

#### **IRVINE MEDICAL CENTER**

#### KAISER FOUNDATION HOSPITAL / SOUTHERN CALIFORNIA MEDGROUP LABORATORIES

Dept.	HEMATOLOGY	POLICY #:	04-150					
POLICY/		EFFECTIVE DATE:	11/09/16					
PROCEDURE	AUTOMATED SEMEN	REVISED DATE:						
TITLE	SQA-VISION	AUTHORIZED	Cindy Schwar	Cindy Schwartz, CLS				
			Area Lab Manager					
	All Hematology Staff		Jana Pindur M.D., Laboratory Director					
PERSONNEL								
COVERED.								
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Purpose	This procedure provides instructions for performing semen analysis using the SQA-VISION automated sperm quality analyzer mixed technology method.					
Scope	This procedure is intended for testing personnel trained in the activities outlined in this procedure.					
Specimen	<ul> <li>Specimen Type: Fresh Semen</li> <li>Specimen Volume: Entire ejaculate is required for determining sample volume</li> <li>Minimum Volume: 0.3 mL</li> <li>Maximum Ejaculation to Test Time: 1 hour</li> </ul>					
Specimen Collection	<ul> <li>Provide the patient with local instructions for semen collection, and verify that they have followed these instructions summarized below:</li> <li>2-7 days abstinence from ejaculation prior to specimen collection</li> <li>Collect sample by masturbation or by special direction from physician</li> <li>Lubricants, spermicides and other contaminants are not to be used.</li> <li>The entire specimen must be collected into a clean container supplied only by the provider's office or laboratory.</li> <li>The specimen container should be clearly labeled with the patient's first and last name, medical record number, and date and time of collection.</li> <li>Keep specimen at room temperature. DO NOT refrigerate or expose to heat.</li> </ul>					
Specimen Transport and Temperature	<ul> <li>Transport the specimen to the laboratory right after collection (within 60 minutes after collection) for an accurate evaluation of sperm motility.</li> <li>During transport to the laboratory, the sample should be kept between 20 °C and 37 °C.</li> <li>Do not heat or cool the sample or the container.</li> </ul>					

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Specimen Stability	<ul> <li>The semen sample must be tested within one hour of collection because motility will decline.</li> <li>Semen samples must be tested by the laboratory on a priority basis upon delivery, and expedited to the testing area.</li> </ul>
Specimen Handling Prior to Testing	<ul> <li>When a patient arrives at the laboratory with his specimen, he is given the Patient Questionnaire Form by the receiving laboratory personnel to fill out. See Procedure for Managing the Semen Analysis – Patient Questionnaire Form and Semen Analysis – Patient Questionnaire Form.</li> <li><i>Important Note:</i> Use the information in the completed Patient Questionnaire Form to result in Cerner.</li> <li>The collection container should remain at room temperature until liquefaction is complete or 45 minutes, whichever is shorter.</li> <li>Some samples will not liquefy within 45 minutes (most will liquefy within 15 minutes).</li> <li>If a specimen is not liquefied, the accuracy of the analysis will be compromised.</li> </ul>
Specimen Rejection	<ul> <li>The following rejection criteria are recommended by the vendor/manufacturer.</li> <li>If testing is greater than 60 minutes but less than 2 hours after sample collection, results are questionable due to age of specimen.</li> <li>If testing is greater than 2 hours after collection, reject the specimen.</li> <li>See procedure block <i>Cerner Resulting</i> to report the required Analysis Time and Analysis Time Comment in Cerner.</li> </ul>
Backup Method	• The backup method in the event that the SQA-Vision is out of service is the

SQA-Vision located at Kaiser Anaheim Medical Center.

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Equipment, Reagents, Materials and Supplies	<ul> <li>SQA-VISION Analyzer and V-Sperm Software</li> <li>QwikCheck Liquefaction Kit (Catalog #0900)</li> <li>QwikCheck Beads (Catalog #0200)</li> <li>QwikCheck Test Strips for Semen Analysis (Catalog #0700) using BioRad Urine Controls</li> <li>QwikCheck Dilution Kit (Catalog #0800)</li> <li>SQA-V Capillaries (Catalog #0402)</li> <li>SQA-V Cleaning Kit (Catalog #0115)</li> <li>QwikCheck Fixed Cover Slip Slides Kit (Catalog # A-CA-01082-00) Medical Electronic Systems, LLC</li> <li>Microscope Slides, Glass, 1" x 3"</li> <li>Coverslips, 22 x 22 mm</li> <li>pH Indicator Paper</li> <li>Vortex Mixer</li> <li>Dilution Container</li> <li>Timer</li> </ul>
Workplace Safety	<ul> <li>All reagents and controls should be handled as though capable of transmitting infectious diseases. Wear appropriate personal protective equipment when running patient samples or performing scheduled maintenance.</li> <li>Refer to Laboratory Policy and Procedure Manual Safety Section (11).</li> </ul>
<b>Preventive</b> <b>Maintenance</b>	<ul> <li>Perform daily and weekly maintenance as described in the Daily Maintenance and Inspection for the SQA-VISION Sperm Quality Analyzer provided by SCPMG Laboratory Technology Services.</li> <li>When to Clean: <ul> <li>Daily when running samples</li> <li>Weekly</li> <li>After every 10-15 tests</li> <li>After ANY spillage</li> <li>If Self-test or any failure occurs</li> <li>If system becomes contaminated with semen</li> </ul> </li> <li>ONLY use the Manufacturer's cleaning kit and cleaning brush or damage will occur to the SQA-VISION film and the system will not operate!</li> </ul>

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Preventive Maintenance, continued Manufacturer's Recommendations:

- Operate the SQA-VISION away from devices that may cause electronic noise or other devices causing vibrations such as centrifuges.
- Turn the SQA-VISION analyzer OFF at the end of the day and leave OFF when not in use for extended period of time.
- Maximum operational humidity is up to 80% for temperatures of up to 31°C with decreasing linearly to 50% at 38°C.
- The system operates in a wide range of ambient temperatures (15-38°C), however the system is calibrated to measure semen samples at room temperature: 20-25°C (68-77°F). Prior to performing patient testing, ensure that the room temperature and humidity are within testing limits. Note: Extreme ambient temperature may impact the accuracy of motility test results because of the known effect of temperature on human semen.

#### **FRONT PANEL**



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# **REAR PANEL**



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#### Add I-Button

Select **ADD TESTS TO COUNTER** from the **SERVICE MENU** or press the **I-Button** key to open the screen below and follow instructions:



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Step	Action
1	Turn on the SQA-VISION by pressing the main switch located on the left side. The SQA-VISION automatically performs a five to seven minutes System stabilization, Auto-Calibration and Self- Test.
	Notes:
	<ul> <li>During this period, do not touch the system, do not insert capillary/slide into the device, and do not use any keyboard functions.</li> </ul>
	• If stabilization or self-test fails, you will receive an error code. See error and warning messages in the SQA-VISION user guide for resolution that includes recalibration and re-stabilization of the system.
2	• The device communication screen will appear when the System Self-Test process is complete as "Ready for Testing. Please Enter Data Into SQA-VISION".
3	• Turn on the SQA-VISION computer (PC). Located on the PC Desktop, double click the SQA-VISION icon to open the SQA-VISION software. Enter the following:
	USER NAME PASSWORD ENTER Remembra is till Statical widh a Sportaul corregint 2013
	<ul> <li>USER NAME: administrator</li> <li>PASSWORD: fertility</li> </ul>
	Note: Once logged in, Home Screen will download Service Data Status

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**Quality Control** Follow the steps below to prepare and run quality controls.

Notes:

- Three levels of latex QwikCheck bead controls are run each day of use.
- With each new lot of QwikCheck beads assayed control to be run, the user <u>must</u> set-up/update the CONTROL settings by following the SQA-VISION User Guide instructions for updating previous CONTROL settings (defaults).
- Verify that quality control results are within acceptable range before testing patients.
- Two levels of controls are run each day of use using Qwik Check Strips for WBC and pH. Use Iris CA and CB controls.

Step	Action
1	Click the QC/Proficiency tab on the left side of the SQA-VISION
	window.
	Note:
	Performing QC is done in the Latex Beads tab; this window will
	show the current Quality Control Data (Status, Lot# in use, Exp.
	Date, Target and Range).
2	Before testing QC, check that all Service Data Parameters have
	passed and current QC lot# in use has the correct expiration date,
	target and date. If all QC information is correct, proceed to <b>Step 5.</b>
	Otherwise, proceed to <b>Step 3</b> to update the CONTROL settings.
3	Click <b>Setup</b> on the lower right of the window to update QC data
	for new lot of QwikCheck beads. This will link to Control Settings
	window.
4	Under Settings >Control Tab, click <b>barcode</b> under Latex Beads to
	scan barcode from the QwikCheck beads QC material box. This
	step will retrieve all the QC Data necessary for QC testing.
	<i>Note:</i> If barcode scanner is not available, enter manually and fill
	up all the fields in the QC settings data. Click <b>Save</b> to keep
	Settings.
5	Before opening the control box, verify that the control lot number
	is the current lot number in use and thoroughly mix the
	QwikCheck beads in the closed container by gently rotating the
	beads by hand (do not use a vortex).
	Notes:
	• It is imperative that the beads are evenly mixed without creating
	bubbles in order to insure accurate results.
	• The negative control does not require extensive mixing.

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# Quality Control, continued

Step	Action
6	• Open and aspirate the beads into a clean SQA-V capillary in the same manner you would fill the capillary for a normal volume specimen, making sure the cuvette section of the capillary is completely full of liquid and free of bubbles. Refer to procedure block <i>Filling the SQA-V Testing Capillary</i> .
	• Immediately and tightly close the control container after withdrawing the sample to avoid evaporation and spillage.
7	Under QC/Proficiency > Latex Beads tab, click <b>TEST NOW</b> on the desired level of QC latex beads to be run to open the sample preparation instructions screen.
8	Follow the SQA-VISION on-screen instructions: "Insert the SQA- V capillary into the testing chamber". Testing will begin automatically.
	<ul> <li><i>Notes:</i></li> <li>Make sure to wipe free of any sample before insertion.</li> <li>Control test results will be displayed on the SQA-VISION screen, and are automatically saved.</li> </ul>

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# Quality Control, continued

Step		Action
9	<ul> <li>The PASS/FAIL results will vs. the target value and +/- r controls whose target range</li> <li>A CORRECTIVE ACTION results. Click this button to problem descriptions and as</li> <li>Implement a corrective action table below if any of the list</li> </ul>	l be displayed based on the test results range (disregard this for non-assayed is set to "0"). I button is displayed for all FAILED open the table which provides a list of sociated corrective actions. on and re-run the test, or follow the ted corrective action does not apply.
	If quality control resultIs outside of acceptablerange (unacceptable)Is still unacceptable whenrepeated with the samecontrol vialIs still unacceptable aftercleaning testing chamberand repeat testing with thesame control vial	<ul> <li>Then</li> <li>Repeat quality control testing with the same control vial.</li> <li>Clean testing chamber and repeat testing with the same control vial.</li> <li>Repeat testing with a new control vial.</li> <li>If results are within acceptable range, proceed with patient testing by following the procedure block <i>FRESH Mode Testing</i>.</li> <li>If results are still unacceptable, notify a manager for further instance.</li> </ul>
10	<ul> <li>Select the problem associate SAVE. It will then be record corrective action noted.</li> <li>If the reason for the failure is reason and the corrective ac field.</li> <li>Click: REPORT to view and</li> </ul>	ed with the test failure and press ded in the QC ARCHIVE with the is not described on the list, note the tion taken in the USER DEFINED d print the test results report.

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# **FRESH Mode**<br/>**Testing**Follow the steps below to perform and enter patient data on the SQA-VISION<br/>Analyzer for FRESH mode testing.

Entering Patient Data	Entering Patient Data							
Step Action								
1 From the Home Screen, select TEST PATIENT.								
2 Under the <b>FRESH</b> Tab, enter the PATIENT/SAMPLE DATA.	Under the <b>FRESH</b> Tab, enter the PATIENT/SAMPLE DATA.							
Some information will be available in the Semen Analysis – Pa	Some information will be available in the Semen Analysis – Patient							
Questionnaire Form submitted with the sample								
PATIENT ID   Patient's medical record								
number.								
<b>PATIENT NAME and D.O.B</b> Patient's Full name and								
Date of Birth, previously								
run patients will auto-								
populate these fields when								
Patient ID is entered.								
ABSTINENCE         Number of days since the								
patient's last ejaculation								
SAMPLE ID/ ACCESSION # Patient's sample accession	L							
number								
<b>COLLECTED Date and Time</b> Sample collection date and	1							
time								
<b>RECEIVED Date and Time</b> Sample received date and								
time								
METHOD COLLECTION Masturbation or special								
direction from physician								
<b>CONTAINER</b> Sterile Cup or other								
Additional information can be entered if applicable by clicking								
ADDITIONAL button on the Test Patient window POST analy	S1S							
SUCH as:								
TDANSDODT ISSUES Enter any transport issue(s)								
SEMEN ADDEADANCE Enter any charmed some								
COMMENT appearance observed								
3 VOLUME								
<ul> <li>Dour specimen into a graduated plastic contribute tube and</li> </ul>								
determine the volume to the nearest 0.1 mJ								
• Enter the volume of the entire specimen (whole cisculate) in								
milliliters								

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	Entering Patient Data
Step	Action
4	WBC CONC
	Follow the package insert instructions for QwikCheck Test Strips
	to test for WBC.
	• <1 M/mL: Any color LIGHTER than the Leukocytes >= $1$ M/mL
	patch on the label is considered Leukocytes <1 M/mL which is
	considered normal.
	• >= $1M/mL$ : When the WBC concentration in semen is >=
	1M/mL, the Leukocytes patch of the QwikCheck test strips
	reacts and reaches or exceeds the darkest color on the color chart
	which is considered abnormal.
5	pH
	Use pH test strip to determine sample pH.
6	APPEARANCE – NORM/ABNORM
	Appearance is based on visual assessment of the specimen.
	• NORMAL – A normal liquefied semen sample has a
	homogeneous, grey-opalescent appearance. It may appear less
	opaque if the sperm concentration is very low; the color may
	also be different, i.e. yellow in a man with jaundice or taking
	certain vitamins or drugs.
	• ABNORMAL – The color of semen may be red-brown when red
	blood cells are present (haemospermia). Abnormal appearance
	may include significant quantities of debris, uric acid
	crystallization, opaque or thick seminal plasma and/or other
	significant abnormalities.

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7	LIQU	EFACTION and VISCOSITY – NORM/ABNORM					
	WBC,	pH and all other chemical tests should be completed before					
	treating sample with QwikCheck Liquefaction kit. Follow these						
	instr	uctions for using the Liquefaction kit:					
	1.	Select one vial of alpha-Chymotrypsin					
	2.	Tap the vial to move the contents to the bottom of the vial					
		prior to opening.					
	3.	Add the entire contents of one vial to a viscous semen					
		sample.					
	4.	Gently mix the sample to dissolve the powder.					
	5.	Once the sample has liquefied (5-10 minutes), immediately					
		perform automated testing.					
	6.	If sample volume is >5 ml and it does not liquefy after					
		adding one vial, add another vial of alpha-Chymotrypsin					
		following steps 1 through 5.					
	• NOI	RM – Sample liquefies within 60 minutes at room					
	temp	perature without using QwikCheck Liquefaction Kit.					
	ABN	NORM –					
	• If	QwikCheck Liquefaction Kit successfully liquefied the					
	sa	mple within 60 minutes.					
	• If	QwikCheck Liquefaction Kit does not successfully liquefy					
	the	e sample within 60 minutes.					

	Entering Patient Data							
Step	Action							
	FRESH Mode Testing							
1	After entering all the Patient and Sample Data, determine the							
	volume of specimen to be tested and prepare for testing.							
	If the volume of Then							
	specimen is							
	Normal (≥0.5 mL)	Prepare a testing capillary for a normal						
		volume specimen. Refer to procedure						
		block Filling the SQA-V Testing						
		Capillary.						

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If the volume of	Then
specimen is Low volume with 0.3 mL - 0.5 mL	<ul> <li>A 1+1 dilution (1:2) is to be performed before testing the semen sample.</li> <li>Use QwikCheck-Dilution Kit and dilute the semen 1:2. <i>Note:</i> Semen sample must be completely liquefied and well mixed prior to dilution.</li> <li>Pipette equal amount (300 uL) of semen sample and QwikCheck Dilution in a wide mouth dilution container provided.</li> <li>Gently rotate the container to evenly distribute the spermatozoa throughout the sample without introducing bubbles To prevent air bubbles from forming, denot shake, or use a pipette to mix, or use a pipette to aspirate.</li> <li>Fill a testing capillary in the usual manner for normal volume specimen. Refer to procedure block <i>Filling the SQA-V Testing Capillary</i>.</li> <li>Highlight the 1+1 DILUTION button by clicking it.</li> </ul>

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	FRESH MODE Testing							
Step	Action							
2	Click <b>TEST NOW</b> to perform patient testing.							
	<i>Note:</i> The system will Self Calibrate. Do not use the keypad or insert a							
	testing capillary/slide at this time.							
	$\mathbf{\hat{k}}$							
	SYSTEM IS CALIBRATING DO NOT INSERT CAPILLARY (FRESH SAMPLE)							
	1. MIX THE SEMEN SAMPLE THOROUGHLY 2. DILUTE SAMPLE (1+1) 3. FILL CAPILLARY WITH SEMEN SAMPLE 4. CLEAN, WIPE AND INSPECT CAPILLARY FOR BUBBLES							

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Cerner Resulting

	FRESH MODE Testing					
Step	Action					
3	• After system calibration, the instrument will instruct to insert testing capillary. Follow VISION instructions:					
	PLEASE INSERT TESTING CAPILLARY					
	(FRESH SAMPLE)					
	1. MIX THE SEMEN SAMPLE THOROUGHLY 2. FILL CAPILLARY WITH SEMEN SAMPLE 3. CLEAN, WIPE AND INSPECT CAPILLARY FOR BUBBLES					
	PRESS ESC ON SQA-VISION TO CANCEL TESTING					
	<ul> <li>A table of results will be displayed after testing Fresh semen samples with normal volume or diluted 1:2 samples.</li> <li>Test Results are automatically saved. Printable test report can be opened by clicking the <b>Report</b> button.</li> </ul>					

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After obtaining report from instrument, proceed to Accession Result Entry in Cerner for final verification of results. Follow steps below before clicking *Verify* of test results.

Step		Action
	DTA	Result
1	Semen Collection Time	Result as obtained from the instrument
2	Days of Abstinence	Result as obtained from the instrument
3	Method of Collection	Result as obtained from the instrument
4	Semen Collection	Result as obtained from the instrument
	Container	
5	Collection Issues	Result as obtained from the instrument
6	Transport Issues	Result as obtained from the instrument
7	Specimen Received	Result as obtained from the instrument
	Time	

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8	Analysis Time	Result as obtained from the instrument
		<ul> <li>Skip Step 9 and proceed to Step 10 if sample is within 1 hour from collection.</li> <li>Proceed to Step 9 if sample is: <ul> <li>&gt;1 hour from collection, or</li> <li>&gt;2 hours from collection AND</li> </ul> </li> </ul>
	A sealessia Times	immotile
9	Analysis Time Comment	<ul> <li>If sample is:</li> <li>&gt;1 hour but &lt;2 hours from collection, then enter drop down selection result of: &gt;1Hr from Collection; Motility results are questionable due to age of specimen.</li> <li>&gt;2 hours from collection AND immotile, then cancel using Cerner cancel message: Stability Exceeded, Test Not Performed. See Lab Informatics procedure for <i>Canceling Test Orders</i> in LabNet or MasterControl.</li> </ul>

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Cerner Resulting Cerner ResultingCerner Resulting, continued

	Step	Action
	DTA	RESULT
10	Semen Appearance	Result as obtained from the instrument
11	Semen Appearance	Result as obtained from the instrument
12	Semen Liquefaction and Viscosity	<ul> <li>Enter drop down selection result:</li> <li>Normal if liquefied within 1 hour without addition of liquefaction kit.</li> </ul>
		<ul> <li>Abn1 if liquefied within 1 hour with addition of liquefaction kit.</li> <li>Abn&gt;1 if not liquefied within 1 hour despite addition of liquefaction kit.</li> </ul>
13	Semen pH	Result as obtained from the instrument
14	Semen WBC	Result as obtained from the instrument
15	Semen Volume	Result as obtained from the instrument
16	Sperm Concentration	<ul> <li>Result as obtained from instrument.</li> <li>proceed to step 17 if result is below instrument reportable range "&lt;2 M/ml"</li> </ul>
		• skip step 17 and proceed to step 18 if result is $\geq 2M/ml$
17	Sperm Concentration Comment	After review of sample microscopically using the visualizer, enter drop down selection result of:
		• Rare Sperm/hpf if sperm is seen.
		• No Sperm/hpf if no sperm is seen.
18	Immotility (IM)	Result as obtained from instrument.
19	Nonprogressive Motility (NP)	Result as obtained from instrument.
20	Progressive Motility (PR)	Result as obtained from instrument.
21	Tot PR Mot Cnt	Result as calculated by Cerner.
22	Norm Morph pct	Result as obtained from instrument.
23	Total Sperm/Ejaculation	Result as obtained from instrument.
24	Tot Motility (PR+NP)	Result as obtained from instrument.

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Filling the<br/>SQA-V Testing<br/>CapillaryFollow the steps below to fill the SQA-V testing capillary with a normal<br/>volume sample.



#### **SQA-V** Testing Capillary

#### Notes:

- Sample volume must be **at least** 0.5 mL.
- Sample container should be wide-necked and deep enough to facilitate inserting the capillary into the sample at the bottom of the container.
- Sample must be completely liquefied and well mixed prior to aspiration. Gently rotate container to fully mix liquefied sample.
   *WARNING:* Do not shake or use a pipette to aspirate and dispense sample in order to mix. Otherwise, air bubbles will form.
- Carefully check that liquefied, fully mixed sample is free of air bubbles (or that there is an adequate amount of sample below the air bubbles) before immersing the capillary into the sample, thus ensuring no air bubbles will be aspirated into the capillary.

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# Filling the SQA-V Testing Capillary, continued

Step	Action
1	<ul> <li>Push the syringe piston in fully. Place only thin part of the capillary into the bottom of the sample while angling the sample container at about 45 degrees.</li> <li>Placing two fingers below the piston head, pull the piston 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.</li> </ul>
2	<ul> <li>Holding the capillary in a vertical position, visually confirm that the sample has completely filled the thin section (without a meniscus) and the cuvette section, and appears in the Luer adaptor.</li> <li>Tap on the syringe to make sure there are no air bubbles in the sample. If, after tapping, some air bubbles appear below the Luer adaptor, dip the capillary into the semen sample again and aspirate a small quantity of semen to draw the air bubbles into the syringe.</li> </ul>

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# Filling the SQA-V Testing Capillary, continued

Step	Action
3	<ul> <li>Quickly (to avoid wicking) and thoroughly wipe the outer surface of the capillary, both top and bottom, with Kimwipe.</li> <li>It is important to remove all semen from the exterior of the capillary to prevent the SQA-VISION optical chamber from becoming clogged.</li> <li>Visually confirm that the capillary chambers are still full following the cleaning process. If some of the sample has been depleted (a meniscus has formed in the thin part of the capillary), fill the capillary part from the cuvette section by slightly pushing in the piston.</li> </ul>
4	<ul> <li>Slowly and carefully push-in the blue separating valve of the testing capillary until it is level with the plastic.</li> <li>Image: The capillary is now ready to be inserted into the SQA-VISION recommendation.</li> </ul>
5	<ul> <li>Insert the testing capillary into the lower measurement compartment with the blue stopper down.</li> <li>Push it in as far as it will go to ensure that the capillary is properly seated in the compartment.</li> </ul>

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Visualization System – Fixed Coverslip and Slide Preparations The SQA-VISION Visualization System (Manual Method in SQA-VISION) is used to view sperm cells, and capture static and dynamic images. The system displays real time videos or pictures of the semen sample on a PC monitor.

The visualization system:

- Accommodates a VISION fixed coverslip slide or a standard slide (both 20micron depth).
- Allows smooth magnification transition from x1188 to x1725 (use Zoom In/Out).

	Fixed Coverslip Preparation				
Step	Action				
1	Mix the semen sample thoroughly and pipette $\sim 3 \mu l$ of semen.				
2	Load the sample in the fixed coverslip as instructed by the arrows. <i>Note:</i> There are two wells on each slide for duplicate counts.				
3	After loading the sample, 'drop' the slide into the slide holder.				
4	Insert the slide holder into the VISION visualization compartment.				



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#### Visualization System - Fixed Coverslip and Slide Preparations, continued

	Standard Slide Preparation
Step	Action
1	Mix the semen sample thoroughly and load 10 $\mu$ l of semen onto the <b>distal</b> end of a standard slide and cover with a 22 mm x 22 mm
	cover-slip (to insure 20 micron depth).
2	Insert the prepared standard slide into the SQA-VISION slide holder and insert into the visualization compartment of the VISION.

The visualization system can be used to review samples that fall below the dynamic range of the SQA-VISION to verify that the count is <2 M/mL.

- See procedure block *Cerner Resulting* to report the required comment for sperm concentration <2 M/mL in the **Sperm Concentration Comment** in Cerner by choosing a drop down result selection of either:
  - *Rare Sperm/hpf* if sperm is seen, or
  - No Sperm/hpf if no sperm is seen.

Continued on next page

Visualization Process – Low Quality and Manual Counters

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# Reportable<br/>RangeThe table below shows the reportable range for SQA-VISION automated<br/>results for FRESH sample type.

Sperm Concentration (M/mL)	<2 - 400
Motility (%)	0 - 100
Progressive PR (%)	$\geq$ 32
Normal Forms Morph (%)	2 - 30
Sperm # (M/ejaculate)	$\geq$ 39

#### Reference Range

- The ranges established by the SQA-VISION are based on WHO 5<sup>th</sup> reference values or MES (for proprietary semen parameters).
- The table below shows the reference ranges for Kaiser Permanente.

Semen Volume	>= 1.5 mL
Semen pH	>= 7.2
Semen WBC	<1 million/mL
Total Sperm/Ejaculation	>= 39 million
Sperm Concentration	>= 15 million/mL
Total Motility (PR+NP)	>= 40%
Progressive Motility (PR)	>= 32%
Morphology Normal Forms	>= 4%

#### Limitations

- Analysis should begin within 60 minutes of collection, otherwise the critical determination of motility and possible other parameters may not be reliable.
- Motility testing is time sensitive and is run FIRST on the SQA-VISION.
- Specimens received more than 60 minutes, but less than 2 hours after collection should be analyzed. Please note that results are questionable due to age of specimen.
- If specimen is not sufficient, report as QNS for those tests that were not completed.

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Regulatory Requirements to be Performed The following regulatory requirements are to be performed:

- **Instrument Calibration Verification**: The analyzer's calibration is checked against the original factory calibration parameters for the following criteria:
  - At complete changes of reagents, unless it can be demonstrated that changing reagent lots does not affect either the range used to report patient test results or the control values
  - When QC materials reflect an unusual trend or shift or are outside acceptable limits, and other means of assessing and correcting unacceptable control values fail to identify and correct the problem
  - After major maintenance or service
  - When recommended by the manufacturer
  - At least every six months
  - System Precision and Lower Limit Detection and Motility Method Verification at least semi-annually:
    - The precision and lower limit detection ability of the SQA-VISION is confirmed by completing an abbreviated validation study.

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Non-Controlled Documents	The following non-controlled documents support this procedure.						
	CAP Laboratory Accreditation Standards Checklist						
	• SQA-VISION User Guide, Version 104.13.2, Catalog # VS-ML-01051-00						
	SQA VISION, February 28, 2016						
	<ul> <li>Product Insert; Medical Electronic Systems, QwikCheck Beads</li> </ul>						
	<ul> <li>Product Insert; Medical Electronic Systems, QwikCheck Test Strips</li> </ul>						
	<ul> <li>Product Insert; Medical Electronic Systems, QwikCheck Liquefaction</li> </ul>						
	<ul> <li>Product Insert; Medical Electronic Systems, QwikCheck Dilution</li> </ul>						
	<ul> <li>Technical Release Bulletin: Semi-Annual (every 6 months) Calibration</li> </ul>						
	Confirmation; Application: Any SQA-V/SPERMALITE Visualization						
	System; Re-Issue date/Distribution: Tuesday, October 22 <sup>nd</sup> , 2013/All						
	SQA-V Users						
	• WHO laboratory manual for the Examination and processing of human						
	semen, 5 Edition						
<b>Documents</b>	The following controlled documents support this procedure.						
	Procedure						
	Semen Analysis Collection from local laboratory or LabNet						
	Procedure for Managing the Semen Analysis – Patient Questionnaire						
	Form						
	Form						
	Semen Analysis – Patient Questionnaire Form						

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Change	Changes Made to SOP – describe	Signature	Med. Dir.	Lab	Date
type: New,	_	responsible	Reviewed/	Manager	change
Major,		person/date	Date	reviewed/	Imp.
Minor etc.		1		date	1 I
Major	New instrument – SQA Vision to	Cindy			
	replace the SQA-V for performing	Schwartz			
	automated semen analysis	10/17/16			