive #340 San Diego, CA 92121, USA

We, the manufacturer, declare under our sole responsibility that the *in vitro* diagnostic device:

Mission® Urinalysis Reagent Strips (U031-XX1)

classified as Others in the directive 98/79/EC,

meets all the provisions of the directive 98/79/EC on in vitro diagnostic medical devices which apply to it

The self-declaration is according to Annex III (excluding Section 6) of the Directive.

Authorized Representative: Medical Device Safety Service GmbH Schiffgraben 41 30175 Hannover, Germany

Signed this 11 day of February, 2020 in San Diego, CA USA

Qiyi Xie, MD, MPH Senior Staff, Regulatory Affairs & Clinical Affairs Acon Laboratories, Inc.

# Mission® Urinalysis Reagent Strips and Urine Analyzers



# **Urinalysis Reagent Strips**

## Simple and Accurate

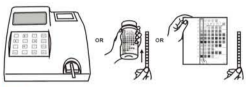
- Analytical sensitivity better than or comparable to market leaders
- · High quality color chart ensures accurate visual reading

- · Compatible for visual and analyzer reading
- · More than 30 different combinations available

## Multiple Packaging Options and Long Shelf Life

- Canister Packaging
   Available in 25, 50, 100 and 150 strips per kit
  - · 2 year shelf life for unopened canisters which offers cost savings and convenience for high volume testing
- 3 month shelf life for strips in opened canisters Pouch Packaging New!
- · Single-strip Pouch
  - Individually packaged strips with 1, 3, 6 and 20 strips and 1 color chart per kit for OTC or low volume testing
- . Unique packaging maintains 2 year shelf life for all strips in the kit compared to 3 months for remaining strips in an opened canister
- Multi-strip Pouch
- · Canister Refill Kits with 25 strips/pouch uniquely packaged to save cost for low volume testing and extended shelf life by using the canister for refills





Ste	Step 1: Immerse strip into urine		Step 2: Remove excess urine				Step 3: Obtain results by analyzer or visual reading																																	
Catalog No. of Type of Strip *		Of the second	Reading Method Analyzer-Read			Parameters																																		
Catalog No.	No. of Parameters	For Visual Reading	For Analyzer Reading (U120/U500)	Strips per Canister	Pouch Packaging*	Visual	U120	U500	Strips: Standard (S) or Additional (A)	ASC	GLU	BIL	KET	sg	BLO	рН	PRO	URO	NIT	LEU	ALB	CRE																		
U031-131	13	13C	NA	100"	<b>✓</b>	1	NA	NA	Α	*	*	*	*	*	*	*	*	*	*	*	*	*																		
U031-111	11		11A	100	4	1	1	1	S	*	*	*	*	*	*	*	*	*	*	*																				
		12	10U	100		4	~	1	S		*	*	*	*	*	*	*	*	*	*																				
U031-101	10		10A		¥	1	1	~	Α	*	*	*	*	*	*	*	*	*	*																					
			10C	100"		1	/	1	S		*		*	*	*	*	*		*	*	*	*																		
U031-091	9		9U	100	✓	<b>~</b>	1	1	S		*	*	*	*	*	*	*	*	*																					
			8U			1	1	1	Α		*	*	*		*	*	*	*	*																					
U031-081	8		8N	100	¥	~	1	1	S		*		*	*	*	*	*		*	*																				
11001 071			8S	100		1	<b>V</b>	1	A		*			*	*	*	*	*	*	*																				
U031-071	7		7N	100	✓	1	1	1	A		*		*		*	*	*		*	*																				
U031-061	6	6N	6NE 6UE	100	✓	4	V	4	A		*	*			*	*	*	100	*	*		$\blacksquare$																		
		6U 5B	5BE	_		4	V	4			*	*	*	*	*	*	*	*	*																					
		5N	5NE	-		4	1	1	1	-	*		*		*	*	*		*	*		Н																		
U031-051	5	5N 5S	5SE	100	~	7	·	· ·	A		*	_	_	*	*	*	*	-		*		$\vdash$																		
		5U	5UE	1		<del>*</del>	9	_	•	*	_	Ĥ	*	-		*	*	*	$\vdash$	$\vdash$																				
		48	4SE			1	1	1			*	_		*	_	*	*	-																						
		4B	4BE			<i>y y y</i>	~				*				*	*	*																							
01022201200	9	4K	4KE	10022				· ·	· ·	· ·	· ·	~	_	1	1			*		*			*	*																
U031-041	4	4G	4GE	100	100								~	~	~	~	~	~	~	~	· ·		1		Α		*				*		*			*				
		4N	4NE		2	1		3						*		*		*	*																					
		4P	4PE			1	1	1			*						*		*	*																				
		3P	3PE			1	<b>V</b>	1			*					*	*																							
U031-031	3	3K	3KE	100		, [	<i>y</i> [	~ [	<i>y</i> [		,			[	, [	1		, [	<i>y</i> [	, [		, [	2	, V	V	1	A		*		*				*					
0031-031		3G	3GE	] 100	*	1	1 1 1	<b>~</b>	^		*		*			*																								
		3N	3NE			4	~	V							*				*	*																				
		2G	2GE			1	1	1			*						*																							
		2K	2KE			1	<b>✓</b>	1			*		*																											
12022017220	020	2N	2NE	1722	N	1	<b>V</b>	1							*					*																				
U031-021	2	2B	2BE	100	<b>*</b>	1	<b>V</b>	1	A		*		*																											
		2U	2UE			4	1	1											*	*																				
		28	2SE	400		1	V	1						*		*																								
		2C	2CE	100*		4	V	4								-					*	*																		
		1B	1BE					4	Y								*							$\vdash$																
11034 044		1P	1PE	100									1	<b>V</b>	¥	,				_	-	-	*	_	-				$\vdash$											
U031-011	1	1G	1GE	100	<b>*</b>	1	<b>V</b>	<b>✓</b>	Α		*			-		-	_					$\vdash$																		
		1K	1KE	-		4	<b>Y</b>	V					*	_	_	_			_	_		$\vdash$																		
		1R	1RE			✓	✓	1				Ц.		_			*					ш																		



Visual Strip Size 1-6 Parameters: 5 mm x 80 mm; 7-11 Parameters: 5 mm x 108 mm;

"E" means extended strip length for 1-6 Parameters

12-13 Parameters: 5 mm x 121 mm

U120/U500 Strip Size 1-11 Parameters: 5 mm x 108 mm;

<sup>▲</sup> Single-strip Pouch available in 1,3, 6 and 20 strip kit
Canister Refill Kit, with 25 strips per pouch or canister, available in 3-pouch and 1- canister kit, or 4-pouch kit



Not available in canisters of 150 strips

Also available in canisters of 25, 50 and 150 strips

# **U120 Urine Analyzer**



- Accurate

   Up to 120 tests/hour in Continuous Test Option
- · Capable of reading 1 strip at a time in Single Test Option
- · Test modes include Routine, STAT and QC
- · Automatic calibration for accurate results and easy operation

- Can read up to 4 Strip combinations with 8, 9, 10, 11 parameters, additional strips with 1-11 parameters available upon request
- · Minimal training required

- Convenient Operation
   Saves and recalls the last 2,000 results automatically
- · Audible beep signals operator to dip strips in urine
- Can print up to 3 copies per test for convenient reviewing and easy record keeping
- · Option to print results on sticker paper for quick and simple record management

## **Easy Data Management**

- Includes RS232C port for easy data transfer to an external computer or LIS
   Optional Barcode Reader to record patient ID

## Unique Lockout Functions new!

- Strip Lockout
  - Prevents using strips of another brand on the U120 Urine Analyzer
  - · Requires barcode reader scan or manual entry of the canister code
- User Lockout

  - Eliminates unapproved users from testing
     Up to 10 lab operators can perform testing, but only the lab administrator can change analyzer settings
- QC Lockout
  - · Prevents testing without passing QC
  - Prevents testing without passing 4C
     QC tests can be performed once every 8 hours, day, week or month
     Analyzer will alert when to run QC test

  - . If QC tests fail, analyzer will switch to STAT mode and list "E" at the end of each test number

## Specifications

Feature	Specifications				
Analyzer Type	Manual				
Methodology	Reflectance Photometry				
Detection	Photosensitive Diode				
Throughput	Single Test Option: 60 tests/hour Continuous Test Option: 120 tests/hour				
Test Modes	Routine, STAT and QC				
Lockout Functions	Strip Lockout: Available Upon Request; Us	er/QC Lockout: Included with option to turn ON/OF			
Memory	Last 2,000 results	-			
Strip Incubation Time	1 Minute				
Wavelength of Monochromatic LED	525 nm and 635 nm				
Standard Strips	8, 9, 10, 11 Parameters (5 mm x 108 mr	n)			
Additional Strips Available	1-11 Parameters (5 mm x 108 mm); see URS Parameters				
Total Combinations Per Analyzer	4 Combinations				
Analyzer Ports	Standard RS232C Port for Barcode Reader or Data Transfer USB Port for Data Transfer 25 Pin Parallel Port for External Printer				
Capabilities	Internal Thermal Printer (included) Optional External Printer (not included)	RS232C Barcode Reader (optional) USB or RS232C Data Transfer Cable (optional)			
Major Readable Barcodes	Code 128, Code 39, Codabar (NW-7), Interleaved 25, UPC-A, UPC-E, EAN 8, EAN 13				
Calibration	Automatic				
Available Languages on the Screen	English and additional language(s)				
Operating Conditions	0-40°C (32-104°F); ≤ 85% RH				
Storage Conditions	-5-50°C (23-122°F); ≤90% RH				
Power Source	100-240 VAC, 50-60 Hz				
Dimensions (L x W x H)	27.2 cm x 26.9 cm x 14.6 cm (10.7" x 10.6" x 5.7")				
Display Dimensions (L x W)	10.8 cm x 5.7 cm (4.2" x 2.2")				
Weight	2.6 kg (5.7 lbs)				

## **Ordering Information**

Product Name	Catalog No.	Co	mponents		Kit Box Dimensions (L x W x H) & Weight	Carton Dimensions (L x W x H) & Weight	Number of Kits/Carton	
U120 Urine Analyzer	/ <b>+</b>	1 Urine Analyzer 1 Strip holder		2 Fuses (2.0A) 1 Power Cord	42.0 cm x 41.5 cm x 3	1 cm; 5.0 kg	4	
O 120 Offile Affalyzer	U111-101 <sup>à</sup>	2 Printer Paper Rolls		1 Quick Start Guide 1 Instruction Manual	16.4" x 16.2" x 12.	1"; 176.4 oz		
U120 Urine Analyzer	U111-111√ <sup>†</sup>	1 Urine Analyzer 1 Strip holder		2 Fuses (2.0A) 1 Power Cord	44.5cm x 44.5cm x 4	0.0cm; 5.5 kg		
with Barcode Reader	om-m	2 Printer Paper Rolls 1 Barcode Reader (RS232C)		1 Serial Splitter Cable (RS232C) 1 Quick Start Guide 1 Instruction Manual	17.5" x 17.5" x 15.	1		
Barcode Reader	U221-111√ <sup>†</sup>	1 Barcode Reader (I	RS232C)	1 Serial Splitter Cable (RS232C)	23.6 cm x 10.8 cm x 7.8 cm; 0.482 kg 9.3" x 4.3" x 3.1"; 17.0 oz	63.0 cm x 37.0 cm x 30.0 cm; 12.0 kg 24.8" x 14.6" x 11.8"; 423.3 oz	22	
Printer Paper Rolls	11101 101	4 Printer Paper Rolls	Thermal Paper (0.06 m x 20 m): 200 results/ro		12.0 cm x 12.0 cm x 6.5 cm; 0.36kg 4.7" x 4.7" x 2.6"; 12.7oz	63.0 cm x 37.0 cm x 30.0 cm; 19.4 kg 24.8" x 14.6" x 11.8"; 684.3 oz	50	
rinter raper itolis	U121-101			per (0.06 m x 9 m): 100 results/roll	12.0 cm x 12.0 cm x 6.5 cm; 0.4 kg 4.7" x 4.7" x 2.6"; 14.1 oz	63.0 cm x 37.0 cm x 30.0 cm; 21.4 kg 24.8" x 14.6" x 11.8"; 684.3 oz; 754.9 oz	kg	
U120 Data Transfer Kit	U221-131 <sup>à</sup>	1 Data Transfer Cable	(RS232C)	1 Package Insert	16.0 cm x 13.0 cm x 3.5 cm; 0.147 kg 6.3" x 5.1" x 1.4"; 5.2 oz	25.0 cm x 21.0 cm x 15.0 cm; 1.36 kg 9.8" x 8.3" x 5.9"; 48.0 oz	8	

# **U500 Urine Analyzer**



Accurate and Efficient

• Up to 500 tests/hour for medium/large volume sample testing
• Professional accuracy equivalent to market leader
• Automatic strip detection and alignment for better efficiency
• Test modes include Routine, STAT and QC

Easy to Operate

Large touch screen LCD offers simple menu navigation

Uniquely designed strip platform/waste tray unit for easy one-step cleaning

CONVENIENT

Automatic calibration and waste disposal reduce hands-on time

Can read strips with 8, 9, 10, 11 parameters, additional strips with 1-11 parameters available upon request

Strip selection of up to 4 combinations for analyzer reading

Stories up to 2,000 records and automatically flags abnormal results

Capable of printing results on sticker paper for quick and easy record management

Data Management Capability
Includes R\$232C port for easy data transfer to an external computer or LIS
Optional Barcode Reader to record patient ID
Unique Lockout Functions Coming Soon!

Strip Lockout
 Prevents using strips of another brand on the U500 Urine Analyzer
 Requires barcode reader scan or manual entry of the canister code

User Lockout

Eliminates unapproved users from testing
 Up to 10 lab operators can perform testing, but only the lab administrator can change analyzer settings.

QC Lockout
 Prevents testing without passing QC

QC tests can be performed once every 8 hours, day, week or month
 Analyzer will alert when to run QC test

If QC tests fail, analyzer will switch to STAT mode and list "E" at the end of each test number

## **Specifications**

Feature	Specifications				
Analyzer Type	Semi-Automatic				
Methodology	Reflectance Photometry				
Detection	Photosensitive Diode				
Throughput	500 tests/hour (Measuring cycle: 7 secon	ds/test)			
Test Modes	Routine, STAT and QC	Herricocourts is			
Lockout Functions	Strip Lockout: Available Upon Request; User,	/QC Lockout: Included with option to turn ON/OFF			
Memory	Last 2,000 Records	**			
Strip Incubation Time	1 Minute				
Wavelength	525 and 635 nm				
Standard Strips	8, 9, 10, 11 Parameters (5 mm x 108 mm)				
Additional Strips Available	1-11 Parameters (5 mm x 108 mm); see URS Parameters				
Total Combinations Per Analyzer	4 Combinations				
Waste Disposal Capacity	Up to 150 Strips				
Analyzer Ports	Standard RS232C Port for Barcode Read 25 Pin Parallel Port for External Printer	er or Data Transfer			
Capabilities	Internal Thermal Printer (included) Optional External Printer (not included)	RS232C Barcode Reader (optional) RS232C Data Transfer Cable (optional)			
Major Readable Barcodes	Code 128, Code 39, Codabar (NW-7), Interle	aved 25, UPC-A, UPC-E, EAN 8, EAN 13			
Calibration	Automatic				
Available Languages on the Screen	English and additional language(s)				
Operating Conditions	0-40°C (32-104°F); ≤85% RH				
Storage Conditions	-5-50°C (23-122°F); ≤90% RH				
Power Source	100-240 VAC, 50-60 Hz				
Dimensions (L x W x H)	36.6 cm x 28.3 cm x 19.5cm (14.4" x 11.1" x 7.7")				
Display Dimensions (LxW)	11.5 cm x 9.0 cm (4.5" x 3.5")				
Weight	4.0 kg (8.8 lbs)				

## **Ordering Information**

Product Name	Catalog No. Components				Kit Box Dimensions (L x W x H) & Weight	Carton Dimensions (L x W x H) & Weight	Number of Kits/Carton	
Settlebusters, terrocommission on a natural	112	1 Urine Analyzer 1 Strip Platform/Waste	a Tray	2 Fuses (2.0A)	51.0 cm x 42.0 cm x 3	8.5 cm; 7 kg		
U500 Urine Analyzer	U211-101√	2 Printer Paper Rolls			20.1" X 16.5" x 15.	2"; 246.9 oz	1	
U500 Urine Analyzer	U211-111√	1 Urine Analyzer 1 Strip Platform/Waste	e Tray	2 Fuses (2.0A) 1 Power Cord	55.0 cm x 55.0 cm x	55.0cm; 9.2 kg	1	
with Barcode Reader	02111111	2 Printer Paper Rolls 1 Barcode Reader (RS232C)		Serial Splitter Cable (RS232C)     Instruction Manual	21.7" x 21.7" x 21.7"; 324.5 oz			
Barcode Reader	U221-111à	1 Barcode Reader (I	RS232C)	1 Serial Splitter Cable (RS232C)	23.6 cm x10.8 cm x 7.8 cm; 0. 482 kg 9.3" x 4.3" x 3.1"; 17.0 oz	63.0 cm x 37.0 cm x 30.0 cm; 12 kg 24.8" x 14.6" x 11.8"; 423.3 oz	22	
Printer Paper Rolls	VAID D7045	a A Delete	4 Printer Paper Rolls	Thermal P	aper (0.06 m x 20 m): 200 results/roll	12.0 cm x 12.0 cm x 6.5 cm; 0.360 kg 4.7" x 4.7" x 2.6"; 12.7 oz	63.0 cm x 37.0 cm x 30.0 cm; 19.4 kg 24.8" x 14.6" x 11.8"; 684.3 oz	50
Filliter Faper Noils	U121-101	4 Filitter Faper Rolls	Sticker Pa	per (0.06 m x 9 m): 100 results/roll	12.0 cm x 12.0 cm x 6.5 cm; 0.40 kg 63.0 cm x 37.0 cm x 30.0 cm; 21.4 4.7" x 4.7" x 2.6"; 14.10z 24.8" x 14.6" x 11.8"; 684.3 oz; 754.9		(18/8)	
U500 Data Transfer Kit	U221-131√	1 Data Transfer Cable	(RS232C)	1 Package Insert	16.0 cm x 13.0 cm x 3.5 cm; 0.147kg 6.3" x 5.1" x 1.4"; 5.2 oz	25.0 cm x 21.0 cm x 15.0 cm; 1.36 kg 9.8" x 8.3" x 5.9"; 48.0 oz	8	

## We also offer other rapid diagnostic and medical products:

Blood Glucose Monitoring Systems, Immunoassay EIA/ELISA and more.

✓ CE Marked for sale in the European Community † Cleared for US 510(k)



ACON Laboratories, Inc., 10125 Mesa Rim Road, San Diego, CA 92121, U.S.A. • Tel: 1-858-875-8000 • Fax: 1-858-200-0729 • E-mail: info@aconlabs.com Please visit our website for details: www.aconlabs.com



## Package Insert

REF U031-011	REF U031-051	REF U031-091	
REF U031-021	REF U031-061	REF U031-101	
REF U031-031	REF U031-071	REF U031-111	English
REF U031-041	REF U031-081		

For rapid detection of multiple analytes in human urine. For in vitro diagnostic use only

## INTENDED USE

The Urinalysis Reagent Strips (Urine) are firm plastic strips onto which several separate reagent areas are affixed. The test is for the qualitative and semi-quantitative detection of one or more of the following analytes in urine: Ascorbic acid, Glucose, Bilirubin, Ketone (Acetoacetic acid), Specific Gravity, Blood, pH, Protein, Urobilinogen, Nitrite and Leukocytes.

### SUMMARY

Urine undergoes many changes during states of disease or body dysfunction before blood composition is altered to a significant extent. Urinalysis is a useful procedure as an indicator of health or disease, and as such, is a part of routine health screening. The Urinalysis Reagent Strips (Urine) can be used in general evaluation of health, and aids in the diagnosis and monitoring of metabolic or systemic diseases that affect kidney function, endocrine disorders and diseases or disorders of the urinary tract.

### PRINCIPLE AND EXPECTED VALUES

Ascorbic acid: This test involves decolorization of Tillmann's reagent. The presence of ascorbic acid causes the color of the test field to change from blue-green to orange. Patients with adequate diet may excrete 2-10 mg/dL daily. After ingesting large amounts of ascorbic acid, levels can be around 200 mg/dL.

Glucose: This test is based on the enzymatic reaction that occurs between glucose oxidase, peroxidase and chromogen. Glucose is first oxidized to produce gluconic acid and hydrogen peroxide in the presence of glucose oxidase. The hydrogen peroxide reacts with potassium iodide chromogen in the presence of peroxidase. The extent to which the chromogen is oxidized determines the color which is produced, ranging from green to brown. Glucose should not be detected in normal urine. Small amounts of glucose may be excreted by the kidney.3 Glucose concentrations as low as 100 mg/dL may be considered abnormal if results are consistent.

Bilirubin: This test is based on azo-coupling reaction of bilirubin with diazotized dichloroaniline in a strongly acidic medium. Varying bilirubin levels will produce a pinkish-tan color proportional to its concentration in urine. In normal urine, no bilirubin is detectable by even the most sensitive methods. Even trace amounts of bilirubin require further investigation. Atypical results (colors different from the negative or positive color blocks shown on the color chart) may indicate that bilirubin-derived bile pigments are present in the urine specimen, and are possibly masking the bilirubin reaction.

Ketone: This test is based on ketones reacting with nitroprusside and acetoacetic acid to produce a color change ranging from light pink for negative results to a darker pink or purple color for positive results. Ketones are normally not present in urine. Detectable ketone levels may occur in urine during physiological stress conditions such as fasting, pregnancy and frequent strenuous exercise. 46 In starvation diets, or in other abnormal carbohydrate metabolism situations, ketones appear in the urine in excessively high concentration before serum ketones are elevated.

Specific Gravity: This test is based on the apparent pKa change of certain pretreated polyelectrolytes in relation to ionic concentration. In the presence of an indicator, colors range from deep blue-green in urine of low ionic concentration to green and yellow-green in urine of increasing ionic concentration. Randomly collected urine may vary in specific gravity from 1.003-1.035.8 Twenty-four hour urine from healthy adults with normal diets and fluid intake will have a specific gravity of 1.016-1.022.8 In cases of severe renal damage. the specific gravity is fixed at 1.010, the value of the glomerular filtrate.

**Blood:** This test is based on the peroxidase-like activity of hemoglobin which catalyzes the reaction of diisopropylbenzene dihydroperoxide and 3,3',5,5'-tetramethylbenzidine. The resulting color ranges from orange to green to dark blue. Any green spots or green color development on the reagent area within 60 seconds is significant and the urine specimen should be examined further. Blood is often, but not invariably, found in the urine of menstruating females. The significance of a trace reading varies among patients and clinical judgment is required in these specimens.

pH: This test is based on a double indicator system which gives a broad range of colors covering the entire urinary pH range. Colors range from orange to yellow and green to blue. The expected range for normal urine specimens from newborns is pH 5-7.9 The expected range for other normal urine specimens is pH 4.5-8, with an average result of pH 6.

**Protein:** This reaction is based on the phenomenon known as the "protein error" of pH indicators where an indicator that is highly buffered will change color in the presence of proteins (anions) as the indicator releases hydrogen ions to the protein. At a constant pH, the development of any green color is due to the presence of protein. Colors range from yellow to yellow-green for negative results and green to green-blue for positive results. 1-14 mg/dL of protein may be excreted by a normal kidney. 10 A color matching any block greater than trace indicates significant proteinuria. Clinical judgment is required to evaluate the significance of trace results.

Urobilinogen: This test is based on a modified Ehrlich reaction between p-diethylaminobenzaldehyde and urobilinogen in strongly acidic medium to produce a pink color. Urobilinogen is one of the major compounds produced in heme synthesis and is a normal substance in urine. The expected range for normal urine with this test is 0.2-1.0 mg/dL (3.5-17 µmol/L). A result of 2.0 mg/dL (35 µmol/L) may be of clinical significance, and the patient specimen should be further evaluated.

Nitrite: This test depends upon the conversion of nitrate to nitrite by the action of Gram negative bacteria in the urine. In an acidic medium, nitrite in the urine reacts with p-arsanilic acid to form a diazonium compound. The diazonium compound in turn couples with 1 N-(1-naphthyl) ethylenediamine to produce a pink color. Nitrite is not detectable in normal urine. The nitrite area will be positive in some cases of infection, depending on how long the urine specimens were retained in the bladder prior to collection. Retrieval of positive cases with the nitrite test ranges from as low as 40% in cases where little bladder incubation occurred, to as high as approximately 80% in cases where bladder incubation took place for at least 4 hours.

**Leukocytes:** This test reveals the presence of granulocyte esterases. The esterases cleave a derivatized pyrazole amino acid ester to liberate derivatized hydroxy pyrazole. This pyrazole then reacts with a diazonium salt to produce a beige-pink to purple color. Normal urine specimens generally yield negative results. Trace results may be of guestionable clinical significance. When trace results occur, it is recommended to retest using a fresh specimen from the same patient. Repeated trace and positive results are of clinical significance

## REAGENTS AND PERFORMANCE CHARACTERISTICS

Based on the dry weight at the time of impregnation, the concentrations given may vary within manufacturing tolerances. The following table below indicates read times and performance characteristics for each parameter.

Reagent	Read Time	Composition	Description
Ascorbic Acid (ASC)	30 seconds	2,6-dichlorophenolindophenol; buffer and non-reactive ingredients	Detects ascorbic acid as low as 5-10 mg/dL (0.28-0.56 mmol/L).
Glucose (GLU)	30 seconds	glucose oxidase; peroxidase; potassium iodide; buffer; non-reactive ingredients	Detects glucose as low as 50-100 mg/dL (2.5-5 mmol/L).
Bilirubin (BIL)	30 seconds	2, 4-dichloroaniline diazonium salt; buffer and non-reactive ingredients	Detects bilirubin as low as 0.4-1.0 mg/dL (6.8-17 μmol/L).
Ketone (KET)	40 seconds	sodium nitroprusside; buffer	Detects acetoacetic acid as low as 2.5-5 mg/dL (0.25-0.5 mmol/L).
Specific Gravity (SG)	45 seconds	bromthymol blue indicator; buffer and non-reactive ingredients; poly (methyl vinyl ether/maleic anhydride); sodium hydroxide	Determines urine specific gravity between 1.000 and 1.030. Results correlate with values obtained by refractive index method within ± 0.005.
Blood (BLO)	60 seconds	3,3',5,5'-tetramethylbenzidine (TMB); diisopropylbenzene dihydroperoxide; buffer and non-reactive ingredients	Detects free hemoglobin as low as 0.018-0.060 mg/dL or 5-10 Ery/µL in urine specimens with ascorbic acid content of < 50 mg/dL.
pH	60 seconds	methyl red sodium salt; bromthymol blue; non-reactive ingredients	Permits the quantitative differentiation of pH values within the range of 5-9.
Protein (PRO)	60 seconds	tetrabromophenol blue; buffer and non-reactive ingredients	Detects albumin as low as 7.5-15 mg/dL (0.075-0.15 g/L).
Urobilinogen (URO)	60 seconds	p-diethylaminobenzaldehyde; buffer and non-reactive ingredients	Detects urobilinogen as low as 0.2-1.0 mg/dL (3.5-17 $\mu$ mol/L).
Nitrite (NIT)	60 seconds	p-arsanilic acid; N-(1-naphthyl) ethylenediamine; non-reactive ingredients	Detects sodium nitrite as low as 0.05-0.1 mg/dL in urine with a low specific gravity and less than 30 mg/dL ascorbic acid.
Leukocytes (LEU)	120 seconds	derivatized pyrrole amino acid ester; diazonium salt; buffer; non-reactive ingredients	Detects leukocytes as low as 9-15 white blood cells Leu/μL in clinical urine.

The performance characteristics of the Urinalysis Reagent Strips (Urine) have been determined in both laboratory and clinical tests. Parameters of importance to the user are sensitivity, specificity, accuracy and precision. Generally, this test has been developed to be specific for the parameters to be measured with the exceptions of the interferences listed. Please refer to the Limitations section in this package insert.

Interpretation of visual results is dependent on several factors: the variability of color perception, the presence or absence of inhibitory factors, and the lighting conditions when the strip is read. Each color block on the chart corresponds to a range of analyte concentrations.

## PRECAUTIONS

- For in vitro diagnostic use only. Do not use after the expiration date.
- The strip should remain in the closed canister until use.
- Do not touch the reagent areas of the strip.
- Discard any discolored strips that may have deteriorated
- All specimens should be considered potentially hazardous and handled in the same manner as an infectious agent
- The used strip should be discarded according to local regulations after testing.

## STORAGE AND STABILITY

Store as packaged in the closed canister either at room temperature or refrigerated (2-30°C). Keep out of direct sunlight. The strip is stable through the expiration date printed on the canister label. Do not remove the desiccant. Remove only enough strips for immediate use. Replace cap immediately and tightly. **DO NOT FREEZE.** Do not use beyond the expiration date

Note: Once the canister has been opened, the remaining strips are stable for up to 3 months. Stability may be reduced in high humidity conditions

## SPECIMEN COLLECTION AND PREPARATION

A urine specimen must be collected in a clean and dry container and tested as soon as possible. Do not centrifuge. The use of urine preservatives is not recommended. If testing cannot be done within an hour after voiding, refrigerate the specimen immediately and let it return to room temperature before testing.

Prolonged storage of unpreserved urine at room temperature may result in microbial proliferation with resultant changes in pH. A shift to alkaline pH may cause false positive results with the protein test area. Urine containing glucose may decrease in pH as organisms metabolize the glucose.

Contamination of the urine specimen with skin cleansers containing chlorhexidine may affect protein (and to a lesser extent, specific gravity and bilirubin) test results.

## MATERIALS Materials Provided

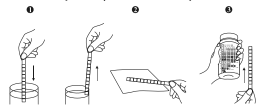
· Package insert

· Specimen collection container Timer

Allow the strip, urine specimen, and/or controls to reach room temperature (15-30°C) prior to testing.

- Remove the strip from the closed canister and use it as soon as possible. Immediately close the canister tightly after removing the required number of strip(s). Completely immerse the reagent areas of the strip in fresh, well-mixed urine and immediately remove the strip to avoid dissolving the reagents. See illustration 1 below.
- While removing the strip from the urine, run the edge of the strip against the rim of the urine container to remove excess urine. Hold the strip in a horizontal position and bring the edge of the strip into contact with an absorbent material (e.g. a paper towel) to avoid mixing chemicals from adjacent reagent areas and/or soiling hands with urine. See illustration 2 below.
- Compare the reagent areas to the corresponding color blocks on the canister label at the specified times. Hold the strip close to the color blocks and match carefully. See illustration 3 below

Note: Results may be read up to 2 minutes after the specified times.



## INTERPRETATION OF RESULTS

Results are obtained by direct comparison of the color blocks printed on the canister label. The color blocks represent nominal values; actual values will vary close to the nominal values. In the event of unexpected or questionable results, the following steps are recommended: confirm that the strips have been tested within the expiration date printed on the canister label, compare results with known positive and negative controls and repeat the test using a new strip. If the problem persists, discontinue using the strip immediately and contact your local distributor.

### QUALITY CONTROL

For best results, performance of reagent strips should be confirmed by testing known positive and negative specimens/controls whenever a new test is performed, or whenever a new canister is first opened. Each laboratory should establish its own goals for adequate standards of performance

## LIMITATIONS

Note: The Urinalysis Reagent Strips (Urine) may be affected by substances that cause abnormal urine color such as drugs containing azo dyes (e.g. Pyridium<sup>®</sup>, Azo Gantrisin<sup>®</sup> Azo Gantanol®), nitrofurantoin (Microdantin®, Furadantin®), and riboflavin.8 The color development on the test pad may be masked or a color reaction may be produced that could be interpreted as false results.

Ascorbic acid: No interference is known

Glucose: The reagent area does not react with lactose, galactose, fructose or other metabolic substances, nor with reducing metabolites of drugs (e.g. salicylates and nalidixic acid). Sensitivity may be decreased in specimens with high specific gravity (>1.025) and with ascorbic acid concentrations of  $\geq$  25 mg/dL. High ketone levels ≥ 100 mg/dL may cause false negative results for specimens containing a small amount of glucose (50-100 mg/dL)

Bilirubin: Bilirubin is absent in normal urine, so any positive result, including a trace positive, indicates an underlying pathological condition and requires further investigation. Reactions may occur with urine containing large doses of chlorpromazine or rifampen that might be mistaken for positive bilirubin. The presence of bilirubin-derived bile pigments may mask the bilirubin reaction. This phenomenon is characterized by color development on the test patch that does not correlate with the colors on the color chart. Large concentrations of ascorbic acid may decrease sensitivity. **Ketone:** The test does not react with acetone or β-hydroxybutyrate. Urine specimens of high pigment, and other substances containing sulfhydryl groups may occasionally give reactions up to and including trace (±).9

Specific Gravity: Ketoacidosis or protein higher than 300 mg/dL may cause elevated results. Results are not affected by non-ionic urine components such as glucose. If the urine has a pH of 7 or greater, add 0.005 to the specific gravity reading indicated on the

Blood: A uniform blue color indicates the presence of myoglobin, hemoglobin or hemolyzed erythrocytes. Scattered or compacted blue spots indicate intact erythrocytes. To enhance accuracy, separate color scales are provided for hemoglobin and for erythrocytes. Positive results with this test are often seen with urine from menstruating females. It has been reported that urine of high pH reduces sensitivity, while moderate to

high concentration of ascorbic acid may inhibit color formation. Microbial peroxidase, associated with urinary tract infection, may cause a false positive reaction. The test is slightly more sensitive to free hemoglobin and myoglobin than to intact erythrocytes.

pH: If the procedure is not followed and excess urine remains on the strip, a phenomenon known as "runover" may occur, in which the acid buffer from the protein reagent will run onto the pH area, causing the pH result to appear artificially low. pH readings are not affected by variations in urinary buffer concentration.

Protein: Any green color indicates the presence of protein in the urine. This test is highly sensitive for albumin, and less sensitive to hemoglobin, globulin and mucoprotein.8 A negative result does not rule out the presence of these other proteins. False positive results may be obtained with highly buffered or alkaline urine. Contamination of urine specimens with quaternary ammonium compounds or skin cleansers containing chlorhexidine may produce false positive results.8 The urine specimens with high specific gravity may give false negative results.

Urobilinogen: All results lower than 1 mg/dL urobilinogen should be interpreted as normal. A negative result does not at any time preclude the absence of urobilinogen. The reagent area may react with interfering substances known to react with Ehrlich's reagent. such as p-aminosalicylic acid and sulfonamides. False negative results may be obtained if formalin is present. The test cannot be used to detect porphobilinogen.

Nitrite: The test is specific for nitrite and will not react with any other substance normally excreted in urine. Any degree of uniform pink to red color should be interpreted as a positive result, suggesting the presence of nitrite. Color intensity is not proportional to the number of bacteria present in the urine specimen. Pink spots or pink edges should not be interpreted as a positive result. Comparing the reacted reagent area on a white background may aid in the detection of low nitrite levels, which might otherwise be missed. Ascorbic acid above 30 mg/dL may cause false negatives in urine containing less than 0.05 mg/dL nitrite ions. The sensitivity of this test is reduced for urine specimens with highly buffered alkaline urine or with high specific gravity. A negative result does not at any time preclude the possibility of bacteruria. Negative results may occur in urinary tract infections from organisms that do not contain reductase to convert nitrate to nitrite; when urine has not been retained in the bladder for a sufficient length of time (at least 4 hours) for reduction of nitrate to nitrite to occur; when receiving antibiotic therapy or when dietary nitrate is absent.

Leukocytes: The result should be read between 60-120 seconds to allow for complete color development. The intensity of the color that develops is proportional to the number of leukocytes present in the urine specimen. High specific gravity or elevated glucose concentrations (≥ 2,000 mg/dL) may cause test results to be artificially low. The presence of cephalexin, cephalothin, or high concentrations of oxalic acid may also cause test results to be artificially low. Tetracycline may cause decreased reactivity, and high levels of the drug may cause a false negative reaction. High urinary protein may diminish the intensity of the reaction color. This test will not react with erythrocytes or bacteria common in urine

## BIBLIOGRAPHY

- Free AH. Free HM. Urinalysis, Critical Discipline of Clinical Science, CRC Crit Rev. Clin. Lab. Sci. 3(4): 481-531, 1972.
- Yoder J. Adams EC. Free. AH. Simultaneous Screening for Urinary Occult Blood. Protein, Glucose, and pH. Amer. J. Med Tech. 31:285, 1965.
- Shchersten B, Fritz H. Subnormal Levels of Glucose in Urine. JAMA 201:129-132 McGarry JD, Lilly. Lecture, 1978: New Perspectives in the Regulation of
- Ketogenesis. Diabetes 28: 517-523 May, 1978. Williamson DH. Physiological Ketoses, or Why Ketone Bodies? Postgrad. Med. J
- (June Suppl.): 372-375, 1971.
- Paterson P, et al. Maternal and Fetal Ketone Concentrations in Plasma and Urine Lancet: 862-865; April 22, 1967. Fraser J, et al. Studies with a Simplified Nitroprusside Test for Ketone Bodies in
- Urine, Serum, Plasma and Milk. Clin. Chem. Acta II: 372-378, 1965. Henry JB, et al. Clinical Diagnosis and Management by Laboratory Methods, 20th Ed.
- Philadelphia, Saunders, 371-372, 375, 379, 382, 385, 2001.
- Tietz NW. Clinical Guide to Laboratory Tests. W.B. Saunders Company. 1976.
- Burtis CA, Ashwood ER. Tietz Textbook of Clinical Chemistry 2<sup>nd</sup> Ed. 2205, 1994.

### Index of Symbols Consult instruction [i Tests per ki lanufacturer or use or in vitro IVD se by o not reuse liagnostic use only tore between LOT REF of Number Catalog # -30°C Authorized EC REP Representative





EC REP MDSS GmbH Schiffgraben 41 30175 Hannover, Germany

Number: 1150310404 Effective date: 2011-03-14



Contract No:Co2403079 Date:09/03/2024

## **Letter of Authorization**

Manufacturer: Atlas Medical GmbH

Ludwig-Erhard-Ring 3,

15827Blankenfelde-Mahlow, Germany

Tel: +49 33 70 83 55 030

Email: amug@atlas-medical.com

Regulatory Office: William James House, Cowley Road, Cambridge, CB4 0WX, UK

Tel: +44 1223 858 910 Fax: +44 1223 858 524 Email: info@atlas-site.co.uk

Middle East Site: Sahab Free Zone Area

P. O. Box 204, Amman 11512, Jordan.

Tel.: +962 6 4026468 Fax: +962 6 4022588

Email: info@atlas-medical.com

Agent: San Medico

Republic of Moldova, city Chisina

+37368228890

Atlas Medical, hereby appoint the above mentioned agent to import, register and distribute Atlas Medical Products in Maldova

**Appointment Conditions:** 

1. This appointment is valid for 3 year from the above mentioned date.

2. Either Party can cancel this appointment by giving the other party a 60 day notice.

On behalf of the Manufacturer General Manager

Haya Amawi





## CERTIFICAT

# CERTIFICATE OF REGISTRATION N° 36655 rev.2

## GMED certifie que le système de management de la qualité développé par

GMED certifies that the quality management system developed by

# ATLAS MEDICAL GmbH Ludwig-Erhard-Ring 3 15827 Blankenfelde-Mahlow GERMANY

pour les activités for the activities

Conception et développement, fabrication et vente de dispositifs médicaux de diagnostic in vitro .

Design and Development, Manufacturing and Sales of in vitro diagnostic medical devices.

réalisées sur le(s) site(s) de performed on the location(s) of

Voir addendum

See addendum

est conforme aux exigences des normes internationales complies with the requirements of the international standards

ISO 13485: 2016

Début de validité / Effective date October 9th, 2023 (included) Valable jusqu'au / Expiry date : October 8th, 2026 (included)

Etabli le / Issued on : October 9th, 2023



GMED N° 36655-2

Ce certificat est délivré selon les règles de certification GMED / This certificate is issued according to the rules of GMED certification

Renouvelle le certificat 36655-1

CERTIFICATION
DE SYSTEMES
DE MANAGEMENT
Accréditation n°4-0608
Let portée disponible su
www.cofrac.fr

GMED
SIÈGE SOC

**GMED** • Société par Actions Simplifiée au capital de 300 000 € • Organisme Notifié/Notified Body n° 0459 Siège social : 1, rue Gaston Boissier - 75015 Paris • Tél. : 01 40 43 37 00 • gmed.fr



Addendum au certificat n° 36655 rev. 2 page 1/1 Addendum of the certificate n° 36655 rev. 2 Dossier / File N°P606647

## Ce certificat couvre les activités et les sites suivants :

This certificate covers the following activities and sites:

## French version:

Conception et développement, fabrication et vente de dispositifs médicaux de diagnostic *in vitro* à usage professionnel et/ ou d'autodiagnostic, dans les domaines du groupage sanguin, de la microbiologie, de la biochimie, de la toxicologie, de l'oncologie, de la cardiologie, de l'histologie, de l'endocrinologie et des maladies infectieuses, dans les techniques d'Agglutination/ ELISA/ Tests rapides/ Colorimétrie/ Disques antibiotiques.

## English version:

Design and Development, Manufacturing and Sales of in vitro diagnostic medical devices for professional use and/or for self-testing, in the field of Immunohematology, Microbiology, Biochemistry, Toxicology, Oncology, Cardiology, Histology, Endocrinology Biosensors and Infectious diseases, in techniques of Agglutination/ELISA/Rapid tests/Colorimetry/Antibiotic disks.

ATLAS MEDICAL GmbH Ludwig-Erhard-Ring 3 15827 Blankenfelde-Mahlow GERMANY

French version:

Siège social, responsable de la mise sur le marché

English version:

Headquarter, legal manufacturer

\*\*\*\*\*\*\*\*\*\*\*\*\*

Sahab Industrial Zone Area King Abdullah II Industrial City Amman 11512 JORDAN

French version:

Conception, fabrication et contrôle final

English version:

Design, manufacture and final control

\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

2 sites / 2 sites

BEATIVE LYS

On behalf of the President Béatrice LYS Technical Director



Declaration Ref No: DC21-0249

Date: 15.10.2021

# **CE Declaration of Conformity**

Name and address of Manufacturer	Atlas Medical GmbH
	Ludwig-Erhard-Ring 3, 15827 Blankenefelde-Mahlow
	Germany .
	Tel: +49(0)33708355030
	Email: info@atlas-medical.com

Atlas Medical GmbH declared our his own responsibility that the following IVD medical devices:

Product Code	Product Name	GMDN code
8.17.003.0300	Atlas Periodic Acid Schiff (PAS) Stain Kit, 3x100ml	43587
8.17.004.0300	Atlas Iron Stain Kit, 3x100ml	43587
8.17.009.1000	Atlas Gram Stain Kit	43733
8.17.010.0750	Atlas ZN (Kinyoun) stain pack , 3x250ml	43587
8.15.144.0250	Atlas ZN Decolouriser, 250 ml /Bottle	43587
8.17.015.0500	Atlas Diff-3 Stain.	43587
8.17.016.1000	Atlas Papanicolau Stain Pack.	43587
8.17.110.0250	Atlas Papanicolau Stain EA35, 250 ml /Bottle.	43587
8.17.111.0250	Atlas Papanicolau Stain EA36, 250 ml /Bottle	43587
8.17.112.0250	Atlas Papanicolau Stain EA65, 250 ml /Bottle.	43587
8.17.114.0250	Atlas Papanicolau Stain EA50, 250 ml /Bottle.	43587
8.17.115.0250	Atlas Papanicolau Stain OG6, 250 ml /Bottle.	43587
8.17.014.1000	Atlas Reticulocytes stain (Methylene Blue) , 1000 ml /Bottle	43587
3.15.037.0250	Atlas Eosin Y (1%) Stain, 250 ml/Bottle	42507
3.15.038.0250	Atlas Eosin Y (5%) Stain, 250 ml/Bottle.	43587
3.15.041.0250	Atlas Field Stain (Solution A), 250ml/Bottle	43587
3.15.042.0250	Atlas Field Stain (Solution B), 250ml/Bottle	43587
.15.043.0750	Atlas Field Stain Kit 3x250ml (250ml Fixing Reagent,	43587
	250ml Eosin Reagent, 250ml Methylene Blue Reagent).	43587
.15.047.0250	Atlas Giemsa Stain, 250 ml/Bottle.	43587
15.059.0250	Atlas Haematoxylin Harris Stain , 250 ml/Bottle	43587
15.069.0250	Atlas Leishman Stain , 250 ml/Bottle.	43587
15.069.1000	Atlas Leishman Stain , 1000 ml/Bottle.	43587
.15.074.0250	Atlas Lugol's Iodine, 250 ml/Bottle.	43587
15.078.0250	Atlas May Grunwald Stain, 250 ml/Bottle.	43587
15.105.0250	Atlas New Methylene Blue for Reticulocytes, 250 ml/Bottle.	43587
.15.143.0250	Atlas Wright's Stain, 250 ml/Bottle.	42507
15.146.0100	Atlas Immersion oil, 100 Bottle/Box	43587
	- TO Source Box	43587



Declaration Ref No: DC21-0249

Date: 15.10.2021

Meets the essential requirments of In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex I

EN ISO 13485 :2016, EN 18113-1, -2,:2011, EN ISO 15223:2016 EN ISO 14971:2019, EN ISO 23640:2015, ISO 2859/1:1999, EN ISO 13612:2002, EN ISO 13641:2002, EN ISO 62366-1+A1:2020.

	Directive 98/79, Other IVDs (Non-annex II, non-self-test).
Conformity Assesment Route	Directive 98/79/EC , Annex III.
Name , Address and Identification number of notified body	N/A

Date of issuance:	15. October.2021
Place	Atlas Medical GmbH
Signed by:	Amani AL-Habahbeh
Position:	Ame
	Regulatory Affairs Manager

Atlas Medical GmbH

Ludwig - Erhard Ring 3

Ludwig - Blankenfelde - Mahlow

15827 Blankenfelde - 355030

Tel. (0049) 33708 - 355030



## **GRAM STAIN PACK**

IVD For in -vitro diagnostic and professional use only



## **INTENDED USE**

Gram Stain used for differentiate between gram positive and gramnegative bacteria.

## INTRODUCTION

Gram staining is used to differentiate bacterial species into two large groups (Gram-positive and Gram-negative) based on the physical properties of their cell walls.

## **PRINCIPLE**

Gram-positive bacteria have a thick mesh-like cell wall made of peptidoglycan (50-90% of cell wall), which stains Blue while gramnegative bacteria have a thinner layer (10% of cell wall), which stains pink. Gram-negative bacteria also have an additional outer membrane which contains lipids, and is separated from the cell wall by the periplasmic space. There are four basic steps of the Gram stain, which include applying a primary stain (crystal violet) to a heat-fixed smear of a bacterial culture, followed by the addition of a trapping agent (Gram's iodine), rapid decolorization with alcohol or acetone, and counterstaining with safranin or basic fuchsin.

Crystal violet (CV) dissociates in aqueous solutions into CV+ and chloride (Cl -) ions. These ions penetrate through the cell wall and cell membrane of both gram-positive and gram-negative cells. The CV+ ion interacts with negatively charged components of bacterial cells and stains the cells Blue.

Iodine (I - or I<sub>3</sub> -) interacts with CV+ and forms large complexes of crystal violet and iodine (CV-I) within the inner and outer layers of the cell. Iodine is often referred to as a mordant, but is a trapping agent that prevents the removal of the CV-I complex and therefore color from the cell.

When a decolorizer such as alcohol or acetone is added, it interacts with the lipids of the cell membrane. A gram-negative cell will lose its outer membrane and the lipopolysaccharide layer is left exposed. The

CV-I complexes are washed from the gram-negative cell along with the outer membrane. In contrast, a gram-positive cell becomes dehydrated from an ethanol treatment. The large CV-I complexes become trapped within the gram-positive cell due to the multilayered nature of its peptidoglycan. The decolorization step is critical and must be timed correctly; the crystal violet stain will be removed from both gram-positive and negative cells if the decolorizing agent is left on too long (a matter of seconds).

After decolorization, the gram-positive cell remains Blue. and the gram-negative cell loses its Blue. color. Counterstain, which is usually positively charged safranin or basic fuchsin, is applied last to give decolorized gram-negative bacteria a pink or red color.

## **MATERIALS**

## **MATERIALS PROVIDED**

- Crystal Violet.
- Gram Iodine.
- Gram Decolouriser.
- Counterstain Safranin O.

Note: This package insert is also used for individually packed reagent.

## Storage and stability

- Store at room temperature.
- Stain Solution is stable up to the printed expiry date.
- Keep the bottles tightly closed to prevent air oxidation.

## **Precautions**

- The reagent may cause eye, skin and respiratory tract irritation; so protective clothing should be worn when handling this reagent.
- The reagent is intended for in vitro diagnostic use only.
- Do not use this reagent if the label is not available or
- Test materials and samples should be discarded properly in biohazards container.
- This reagent is considered toxic, so do not drink or eat beside it.
- Wash hands and test table top with water and soap once the testing is done.

## **PROCEDURE**

- 1. immerse the heat fixed smears with Crystal Violet and allow to stain for up to 1 minute.
- 2. Wash with tap water.
- 3. Flood the smear with Gram Iodine for 2 minutes.
- 4. Wash with tap water.
- 5. Decolorize the smear for few second only.
- 6. Wash thoroughly with tap water.
- 7. Counterstain with Safranin O for up to 2 minutes.
- 8. Wash and allow to dry.
- 9. Examine under microscope using oil immersion objective

## **RESULTS**

- Gram positive organisms (Blue).
- Gram negative organisms (Red).

ATLAS Medical Ludwig-Erhard Ring 3 15827 Blankenfelde-Mahlow Germany

Tel: +49 - 33708 - 3550 30 Email: Info@atlas-medical.com Website: www.atlas-medical.com

## PPI2112A01 Rev B (08.10.2020)

REF	Catalogue Number	1	Temperature limit
IVD	<i>In Vitro</i> diagnostic medical device	$\triangle$	Caution
Σ	Contains sufficient for <n> tests and Relative size</n>		Consult instructions for use (IFU)
LOT	Batch code		Manufacturer
Ī	Fragile, handle with care		Use-by date
1	Manufacturer fax number	( <u>(</u>	Do not use if package is damaged
<b>3</b>	Manufacturer telephone number	3	Date of Manufacture
类	Keep away from sunlight	学	Keep dry
<b>®</b>	Flammable		



## 浙江东方基因生物制品股份有限公司 Zhejiang Orient Gene Biotech Co.,LTD

## **STATEMENT**

We, Zhejiang Orient Gene Biotech Co., Ltd , having a registered office at 3787#, East Yangguang Avenue, Dipu Street Anji 313300, Huzhou, Zhejiang, China assign SRL SANMEDICO having a registered office at A. Corobceanu street 7A, apt. 9, Chişinău MD-2012, Moldova, as non-exclusive authorized representative for Orient Gene Brand product in correspondence with the conditions of directive 98/79/EEC.

We declare that the company mentioned above is authorized to register, notify, renew or modify the registration of medical devices on the territory of the Republic of Moldova.

This Statement letter will be valid from Feb.21th,2023 to Feb.20th, 2024.

Zhejiang Orient Gene Biotech

General Manager

Date: 2023/2/21

电话 Tel:+86-572-5226111







## **Product Service**

# **Certificate**

No. Q5 092305 0001 Rev. 01

**Holder of Certificate: Zhejiang Orient Gene Biotech Co., Ltd.** 

3787#, East Yangguang Avenue, Dipu Street Anji

313300 Huzhou, Zhejiang

PEOPLE'S REPUBLIC OF CHINA

**Certification Mark:** 



Design and Development, Production and Distribution Scope of Certificate:

of In Vitro Diagnostic Reagent and Instrument for the **Detection of Drugs of Abuse, Fertility, Infectious** Diseases, Oncology, Biochemistry, Cardiac Diseases, Allergic Disease based on Rapid Test, PCR and Liquid

**Biochip Method.** 

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:Q5 092305 0001 Rev. 01

Report No.: SH2198802

Valid from: 2022-04-11 Valid until: 2024-03-16

Christoph Dicks 2022-04-11 Date,

Head of Certification/Notified Body



# **Certificate**

No. Q5 092305 0001 Rev. 01

**Applied Standard(s):** EN ISO 13485:2016

Medical devices - Quality management systems -

Requirements for regulatory purposes

(ISO 13485:2016) DIN EN ISO 13485:2016

Facility(ies): Zhejiang Orient Gene Biotech Co., Ltd.

3787#, East Yangguang Avenue, Dipu Street Anji, 313300 Huzhou, Zhejiang, PEOPLE'S REPUBLIC OF CHINA

See Scope of Certificate





## 浙江东方基因生物制品股份有限公司 Zhejiang Orient Gene Biotech Co., LTD



CE-DOC-OG038 Version 2.0

# **EC** Declaration of Conformity

In accordance with Directive 98/79/EC

Legal Manufacturer: Zhejiang Orient Gene Biotech Co., Ltd

**Legal Manufacturer Address:** 3787#, East Yangguang Avenue, Dipu Street,

Anji 313300, Huzhou, Zhejiang, China

Declares, that the products Product Name and Model(s)

Troponin I Rapid Test Cassette (Whole Blood/Serum/Plasma)	GDTRO-402a
Troponin I Rapid Test Cassette (Whole Blood/Serum/Plasma)	GDTRO-402b

Classification: Other

Conformity assessment route: Annex III (EC DECLARATION OF CONFORMITY)

We, the Manufacturer, herewith declare with sole responsibility that our product/s mentioned above meet/s the provisions of the Directive 98/79/EC of the European Parliament and of the Council on In-Vitro Diagnostic Medical Devices.

We hereby explicitly appoint

**EC Representative's Name:** Shanghai International Holding Corp. GmbH (Europe)

**EC Representative's Address:** Eiffestrasse 80, 20537 Hamburg, Germany

to act as our European Authorized Representative as defined in the aforementioned Directive.

I, the undersigned, hereby declare that the medical devices specified above conform with the directive 98/79/EC on in vitro diagnostic medical devices and pertinent essential requirements

Date Signed: August 11, 2020

Name of authorized signatory: Joyce Pang Position held in the company: Vice-President

Tyle Py.











Product Description	Format	Cut-off Value	Qualification
Acetaminophen (ACE) Test	Strip/Cassette/Dip Card/Cup	5000 ng/mL	CE
Amphetamine (AMP) Test	Strip/Cassette/Dip Card/Cup	2000/1000/500/300/250 ng/mL	CE 510(k)
Barbiturates (BAR) Test	Strip/Cassette/Dip Card/Cup	2000/600/300/200 ng/mL	CE 510(k)
Benzodiazepines (BZO) Test	Strip/Cassette/Dip Card/Cup	600/400/300/200/100 ng/mL	CE 510(k)
Buprenorphine (BUP) Test	Strip/Cassette/Dip Card/Cup	10/5 ng/mL	CE 510(k)
Caffeine (CAF) Test	Strip/Cassette/Dip Card/Cup	6000 ng/mL	/
Carisoprodol (SOMA) Test	Strip/Cassette/Dip Card/Cup	1000 ng/mL	CE
Clonazepam (CLO) Test	Strip/Cassette/Dip Card/Cup	500/100 ng/mL	CE
Cocaine (COC) Test	Strip/Cassette/Dip Card/Cup	600/300/150/100 ng/mL	CE 510(k)
Codeine (COD) Test	Strip/Cassette/Dip Card/Cup	2000 ng/mL	CE
Cotinine (COT) Test	Strip/Cassette/Dip Card/Cup	400/300/200/100/50 ng/mL	CE
Ecstasy (MDMA) Test	Strip/Cassette/Dip Card/Cup	2000/1000/500/300/250/150 ng/mL	CE 510(k)
Ethyl Glucuronide (EtG) Test	Strip/Cassette/Dip Card/Cup	500/300ng/mL	CE
Fentanyl (FEN) Test	Strip/Cassette/Dip Card/Cup	300/200/100/50 ng/mL	CE
Norfentanyl (FEN) Test	Strip/Cassette/Dip Card/Cup	200/50/20/10/5 ng/mL	CE
Gabapentin (GAB) Test	Strip/Cassette/Dip Card/Cup	3750/2000/1000 ng/mL	CE
Hydrocodone (HCD) Test	Strip/Cassette/Dip Card/Cup	300/10 ng/mL	CE
Hydromorphone (HMO) Test	Strip/Cassette/Dip Card/Cup	300 ng/mL	CE
Ketamine (KET) Test	Strip/Cassette/Dip Card/Cup	3000/2000/1000/500/100 ng/mL	CE
Kratom (KRA) Test	Strip/Cassette/Dip Card/Cup	250/150/100 ng/mL	CE
ysergic acid diethylamide (LSD) Test	Strip/Cassette/Dip Card/Cup	20 ng/mL	CE
Marijuana (THC) Test	Strip/Cassette/Dip Card/Cup	600/300/200/150/100/50/40/25/20/18/15 ng/mL	CE 510(k)
Methadone Metabolite (EDDP) Test	Strip/Cassette/Dip Card/Cup	300/100 ng/mL	CE 510(k)
Methadone (MTD) Test	Strip/Cassette/Dip Card/Cup	1000/600/300/200/50 ng/mL	CE 510(k)
Methamphetamine (MET) Test	Strip/Cassette/Dip Card/Cup	2000/1000/500/300/250 ng/mL	CE 510(k)
Methaqualone (MQL) Test	Strip/Cassette/Dip Card/Cup	300/1000 ng/mL	CE
Methcathinone (MTC) Test	Strip/Cassette/Dip Card/Cup	500/300 ng/mL	CE
3,4-Methylenedioxypyrovalerone (MDPV) Test	Strip/Cassette/Dip Card/Cup	1000/500/300 ng/mL	CE
Methylphenidate (MPD) Test	Strip/Cassette/Dip Card/Cup	300 ng/mL	CE
6-Monoacetylmorphine (6-MAM) Test	Strip/Cassette/Dip Card/Cup	20/10 ng/mL	CE
Morphine (MOP) Test	Strip/Cassette/Dip Card/Cup	2000/600/300/150/100 ng/mL	CE 510(k)
Opiate (OPI) Test	Strip/Cassette/Dip Card/Cup	2000/300/100 ng/mL	CE 510(k)
Dxycodone (OXY) Test	Strip/Cassette/Dip Card/Cup	300/100 ng/mL	CE 510(k)
Phencyclidine (PCP) Test	Strip/Cassette/Dip Card/Cup	50/25 ng/mL	CE 510(k)
Pinaca Ab (K3) Test	Strip/Cassette/Dip Card/Cup	10 ng/mL	CE
Pregabalin (PGB) Test	Strip/Cassette/Dip Card/Cup	2000/1000/500 ng/mL	CE
Propoxyphene (PPX) Test	Strip/Cassette/Dip Card/Cup	600/300 ng/mL	CE 510(k)
Synthetic Marijuana (K2) Test	Strip/Cassette/Dip Card/Cup	75/50/25/20/10 ng/mL	CE
Framadol (TRA) Test	Strip/Cassette/Dip Card/Cup	200/100 ng/mL	CE
Fricyclic Antidepressants (TCA) Test	Strip/Cassette/Dip Card/Cup	1000/300 ng/mL	CE 510(k)
JR-144 Test	Strip/Cassette/Dip Card/Cup	50 ng/mL	CE
Zolpidem (ZOL) Test New	Strip/Cassette/Dip Card/Cup	50 ng/mL	/
Zopiclone (ZOP) Test New	Strip/Cassette/Dip Card/Cup	50 ng/mL	/
·			









U	

<b>Product Description</b>	Format	Cut-off Value	Qualification
7-Aminonclnozepam (ACL) Test New	Device	100 ng/mL	/
Amphetamine (AMP) Test	Device	50/40 ng/mL	CE
Barbiturates (BAR) Test	Device	300/50/30 ng/mL	CE
Benzodiazepines (BZO) Test	Device	50/20/10 ng/mL	CE
Buprenorphine (BUP) Test	Device	10/5 ng/mL	CE
Carisoprodol (SOMA) Test	Device	300 ng/mL	/
Cocaine (COC) Test	Device	50/20 ng/mL	CE
Codeine (COD) Test	Device	10 ng/mL	CE
Cotinine (COT) Test	Device	50/30/10 ng/mL	CE
Diphenhydramine (DIP) Test New	Device	150/100 ng/mL	/
Ecstasy (MDMA) Test	Device	60/50 ng/mL	CE
Ethyl Glucuronide (EtG) Test New	Device	150/100 ng/mL	/
Fentanyl (FEN) Test	Device	10 ng/mL	CE
Hydrocodone (HCD) Test New	Device	10 ng/mL	/
Hydromorphone (HMO) Test New	Device	300/150 ng/mL	/
Ketamine (KET) Test	Device	100/50 ng/mL	CE
Lysergic acid diethylamide (LSD) Test	Device	25/10 ng/mL	CE
Marijuana (THC) Test	Device	50/40/30/25/15/12/10/5/4/3 ng/mL	CE
Methadone Metabolite (EDDP) Test	Device	20 ng/mL	CE
Mephedrone (MEP) Test New	Device	50 ng/mL	1
Methadone (MTD) Test	Device	75/50/30 ng/mL	CE

Methamphetamine (MET) Test	Device	50 ng/mL	CE
Methaqualone (MQL) Test	Device	150/100 ng/mL	CE
Methcathinone (MTC) Test	Device	50 ng/mL	/
3,4-Methylenedioxypyrovalerone (MDPV) Test	Device	200/100/50 ng/mL	CE
Methylphenidate (MPD) Test	Device	50 ng/mL	/
6-Monoacetylmorphine (6-MAM) Test	Device	25/15/10/5/4 ng/mL	CE
Morphine (MOP) Test	Device	15 ng/mL	CE
Opiate (OPI) Test	Device	50/40 ng/mL	CE
Oxycodone (OXY) Test	Device	50/40/20 ng/mL	CE
Phencyclidine (PCP) Test	Device	10 ng/mL	CE
Phenytoin (PHEN) Test New	Device	150/100 ng/mL	/
Pinaca Ab (K3) Test New	Device	10 ng/mL	/
Pregabalin (PGB) Test New	Device	100 ng/mL	/
Propoxyphene (PPX) Test	Device	50/20 ng/mL	CE
Synthetic Marijuana (K2) Test	Device	25/10/5 ng/mL	CE
Tramadol (TRA) Test	Device	100/50 ng/mL	CE
Tricyclic Antidepressants (TCA) Test	Device	100 ng/mL	CE
XLR-11 Test New	Device	100 ng/mL	/
Zolpidem (ZOL) Test New	Device	25 ng/mL	/
Zopiclone (ZOP) Test New	Device	25 ng/mL	/
Alcohol (ALC) Test	Device	0.05/0.02%	CE

# Toxicology Hair Test





<b>Product Description</b>	Format	Label	Cut-off Value	Qualification
Amphetamine (AMP) Test	Cassette	Fluorescence	0.5/0.2 ng/mg	CE
Amphetamine (AMP) Test	Cassette	Gold	5 ng/mg	/
Benzodiazepines (BZO) Test	Cassette	Fluorescence	0.2 ng/mg	/
berizodiazepiries (bzO) Test	Cassette	Gold	1 ng/mg	/
Cocaine (COC) Test	Cassette	Fluorescence	0.5/0.2 ng/mg	CE
Cocame (COC) Test	Cassette	Gold	5/2 ng/mg	CE
Ecstasy (MDMA) Test	Cassette	Fluorescence	0.2 ng/mg	CE
Lestady (MIDIMIN) Test	Casselle	Gold	5 ng/mg	/
2-Fluorodeschloroketamin (FKE) Test	Cassette	Fluorescence	0.2 ng/mg	/
Ketamine (KET) Test	Cassette	Fluorescence	0.2 ng/mg	CE
Returnine (RET) Test	Gussette	Gold	2/1/0.5 ng/mg	CE
Marijuana (THC) Test	Cassette	Fluorescence	0.05 ng/mg	CE
Manjaana (1110) Test	Cussotto	Gold	2/1.5 ng/mg	CE
Methamphetamine (MET) Test	Cassette	Fluorescence	0.5/0.2 ng/mg	CE
,	Custotic	Gold	5/2/1 ng/mg	CE
Methcathinone (MTC) Test	Cassette	Fluorescence	0.2 ng/mg	CE
6-Monoacetylmorphine (6-MAM) Test	Cassette	Fluorescence	0.2 ng/mg	CE
o worlddooty arios prairio (o wir awr, 100t	Custotic	Gold	2 ng/mg	CE
Morphine (MOP) Test	Cassette	Fluorescence	0.2 ng/mg	CE
wierprinie (wier / rest	Cussotto	Gold	5/2/0.5 ng/mg	CE
Oxycodone (OXY) Test	Cassette	Fluorescence	0.2 ng/mg	CE
onycouchio (ent.) tool	Cuccotto	Gold	4 ng/mg	/
Phencyclidine (PCP) Test	Cassette	Fluorescence	0.3 ng/mg	CE
. Heriofelianie (Ferri Foot	Guestie	Gold	1 ng/mg	CE
Pinaca Ab (K3) Test	Cassette	Fluorescence	0.2 ng/mg	CE
	22230110	Gold	0.5 ng/mg	/
Synthetic Marijuana (K2) Test	Cassette	Fluorescence	0.2 ng/mg	CE
•		Gold	1 ng/mg	/
Tramadol (TRA) Test	Cassette	Fluorescence	0.2 ng/mg	/
LIR-144 Test	Cassette	Fluorescence	0.05 na/ma	/





oduct Description	Specimen	Catalog No.	Format	Cut-off Value	Kit Size
enovirus Antigen Test	Swab	GCADE-502a√	Cassette	/	20 Tests/Kit
enovirus Test	Feces	GCADE-602a√	Cassette	/	20 Tests/Kit
icella Antibody Test	WB/S/P	GCBRU-402a√	Cassette	/	25 Tests/Kit
ndida albicans Test	Vaginal Secretion	GCCA-502a√	Cassette	10 <sup>5</sup> CFU/mL	20 Tests/Kit
	S/P	GCCHA-302a√	Cassette	/	25 Tests/Kit
agas Antibody Test -	WB/S/P	GCCHA-402a√	Cassette	/	25 Tests/Kit
ostridium difficile GDH Test	Feces	GCCD(GDH)-602a√	Cassette	2 ng/mL	20 Tests/Kit
ostridium difficile Toxin A/B Test	Feces	GCCD(Toxin A/B)-602a√	Cassette	Toxin A: 2 ng/mL Toxin B: 2 ng/mL	20 Tests/Kit
ostridium difficile GDH & kin A/B Combo Test	Feces	GCCD-625a√	Cassette	GDH: 2 ng/mL Toxin A: 2 ng/mL Toxin B: 2 ng/mL	20 Tests/Kit
ikungunya IgM Test -	S/P	GCCHK(IgM)-302a√	Cassette	/	25 Tests/Kit
ikuriguriya igivi rest	WB/S/P	GCCHK(IgM)-402a√	Cassette	/	25 Tests/Kit

nikungunya IgG/IgM Test	WB/S/P	GCCHK(lgG/lgM)-402a	Cassette	/	25 Tests/Kit
nlamydia Test	Swab/Urine	GCCHL-502a√	Cassette	4.8×10 <sup>3</sup> IFU/mL	20 Tests/Kit
MV IgG Test	S/P	GCCMV(IgG)-302a	Cassette	/	25 Tests/Kit
ii iga rost	WB/S/P	GCCMV(IgG)-402a	Cassette	/	25 Tests/Kit
MV IgM Test —	S/P	GCCMV(IgM)-302a	Cassette	/	25 Tests/Kit
•	WB/S/P	GCCMV(IgM)-402a	Cassette	/	25 Tests/Kit
MV lgG/lgM Test —	S/P	GCCMV(lgG/lgM)-302a	Cassette	1	25 Tests/Kit
	WB/S/P	GCCMV(lgG/lgM)-402a	Cassette	/	25 Tests/Kit
OVID-19 IgM/IgG Test	WB/S/P	GCCOV(NAb) 402b /	Cassette Cassette	/	25 Tests/Kit
OVID-19 Neutralizing Antibody Test	WB/S/P	GCCOV(NAb)-402b√	Cassette		25 Tests/Kit
	Nasopharyngeal Swab	GCCOV-502a√ GCCOV-502Ca√	Cassette	/	20 Tests/Kit 20 Tests/Kit
_		GCCOV-502Ca√	Strip	/	20 Tests/Kit
DVID-19 Antigen Test	Nasal Swab	GCCOV-501a√	Cassette	/	1/2/3/5/7/10/15/20 Test(s)/k
JVID-19 Altitigen Test	ivasat Swab	GCCOV-502a-NAV	Device	/	1/2/5/10 Tests/Kit
_	NA & NP Swab	GCCOV-502a-NN√	Cassette	/	20 Tests/Kit
_	Oral Fluid	GCCOV-702a√	Cassette	/	20 Tests/Kit
		GCCOV-502a-Hxx√	Cassette	/	1/2/3/5/7/10/15/20 Test(s)/k
OVID-19 Antigen Self-Test	Nasal Swab	GCCOV-502a-HxxOGE√	Cassette	/	1/2/3/5/7/8/10/15/20/25 Test(s)/k
_	Oral Fluid	GCCOV-702a-Hxx√New	Cassette	/	1/2/3/5/7/10/15/20 Test(s)/k
gital COVID-19 Antigen Test	Nasal Swab	GCCOV-D503a√ New	Reader	/	1/2/3/5/7/10/15/20 Test(s)/k
OVID-19 Antigen & B.1.1.7 Mutant Strain Combo Test	Nasal Swab	GCCOV(B117)-525a√	Cassette	/	20 Tests/Kit
OVID-19/Flu A&B /RSV Antigen Combo Test	Nasal Swab	GCFCR-T525a√ New	Cassette	/	20 Tests/Kit
RS-CoV-2 Delta-series Mutant Strain Antigen Test	Nasal Swab	GCCOV(Del)-T502a√	Cassette	/	20 Tests/Kit
RS-CoV-2 Ag Fluorescence Rapid Test	Nasal Swab	FCCOV-502a√ New	Cassette	1	20 Tests/Kit
engue IgG/IgM Antibody Test	WB/S/P	GCDEN(ab)-402c√	Cassette	1	25 Tests/Kit
engue NS 1 Antigen Test	WB/S/P	GCDEN(NS)-402c√	Cassette	/	25 Tests/Kit
engue NS1 & IgG/IgM Combo Test	WB/S/P	GCDEN-425a√	Cassette		20 Tests/Kit
71 IgM Test	S/P	GCEV71(IgM)-302a√	Cassette	/	25 Tests/Kit
<u> </u>	WB/S/P	GCEV71(IgM)-402a√	Cassette	/	25 Tests/Kit
ardia lamblia Test	Feces	GCGIA-602a√	Cassette	/	20 Tests/Kit
onorrhoeae Test	Swab	GCGON-502b	Cassette	1.0E*7	20 Tests/Kit
AV IgM Test	S/P	GCHAV(IgM)-302Ba√	Cassette	/	25 Tests/Kit
NV IgG/IgM Test	WB/S/P	GCHAV(IgG/IgM)-402a√	Cassette	/	25 Tests/Kit
NV AntigenTest	Feces	GCHAV-602a√	Cassette	1	25 Tests/Kit
	S/P	GCHBcb-302a	Cassette	2 NCU	25 Tests/Kit
BcAb Hepatitis B Core Antibody Test		GCHBcb-302b	Cassette	8 NCU	25 Tests/Kit
	WB/S/P	GCHBcb-402a	Cassette	2 NCU	25 Tests/Kit
	S/P	GCHBeb-302a	Cassette	2 NCU	25 Tests/Kit
BeAb Hepatitis B Envelope Antibody Test		GCHBeb-302b	Cassette	8 NCU	25 Tests/Kit
	WB/S/P	GCHBeb-402a	Cassette	2 NCU	25 Tests/Kit
BeAg Hepatitis B Envelope Antigen Test	S/P	GCHBeg-302a	Cassette	0.5 NCU	25 Tests/Kit
	WB/S/P	GCHBeg-402a	Cassette	0.5 NCU	25 Tests/Kit
	S/P	GCHBsb-301a	Strip	30 mIU/mL	50 Tests/Kit
		GCHBsb-302a	Cassette	30 mIU/mL	25 Tests/Kit
BsAb Hepatitis B Surface Antibody Test		GCHBsb-401a	Strip	30 mIU/mL	50 Tests/Kit
	WB/S/P	GCHBsb-402a	Cassette	30 mIU/mL	25 Tests/Kit
		GCHBsb-402b	Cassette	20 mIU/mL	25 Tests/Kit
	S/P	GCHBsg-301a	Strip	1 ng/mL	50 Tests/Kit
BsAg Hepatitis B Surface Antigen Rapid Test —		GCHBsg-302a	Cassette	1 ng/mL	25 Tests/Kit
Constitution of the second of	WB/S/P	GCHBsg-401a	Strip	1 ng/mL	50 Tests/Kit
		GCHBsg-402a	Cassette	1 ng/mL	25 Tests/Kit
BsAg/HCV Combo Test	WB/S/P	GCHBC-402a	Cassette	/	25 Tests/Kit
BsAg/HCV/HIV/Syphilis Combo Test	S/P	GCHBCISY-345a	Cassette	1	20 Tests/Kit
•	WB/S/P	GCHBCISY-445a	Cassette	/	20 Tests/Kit
BY HBcAb/HBeAb/HBeAg/HBsAb	S/P	GCHBV-355a	Cassette	/	20 Tests/Kit
BsAg Combo Test	WB/S/P	GCHBV-455a	Cassette	/	20 Tests/Kit
	S/P	GCHCV-301a	Strip	1	50 Tests/Kit
CV Hepatitis C Virus Test —		GCHCV-302a√	Cassette		25 Tests/Kit
•	WB/S/P	GCHCV-401a	Strip	/	50 Tests/Kit
		GCHCV-402a√	Cassette	/	25 Tests/Kit
CV/HIV Combo Test	WB/S/P	GCHCI-402a	Cassette		25 Tests/Kit
V Hepatitis E Virus IgM Test	S/P	GCHEV-302a√	Cassette	/	25 Tests/Kit
	S/P	GCHIV-301a	Strip	1	50 Tests/Kit
V 1/2 Antibody Test		GCHIV-302a√	Cassette	1	25 Tests/Kit
,	WB/S/P	GCHIV-401a	Strip	1	50 Tests/Kit
		GCHIV-402a√	Cassette		25 Tests/Kit
V 1/2 Antibody Tri-line Test	WB/S/P	GCHIV-GT402a	Cassette	/	25 Tests/Kit
V 1/2/O Antibody Test	S/P	GCHIV-T302b	Cassette	/	25 Tests/Kit
·	WB/S/P	GCHIV-T402a	Cassette	/	25 Tests/Kit
V Antigen/Antibody Combo Test	WB/S/P	GCHIV(Ag/Ab)-402a	Cassette	/	25 Tests/Kit
SV IgG Test —	S/P	GCHSV(IgG)-302a√	Cassette	1	25 Tests/Kit
J	WB/S/P	GCHSV(IgG)-402a√	Cassette	1	25 Tests/Kit
SV IgM Test —	S/P	GCHSV(IgM)-302a√	Cassette	/	25 Tests/Kit
<u> </u>	WB/S/P	GCHSV(IgM)-402a√	Cassette	/	25 Tests/Kit
SV IgG/IgM Test	S/P	GCHSV(IgG/IgM)-302a	Cassette	1	25 Tests/Kit
	WB/S/P	GCHSV(lgG/lgM)-402a	Cassette	/	25 Tests/Kit
	S/P	GCHP-301a√	Strip	1	50 Tests/Kit
pylori Antibody Test —	O/ F	GCHP-302a√	Cassette	1	25 Tests/Kit
pyton milibody rost —	W/R/S/D	GCHP-401a√	Strip	1	50 Tests/Kit
	WB/S/P	GCHP-402a√	Cassette	/	25 Tests/Kit

		GCHP-601a√	Strip	/	25 Tests/Kit
II mulari Antinan Tast	F	GCHP-601Ca√	Strip	/	25 Tests/Kit
H. pylori Antigen Test	Feces	GCHP-602a√	Cassette	/	20 Tests/Kit
		GCHP-602Ca√	Cassette	/	20 Tests/Kit
		GCFLU(A)-501a√	Strip	1.5 x 10 <sup>4</sup> TCID <sub>50</sub>	25 Tests/Kit
Influenza A Antigen Test	Nasal/Throat Swabs	GCFLU(A)-502a√	Cassette	1.5 x 10 <sup>4</sup> TCID <sub>50</sub>	20 Tests/Kit
		GCI LO(A)-302ay	Cassette		20 Tests/Kit
		GCFLU(A/B)-501a√	Strip	1.5x 10⁴ TCID <sub>50</sub> / 1.5 x 10⁵ TCID <sub>50</sub>	25 Tests/Kit
Influenza A/B Antigen Test	Nasal/Throat Swabs	GCFLU(A/B)-502a√	Cassette	1.5x 10 <sup>4</sup> TCID <sub>50</sub> / 1.5 x 10 <sup>5</sup> TCID <sub>50</sub>	20 Tests/Kit
		GCFLU(A/B)-502Ca√	Cassette	1.5x 10 <sup>4</sup> TCID <sub>50</sub> / 1.5 x 10 <sup>5</sup> TCID <sub>50</sub>	20 Tests/Kit
	Necesban massl Curch	CCCC ESE	Cassatta	/	20 Tooto/Vit
_	Nasopharyngeal Swab	GCFC-525a√	Cassette		20 Tests/Kit
	NA & NP Swab	GCFC-525a-NN√	Cassette	/	20 Tests/Kit
Influenza & COVID-19 Antigen Combo Test		GCFC-525a-NA√	Cassette	/	20 Tests/Kit
	Nasal Swab	GCFC-T502a√ New	Cassette		1/5/20 Tests/Kit
		GCFC-T503a√ <sup>New</sup>	Device	/	1/2/5/10 Test(s)/Kit
Flu, COVID-19, RSV & Adeno Antigen Combo Test —	Nasopharyngeal Swab	GCFCRA-545a√	Cassette		20 Tests/Kit
rta, oovid 13, nov a raciio rinigen combo rest	Nasal Swab	GCFCRA-T525a√ New	Cassette	/	20 Tests/Kit
	S/P	GCKal-301a	Strip	/	50 Tests/Kit
	5/P	GCKal-302a	Cassette	/	25 Tests/Kit
Leishmania Antibody Test		GCKal-401a√	Strip	/	50 Tests/Kit
, , , , , , , , , , , , , , , , , , , ,	WB/S/P	GCKal-402a	Cassette	/	25 Tests/Kit
		GCKal-T402a√	Cassette	/	25 Tests/Kit
Malaria Pan Antigen Test	Whole Blood	GCMAL(pan)-402a√	Cassette	200 parasites	25 Tests/Kit
•	Whole Blood	GCMAL(pf)-402a√	Cassette	200 parasites 200 parasites	25 Tests/Kit
Malaria P.f. Antigen Test		4			
Malaria P.f./Pan Antigen Test	Whole Blood	GCMAL(pf/pan)-402a√	Cassette	200 parasites	25 Tests/Kit
Malaria P.f./P.v. Antigen Test	Whole Blood	GCMAL(pf/pv)-402a√	Cassette	200 parasites	25 Tests/Kit
Malaria P.f./P.v. Antibody Test	S/P	GCMAL(pf/pv Ab)-302a√	Cassette		25 Tests/Kit
matana i miji ivi i mabbay i bot	WB/S/P	GCMAL(pf/pv Ab)-402a√	Cassette	/	25 Tests/Kit
Monkeypox IgG/IgM Antibody Test	WB/S/P	GCMKP-402a√ New	Cassette	/	25 Tests/Kit
Monkeypox Antigen Test	WB/S/P or Throat swab	GCMKP-502a√ New	Cassette	/	25 Tests/Kit
	S/P	GCMON-325a√	Cassette	/	25 Tests/Kit
Mononucleosis Test		GCMON-402a√	Cassette	/	25 Tests/Kit
	WB/S/P	GCMON-425a√	Cassette	/	25 Tests/Kit
M. pneumonia IgM Test	S/P	GCMP(IgM)-302a√	Cassette	1	25 Tests/Kit
Respiratory Syncytial Virus Antigen Test	Swab	GCRSV-502a√	Cassette	1	20 Tests/Kit
Rotavirus Test	Feces	GCROA-602a√	Cassette	,	25 Tests/Kit
	reces		Cassette	/	
Rotavirus/Adenovirus Test	0/0	GCROA/ADE-602a√			25 Tests/Kit
Rubella IgG Test -	S/P	GCRUB(IgG)-302a	Cassette		25 Tests/Kit
	WB/S/P	GCRUB(IgG)-402a	Cassette	/	25 Tests/Kit
Rubella IgM Test	S/P	GCRUB(IgM)-302a	Cassette	/	25 Tests/Kit
nabotta igini root	WB/S/P	GCRUB(IgM)-402a	Cassette	/	25 Tests/Kit
	S/P	GCRUB(IgG/IgM)-302a	Cassette	/	25 Tests/Kit
Rubella IgG/IgM Test	WB/S/P	GCRUB(IgG/IgM)-402a	Cassette	/	25 Tests/Kit
	VV B/ S/ P	GCRUB(IgG/IgM)-T402a	Cassette	/	25 Tests/Kit
		GCSTR-501a√	Strip	/	25 Tests/Kit
		GCSTR-501Caà	Strip	/	25 Tests/Kit
Strep A Test	Throat Swab	GCSTR-502a√	Cassette	/	20 Tests/Kit
		GCSTR-502Ca√	Cassette	/	20 Tests/Kit
		GCSYP-301a√	Strip	1	50 Tests/Kit
	S/P	GCSYP-302a√	Cassette		
Syphilis Test —		GCSYP-401a√	Strip		25 Tests/Kit
	WB/S/P	<u>_</u>			50 Tests/Kit
a	0.15.15	GCSYP-402a√	Cassette	,	25 Tests/Kit
S. typhi Antigen Test	S/P/Feces	GCSAL(ST)-602a√	Cassette	/	20 Tests/Kit
TOXO IgG Test -	S/P	GCTOX(IgG)-302a√	Cassette	/	25 Tests/Kit
	WB/S/P	GCTOX(IgG)-402a	Cassette	/	25 Tests/Kit
TOXO IgM Test	S/P	GCTOXO(lgM)-302a√	Cassette	/	25 Tests/Kit
TONO Igivi Test	WB/S/P	GCTOXO(IgM)-402a	Cassette	/	25 Tests/Kit
	C/D	GCTOX-302b	Cassette	/	25 Tests/Kit
Toxo IgG/IgM Test	S/P	GCTOX(IgG/IgM)-302a√	Cassette	/	20 Tests/Kit
	WB/S/P	GCTOX-402b	Cassette	/	25 Tests/Kit
ToRCH Toxo/Rubella/CMV/HSV IgG Combo Test	S/P	GCTOG-345a	Cassette	/	20 Tests/Kit
Torch Toxo/Rubella/CMV/HSV IgM Combo Test	S/P	GCTOM-345a	Cassette	/	20 Tests/Kit
Trichomonas vaginalis Test	Vaginal Swab	GCTV-502a√	Cassette	/	20 Tests/Kit
	S/P	GCTV-302a√	Cassette	/	25 Tests/Kit
Tuberculosis IgG/IgM Test —	WB/S/P		Cassette		
	VV D/ 3/ F	GCTVP 2010		/	25 Tests/Kit
Typhoid IgG/IgM Test	S/P	GCTYP-301a	Strip		50 Tests/Kit
		GCTYP-302a√	Cassette	/	25 Tests/Kit
V. cholerae O1 Antigen Test	Feces	GCVCH(O1)-602a√	Cassette	/	25 Tests/Kit
V. cholerae O1/O139 Antigen Test	Feces	GCVCH(O1/O9)-602a√	Cassette	/	25 Tests/Kit
ZIKA IgM Test	WB/S/P	GCZIK(IgM)-402a	Cassette	/	25 Tests/Kit
ZIKA IgG Test	WB/S/P	GCZIK(IgG)-402a	Cassette	/	25 Tests/Kit
ZIKA NS1 Test	WB/S/P	GCZIK(NS1)-402a	Cassette	/	25 Tests/Kit







<b>Product Description</b>	Specimen	Catalog No.	Format	Cut-off Value	Kit Size
		GAHCG-101aà	Strip	25 mIU/mL	100 Tests/Kit
		GAHCG-101b√	Strip	10 mIU/mL	100 Tests/Kit
		GAHCG-101d√	Strip	20 mIU/mL	100 Tests/Kit
		GAHCG-102aà	Cassette	25 mIU/mL	25 Tests/Kit
	Urine -	GAHCG-102b√	Cassette	10 mIU/mL	25 Tests/Kit
	onne -	GAHCG-102d√	Cassette	20 mIU/mL	25 Tests/Kit
h OO Brannan ay Taat		GAHCG-103aà	Midstream	25 mIU/mL	1/2 Test(s)/Kit
hCG Pregnancy Test		GAHCG-103b√	Midstream	10 mIU/mL	1/2 Test(s)/Kit
		GAHCG-103d√	Midstream	20 mIU/mL	1/2 Test(s)/Kit
Urine/Se		GAHCG-105a	Panel	25 mIU/mL	25 Tests/Kit
	Urine/Serum -	GAHCG-201a√	Strip	25 mIU/mL	100 Tests/Kit
		GAHCG-201b√	Strip	10 mIU/mL	100 Tests/Kit
		GAHCG-202a√	Cassette	25 mIU/mL	25 Tests/Kit
		GAHCG-202b√	Cassette	10 mIU/mL	25 Tests/Kit
Digital Pregnancy Test	Urine	GAHCG-D103a√	Midstream	25 mIU/mL	1/2 Test(s)/Kit
		GALH-101a√	Strip	25 mIU/mL	100 Tests/Kit
		GALH-101b√	Strip	40 mIU/mL	100 Tests/Kit
		GALH-101d	Strip	30 mIU/mL	100 Tests/Kit
LH Ovulation Test	Urine -	GALH-102a√	Cassette	25 mIU/mL	25 Tests/Kit
LH Ovulation Test	Uline -	GALH-102b√	Cassette	40 mIU/mL	25 Tests/Kit
		GALH-103a√	Midstream	25 mIU/mL	1/5 Test(s)/Kit
		GALH-103b√	Midstream	40 mIU/mL	1/5 Test(s)/Kit
		GALH-103d	Midstream	30 mIU/mL	1/5 Test(s)/Kit
FSH Menopause Test	Urine -	GAFSH-101a√	Strip	25 mIU/mL	100 Tests/Kit
ran wenopause test	Ullile	GAFSH-102a√	Cassette	25 mIU/mL	25 Tests/Kit
IGFBP-1 PROM Test	Cervical Secretion	GAIGF1-501a√	Strip	25 ng/mL	25 Tests/Kit
IGFBF-1 FNOW TEST	Cervical Secretion	GAIGF1-502a√	Cassette	25 ng/mL	20 Tests/Kit
Male Fertility Test	Semen	GASPE-902a√	Cassette	15M/mL	1 Test/Kit



<b>Product Description</b>	Specimen	Catalog No.	Format	Cut-off Value	Kit Size
CK-MB Test	S/P	GDCKM-302a√	Cassette	5 ng/mL	25 Tests/Kit
CK-IVIB TEST	WB/S/P	GDCKM-402a√	Cassette	5 ng/mL	25 Tests/Kit
CRP C-Reactive Protein Semi	WB/S/P -	GDCRP-402a√	Cassette	1~3~10 mg/L	25 Tests/Kit
-Quantitative Test	VVD/3/P -	GDCRP-T402b√	Cassette	10~40~80 mg/L	25 Tests/Kit
D-dimer Test	WB/P	GDDDI-402b√	Cassette	500 ng/mL	25 Tests/Kit
Myoglobin Test	WB/S/P	GDMYO-402a√	Cassette	50 ng/mL	25 Tests/Kit
Procalcitonin Test	WB/S/P	GDPCT-T402a√	Cassette	0.5~2~10 ng/mL	25 Tests/Kit
	S/P	GDTRO-302a√	Cassette	0.5 ng/mL	25 Tests/Kit
Troponin I Test	MID/C/D	GDTRO-402a√	Cassette	0.5 ng/mL	25 Tests/Kit
	WB/S/P —	GDTRO-402b√	Cassette	0.5 ng/mL	25 Tests/Kit
Cardiac Myoglobin/CK - MB/cTnl Combo Test	S/P	GDCAR-335a√	Cassette	50/5/0.5 ng/mL	25 Tests/Kit
	W/D/C/D	GDCAR-435a√	Cassette	50/5/0.5 ng/mL	25 Tests/Kit
Wib/Citil Combo lest	WB/S/P —	GDCAR-W435a√	Cassette	50/5/0.5 ng/mL	20 Tests/Kit



				and all
Product Description	Specimen	Format	Cut-off Value	Kit Size
Ascorbateà	Urine	Strip	0.5-0.6 mmol/L	100 Tests/Canister
Bilirubinà	Urine	Strip	8.6-17 μmol/L	100 Tests/Canister
Bloodà	Urine	Strip	5-15 Ery/μL	100 Tests/Canister
Ca√	Urine	Strip	2.5 mmol/L	100 Tests/Canister
Creatinine√	Urine	Strip	50 mg/dL	100 Tests/Canister
Gluoseà	Urine	Strip	2.8~5.5 mmol/L	100 Tests/Canister
Ketoneà	Urine	Strip	0.5~1.0 mmol/L	100 Tests/Canister
Leukocytesà	Urine	Strip	5-15 Leuko/μL	100 Tests/Canister
Micro Albumin√	Urine	Strip	0.08~0.15 mg/dL	100 Tests/Canister
Nitriteà	Urine	Strip	13~22 μmol/L	100 Tests/Canister
pHà	Urine	Strip	0.5	100 Tests/Canister
Proteinà	Urine	Strip	0.15~0.3 g/L	100 Tests/Canister
Specific Gravityà	Urine	Strip	0.005	100 Tests/Canister
Urobilinogenà	Urine	Strip	3.3-16 μmol/L	100 Tests/Canister
Urinary Tract Infection Test Strip	Urine	Strip	LEU: 10-15 Leuko/μL NIT: 13~22 μmol/L	3 Tests/Kit



Product Description	Specimen	Catalog No.	Format	Cut-off Value	Kit Size
AFP Alpha Fetal Protein Test	S/P	GEAFP-301a	Strip	20 ng/mL	50 Tests/Kit
	3/P	GEAFP-302a√	Cassette	20 ng/mL	25 Tests/Kit
	WB/S/P	GEAFP-401a√	Strip	20 ng/mL	50 Tests/Kit
	WD/3/P	GEAFP-402a√	Cassette	20 ng/mL	25 Tests/Kit
	S/P	GECEA-301a	Strip	5 ng/mL	50 Tests/Kit
CEA Carcinoembryonic Antigen Test	3/P	GECEA-302a	Cassette	5 ng/mL	25 Tests/Kit
CEA Carcinoembryonic Antigen Test	WB/S/P	GECEA-401a√	Strip	5 ng/mL	50 Tests/Kit
	WD/3/P	GECEA-402a√	Cassette	5 ng/mL	25 Tests/Kit
		GEFOB-601bà	Strip	50 ng/mL	25 Tests/Kit
		GEFOB-601Cb√	Strip	50 ng/mL	25 Tests/Kit
		GEFOB-601c√	Strip	100 ng/mL	25 Tests/Kit
		GEFOB-601d	Strip	200 ng/mL	25 Tests/Kit
FOB Fecal Occult Blood Test	F	GEFOB-602bà	Cassette	50 ng/mL	20 Tests/Kit
-OB Fedal Occult Blood Test	Feces	GEFOB-602Cb√	Cassette	50 ng/mL	20 Tests/Kit
		GEFOB-602c√	Cassette	100 ng/mL	20 Tests/Kit
		GEFOB-602d	Cassette	200 ng/mL	20 Tests/Kit
		GEFOB-602h	Cassette	150 ng/mL	20 Tests/Kit
		GEFOB-602j√	Cassette	10 ng/mL	20 Tests/Kit
FOB /Transferrin Combo Test	Feces	GEFOB/TF-602a√	Cassette	50/10 ng/mL	20 Tests/Kit
Nuclear Matrix Protein 22 Test	Urine	GENMP22-102a√ New	Cassette	10 U/mL	25 Tests/Kit
	0/5	GEPSA-301a√	Strip	4 ng/mL	50 Tests/Kit
DSA Brostato Spooifio Antigon Tost	S/P	GEPSA-302a√	Cassette	4 ng/mL	25 Tests/Kit
PSA Prostate Specific Antigen Test	WB/S/P	GEPSA-401a√	Strip	4 ng/mL	50 Tests/Kit
	VV D/ O/ I	GEPSA-402a√	Cassette	4 ng/mL	25 Tests/Kit
PSA Prostate Specific Antigen	S/P	GEPSA-302b	Cassette	4 ng/mL, 10 ng/mL	25 Tests/Kit
Semi-QuantitativeTest	WB/S/P	GEPSA-402b	Cassette	4 ng/mL, 10 ng/mL	25 Tests/Kit
Transferrin Test	Feces	GETF-601a√	Strip	10 ng/mL	25 Tests/Kit
Hallstellill Lest	reces	GETF-602a√	Cassette	10 ng/mL	20 Tests/Kit

# Veterinary

Product Description	Specimen	Catalog No.	Format	Label	Cut-off Value	Kit Size
Canine Adenovirus (CAV) Antigen Test	Secretions	GFCAV-502a	Cassette	Gold	/	10 Tests/Kit
Canine Coronavirus (CCV) Antigen Test	Feces	GFCCV-602a	Cassette	Gold	/	10 Tests/Kit
		FFCCV-602a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Canine Coronavirus (CCV) &	Feces	GFCCP-T602a	Cassette	Gold	/	10 Tests/Kit
Parvovirus (CPV) Antigen Combo Test		FFCCP-T602a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Canine C-Reactive Protein (cCRP) Test	WB/S/P	FFCCR-402a	Cassette	Fluorescence	10 mg/L	10 Tests/Kit
Canine Distemper Virus (CDV) Antigen Test	Secretions	GFCDV-502a	Cassette	Gold	/	10 Tests/Kit
	OGGIGUOII3	FFCDV-502a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Canine Distemper Virus (CDV), Influenza Virus (CIV) & Adenovirus (CAV) Antigen Combo Test	Secretions	GFCDIA-532a	Cassette	Gold	/	10 Tests/Kit
Canine Influenza Virus (CIV) Antigen Test	Secretions	GFCIV-502a	Cassette	Gold	/	10 Tests/Kit
Canine Parvovirus (CPV) Antigen Test	Feces	GFCPV-602a	Cassette	Gold	/	10 Tests/Kit
Canine Parvovirus (CPV) Antigen Test		FFCPV-602a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Canine Progesterone (cProg) Test	WB/S/P	FFCPR-402a	Cassette	Fluorescence	15 ng/mL	10 Tests/Kit
Feline Calicivirus (FCV) Antigen Test	Secretions	GFFCV-502a	Cassette	Gold	/	10 Tests/Kit
retifie Caticivitus (FCV) Artitigeri Test		FFFCV-502a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Feline Coronavirus (FCoV) Antigen Test	Feces	GFFCO-602a	Cassette	Gold	/	10 Tests/Kit
Feline Herpes Virus (FHV) Antigen Test	Secretions	GFFHV-502a	Cassette	Gold	/	10 Tests/Kit
retifie herpes virus (FHV) Affiligen Test		FFFHV-502a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Feline Parvovirus (FPV) Antigen Test	Feces	GFFPV-602a	Cassette	Gold	1	10 Tests/Kit
retifie ratvovitus (FPV) Affiligen Test		FFFPV-602a	Cassette	Fluorescence	10 IU	10 Tests/Kit
Feline Parvovirus (FPV) & Coronavirus FCoV) Antigen Combo Test	Feces	GFFPC-622a	Cassette	Gold	1	10 Tests/Kit
Feline Serum Amyloid A (fSAA) Test	WB/S/P	FFFSA-402a	Cassette	Fluorescence	5 mg/L	10 Tests/Kit
Toxoplasma (Toxo) IgG/IgM Test	WB/S/P	GFTOX-402a	Cassette	Gold	1	10 Tests/Kit

# Non-Infectious Disease

<b>Product Description</b>	Specimen	Catalog No.	Format	Cut-off Value	Kit Size
Micro-Albumin Test	Urine	GIHSA-101a√	Strip	20 μg/mL	100 Tests/Kit
	Office	GIHSA-102a	Cassette	20 μg/mL	25 Tests/Kit
Vaginal pH Test	Vaginal Secretion	VPH-501a New	Strip	3.8-4.4	100 Tests/Canister

## Autoimmunity

<b>Product Description</b>	Specimen	Catalog No.	Format	Kit Size	
Rheumatoid Factor IgM Test	S/P	GCRF(IgM)-302a	Cassette	25 Tests/Kit	
Total IgE Test	S/P	GCIGE-302a	Cassette	25 Tests/Kit	





<b>Product Description</b>	Model
Urine Analyzer	Healgen 500√
Urine Analyzer	Healgen 501√
Colloidal Gold Test Reader	OG-D180
Handheld Oral Fluid Drug Test Reader	OG-D200
Multi-Function Colloidal Gold Test Reader	OG-D600
Fluorescence Immunoassay Analyzer	OG-G200
Handheld Fluorescence Immunoassay Analyzer	OG-G300
Mini Immunofluorescence Analyzer	OG-H100√
Veterinary Fluorescence Immunoassay Analyzer	OG-V100
/CE Marked †Cleared for US 510(k)	In Specimen column: WB: Whole Blood S: Serum P: Plasma



Zhejiang Orient Gene Biotech Co., Ltd was founded in December 2005 and listed on the SEE STAR Market on February 5, 2020 (securities code: 688298).

Orient Gene specializes in R&D, production and sales of in vitro diagnostic products, mainly covering infectious diseases (including COVID-19 test series), toxicology, tumor markers, cardiac markers and fertility testing, etc. Through 16 years of technology accumulation and continuous investment in R&D, the Company has independently developed hundreds of products. The company own more than 200 authorized patents, and has obtained more than 500 product medical device certifications at home and abroad. The Company's sales network covers more than 100 countries, products are mainly sold to Europe, America and other developed countries.

Healgen Scientific LLC, a wholly owned subsidiary of Zhejiang Orient Gene Biotech Co., Ltd develops, manufactures and commercializes in vitro diagnostic test systems worldwide. Our product portfolio spans multiple testing categories and analytes to meet various clinical and laboratory needs.

> Healgen Scientific Limited Liability Company Add: 3818 Fuqua Street, Houston, TX77047, USA. Tel: +1 713-733-8088 Toll free: 866 982 3818 Fax: +1713-733-8848

E-mail: <u>Healgensales@healgen.us</u> (For South America and North America) Web: http://www.healgen.com

> Zhejiang Orient Gene Biotech Co., Ltd Add: 3787#, East Yangguang Avenue, Dipu Street, Anji, Huzhou, Zhejiang, China. P.C.: 313300 Tel: +86-572-5303755/5303756 Fax: +86-572-5226222

E-mail: <a href="mailto:sales@orientgene.com">sales@orientgene.com</a> (For rest of world) Web: <a href="http://www.orientgene.com">http://www.orientgene.com</a>

Rev.08/2022

# **PRODUCT** CATALOG

# Enhancing Global Health





## **Troponin I**

# Troponin I Rapid Test Device (Whole Blood/Serum/Plasma)

## Package Insert

A rapid visual immunoassay for the qualitative presumptive detection of cardiac Troponin I in human whole blood, serum, or plasma specimens.

For professional in vitro diagnostic use only.

## INTENDED USE

The Troponin I Rapid Test Device (Whole Blood/Serum/Plasma) is a rapid visual immunoassay for the qualitative presumptive detection of cardiac Troponin I in human whole blood, serum, or plasma specimens. This kit is intended to be used as an aid in the diagnosis of myocardial infarction (MI).

## SUMMARY

Cardiac Troponin I (cTnI) is a protein found in cardiac muscle with a molecular weight of 22.5 kDa. Troponin I is part of a three subunit complex comprising of Troponin T and Troponin C. Along with tropomyosin, this structural complex forms the main component that regulates the calcium sensitive ATPase activity of actomyosin in striated skeletal and cardiac muscle. After cardiac injury occurs, Troponin I is released into the blood 4-6 hours after the onset of pain. The release pattern of cTnI is similar to CK-MB, but while CK-MB levels return to normal after 72 hours, Troponin I remains elevated for 6-10 days, thus providing for a longer window of detection for cardiac injury. The high specificity of cTnI measurements for the identification of myocardial damage has been demonstrated in conditions such as the perioperative period, after marathon runs, and blunt chest trauma. CTnI release has also been documented in cardiac conditions other than acute myocardial infarction (AMI) such as unstable angina, congestive heart failure, and ischemic damage due to coronary artery bypass surgery. Because of its high specificity and sensitivity in the myocardial infarction.

## PRINCIPLE

The Troponin I Rapid Test Device (Whole Blood/Serum/Plasma) has been designed to detect cardiac Troponin I through visual interpretation of color development in the strip. The membrane was immobilized with anti-cTnI antibodies on the test region.

During the test, the specimen is allowed to react with colored anti-cTnl antibodies colloidal gold conjugates, which were precoated on the sample pad of the test. The mixture then moves on the membrane by a capillary action, and interact with reagents on the membrane. If there were enough cTnl in specimens, a colored band will form at the test region of the membrane.

Presence of this colored band indicates a positive result, while its absence indicates a negative result. Appearance of a colored band at the control region serves as a procedural control. This indicates that proper volume of specimen has been added and membrane wicking has occurred.

## **PRECAUTIONS**

- 1. For professional In Vitro diagnostic use only.
- Warning: the reagents in this kit contain sodium azide which may react with lead or copper plumbing to form potentially explosive metal azides. When disposing of such reagents, always flush with large volumes of water to prevent azide build-up.
- Do not use it if the tube/pouch is damaged or broken.
- . Test is for single use only. Do not re- use under any circumstances.
- Handle all specimens as if they contain infectious agents. Observe established standard procedure for proper disposal of specimens
- Wear protective clothing such as laboratory coats, disposable gloves and eye protection when specimens are assay.
- 7. Humidity and temperature can adversely affect results

## STORAGE AND STABILITY

All reagents are ready to use as supplied. Store unused test device unopened at 2°C-30°C. If stored at 2°C-8°C, ensure that the test device is brought to room temperature before opening. The test is not stable out off the expiration date printed on the sealed pouch. Do not freeze the kit or expose the kit over 30°C.

## SPECIMEN COLLECTION AND PREPARATION

- The Troponin I Rapid Test Device (Whole Blood/Serum/Plasma) is intended only for use with human whole blood, serum, or plasma specimens.
- Only clear, non-hemolyzed specimens are recommended for use with this test.
- Serum or plasma should be separated with soonest possible opportunity to avoid hemolysis.
- Perform the testing immediately after the specimen collection. Do not leave the specimens at room temperature for prolonged periods. Specimens may be stored at 2-8°C for up to 3 days. For long term storage, specimens should be kept below -20°C.
- Bring specimens to room temperature prior to testing. Frozen specimens must be completely thawed and mixed well prior to testing. Avoid repeated freezing and thawing of specimens.

- Pack the specimens in compliance with applicable regulations for transportation of etiological agents, in case they need to be shipped.
- Icteric, lipemic, hemolysed, heat treated and contaminated sera may cause erroneous results.
- There is a slight possibility that some whole blood specimens with very high viscosity
  or which have been stored for more than 2 days may not run properly on the test
  device. Repeat the test with a serum or plasma specimen from the same patient using
  a new test device.

### **MATERIALS**

### Materials Provided

Test devicesBuffer

- Disposable Droppers
- Package insert

## Materials Required But Not Provided

- Specimen collection containers
   Centrifuge (for plasma only)
- Clock or Timer

## DIRECTIONS FOR USE

Allow test device, specimen, buffer and/or controls to equilibrate to room temperature (15-30°C) prior to testing.

- Remove the test from its sealed pouch, and place it on a clean, level surface. Label
  the device with patient or control identification. To obtain a best result, the assay
  should be performed within one hour.
- Transfer 2 drops of serum or plasma to the specimen well of the device with a disposable pipette provided in the kit, and then start the timer.

Transfer 3 drops of whole blood specimen to the specimen well of the device with a disposable pipette provided in the kit, then add 1 drop of buffer, and start the timer.

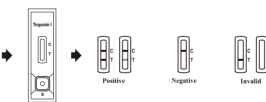
Allow 3 hanging drops of fingerstick whole blood specimen to fall into the center of the specimen well (S) on the device, then add 1 drop of buffer, and start the timer. Avoid trapping air bubbles in the specimen well (S), and do not drop any solution in observation window.

As the test begins to work, you will see color move across the membrane.

3. Wait for the colored band(s) to appear. The result should be read at 10 minutes. If

Wait for the colored band(s) to appear. The result should be read at 10 minutes. Do not interpret the result after 20 minutes.





## INTERPRETATION OF RESULTS

(Please refer to the illustration above)

**POSITIVE**: Two colored bands appear on the membrane. One band appears in the control region (C) and another band appears in the test region (T).

**NEGATIVE**:Only one colored band appears in the control region (C). No apparent colored band appears in the test region (T).

**INVALID**:Control band fails to appear. Results from any test which has not produced a control band at the specified reading time must be discarded.

Please review the procedure and repeat with a new test. If the problem persists, discontinue using the kit immediately and contact your local distributor.

- The intensity of the color in test region (T) may vary depending on the concentration
  of aimed substances present in the specimen. Therefore, any shade of color in the
  test region should be considered positive. Besides, the substances level can not be
  determined by this qualitative test.
- Insufficient specimen volume, incorrect operation procedure, or performing expired tests are the most likely reasons for control band failure.

## QUALITY CONTROL

Internal procedural controls are included in the test. A colored band appearing in the control region (C) is considered an internal positive procedural control. It confirms sufficient specimen volume and correct procedural technique.

External controls are not supplied with this kit. It is recommended that positive and negative controls be tested as a good laboratory practice to confirm the test procedure and to verify proper test performance.

### LIMITATIONS

- The Troponin I Rapid Test Device (Whole Blood/Serum/Plasma) is for professional in vitro diagnostic use, and should be used for the qualitative detection of cardiac Troponin I only. There is no meaning attributed to linen color intensity or width.
- The Troponin I Rapid Test Device (Whole Blood/Serum/Plasma) will only indicate the
  presence of Troponin I in the specimen and should not be used as the sole criteria for
  the diagnosis of tuberculosis.
- 3. If the test result is negative and clinical symptoms persist, additional testing using other clinical methods is recommended. The test cannot detect less than 0.5 ng/mL of cTnI in specimens. Thus, a negative result does not at anytime rule out the existence of Troponin I in blood, because the antibodies may be absent or below the minimum detection level of the test.
- Like with all diagnostic tests, a confirmed diagnosis should only be made by a physician after all clinical and laboratory findings have been evaluated.
- Some specimens containing unusually high titers of heterophile antibodies or rheumatoid factor (RF) may affect expected results. Even if the test results are positive, further clinical evaluation should be considered with other clinical information available to the physician.

## PERFORMANCE CHARACTERISTICS

Table: Trononin I Ranid Test vs. FIA

Table: Hopoliili Frapia Test vs. EIA						
Method		Tropo Rapid Te	Total Results			
EIA	Results	Positive	Negative	Results		
	Positive	138	2	140		
	Negative	1	315	316		
Total Results		139	317	456		

Relative Sensitivity: 98.6% (94.9%-99.8%)\*
Relative Specificity: 99.7% (98.3%-99.9%)\*

Relative Specificity, 99.7% (90.5%-99.9%

Overall Agreement: 99.3% (98.1%-99.9%)\*

\*95% Confidence Interval

## **BIBLIOGRAPHY**

- Adams, et al. Biochemical markers of myocardial injury, Immunoassay Circulation 88: 750-763, 1993.
- Mehegan JP, Tobacman LS. Cooperative interaction between troponin molecules bound to the cardiac thin filament. J.Biol.Chem. 266:966, 1991.
- Adams, et al. Diagnosis of Perioperative myocardial infarction with measurements of cardiac troponin I. N.Eng.J.Med 330.670, 1994.
   Hossein-Nia M. et al. Cardiac troponin I release in heart transplantation. Ann. Thorac.
- Hossein-Nia M, et al. Cardiac troponin i release in heart transplantation. Ann. Thorac Surg. 61: 227, 1996.
- Alpert JS, et al. Myocardial Infarction Redefined, Joint European Society of Cardiology American College of Cardiology. J. Am. Coll. Cardio., 36(3):959, 2000.

B20570-01