

Specification

Feature	Specification	MS		
Technology	Biosensor/Electrochemical, Glucose oxidase (GOD)	The state of the s		
Result Calibration	Plasma-equivalent			
Test Time	10 seconds			
Sample Size	0.5 μL			
Sample Type	Fresh capillary whole blood			
Hematocrit Range	25 - 60%			
Glucose Test Range	20 - 600 mg/dL (1.1 - 33.3 mmol/L)			
Memory Storage	300 results with date and time	300 results with date and time		
Test Averaging	7, 14, 30-day averages			
Data Transfer	USB			
Control Solution	3 levels			
Audio Feature	Optional beep for sample detection, error messages			
Automatic Shutoff	2 minutes after last action			
Battery	One (1) CR 2032 3.0V coin cell battery			
Battery Life	1,000 measurements			
Operating Conditions	41 - 113 °F (5 - 45°C) and 10 - 90% relative humidity			
Strip Storage Temperature	2-35°C			
Expiration Date	24 months (6 months after first opening)			

Catalog

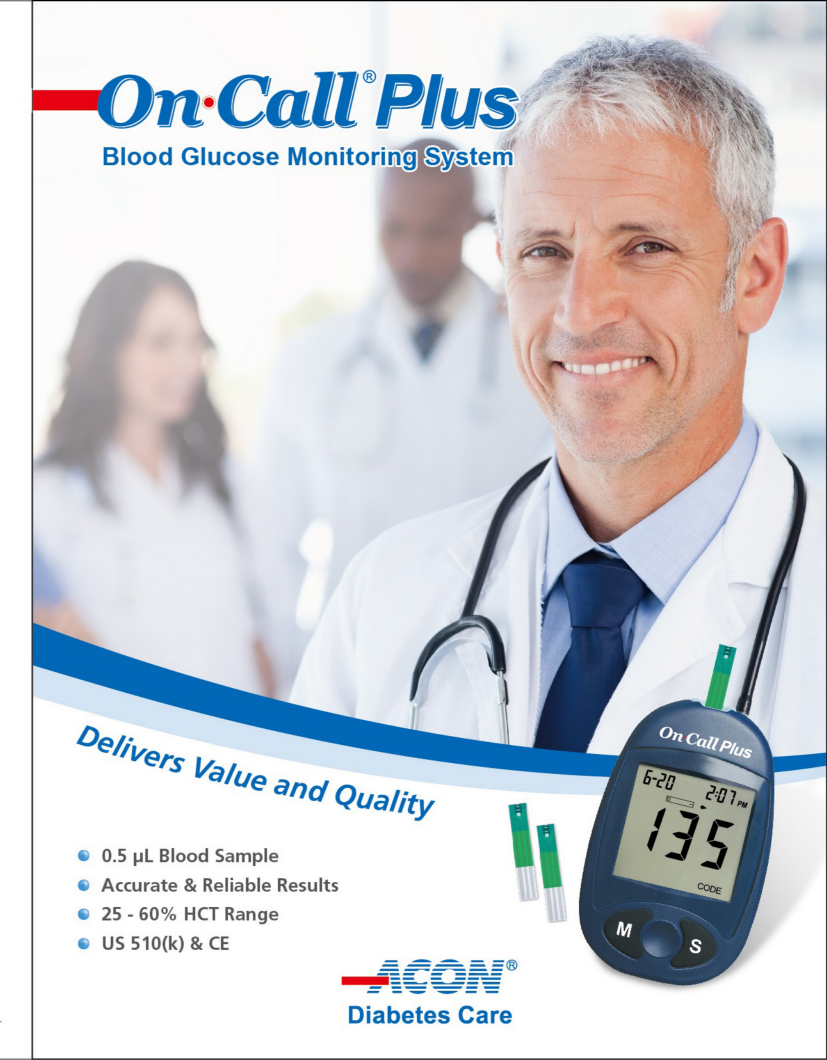
Product Name	Catalog No.			Conte	ents		
On Call® Plus Blood Glucose Monitoring System	G113-111 √ †	1 Meter 1 Manual 10 Lancets	10 Test St 1 Carrying 1 Code Ch	Case		lution 1 erence Guide (for testing on forear	1 Lancing Device 1 Warranty Card m and palm)
<i>On Call® Plus</i> Blood Glucose Meter	G113-211 √ †	1 Meter 1 Manual		rol Solution 1 ranty Card		arrying Case uick Reference Guide)
	G113-214 V	1 Meter 1 Manual 10 Lancets	1 Carry	ing Device ying Case ranty Card	10	ontrol Solution 1 Juick Reference Guide lear Cap (for testing o	e on forearm and palm)
On Call® Plus Blood Glucose Test Strips	0400 444 44	50 Test Strips (25/vial)		1 C	ode Chip	1 Package Insert
	G133-111 √ †	50 Test Strips (50/vial)		1 C	ode Chip	1 Package Insert
	G133-112 √	100 Test Strips	(25/vial)		1 C	ode Chip	1 Package Insert
	G133-114 V	10 Test Strips (10/vial)		1 C	ode Chip	1 Package Insert
	G133-115 √	25 Test Strips (Individually I	Foil Wrapped)	10	ode Chip	1 Package Insert
	G133-117 √	50 Test Strips (Individually I	Foil Wrapped)	10	ode Chip	1 Package Insert
	G133-118 √	25 Test Strips (25/vial)		1 C	ode Chip	1 Package Insert
On Call® Plus Blood Glucose Test Strips and Lancets	G133-211 √	50 Test Strips (25/vial)	50 Lancets (2	25/bag)	1 Code Chip	1 Package Insert
On Call® Plus Blood Glucose Control Solution	G123-311 à	1 Control Solut	tion 0 1	Control Solution	on 1 1	Control Solution 2	1 Package Insert
On Call® Lancets	G124-10A à	100 Lancets (2	5/bag)				
On Call® Lancing Device	G124-11AV	1 Lancing Devi	ce		1 Package In	sert	
On Call® Diabetes Management Software Kit	G124-13A†	1 USB Data Tra	nsfer Cable		1 Installation	n Disk	

v CE Marked for sale in the European Community (6 0123



† US 510(k) Cleared and CLIA Waived

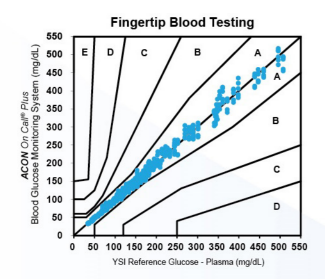




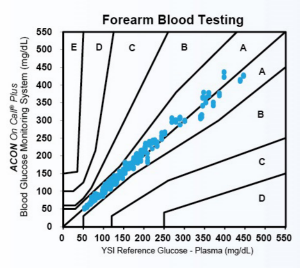


Accurate and Reliable

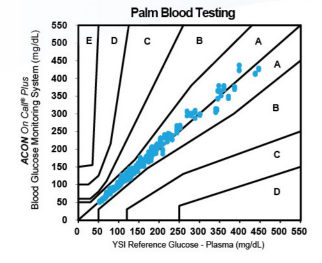
Extensive clinical studies proved the accuracy of *On Call® Plus* Blood Glucose Monitoring System with fresh capillary blood samples, which can comply with EN ISO 15197: 2015.



Clinical Trial -	onsensus Error Grid Ana Fingertip Capillary Bloo lus Blood Glucose Monit	d, by Technican		
System Accuracy Res	System Accuracy Results for Glucose Concentration \geq 100 mg/dL			
Within ±5%	Within ±10%	Within ±15%		
290 / 462 (62.8%)	432 / 462 (93.5%)	462 / 462 (100.0%)		
System Accuracy Results for Glucose Concentration <100 mg/dL				
Within ±5 mg/dL	Within ±10 mg/dL	Within ±15 mg/dL		
145 / 198 (73.2%)	193 / 198 (97.5%)	198 / 198 (100.0%)		
System Accuracy Results for both Glucose Concentration ≥ 100 mg/dL and < 100 mg/dL				
Within ±15% or ±15 mg/dL				
658 / 660 (99.7%)				



Clinical Trial	onsensus Error Grid Ana - Forearm Capillary Bloo <i>lus</i> Blood Glucose Monit	d, by Technican	
System Accuracy Res	sults for Glucose Conce	ntration ≥ 100 mg/dL	
Within ± 5%	Within ± 10%	Within ± 15%	
202 / 444 (45.5%)	375 / 444 (84.5%)	440 / 444 (99.1%)	
System Accuracy Results for Glucose Concentration <100 mg/dL			
Within ±5 mg/dL	Within ±10 mg/dL	Within ±15 mg/dL	
110 / 168 (65.5%)	154 / 168 (91.7%)	168 / 168 (100.0%)	
System Accuracy Results for both Glucose Concentration ≥ 100 mg/dL and < 100 mg/dL			
V	Vithin ±15% or ±15 mg/o	dL	
	608 / 612 (99.3%)		



Clinical Tria	onsensus Error Grid Ana al - Palm Capillary Blood, <i>lus</i> Blood Glucose Monit	by Technican	
System Accuracy Re	sults for Glucose Conce	ntration ≥ 100 mg/dL	
Within ±5%	Within ±10%	Within ±15%	
219 / 444 (49.3%)	395 / 444 (89.0%)	441 / 444 (99.3%)	
System Accuracy Results for Glucose Concentration < 100 mg/dL			
Within ±5 mg/dL	Within ±10 mg/dL	Within ±15 mg/dL	
130 / 168 (77.4%)	166 / 168 (98.8%)	168 / 168 (100.0%)	
System Accuracy Results for both Glucose Concentration ≥ 100 mg/dL and < 100 mg/dL			
١	Vithin ±15% or ±15 mg/o	dL	
	609 / 612 (99.5%)		



Key Features



Authority Certificate







CE certificate

USFDA CFG certificate

Health Canada 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. 01

Manufacturer: ACON Laboratories, Inc.

5850 Oberlin Drive, #340 San Diego CA 92121

USA

Product Category(ies): In Vitro diagnostics for the detection of

human infections and tumor markers, blood glucose measuring self-testing systems,

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. See also notes overleaf.

Report no.: SH1974310

 Valid from:
 2019-10-24

 Valid until:
 2022-09-12

Date, 2019-10-24

Stefan Preiß

1. Pumil

Head of Certification/Notified Body

Page 1 of 4
TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123



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. 01

Model(s): For Detail Models see attachment

Facility(ies):

ACON Laboratories, Inc.
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





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. 01

For the product(s)/product category (ies):

On Call Plus Blood Glucose Monitoring System,

On Call Plus Blood Glucose Test Strips,

On Call EZ II Blood Glucose Monitoring System,

On Call Redi Blood Glucose Monitoring System,

On Call Redi II Blood Glucose Test Strips,

On Call Advanced Blood Glucose Monitoring System,

On Call Advanced Blood Glucose Test Strips,

On Call Platinum Blood Glucose Monitoring System,

On Call Platinum Blood Glucose Test Strips,

On Call Chosen Blood Glucose Monitoring System,

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 Vivid Pal Blood Glucose Monitoring System (OGM-102),

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-171),

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),

D-ONE Blood Glucose Monitoring System,

D-ONE Blood Glucose Test Strips,

Urinalysis Reagent Strips (Urine),

UTI Urinary Tract Infection Test Strips,

Toxoplasma IgG EIA Test Kit,

Toxoplasma IgM EIA Test Kit,

Rubella IgG EIA Test Kit,

Rubella IgM EIA Test Kit,

CMV IgG EIA Test Kit,

Page 3 of 4

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



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. 01

CMV IgM EIA Test Kit,

Total PSA EIA Test Kit,

PT Coagulation Monitoring System (CCM-121),

PT Coagulation Test Strips (CCS-121),

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)

On Call GU Dual Blood Glucose & Uric Acid Monitoring 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









Full Quality Assurance System
Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4)
(Devices in Class IIa, IIb or III)

No. G1 104507 0002 Rev. 01

Manufacturer: ACON Laboratories, Inc.

5850 Oberlin Drive, #340 San Diego CA 92121

USA

Product Category(ies): Lancets, Safety Lancets

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 categories in accordance with MDD Annex II. This quality assurance system conforms to the requirements of this Directive and is subject to periodical surveillance. For marketing of class III devices an additional Annex II (4) certificate is mandatory. See also notes overleaf.

Report No.: SH1974310

 Valid from:
 2019-10-24

 Valid until:
 2023-09-06

Date, 2019-10-24

Stefan Preiß
Head of Certification/Notified Body

1. Punil





Full Quality Assurance System Directive 93/42/EEC on Medical Devices (MDD), Annex II excluding (4) (Devices in Class IIa, IIb or III)

No. G1 104507 0002 Rev. 01

Facility(ies): ACON Laboratories, Inc.

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



10125 Mesa Rim Road. · San Diego, CA 92121 · USA Tel: (858) 875-8000 · Fax: (858) 875-8099 · E-mail: info@aconlabs.com

November 11th 2016

CERTIFICATION LETTER

This letter is to certify that, Vitalie Goreacii, employed by Sanmedico SRL located at: Republic of Moldova, city Chisinau, str. Petricani 88/1 of. 10, MD-2059, have received all required training and is enabled and authorized to provide services with installation, commissioning, and maintenance to the products listed below:

Mission® U120 Urine Analyzer

Mission® U120 Ultra Urine Analyzer

Mission® U500 Urine Analyzer

Mission® PT/INR Coagulation Monitoring System

Mission® Cholesterol Monitoring System

Mission® Ultra Cholesterol Monitoring System

Mission® HB Hemoglobin Testing System

Mission® Plus HB Hemoglobin Testing System

OnCall® Glucose Meter

For further questions or inquiries regarding this matter, please refer to the contact information below.

Sincerely

Jassy Alvarenga

International Account Manager

ACON Laboratories, Incs.A.

jalvarenga@aconlabs.com

+1 858 875 8085

Letter of Declaration

To whom it may concern:

We *Acon Laboratories,Inc.*, who is the legal manufacturer of Blood Glucose Monitoring System (Including Glucose Meter, Glucose test strip, Control Solution, Lancet and lancing device etc, to test the glucose level of human blood),have registered office at 10125 Mesa Rim Road, San Diego, CA 92121 USA, here to declare that:

- On Call® Plus Strips correspond with On Call® Plus Blood Glucose Monitoring System.
- We currently have in stock the tender required quantity of Meters, Strips and Lancets (1000/50000/50000).

This clarification letter will only be used for product registration, tender submission, sales and marketing of *On Call® Plus* Blood Glucose Monitoring System in **Moldova** it should not be used for any other business or non-business purposes.

Sincerely yours,

Eddie.SA

International Sales Warketing Sales Manager

Diabetes Care

Acon Laboratories,Inc.







Certificate

No. Q5 104507 0001 Rev. 01

Holder of Certificate: ACON Laboratories, Inc.

5850 Oberlin Drive, #340 San Diego CA 92121

USA

Certification Mark:



Scope of Certificate: Design and Development,

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

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). See also notes overleaf.

Report No.: SH1974310

 Valid from:
 2019-10-24

 Valid until:
 2022-09-06

Date, 2019-10-24

Stefan Preiß

Head of Certification/Notified Body





Certificate

No. Q5 104507 0001 Rev. 01

EN ISO 13485:2016 Applied Standard(s):

Medical devices - Quality management systems -

Requirements for regulatory purposes

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

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

ACON Laboratories, Inc.

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

AZURE Institute, Inc.

10125 Mesa Rim Road, San Diego CA 92121, USA



STATEMENT

We, ACON Laboratories, Inc. having a registered office at 5850 Oberlin Drive #340, San Diego, CA 92121 assign SRL Sanmedico having a registered office at A. Corobceanu street 7A, apt. 9, Chisinău, MD-2012, Moldova, as authorized representative in correspondence with the conditions of directive 98/79/EC.

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 authorization will be valid for one year after the date of this statement.

Date: June 1, 2021

Signature:

Qiyi Xie, Md, MPH

Sr. Officer, Regulatory & Clinical Affairs

ACON Laboratories, Inc.

Ph: 858-875-8011

Email: gxie@aconlabs.com