#### TRAINING UPDATE

Lab Location:GECDate Distributed:7/13/2015Department:CoreDue Date:8/9/2015Implementation:8/10/2015

## **DESCRIPTION OF PROCEDURE REVISION**

# Name of procedure:

QuikLYTE Na+/K+/Cl- by Dimension® Xpand Chemistry Analyzer GEC.C03 v3

# **Description of change(s):**

Section	Reason	
1, 3, 5, 9, 10, 11, 14	Remove testing criteria for urine specimens (not performed at this site)	
10.2	Add reporting potassium to one decimal	

This revised SOP will be implemented on August 10, 2015

Document your compliance with this training update by taking the quiz in the MTS system.

# **Approve draft for training (version 3)**

## **Technical SOP**

Title	QuikLYTE Na <sup>+</sup> / K <sup>+</sup> / Cl <sup>-</sup> by Dimension® Xpand Chemistry Analyzer	
Prepared by	Ashkan Chini	Date: 5/11/2011
Owner	Robert SanLuis	Date: 4/13/2015

Laboratory Approval	Local Effective Date:	
Print Name and Title	Signature	Date
Refer to the electronic signature		
page for approval and approval		
dates.		

Review		
Print Name	Signature	Date

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## 1. TEST INFORMATION

Assay	Method/Instrument	Local Code
Sodium, Plasma/Serum		SOD
Sodium, Random Urine		<del>UNAR</del>
Sodium, 24 hour Urine	D' ' @ V 1	UNA24
Potassium, Plasma/Serum	Dimension® Xpand Chemistry Analyzer	K
Potassium, Random Urine	Chemistry Anaryzer	<del>UKR</del>
Potassium, 24 hour Urine		UK24
Chloride, Plasma/Serum		CL

# Synonyms/Abbreviations

Sodium / Na<sup>+</sup>, Potassium/ K<sup>+</sup>, Chloride/ Cl<sup>-</sup>, Lytes

Sodium, Potassium, and Chloride are part of batteries BMP, COMP, LYTE, AND RENP

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Chemistry

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#### 2. ANALYTICAL PRINCIPLE

The sodium, potassium and chloride (Na/K/Cl) methods use indirect sample sensing with the QuikLYTE® Integrated Multisensor Technology (IMT) to develop an electrical potential proportional to the activity of each specific ion in the sample.

There are five electrodes used to measure electrolytes on the Dimension® system. Three of these electrodes are incorporated into the QuikLYTE® Integrated Multisensor and are ion selective for sodium, potassium and chloride. A reference electrode is also incorporated in the multisensor. After a diluted sample is positioned in the sensor, Na<sup>+</sup>, K<sup>+</sup> or Cl<sup>-</sup> ions establish an equilibrium with the electrode surface. A potential is generated proportional to the logarithm of the analyte activity in the sample. The electrical potential generated on a sample is compared to the electrical potential generated on a standard solution, and the concentration of the desired ions is calculated by use of the Nernst equation.

## 3. SPECIMEN REQUIREMENTS

## 3.1 Patient Preparation

Component	Special Notations
Fasting/Special Diets	N/A
Specimen Collection and/or Timing	Normal procedures for collecting and storing serum and plasma may be used for samples to be analyzed by this method. For potassium measurements, avoid having the patient make a fist during collection.  Random Urine: Clean catch specimen. Deliver to laboratory promptly.  24 Hour Urine: Record duration of collection on requisition and container
Special Collection Procedures	N/A 24 Hour Urine: Inpatient: See Laboratory Test Directory (electronic) for details. No preservative should be added. Refrigerate during collection. Outpatient: Provide patient with prepared instruction sheet and container.
Other	N/A

### 3.2 Specimen Type & Handling

	Criteria	
Type	-Preferred	Plasma (Lithium Heparin), Urine
	-Other Acceptable	Serum

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Criteria			
<b>Collection Container</b>	Plasma: Mint green top tube		
	_	tube, Serum separato	
	24 Hour Urine:	24 hour container, r	<del>10 additives or</del>
	<del>preservatives.</del>		
		Urine collection cur	
Volume - Optimum	Plasma/serum:	<del>24 hr. Urine:</del>	Random urine:
	1.0 mL	Total voided in	<del>10 mL</del>
		<del>24 hours</del>	
- Minimum	Plasma/serum:	<del>24 hr. urine:</del>	Random urine:
	0.5 mL	<del>N/A</del>	<del>5mL</del>
Transport Container and		Collection container	or Plastic vial at
Temperature	room temperatur		
	· · · · · · · · · · · · · · · · · · ·	Collection kit or con	
	<u> </u>	mitted within 2 hour	
		Collection container	<del>at room</del>
G. Int. G. G.	temperature		
Stability & Storage	Room Temperature: Plasma/serum: 1 week		
Requirements	Urine: Not recommended   Refrigerated: Plasma/serum: 1 week		
	Refrigerated:		
	(2-8°C)	Urine: 24 hou	
	Frozen:	Plasma/serun	
	(-20°C or colder	,	
<b>Timing Considerations</b>	Plasma/serum should be separated from the cells within one hour.		
<b>Unacceptable Specimens</b>	Specimens that are unlabeled, improperly labeled, or		
& Actions to Take	those that do not meet the stated criteria are		
	-	equest a recollection	
	with the appropriate LIS English text code for "test not		
		sage. Examples: Qua	
		llection-UNAC. Doc	cument the request
	for recollection i		
<b>Compromising Physical</b>		<ol> <li>Reject sample and</li> </ol>	•
Characteristics		edit the test with the	appropriate LIS
	English text code.		
Other Considerations	Allow to clot co	mpletely prior to cen	trifugation.

#### 4. REAGENTS

Refer to the Material Safety Data Sheet (MSDS) supplied with the reagents for complete safety hazards. Refer to the section in this procedure covering "SAFETY" for additional information.

# 4.1 Reagent Summary

Reagents Supplier & Catalog Number	
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Sample Diluent	Dimension® clinical chemistry system, REF791092901
Dilution Check	Dimension® clinical chemistry system, REF S640
QuikLYTE®	Dimension® clinical chemistry system, REF S600
Integrated Multisensor	
Flush Solution	Dimension® clinical chemistry system, REF S630

## 4.2 Reagent Preparation and Storage

NOTES: Date and initial all reagents upon opening. Each container must be labeled with (1) substance name, (2) lot number, (3) date of preparation, (4) expiration date, (5) initials of tech, (6) any special storage instructions; check for visible signs of degradation.

Refer to the Material Safety Data Sheet (MSDS) for a complete description of hazards. If a specific hazard is present, it will be noted in this procedure when the hazard is first encountered in a procedural step.

## Irritant. Contains 2-chloroacetamide. May cause sensitization by skin contact.

Reagent	Sample Diluent
Container	Manufacturer supplied vial
Storage	Store at 2-8° C
Stability	Sample diluent, opened or unopened product, is stable until the
	expiration date stamped on the vial.
Preparation	Sample diluent is ready for use. No preparation is required.

Reagent	Dilution Check	
Container	Manufacturer supplied bottle	
Storage	Store at 2-8°C	
Stability	<ul> <li>Unopened vial is stable until the expiration date stamped on the bottle.</li> <li>Opened vial is stable for 6 months after first use.</li> </ul>	
Preparation	Reagents are supplied ready for use. No additional preparation is required.	

Reagent	Flush Solution	
Container	Manufacturer supplied reagent bag	
Storage	Store at 2-30°C	
Stability	<ul> <li>Unopened reagent bag is stable until the expiration date stamped on the label.</li> <li>Opened reagent bag is stable for 21 days after first use.</li> </ul>	
Preparation	Reagents are supplied ready for use. No additional preparation is required.	

Sensor	QuikLYTE® Integrated Multisensor
Container	Manufacturer sensor
Storage	Store at 2-8°C
Stability	Sensor is stable for 5 days or 1000 samples after first use.
Preparation	Sensors are supplied ready for use. No additional preparation is required.

Reagent	Salt Bridge Solution	
Container	Manufacturer supplied bottle	
Storage	Store at 2-30° C	
Stability	<ul> <li>Unopened bottle is stable until the expiration date stamped on the bottle.</li> <li>Opened Salt Bridge Solution is stable for 21 days after first use.</li> </ul>	
Preparation	Salt Bridge Solution is ready for use. No preparation is required.	

#### 5. CALIBRATORS/STANDARDS

#### 5.1 Calibrators/Standards Used

Calibrator	Supplier and Catalog Number
QuikLYTE® Standard A solution	Siemens Dimension®, REF S620
QuikLYTE® Standard B solution	Siemens Dimension®, REF S625

## 5.2 Calibrator Preparation and Storage

NOTE: Date and initial all calibrators upon opening. Each container must be labeled with (1) substance name, (2) lot number, (3) date of preparation, (4) expiration date, (5) initials of tech (6) any special storage instructions; check for visible signs of degradation.

Calibrator	QuikLYTE® Standard A & B solutions		
Preparation	Calibrators are supplied ready for use. No additional preparation		
	is required.		
Storage/Stability	• Store at 2-30°C		
	Unopened reagent bag is stable until the expiration date		
	stamped on the label.		
	• Opened reagent bag is stable for 21 days after first use.		

#### **5.3** Calibration Parameter

Criteria	Special Notations
Criteria	Special Modernis

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Reference Material	QuikLYTE® Standard A & B solutions			
Frequency	The IMT system will routinely perform a one point			
	calibrat	tion with eac	h sample measureme	nt. In addition, the
	system	system performs a two point automatic calibration in		
	duplica	ite every 2 ho	ours. If no analysis is	in progress. Auto-
	calibrat	calibration also occurs shortly after turn-on, with the		
	changii	changing of standards A, B, or a sensor and when reset.		
			Serum/Plasma	<del>Urine</del>
Aggay Danga	Sodiun	1	50 - 200 mmol/L	5-300 mmol/L
Assay Range	Potassi	um	1 - 10 mmol/L	1-300 mmol/L
	Chloric	le	50 - 200 mmol/L	<del>10 - 330 mmol/L</del>
<b>Assigned Coefficients</b>		Sodium	Potassium	Chloride
	$C_0$	1.5	- 0.2	- 10.0
	$C_1$	1.01	1.05	1.15

#### 5.4 **Calibration Procedure**

1. From Operating Menu

press F4: System Prep

press F3: IMT

press F2: Calibration (calibration gets done automatically, no further action is required, unless calibration fails)

#### 5.5 **Tolerance Limits**

IF	THEN
If result fall within assay-specific specification,	proceed with analysis
and QC values are within acceptable limits,	
If result falls outside assay-specific specification,	troubleshoot the assay and/or
or QC values are out of Acceptable limits,	instrument and repeat calibration

#### **QUALITY CONTROL** 6.

#### 6.1 **Controls Used**

Controls	Supplier and Catalog Number
Liquichek <sup>TM</sup> Unassayed Chemistry Control	Bio-Rad Laboratories
Levels 1 and 2	Cat. No. 691 and 692

#### 6.2 **Control Preparation and Storage**

**NOTE:** Date and initial all controls upon opening. Each container should be labeled with (1) substance name, (2) lot number, (3) date of preparation, (4)

# expiration date, (5) initials of tech, and (6) any special storage instructions; check for visible signs of degradation.

Control	Liquichek Unassayed Chemistry Controls Levels 1 & 2
Preparation	Allow the frozen control to stand at room temperature (18-25°C) until completely thawed. Swirl the contents gently to ensure homogeneity. (Do not use a mechanical mixer) Use immediately. After each use, promptly replace the stopper and return to 2-8°C storage.
Storage/Stability	Open controls are stable for 15 days at 2-8°C. Unopened controls are stable until the expiration date at -20 to -70°C.

#### 6.3 **Frequency**

Analyze all levels of QC material after every calibration and each day of testing.

Refer to the Dimension Xpand® QC Schedule in the Laboratory policy Quality Control Program and in the Dimension X-pand® Quick Reference Guide.

#### **6.4 Tolerance Limits**

Step	Action		
1	Acceptable ranges for QC are programmed into the instrument's Quality Control software system and Unity Real Time, and may be posted near the instrument for use during computer downtime.		
2	<ul> <li>Run Rejection Criteria</li> <li>Anytime the established parameters are exceeded (if one QC result exceeds 2 SD), the run is considered out of control (failed) and patient results must not be reported.</li> <li>The technologist must follow the procedure in the Laboratory QC Program to resolve the problem.</li> </ul>		
3	<ul> <li>Corrective Action:         <ul> <li>All rejected runs must be effectively addressed through corrective action. Steps taken in response to QC failures must be documented. Patient samples in failed analytical runs must be reanalyzed according to the Laboratory QC Program. Supervisors may override rejection of partial or complete runs only with detailed documentation and criteria for overrides that are approved by the Medical Director. Consult corrective action guidelines in Laboratory QC Program. Follow corrective action guidelines in the Laboratory QC Program.</li> </ul> </li> <li>Corrective action documentation must follow the Laboratory Quality Control Program.</li> </ul>		

Step	Action			
4	Review of QC			
	QC must be reviewed weekly by the Group Lead or designee and monthly by the Supervisor/Manager or designee.			
	• If the SD and/or CV are greater than established ranges, investigate the cause for the imprecision and document implementation of corrective actions.			

#### 6.5 Review Patient Data

Technologist must review each result with error messages. Refer to the Dimension Xpand® system manual "Error messages" section for troubleshooting. Check for unusual patterns, trends, or distributions in patient results (such as an unusually high percentage of abnormal results). Resolve any problems noted before issuing patient reports.

#### 6.6 Documentation

- QC tolerance limits are programmed into the instrument and Unity Real Time; it calculates cumulative mean, SD and CV and stores all information for easy retrieval.
- Quality control records are reviewed daily at the bench, weekly by the Group Lead or designee, and monthly by the Supervisor/Manager or designee.
- Refer to complete policies and procedures for QC documentation and for record retention requirements in the Laboratory QC Program.

#### **6.7 Quality Assurance Program**

- Each new lot number of reagent or new shipment of the same lot of reagent must be tested with external control materials and previously analyzed samples. Performance of the new lot must be equivalent to the previous lot; utilize published TEA for acceptability criteria.
- Training must be successfully completed and documented prior to performing this
  test. This procedure must be incorporated into the departmental competency
  assessment program.
- The laboratory participates in CAP proficiency testing. All proficiency testing materials must be treated in the same manner as patient samples.
- Monthly QC must be presented to the Medical Director or designee for review and signature.
- Monthly QC mean and SD are sent to Bio-Rad Laboratories for peer group comparison.
- Consult the Laboratory QC Program for complete details.

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# 7. EQUIPMENT and SUPPLIES

## 7.1 Assay Platform

Dimension Xpand® System

# 7.2 Equipment

- Refrigerator capable of sustaining 2–8°C.
- Freezer capable of sustaining range not to exceed -20 to -70°C.
- Centrifuge

## 7.3 Supplies

- Plastic serum tubes and serum cups
- Purified water (Millipore® or equivalent)
- Calibrated pipettes and disposable tips

#### 8. PROCEDURE

All necessary reagents are required to perform this test.

QuikLYTE® is performed on the Dimension Xpand® System after the method is calibrated (see Reference Material in Calibration section) and Quality Controls are acceptable.

NOTE: For all procedures involving specimens, buttoned lab coats, gloves, and face protection are required minimum personal protective equipment. Report all accidents to your supervisor.

The package insert for a new lot of kits must be reviewed for any changes before the kit is used. A current Package Insert is included as a Related Document.

8.1	Instrument Set-Up Protocol			
1.	For instrument set up and operation: Refer to Startup and Maintenance, Siemens Dimension® Xpand procedure.			
2.	Check reagent inventory			
3.	Sampling, reagent delivery, mixing, processing, and printing of results are automatically performed by the Dimension® Xpand system. For details of the automated parameters, see below under "Test conditions."			

8	8.2	Specimen/Reagent Preparation			
	1.	Centrifuge the specimens.			

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8.2	Specimen/Reagent Preparation			
2.	Specimens are placed in Dimension <sup>®</sup> Xpand segments for analysis by the instrument. Refer to the Sample Processing, Siemens Dimension <sup>®</sup> Xpand procedure. The sample			
	container (if not a primary tube) must contain sufficient quantity to accommodate the			
	sample volume plus 50 µL of dead volume. Precise container filling is not required.			

8.3	Specimen Testing			
1.	For QC placement and frequency, refer to the Dimension <sup>®</sup> Xpand QC Schedule in the Laboratory QC Program.			
2.	Follow the instructions, outlined in the Dimension® Xpand Operators Manual			
3.	The instrument reporting system contains error messages to warn the user of specific malfunctions. Results followed by such error messages should be held for follow-up. Refer to the Dimension® Xpand system manual "Error messages" section for troubleshooting.			
4.	Follow protocol in Section 10.5 "Repeat criteria and resulting" for samples with results above or below the Analytical Measurement Range (AMR).  Repeat critical values and document according to Critical Values procedure.  Investigate any failed delta result and repeat, if necessary.			
5.	Append the appropriate English text code qualifier messages to any samples requiring a comment regarding sample quality and/or any other pertinent factors.			

Test Conditions		
Sample Size: Na <sup>+</sup> , K <sup>+</sup> , Cl <sup>-</sup>	45 μL	
Temperature:	18-29° C	
Type of Measurement:	Indirect Potentiometric	

### 9. CALCULATIONS

The instrument automatically calculates and prints the concentration of Na<sup>+</sup>, K<sup>+</sup>, Cl<sup>-</sup> in mmol/L.

For 24 hour urine **Sodium**, the LIS will calculate the total mmol/L of Sodium/24hrs if the Sodium result from the aliquot is within the CRR. **CRR for urine Sodium is 5-300** mmol/L.

If below 5 mmol/L, the total mmol/24hrs is manually calculated as follows:



A "less than" character (<) is placed in front of the numerical value when reporting.

For 24 hour urine **Potassium**, substitute the number **1** instead of the **5** in the above calculation. **CRR for urine Potassium is 1-300** mmol/L.

For values **above** the CRR for both Sodium and Potassium (300 mmol/L), the same calculation is used as above except substitute **300** for **5** and use the "greater than" character (>) in front of the numerical answer from the calculation.

#### 10. REPORTING RESULTS AND REPEAT CRITERIA

#### 10.1 Interpretation of Data

None required

#### 10.2 Rounding

No rounding is necessary. Instrument reports sodium and chloride results as a whole number and potassium results to one decimal place.

#### 10.3 Units of Measure

mmol/L

#### 10.4 Clinically Reportable Range (CRR)

	Serum/Plasma	<del>Urine</del>
Sodium	50-200 mmol/L	5-300 mmol/L
Potassium	1-10 mmol/L	1-300 mmol/L
Chloride	50-200 mmol/L	

#### 10.5 Repeat Criteria and Resulting

All repeats must replicate the original result within the total allowable error (TEa) of the assay. Refer to TEa listing for specific information.

Values that fall within the AMR or CRR may be reported without repeat. Values that fall outside these ranges must be repeated

IF the result is		THEN
Plasma/serum		
		Repeat the assay using the primary sample. If results are still
Na	>200 mmol/L	greater than the CRR, consult supervisor before releasing
K	>10.0 mmol/L	results.
Cl	>200 mmol/L	

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IF the result is	THEN
Urine results	Repeat the assay using a freshly prepared aliquot from the
	primary sample. If results are still greater than the CRR,
Na >300 mmol/L	consult supervisor before releasing results. See Section 9 for
K >300 mmol/L	Calculation instructions for 24 hour urines.

Message	Code	
Verified by repeat analysis	Append –REP to the result.	

## 11. EXPECTED VALUES

# 11.1 Reference Ranges

# Plasma/Serum:

Age	Sodium	Potassium	Cloride
Adult (>18 years):	135-145 mmol/L	3.5-5.1 mmol/L	98-107 mmol/L
Pediatric:			
0 – 6 days	131-144	3.5-5.7	97-108
7 – 30 days	132-142	3.4-6.1	97-108
1 – 5 months	132-140	3.5-5.8	97-108
6 – 12 months	131-140	3.5-6.1	97-106
13 – 24 months	132-141	3.3-4.7	97-107
> 2 years	135-145	3.5-5.1	98-107

## Random Urine:

Age	<del>Sodium</del>	<b>Potassium</b>
All	<del>20-110 mmol/L</del>	<del>12-62 mmol/L</del>

#### 24 hour Urine:

Age	Sodium	Potassium	
All	40-200 mmol/24hr	25-125 mmol/24hr	

## 11.2 Critical Values

## Plasma/serum

Analyte	Low Critical Values	High Critical Values
Na	< 120 mmol/L	> 160 mmol/L
K	< 3.0 mmol/L	> 6.1 mmol/L
Cl	< 75 mmol/L	> 126 mmol/L

# 11.3 Priority 3 Limit(s)

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#### 12. CLINICAL SIGNIFICANCE

Chloride may be decreased in the following cases: Hypoventilation, Protracted Vomiting, Chronic Diarrhea, Diabetic Ketoacidosis, Lactic Acidosis, Adrenal Disease, and/or Renal Failure. Increased Chloride may occur in the following: **Hyperventilation:** Excess breathing results in the reduction of carbonic acid content of plasma and therefore a fall in bicarbonate ion concentration. There are many causes of excess ventilation: they include many diverse diseases, drugs which stimulate the respiratory center, anxiety, fear, and decreased oxygen tension or increased CO2 tension in the blood. **Drugs:** Large doses of ammonium or potassium chloride may produce hyperchloremia. **Dehydration:** A decrease in plasma water will necessarily result in an increase in the chloride concentration.

Some causes of increased potassium may include anuria, tissue damage (crush injuries, with damage to large volumes of muscle tissue, and massive hemolysis, are examples), violent muscle contraction (vigorous exercise may cause a temporary elevation in plasma potassium), certain seizures, Addison's disease (Primary Adrenal Insufficiency), and Diabetes mellitus.

Decreased potassium levels may occur in prolonged diarrhea or vomiting, diuretic administration, and mineralocorticoid excess.

Increased Sodium may occur in simple dehydration, diabetes insipidus, hypothalamic disease, osmotic loading, excessive sodium intake, steroid therapy, excessive sweating, or Cushing's disease. Decreased sodium levels are more common and may be due to diuretics, sweating, kidney disease, congestive heart failure, severe diarrhea and vomiting, primary adrenal insufficiency, hepatic cirrhosis, diabetes mellitus, or inappropriate antidiuretic hormone secretion.

#### 13. PROCEDURE NOTES

FDA Status: FDA Approved/clearedValidated Test Modifications: None

The instrument reporting system contains error messages to warn the operator of specific malfunctions. Any report slip containing such error messages should be held for follow-up. Refer to your Dimension Operator's Guide.

A system malfunction may exist if the following 5-test precision is observed:

	Concentration	S.D.
Sodium	120-160 mmol/L	> 2.0  mmol/L
Potassium	2-6 mmol/L	> 0.15 mmol/L
Chloride	95-128 mmol/L	> 2.0  mmol/L

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#### 14. LIMITATIONS OF METHOD

#### 14.1 Analytical Measurement Range (AMR)

	Serum/Plasma	<del>Urine</del>
Sodium	50-200 mmol/L	5-300 mmol/L
Potassium	1-10 mmol/L	1-300 mmol/L
Chloride	50-200 mmol/L	

#### 14.2 Precision

	Mean	Standard Deviation (%CV)	
Material	mmol/L	Within-run	Total
Sodium Normal QC	130	0.7	2
Sodium Elevated QC	148	0.9	1.5
Sodium Urine QC	148	0.8	1.7
Potassium Normal QC	4.2	0.02	0.05
Potassium Elevated QC	5.8	0.05	0.07
Potassium Urine QC	66	0.4	1.6
Chloride Normal QC	96	0.6	1.4
Chloride Elevated QC	110	0.6	1.1
Chloride Urine QC	173	1.7	1.9

#### 14.3 Interfering Substances

Samples exposed to Benzalkonium salts present in certain blood catheter devices will cause falsely elevated sodium and potassium measurements.

Citrate at a test concentration of 52.9 mmol/L decreases sodium by 38 mmol/L, decreases potassium by 0.6 mmol/L and increases chloride by 57 mmol/L. Thiopental increases sodium results by as much as 8 mmol/L at 14 mg/dL of thiopental and up to 4 mmol/L at 2.8 mg/dL of tipental.

#### 14.4 Clinical Sensitivity/Specificity/Predictive Values

Not available.

#### 15. SAFETY

The employee has direct responsibility to avoid injury and illness at work. Nearly all harmful exposures to infectious substances and chemicals, and other injuries, can be avoided with effective training and consistent safe work practices.

Become familiar with the Environmental Health and Safety (EHS) Manual to learn the requirements on working safely and protecting the environment from harm. Although lab work typically focuses on the hazards of working with specimens and chemicals, we must also control other important hazards.

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- Slips, trips, and falls cause many serious injuries. Please ensure that spills are cleaned quickly (to avoid slippery floors) and that you can see and avoid obstacles in your path.
- Ergonomic injuries result from performing tasks with too much repetition, force, or awkward position. Ergonomic injuries include strains and back injuries. Learn about ergonomic hazards and how to prevent this type of injury.
- Scratches, lacerations, and needlesticks can result in serious health consequences. Attempt to find ways to eliminate your risk when working with sharp materials.

Report all accidents and injuries <u>immediately</u> to your supervisor or the business unit Environmental Health and Safety Manager or Specialist.

#### 16. RELATED DOCUMENTS

- 1. Dimension Xpand® Clinical Chemistry System Operator's Manual
- 2. Calibration / Verification Siemens Dimension® Xpand procedure
- 3. Dimension Xpand® Cal Accept Guidelines
- 4. Dimension Xpand<sup>®</sup> Calibration summary
- 5. Sample Processing, Siemens Dimension® Xpand procedure
- 6. Start up and Maintenance, Siemens Dimension® Xpand procedure
- 7. Laboratory Quality Control Program
- 8. QC Schedule for Siemens Dimension Xpand®
- 9. Laboratory Safety Manual
- 10. Material Safety Data Sheets (MSDS)
- 11. Siemens Dimension Xpand<sup>®</sup> Limits Chart (AG.F143)
- 12. Quest Diagnostics Records Management Procedure
- 13. Dimension Xpand<sup>®</sup> System Error Messages Chart
- 14. Centrifuge Use, Maintenance and Functions Checks (Lab policy)
- 15. Hemolysis, Icteria and Lipemia Interference (Lab policy)
- 16. Repeat Testing Requirements (Lab policy)
- 17. Critical Values (Lab policy)
- 18. Current Allowable Total Error Specifications at <a href="http://questnet1.qdx.com/Business">http://questnet1.qdx.com/Business</a> Groups/Medical/qc/docs/qc\_bpt\_tea.xls
- 19. Current package insert QuikLYTE<sup>®</sup> Integrated Multisensor REF S600

#### 17. REFERENCES

- 1. Ghoshal, Amit K. and Soldin, Steven J., Evaluation of the Dade Behring Dimension<sup>®</sup> RxL: Integrated chemistry system-pediatric reference ranges. Clinica Chimica Acta 2003; 331:144
- 2. Package Insert, QuikLYTE<sup>®</sup> Integrated Multisensor REF S600, Siemens Healthcare Diagnostics Inc., 08/25/2011.
- 3. Package Insert, Liquichek Unassayed Serum Chemistry Controls, Bio-Rad Laboratories, 05/2014.
- 4. Package Insert, Sample diluent REF791092901, 01/2010.

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## 18. REVISION HISTORY

Version	Date	Section	Reason	Reviser	Approval
			Supersedes SOP C106.001		
000	7/30/2012	1	Add analyzer name	L Barrett	J Buss, RSL
000	7/30/2012	3.1	Add electronic directory	L Barrett	J Buss, RSL
000	7/30/2012	6.7	Add use of TEA for lot to lot runs	L Barrett	J Buss, RSL
000	7/30/2012	9	Add manual calculation for 24hr urine	J Buss	J Buss, RSL
000	7/30/2012	10.5	Remove code QNSR	L Barrett	J Buss, RSL
000	7/30/2012	11.3	Removed SGAH specific preop value	L Barrett	J Buss, RSL
000	7/30/2012	15	Update to standard wording	L Barrett	J Buss, RSL
001	4/13/2015		Update owner	L Barrett	R SanLuis
001	4/13/2015	3.2	Specify lithium heparin anticoagulant	L Barrett	R SanLuis
001	4/13/2015	7.1	Add analyzer name	L Barrett	R SanLuis
001	4/13/2015	6.4, 6.6	Replace LIS with Unity Real Time	L Barrett	R SanLuis
001	4/13/2015	11.2	Standardize low K+ critical value <3.0	L Barrett	R SanLuis
001	4/13/2015	Footer	Version # leading zero's dropped due to new EDCS in use as of 10/7/13.	L Barrett	R SanLuis
2	7/1/2015	1, 3, 5, 9, 10,11,14	Remove testing criteria for urine specimens (not performed at this site)	L Barrett	R SanLuis
2	7/1/2015	10.2	Add reporting potassium to one decimal	L Barrett	R SanLuis

# 19. ADDENDA

None

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