#### TRAINING UPDATE

**Lab Location: Department:** 

GEC, SGMC & WAH Core

Date Distributed:
Due Date:
Implementation:

8/3/2016 8/24/2016 **8/4/2016** 

#### **DESCRIPTION OF PROCEDURE REVISION**

# Name of procedure:

# **D-Dimer SGAH.G04 v5**

Note: this SOP has been converted to a system SOP

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

Section	Reason
5.1	Removed Bio-Rad QC information
6.1, 6.2	Replace Bio-Rad QC with STA Coag Controls
6.7	Remove QC submission to Bio-Rad
8.3	Add QC/Reagent Loading Instructions
11.3	Move report comment from 10.5
17	Remove Bio-Rad insert

This revised SOP will be implemented on August 4, 2016

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

Site: Shady Grove Medical Center, Washington Adventist Hospital, Germantown Emergency Center

**Technical SOP** 

Title	D-Dimer		
Prepared by	Ashkan Chini	Date:	4/7/2011
Owner	Robert SanLuis	Date:	6/3/2014

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

Signature	Date
	Signature

SOP ID: SGAH.G04 SOP Version # 5

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#### 1. TEST INFORMATION

Assay	Method/Instrument	Local Code
D - Dimer Quantitative	Immunoturbidometric / STA® Compact	DDIMER

Synonyms/Abbreviations	
D - Dimer	

Department	
Coagulation	

#### 2. ANALYTICAL PRINCIPLE

This assay is based on the change in turbidity of a microparticle suspension that is measured by photometry. A suspension of latex microparticles, coated by covalent bonding with monoclonal antibodies specific for D-dimer, is mixed with the test plasma whose D-dimer level is to be assayed. An antigen-antibody reaction takes place, leading to an agglutination of the latex microparticles which induces an increase in turbidity of the reaction medium. This increase in turbidity is reflected by an increase in absorbance, the latter being measured photometrically. The increase in absorbance is a function of the D-dimer level present in the test sample.

#### 3. SPECIMEN REQUIREMENTS

#### 3.1 Patient Preparation

Component	Special Notations	
Fasting/Special Diets	N/A	
Specimen Collection and/or Timing	Normal procedures for collecting plasma may be used for samples to be analyzed by this method.  Vacutainer tube must be filled to the line to ensure the proper ratio of blood to anticoagulant.	
Special Collection Procedures	If hematocrit >55%, refer to appendices A and B for collection instructions.	
Other	N/A	

#### 3.2 Specimen Type & Handling

Criteria		
Type -Preferred	PLT Poor Plasma (sodium citrate)	
-Other Acceptable	None	
Collection Container	Light blue top tube (3.2% sodium citrate)	
	Citrated blood 9:1 (blood to anticoagulant)	
Volume - Optimum	2.7 mL (9:1 blood to anticoagulant) in a 2.7 ml tube	
- Minimum	2.4 mL (9:1 blood to anticoagulant) in a 2.7 ml tube	
- Optimum - Minimum	1.8 mL (9:1 blood to anticoagulant) in a 1.8 mL tube	
The line (5.11 blood to distrebuguish) in a file line tag		
Transport Container and Temperature	Light blue vacutainer (as above) or a clean plastic screw capped vial at room temperature.	
Stability & Storage	Room Temperature: 8 hours	
Requirements	$(20 \pm 5^{\circ} \text{ C})$	
	Refrigerated: Not recommended	
	Frozen plasma: 1 month at -20 C.	

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Criteria	
Specimen preparation	Centrifuge whole blood for specified time /speed
	documented on each centrifuge for preparing platelet-poor
	plasma.
Unacceptable Specimens	Specimens that are unlabeled, improperly labeled, or those
& Actions to Take	that do not meet the stated criteria are unacceptable.
	Clotted or under-filled tubes are not accepted.
	Request a recollection and credit the test with the
	appropriate LIS English text code for "test not performed"
	message.
Compromising Physical	Moderate to gross hemolysis. Reject sample and request a
Characteristics	recollection. Credit the test with appropriate LIS English
	text code HMM (Specimen moderately hemolyzed) or
	HMT (Specimen markedly hemolyzed)
	Lipemia: Acceptable
	Icterus: Acceptable
Other Considerations	None

#### 4. REAGENTS

Refer to the Safety Data Sheet (SDS) for complete safety hazards. Refer to the section in this procedure covering "SAFETY" for additional information.

#### 4.1 Reagent Summary

Reagents	Supplier & Catalog Number
STA - LIATEST® D – DI: Buffer & Latex	Diagnostic Stago (REF 00515)
STA – Owren-Koller Buffer	Diagnostic Stago (REF 00360)

#### 4.2 Reagent Preparations 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 Safety Data Sheet (SDS) 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.

Reagent 1 & 2	STA - LIATEST® D – DI: Buffer and Latex	
Container Manufacturer supplied vial		
Storage	2 - 8°C	

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Stability	Unopened reagents are stable until expiration date indicated on the box label.  With the STA-mini Reducer and perforated cap in place the stability of Reagents 1 and 2 after opening and in their original vials is 15 days on the Stago.
Preparation	Allow the reagents 1 and 2 to stand at room temperature (18-25°C) for 15 minutes before use. Mix the reagents by gentle swirling of the vials without creating any bubbles. Then place the perforated cap on each vial.

Reagent 3	STA – Owren-Koller Buffer	
Container	Manufacturer supplied vial	
Storage	2 - 8°C	
Stability  The buffer solution in intact bottles is stable until the expiration date indicated on the box label.  After opening it remains stable for 3 days.		
Preparation Allow it to stand at room temperature (18-25°C) for 30 m before use.		

#### 5. CALIBRATORS/STANDARDS

#### **5.1** Calibration Procedure

The pre-calibrated D-dimer values are identical for all the vials of each lot. To enter the calibration data on the analyzer, scan the barcode printed on the assay value insert across the instrument barcode reader.

The calibration data will be validated for the lot being used once the Stago D-dimer controls are run and tested.

The calibration curve is considered verified for the new reagent lot when both the  $STA^{\$}$ -LIATEST Control N + P are within acceptable range. The acceptable  $STA^{\$}$ -LIATEST Control N + P range is supplied by Stago. QC ranges must fall within the acceptable range which is established utilizing the peer group data in combination with our current/historic analytic performance.

To examine calibration curve on screen:

- Through the MAIN MENU under CALIB/CONTROL select CALIBRATION.
- Move the cursor to D-Dimer and press Enter ← Curve will appear on STA<sup>®</sup> Compact screen.

To print calibration curve:

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• While examining the curve on the STA® Compact screen, press ESC key for options.

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- Select print Option Enter ←. Select Execute Enter ←.
- The curve will print along with the information on all reagents and control lot numbers. Also included are control results and ranges.

#### 6. QUALITY CONTROL

#### **6.1** Controls Used

Controls	Supplier and Catalogue Number
STA - LIATEST Control N + P	Stago Diagnostic, Cat. No. 00526

#### **6.2** Control Preparations 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) time prepared, (5) expiration date and time, (6) initials of tech, and (7) any special storage instructions; check for visible signs of degradation.

Control	STA - LIATEST Control N + P	
Preparation	Reconstitute each vial of Reagent 1 or 2 with exactly 1 mL of Reagent Grade water. Allow the reconstituted material to stand at room temperature (18 - 25° C) for 30 minutes. Then, swirl the vial gently before use.	
Storage	2-8° C	
Stability  The reagents in intact vials are stable until the expiration do indicated on the box label, when stored at 2 - 8° C.  Once reconstituted, Reagents 1 and 2 remain stable for 8 he		

- 1. QC can be run automatically at pre-set intervals (in Test Set-up) or by ordering manually from the Quality Control Menu.
- 2. All control ranges are monitored automatically by the STA® Compact. If any controls are outside the ± 2 SD range, the instrument will audibly and visually alarm the operator. Otherwise, the results can be found in the individual QC files. Control results are automatically filed in the STA® Compact QC file. All results for a 24-hour period are converted to a "mean" value at midnight. This mean is used in the statistical data and is plotted on the Levy-Jennings chart as a daily mean.
- 3. To print all the QC data points for the D-Dimer test, perform the following procedure prior to midnight. From the MAIN MENU under CAL. /CONTROL select QUALITY CONTROL press Enter ← Cursor to the D-Dimer test and press Enter ← to view the Levy-Jennings chart. Press F1 to view the results in tabular form. Press F6, select Execute then press Enter ← to print the individual values under current controls. Press ESC key to exit (back to graph). Press F2 or F3 to view other levels and continue with F1 to view the result list.

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Controls are run at the beginning of each shift, every 4 hours after, and with the change of any reagent used in test performance.

Controls are run after any maintenance is performed on the analyzer.

#### **6.4** Tolerance Limits

Step	Action
1	The established QC ranges are in the QC file of the STA Compact. The quality control results from the instrument are transmitted to Unity Real
	Time and can be viewed in that program. Any out-of-range QC results will be flagged.
2	If all controls are within QC parameters all sample results can be reported.
3	Rejected runs must be effectively addressed by corrective action. Steps taken in response to QC failures must be documented. Patient samples in failed analytical runs must be reanalyzed. Supervisor may override rejection of partial or complete runs only with detailed documentation that follows criteria that is approved by the Medical Director.
4	Corrective action documentation must include the following: QC rule(s) violated, the root cause of the problem, steps taken to correct the problem, how patient samples were handled, and the date and initials of the person recording the information. See the QC/QA SOP "QC Responsibilities" for more detail.
5	If the assay is down and results will not be reported in the scheduled turnaround time, clients will be notified of the situation.

#### 6.5 Review Patient Data

Each result is reviewed for error messages. Refer to the STA® Compact system manual "Error messages" section for troubleshooting. Resolve any problems noted before issuing patient reports.

#### **6.6** Documentation

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- 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.

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#### 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.
- Linearity must be verified with each new lot and at least every six months.
- 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.
- Consult the Laboratory QC Program for complete details.

#### 7. EQUIPMENT and SUPPLIES

#### 7.1 Assay Platform

STA® Compact – Analyzer

#### 7.2 Equipment

- Refrigerator capable of sustaining 2–8°C.
- Freezer capable of sustaining range not to exceed -20 to -70°C.
- Centrifuge calibrated for preparing platelet-poor plasma

#### 7.3 Supplies

- Cuvette Roll Diagnostic Stago
- STA brass adapters
- Plastic micro cups
- STA Mini-Reducer
- Plastic transfer pipettes
- Micro adapter
- Glass micro cups

#### 8. PROCEDURE

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

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Form revised 10/31/02

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# 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	At the start of each shift, verify instrument temperatures and availability of cuvettes and cleaner solution by accessing the System Status screen from the main bar.
2	Record the temperatures on the maintenance sheet. If the reagent arm 2, measuring block, or reagent drawer temperatures are out of range, corrective action must be taken prior to patients being run.
3	Make sure that there is an adequate supply of reagents in the analyzer, and they are in date.
4	Load cuvettes and cleaner/wash solution on the analyzer if needed.

8.2	Analytical Procedure
1	Refer to START-UP procedure for STA® Compact before running patient
	specimens on the STA® Compact at the start of each shift.
2	Request quality control. Through MAIN MENU under CALIB. /CONTROL select
	QUALITY CONTROL and press <b>Enter ←</b> . Cursor to the D-Dimer test.
	Select D-Dimer by pressing <b>F1</b> and then <b>F10.</b> Type in your Access Code to run the QC.
3	Load patients' samples: Access the sample drawer(s) through the MAIN MENU,
	under LOADING, Select Sample, press <b>Enter</b> ←. After the drawer opens, identify the
	type of specimen, such as micro sample (press <b>F8</b> ), or stat (press <b>F12</b> ). Identify the
	sample by bar coding or manually entering on the keyboard the patient identification
	number and then placing the specimen into the drawer.
4	In MANUAL MODE, the operator must order the test(s) from the Selection menu or
	from the Recorded Profile/s Cursor to the test and press <b>Enter</b> to select. When all
	tests are ordered, press F10 to save.
5	In AUTO MODE, the STA®/STA® Compact will automatically order the test(s)
	selected in the AUTO MODE profile.
6	If TELELOADING is selected as the AUTO MODE profile, the STA®/STA® Compact
	will query the host computer and download the test(s) as well as assign the status (i.e.
	stat).
7	As soon as the sample drawer closes, the TEST STATUS screen will appear. If
	there is not enough reagent(s) to run the test(s), the suspect reagent(s) will appear in red
	with the amount of depletion. This depletion of reagent will BLOCK the SAMPLE
	PIPETTING. When this occurs, add the necessary reagent(s) to run the samples by responding N (NO) to the warning message 'NEW TESTS ARE DELAYED -
	REACTIVATE?' Reagents can then be loaded in the drawer. By responding
	Y (YES) to the warning message 'NEW TESTS ARE DELAYED -
	REACTIVATE?', the instrument will continue to perform all tests for which there
	is sufficient reagent (i.e. while waiting for reagents to stabilize after reconstitution)

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8.2	Analytical Procedure	
8	If the patients' results fall outside the assay reportable range <b>4.0 µg/mL</b> the instrument will automatically do a 1:5 dilution on the samples in question. This auto dilution will let the instrument report results up to <b>20.0 µg/mL</b> .	
9	All patient results are displayed on the TEST PANEL screen and automatically print out and transmit if selected on the system status menu.	
10	For results in question that need operator intervention, cursor to the identification number in the TEST PANEL screen and press enter. This will display the FILE PROCESSING screen. Follow the options on the left-hand side of the screen (i.e. <b>F3</b> - rerun test).	

8.3	Reagent and QC Loading Instructions	
1	When Reagent/QC material is reconstituted and ready for use proceed to step 2	
2	From Home Page select <b>Loading</b> – <b>Products</b>	
3	Scan the Reagent/QC	
4	The Instrument will ask whether the volume is correct, or it needs to be modified.	
5	Accept or modify the volume then press <b>Enter</b>	
6	Load the Reagent/QC which was just scanned.	
	<b>Note:</b> Neoplastine CI Plus reagent vial requires to be sitting in the position which is systematically stirred by a lateral movement.	

#### 9. CALCULATIONS

- 1. The STA $^{\otimes}$  Compact automatically plots the results in delta OD off of a standard curve and converts the results to  $\mu g/ml$  FEU.
- 2. The assay uses the sample undiluted. If the result is greater than the reportable range, 4.0, a dependent test with a 1:5 dilution will be ordered to take the reportable range to 20.0. The STA® Compact automatically corrects the result for the dilution change.

#### 10. REPORTING RESULTS AND REPEAT CRITERIA

#### 10.1 Interpretation of Data

N/A

#### 10.2 Rounding

No rounding is necessary. The instrument reports results up to two decimal points.

#### 10.3 Units of Measure

μg/mL FEU

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#### 10.4 Clinically Reportable Range (CRR)

 $0.27 - 20.0 \,\mu g/mL \, FEU$ 

#### 10.5 **Repeat Criteria and Resulting**

The printout from the STA Compact is reviewed for repeat criteria and samples are repeated if needed. Results will be transmitted to the LIS and released using the OEM function.

IF the result is	THEN
< Mmin	Repeat, check for clots. If result is still <mmin, as<="" report="" td=""></mmin,>
	<0.27 μg/mL FEU-REP
> Mmax	Check for clots, repeat using the D-Di 1:5 test
If D-Di 1:5 is quantifiable	Report the result with comment REP
If D-Di 1:5 is > Mmax	Repeat. Report the result as > 20.0 µg/mL FEU-REP

For any of the above situations, be sure the specimen is not under-filled or over-filled, then check the Hematocrit (HCT) result. If the HCT is greater than 55%, refer to appendices A and B for special tube preparation.

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

#### 11. **EXPECTED VALUES**

#### 11.1 **Reference Ranges**

 $\leq 0.5 \,\mu \text{g/mL FEU}$ 

#### 11.2 **Critical Values**

None established

#### 11.3 **Standard Required Messages**

The following comment is automatically added to the report by the LIS:

Less than  $\leq 0.50 \mu g/mL$  FEU = Negative Greater than  $>0.50 \mu g/mL$  FEU = Positive

Positive results are non-specific and are seen in a variety of conditions including DVT, pulmonary embolism, recent surgery, cancer, trauma and pregnancy. Values

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greater than  $0.50 \,\mu\text{g/mL}$  FEU may also be seen in otherwise healthy patients >70 years of age.

Reference:

van der Graaf F, et. al.

Exclusion of Deep Venous Thrombosis with D-Dimer Testing

Thromb Haemost. 2000;83:191-198

#### 12. CLINICAL SIGNIFICANCE

D-Dimer quantitative assay detects the presence of Disseminated Intravascular Coagulation (DIC). In DIC the fibrinolytic system is activated and therefore the D-Dimer level increases. D-Dimer assays can help in the diagnosis of DIC in these cases. It is established that a normal D-Dimer level is an important factor to rule out the diagnosis of deep venous thromboses (DVT) or pulmonary embolisms (PE). The decrease of D-Dimer levels during heparin therapy for a DVT allows the monitoring of evolution and prognosis of the thrombosis. This decrease reflects the quality of the endogenic thrombolysis. The D-Dimer level increases during the activation states of coagulation because they induce the production of thrombin which is followed by the formation of fibrin and leads to fibrinolysis, the latter being most frequently reactive. The D-Dimer level thus increases following coagulation activation.

Increased levels of D-Dimer have been reported in post-operative period, cancers, cirrhosis, and hemorrhages.

#### 13. PROCEDURE NOTES

• **FDA Status:** Approved/cleared

• Validated Test Modifications: None

- 1. The detection threshold of the STA® Liatest® D-Dimer on the STA® Compact is 0.27  $\mu$ g/mL FEU. The printout limits are pre-set at 0.27 4.00  $\mu$ g/mL FEU. When a dependent test is set-up to extend the reportable range of the main test, the printout limit should be extended to 20.00  $\mu$ g/mL FEU.
- 2. The STA<sup>®</sup> Liatest<sup>®</sup> D-Dimer results are expressed in FEU, Fibrinogen Equivalent Units. By definition, an FEU is the quantity of fibrinogen initially present that leads to the observed level of D-Dimer. In general, the actual quantity of D-Dimer is approximately half of an FEU.
- 3. A >Mmax result on the primary assay dilution (1:1 dilution) indicates a result that is greater than  $4.00 \mu g/mL$  FEU. In this case the analyzer will automatically do a 1:5 dilution to obtain the result.

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#### 14. LIMITATIONS OF METHOD

#### 14.1 Analytical Measurement Range (AMR)

 $0.27 - 4.00 \,\mu g/mL \; FEU$ 

#### 14.2 Precision

Different plasmas were used for the intra assay and inter assay reproducibility studies on the STA® Compact.

	Repeat	tability	Within-laboratory Precision		
Sample	Sample 1	Sample 2	Sample 1	Sample 2	
Mean (µg/mL)	0.67	2.20	0.67	2.20	
SD (µg/mL)	0.042	0.049	0.049	0.085	

#### 14.3 Interfering Substances

- 1. Cloudy plasmas may lead to an under-estimation of the D-Dimer level. Ensure that the absorbance value at 540 nm of the plasma diluted 1:6 with STA® Owren-Koller buffer is < 0.35.
- 2. Concentration of Fibrinogen Degradation Products greater than 15  $\mu$ g/ml may lead to an over-estimation of the D-Dimer level.
- 3. The presence of rheumatoid factor at a level greater than 50 IU/ml may lead to an over-estimation of the D-Dimer level.
- 4. The STA<sup>®</sup> Liatest<sup>®</sup> D-Dimer is insensitive to fibrinogen and the E fragment. A low cross-reactivity is observed with the D fragment.
- 5. The STA® Liatest® D-Dimer is insensitive to the following substances: hemoglobin (up to 2 g/l); conjugated bilirubin (up to 290 mg/L); unconjugated bilirubin (up to 200 mg/L); unfractionated heparin (up to 2 IU/mL; LMWH (up to 2 anti-Xa IU/ml)

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

N/A

#### 15. SAFETY

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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

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work typically focuses on the hazards of working with specimens and chemicals, we must also control other important hazards.

- 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. Laboratory Quality Control Program
- 2. Laboratory Safety Manual
- 3. Safety Data Sheets (SDS)
- 4. Hemolysis, Icteria and Lipemia Interference (Lab policy)
- 5. Repeat Testing Requirements (Lab policy)
- 6. STA Compact Operating Instructions, Coagulation procedure
- 7. Verification of Platelet Poor Plasma, Coagulation procedure
- 8. Current package insert for STA® LIATEST D-DIMER

#### 17. REFERENCES

- 1. van der Graaf F, et. al., Exclusion of Deep Venous Thrombosis with D-Dimer Testing, Thromb Haemost. 2000;83:191-198
- 2. Diagnostic Stago STA® LIATEST D-DIMER package insert: Revised March 2015.
- 3. STA LIATEST Control N + P (REF 00526): citrated control plasmas normal and abnormal levels; Control Plasmas for Assays of Coagulation Parameters on STA<sup>®</sup>, Revised 03/2015.
- 4. STA<sup>®</sup> Compact Operators Manual. STA<sup>®</sup> DSI-TSD-SM August 2004, STA<sup>®</sup> DSI-TSD-US April 2003, and V1.3 revised February 2003.
- 5. Diagnostic Stago STA® Owren-Koller Buffer Solution for Coagulation Tests. Revised: May 2014.

#### 18. REVISION HISTORY

Version	Date	Section	Reason	Reviser	Approval
			Supersedes G003.006		
000	6/11/12	2.0	Update to match package insert	J.Buss	J. Buss, RSL
000	6/11/12	3.2	Add frozen temperature	J.Buss	J. Buss, RSL

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000	6/11/12	4.1	Remove Millipore water	J. Buss	J. Buss, RSL
000	6/11/12	4.2	D-DI reagent open stability edited	J.Buss	J. Buss, RSL
000	6/11/12	6.3	Add QC performed after maintenance	J.Buss	J. Buss, RSL
000	6/11/12	15	Update to standard wording	L. Barrett	J. Buss, RSL
001	6/3/14		Update owner	L Barrett	R SanLuis
001	6/3/14	3.1	Add reference to Appendices	A Chini	R SanLuis
001	6/3/14	3.2	Update tube volumes	A Chini	R SanLuis
001	6/3/14	4.2	Change storage temp and prep for buffer	A Chini	R SanLuis
001	6/3/14	6.2	Add step to print QC	A Chini	R SanLuis
001	6/3/14	10.4	Change CRR lower value	A Chini	R SanLuis
001	6/3/14	10.5	Add instruction for Hct >55	A Chini	R SanLuis
001	6/3/14	13, 14.1	Change lower value of analytical range	A Chini	R SanLuis
001	6/3/14	14.3	Update to match package insert	A Chini	R SanLuis
001	6/3/14	16	Update titles	L Barrett	R SanLuis
001	6/3/14	19	Add Appendix A and B	A Chini	R SanLuis
001	6/3/14	Footer	Version # leading zero's dropped due to new EDCS in use as of 10/7/13.	L Barrett	R SanLuis
2	4/7/15	6.4, 6.6	Replace LIS with Unity Real Time	L Barrett	R SanLuis
2	4/7/15	10.5	Correct <mmin reporting="" td="" value<=""><td>L Barrett</td><td>R SanLuis</td></mmin>	L Barrett	R SanLuis
3	1/12/16	3.2	Change whole blood to plasma	A Chini	R SanLuis
3	1/12/16	4	Delete reagent grade water	A Chini	R SanLuis
3	1/12/16	5	Add explanation for Liatest D-dimer and Bio-Rad QC. Add Liatest QC info.	A Chini	R SanLuis
3	1/12/16	6.1, 6.2	Update to Bio-Rad QC	A Chini	R SanLuis
3	1/12/16	6.2	Add instruction for loading onboard	A Chini	R SanLuis
3	1/12/16	6.5	Update to match auto verification system	A Chini	R SanLuis
3	1/12/16	6.7	Add TEa criteria, linearity check with log change, QC sent to Bio-Rad monthly	A Chini	R SanLuis
3	1/12/16	7.3	Add micro adapter and glass micro cups	A Chini	R SanLuis
3	1/12/16	14.2	Update precision data	A Chini	R SanLuis
3	1/12/16	17	Add Bio-Rad QC	A Chini	R SanLuis
4	7/25/16	Header	Add other sites	L Barrett	R SanLuis
4	7/25/16	5.1	Removed Bio-Rad QC information	A Chini	R SanLuis
4	7/25/16	6.1, 6.2	Replace Bio-Rad QC with STA Coag Controls	A Chini	R SanLuis
4	7/25/16	6.7	Remove QC submission to Bio-Rad	A Chini	R SanLuis

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Germantown Emergency Center

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Version	Date	Section	Reason	Reviser	Approval
4	7/25/16	8.3	Add QC/Reagent Loading Instructions	A Chini	R SanLuis
4	7/25/16	11.3	Move report comment from 10.5	L Barrett	R SanLuis
4	7/25/16	17	Remove Bio-Rad insert	A Chini	R SanLuis

### 19. ADDENDA

- A. Instructions for Preparing Collection Tube for Hematocrit >55%
- B. Phlebotomist Instructions for Blood Collection

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## Appendix A

#### **Instructions for Preparing Collection Tube for Hematocrit >55%**

#### **Explanation**:

Polycythemia is a disease state in which the proportion of blood volume that is occupied by red blood cells increases - basically when Hematocrit (HCT) is greater than 55%. It can cause prolonged coagulation results.

When a prolonged coagulation result is obtained, check the specimen for a clot first.

If the specimen is not clotted, be sure the specimen is not under-filled or over-filled, then check the HCT result.

If a HCT result of greater than 55% is obtained, immediately notify the doctor or attending nurse and ask for a redraw using a special tube prepared by the lab.

#### To prepare a special tube in the lab use the following instructions and formula:

The anticoagulant volume in the collection tube must be adjusted to obtain a 9:1 ratio of blood to Sodium Citrate. Under or over-filling of the specially prepared collection tube is not acceptable. The vacuum in the collection tube will be broken to adjust the volume of collection anticoagulant. Because of this special collection technique, the stability for these whole blood specimens is reduced to four (4) hours after collection.

#### Formula to calculate the anticoagulant volume is:

Anticoagulant in  $mL = [(100 - HCT) / (595 - HCT)] \times Volume of blood$ 

Example 1: Specimen with a 70% HCT in a 2.7 mL tube:

Patient with HCT of 70%
Using a 2.7 mL tube
Anticoagulant in mL = $[(100 - 70) / (595 - 70)] \times 2.7 = 0.15 \text{ mL or } 150 \text{ uL}$
Pipette a 2.7 mL tube in a way to leave only 150 uL of anticoagulant in there.
A 2.7 mL tube contains 0.3mL anticoagulant; therefore remove 0.15mL

Example 2: Specimen with a 70% HCT in a 1.8 mL tube:

Patient with HCT of 70%
Using a 1.8 mL tube
Anticoagulant in mL = $[(100 - 70) / (595 - 70)] \times 1.8 = 0.1 \text{ mL or } 100 \text{ uL}$
Pipette a 1.8 mL tube in a way to leave only 100 uL of anticoagulant in there.
A 1.8 mL tube contains 0.2mL anticoagulant; therefore remove 0.1mL

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# Example 3: Specimen with a 60% HCT in a 2.7 mL tube:

Patient with HCT of 60%		
Using a 2.7 mL tube		
Anticoagulant in $mL = [(100 - 60) / (595 - 60)] \times 2.7 = 0.2 \text{ mL or } 200 \text{ uL}$		
Pipette a 2.7 mL tube in a way to leave only 200 uL of anticoagulant in there.		
A 2.7 mL tube contains 0.3mL anticoagulant; therefore remove 0.1mL		

Example 4: Specimen with a 60% HCT in a 1.8 mL tube:

Patient with HCT of 60%
Using a 1.8 mL tube
Anticoagulant in mL = $[(100 - 60) / (595 - 60)] \times 1.8 = 0.13 \text{ mL or } 130 \text{ uL}$
Pipette a 1.8 mL tube in a way to leave only 130 uL of anticoagulant in there.
A 1.8 mL tube contains 0.2mL anticoagulant; therefore remove 0.07mL

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#### Appendix B

#### **Phlebotomist Instructions for Blood Collection**

The technologist will prepare a special tube in which the anticoagulant has been adjusted, therefore the tube is not vacuumed. The technologist will inform the phlebotomist of the exact amount of blood needed to fill the tube.

#### **Equipment and Supplies**

Latex gloves

Latex free tourniquet

Latex free Band Aid or Tape

Alcohol Prep (70% alcohol)

2x2 sterile gauze

Collection tube

Blood Collection Set 21 or 23 gauge winged set

**Blood Transfer Device** 

3mL syringe

Biohazard bag

Biohazard sharps container

LIS collection list and label/Lab requisition

#### **Collection Steps**

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- 1. Introduce yourself to the patient by stating your first and last name.
- 2. Positively identify the patient according to the SOP 'Patient Identification', Phlebotomy procedure manual.
- 3. Wash hands. Apply gloves.
- 4. Explain the procedure to the patient and obtain patient's consent to draw blood.
- 5. Collect equipment and correct technologist-provided collection tube.
- 6. Assemble equipment and break needle and syringe seals in the presence of the patient.
- 7. Apply tourniquet about midway between the elbow and the shoulder 3-4 inches above the venipuncture site). Place patient's arm in a downward position to prevent reflux of 'backflow' of blood from the tube into the venous system. Ask the patient to close hand gently.
- 8. Palpate/feel for vein locating a vein that will flow fast (reducing the possibility of the blood clotting).
- 9. Clean the area for venipuncture with a 70% alcohol pad decontaminating the collection site.
- 10. Allow the area to air-dry completely.
- 11. Assemble the 21 or 23 gauge winged set to the 3mL syringe. Pull back the plunger to dispel all the air out of the syringe.
- 12. With the bevel up, align the needle with the vein while holding the skin taut. Insert the needle at a 15-30 degree angle with the skin. Remove your hand from drawing the skin taut. Grasp the syringe and draw back bringing the plunger tip to the exact amount of blood requested by the technologist.
- 13. Release the tourniquet. Ask the patient to open hand.

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- 14. Place gauze above the puncture site and remove the needle while simultaneously applying pressure on the puncture sit. Firmly activate needle safety shield, a click must be heard to ensure that the safety shield is secure.
- 15. Remove 21 or 23 gauge winged set from syringe.
- 16. Attach the blood-filled syringe to the Blood Transfer Device.
- 17. Connect the Blood Transfer Device to the un-vacuumed tube, provided by the technologist, and slow and gently fill the collection tube. DO NOT FORCE blood into tube. Pressure can lead to tube explosion and blood exposure.
- 18. Place the cap on the tube and invert a few times to make sure the anticoagulant is mixed with blood.
- 19. Dispose of all blood collection equipment into the nearest sharps container. DO NOT disassemble the syringe from the Blood Transfer Device.
- 20. Dispose of all other used materials in appropriate container and wash hands.
- 21. Label the sample with the LIS collection label and write the time, date, and your tech code.
- 22. Transport specimen to the Lab.

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