

SQA-iC User Manual

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SECTION 1: System Specifications and Requirements

The SQA-iO is a high-performance PC-based semen analyzer. The device works with a computer app that interfaces with the device to guide the user through sample testing and results archiving on the cloud. The SQA-iO device is intended for prescription use only.

Device Hardware: Houses a measurement compartment for testing and USB connection for connectivity.

Specifications

- Dimensions: 8 X 9.5 X 10.5 cm / Weight: 0.350 Kg
- Analysis Time: 75 seconds
- Power supply: 5V DC (USB)
- Noise level: 0 [dBA]
- Device power consumption: 1.7 [BTU/hour] = 0.5 [Watts]
- Sources of radiant energy: Two LEDs (motility and concentration channels)
- Detector system: Two photo detectors (Motility and Optical Density)
- Software: Resides on flash memory and on a secure server on the Cloud
- Motility channel input signal: Analog, up to 5V
- Recommended browsers for optimal performance: Chrome, Microsoft Edge

SQA-iO Minimum Requirements

- PC: Intel Core i5 M520 2.4GHz or equivalent
- RAM: 4GB
- Monitor Screen: Color, Wide screen minimum resolution 1024 x 768
- Operating system compatibility: Windows 7 Professional or above
- Communication Ports: one USB port
- Internet Access: 5mb/second

Operating/Sample Temperature, Humidity and Altitude

- Operates in ambient temperature (15-38°C). Calibrated at room temperature: 20-25°C (68-77°F).
- Maximum operational humidity up to 80% for temperatures up to 31°C. Linearity decreased 50% at 38°C.
- Intended for indoor use at a maximum altitude of 2000m, mains supply fluctuations ±10%,
- Overvoltage Category II, Pollution Degree II.

Quality Control/Calibration

• Internal: Electronic Self-Test/Auto-Calibration runs @ start-up. Reference values verified prior to each test.

Sample Testing

- Calibrated to test samples at room temperature 20-25°C (68-77°F) within one hour of sample collection.
- Test only liquefied human semen samples. QwikCheck Liquefaction vials (available from MES and sold separately) can be used to liquify semen sample prior to aspirating the sample into the testing capillary, when needed.

Accessory (optional) devices:

- <u>SQA-VU Visualization system</u> works only with the SQA-iO to visualize sperm samples and capture Motility videos and Morphology images. Additional information can be found in Appendix 7.
- SQA-iO Docking station enables both the SQA-iO and SQA-Vu to connect to one power source and maintain a small laboratory footprint.





SECTION 2: Semen Parameters and Reportable Range

The SQA-iO is a high-performance PC-based analytical medical device that tests FRESH and WASHED semen samples. The device works with a computer application that contains the device, patient, sample, test results and facility information.

After collection and preparation, a semen sample is withdrawn into an SQA testing capillary and inserted into the SQA-iO where the sample testing is performed. Test results are available in 75 seconds.

SQA-iO Reportable Range							
WHO 5 th	Range	WHO 6th	Range				
CONCENTRATION (M/ml)	<2-400	CONCENTRATION (M/ml)	<2-400				
TOTAL MOTILITY (PROG + NON PROG) (%)	0-100	TOTAL MOTILITY (PROG + NON PROG) (%)	0-100				
PROGRESSIVE (%)	0-100	PROGRESSIVE (RAPID + SLOW) (%)	0-100				
		RAPIDLY PROGRESSIVE (%)	0-100				
		SLOWLY PROGRESSIVE (%)	0-100				
NON-PROGRESSIVE (%)	0-100	NON-PROGRESSIVE (%)	0-100				
IMMOTILE (%)	0-100	IMMOTILE (%)	0-100				
NORMAL FORMS (%)	0-30	NORMAL FORMS (%)	0-30				
MOTILE SPERM CONC.* (M/ml)	<0.2-250	MOTILE SPERM CONC.* (M/ml)	<0.2-250				
PROG. MOTILE SPERM CONC.* (M/ml)	0-200	PROG. MOTILE SPERM CONC.* (M/ml)	0-200				
		RAPID PR. MOTILE SPERM CONC.* (M/ml)	0-100				
		SLOW PR. MOTILE SPERM CONC.* (M/ml)	0-100				
FUNCTIONAL SPERM CONC.* (M/ml)	0-120	FUNCTIONAL SPERM CONC.* (M/ml)	0-120				
VELOCITY (VCL)** (mic/sec)	0-100	VELOCITY (VCL)** (mic/sec)	0-100				
SPERM MOTILITY INDEX**	0-500	SPERM MOTILITY INDEX**	0-500				
SPERM # (M/ejac)	0-900	SPERM # (M/ejac)	0-900				
MOTILE SPERM* (M/ejac)	0-800	MOTILE SPERM* (M/ejac)	0-800				
PROG. MOTILE SPERM* (M/ejac)	0-700	PROG. MOTILE SPERM* (M/ejac)	0-700				
FUNCTIONAL SPERM* (M/ejac)	0-150	FUNCTIONAL SPERM* (M/ejac)	0-150				
MORPH NORMAL SPERM** (M/ejac)	0-260	MORPH NORMAL SPERM** (M/ejac)	0-260				

*MES parameters are indicated by an asterisk. ** These parameters are not reported in the US market







Testing Capillary

- Disposable, plastic, testing capillary. Requires 500 μ l of sample for normal volume testing, 10 μ l for low volume testing.
- Designed to collect and test samples in a biologically safe manner. Use only manufacturers' certified testing capillaries.

Cuvette Section (Concentration assessment)

• Millions of sperm cells are analyzed in the 'tall' cuvette section of the testing capillary based on spectrophotometry analysis of the semen sample and application of proprietary algorithms.

Motility Section (Motility parameter assessment)

- Tens of thousands of sperm cells are analyzed in the 'thin' motility section of the testing capillary as they move through a light beam in the device.
- Light disturbances are then converted into analog signals and analyzed by proprietary algorithms.

Inserting the Testing Capillary into the SQA-iO

• After filling the testing capillary (see Appendix Section for guidelines), insert the SQA testing capillary all the way into the SQA-iO measurement chamber with the BLUE PISTON facing down.

SECTION 4: Getting Started

First time connecting: Follow the **SQA-iO Quick Start Guide** instructions or directly download the software from <u>www.sqa-io.com</u> and register your account following the on-screen instructions. This is the time to set all preferred testing and your facility defaults and to load test credits.

Connecting to the SQA-iO for testing:

- Connect the SQA-iO to the computer using the supplied USB cable.
- Go to: <u>www.sqa-io.com</u> or click on the desktop icon **o** to activate the SQA-iO interface.
- Enter the unique 8-digit registration number located inside your device kit
- If you are asked to authorize the download of a driver required to run the SQA-iO app, please accept.
- Log into the SQA-iO using your username and password.
- The SQA-iO will now go through a calibration check, wait until it finishes.
- The device is now ready for sperm testing

SECTION 5: SQA-iO Navigation and Testing

- **Navigation:** The SQA-iO navigation bar is always available. Click on one of the options on the SQA-iO navigation bar to select where you want to go.
- **Security:** After 15 minutes of SQA-iO inactivity, a timeout warning will be displayed. If the device/app is not used for another 5 minutes, the SQA-iO will shut down. Log back in when ready to start testing.



The Home Screen provides the following information:

- This icon is **GREEN** when the device is connected and **RED** when disconnected.
- This icon is **BLACK** when the internet connection is stable, **RED**! when the internet is slow, and will display **GREY** bars when there is no internet connection.
- Service Data Key Parameters: Displays the SQA-iO calibration and self-test parameters. A
 GREEN checkmark icon indicates everything is within normal limits, YELLOW indicates borderline
 limits and RED indicates out of range. Click on the checkmark for details and the REPORT button to
 run a calibration report for your records.
- **Test Status:** Provides the current status of the # of remaining test as well as a graph of tests run per week.



Test Patient

Select the TEST TYPE based on the following sample definitions/options:

- FRESH Sample is not enriched, diluted or treated and is within 1 hour of collection. Required testing volume is ≥0.6 ml (the entire testing capillary needs to be filled) or, if less sample is available, the sample can be diluted 1:2 [1+1] for a full report of all semen parameters. A 10-microliter sample can be loaded into just the thin capillary section for a limited report of just motility parameters.
- WASHED Sample is enriched or prepared for artificial insemination by centrifugation using a washing media to replace the seminal plasma. Testing volume required is ≥ 0.6 ml (the entire testing capillary needs to be filled) or, if less sample is available, the sample can be diluted 1:2 [1+1] for a full report of all semen parameters. A 10-microliter sample can be loaded into just the thin capillary section for a limited report of just motility parameters.

Enter patient and sample data in the TEST PATIENT screen seen below. Mandatory fields are indicated by an asterisk *, and an error message will appear if empty. The Collected/Received Date and time will be filled automatically according to the current Date and time of the test and can be edited.

PLEASE NOTE: Although sample volume is not mandatory, semen parameters related to sample volume will not be presented if the volume is not entered. The accuracy of the operator is relied upon to correctly measure sample volume.

SQAiO	≡ •←			o 🤰 🗈 0 4 🎳 틀 🛛
	Test Patient & / Home / Test Proce	255		
	FRESH WASHED			
Dr. J. Smith	PATIENT INFORMATION			
😤 номе	PATIENT ID * 4435353 ~	FIRST NAME John	LAST NAME Doe	SAMPLE ID 454546
	AGE 18	PHONE NUMBER N/A	ABSTINENCE (days) 7	REFERRING DOCTOR
	SAMPLE INFORMATION			
T ARCHIVE	COLLECTED DATE AND TIME	RECEIVED DATE AND TIME	VOLUME (ml)	WBC CONC. (M/ml) *
PATIENT INFO	13/02/2022	13/02/2022	5	*
⋛ SERVICE	рН 7.2	APPEARANCE Other +	NORMAL	LIQUEFACTION O-30 Minutes
🔅 SETTINGS	OPTIONAL 1	OPTIONAL 2		
	TESTER INFORMATION			
	TESTER NAME John Doe	TITLE (DESIGNATION)	COMMENTS	
			VIEW	7 SAMPLE 1+1 DILUTION 10 µJ TEST NOW

OPTIONAL 1 and OPTIONAL 2 are 'open' fields to input any information desired.

After entering patient data, select the type of test to be run:

Sample Handling and Testing Options:

- **Sample Handling:** Samples need to be completely liquefied and run within one hour of collection so that motility parameters are reported accurately. Always maintain samples at room temperature prior to and during testing; Excessive heat and/or cold will shock sperm cells and affect motility. See Appendix section for semen collection, capillary filling, and sample testing guidelines.
 - <u>Temperature Control</u>: Maintain sample at room temperature (20-25°C / 68-77°F). Do not heat as excessive heat will deplete sperm resources and cold will shock sperm cells and affect motility.



- <u>Sample Collection</u>: See Appendix section for guidelines about semen sample collection and for instructions on how to fill the testing capillary and insert it into the SQA-iO.
- <u>Sample Liquefaction</u>: Samples need to be completely liquefied and run within one hour of collection as motility parameters can decline over time. QwikCheck Liquefaction vials can be used to liquify semen samples prior to aspirating into the testing capillary, if needed.

• Measuring Sample Volume:

- \circ $\,$ When to measure: After sample liquefaction and before testing, measure according to laboratory procedures.
- Entering sample volume into the SQA-iO: Enter volume in the "Test Patient" screen of the SQA-iO.
- WBC / pH: Assess pH and WBC prior to testing using QwikCheck WBC/pH test strips.
- **1:2 (1+1) Dilution:** 0.3 to 0.5 ml of sample required. Dilute sample 1:2 (1+1) using the QwikCheck[™] Dilution kit reagent. 1+1 dilution requires equal amounts of sample and diluent (i.e. If the total sample volume is 0.4 ml, add 0.4 ml of dilution media). Operator sample dilution errors will result in inaccurate results.
- Low Volume Sample /10 microliter: Fill only the tip (motility channel) of the capillary using 10µl of sample. A limited test report with motility parameters only will be provided.
- WASHED samples: select to run NORMAL or LOW VOLUME (10µl samples).

Test Patient – Test Results

Click **TEST NOW** and insert the testing capillary when prompted. 0.6 ml of sample is required. Do not move the device during testing. After approximately 75 seconds, all sperm parameter results will be displayed. An indicator arrow will appear if the results are high or low based on the laboratory's approved reference values and protocols for results interpretation. If there is no arrow, the test results are either in the normal range or there is no reference value for the parameter.

TEST RESULTS				SAMPLE INFORMATION	
PARAMETER	RESULT	REF VALUE	STATUS	TEST TYPE:	WAHSED
CONCENTRATION (M/ml)	64.4	>= 16		SAMPLE ID:	454546
MOTILITY (%)	34	>= 42		COLLECTED DATE TIME:	20/05/2020 10:00
PROGRESSIVE (%)	19	>= 30		RECEIVED DATE TIME:	20/05/2020 10:20
RAPIDLY PROGRESSIVE (%)	3			TEST DATE LTIME:	10/05/2020 11:00
SLOWLY PROGRESSIVE (%)	16				10.00.2020 11.00
NON-PROGRESSIVE (%)	15	<= 1		CRITERIA:	WHO 6"
IMMOTILE (%)	66	<= 20		SAMPLE TESTED:	Full Volume
NORMAL FORMS (%)	13	>= 4		VOLUME (ml):	0.50
MOTILE SPERM CONC. (M/ml)	9.0			WBC CONC. (M/ml):	<1
PROG. MOTILE SPERM CONC. (M/ml)	9.0			OPTIONAL 1:	Clear seminal plasma
RAPID PR. MOTILE SPERM CONC. (M/ml)	5.8			OPTIONAL 2	
SLOW PR. MOTILE SPERM CONC. (M/ml)	3.2				laha Daa
FUNCTIONAL SPERM CONC. (M/ml)	NA			TESTER NAME:	John Doe
VELOCITY (mic/sec)	NA			TITLE (DESIGNATION):	Lab Manager
SPERM MOTILITY INDEX	58			COMMENTS: Send results to John Doe	SAV
SPERM # (M/ejac)	32.2	>= 39	1		
MOTILE SPERM (M/ejac)	4.8				
PROG. MOTILE SPERM (M/ejac)	3.2				
FUNCTIONAL SPERM (M/ejac)	NA				
MORPH. NORMAL SPERM (M/ejac)	6.1				
MES parameters are indicated by an asterisk					
Please note, some results were manually valid	lated by the low	-quality counter.			
LOW QUALITY COUNTER					
MOTILITY GRAPH					
(15%)					
(16%)-					
(24)					
(3%)-(66%)					
Immotile (%) Rapidly Progressive (%)					



Test Results: The table above will be displayed after testing **FRESH** and **WASHED** semen samples with normal testing volume, 10 µl or diluted 1:2 (1+1). Five navigation options are available from the TEST RESULTS screen:

- **RETEST:** Select to run a second test on the same patient.
- **VIEW REPORT:** Click to view the patient's test report.
- DOWNLOAD REPORT: Click to download and print the patient's test report.
- **CAPTURE (Requires SQA-VU):** Attach up to 10 images to the report. The capture option allows image/video viewing, deleting, downloading.
- MORPHOLOGY (Normal Forms): Connect the SQA-VU to manually assess Normal/Abnormal sperm.
- **REPORT EDITS:** After testing, click on PATIENT NAME/REFERRING DOCTOR/BIRTHDATE or AGE to edit.

EDIT PATIENT INFORMATION	
First Name: John	
Last Name: _Doe	
Ref Doctor: J. Smith	
Birth Date: 17/03/1983	

Low Quality – Test Results

Low quality test results may be reported as < (less than) or > (greater than) when one or more of the parameters falls below the SQA-iO dynamic range. Only Sperm Concentration, Total Motile, Motile Sperm Concentration and SMI values will be reported automatically due to the limited number of sperm cells, very low motility and/or poor morphology. Manual results can be entered to provide a full report if desired.

TEST RESULTS					SAMPLE INFORMATION	
PARAMETER	RESULT	REF VALUE	STATUS		TEST TYPE:	FRESH
CONCENTRATION (M/ml)	< 2.0	>= 16			SAMPLE ID:	454546
MOTILITY (%)	0	>= 42			COLLECTED DATE TIME:	20/05/2020 10:
RAPIDLY PROGRESSIVE (%)	N/A				RECEIVED DATE TIME:	20/05/2020 10:
SLOWLY PROGRESSIVE (%)	N/A				TEST DATE TIME:	10/05/2020 11:0
NON-PROGRESSIVE (%)	N/A					WILO STU
IMMOTILE (%)	N/A	<= 20			CRITERIA:	WHO 6TH
NORMAL FORMS (%)	N/A	>= 4			SAMPLE TESTED:	10 µl
MOTILE SPERM CONC. (M/ml)	< 0.2				VOLUME (ml):	2
RAPID PR. MOTILE SPERM CONC. (M/ml)	N/A				WBC CONC. (M/ml):	<1
SLOW PR. MOTILE SPERM CONC. (M/ml)	N/A				pH:	7.5
FUNCTIONAL SPERM CONC. (M/ml)	N/A				APPEARANCE:	Normal
VELOCITY (mic/sec)	N/A	>= 5			VISCOSITY:	Abnormal
SPERM MOTILITY INDEX	0				HOUSEACTION	blassal
SPERM # (M/ejac)	N/A	>= 39			LIQUEFACTION:	Normai
MOTILE SPERM (M/ejac)	N/A				ABSTINENCE (Days):	3
PROG. MOTILE SPERM (M/ejac)	N/A				OPTIONAL 1:	Clear seminal plas
FUNCTIONAL SPERM (M/ejac)	N/A				OPTIONAL 2:	
MORPH. NORMAL SPERM (M/ejac)	N/A					John Dee
*Automated test results indicate a low-qua a manual analysis or refer the patient for a	lity sample. For a co full laboratory sem	omplete report, we en analysis.	recommend per	forming	TITLE (DESIGNATION):	Lab Manager
SUBMIT MANUAL RESULTS					COMMENTS:	
MOTILITY GRAPH						
Motility graph is not available due to the low qualit	v of the sample.					



Low Quality – Manual Results

Manual results can be added to the test report to supplement the motility values reported in the automated lowquality test. A proficient semen analysis laboratory equipped with apparatus for sperm concentration, motility, and (optional) morphology testing, is required. Please note that the accuracy and precision of the manual results will rely on the proficiency of the operator and accurate reporting is the operator's responsibility.

ATENTID: 4453535 PATEN	INAME. JOIN DOE BIRTH D	TE / AGE. 1//03/1386 52 PROVE NOWIDER: 813 3041 423 REPERRING DOCTOR: JUIN DUE	
SUBMIT MANUAL RESULTS C	Criteria: WHO 6th EDITION	MANUAL TESTING GUIDE	ons
CONCENTRATION (M/ml)	RESULTS	When do I have the option to add manual sperm results?	^
CONCENTRATION *	10	 When the SQA-iO reports very low-quality automated results: Concentration < 2.0 M/ml and/or Motile Sperm Concentration (MSC) < 0.2 M/ml. 	1
IOTILTY PARAMETERS (%)		- For a more complete sement analysis report.	
/IOTILITY *	20	Will I be able to change my results after exiting this screen?	\sim
APIDLY PROGRESSIVE *	15	How do I know if my manual count is accurate?	\sim
LOWLY PROGRESSIVE *	1	TOTAL MOTH ITY (RAPIDLY PROG + SLOWLY PROG + NP): RAPIDLY AND SLOWLY PROG quidelines for manual	~
IORPHOLOGY (%)		testing	
IORMAL FORMS	13	MORPHOLOGY NORMAL FORMS guidelines for manual testing	\sim
No sperm seen		CONCENTRATION guidelines for manual testing	\sim
		Show less A	
ese semen parameters are requ	ired for a complete report	SUBIVITI RESULTS CLEAR SKIP TO AUTO RES	ULI

NORMAL FORMS (Morphology) are not included in the LOW QUALITY report unless manually assessed.

TEST RESULTS	SAMPLE INFORMATION				
PARAMETER	RESULT	REF VALUE	STATUS	TEST TYPE:	FRESH
CONCENTRATION (M/ml)	10.2	>= 16	4	SAMPLE ID:	454546
MOTILITY (%)	20	>= 42	4	COLLECTED DATE AND TIME	20/05/2022 10:00
PROGRESSIVE (%)	5	>= 30	4	RECEIVED DATE AND TIME	20/05/2022 10:20
RAPIDLY PROGRESSIVE (%)	2				20/03/2022 10:20
SLOWLY PROGRESSIVE (%)	3			TEST DATE AND TIME:	10/05/2022 11:00
NON-PROGRESSIVE (%)	15	<= 1		CRITERIA:	WHO 6TH
IMMOTILE (%)	80	r= 20		SAMPLE TESTED:	10 µl
NORMAL FORMS (%)	N/A	<u>2</u> 0		VOLUME (ml):	0.5
MOTULE SPERM CONC * (M/ml)	9.7			WBC CONC. (M/ml):	<1
PROG_MOTILE SPERM CONC.* (M/ml)	6.4			pH:	7.5
RAPID PR. MOTILE SPERM CONC. (M/ml)	1.3			pro	1.0
SLOW PR. MOTILE SPERM CONC.* (M/ml)	5.1			APPEARANCE:	Normal
FUNCTIONAL SPERM CONC.* (M/ml)	N/A			VISCOSITY:	Abnormal
VELOCITY (VCL)* (mic/sec)	N/A	>= 5		LIQUEFACTION:	Normal
SPERM MOTILITY INDEX*	0			ABSTINENCE (Days):	3
SPERM # (M/ejac)	32.2	>= 39	1	OPTIONAL 1:	Clear seminal plasm
MOTILE SPERM* (M/ejac)	16.1				
PROG. MOTILE SPERM* (M/ejac)	23.4			OPTIONAL 2:	
FUNCTIONAL SPERM* (M/ejac)	N/A			TESTER NAME:	John Doe
MORPH. NORMAL SPERM* (M/ejac)	N/A			TITLE (DESIGNATION):	Lab Manager
MES parameters are indicated by an asterisk				COMMENTS:	
Please note, some results were manually validated					SAV

* Motility and Concentration results cannot be submitted after leaving the manual assessment or test results page. Normal forms can be added at any time from the Patient Data Archive if manual results were entered for the other parameters.

Test Patient – Semen Analysis Report

Test reports format options are available in SETTINGS:

- Graph report: Two-page report with Motility Graph, editable header/footer and signature section with the option to include additional information and edit or remove the email address.
- Standard report: One-page report with editable header/footer re-sizing and the option to edit or remove the email address.
- Flexible report Can be customized by downloading and modifying an HTML template.



Graph Report - page 1

CA 90000 University Laboratory					EMAIL: mes@gmail.	
		~=~~=				
SQA-IO AUTO SC	A-iO MES - S	SEIVIE ignal Proc	N AN essing 1	ALY:	SIS RESULIS logy	
PATIENT INFORMATION						
FIRST NAME: John			LAST NA	ME:	Doe	
PATIENT ID: 4435353					32	
REFERRING DOCTOR. J Smith			PHONE	NUMBER	R: 546-6784-222	
SAMPLE INFORMATION						
SAMPLE ID: 454546			pH:		7.5	
SAMPLE TYPE: FRESH				APPEARANCE: NORMAL		
COLLECTED DATE / TIME: 13/02/2022 10:30				VISCOSITY: NORMAL		
RECEIVED DATE / TIME: 13/02/2022 11:00				LIQUEFACTION: 0-30 Minutes		
TEST DATE / TIME: 13/02/2022 11:19				ABSTINENCE (days): 7		
				ays): 7		
GRITERIA: WHO 6TH			OPTIONA	L 1:	ays): 7 Very clear seminal plasma	
VOLUME (ml): 5				L 1: L 2:	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction	
CRITERIA: WHO 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1			OPTIONA OPTIONA	NL 1: NL 2:	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction	
URITERIA: WHO 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT	UNITS	OPTIONA OPTIONA REF V	AL 1: AL 2: ALUE	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WHO 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7	UNITS M/ml	OPTIONA OPTIONA REF V >=16	AL 1: AL 2: ALUE	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28	UNITS M/ml %	OPTIONA OPTIONA REF V >=16 >=42	AL 1: AL 2: ALUE	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (8%)	
URITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20	UNITS M/ml %	OPTIONA OPTIONA REF V >=16 >=42	AL 1: AL 2: ALUE	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (8%) (12%)	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8	UNITS M/ml % %	OPTIONA OPTIONA REF V >=16 >=42	LL 1: LL 2: ALUE ↓	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (8%) (12%)	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12	UNITS M/ml % % %	OPTIONA OPTIONA REF V >=16 >=42	LL 1: LL 2: ALUE	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8	UNITS M/ml % % % %	OPTIONA OPTIONA REF V >=16 >=42	AL 1: AL 2: ↓	ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72	UNITS M/ml % % % % % % % % % % % %	OPTIONA OPTIONA REF V >=16 >=42	AL 1: AL 2: ↓ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72	UNITS M/ml %	OPTIONA OPTIONA PTIONA PTIONA REF V >=16 >=42 <=1	ALUE ↓ ↑ ↑	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3	UNITS M/ml % % % % % %	OPTIONA OPTIONA REF V >=16 >=42	ALUE ↓ ↑ ↑	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3 7.0	UNITS M/ml % % % % % % % % %	OPTIONA OPTIONA REF V >=16 >=42 <=1	AL 1: AL 2: ↓ ↓ ↑ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3 7.0 4.9	UNITS M/ml %<	OPTIONA OPTIONA REF V >=16 >=42 <=1	ALUE ↓ ↑ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (72%) Rapidly Progressive (%) Slowly Progressive (%)	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3 7.0 4.9 1.9	UNITS M/ml %<	OPTIONA OPTIONA REF V >=16 >=42 <=20	ALUE ↓ ↑ ↑ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (72%) Rapidly Progressive (%) Slowly Progressive (%) Non-Progressive (%)	
CRITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3 7.0 4.9 1.9 3.1	UNITS M/ml %<	OPTIONA OPTIONA >=16 >=42 <=1 <=20 >=4	AL 1: AL 2: ↓ ↓ ↑ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (72%) Rapidly Progressive (%) Slowly Progressive (%) Non-Progressive (%) Immotile (%)	
CKITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3 7.0 4.9 1.9 3.1 0.4	UNITS M/ml % <td>OPTIONA OPTIONA REF V >=16 >=42 <=1</td> <=20	OPTIONA OPTIONA REF V >=16 >=42 <=1	ALUE ↓ ↑ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (12%) (72	
CKITERIA: WH0 6TH VOLUME (ml): 5 WBC CONC. (M/ml): <=1	RESULT 24.7 28 20 8 12 8 72 3 7.0 4.9 1.9 3.1 0.4 31	UNITS M/ml % <td>OPTIONA OPTIONA REF V >=16 >=42 <=20</td> >=4 >=4 >=5	OPTIONA OPTIONA REF V >=16 >=42 <=20	ALUE ↓ ↑ ↑ ↓	Ays): 7 Very clear seminal plasma QwikCheck used for liquefaction MOTILITY GRAPH (12%) (72%) Rapidly Progressive (%) Slowly Progressive (%) Slowly Progressive (%) Non-Progressive (%) Immotile (%)	

SQA-iO Device SR: 10111 | Conc. Standard 1 | 20/05/2020 11:30:01 | AVG 55.81 | AW 15427 | CNT 330 | OD 1.126



Graph Report – page 2

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REPORT CONTINUED | PATIENT ID: 134145 | TEST DATE / TIME: 13/02/2022 | 11:19

TOTALS PER EJACULATE	RESULT	UNITS	REF V	ALUE
SPERM #	74.0	M/ejac	>=39	
MOTILE SPERM*	21.0	M/ejac		
PROG. MOTILE SPERM*	14.8	M/ejac		
FUNCTIONAL SPERM*	1.1	M/ejac		
MORPH. NORMAL SPERM*	2.2	M/ejac		

COMMENTS: Very clear seminal plasma, QwikCheck device used for liquefaction of this sample.

SMI (Sperm Motility Index) A CONTRIBUTING PREDICTOR OF MALE FERTILITY

What is SMI?

• Sperm Motility Index (SMI) is an MES SQA automatically generated index integrating motile sperm concentration and curvilinear velocity (VCL) into one value. It is reported in whole numbers without units.

• The positive correlation of SMI to male fertility was established and described in a number of publications.

SMI Grading and Male Fertility Potential:

• Studies have shown the correlation of SMI to male fertility potential as follows: 'LOW' (SMI < 80), 'MODERATE'

(SMI = 80-160) and 'HIGH' (SMI > 160) with an SMI reference value (cut-off) of 80.

• Conclusions: In conjunction with other semen parameters, SMI can be utilized to assess/grade male fertility potential.

Ant + Signature:

Tester Name: John Doe

Title (Designation): Lab Technician

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Standard report

PHONE: EMAIL: WEBSITE:	837-6029-686 MES@gmail.c www.mes-glob	<u>om</u> al.com	MES GL 5000 West CA 90000 Unive	OBAL Street LA. rsity Laboratory	Hedical Electronic Sys	Stems P	Page 1 d
PATIENT INF	FORMATION	SQA-iO AUTO	MATED SEN	/IEN ANALYSIS R	ESULTS		
FIRST NAME:		John		LAST NAME:	Doe		
PATIENT ID:		4435353		BIRTH DATE AGE:	17/03/1988 32		
SAMPLE INF	ORMATION						
SAMPLE ID:		454546		TEST RUN BY:	J. Smith		
IEST TYPE:		FRESH		APPEARANCE:	NORMAL		
	DATE / TIME:	20/05/2023 10:00		VISCOSITY:	ABNORMAL		
	ALE / HIME:	20/05/2023 10:20		LIQUEFACTION:	0-30 Minutes		
	(deve)	20/05/2023 11:00			WHO 6th	мп	
ABSTINENCE	(days):	3		SAMPLE TESTED:	NORMAL VOLU		
OPTIONAL 1:		QwikCheck used for lic	uefaction	OPTIONAL 2:	Very clear semir	nal plasma	
PARAMETEI	R		RESULT	UNITS	REF.VALUE	STATUS	
VOLUME			6	ml			
pH:			4				
WBC CONC.			<1	M/ml			
CONCENTRA	TION		6.0	M/ml	>=16	<u> </u>	
MOTILITY			34	%	>=42	Ļ	
PROGRE	SSIVE		11	%	>=30		
RAPIE	DLY PROGRESSI	VE	3	%			
SLOW	ILY PROGRESSI	/E	16	%			
NON-PRO	GRESSIVE		15	%	<=1		
IMMOTILE			66	%	<=20		
NORMAL FOR	RMS		3	%	>=4	Ļ	
MOTILE SPER	RM CONC.*		2.0	M/ml			
PROG. MO	OTILE SPERM CO	DNC.*	1.0	M/ml			
RAPIE	PR. MOTILE SP	ERM CONC.*	0.2	M/ml			
SLOW	PR. MOTILE SP	ERM CONC.*	1.0	M/ml			
FUNCTIONAL	SPERM CONC.*		N/A	M/ml			
VELOCITY (V	CL)*		N/A	mic/sec	>=5		
SPERM MOTI	LITY INDEX*		0				
SPERM #			18.0	M/ejac	>=39	Ļ	
	RM*		6.1	M/ejac			
PROG. MOTI	_E SPERM*		3.4	M/ejac			
FUNCIONAL S	SPERM*		N/A	M/ejac			
				•			

*MES parameters are indicated by an asterisk

COMMENTS: Very clear seminal plasma, QwikCheck device used for liquefaction of this sample.

FAC ID#: Z9XQWR | SN#: 10111 | [MA] | Conc. Standard 1 | 20/05/2020 11:30:01 | AVG 55.81 | AW 15427 | CNT 330 | OD 1.126

Patient Information

- **ADD NEW** patients by accessing the PATIENT INFORMATION screen.
- **Click ACTION** to edit or delete patient information.
- **SORT** by clicking on the column header.

v 10 ¢ entries Sea								arch:		
ACTIONS	PATIENT ID	FIRST NAME	LAST NAME	BIRTH DATE	PHONE NUMBER	MOST RECENT	WEIGHT (kg)	HEIGHT (cm)		
∕ ∎ 4	12345	Patient	One	03/07/1988	184732563	29/07/2020 09:31	78	187		
/ = 4	12912	Patient	Two	13/08/1987	2243254354	03/08/2020 09:39	92	165		
/ = 4	15774	Patient	Three	17/03/1983	954115651 <mark>1</mark>	Not entered	85	178		
/ = 4	18975	Patient	Four	12/03/1970	8896870070	14/08/2020 06:54	63	170		
/ = 4	19971	Patient	Five	08/02/1980	7063790690	18/08/2020 15:38	60	175		
∕∎ 4	20231	Patient	Six	17/10/1971	3123213123	05/10/2020 14:42	80	181		
/ = 4	22229	Patient	Seven	11/06/1987	23415325	Not entered	71	165		

Archive

- Click ARCHIVE for a list of all patient's test results.
- **SORT** by selecting the patient and then clicking on the ACTION button for date range, view, delete, or reports.

SQA-IO	≡ •<					و ©	D 0 4 🍈 틀 :
Dr. Yamir Sharma	Archive A Home /	Archive vikCheck QC BEADS	PROFICIENCY TESTS	FOR	USER GUIDE		
A HOME	SELECT DATE RANGE	T0 <u>18/09/2</u>	2023 APF	CLEAR			Search:
	ACTIONS	PATIENT ID	PATIENT NAME	TEST DATE TIME	TEST TYPE 10	SAMPLE ID	TESTER NAME
archive	⊙ ∎ <u>D</u> ± 0	4435353 321116	Patient One Patient Two	10/05/2023 11:00 17/09/2023 09:36	FRESH (DILUTED 1+1)	6464 454547	John Doe John Doe
PATIENT INFO	• 🖬 🚺 🚣	8787867	Patient Three	18/09/2023 10:06	WASHED (10 µl)	454548	John Doe
⋛ SERVICE	o î 🖟 🛓	54534	Patient Four	07/06/2023 15:43	WASHED	454546	John Doe
🔅 SETTINGS	o i 🏼 🕹 🗅	433447	Patient Five	06/04/2023 11:00	WASHED (10 µl)	32323	John Doe
	o 🕯 🔓 🕹 🙆	65656	Patient Six	01/07/2023 13:00	FRESH	545353	John Doe
	o i 🎝 🕹	766590	Patient Seven	10/05/2023 10:34	FRESH (DILUTED 1+1)	42434	John Doe
	o i 🖟 🛃	343412	Patient Eight	23/05/2023 11:30	WASHED (DILUTED 1+1)	4544346	John Doe
	Showing 1 to 8 of 8 entries	i					PREVIOUS 🚺 NEXT

SECTION 6: QC / CONTROLS and Proficiency

Select QC/Proficiency from the navigation panel to run three levels of QwikCheck Beads quality control samples or perform Proficiency testing. When running QwikCheck Beads controls or Proficiency samples please follow the instructions in the package insert. Also, be sure to:

- Use a separate, new capillary for each beads level.
- Mix the samples gently before aspirating into the testing capillary.
- Do not return beads solution to the container after testing this will contaminate the samples AND beads adhere to the capillary walls so the concentration of the beads will be altered.

Testing QC Controls

- The screen below will be activated when entering QC/Proficiency > QC from the navigation panel. If controls have never been run, all TEST RESULTS and SAMPLE INFORMATION will be shown as PENDING.
- **SELECT BATCH NUMBER:** From the drop-down menu, find the batch number that corresponds to the batch number on the outside label of the QwikCheck beads box that will be tested.
- **SAMPLE INFORMATION:** All three levels of beads will automatically be filled when the batch is selected.
- LAST RUN: If previous tests have been run, the last test date and time is shown.
- **TEST NOW:** Select TEST NOW when the testing capillaries are prepared for each test.

QwikCheck QC BEADS PROFICIENCY TESTING LAST RUN: 03-MAY-2020 10:43 O						
SELECT BATCH NUMBER: 011217 ▼ 0						
LEVEL 1	LEVEL 2	NEGATIVE CONTROL				
TEST NOW	TEST NOW	TEST NOW				
TEST RESULTS	TEST RESULTS	TEST RESULTS				
CONC (M/mi): PENDING STATUS: PENDING RUN DATE: PENDING	CONC (M/ml): PENDING STATUS: PENDING RUN DATE: PENDING	CONC (M/ml): PENDING MSC (M/ml): PENDING STATUS: PENDING RUN DATE: PENDING				
SAMPLE INFORMATION	SAMPLE INFORMATION	SAMPLE INFORMATION				
LOT #: 011217001	LOT #: 011217002	LOT #. 011217003				
EXP DATE: DEC / 2018	EXP DATE: DEC / 2018	EXP DATE: DEC / 2018				
TARGET (M/ml): 44	TARGET (M/ml): 22	TARGET (M/ml): 0.0				
VALUE (+/-): 6.2	VALUE (+/-): 4.4	VALUE (+/-): 0.0				
PASS RANGE: 37.8 - 50.20	PASS RANGE: 17.6 - 26.4	PASS RANGE: 0.0				
*Please check sample information with the beads contained	er label before running a test	QC ARCHIVE REPORT				

Follow the on-screen instructions for capillary preparation and insertion.

UTO CALIBRATION	INSERT CAPILLARY
()	×
SYSTEM IS CALIBRATING DO NOT INSERT CAPILLARY!	LOT #011214001 / LEVEL 1
MIX THE SEMEN SAMPLE THOROUGHLY	INSERT THE CAPILLARY NOW PRESS "TEST NOW" TO START THE ANALYSIS
FILL THE TESTING CAPILLARY CLEAN, WIPE AND INSPECT THE CAPILLARY FOR BUBBLES	TEST NOW CANCEL



Results and Corrective Action:

- **RESULTS:** Control testing takes about 20 seconds per test. Results are displayed automatically and, if out of range, a CORRECTIVE ACTION alert will be shown. Select the CORRECTIVE ACTION button to identify what caused the out-of-range results.
- **RE-TEST:** This button will appear after the first test was performed. Select it to test the sample again with no extra charge of credit code. The re-test option is time limited.

SELECT BATCH NUMBER: 061122 - 0		
LEVEL 1	LEVEL 2	NEGATIVE CONTROL
RE-TEST	RE-TEST	RE-TEST
TEST RESULTS CONC (M/ml): 31 STATUS: FAIL RUN DATE: 03-FEB-2023 10:30	TEST RESULTS CONC (Mimi): 28.3 STATUS: PASS ⊘ RUN DATE: 03-FEB-2023 10.39	TEST RESULTS CONC (M/ml): 0.0 MSC (M/ml): 0.0 STATUS: PASS ♥ RUN DATE: 03-FEB-2023 10.43
SAMPLE INFORMATION	SAMPLE INFORMATION	SAMPLE INFORMATION
LOT #: 061122001	LOT #: 061122002	LOT #: 061122003
EXP DATE: NOV / 2023	EXP DATE: NOV / 2023	EXP DATE: NOV / 2023
TARGET (M/ml): 47	TARGET (M/ml): 26	TARGET (M/ml): 0
VALUE (+/-): 6.6	VALUE (+/-): 5.2	VALUE (+/-): 0
PASS RANGE: 40.4 - 53.6	PASS RANGE: 20.8 - 31.2	PASS RANGE: 0.0 - 0.0
CORRECTIVE ACTION		

• **CORRECTIVE ACTIONS** are listed below and once selected, will appear on the QC Report and will be saved in the QC archive. Use the USER DEFINED option if none of the actions listed describe the problem.

RRECTIVE ACTION		
PROBLEM DESCRIPTION	CORRECTIVE ACTION	
SYSTEM REQUIRES CLEANING	CLEAN SYSTEM: RE-TEST	
CONTROL MATERIAL EXPIRED	RUN NEW BATCH OF CONTROLS	
SAMPLE HANDLING / MIXING	HOMOGENEOUSLY MIX: RE-TEST	
CONTROL IMPROPERLY STORED	RUN NEW BATCH	
WRONG LEVEL TESTED	RUN CURRECT LEVEL	
USER DEFINED	Maximum 30 characters	X

• **QC ARCHIVE:** Select from the TESTING or ARCHIVE screen to view all QC tests. Many options for selecting and presenting results are available from this screen and results can be exported.

PATIEN	TIENT DATA QwikCheck QC Beads PROFICIENCY TESTS		/ TESTS								
ILTER E	BY BATCH FILTER	BY LEVEL	FILTER BY	STATUS	FILTER B Please cl	Y DATE		To Pleas	e choose a da	ite	APPLY CLEAR
	RUN DATE / TIME	LEVEL	LOT #	EXP DATE	TARGET (M/ml)	VALUE (+/-)	PASS RANGE	CONC (M/ml)	MSC (M/ml)	STATUS	CORRECTIVE ACTION
	15-NOV-2020 10:06	1	230920001	OCT / 2021	46.0	6.2	41.3 - 54.7	50.3	NA	PASS	
	15-NOV-2020 10:15	2	230920002	OCT / 2021	22.1	4.1	20.0 - 30.0	23.3	NA	PASS	
	15-NOV-2020 10:20	NEG. CONTROL	230920003	OCT / 2021	0.0	0.0	0.0	0.0	0.0	PASS	
	01-APR-2020 09:15	1	051119001	NOV / 2020	24.1	6.5	20.0 - 30.0	23.3	NA	PASS	
	15-JUN-2020 15:43	1	230919001	AUG / 2019	48.0	4.4	41.3 - 54.7	30.7	NA	FAIL	RUN NEW BATCH
	18-OCT-2020 18:09	NEG. CONTROL	050319003	MAY / 2019	0.0	0.0	0.0	5	1.5	FAIL	CLEAN SYSTEM: RE-TEST
	07-DEC-2019 10:20	2	210918002	SEP / 2018	24.1	6.5	20.0 - 30.0	15.7	NA	FAIL	RUN CURRENT LEVEL



• **QC/Controls Test Report:** After testing, select REPORT to print a final report with QC results and graph.





Testing Proficiency Samples

- Select QC/Proficiency from the navigation panel, then activate the Proficiency Testing tab to view the screen displayed below.
- SCHEMES: There are four different schemes available to select from:
 - o NEQAS
 - o QuaDeGa
 - CAP/API
 - o iPRO
- **SELECT SCHEME:** From the drop-down menu, select the scheme in which the lab is enrolled.
- **SELECT DISTRIBUTION NUMBER:** For NEQAS and QuaDeGa, the distribution number can be found on the box labeling. Select the corresponding distribution number from the drop-down menu.
- **ENTER ISSUE DATE/BATCH NUMBER:** For CAP/API and iPRO, the issue date/Batch number can be found on the box labeling. Enter the information in the provided field.
- **SAMPLE INFORMATION:** NEQAS and QuaDeGa sample ID's will be shown automatically when the distribution number is selected. For CAP/API, manually enter the sample IDs found on the box labeling.
- LAST RUN: If previous tests have been run, a notification of the last date and time is shown.
- **TEST NOW:** Select TEST NOW when the testing capillaries are prepared for each test. Follow the onscreen instructions for capillary insertion.
- **RE-TEST:** This button will appear after the first test was performed. Select it to test the sample again with no extra charge of credit code.
- SUBMISSION DEADLINE: The date the proficiency results must be reported.
- **NOTE:** Enter sample testing notes after testing, if desired. Click SAVE to display notes on the report/archive or CLEAR to remove the notes.

QwikCheck QC Beads Proficiency Test			LAST RUN: 03 / FEB / 2023 10:43 ①
SELECT SCHEME: NEQAS	SELECT DISTRIBUTION NUMBER: 109	-	
SAMPLE #S433	SAMPLE #S434	SAMPLE #S435	SAMPLE #S436
TEST NOW	TEST NOW	RE-TEST	RE-TEST
TEST RESULTS	TEST RESULTS	TEST RESULTS	TEST RESULTS
CONC (M/ml): PENDING	CONC (M/ml): PENDING	CONC (M/ml): 26.7	CONC (M/ml): 18.7
RUN DATE: PENDING	RUN DATE: PENDING	RUN DATE: 03 / FEB / 2023 10:57	RUN DATE: 03 / FEB / 2023 11:05
SUBMISSION DEADLINE: 20 / FEB / 2023	SUBMISSION DEADLINE: 20 / FEB / 2023	SUBMISSION DEADLINE: 20 / FEB / 2023	SUBMISSION DEADLINE: 20 / FEB / 2023
NOTE:	NOTE:	NOTE: It is required to retest this sample after cleaning SAVE CLEAR	NOTE: SAVE CLEAR
* A note can be added after the proficiency test is	s performed.		PROFICIENCY ARCHIVE REPORT



Proficiency Results:

- **RESULTS:** Proficiency testing takes about 20 seconds per test. Concentration Results are displayed automatically. If the results indicate the SQA-iO was not cleaned effectively before testing, the results will be displayed in red, and a re-test option will be available after cleaning the device.
- **PROFICIENCY ARCHIVE:** Select this option from the TESTING or ARCHIVE screen to view all Proficiency Tests. Options for filtering, presenting/deleting or exporting results are available.

PATIEN	T DATA QwikCheck	QC Beads	PROFICIENCY	TESTS			
SCHEMI CAP/AP	E FILTER BY ISSUE	DATE	FILTER BY SAMPLE	E ID FILTER BY DATE	. To Please	e choose a date	APPLY CLEAR
	RUN DATE / TIME	SCHEME	ISSUE DATE	SAMPLE ID	CONCENTRATION (M/ml)	NOTE	
	09-DEC-2019 10:06	CAP/API	09-DEC-2019	11223344556677889900	42.0	Proficiency #1	
	09-DEC-2019 10:40	CAP/API	09-DEC-2019	11223344556677889911	39.1	Proficiency #2	
	30-OCT-2020 09:20	CAP/API	30-OCT-2020	71225332356677453300	PENDING		
							PREVIOUS 1 NEXT
						CREATE REPORT EXF	PORT DELETE

• **TEST REPORT:** After completing a test, select the REPORT button to view the final Report.

PROFICIENCY TESTING REPORT SQA-iO MES - Signal Processing Technology						
PROFICIENCY TESTI	NG INFORMTATIO	DN				
SCHEME: NEQAS			REF	PORT DATE / TIME	: 04-0CT-2022 11:05	
RUN DATE: 04-0CT-2022						
RUN DATE / TIME	DISTRIBUTION NUMBER	SAMPLE	RESULTS (M/ml)	SUBMISSION DEADLINE	NOTE	
RUN DATE / TIME 04-0CT-2022 10:47	DISTRIBUTION NUMBER 109	SAMPLE #S433	RESULTS (M/ml) 53.6	SUBMISSION DEADLINE 05 / OCT / 22	NOTE Retest this sample again	
RUN DATE / TIME 04-OCT-2022 10:47 04-OCT-2022 10:53	DISTRIBUTION NUMBER 109 109	SAMPLE #S433 #S434	RESULTS (M/ml) 53.6 23.0	SUBMISSION DEADLINE05 / OCT / 2205 / OCT / 22	NOTE Retest this sample again	
RUN DATE / TIME 04-OCT-2022 10:47 04-OCT-2022 10:53 04-OCT-2022 10:57	DISTRIBUTION NUMBER 109 109 109	SAMPLE #S433 #S434 #S435	RESULTS (M/ml) 53.6 23.0 26.7	SUBMISSION DEADLINE 05 / 0CT / 22 05 / 0CT / 22 05 / 0CT / 22	NOTE Retest this sample again	



SECTION 7: SQA-iO Test Credits

The SQA-iO cannot operate without test credits. Each new test kit or SQA Testing Capillaries box contains a unique TEST CREDIT CODE. Enter this code into the SQA-iO when opening a new test kit or when you receive an alert that test credits are low.

From the Home page SELECT:

- **ORDER SUPPLIES** to request SQA-iO supplies from your distributor.
- Click on **CREDIT CODES** if you need to load more tests.



SQA-iO supplies can also be ordered through **CONTACT US** using the convenient drop-down menu **by direct contact with your local distributor.**

Required service				
Order supplies			•	
Product	Un	its	Total	
SQA-iO Test Kit (50 tests)	• 3	¢	SQA-iO Test Kit (50 tests) X3	Х
SQA-iO Test Kit (50 tests)	1	÷	SQA-iO Test Kit (50 tests) X1	×
SQA-iO Device				
SQA-iO Cleaning Kit				
SQA Testing Capillaries				
QwikCheck Liquefaction kit	necting.			
QwikCheck Beads				
SQA-VU Device				
Other			Send	

SECTION 8: Set-up the SQA-iO Default Settings

Various levels of default SETTINGS can be implemented in the SQA-iO based on the user's permission status. Each user will have different permissions and his own login credentials (email and password).

TYPES OF USERS: Three types of users are described below along with their permission rights.

- BASIC Can view and modify the User Profile.
- **EDITOR** Can change the User Profile, Reference Values and Test Patient defaults.
- **ADMIN** Has access rights and can view/modify all Settings options and can add new, remove, or edit other user accounts. Admin users are limited to one per account.



TEST PATIENT (Editor Level Permission): Select **Settings -> Test Patient** to define the defaults for sample testing.

- **CONC. STANDARD**: Select "Standard 1" for 10-20-micron counting chambers (Makler) that do not require sample dilution; Select "Standard 2" for hemacytometers OR Neubauer.
- **APPEARANCE**: Use to select the color or Normal/Abnormal sample appearance.
- **MORPHOLOGY RANGE**: Set Morphology Upper Range Limit as an integer number from 10% through 30% based on the lab morphology assessment data. The default limit is 20%.
- **DEBRIS ASSESSMENT**: Select whether a manual Debris Assessment will be activated for every sample. See the Debris Assessment Protocol in the Appendix section of this guide.
- **OPTIONAL FIELDS**: Enter any labeling desired in any one of these fields. They will appear as labeled on the test report and on the data entry/patient testing screen.

Settings A / Home / Settings				
TEST PATIENT SYSTEM REF VALUE FACILITY PROFILE USER PROFIL	E USER MANAGEMENT			
SAMPLE INFORMATION	PATIENT INFORMATION			
CONC. STANDARD:	Patient Age O Patient Birth Date			
Conc. Standard 1 (Makler) Conc. Standard 2 (Hemocytometer/Neubauer)	◯ ft, in ● cm			
APPEARANCE:	🔿 lb 🛛 🕒 kg			
Specify sample color				
O Normal / Abnormal	WHO 6" TEST REPORT			
MORPHOLOGY RANGE (WHO 6 th only): Set upper limit (10% - 30%). 20	FIRST PAGE SECOND PAGE			
DEBRIS ASSESSMENT	Display page number			
Include Debris Assessment (Requires visual assessment)	Remove Signature			
Optional Field 1 [©]	Remove Tester Name and Title (Designation) Remove Header Blank Header space:mm			
Optional Field 2 $^{\odot}$	Remove Footer Blank Footer space:mm			
	Edit Email Address: O Remove Change:Dr.Dave@facility.com			
	DEFAULTS SAVE			

SYSTEM (Editor Level Permission): Select Settings -> System to define the system defaults.

- BEEP SOUND: Turn on or off beep that indicates when to insert capillary after auto calibration.
- **ARCHIVE**: The **page** of the last test run/reviewed will appear first when opening the Archive and the last run/reviewed **test** will be highlighted.
- **PHONE NUMBER:** Remove the patient phone number field from the TEST PATIENT and PATIENT INFO categories.
- **AUTOMATIC LOG-OUT**: Define time for the automatic log-out up to 12 hours max.
- SQA-VU SETTINGS: Any operator can adjust:
 - <u>Image format</u>: Change the IMAGE download format from PNG (default) to JPEG.
 - Low Quality Counter: Select to automatically open a manual counting screen for all low-quality samples.

TEST PATIENT	SYSTEM	REF VALUE	FACILITY PROFILE	USER PROFILE	USER MANAGEMENT
SYSTEM SETTING	SS				SQA-VU USERS
 Play a beep so Last test revie Remove patie Automatic Logout a 	ound when the c wed will be first nt phone numbe after: 00 hr : 30	apillary is ready to and highlighted in or mins 🗘 (Max 12 hi	be inserted into the chamt the Archive table (Patient I) ^①	ber Data tab)	Set image download format: PNG JPEG Open the Low-quality counter automatically



REF. VALUE (Editor and Admin Permission): Select WHO 5th of 6th edition testing criteria for reference values. The manufacturer's factory defaults are pre-set to WHO 6th criteria. Or, set custom reference values by un-checking the box.

FACILITY PROFILE (Administrator Permission): Select to customize the SQA-iO test report logo and facility information.

USER PROFILE (All Users): View personal profile information, change password, set-up test report signature and upload a personal profile picture.

SECTION 9: Service

Enter this screen to view/access the:

- **DISTRIBUTOR**: Link to your distributor for service and support by their unique ID number.
- MAINTENANCE CHECKLIST: Document and track the device maintenance and cleaning schedule.
- MAINTENANCE REPORT: Displays the most recent maintenance checklist.
- **SERVICE REPORT**: Provides technical information about the device.
- **SERVICE DATA/KEY PARAMETERS**: Check to confirm that the SQA-iO device is ready for testing.
- User Guide, Service Manual and Troubleshooting Guide: Links provided to review or download.

MAINTENANCE CHECKLIST		SERVICE DATA		
AST PERFORMED: 05-NOV-2023		KEY PARAMETERS:		ACCEPTABLE RANGE:
VERIFY SUPPLY INVENTORY	\checkmark	REFERENCE 1 (mV):	200 🥏	150mV - 350mV
CLEAN TESTING CHAMBER	v	LED CURRENT 1 (mA):	15 📀	5mA - 20mA
		REFERENCE 2 (mV):	3000 📀	2500mV - 3500mV
		LED CURRENT 2 (mA):	20 📀	10mA - 32mA
DUST TESTING CHAMBER		ZERO LEVEL:	515 📀	500 - 525
CONFIRM SYSTEM PASSED SELF-TEST	\checkmark	AMPLITUDE (mV):	80 📀	50mV - 100mV
PERFORMED BY:		SELF-TEST STATUS: PAS CALIBRATION AND STAE	SS BILIZATION: PASS	SELF-TEST SERVICE REPORT
QC CONTROLS: QwikCheck QC Beads	Last Run: 01/02/2021	RESOURCES/GUIDES		
PROFICIENCY: CAP/API Last Run 11223344556677889910 53.6 (11223344556677889911 23.0 (11223344556677889912 26.7 (: 16/05/2021 ///ml) ///ml)	<u>View User Guide</u> <u>View Service Manual</u> <u>View Troubleshooting</u>	Guide	ADD TEST CREDITS

SECTION 10: Help Center / Contact Us

TRAINING VIDEOS: Provide step-by-step instructions on the different features and processes of the SQA-iO. **FAQ**: Presents different troubleshooting questions and answers to solve technical problems.

GUIDES: Displays all SQA-iO guides for viewing or downloading.

CONTACT US: Click the phone icon at the top righthand corner of the screen or access from the Help Center to order new test kits or request support. Use the drop-down menu and message box to contact your local distributor.



APPENDIX 1: Filling the Testing Capillary with a Normal Volume Sample



Sample size, collection and preparation instructions:

- 1. A minimum of .6 ml. of semen is required for the SQA testing capillary.
- 2. Self-collected the sample without using lubricants/creams or partners.
- 3. Test the sample after liquefaction and within 1 hour of collection for optimal results.
- 4. Maintained at room temperature 20-25°C / 68-77°F (do not heat or refrigerate).
- 5. Measure sample volume according to laboratory protocols.
- 6. Before filling the capillary, mix the liquefied sample gently by rotating the sample collection container.
- 7. *WARNING:* Do not shake or use a pipette to mix the sample otherwise air bubbles will form and test results will be inaccurate.
- 8. Carefully check that the liquefied, fully mixed semen is **free** of air bubbles.

Filling the capillary... Ready to test:

- 1. Push the syringe pump fully into the syringe and then place only thin part of the capillary into the bottom of the sample (Fig 1).
- 2. Pull the syringe pump back slowly while keeping the tip of the capillary well below the sample level and below any surface bubbles. Continue to aspirate the sample until it appears in the Luer adaptor (Fig. 1 & 2).
- 3. Check the capillary after filling (Fig. 2), visually confirm that the sample has completely filled the cuvette and thin section of the capillary (without a meniscus). Tap on the syringe to make sure there are no air bubbles in the sample. If air bubbles still appear below the Luer adaptor, fill again with a small quantity of semen to draw the air bubbles into the syringe.
- 4. Wipe the tip of the capillary with a **Kimwipe** quickly (to avoid wicking) (Fig. 3). Also wipe the exterior of the capillary if any spillage occurred, in order to keep the SQA-iO clean. Visually **confirm** that the capillary chambers are still full after cleaning. If not, slightly push in the piston of the syringe to re-fill the capillary section.
- 5. Slowly push in the blue separating valve until it is level with the plastic (Fig. 4).
- 6. Insert the testing capillary into the SQA-iO **all the way** with the blue valve down (Fig 5)







Fig. 5: Insert capillary into SQA-iO



Fig. 1: Filling



Fig. 2: Inspect for bubbles



Fig. 3: Wipe the tip



APPENDIX 2: Filling the Testing Capillary with a LOW Volume Sample

Sample size and preparation:

- 1. A **minimum** of 10 microliters of semen can be tested by filling ONLY the thin section of the testing capillary. Only semen motility parameters will be reported.
- 2. The sample must be maintained at room temperature (do not heat or refrigerate), tested within 1 hour of collection and be fully liquified.
- 3. After liquefaction, gently mix the sample by rotating it in the container.
- 4. Carefully check that the liquified, fully mixed semen is free of air bubbles.

WARNING: Do not shake or use a pipette to mix the sample otherwise air bubbles will form and test results will be inaccurate.

Fill the SQA-iO testing capillary:

- 1. **Push the syringe piston in fully**. Place only the thin part of the capillary into the bottom of the sample (Figure 1).
- 2. **Pull the piston back slowly** without withdrawing the capillary from the sample.
- 3. **Fill only the (thin) capillary chamber** with 10 microliters of semen (Figure 1). Aspirate the sample until it just appears in the cuvette section while keeping the tip of the capillary well below the sample level and well below the level of any bubbles covering the liquid.
- 4. Withdraw the capillary tip from the semen sample and visually inspect to ensure that the sample has completely filled the thin section (no meniscus).
- 5. Wipe the tip of the capillary with a **Kimwipe** quickly (to avoid wicking). Also wipe the exterior of the capillary if any spillage occurred, in order to keep the SQA-iO clean.
- 6. Visually **confirm** that the thin section of the capillary is still full after cleaning. If not, **slightly** push in the piston of the syringe until a small drop appears on the capillary tip and then re-fill the capillary tip with more sample.

Remove the blue separating valve:

- Detach the entire syringe from the hub (Figure 2)
- Use the syringe or capillary jig to push-out the blue separating valve from the capillary (Figure 3)
- Completely remove the blue separating valve (Figure 4)
- Insert the testing capillary into the SQA-iO



Fig 2:Detach the syringe



Fig 3:Push the valve out



Fig 4:Remove the blue

PLEASE NOTE: Test Low Volume samples as soon as the capillary is filled.



Fig. 1: Fill the tip



APPENDIX 3: Cleaning the SQA-iO

When to clean: WEEKLY

- Or if SELF-TEST or any other failure occurs
- Or if System becomes contaminated with semen

Cleaning kit components:

Long cleaning brush (provided in the SQA-iO TEST KIT) Fibrous material cleaning paddles (single use) Sponge-tipped drying paddles (single use) Cleaning fluid (single drop dispenser)

CLEANING: STEP 1

- Insert the long brush supplied in your TEST KIT (bristle side down) into the chamber of the SQA-iO in the same way a testing capillary would be inserted (Fig 1 and 2).
- Pull the brush out, applying downward pressure to sweep or 'dust off' the optics (you will feel a 'shelf' in the back/top section of the chamber)
 – (Fig 2 and 3)

CLEANING: STEP 2

- 1. Use a **Fibrous material** cleaning paddle (Fig 4) supplied in your TEST KIT.
 - Moisten with only ONE drop of cleaning fluid.
 - Shake off excess fluid.
 - Insert into the measurement compartment fibrous material facing **down** and move the cleaning paddle in and out 5 times (Fig 5).
 - Then, insert into the measurement compartment fibrous material facing **up** and move the cleaning paddle in and out 5 times (Fig 5).
- 2. Dry the testing chamber using a sponge-tipped drying paddle that is supplied in your TEST KIT.
 - Insert it into the testing chamber and leave it for 10 15 seconds (Fig 6).
 - Leave the drying paddle in place, DO NOT move it in and out.





Fig. 2 Clean the chamber



Fig. 3 "Dust off"



Fig. 4 Fibrous cleaning paddle



Fig. 5 Insert cleaning paddle down and up



Fig. 6 Dry the testing chamber with sponge



APPENDIX 4: Reference Range Values of Semen Parameters

WHO 5 th		WHO 6 th		
SEMEN PARAMETER	REFERENCE RANGE*	SEMEN PARAMETER	REFERENCE RANGE*	SOURCE
CONCENTRATION (M/ml)	≥ 15	CONCENTRATION (M/ml)	≥ 16	WHO
TOTAL MOTILE PR + NP (%)	≥ 40	TOTAL MOTILITY (%)	≥ 42	WHO
PROGRESSIVE PR (%)	≥ 32	PROGRESSIVE (%) (RAPIDLY + SLOW PROG)	≥ 30	WHO
NON-PROGRESSIVE NP (%)	N/A	NON-PROGRESSIVE (%)	≤1	WHO
IMMOTILE IM (%)	N/A	IMMOTILE (%)	≤ 20	WIIO
MOTILE SPERM CONC. (M/ml)	≥ 6	MOTILE SPERM CONC. (M/ml)	≥ 7	MES
PROG. MOTILE SPERM CONC. (M/ml)	≥ 5	PROG. MOTILE SPERM CONC. (M/ml) (RAPIDLY + SLOW)	≥ 5	MES
NORMAL FORMS (%)	≥ 4	NORMAL FORMS (%)	≥ 4	WHO
CDEDM MOTH ITV INDEV**	> 80	FUNCTIONAL SPERM CONC. (M/ml)	≥ 0.2	WHO
SPERM MOTILITY INDEX	≥ 80	SPERM MOTILITY INDEX**	≥ 80	MES
SPERM # (M/ejac)	≥ 39	SPERM # (M/ejac)	≥ 39	MES
MOTILE SPERM (M/ejac)	≥ 16	MOTILE SPERM (M/ejac)	≥ 16	MES
		PROG. MOTILE SPERM (M/ejac)	≥ 12	MES
		FUNCTIONAL SPERM (M/ejac)	≥ 0.5	MES
		MORPH NORMAL SPERM (M/ejac)**	≥ 2	MES
		VELOCITY** (VCL) (mic/sec)	≥ 5	MES

* The reference values established above are based on WHO 5th/6th edition manual data or MES (for proprietary semen parameters). Each laboratory/clinic can establish their own requirements and cut-offs for semen parameters.

** Semen parameters not reported in U.S. market



APPENDIX 5: Product Performance Data:

Accuracy:

The SQA-iO WHO 6th accuracy vs. the SQA-V PREDICATE is established using Passing-Bablok regression analysis. The trendline slope, intercept and correlation Accuracy results are shown in Table 1 below.

Parameter	Intercept	CI	Slope	CI	Correlation	CI
CONCENTRATION, M/ml	-1.5	-2.0 to -0.7	1.0	1.0 to 1.0	1.0	0.98 to 0.99
MOTILITY, %	-3.0	-3.1 to -1.7	1.0	1.0 to 1.0	1.0	0.95 to 0.97
PROGRESSIVE MOTILITY, %	-0.8	-1.0 to 0.0	0.9	0.9 to 1.0	1.0	0.97 to 0.98
RAPIDLY PROGRESSIVE, %	0.1	0.0 to 0.3	1.0	0.9 to 1.0	0.9	0.90 to 0.94
SLOWLY PROGRESSIVE, %	-0.8	-1.0 to 0.0	1.0	0.9 to 1.0	0.9	0.86 to 0.93
NON-PROGRESSIVE, %	-1.9	-3.0 to -1.0	1.2	1.0 to 1.3	0.8	0.71 to 0.83
IMMOTILE, %	3.0	1.0 to 5.0	1.0	1.0 to 1.0	1.0	0.95 to 0.97
MSC, M/ml	-0.9	-1.7 to -0.6	1.0	1.0 to 1.0	1.0	0.98 to 0.99
PMSC, M/ml	-0.4	-0.7 to -0.3	1.0	0.9 to 1.0	1.0	0.99 to 1.00
RAPID PMSC, M/ml	0.0	-0.1 to 0.0	1.0	1.0 to 1.0	1.0	0.96 to 0.98
SLOW PMSC, M/ml	-0.1	-0.4 to -0.1	1.0	0.9 to 1.0	1.0	0.98 to 0.99
MORPHOLOGY, % (n = 155)	0.0	0.0 to 0.1	1.0	0.9 to 1.0	1.0	0.96 to 0.98
FSC, M/ml (n = 155)	-0.1	-0.1 to 0.0	0.9	0.9 to 1.0	1.0	0.97 to 0.99

Table 1. SQA-iO Intended User vs. SQA-V Expert User (n = 165)

Precision:

Table 1: SQA-iO Sperm Concentration Precision

Concentration		Within-Run		Between- Run		Between- Day		Between- Operator/ Lot/Instrumen t		Total		
Sample	N	Mean	SD	%CV	SD	%CV	SD	%CV	SD	%CV	SD	%CV
1	40	8.5	0.63	7.4%	0.61	7.2%	0.25	2.9%	0.60	7.1%	0.62	7.3%
2	40	34.5	1.66	4.8%	1.70	4.9%	0.77	2.2%	1.31	3.8%	1.76	5.1%
3	40	45.4	3.25	7.2%	3.30	7.3%	1.66	3.7%	3.09	6.8%	3.46	7.6%
4	40	58.5	3.12	5.3%	3.07	5.2%	1.04	1.8%	2.11	3.6%	3.04	5.2%
5	40	62.2	2.42	3.9%	2.38	3.8%	1.42	2.3%	2.30	3.7%	2.64	4.2%
6	40	181.6	5.25	2.9%	5.35	2.9%	3.42	1.9%	3.83	2.1%	5.87	3.2%
7	40	227.6	5.87	2.6%	6.25	2.7%	5.45	2.4%	3.48	1.5%	7.58	3.3%
8	40	212.9	3.74	1.8%	4.42	2.1%	4.87	2.3%	2.67	1.3%	5.79	2.7%

Table 2: SQA-iO Motility Precision

Motility		Within-Run		Between- Run		Between- Day		Between- Operator/ Lot/Instrument		Total		
Sample	Ν	Mean	SD	%CV	SD	%CV	SD	%CV	SD	%CV	SD	%CV
1	40	0.0	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%
2	40	77.0	2.82	3.7%	2.74	3.6%	1.20	1.6%	2.59	3.4%	2.87	3.7%
3	40	62.3	2.62	4.2%	2.59	4.2%	0.74	1.2%	2.27	3.7%	2.54	4.1%
4	40	80.6	0.99	1.2%	1.00	1.2%	0.46	0.6%	0.83	1.0%	1.01	1.3%
5	40	58.0	3.83	6.2%	4.65	7.7%	3.23	5.6%	2.60	4.5%	6.99	12.1%
6	40	43.9	1.81	4.1%	1.99	4.5%	1.18	2.7%	1.37	3.1%	2.04	4.6%
7	40	30.7	2.29	7.5%	2.52	8.3%	2.22	7.2%	0.94	3.1%	3.03	9.9%
8	40	49.9	1.52	3.0%	1.77	3.5%	1.52	3.0%	1.28	2.6%	2.05	4.1%

	MSC		Within-Run		Between- Run		Betwe	en-Day	Between- Operator/ Lot/Instrument		Тс	otal
Sample	N	Mean	SD	%CV	SD	%CV	SD	%CV	SD	%CV	SD	%CV
1	40	2.0	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%
2	40	26.5	1.31	5.0%	1.36	5.1%	1.05	4.0%	0.68	2.6%	1.60	6.0%
3	40	27.9	1.40	5.0%	1.55	5.5%	1.03	3.7%	1.08	3.9%	1.67	6.0%
4	40	47.0	2.99	6.4%	2.99	6.4%	1.13	2.4%	2.27	4.8%	2.97	6.3%
5	40	35.5	1.42	4.0%	1.56	4.4%	0.77	2.2%	1.27	3.6%	1.54	4.3%
6	40	79.4	2.87	3.6%	3.54	4.5%	2.41	3.0%	1.09	1.4%	3.60	4.5%
7	40	69.3	4.26	6.2%	5.05	7.3%	4.29	6.2%	1.37	2.0%	5.85	8.4%
8	40	106.2	3.43	3.2%	4.48	4.2%	5.30	5.0%	2.18	2.1%	6.12	5.8%

Table 3: SQA-iO Motile Sperm Concentration (MSC) Precision

Table 4: SQA-iO Progressively Motile Sperm Concentration (PMSC) Precision

PMSC		Within-Run Run		Between-Day		Between- Operator/ Lot/Instrument		Total				
Sample	N	Mean	SD	%CV	SD	%CV	SD	%CV	SD	%CV	SD	%CV
1	40	0.0	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%
2	40	23.2	1.11	4.8%	1.14	4.9%	0.94	4.1%	0.74	3.2%	1.38	6.0%
3	40	24.2	1.27	5.2%	1.35	5.6%	0.83	3.4%	0.90	3.7%	1.41	5.8%
4	40	42.2	2.80	6.6%	2.81	6.7%	1.16	2.8%	2.11	5.0%	2.82	6.7%
5	40	31.5	1.78	5.6%	1.86	5.9%	0.76	2.4%	1.11	3.5%	1.92	6.1%
6	40	70.3	2.64	3.8%	3.34	4.8%	2.34	3.3%	0.92	1.3%	3.40	4.8%
7	40	51.0	4.60	9.1%	5.34	10.6%	5.20	10.2%	2.51	4.9%	6.54	12.8%
8	40	93.4	3.58	3.8%	4.39	4.7%	5.32	5.7%	2.21	2.4%	6.14	6.6%

Table 5: SQA-iO Normal Morphology Precision

Normal Morphology		Within-Run		Between- Run		Between- Day		Between- Operator/ Lot/Instrument		Total		
Sample	N	Mean	SD	%CV	SD	%CV	SD	%CV	SD	%CV	SD	%CV
1	40	0.0	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%	0.00	0.0%
2	40	15.4	0.87	5.7%	0.87	5.7%	0.33	2.2%	0.78	5.1%	0.92	6.0%
3	40	11.2	1.00	9.0%	1.00	8.9%	0.25	2.2%	0.89	8.0%	0.98	8.8%
4	40	16.5	0.78	4.7%	0.83	5.0%	0.37	2.2%	0.59	3.6%	0.85	5.1%
5	40	10.2	0.58	5.7%	0.61	6.0%	0.41	4.0%	0.45	4.4%	0.66	6.5%
6	40	7.2	0.35	4.8%	0.39	5.4%	0.19	2.6%	0.26	3.6%	0.41	5.6%
7	40	3.6	0.42	11.9%	0.46	13.0%	0.39	10.7%	0.22	6.2%	0.55	15.1%
8	40	8.5	0.48	5.6%	0.53	6.3%	0.51	6.0%	0.35	4.2%	0.68	8.0%

Analytical sensitivity (limits of blank and detection/quantitation):

The defined limit of blank (LoB), Limit of Detection (LoD) and limit of Quantitation (LoQ) of the SQA-iO system for sperm concentration is as follows:

- Limit of Blank (LoB) = 0 M/mL
- Limit of Detection (LoD) = 1.73 M/mL
- Limit of Quantitation (LoQ) = 6.8 M/mL



APPENDIX 6: SQA-iO Warranty

Warranty Sperm Quality Analyzer SQA-iO

Medical Electronic Systems ("MES") warrants that the SQA-iO Sperm Quality Analyzer will be free from defects in workmanship and materials for a period of twelve (12) months from date of the first, initial installation. If a device is resold or re-installed after the first, initial installation, the warranty will continue (or expire) based on the first, initial installation date.

If, during the one-year warranty period, the device is shown to MES's reasonable satisfaction to be defective, MES shall, at its option, replace or repair such a device without charge for parts or labor. The foregoing remedy shall be purchaser's sole and exclusive remedy under this warranty.

The warranty is subject to the following conditions:

- Proper cleaning is followed based on the manufacturer's guidance AND evidence of such scheduled cleaning (weekly) and proper maintenance of the device per the manufacturer's guidelines is provided from the system records.
- No modifications or alterations are made to the SQA-iO device or related testing supplies.
- The SQA-iO is not used, operated, opened by anyone other than the purchaser.
- The SQA-iO is not serviced by anyone or any other entity other than MES or its designee.
- The SQA-iO is used, as labeled for human semen testing only, transported in its original box, stored in the proper temperature range and only manufacturer supplied testing supplies are used for testing, service and maintenance.

If the above conditions are not met or proper maintenance/cleaning records are not provided, this warranty shall be void and of no further force or effect. EXCEPT FOR THE FOREGOING WARRANTIES, THE PRODUCTS ARE SOLD AS-IS AND WITHOUT ANY OTHER WARRANTY OF ANY NATURE WHATSOEVER. MES HAS NOT MADE AND DOES NOT MAKE ANY OTHER REPRESENTATION, WARRANTY, GUARANTY, OR COVENANT, EXPRESS OR IMPLIED, WITH RESPECT TO THE DESIGN, CONDITION, DURABILITY, SUITABILITY, FITNESS FOR USE, FITNESS FOR A PARTICULAR PURPOSE, OR MERCHANTABILITY OF THE SQA IN ANY RESPECT. UNDER NO CIRCUMSTANCES AND IN NO EVENT, WHETHER AS A RESULT OF BREACH OF CONTRACT OR WARRANTY, TORT (INCLUDING NEGLIGENCE AND STRICT LIABILITY) OR OTHERWISE, INCLUDING BUT NOT LIMITED TO INACCURATE RESULTS OR OPERATOR ERROR, SHALL MES BE LIABLE FOR ANY SPECIAL, INCIDENTAL OR CONSEQUENTIAL DAMAGES. IN NO EVENT SHALL MES'S LIABILITY WITH RESPECT TO THE PRODUCT EXCEED THE PURCHASE PRICE FOR SUCH PRODUCT.



APPENDIX 7: SQA-VU Visualization Device

SECTION 1: Overview

The SQA-VU visualization system works specifically with the SQA-iO sperm quality analyzer to visualize sperm samples and capture Motility videos and Morphology images for manual assessment and integration into the test report and SQA-iO patient archive. It is not a standalone device.

SECTION 2: Connect and Operate the SQA-VU

Register / Login to your SQA-iO account: www.sqa-io.com

- 1. Connect the SQA-VU to the same computer as the SQA-iO, using the supplied USB cable.
- Click ALLOW to permit the SQA-VU to access the camera (the SQA-VU will not work without this permission).



- 3. Access the SQA-VU visualization displays from the:
 - **TEST PATIENT** page click the **VIEW SAMPLE** button.
 - **ARCHIVE** click the camera icon **o** for a specific test/patient.
 - TEST RESULTS click the CAPTURE or MORPHOLOGY button.
- 4. Prepare a semen sample using a standard slide and 22X22mm coverslip or an SQA-Vision fixed coverslip slide (for optimal quality).
- 5. Place the slide into the SQA-VU slide adaptor. Insert into the **Viewing Chamber** of the SQA-VU device.
- Use the Focus Knob to visualize the sample clearly. Use the Stage Knob to move to additional fields of view.
- 7. These options are available for assessing the sample:
 - **GRID ON** for easier counting
 - **REAL-TIME** for viewing the sample on the screen.
 - **FREEZE** to accurately count the total number of sperm cells.
 - FULL SCREEN to view the sample on a larger display.
 - **SETTINGS** to adjust the video settings to your preference.
 - **NO SPERM SEEN** can be checked if no spermatozoa were found in all fields of view.
- 8. Capture images and Videos
 - Click the icon on the image to attach it to the REPORT (up to 10 can be attached).
 - Click the MANAGE VIDEOS / MANAGE IMAGES header to view, delete or download.
- 9. Remove the slide adaptor and unplug the SQA-VU from the computer when not in use.





SECTION 3: Device Specifications, Operating Conditions and Cautions

Device Specifications:

- Dimensions: 20 X 16 X 11 cm
- Weight: 1.40 kg
- Power supply: USB powered 5 VDC
- SQA-VU device power consumption: 2.5 [Watt] max
- Recommanded browsers for optimal performance: Chrome, Microsoft Edge

System Requirements:

- Recommended browsers for optimal performance: Chrome, Microsoft Edge
- Operating System: PC with WIN 8 Professional x 32 or above
- Recommended Hardware:
 - CPU: Intel Core I5 & Above
 - o RAM: 8GB
 - Video card: Powerful graphics card to support HD resolution (1280x960)
 - Screen resolution: 1280x960
 - Hard drive: 400GB of free space to store downloaded videos & images
- One free available USB port
- Internet Connection: 5mb per second

Visualization Compartment:

- White LED illumination system with luminous intensity 35000 mcd
- Objective: Standard, x20, chromatic aberration correction
- Focus knob
- Digital CCD
- Field of View Stage knob

Video/image resolution:

- Video: 1280 x 960 pixels, 40 FPS capture of high-resolution videos
- Image: 2560 x 1920 pixels

Operating Temperature and Humidity:

The SQA-VU is designed to operate at the WHO recommended ROOM TEMPERATURE controlled environment of 20-25°C (68-77°F), which is optional for semen testing.

Note: Although the SQA-VU can operate at a higher ambient temperature range of (15-38°C), extreme ambient temperature may impact the accuracy of the semen test results.

Operational Environmental conditions:

The SQA-VU system is intended for indoor use, mains supply fluctuations $\pm 10\%$, Overvoltage Category I, Pollution Degree II.

Caution when device is not in use:

Remove the slide adaptor and unplug the SQA-VU from the computer when not in use.



APPENDIX 8: Assessing Debris/Round Cells in Semen Samples

OVERVIEW:

Grading the degree of debris/round cells in semen samples is important because these components (that are the size of sperm heads or larger) can influence the accuracy of reporting automated concentration. This technical bulletin provides guidance for assessing/grading the % of sample debris/round cells by category.

ASSESSMENT TECHNIQUE:

- 1. Debris/round cells are graded as a percentage in proportion to the number of sperm cells and then divided into 1-4 ranges. Assessing several images may be required to assign a range.
- 2. Count debris/round cell particles without tails that are the size of sperm heads or larger.
- 3. Count the # sperm cells in the image.
- 4. Calculate the % debris: Divide the # debris by the # sperm cells then multiply by 100 for %.
- 5. The absolute number of debris/round cells is only important for determining the % of debris vs. sperm in order to classify the debris level by category (refer to table below).

#	% Range of Debris/Round Cellsvs Sperm	Example	Debris Category in SQA-iO
1	1 Less than 10%	# Sperm 50 and # Debris 1 = 2%	None/Few< 10%
2	11 to 30%	# Sperm 50 and # Debris 10 = 20%	Moderate 11%-30%
3	31 to 99%	# Sperm 50 and # Debris 30 = 60%	Many 31%-99%
4	≥ 100%	# Sperm 50 and # Debris 60 = 120%	Gross >=100%



Appendix 6: Warnings and Regulatory Information

Warnings and Precautions:

- Maintenance Schedule: Clean the measurement compartment weekly using ONLY manufacturer's cleaning supplies provided in the test kit.
- Semen is considered a biologically hazardous material and is subject to laboratory protocols for handling and disposing of such materials in specially marked hazardous waste containers.
- Indoor Use

Cybersecurity Controls:

- Operate the SQA-iO software interface in a controlled environment of the laboratory, accessible to trusted, authorized personnel only.
- Carefully read the entire SQA-iO IFU before initial use to ensure optimal results.
- The SQA-iO USB port is intended to connect the SQA-iO device only. Do not connect any USB devices such as a mouse or a keyboard to the USB port of the SQA-iO.

EMC Related Information

- Intended Use: the SQA-iO is designed and tested to comply with applicable Electromagnetic Compatibility (EMC) standards for use in the electromagnetic environment specified below.
- EMC Compliance: the SQA-iO complies with the requirements of IEC 60601-1-2 general requirements for basic safety and essential performance related with electromagnetic compatibility of Medical Devices. Compliance has been verified through testing under specific conditions. To maintain compliance, follow the guidelines provided in this Instruction for Use.
- No SQA-iO deviations were found from the reference standard or allowances during the SQA-iO EMC testing.
- Electromagnetic Environment: The SQA-iO is intended for use in an indoor environment where radiated RF disturbances are controlled. The intended user of the SQA-iO device should ensure that it is used in such an environment.
- Operate the device away from any source of vibrations such as a centrifuge.
- Use of Accessories: Only use accessories and cables provided or approved by the manufacturer. The use of unauthorized accessories may result in increased emissions or decreased immunity of the device. Specifications of the accessories (PC) required for the safe performance of the SQA-iO are included in Section 1 of the Instruction for Use.
- Interference Caution and Reporting: The user should be aware that electromagnetic emissions from nearby equipment or devices may affect the proper operation of the SQA-iO.
- If electromagnetic interference is suspected to impact the performance of the SQA-iO, report the issue to the manufacturer through CONTACT US, and to the relevant regulatory authority (such as US FCC-Federal Communication Committee). Provide details of the interference, equipment involved, and operating conditions.
- The SQA-iO complies with both emission and immunity requirements.
- The SQA-iO device communicates with the user's PC via a single USB port. There are no RF wireless functions applied by the SQA-iO device.
- Maintenance instructions to ensure that the SQA-iO remains safe and performs to EM disturbances as intended: Disconnect the device if not in use for an extended period of time.
- FCC warning: The SQA-iO operator is required to cease operating the device if the Commission or its representative find that the device is causing harmful interference. Operation cannot resume until the condition causing the harmful interference has been corrected.



• NOTE: "Harmful interference" is defined in 47 CFR §2.122 by the FCC as follows: Interference which endangers the functioning of a radionavigation service or of other safety services or seriously degrades, obstructs, or repeatedly interrupts a radio communication service operating in accordance with the [ITU] Radio Regulations.

Symbols:



CE mark



R

Symbol for **"IN VITRO DIAGNOSTIC MEDICAL DEVICE"**

Symbol for "The intended use of a prescription IVD product"