### TRAINING UPDATE

Lab Location: Department: GEC, SGMC & WAH Core Lab 
 Date Distributed:
 5/22/2019

 Due Date:
 6/28/2019

 Implementation:
 6/2/2019

## **DESCRIPTION OF PROCEDURE REVISION**

Name of procedure:

## CSF Cell Count and Differential, Manual Method SGAH.H08 v6

**Description of change(s):** 

## TWO major changes to SOP:

Section	Reason
2	Modified local test codes
8.1	Specify reporting for appearance & color
	<ul> <li>Appearance evaluated before centrifugation, color after</li> <li>Added codes to match Sysmex/DI reporting</li> </ul>
8.3	Specified diff is only required when cell count is greater than 5 (no diff is needed if cell count is less than/equal to 5)

This revised SOP will be implemented on June 2, 2019

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

Technical SOP

Title	CSF Cell Count and Differen	itial, Manual N	Iethod
Prepared by	Cynthia Reidenauer	Date:	3/21/2011
Owner	Robert SanLuis	Date:	11/26/2013

Laboratory Approval	Local Effective Date:	
Print Name	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
Cell Counts, Total RBC and Total Nucleated Cells, CSF (tube specific)	Manual/Microscopic	CT1, CT2, CT3, CT4

#### Synonyms/Abbreviations

CSF Count, Cerebrospinal Fluid Cell Count, Spinal Fluid Count

#### Department

Hematology

### 2. ANALYTICAL PRINCIPLE

Gross examination of the specimen is performed to determine the appearance. A microscopic examination is performed for the Total Nucleated Cell count (TNC) and Red Blood Cell count (RBC). Smears for cell identification are prepared using cyto-centrifuge or conventional centrifuge. Nucleated cell identification/ differential counts are done on Wright's Stained smears prepared using a cyto-centrifuge or smeared sediment from clinical centrifugation.

### **3. SPECIMEN REQUIREMENTS**

#### **3.1** Patient Preparation

Component	Special Notations
Fasting/Special Diets	Not Applicable
Specimen Collection and/or Timing	Not Applicable
Special Collection Procedures	<ul> <li>Specimens are collected in sterile tubes labeled in the order in which they are withdrawn (1, 2, 3, 4).</li> <li>Tube 1 is used for color, appearance, cell count and Chemistry tests</li> <li>Tube 2 is used for Serology tests</li> <li>Tube 3 is used for color, appearance, cell count and diff</li> <li>Tube 4 is used for Microbiology</li> <li>Note: If there is a Cytology order, process core lab testing per 3 tube protocol and use tube 4 for Cytology.</li> <li>If 3 tubes are received:</li> <li>Tube 1 is used for Serology tests, color, appearance, cell count and Chemistry tests</li> <li>Tube 2 is used for Serology tests, color, appearance, cell count and the chemistry tests</li> <li>Tube 3 is used for Microbiology</li> <li>Note: If there is a Cytology order, after Microbiology has taken their sample from tube 3, send remainder of tube 3 for Cytology.</li> <li>If less than 3 tubes are received, contact the physician for specific tests to be performed.</li> </ul>
Other	Not applicable

Criteria			
Type -Preferred	Tube #1 and #3 (See section 3.1 if less than 4 tubes)		
-Other Acceptable	None		
<b>Collection Container</b>	Sterile Plastic Co	onical Tube	
Volume - Optimum	2.0 mL		
- Minimum	0.5 mL		
Transport Container and Temperature	Sterile Plastic Conical Tube at room temperature		
Stability & Storage	Room	Process Immediately. Rapid	
Requirements	Temperature:	deterioration and cell lysis occurs on	
		prolonged standing in CSF.	
	Refrigerated:	Same as above.	
	Frozen:	Unacceptable	
Timing Considerations	Not Applicable		
Unacceptable Specimens	<b>Clotted specime</b>	ns: Perform counts and append the code	
& Actions to Take	SCLOT (Specimen contains clots, counts may not be		
	accurate).		
	Specimens received after 24 hours: Perform counts and		
	append the code SAGE (Counts may not be accurate due		
	to the age of the specimen).		
	Due to nature of	specimen, do not reject, unless frozen.	
	If the specimen is received frozen: Cancel the test with		
	the reason code SFRZ (Specimen unsuitable for assay;		
	received frozen).	Notify a caregiver and document in the	
	LIS.		
Compromising Physical	None defined		
Characteristics			
Other Considerations	None defined		

#### **3.2** Specimen Type & Handling

NOTE: Labeling requirements for all reagents, calibrators and controls include: (1) Open date, (2) Substance name, (3) Lot number, (4) Date of preparation, (5) Expiration date, (6) Initials of tech, and (7) Any special storage instructions. Check all for visible signs of degradation.

#### 4. **REAGENTS**

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.

#### 4.1 Reagent Summary

Reagents	Supplier & Catalog Number
Rinse	ELITechGroup SS-071 A
Thiazin	ELITechGroup SS-071 B

Reagents	Supplier & Catalog Number
Eosin	ELITechGroup SS-071 C
Methanol	ELITechGroup SS-MEOH
Aerofix (Additive for Methanol)	ELITechGroup SS-148
0.9% Saline	Thermo 0.9% Saline cat # 23535435
22% Albumin (Obtain from Blood Bank)	Immucor CE 0088
RAL Diff-Quik Stain Pak	RAL Diagnostics #720555-0000
0.005% Methylene Blue Diluting Fluid	Chantilly reagent room

## 4.2 Reagent Preparation and Storage

Reagent A	ELITechGroup Aerospray Rinse
Reagent B	ELITechGrout Aerospray Thiazin
Reagent C	ELITechGroup Aerospray Eosin
Container	Plastic Bottle
Storage	5-30°C
Stability	Manufacturer's expiration date
Preparation	Ready to use

Reagent	Wescor Aerospray Aerofix
Container	Plastic Bottle
Storage	15-30°C
Stability	Manufacturer's expiration date
Preparation	Add 10 mL to Methanol and mix well prior to use.

Reagent	0.9% Saline (Obtain fresh daily from Blood Bank)	
Container	Plastic Bottle	
Storage	15-30°C	
Stability	24 hours, working supply in hematology. Open expiration on container in Blood Bank is 30 days.	
Preparation	Ready to use	

Reagent	22% Bovine Albumin
Container	Glass Bottle 10 mL
Storage	1 - 10C for long term storage
Stability	Stable until expiration date on the bottle. If turbid, discard.
Preparation	Ready to use

Reagent	RAL Diff-Quik Stain Pack
Container	Plastic Bottle
Storage	15 - 25C

Stability	Unopened: Until expiration date on box label.	
	<b>Opened:</b> Remains stable 2 months after opening.	
	Keep away from light.	
Preparation	Ready to use	
	1	
Reagent	0.005% Methylene Blue Diluting Fluid. Obtain when needed	
	from the reagent room in Chantilly.	
Container	Brown Glass Bottle	
Storage	15-30°C	
Stability	Manufacturer's expiration date.	
	Aliquot small amount to use when needed. Stability of aliquot is 24	
	hours.	
Preparation	Ready to use	

## 5. CALIBRATORS/STANDARDS

Not applicable

## 6. QUALITY CONTROL

#### 6.1 Controls Used

Control	Supplier & Catalog Number
Cell-Chex L1-UC, L1-CC and L2 (2mL each)	Streck Laboratories, Inc. Cat # 212431
Cell-Chex L1-UC, L2	Streck Laboratories, Inc. Cat # 212420
Cell-Chex L1-CC	Streck Laboratories, Inc. Cat # 212430

## 6.2 Control Preparation and Storage

Control	Cell-Chex Level L1-UC, L1-CC and L2	
Preparation	None required. It is not necessary to warm the controls to room	
	temperature before using.	
Storage/Stability	• Store upright at 2-10°C	
	Closed-vial stability 180 days	
	Open-vial stability 30 days	

### 6.3 Frequency

• Cell Count and Cytocentrifuge QC is performed every 8 hours of patient testing for manual body fluid counting and per technologist.

QC menu each level of controls is as follows:

- L1-UC perform cell count
- L1-CC perform a cytospin differential
- L2 perform cell count

- Automated or Manual stain method comparison is performed once per day. A smear must be reviewed on a daily basis to verify that the staining is adequate for differential of the various cells. The result of this review is documented in the manual Hematology QC book.
- **Diluting fluid** must be checked daily for contamination and documented. Refer to section 8.2

## 6.4 **Tolerance Limits**

## a) Cell count by Manual Hemacytometer:

QC values for Manual Hemacytometer are lot specific so check package insert for lot number and expiration date. The lot number and ranges for each lot in use will be available on the Cell Chex Log.

### b) **Differential %:**

QC values for Differential % are lot specific so check package insert. The lot number and ranges for each lot in use will be available on the Cell Chex Differential Log.

## c) Corrective Action:

- 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 <u>reanalyzed</u>.
- Corrective action documentation must include the following: The QC rule(s) (or specific QC criteria) 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.

## d) Review of QC

- Refer to SOP Laboratory Quality Control Program for more details.
- Upon weekly and monthly review of QC, if the SD's or CV's are greater than the above maximums, investigate the cause for the imprecision and document implementation of corrective actions.

## 6.5 Documentation

QC results are recorded on the Cell Chex QC log sheets.

## 6.6 Quality Assurance Program

The laboratory participates in CAP proficiency testing.

## 7. EQUIPMENT and SUPPLIES

## 7.1 Assay Platform

Not applicable

## 7.2 Equipment

Microscope Aerospray Cytocentrifuge CytoTek centrifuge (GEC only)

## 7.3 Supplies

Disposable Pipettes Hemacytometer (disposable) C-CHIP MLA pipette and tips Disposable tubes Cover glass Microscope slides Petri dish Applicator sticks Cytopro Cyto-Tek 2500 (GEC)

## 8. **PROCEDURE**

# 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.

PROMPT examination after receipt of CSF in the laboratory is ESSENTIAL for accurate results. Cellular disintegration may occur if there is a delay in testing. If delay is unavoidable, refrigerate until testing can commence.

## 8.1 Color and Appearance

Step	Exa	mination for Ap	pearance and Color	
1.	Examine the CSF for appearance and color.			
2.	Appearance: Indicate	Appearance: Indicate what the fluid looks like before centrifugation; use the		
	following codes:			
	Description	Code	Description	Code
	Clear	CLEAR	Turbid	TURB
	Cloudy	CLDY	Bloody	BLDY
	Slightly Cloudy	SLCLDY		
3.	<b>Color</b> : Centrifuge an aliquot for the time and speed posted on centrifuge to remove the cellular elements. Examine the supernatant and report the color using the following descriptions and codes:			
	Description	Code	Description	Code
	Colorless	COLR	Brown	BRWN
	Yellow	YEL	Red	RED
	Pink	PINK		

## 8.2 Concentration

Step	Specimen Preparation
1.	Place a drop of Methylene Blue diluting fluid on a slide and coverslip. Examine under 100X for contamination with artifacts, crystals or bacteria,
	replace fluid if necessary. Record the examination on the Cell Count Worksheet. If the diluting fluid is acceptable to, proceed to specimen dilution.
2.	<ul> <li>Inspect specimen to determine the appropriate dilution.</li> <li>a. All specimens will be diluted with 0.005% Methylene Blue Diluting fluid.</li> <li>b. The minimum dilution is 1:2. This will ensure distinction between RBC and TNC. Red Cells will not pick up the methylene blue stain and will appear agranular. Methylene Blue allows the visual distinction of nucleated</li> </ul>
	cells by staining the granules a faint blue.
3.	Mix specimen well and make the appropriate dilution. Refer to dilution tables below.

Step	1:2 Dilution
1.	Perform the diluting fluid check as described above. If the diluting fluid is
	acceptable to use, proceed to dilution of the specimen.
2.	Mix specimen well. Using a 100µL pipette, add 100µL of body fluid to 100µl
	of Methylene Blue Diluting Fluid. Mix dilution well. Let sit 10-15 minutes.
	Dilution Factor is 2
3.	Charge the two chambers of the hemacytometer by touching the tip of the
	pipette to the coverslip edge where it meets the chamber floor. The chamber
	will fill by capillary action if the hemacytometer and coverslip are clean.
4.	If the hemacytometer is overcharged, it must be discarded and a fresh one
	used.
5.	Place the charged hemacytometer in a humidified Petri dish for 10 minutes to
	allow the cells to settle.
6.	Place the hemacytometer on the microscope and examine. The area to be
	counted is adjusted according to the sample.
	• If less than 20 cells are present in one square, count all the squares.
	• If greater than 20 and less than 200 cells are present in one square,
	count the four corner squares only.
	• If greater than 200 cells are present in one square count 5 of the 25
	squares in the middle square.
	ALWAYS USE THE AVERAGE COUNT FROM BOTH SIDES OF THE
	CHAMBER IN THE FORMULA. Count the total number of rbcs and
	nucleated cells present on both sides. The sides should agree within 20%.
7.	Calculate the total number of RBCs and nucleated cells. Follow instructions
	on the Cell Count Worksheet to calculate results.
8.	All calculations must be recorded on worksheet.

Step	Diluted Specimen 1:10
1.	Perform the diluting fluid check as described above. If the diluting fluid is
	acceptable to use, proceed to dilution of the specimen.
2.	Mix specimen well. Using a 100µL pipette, add 100µL of body fluid to 900µl
	of Methylene Blue Diluting Fluid. Mix dilution well. Let sit 10-15 minutes.
	Dilution Factor is 10
3.	Charge a counting chamber (one pipette per side), using proper technique.
4.	Place in a Petri dish for about 10 minutes to let the cells settle.
5.	For counting guidelines, follow steps 6 through 8 for 1:2 Dilution

Step	Diluted Specimen 1:20
1.	Perform the diluting fluid check as described above. If the diluting fluid is
	acceptable to use, proceed to dilution of the specimen.
2.	Mix specimen well. Using a 50µL pipette, add 50µL of body fluid to 950µl of
	Methylene Blue diluting fluid. Mix dilution well. Let sit 10-15 minutes.
	Dilution Factor is 20
3.	Charge a counting chamber (one pipette per side), using proper technique.
4.	Place in a Petri dish for about 10 minutes to let the cells settle.
5.	For counting guidelines, follow steps 6 through 8 for 1:2 Dilution

Step	Diluted Specimen 1:50
1.	Perform the diluting fluid check as described above. If the diluting fluid is
	acceptable to use, proceed to dilution of the specimen.
2.	Mix specimen well. Using a 20µL pipette, add 20µL of body fluid to 980µl of
	Methylene Blue Diluting Fluid. Mix dilution well. Let sit 10-15 minutes.
	Dilution Factor is 50
3.	Charge a counting chamber (one pipette per side), using proper technique.
4.	Place in a Petri dish for about 10 minutes to let the cells settle.
5.	For counting guidelines, follow steps 6 through 8 for 1:2 Dilution

Step	Diluted Specimen 1:100
1.	Perform the diluting fluid check as described above. If the diluting fluid is
	acceptable to use, proceed to dilution of the specimen.
2.	Mix specimen well. Using a 10µL pipette, add 10µL of body fluid to 990µl of
	Methylene Blue Diluting Fluid. Mix dilution well. Let sit 10-15 minutes.
	Dilution Factor is 100
3.	Charge a counting chamber (one pipette per side), using proper technique.
4.	Place in a Petri dish for about 10 minutes to let the cells settle.
5.	For counting guidelines, follow steps 6 through 8 for 1:2 Dilution

### 8.3 Differential Count

IF	THEN
Cell count is $\leq 5$	Do not perform differential. Result with NOTP-; due to an
	insufficient number of cells in the sample.

Form revised 7/01/01

IF	THEN				
Cell count is $>5$	Perform a 5 part differential of 100 cells on a cytocentrifuged				
	specimen using Wescor slide stainer, or a manual stain (GEC).				
	The nucleated cells are classified and reported as a percentage.				
	Examine smear for the presence of immature or abnormal cells,				
	crystals and bacteria. If abnormal or immature cells are noted,				
	second technologist must also perform a differential and then				
	refer slide(s) to a Pathologist for review.				

## 8.4 Cytospin

Step	Cytospin						
1.	Assemble sample of	hamber and gl	hamber and glass microscope slide in the Wescor				
	Aerospray cytocentr	ifuge carousel. A	t GEC, follow Cytopsin procedure.				
2.	IF		THEN				
	Nucleated cell	Place 3-5 drops	s of fluid plus 1 drop of albumin into a				
	count is <300	disposable cyte	ofunnel and place into the Cytospin				
		centrifuge. Th	ne albumin is used to make the cells				
		adhere to the sli	de better before the staining procedure.				
	Nucleated cell	Cells/ µL	Dilution				
	count is >300	301-700	1:2 (5 drops CSF + 5 drops saline)				
		701-1500	1:5 (2 drops CSF + 10 drops saline)				
		1501-3000	1:10 (2 drops CSF + 20 drops saline)				
		>3000	1:20 (2 drops CSF + 40 drops saline)				
		Mix dilution we	ell and place 3-5 drops into the Cytospin				
		funnel. Add 1 c	lrop of albumin.				
3.	Centrifuge Sample:						
	See procedure Aerospray Hematology Slide Stainer Cytocentrifuge						
	(SGMC/WAH) or Cy	Cytospin CSF/Body Fluid Slide Preparation (GEC) as					
	appropriate.						
4.	Stain slide using the	Aerospray stainer	or Diff Quick Stain Pack as appropriate				

## 9. CALCULATIONS

Formula for Hemacytometer

 $\frac{\text{Cells Counted}}{\# \text{ of squares counted}} \times 10 \times \text{dilution}$ 

## **10. REPORTING RESULTS AND REPEAT CRITERIA**

## **10.1** Interpretation of Data

None required

### 10.2 Rounding

Results for cell counts are rounded to whole numbers.

#### **10.3** Units of Measure

Parameter	Units
RBC	Cells/µL
TNC (WBC-BF)	Cells/µL
Differential Counts	%

#### **10.4** Clinical Reportable Range

Not applicable

### **10.5** Review Patient Data

Since only a few patient samples may be tested in one day, daily review for trends may not be applicable.

### 10.6 Repeat Criteria and Resulting

Any duplicate counts not agreeing within 20% must be repeated.

All CSF counts must be reviewed by a second technologist prior to resulting. Calculations must be rechecked and proper placement and documentation of cell counts on the worksheet must be verified. In addition, once typed into the computer a second tech must verify the proper placement of the counts **PRIOR to accepting the results**.

### Second tech review for Germantown Emergency Center ONLY:

Due to the fact that there is only one person working per shift, if a CSF cell count is performed then it will be the first duty of the next shift tech to review the cell count worksheet and compare it to the results entered into the computer. The reviewing tech will initial that the second tech review was performed.

### **Pathology Review:**

All fluids needing a pathology review are to be taken to the pathologist on call for Hematology. All slides are to be accompanied by an IRA report from the LIS and the Pathologist Slide Review form.

### **Resulting:**

Refer to the addendum *Fluid Keyboard: Accessing Differential Result Entry for CSF* for details to result via the SQ keyboard.

Note: Manual differentials performed due to TEa failures on Sysmex (difference between TC-BF and WBC-BF exceeds the TEa of 20% during Sysmex testing) must be reported via the CSF Cell Counter in DI.

## 11. EXPECTED VALUES

#### **11.1 Reference Ranges**

Parameter / Units of	Both Male and Female				
Measurement	Neonate	Adult			
Color	Colorless				
Appearance	Clear				
RBC - BF cells/µL	None	None			
WBC - BF cells/µL	0 - 30	0 - 5			
Lymphocyte %	<70	<70			
Monocyte %	<30	<30			
Eosinophil %	<10	<10			

Note: TNC are reported in LIS as 'WBC-BF' to align with automated method.

#### 11.2 Critical Values

None established

## 11.3 Standard Required Messages

None established

### **12.** CLINICAL SIGNIFICANCE

	CSF Appearance					
Appearance	Cause	Most Significance				
Crystal Clear		Normal				
Hazy, turbid, cloudy	WBC's; RBC's	Meningitis, Hemorrhage, Traumatic tap				
	Microorganisms	Meningitis				
	Protein	Disorders that affect blood-brain barrier,				
		Productions of IgG within CNS				
Oily	Radiographic					
	Contrast					
	Material					
Bloody	RBC's	Hemorrhage				
Xanthochromic (color)	Hemoglobin	Old Hemorrhage				
		Lysed cells from traumatic tap				
	Bilirubin	RBC Breakdown				
		Elevated serum bilirubin				
	Merthiolate	Contamination				
	Carotene	Increased serum levels				
	Protein	See above				

The CSF is the third major fluid of the body. It provides a physiologic system to supply nutrients to the nervous system, remove metabolic wastes and produce a mechanical barrier to cushion the brain and spinal cord against trauma. Identification of cell types present in the CSF has become a valuable diagnostic aid most frequently associated with meningitis. High WBC counts with neutrophilic majority are associated with bacterial meningitis while lymphocyte/monocyte predominance indicates viral, tubercular, etc., origin. The differential can impart diagnostic information based on abnormal cell types found indicating metastatic carcinoma, central nervous system involvement of leukemia or parasitic infections. Refer to the table below for a more complete list.

Predominant Cells Seen in CSF						
Type of Cell	Major Clinical Significance	Microscopic Findings				
Lymphocyte	Normal	All stages of development may be				
	Viral, tubercular and fungal	found.				
	meningitis					
	Multiple Sclerosis					
Neutrophil	Bacterial meningitis	Granules may be less prominent than				
	Early cases of viral, tubercular, or	in blood.				
	fungal meningitis					
	Cerebral hemorrhage	Cells disintegrate rapidly.				
Monocyte	Chronic bacterial meningitis	Found mixed with lymphocytes and				
	Viral, tubercular, and fungal	neutrophils.				
	meningitis					
	Multiple Sclerosis					
Eosinophil	Parasitic infections	Same appearance as seen in blood.				
	Allergic reactions					
	Intracranial shunts (hydrocephalus)					
Macrophages	Viral and tubercular meningitis	May contain phagocytized RBCs				
	RBC's in spinal fluid	appearing as empty vacuoles or				
		ghost cells and hemosiderin				
		granules.				
Pia arachnoid	Normal, mixed reactions, including	Resemble young monocytes with a				
mesothelial (PAM)	neutrophils, lymphocytes, monocytes	round, not indented, nucleus.				
cells	and plasma cells					
Blast forms	Acute leukemia	Lymphocytes or myeloblasts.				
Plasma cells	Multiple Sclerosis	Transitional and classic forms seen.				
	Lymphocyte reactions					
Ependymal Cells	Normal trauma	Seen in clusters with distinct nuclei				
Choroidal Cells	Diagnostic procedures	and distinct cell walls.				
Malignant Cells	Metastatic carcinoma	Seen in clusters with fusing of cell				
		borders and nuclei.				

### **13. PROCEDURE NOTES**

- FDA Status: Laboratory Developed Test (LDT) without message
- Validated test modifications: not applicable

- Perform cell counts as soon as possible since cells deteriorate with time.
- If there is a clot, perform count on available liquid and make notation in the report. Counts on partially clotted samples may be affected depending whether or not cells are trapped in the clot.
- Low power scanning should be performed on smear to evaluate cell distribution and evaluate for presence of malignant cells.

## 14. LIMITATIONS OF METHOD

Not applicable

### **15. SAFETY**

Refer to your local and corporate safety manuals and Safety Data Sheet (SDS) for detailed information on safety practices and procedures and a complete description of hazards.

### **16. RELATED DOCUMENTS**

- 1. Laboratory Quality Control Program
- 2. Hematology Slide Stainer Cytocentrifuge, Aerospray® Model 7151, SGMC / WAH Hematology SOP
- 3. Cytospin CSF/Body Fluid Slide Preparation, GEC Hematology SOP
- 4. RAL Diff-Quik Stain Kit, Hematology SOP
- 5. Cell Count Worksheet (AG.F12)
- 6. Cell Chex Control Log (AG.F87)
- 7. Pathologist Slide Review Request (AG.F127)

## **17. REFERENCES**

- 1) Body Fluids, Third Edition, Kjeldsberg, C.R., and Knight, J.A., American Society of Clinical Pathologists Press, Chicago, 1993.
- 2) Clinical Hematology and Fundamentals of Hemostasis, Second Edition, Harmening, Denise M., F.A. Davis Company, Philadelphia, 1992.
- 3) Urinalysis and Body Fluids, Edition 2, Strasinger, S.K., F.A. Davis Company, 1989
- 4) Defining CSF WBC Count Reference Values in Neonates and Young Infants, Kestenbaum Ebberson et al Pediatrics 2010;125;257-264
- 5) CSF Analysis, D. Seehusen et al American Family Physician September 15,2003; Vol. 68; Number 6, 1103-1108

### **18. REVISION HISTORY**

Version	Date	Section	Reason	Reviser	Approval
			Supersedes SOP SGAH-WAH H019.000		
000	06/06/12		Update owner	L Barrett	J Buss, RSL

Form revised 7/01/01

#### Adventist HealthCare

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

## Title: CSF Cell Count and Differential, Manual Method

Version	Date	Section	Reason	Reviser	Approval
000	06/06/12	6.1, 6.7	Add diluting fluid check to match Cell Count J Buss . Worksheet		J Buss, RSanLuis
001	11/26/13		Update owner	L Barrett	R SanLuis
001	11/26/13	4	Add Methylene Blue diluting fluid and stain components	L Barrett	R SanLuis
001	11/26/13	6	Update QC material, frequency clarified	L Barrett	R SanLuis
001	11/26/13	7, 8	Remove use of alcohol swabs, filter paper and non disposable hemacytometer,	L Barrett	R SanLuis
001	11/26/13	8.2	Add Methylene Blue as diluting fluid, add process to make each dilution	L Barrett	R SanLuis
001	11/26/13	10.5	Add second review process for GEC	L Barrett	R SanLuis
001	11/26/13	13	Add handling for clots	L Barrett	R SanLuis
001	11/26/13	15	Update to standard wording	L Barrett	R SanLuis
001	11/26/13	16	Add forms, update SOP titles	L Barrett	R SanLuis
001	11/26/13	19	Remove forms	L Barrett	R SanLuis
001	11/26/13	Footer	Version # leading zero's dropped due to new EDCS in use as of 10/7/13.	L Barrett	R SanLuis
2	3/12/14	8.3	Correct 1:1 dilution to 1:2. Add dilution factors	C Reidenauer	R SanLuis
2	3/12/14	8.4	Change Isoton to saline	C Reidenauer	R SanLuis
3	3/30/16		Change SGAH to SGMC throughout	L Barrett	R SanLuis
3	3/30/16	8.1	Replace specific centrifugation instruction with L Barrett referral to posted instruction		R SanLuis
4	10/15/18	Header	Add other sites, update title to include method	L Barrett	R SanLuis
4	10/15/18	1	Update assay name & codes to match LIS	L Barrett	R SanLuis
4	10/15/18	3.1	Revise tube processing order to match standard protocol	Revise tube processing order to match standard L Barrett R protocol	
4	10/15/18	3.2	Revise tube numbers, add codes for comments	L Barrett	R SanLuis
4	10/15/18	4,6	Remove individual section labeling instructions and add general one	L Barrett	R SanLuis
4	10/15/18	4	Update automated stain and Diff-Quik info	D Collier	R SanLuis
4	10/15/18	6	Update product numbers & storage temp	D Collier	R SanLuis
4	10/15/18	8.3	Add second tech review for abnormal cells	L Barrett	R SanLuis
4	10/15/18	10.5	Moved review from section 6	L Barrett	R SanLuis
4	10/15/18	10.6	Added reporting section	L Barrett	R SanLuis
4	10/15/18	11.1	Updated RBC & WBC to match automated L Barrett F method		R SanLuis
4	10/15/18	12	Updated appearance to match reporting practice, D Collier Removed extraneous info		R SanLuis
4	10/15/18	15	Update to new standard wording	L Barrett	R SanLuis
4	10/15/18	16	Update SOP title	L Barrett	R SanLuis
4	10/15/18	19	I Barrett	R SanLuis	

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Adventist HealthCare

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

## Title: CSF Cell Count and Differential, Manual Method

Version	Date	Section	Reason	Reviser	Approval
5	5/16/19	2	Updated local test codes	L Barrett	R SanLuis
5	5/16/19	8.1	Specify reporting for appearance & color	H Genser	R SanLuis
5	5/16/19	8.3	Add no diff performed if count is less than 5	L Barrett	R SanLuis

## **19. ADDENDA**

A: Fluid Keyboard: Accessing Differential Result Entry for CSF

## Addenda A

## Fluid Keyboard: Accessing Differential Result Entry for CSF

1. Log into the Sunquest GUI application.



2. Click on Differential Result Entry.

S Sunquest Laboratory							= 6 x	
Sunquesta Smartan Produit List Messages Help Settings Ext Legout						egout LOGO		
Elood Bank 🗿 Microbiolo	ogy <u>M</u> aintenance					Tech Id: Area: Lab Locat	4072 LAB1 ion: WS1	CPU: A Version: 7.1 Build: 7.1.2001
My (Logged in as: CHINI,ASHKAN View By	All( <u>Z</u> )						Vi	ew By 🔡 📰 🔛
Name         Description           Image: Constant of the second secon	Administrative Billin	Elood Order	Callback	Differential	Message Filter	Micro	Micro Inquiry	Micro Result
Laboratory Inquiry Sunquest Laboratory Inquiry	Micro Tech final	is		Result Endy	ounty	Growth		endy
	Release Result É	try	Ι	Differe	ntial R	esult E	Entry	

- 3. Under "Information Dialog" screen, click on the down arrow and select the type of fluid.
  - Note: For CSF select the keyboard that is associated with the CSF tube number. *Example*: if diff is being performed on tube 1 then select C1DIFF.

Information Diak	og	
Tech Code(s)	102	
<u>K</u> eyboard	FLDIFF	•
	C1DIFF C2DIFF	-
Shift	C3DIFF	

4. Enter the accession number and press enter. When patient information is displayed, verify it matches the specimen tested. Once patient identification is confirmed, click on **Count**.

T41838 TEST,AML ONE				inul	Differential Resulting				
	oung the morp	IOIO Y	U UA KE	lew				1000 - 100 - Million Text Provide	Save
<u>A</u> cc # <mark>T41838</mark> Age 67Y	Name	TES on 03/2	T,AML ONE 9/2016 - 0	600	Co	unt	ST-1	Hospital WAH	Save/Rpt Blk
DOB 12/12/194	8 Physicia	an Trot			Spec C	omment			Hold
Sex M	Diagno	S. TEST			Order C Order C	Comment	CCD		Reject
Hemogram		∖							Close
Accession # [	Date	Time	FAPP	FC	OL	FRBC	FTNC	FTYP23	Workload
			•	Acc	essic	n Nu	imber	F	Retain Count y
Count Accession # [	Date	Time	FPOL	FLYMP	FMAM	FEOS	COMM2		
T41838	03/29/2016	0600	-	-		9 <b>—</b>	-		
Morphology									
Accession # [	Date	Time	1						
					Com	iment	]		

5. Examine the slide and adjust the "Total Count Limit" accordingly. To adjust the Total Count Limit, click on **Set Count** and then choose one of the options.



6. Enter the differential count. When finished click on QA Review.

#### Adventist HealthCare

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

T41838 T	EST, AML ON	IE			Differential Resulting		
History	Count 🖸 Morp	hology 🗹 🤇	A Review				
FPOL F	LYMP FMAM 0 0	FEOS 0	COMM2 0		QA Review	New Count Sgt Count Temp Sgve Count Manual	Save/Rpt Blk Hold Reject
Hemogran FAPP –	FCOL	FRBC -	FTNC –	FTYP23 ¥	Total 0 Count Limit: 100 FVOL	Comment	Close Workload Retain Morph Retain Count <u>v</u>
	Q TERM Z FPOL F	X C LYMP FMAM	V B FEOS COMM2				

## 7. Review the QA report, then click **SAVE**.

1838 T	TEST,AML				Differential Resulting	1		
History	Count 🔯	Morphology	QA Review					
								Save
POL	15							Save/Pot B
JAMP JAM	22							
SOS	16					SAVE		Hol <u>d</u>
DMM2	28					DIAVL		
					L		-	Reject
								Close
								workload.
								Retain Mor
								Retain Mor
								Retain Mor
								Retain Mor
		Order Prio	ority Codes and C	comments				Retain <u>M</u> or
		Order Prio	ority Codes and C	Comments	Filed			Retain Mor
		Order Prio	prity Codes and C Appended Code/Comment	Comments	Eiled Order Ode/Comme	int		Retain Mor
		Order Prio	<b>Prity Codes and C</b> Appe <u>n</u> ded Code/Comment	Comments	Eiled Order Code/Comme	int		Retain Mor
		Order Prio	ority Codes and C Appe <u>n</u> ded Code/Comment	Comments	Eiled Order Code/Comme	int		Retain <u>M</u> or
		Order Prio	Prity Codes and C Appended Code/Comment	Comments Add Remove	Eiled Order Code/Comme	int		Retain <u>M</u> or
		Order Prio	Appended Code/Comment	Comments Add Remove	Eiled Order Code/Comme	int		Retain <u>M</u> or
		Order Prio	Appended Appended Code/Comment	Comments Add Remove	Eiled Order Code/Comme	int		Retain <u>M</u> or
		Order Prio	Prity Codes and C Appended Code/Comment	Comments	Eiled Order Code/Comme	int y		Retain Mor
		Order Prio	FCCD	Comments Add Remove Comment Te <u>x</u> t	Eiled Order Code/Comme	int y		Retain Mor

- 8. If the slide requires a second tech review, click on HOLD.
  - a. A second tech will perform the differential using a manual cell counter or the off-mode in Sunquest and record results on the Pathologist Slide Review Request form.
  - b. The criteria specified on the Automated Stainer Differential Comparison and Stain Quality Log is used to compare results.
    - If the result comparison meets the criteria, then the original differential is reported.
    - If the differential results do NOT correlate, then supervisor/tech in charge will review the diff and decide which results to report.

- 9. If the slide requires a pathologist review, then check to see if there is a Cytology order.
  - a. If there is a cytology order
    - The slide does NOT need to be submitted to pathology for review.
    - Append English Text code **SCYT** (translates to See Cytology Report) to one of your cells counts. Choose a cell type that was observed because the English Text code will not post if you append it to a cell count with a result of 0.

### Example:

Click in the CLYMP3 count box, and then click on the Comment button.

History	Coun <u>t</u>	Morph	iology 🛛 🗹	QA Review		
CPOL3	CLYMP3	СММЗ	ССОМЗ 0	CEOS3		New Count
						S <u>e</u> t Count
						Temp S <u>a</u> ve
						Count Manual
					T-4-1	Comment
					100 Count Limit: 100	

Another window opens. In the Text Code box type in **SCTY** and then click **ADD**. Text Code is translated in the box to the right. Click **OK** to save.

Cor	nments	]
	Acc#         H2433         Test         CLYMP3           Date         10/11/2018         Time         0943	O <u>K</u>
	Selected Entries	
	Add SCYT See cytology report	
	Dept codes only	
	Allow Formatting	
	Correction <u>S</u> tatements (Read Only)	

- Note: The comment will append to the cell type you selected and can be seen in the QA Review tab.
- b. If there is no Cytology order -
  - The slide will need to be submitted to pathology for review. The next steps describe how to document sending it for review, sections 9.c and 9.d outline how to order Path Review and result it.
  - Append English Text code **SFPR** (translates to Submitted for Path Review) to one of your cells counts. Choose a cell type that was observed because the English Text code will not post if you append it to a cell count with a result of 0.

### Example:

Click in the CLYMP3 count box, and then click on the Comment button.

POL3	CLYMP3	CMM3	CCOM3	CEOS3		New Cou
εu	194	110	lo	110		Sgt Count
						Temp Sav
						Count Ma
						Comment
					T	Morphology

Another window opens. In the Text Code box type in **SFPR** and then click **ADD**. Text Code is translated in the box to the right. Click **OK** to save.

Comments		×
Acc# H2433 Test CLYMP3 Date 10/11/2018 Time 0943	O <u>K</u> Cancel	
T <u>e</u> xt Code Add Dept codes only Allow Formatting	Selected Entries Code <u>T</u> ranslation SFPR Submitted for path review	

Note: The comment will append to the cell type you selected and will be seen in the QA Review tab. *Example:* 

6	✓ H <u>i</u> story	Coun <u>t</u> 🔯	Morpholog <u>v</u>	☑ <u>Q</u> A Rev	view		
	CPOL3	20					
	CLYMP3	54-sfpr	Su	bmitted	for	path	review
	СММЗ	16					
	CCOM3	0					
	CEOS3	10					

- c. To submit slides for path review -
  - Add order code CPATH to the Accession via REI or GUI Order Entry.
  - Complete Pathologist Slide Review Request form.
  - Give slide(s) and review form to the pathologist.

d. When the Pathologist Slide Review form and slide(s) are returned to the lab, enter results into the LIS via SmartTerm. Note: This should also include the pathologist's comments or assessment regarding the diff count which has already been reported in SmartTerm. There original reported diff does NOT need to be corrected.

Example:						
nee no	nnne	FIL. 1231-30	HOE/ JE	A LUC	FIIIJICI	
H2433	TEST, MARIE		3M F	TEST	CACCIAE	BEVE N
		DOB:	06/26/2018	COLL:	10/11/2018	09:43
Fatas Taut						
Pocult · P						
Result . L	LUN					
Positive	for malignar	t cells. Possiblu	lumphoma/leuk	emia.	Flow	
cutometry	analysis is	recommended. Findi	ngs were disc	ussed w	ith Dr.	
No on 10/1	1/18 at 0938	. Pathologist: Dr.	Pathologist	10/11/1	8	
		Ū	Ū			
Window: M	IAIN Fi	le: NONE		М	rap: 70	Inser
SAVE and E	XIT: Are you	sure? ( <y>/N)</y>				

Example of display in Sunquest Inquiry:

COLL: 10/11/2018 09:43 REC: 10/11/2018 09:53 PHYS: CACCIABEVE MD, H2433 Req. No.:

CSF Path Review **CSF** Path Review See below (See Below)

Positive for malignant cells. Possibly lymphoma/leukemia. Flow cytometry analysis is recommended. Findings were discussed with Dr. No on 10/11/18 at 0938. Pathologist: Dr. Pathologist 10/11/18

CSF Tube 3				
Apperance tube 3	Clo	oudy	[CLEAR]	
Color tube 3	Col	lorless	[COLR]	
CSF WBC Tube 3	2		[0-5]	cell/mcL
CSF RBC Tube 3	12			cells/mcL
CSF Polys tube 3	20		%	
CSF Lymph tube 3	54		%	
Submi	tted	for path	review	
CSF Macro/Mono tube	3	16	%	
CSF EOS tube 3		10	%	