

TRAINING UPDATE

Lab Location: GEC, SGMC & WAH
Department: Core Lab

Date Distributed: 3/28/2018
Due Date: 4/18/2018
Implementation: 4/4/2018

DESCRIPTION OF PROCEDURE REVISION

Name of procedure:									
Sysmex XN Series Operation for CBC and Reticulocytes SGAH.H963 v2									
Description of change(s):									
<table border="1"><thead><tr><th>Section</th><th>Reason</th></tr></thead><tbody><tr><td>10.6</td><td>Add physician request for manual diff</td></tr><tr><td>19</td><td>Add addendum 8 (<i>steps to upload or change instrument rules</i>)</td></tr><tr><td>Add 1</td><td>Add peds values for IG %, extend adult age to include 13-18 yrs.</td></tr></tbody></table>		Section	Reason	10.6	Add physician request for manual diff	19	Add addendum 8 (<i>steps to upload or change instrument rules</i>)	Add 1	Add peds values for IG %, extend adult age to include 13-18 yrs.
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Add 1	Add peds values for IG %, extend adult age to include 13-18 yrs.								
<p>This revised SOP will be implemented on April 4, 2018</p>									

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

Technical SOP

Title	Sysmex XN Series Operation for CBC and Reticulocytes	
Prepared by	Ashkan Chini	Date: 5/17/2017
Owner	Robert SanLuis	Date: 5/17/2017

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

Review		
Print Name	Signature	Date

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1. TEST INFORMATION

Assay	Method/Instrument	Local Code
Hemogram (<i>WBC, RBC, HGB, HCT, MCV, MCH, MCHC, RDW, PLT, MPV</i>)	Sysmex XN Series (1000 or 3000)	CBCND
Hemogram & diff (<i>WBC, RBC, HGB, HCT, MCV, MCH, MCHC, RDW, PLT, MPV, differential</i>)		CBC
Differential count only		DIFF
Platelet Count		PLTC
Reticulocyte (<i>Percent, Absolute, RET-He</i>)		RCOUNT

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Abbreviation	Term	Abbreviation	Term
WBC	White Blood Cell	MCHC	Mean Corpuscular Hemoglobin Concentration
RBC	Red Blood Cell		
HGB	Hemoglobin	RDW	Red Cell distribution Width
HCT	Hematocrit	DIFF	Differential Count
MCV	Mean Cell Volume	PLT	Platelet
MCH	Mean Corpuscular Hgb	MPV	Mean Platelet Volume
RETIC	Reticulocyte Count	IPF	Immature Platelet Fraction
RET-He	Reticulocyte Hgb Equivalent	IG	Immature Granulocytes

Department
Hematology

2. ANALYTICAL PRINCIPLE

The RBC detector counts the RBC and PLT via the Hydro Dynamic Focusing. At the same time, the hematocrit is calculated via the RBC pulse height detection method.

The RBC is calculated as a particle count between lower and upper discriminators, which are automatically setup in the ranges of 25 – 75 fL and 200 – 250 fL. The particle size distribution is checked for abnormal relative frequencies at each discriminator level existence of more than one peak and abnormal distribution width.

The PLT count is calculated as a particle count between lower and upper discriminator, which are automatically set up in the ranges of 2 – 6 fL and 12 – 30 fL. PLT particular size distributions are checked for abnormalities, including abnormal relative frequencies at the lower discriminator, abnormal distribution widths, and existence of more than one peak.

Flow Cytometry is used to analyze physiological and chemical characteristics of cells and other biological particles. It is also used to analyze those cells and particles as they are passed through extremely small flow cells.

The WNR Channel uses flow cytometry to create a scatter gram and is primarily used to count the white blood cells and nucleated RBC. This scatter gram displays groups of nucleated RBC, basophil, non-basophil WBC, hemolyzed RBC and platelets.

The WDF Channel uses flow cytometry to create a scatter gram and is primarily used for classifying WBCs. This scatter gram displays groups of lymphocytes, monocytes, eosinophils, basophils and neutrophils.

The WPC Channel uses flow cytometry to create a scatter gram and is used for detecting immature WBCs such as myeloblasts and abnormal lymphocytes. This scatter gram displays groups of immature/abnormal WBCs and mature WBCs.

Sysmex XN does not use the formula $HCT = (RBC \times MCV)/10$; instead it directly measures hematocrit by adding up the cumulative number and heights of the pulses determined during the RBC counting process.

3. SPECIMEN REQUIREMENTS

3.1 Patient Preparation

Component	Special Notations
Fasting/Special Diets	N/A
Specimen Collection and/or Timing	N/A
Special Collection Procedures	N/A

3.2 Specimen Type & Handling

Criteria	
Type -Preferred -Other Acceptable	K ₃ EDTA or K ₂ EDTA Whole Blood Sodium Citrate – for platelet count only
Collection Container	Lavender Top Tube or Microtainer Blue Top Tube (Sodium Citrate)
Volume - Optimum - Minimum	Full Tube Adult: 1.0 mL, Microtainer: 0.5 mL
Transport Container and Temperature	Collection container at room temperature
Stability & Storage Requirements	Room Temperature: 48 Hours
	Refrigerated: 48 Hours
	Frozen: N/A
Timing Considerations	N/A
Unacceptable Specimens & Actions to Take	Specimens that are unlabeled, improperly labeled, or those that do not meet the stated criteria are unacceptable. Notify the attending nurse or physician and request a recollection and credit the test with the appropriate LIS English text code for “test not performed” message. Example Quantity no sufficient-QNS; Wrong collection-UNAC. Document the request for recollection in the LIS. Refer to section 13.11 for Icterus & Lipemic specimens
Compromising Physical Characteristics	Gross hemolysis: Refer to section 13.11
Other Considerations	Cancel clotted specimens, notify the attending nurse or physician and request a redraw.

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.

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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 / Kits	Supplier & Catalog Number
Cell Clean Auto	Sysmex Corporation, Cat. No. CF579595
Cell Pack DCL	Sysmex Corporation, Cat. No. DCL-300A
Cell Pack DFL	Sysmex Corporation, Cat. No. BT965910
Fluorocell PLT	Sysmex Corporation, Cat. No. CD994563
Fluorocell RET	Sysmex Corporation, Cat. No. BN337547
Fluorocell WDF	Sysmex Corporation, Cat. No. CV377552
Fluorocell WNR	Sysmex Corporation, Cat. No. CP066715
Lysercell WDF	Sysmex Corporation, Cat. No. ZA900001
Lysercell WNR	Sysmex Corporation, Cat. No. ZA900002
Sulfolyser SLS	Sysmex Corporation, Cat. No. BJ350971
Wright-Giemsa Stain	Sysmex Corporation, Cat. No. ACC-SP5741
Phosphate Buffer Solution, pH6.8	Sysmex Corporation, Cat. No. ACC-SP5548
Methyl Alcohol Absolute	Medical Chemical corporation, Cat. No. 107B
NERL Reagent Grade Water	Thermo Fisher Scientific, Cat. No. 9800-4

4.2 Reagent Preparation and Storage

Reagent	Cell Clean Auto
Storage	Store at 1 - 30°C. Avoid exposing to direct sunlight
Stability	This reagent is for single use only. Once the product stopper is punctured, the remaining reagent must be thrown away after each use.
Preparation	None

Reagents	Cell Pack DCL, Cell Pack DFL
Storage	Store at 2 - 35°C. Avoid exposing to direct sunlight
Stability	Once in use, these reagents remain stable for 60 days.
Preparation	None

Reagents	Fluorocell PLT, Fluorocell RET, Fluorocell WDF, Fluorocell WNR, Lysercell WDF
Storage	Store at 2 - 35°C. Avoid exposing to direct sunlight
Stability	Once in use, this reagent remains stable for 90 days.
Preparation	None

Reagent	Lysercell WNR
Storage	Store at 2 - 35°C. Avoid exposing to direct sunlight
Stability	Once in use, this reagent remains stable for 60 days.
Preparation	None

Reagent	Sulfolyser SLS 1.5 L
Storage	Store at 2 - 30°C. Avoid exposing to direct sunlight
Stability	Once in use, this reagent remains stable for 60 days.
Preparation	None

Reagent	Wright – Giemsa Stain
Storage	Store at 15 - 30°C
Stability	Reagent is stable unopened and opened until the expiration date on the container.
Preparation	None

Reagent	Phosphate Buffer Solution, pH 6.8
Storage	Store at 15 - 30°C
Stability	Reagent is stable unopened and opened until the expiration date on the container.
Preparation	None

Reagent	Methyl Alcohol Absolute
Storage	Store at 15 - 35°C
Stability	Reagent is stable unopened and opened until the expiration date on the container.
Preparation	None

Reagent	NERL Reagent Grade Water
Storage	Store at room temperature
Stability	Reagent will remain stable for 30 days after opening
Preparation	None

Auto Rinse needs to be run after every reagent change; to run an Auto Rinse:

- Click the **Analyzer Menu** button
- Select **Auto Rinse**
- Once complete, [Auto Rinse] disappears and the background check begins
- Background check performs analysis without aspirating the sample to verify the effects of the auto rinse
- When Background check is finished, from the **Main Menu**
- Select **Sample Explorer**

- High light the background check which was just run
- Click **Validate**
- Report prints
Note: There should be no values on the report; only zeros and dashes should appear. If any value other than zero appears on the report, that means the background check has failed and the instrument will flag the failure.
- Review and sign the printed report and place in the Sysmex Maintenance binder.

To get reagent details, such as when it was placed onboard, lot number, expiration date and volume; From **Main Menu** select **History**, then click on **Reagent Replacement Log** tab.

To scan a reagent:

- From **Main Menu** either click on the reagent picture or click on **Analyzer Manu** then select **Reagent Replacement**.
- Select the desired reagent
- Select **Replace the Reagent**
- Click on the blank space below **Reagent Code**, make sure the cursor is blinking, then scan the barcode of the reagent
- Click on **Execute**

5. CALIBRATORS/STANDARDS

5.1 Calibrators/Standards Used

Calibrator	Supplier& Catalog Number
XN CAL™	Sysmex 213527
XN CAL™ PF	Sysmex 213536

5.2 Calibrator Preparation and Storage

Calibrator	XN CAL™, XN CAL™ PF
Storage	Store at 2-8°C in a dark refrigerator.
Stability	Unopened: Manufacturer’s Expiration Date Opened: 4 hours
Preparation	Calibrator is supplied ready to use. Bring to room temperature prior to testing. Mix per manufacturer’s guidelines.

5.3 Calibration Criteria and Procedure

Criteria	Special Notations
Frequency	<ul style="list-style-type: none"> • Assay calibration must be performed when the instrument is first placed in service and at least every 6 months thereafter. This process is managed by Sysmex Customer Support Center. <ol style="list-style-type: none"> 1) First half of the year: Field Service Engineer (FSE) will perform the calibration. 2) Second half of the year: Customer Support Center will ship the calibrator ahead of the time. They will contact the Insight account holders prior to the calibration due date to inform them of the date and time for the calibration. • In addition, calibration verification is required (regardless of the length of time since last performed) immediately if any of the following occurs: <ol style="list-style-type: none"> 1) When control data indicates a significant shift in assay results and cannot be corrected by maintenance or troubleshooting. 2) After major maintenance, service or replacement of critical parts. 3) When advised by Sysmex Field Service Representative.
Tolerance Limits	<p>Sysmex will provide a calibration report:</p> <ol style="list-style-type: none"> 1) Review Reagents Information and Calibration Materials to verify all products used for calibration are in-date. 2) On the Background Counts chart, the Count for each parameter must be less than the Limit. 3) On the Precision Open Mode chart, the CV% of each parameter must not exceed the % Limit. 4) On the Sensitivity Verification, each parameter's Mean must fall between Lower and Upper Limit. 5) On the Calibration of Directly Measured Parameters – <ol style="list-style-type: none"> a. The first run is always eliminated. b. The Mean of each parameter must fall between Lower and Upper Limit. c. If the current coefficient of variation is not the same as the new one, 10 points of QC must be run and reviewed and the QC ranges adjusted if necessary.
Procedure	<ul style="list-style-type: none"> • Calibration is performed by Sysmex FSE or • Calibration is performed by Sysmex through remote access. Customer must contact Sysmex to order calibration material, for any calibrations that are done outside of major maintenance by FSE or the Sysmex calibration schedule. • Refer to Chapter 12 of the Instructions for Use manual – Performing Calibration – The Customer will be guided by the Sysmex representative to run the calibrators and the values will be updated by the Sysmex representative. • If calibration factors (compensations rates) have been changed,

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Criteria	Special Notations
	calibration verification must be performed by running all levels of commercial control. <ul style="list-style-type: none"> At the completion of the calibration, a certificate of calibration will be printed by the Sysmex FSE that must be reviewed by a Group Lead/ Supervisor or designee.

6. QUALITY CONTROL

6.1 Controls Used

Controls	Supplier and Catalog Number
XN CHECK, Levels 1, 2 & 3	Sysmex Corporation, Cat. No. 213499

6.2 Control Preparation and Storage

Control	XN CHECK
Preparation	None
Storage	Store at 2 - 8°C
Stability	Unopened: manufacturer’s expiration date Opened: 7 days when stored at 2 - 8° C after each use.

6.3 Frequency

All three levels of control must be run on all Sysmex XN instruments every 4 hours of patient testing. QC must also be performed after shutdown, maintenance or instrument repairs.

Refer to addenda 4 for instructions to perform parallel testing for new lots of QC materials.

6.4 Tolerance Limits and Criteria for Acceptable QC

A. Tolerance Limits

The Hematology QC program is monitored in the instrument and should be set up using the Evidence-based QC Limit % Range specific for XN analyzers. These limits are provided by Sysmex and are intended to ensure reasonable error detection capability and minimal false rejection rates. Target values for each level of control will be calculated based on the data collected in the new lot evaluation.

B. Criteria for Acceptable QC

- All Controls must be within the acceptable range.

- Controls and patient data must be reviewed for acceptability and for atypical or unexpected results or trends prior to reporting patient results.
- DO NOT release results from runs with unacceptable controls or with unusual patterns, trends or distribution in patient values.

C. Corrective Action

- All rejected runs must be effectively addressed and include the following documentation:
 - Control(s) that failed and/or atypical or unexpected patient results
 - Actions taken
 - Statement of what was done with the patient samples from the affected run/batch,
 - Date and initials of the person recording the information.
- Patient samples in failed analytical runs must be reanalyzed.
- **Precision Statistics:** When there is a significant shift/bias on QC data, the root cause of the increased imprecision must be investigated and a resolution needs to be considered immediately. All of these actions must be documented including an evaluation of whether or not this affected patient care.

NOTE: The laboratory director or designee may override rejection of partial or complete runs. Justification for the override must be documented in detail.

6.5 Documentation

- QC tolerance limits are programmed on the instrument; 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 Lead Technologist 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.

6.6 Quality Assurance Program

- 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.
- QC is submitted to Sysmex for peer group comparison as it is run. Refer to addenda 4 and 5
- Consult the Laboratory QC Program for complete details.

7. EQUIPMENT and SUPPLIES

7.1 Assay Platform

Sysmex XN Series 1000/3000
Sysmex Automated Slide Preparation Unit SP – 10 (SG/WAH only)

7.2 Equipment

Microscope
Refrigerator

7.3 Supplies

Immersion Oil
Applicator sticks
Glass Slides
Lens Paper
Sysmex Glass Slides for Stainer (SG/WAH only)
Ribbon for Stainer (SG/WAH only)
Glass Plate Spreader (SG/WAH only)

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.

8.1	Maintenance
1	Perform required instrument maintenance. Refer to addendum 2 for details.
2	Perform required stainer maintenance (SG/WAH). Refer to addendum 3 for details.

8.2	Test Run
1	Place the racks in the right side feed conveyor. Pre- mixing of samples is not required.
2	Racks will automatically feed to the analyzer. Tubes are rotated to read barcode.
3	On-board IPU rules will determine repeat or reflex testing. Rack will run in reverse to perform the repeat or reflex test.
4	When load is completed, remove racks from left side feed conveyor.

8.3	Special Handling
1	To load a STAT specimen while the instrument is analyzing other samples: Press the mode switch button, the tube holder (for manual/pediatric samples) slides out forward; then run the STAT specimen in manual mode.

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8.3	Special Handling
2	Aspiration sensor must remain ON all the time. The only times that Aspiration Sensor is turn off are: <ul style="list-style-type: none"> • When hemoglobin is < 4.0 • When Platelet Poor Plasma study is in progress

NOTE: In the event that the test system becomes inoperable, notify supervision or designee for further direction. Patient specimens must be stored in a manner that maintains the integrity of the specimen.

9. CALCULATIONS

The following calculations are performed automatically by the instrument. The formulas used to calculate MCV, MCH and MCHC are:

- MCV: (HCT/RBC) x 10
- MCH: (HGB/RBC) x 10
- MCHC: (HGB/HCT) x 100

The following calculations are performed when screening a slide and/or performing a differential (refer to Sections 10.6 and 13 for details)

WBC Estimate:

Calculate the average WBC in 10 fields using the 50X objective and multiply by 3,000. If WBC estimate does not agree with what the instrument has reported within ± 20%, repeat the estimate focusing on the feathered edge of the smear. If counts still do not agree, consult with the supervisor or tech in charge.

Platelet Estimate:

Calculate the average PLT in 10 fields using the 100X objective and multiply by 20,000. If PLT estimate does not agree with what the instrument has reported within ± 20%, repeat the estimate focusing on the feathered edge of the smear. If counts still do not agree, consult with the supervisor or tech in charge.

Megakaryocytes could potentially interfere with the WBC count.

If the WBC estimate does not correlate with the automated WBC count in the presence of five or more megakaryocytes, then correct the WBC count use the formula below. To get the uncorrected WBC, **Main Menu – Data Browser – select Service tab – select WNR.**
TNC – N has the uncorrected WBC count.

$$\text{Corrected WBC} = (\text{Uncorrected WBC} \times 100) / (100 + \# \text{ of megakaryocytes})$$

10. REPORTING RESULTS AND REPEAT CRITERIA

10.1 Interpretation of Data

The relationship of the RBC, HGB and HCT to MCV, MCH and MCHC			
Parameter Affected	MCV	MCH	MCHC
RBC ↓	↑	↑	No change
RBC ↑	↓	↓	No change
HGB ↓	No change	↓	↓
HGB ↑	No change	↑	↑
HCT ↓	↓	No change	↑
HCT ↑	↑	No change	↓

10.2 Rounding

Any result rounding is performed at the interface level.

10.3 Units of Measure

Parameter	Units	Parameter	Units	Parameter	Units
WBC	x10(3)/mcL	RBC	10 ⁶ /μL	HGB	g/dL
HCT	%	MCV	fL	MCH	pg
MCHC	g/dL	PLT	x10(3)/mcL	MPV	fL
RDW	%	Diff count	%	Reticulocyte	%
RET-He	pg	IPF	%	IG	%

10.4 Analytical Measurement Range (AMR)

Parameter	Sysmex XN Series
WBC	0 - 440 x 10 ³
RBC	0.00 - 8.60 x 10 ⁶
HGB	0 - 25 g/dL
HCT	0 - 75%
PLT, PLT-F	0 - 5,000 x 10 ³
RET%	0 - 30.0%

10.5 Review Patient Data

- Review patient results for unusual patterns, trends or distribution.
- Report atypical or unexpected results or trends for this test to appropriate supervisory personnel, prior to releasing results.

10.6 Repeat Criteria and Resulting

Dilution:

Sysmex XN is not capable of doing any dilutions. Refer to WBC and RBC sections for dilution details, and then follow these steps to program them on the instrument:

- Manually program the accession number by typing the corresponding alpha character equivalent of the day of the week then followed by the number. **Verify the characters are set in a way that the DI will not auto verify the result.**
- Run the sample in manual mode.
- When the sample is resulted, multiply the value by the dilution factor used to make the dilution.

WBC		
Condition	Action 1	Action 2
< 0.5	Check for a clot, re-analyze sample in LW (Low WBC) mode.	SCAN
≤ 2.0	<ul style="list-style-type: none"> • Check Sample for a clot • If unable to evaluate 100 cells, do a 50 cell Diff and multiply results by 2. • Re-analyze and verify the count is within ± 15%. Add comment code RVT (reviewed by Technologist) 	DIFF
≥ 30.0	Scan to verify count and rule out the increase due to presence of Giant platelets or abnormal protein/cryoglobulin	SCAN
≥ 440.0	Re-analyze using dilution factor 2 (use DCL Cell Pack as diluent); if the diluted result is still > 440.0, then report as > 880.0, refer to ≥30.0, add comment code REP (results confirmed, test repeated)	SCAN

RBC		
Condition	Action 1	Action 2
≥ 8.60	Re-analyze using dilution factor 2 (use DCL Cell Pack as diluent); if the diluted result is still > 8.60 then report as > 17.2, scan to verify morphology and add comment code REP	SCAN

HGB		
Condition	Action 1	Action 2
< 4.0	Refer to section 8.3	N/A
≤ 6.0	Check for a clot	N/A
≥ 20.0 Excludes Neonates	Check coagulation sample if HCT ≥ 55, add REP comment.	N/A
≥ 25.0	Re-analyze using dilution, check coagulation sample if HCT ≥ 55, add comment	N/A

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MCV		
Condition	Action 1	Action 2
≤ 50.0	Check for a clot, re-analyze and verify morphology, add RVT comment.	SCAN
≥ 130.0	Warm specimen to 37°C for 30 minutes then re-analyze, verify morphology, denote any rouleaux or RBC agglutinins	SCAN

Platelet		
Condition	Action 1	Action 2
< 50	Check for a clot, re-analyze using optical mode if not already performed.	N/A
> 5000	Re-analyze using dilution, perform platelet estimate, add REP comment.	SCAN

Lymphocyte		
Condition	Action 1	Action 2
≥ 70	If differential agrees with auto count, Pathologist review is required	DIFF
Note: For patients <12 years old perform DIFF if the Lymphocyte count is greater than Neutrophil		

Neutrophil		
Condition	Action 1	Action 2
≥ 90 No flags	Scan for any bands, hyper segmentation or toxic granulation	SCAN
≥ 90 With flags	Rule out immature Neutrophils	DIFF

Monocyte		
Condition	Action 1	Action 2
> 25	Rule out immature Monocytes	DIFF

Eosinophil		
Condition	Action 1	Action 2
≥ 35	Scan to verify Eosinophils. Rule out presence of parasites.	SCAN

Basophil		
Condition	Action 1	Action 2
≥ 3.5	Mix for 5 minutes then re-analyze, if still elevated scan to verify cells	SCAN

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Reticulocyte		
Condition	Action 1	Action 2
> 30.0%	<ul style="list-style-type: none"> • Check Sample for a clot, Vortex the sample for 1 – 2 minutes and reanalyze • If same result is obtained with no flag - check patient's history, check with the attending physician/nurse to confirm the situation and result, inform the supervisor/tech in charge and with their permission report as > 30.0 %. • If same result is obtained with a flag, refer to section 13.9 	Repeat

Differential Timing
<p>If a patient had a differential done in the past 48 hours while remaining admitted, there is no need to do another differential if CBC parameters are improving (moving toward normal). Exceptions are:</p> <ul style="list-style-type: none"> • If blasts flag • Physician request • If Asterisk appears next to results

H and H Mismatch
<p>On samples where hemoglobin and hematocrit results differ significantly (for example HGB value of 12.5 and HCT value of 1.6), do the following:</p> <ul style="list-style-type: none"> • Check for a clot • Incubate the sample for 30 minutes at 37 °C and reanalyze. If the H and H mismatch remains proceed to next step • Spin the sample and check for hemolysis, lipemia or icteric. If either one is detected, refer to section 13.11 • Contact attending physician / nurse and ask for patient history, medication and a redraw if the current result seems questionable.

Supervisor (or designee) / Pathologist slide review		
Abnormality	Supervisor	Pathologist
Prolymphs > 5%	X	
Reactive and/or atypical lymphocytes >20%	X	
Bands > 25%	X	
Meta/Myelos/Promyelo >10%	X	
Any blast cell	X	X
Any unidentifiable cell	X	X
Any parasite or microorganism (reviewed by microbiology also)	X	
Lymphocyte > 75% in patients < 17 years of age	X	
Lymphocyte > 70% in patients > 17 years of age	X	X

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NOTE: The above guidelines are for new and recurring patients performed initially and over each subsequent hospital encounter (ED visit, OP visit or admission).

Handling and Resulting Pathologist Reviewed Slides

- A. Technician/Technologist will submit slides for pathologist review as follows:
 1. Ensure slide is of acceptable quality for pathology review; appropriate smear, adequate staining, and properly labeled.
 2. Cover-slip the slide
 3. Complete Pathologist Slide Review Request form
 4. Attach analyzer print-out

- B. Technician/Technologist will enter Pathologist comments in DI. Refer to Data Innovations Instrument Manager procedure for detailed instructions.

Physician Request for Manual Differential

If a physician requests a manual differential, then a new order (accession number) must be created. Utilize the same collect date and time as the sample and order test code DIFF.

11. EXPECTED VALUES

11.1 Reference Ranges

See Addendum 1.

11.2 Critical Values

Parameter	Age	Critical Low	Critical High
HGB	1 month and older	< 6.1	> 19.9
HGB	0-29 days	< 6.1	> 23.9
WBC	all ages	< 2.1	> 29.9
Platelet	all ages	< 31	> 899

11.3 Standard Required Messages

SGMC only: If patient is in NICU (3C), then call platelet count $\leq 75 \times 10^3$ /mCL

12. CLINICAL SIGNIFICANCE

- **CBC** – The quantitative and qualitative analysis of the cellular elements of blood will identify imbalance between cell production, cell release, cell survival, or cell loss. This information increases the accuracy and specificity of diagnosis based on pathogenesis and is also used to monitor the effectiveness of therapy.
- **Automated Differential** – The Differential distribution of white blood cells will, when correlated with absolute white cell count, identify imbalances between cell production,

cell release, cell survival and/or cell loss. This information increases the accuracy and specificity of diagnosis based on pathogenesis and is also used to monitor the effectiveness of therapy.

- **Platelet Count** – Platelets must be present in adequate numbers and have proper function to aid in hemostasis. A normal bleeding time is dependent on adequate platelet number and function.
- **Reticulocyte Count** - The enumeration of reticulocytes provides an effective means of determining red cell production and regeneration. Elevation is seen in patients with hemolytic anemia, hemorrhage (acute and chronic), treatment of iron-deficiency anemia and megaloblastic anemias and uremia. Decreased counts may be seen in aplastic anemia, aplastic crisis of hemolytic anemias and ineffective erythropoiesis as seen in thalassemia, pernicious anemia and sideroblastic anemia.
- **IPF – Immature Platelet Fraction:** Platelets are produced in the bone marrow and are normally not released into the bloodstream until they have matured. When platelet numbers in the blood are low (thrombocytopenia), it stimulates the bone marrow to produce platelets faster. When the need is great and when production cannot keep up with demand, then an increased number of immature platelets will be released into the blood stream. The IPF may be used to help a healthcare provider determine the likely cause of a person's thrombocytopenia, that is, decrease in production by the bone marrow (IPF is low) versus increased loss of platelets in the blood (IPF is higher).
- **RET-He – Reticulocyte Hemoglobin Equivalent:** It is one way to measure the hemoglobin inside of reticulocyte. Reticulocytes are "young" red blood cells that are released by the bone marrow before they become fully mature. The amount of hemoglobin inside of reticulocytes can help determine if there has been enough iron available, to be incorporated into hemoglobin production and then into red blood cell production in the bone marrow. This makes the test useful in identifying functional iron deficiency.
- **IRF – Immature Reticulocyte Fraction:** It is the ratio of immature reticulocytes to the total number of reticulocytes. This parameter provides a very early and sensitive index of marrow erythropoietic (RBC production) activity.
- **Immature Granulocytes (IG):** This instrument has a 6-part differential that is comprised of Neutrophil, Lymphocyte, Monocyte, Eosinophil, Basophil and Immature Granulocyte. The Immature Granulocyte results include metamyelocyte, myelocyte and promyelocyte.
- **Platelet Fluorescent:** The platelet measurement is done using a nucleic acid stain specific for platelet organelles and flow cytometry. The PLT-F result will have “&F” to the left of the result indicating the result was obtained in the PLT-F channel.

13. PROCEDURE NOTES

- **FDA Status:** FDA Exempt/Cleared or Approved with modification(s).
- **Validated Test Modifications:** Sample room temperature stability extended. Quest Corporate Validation on file, see the Technical document index.

This section explains the Interpretive Program (IP) messages generated by the Sysmex XN analyzer and the corrective actions.

13.1	WBC Abn Scattergram
<p>Cause: Clustering in the WNR or WDF scattergrams is abnormal; meaning analyzer cannot separate the cell population with confidence.</p>	
<p>Corrective Action:</p> <ol style="list-style-type: none"> 1. If dashes appear in place of data: <ol style="list-style-type: none"> a. Repeat the sample b. If dashes still remain, perform a manual differential count 2. If Asterisk appears next to data: <ol style="list-style-type: none"> a. Scan the slide for abnormal cells and NRBC b. Perform a manual differential if abnormal cells are observed c. If no abnormal cells are found, then report the result 	
13.2	NRBC Present
<p>The analyzer identifies and counts NRBCs simultaneously while counting WBCs (both are counted separately). No further correction of the WBC count is required. If NRBCs are greater than 0.01/100 WBC, the lymphocyte count is automatically corrected.</p>	
13.3	IG Present (>5% flag)
<p>Cause: Presence of cells (metamyelocyte, myelocyte and promyelocyte) accurately quantitated by the analyzer</p>	
<p>Corrective Action:</p> <p>For IG >5%, this flag appears in red. Perform a manual differential count and scan the peripheral smear for the presence of the following:</p> <ul style="list-style-type: none"> • Promyelocytes, myelocytes and metamyelocytes • Band cells in increased numbers • Toxic granulation or vacuolation of neutrophils • Other abnormal cells 	
13.4	Blast/Abn Lymph
<p>Cause: The analyzer has detected abnormal clustering in the region for blasts and abnormal lymphocytes in the WDF (differential) scattergram.</p>	
<p>Corrective Action:</p> <ol style="list-style-type: none"> 1. If dashes appear in place of data: <ol style="list-style-type: none"> a. Repeat the sample b. If dashes still remain, perform a manual differential count 2. If Asterisk appears next to data, perform a manual differential count and scan the peripheral smear for the presence of the following: <ol style="list-style-type: none"> a. Blasts – lymphoblasts, myeloblasts, and myelomonoblasts b. Immature Granulocytes – promyelocytes, myelocytes, and metamyelocytes c. Atypical or immature lymphocytes d. Other abnormal cells 	
13.5	Left Shift
<p>Cause: The analyzer has detected abnormal clustering in the region for the left shift (bands) in the WDF (differential) scattergram.</p>	

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13.5	Left Shift
Corrective Action:	
<ol style="list-style-type: none"> 1. If dashes appear in place of data: <ol style="list-style-type: none"> a. Repeat the sample b. If dashes still remain, perform a manual differential count 2. Perform a manual differential count and scan the peripheral smear for the presence of the following: <ol style="list-style-type: none"> a. Band cells in increased numbers b. Toxic granulation, hyper-segmentation or vacuolation of neutrophils c. Other abnormal cells 	

13.6	Atypical Lymph
Cause: The analyzer has detected significant clustering in the region for atypical lymphocytes that is located in the upper left lymphocyte region on the WDF (differential) scattergram. An asterisk appears next to the Neutrophil, Lymphocyte, Monocyte, Eosinophil and Immature Granulocyte % and #.	
Corrective Action:	
Perform a manual differential count and scan the peripheral smear for the presence of the following:	
<ul style="list-style-type: none"> • Atypical or variant lymphocytes • Abnormal or atypical monocytes • Immature lymphocytes • Immature monocytes • Smudge cells • Other abnormal cells 	

13.7	RBC Abn. Distribution
Cause: When the histogram pattern from the RBC channel is abnormal or when RBC is $< 0.50 \times 10^6/\mu\text{L}$. Dashes may appear in place of results for the RDW-CV. This message may cause certain RBC parameters to be marked with an asterisk.	
Corrective Action:	
Scan the peripheral smear for the presence of abnormal RBC morphology such as those listed. (Only note if findings are $\geq 2+$):	
<ul style="list-style-type: none"> • Increased anisocytosis • Multiple RBC population • Fragmented RBC • Poikilocytosis • Rouleaux or RBC agglutination 	
If the RBC morphology is normal and MCHC is abnormal (<30 or >37.5 g/dL) refer to guidelines for the HGB/Turbidity Interference? IP Message, section 13.11	
If RDW remains uncalculated, result as UNC (unable to calculate)	

Note: For consistent morphological reporting, the following criteria are recommended. Review 10 fields using the 100X objective to calculate this mean range.

Variation	Normal	1+	2+	3+
Poikilocytosis	0	1 – 5	6 – 15	> 15
Microcytosis or Macrocytosis	0	1 – 5	6 – 15	> 15

Variation	Normal	1+	2+	3+
Anisocytosis	0 – 5	6 – 15	15 – 30	> 30
Polychromasia	0 – 2	3 – 4	5 – 6	> 6
Hypochromia	0 – 5	6 – 15	16 – 30	> 30

Cell Type	Normal	1+	2+	3+
Spherocyte, Acanthocyte, Sickle cell, Rouleaux	0	1 – 5	6 – 15	> 15
Helmet cell	0 – 1	1 – 5	6 – 15	> 15
Tear drop, Target cell, Schistocyte, Ovalocyte, Elliptocyte, Burr cell, Stomatocyte, Blister cell	0 – 2	3 – 4	5 – 6	> 6

13.8	Dimorphic Population
Cause: When there are multiple peaks in the RBC histogram pattern. Refer to section 13.7 for corrective action.	

13.9	RET Abn Scattergram
Cause: The analyzer cannot separate the cell population with confidence. There is a possibility of cell overlap. Asterisks appear next to the RET%, RET#, IRF and RET-He parameters.	
Corrective Action:	
<ol style="list-style-type: none"> 1. Vortex 1-2 minutes and repeat. 2. If the flag is not eliminated, or the RBC count is $<0.50 \times 10^6/\mu\text{L}$ review the peripheral smear for the presence of polychromasia, parasites, NRBCs, Howell-Jolly Bodies, Heinz bodies or basophilic stippling. If any of those abnormal cells are present, report the results with comment “results may be affected by the presence of interfering substances”. 	

13.10	RBC Agglutination
Corrective Action:	
<ol style="list-style-type: none"> 1. Warm the sample at 37° C for 15 - 30 minutes. Reanalyze the warmed sample in the manual mode after mixing by manual inversion 10 times. Make a new peripheral smear from the warmed sample if agglutination is severe and WBCs and PLTs cannot be accurately measured. See section 9 for WBC and PLT estimate calculations. 2. Sometimes agglutination can be so severe that warming the sample does not enable accurate analysis. In this case perform a plasma replacement: <ol style="list-style-type: none"> a. Centrifuge an aliquot of blood from the primary tube to separate the cells from the plasma. b. Using a pipette, remove a measured amount of plasma removing as much plasma as possible without disturbing the buffy coat. c. Add back the same amount of CELLPACK DCL as the volume of plasma removed in step “b”. d. Cap the tube and mix the sample by manual inversion until the cells are fully re-suspended in the CELLPACK DCL. e. Reanalyze the sample in the manual mode. 	

13.11	Turbidity/HGB Interference
<p>If the MCHC is ≤ 30.0 or ≥ 37.5, repeat to rule out random error.</p> <ul style="list-style-type: none"> • If MCHC is ≤ 30.0 a slide should be made and scanned to look for potential causes of spuriously low MCHC, example: marked sickle cells or target cells. 	

13.11	Turbidity/HGB Interference
<ul style="list-style-type: none"> • If the MCHC is greater than 37.5, a slide should be made and examined as well as visual inspection of the sample to determine the integrity of the specimen. The smear review / visual inspection should indicate to the technologist which category the specimen falls into – cold agglutinin, lipemia, hemolysis, icterus or the situation where the results are accurate due to the presence of spherocytes. 	
<p>Corrective Action:</p> <ol style="list-style-type: none"> 1. If Spherocytes are noted on the slide: <ol style="list-style-type: none"> a. Report the MCHC with a comment reflecting the presence of spherocytes as 1+, 2+ or 3+ 2. If significant RBC agglutination is noted on the slide, warm specimen in a 37°C heat block for 30 minutes and rerun. If not resolved, continue warming and rerun every 15 minutes continuing incubation after each run, not to exceed one hour. If necessary, make a warmed slide for morphology evaluation. <ol style="list-style-type: none"> a. If MCHC is within normal range, Report results with the LIS code SWCG which translates to “Specimen was prewarmed to 37°C to obtain results. Cold agglutinin/cryoglobulin suspected.” b. If MCHC is still greater than 37.5 after one hour of incubation, perform plasma replacement. Refer to section 13.10 3. If hemolysis is suspected, examine the specimen for visual hemolysis. If gross hemolysis is observed, cancel the specimen with the appropriate comment: -HMT 4. If lipemia or icterus is suspected, perform plasma replacement. Refer to section 13.10 	
13.12	Fragments
<p>Cause: Due to size comparison of certain RBC or Platelet population, the analyzer cannot separate the cell population with confident. There is a possibility of cell overlap.</p>	
<p>Corrective Action: Scan the peripheral smear for the presence of fragmented RBCs and other poikilocytosis</p>	
13.13	PLT Abn Distribution
<p>Cause: Indicates that the analyzer has detected abnormal size and population of platelets. Dashes may appear in place of data for the MPV or the MPV may be marked with an asterisk.</p>	
<p>Corrective Action:</p> <p>GEC: Vortex the specimen for 1 – 2 minutes and repeat. If the same flag appears, then scan the peripheral smear to estimate the platelet count and review for the presence of abnormal RBC or PLT morphology such as:</p> <ol style="list-style-type: none"> a. Giant platelets b. Platelet clumps c. Fragmented RBCs d. Microcytic RBCs e. Parasites <p>If platelet estimate confirms accuracy of analyzer count, report the result. If MPV remains uncalculated, result as UNC (unable to calculate)</p> <p>SGMC & WAH: Vortex the specimen for 1 – 2 minutes and reanalyze in PLT-F mode. If the same flag appears, then scan the peripheral smear to estimate the platelet count and review for the presence of abnormal RBC or PLT morphology such as:</p>	

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13.13	PLT Abn Distribution
<ul style="list-style-type: none"> a. Giant platelets b. Platelet clumps c. Fragmented RBCs d. Microcytic RBCs e. Parasites <p>If platelet estimate confirms accuracy of analyzer count, report the result. If MPV remains uncalculated, result as UNC (unable to calculate)</p>	

13.14	PLT Abn Scattergram (SGMC & WAH Only)
<p>Cause: When clustering in the platelet and IPF area on the PLT – F Scattergram is abnormal. The PLT – F, IPF %, and sometimes MPV are reported with an asterisk. Dashes may appear in place of data for the MPV.</p> <p>Corrective Action: Follow the steps in section 13.13 under SGMC and WAH.</p>	

13.15	PLT Clumps
<p>Cause: Abnormal clustering of platelets in the WNR, WDF or PLT – F scattergrams.</p> <p>Corrective Action:</p> <ol style="list-style-type: none"> 1. Check the sample for the presence of clots: <ul style="list-style-type: none"> • If a clot is detected, cancel the current specimen and request a redraw. • If no clot is detected, <ul style="list-style-type: none"> SGMC & WAH: vortex the specimen for 1 – 2 minutes and repeat in PLT – F mode. GEC: vortex the specimen for 1 – 2 minutes and repeat. 2. If the flag appears again, then scan the peripheral smear for the presence of abnormal morphology including: <ul style="list-style-type: none"> • Fibrin strands • Platelet clumps <p>Note: If either fibrin or clumps are present, verify the WBC and PLT by a manual slide estimate (see section 9). If the WBC and PLT estimates match the analyzer counts, report the results. If the estimates do not match the analyzer counts, proceed to next step.</p> 3. Recollect a new sample in sodium citrate. The platelet and/or WBC count obtained from the sodium citrate tube must be multiplied by 1.1 to account for the different blood to anticoagulant ratio in the citrate tube. The MPV from the citrate tube is also reported, but no correction factor is applied because the MPV is not affected by dilution. 4. If the platelet clumps still remain and the platelet count is ≤ 130, remove the platelet count number and result with these two comments: <ul style="list-style-type: none"> • Unable to report due to significant platelet clumping • Platelet estimate comment (decreased, increased or normal) 	

13.16	Giant Platelets – GEC only
<p>Corrective Action: Refer to section 13.15 steps 1 and 2 If PLT count is ≤ 130 with significant giant platelets found during morphology review, then release the</p>	

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13.16	Giant Platelets – GEC only
<p>result and append the LIS code GPINF which translates to “The WBC count and platelet count may be altered due to interferences caused by the presence of significant numbers of large giant platelets”.</p> <p>Note: Fluorescent platelet functionality will resolve this at SGMC and WAH.</p>	

14. LIMITATIONS OF METHOD

14.1 Precision

Precision is assessed by analysis of peripheral blood. The data appears consistent and all parameters have a low CV%.

14.2 Interfering Substances

Parameter	Causes of Spurious Increase	Causes of Spurious Decrease
WBC	PLT clumps, Cryoprotein, Cryoglobulin, Fibrin, Giant platelets (Platelets > 1,000,000/ μ L)	Leukocyte aggregation
RBC	Leukocytosis (>100,000/ μ L) Giant platelets (Platelets > 1,000,000/ μ L)	Erythrocyte aggregation Microerythrocytes Fragmented RBCs
HGB	Leukocytosis (>100,000/ μ L) Lipemia, Abnormal protein	
HCT	Leukocytosis (>100,000/ μ L) Severe diabetes, Uremia, Spherocytosis	Erythrocyte aggregation, Microerythrocytes, Fragmented RBCs
MCV	Spuriously increased HCT Spuriously decreased RBC	Spuriously decreased HCT Spuriously increased RBC
MCH	Spuriously increased HGB Spuriously decreased RBC	Spuriously decreased HGB Spuriously increased RBC
MCHC	Spuriously increased HGB Spuriously decreased HCT	Spuriously decreased HGB Spuriously increased HCT
Platelets	Cryoglobulin, Cryoprotein, Fragmented WBCs, Fragmented RBCs, Microerythrocytes	Giant PLT, PLT Clumping PLT Satellitosis
Retic	Erythrocyte aggregation, giant platelets, clumped platelets, fragmented leukocytes, malaria, Howell –Jolly bodies	None specified

14.3 Clinical Sensitivity/Specificity/Predictive Values

Patient results may vary with sample condition and setting of instrument to review criteria established by the laboratory. Results may also vary due to disease abnormalities, overall patient population and institutional review criteria.

15. SAFETY

Cell Clean Auto reagent causes severe skin burns and eye damage.

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

- Safety Data Sheets
- Sysmex XN Reference Manual
- Critical Values (Lab policy)
- Quality Control Program policy
- Quest Diagnostics Records Management Program
- Laboratory Safety Manual
- Data Innovations Instrument Manager, Laboratory Policy
- Current Allowable Total Error Specifications at http://questnet1.qdx.com/Business_Groups/Medical/qc/docs/qc_bpt_tea.xls
- Pathologist Slide Review Request (AG.F127)
- Sysmex XN Maintenance Log (AG.F377)
- Sysmex Stainer SP-10 Maintenance Log (AG.F378)
- Automated Stainer Differential Comparison and Stain Quality Log (AG.F379)

17. REFERENCES

1. Quest Diagnostics Best Practice Sysmex XN Series Operation for CBC SOP, revised 04/03/2017
2. Quest Diagnostics Best Practice Sysmex XN Series Operation for Reticulocytes SOP, revised 01/09/2017
3. Sysmex Hematology Analyzer XN Series Instruction for use, revised 07/2015
4. Sysmex Stainer SP – 10 Series Instruction for use, revised 02/2013
5. Sysmex XN – 3000 Automated Hematology System Quick Guide, revised 01/2013
6. Sysmex XN Check Quality Control Package Insert, revised 12/2014
7. Roehrl, M.H., et al. Age-Dependent Reference Ranges for Automated Assessment of Immature Granulocytes and Clinical Significance in an Outpatient Setting. Arch Pathol Lab Med. 2011, 135(4):471-7.
8. Kickle, T.S., et al. A Clinical Evaluation of High Florescent Platelet Fraction Percentage in Thrombocytopenia. Am J Clin Pathol 2006;125:282-287
9. Reticulocyte hemoglobin content. Alan Mast, Morey Blinder, Dennis Dietzen. Blood: Vol 83, Issue 4, 307-310

18. REVISION HISTORY

Version	Date	Section	Reason	Reviser	Approval
0	8/25/17	8.2	Add stainer maintenance	L Barrett	R SanLuis
0	8/25/17	10.3	Add missing parameters	L Barrett	R SanLuis
0	8/25/17	10.4	Correct upper limit for Hgb	A Chini	R SanLuis
0	8/25/17	10.6	Update repeat criteria	R SanLuis	R SanLuis
0	8/25/17	13.7	Add micro & macrocytosis criteria	R SanLuis	R SanLuis
0	8/25/17	13.9.13.11	Delete dilution	R SanLuis	R SanLuis
0	8/25/17	19	Add addenda 6 and 7	L Barrett	R SanLuis
0	8/25/17	Add 1	Standardize decimal places	L Barrett	R SanLuis
0	8/25/17	Add 3	Add QC requirements	L Barrett	R SanLuis
0	8/25/17	Add 4	Add QC outlier documentation steps	Z Morrow	R SanLuis
0	8/25/17	Add 5	Add detail to data management	Z Morrow	R SanLuis
0	8/25/17	Footer	Corrected pages 28-36	L Barrett	R SanLuis
1	3/13/18	10.6	Add physician request for manual diff	L Barrett	R SanLuis
1	3/13/18	19	Add addendum 8	A Chini	R SanLuis
1	3/13/18	Add 1	Add peds values for IG %, extend adult age to include 13-18 yrs.	L Barrett	R SanLuis

19. ADDENDA

Addendum	Title
1	Reference Ranges
2	Sysmex XN Maintenance and Quality Control
3	Sysmex Stainer SP-10 Maintenance
4	QC Instructions on the Sysmex XN
5	QC Instructions on Insight
6	Smear Review and Manual Differential
7	CBC DIFF/SCAN Action and Repeat Criteria
8	Sysmex Rules Management

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Addendum 1:

Adult Reference Ranges

Parameter / Units of Measure	Female		Male	
	13 – 18 years	> 18 years	13 – 18 years	> 18 years
WBC / x10(3)/mcL	4.5 – 13.0	4.0 – 10.0	4.5 – 13.0	4.2 – 9.1
RBC / 10 ⁶ /µL	4.10 – 5.10	3.93 – 5.22	4.50 – 5.30	4.63 – 6.08
HGB / g/dL	12.0 – 16.0	11.2 – 15.7	13.0 – 16.0	13.7 – 17.5
HCT / %	36.0 – 46.0	34.1 – 44.9	37.0 – 49.0	40.1 – 51.0
MCV / fL	78.0 – 102.0	79.4 – 94.8	78.0 – 102.0	79.0 – 92.2
MCH / pg	25.0 – 35.0	25.6 – 32.2	25.0 – 35.0	25.7 – 32.2
MCHC / g/dL	32.0 – 37.0	32.2 – 35.5	32.0 – 37.0	32.3 – 36.5
RDW / %	11.5 – 14.0	11.7 – 14.4	11.5 – 14.0	11.6 – 14.4
PLT / x10(3)/mcL	150 – 450	182 – 369	150 – 450	163 – 337
MPV / fL	7.2 – 11.1	9.4 – 12.3	7.2 – 11.1	9.4 – 12.4
Absolute Neutrophil / x10(3)/mcL	2.10 – 11.52	1.56 – 6.13	2.10 – 11.52	1.78 – 5.38
Absolute Lymphocyte / x10(3)/mcL	0.77 – 5.85	1.18 – 3.74	0.77 – 5.85	1.32 – 3.57
Absolute Monocytes / x10(3)/mcL	0.14 – 1.30	0.24 – 0.86	0.14 – 1.30	0.30 – 0.82
Absolute Eosinophil / x10(3)/mcL	0 – 0.78	0.04 – 0.36	0 – 0.78	0.04 – 0.54
Absolute Basophil / x10(3)/mcL	0 – 0.26	0.01 – 0.08	0 – 0.26	0.01 – 0.08
NRBC / 100 WBC	0	0	0	0
IG / %	0.00 – 0.75	0.00 – 0.75	0.00 – 0.75	0.00 – 0.75
RETIC / %	0.6 – 2.7	0.5 – 1.7	0.6 – 2.7	0.5 – 1.8
Absolut RETIC / x10(6)/mcL		0.0164 – 0.0776		0.0260 – 0.0950
RET-He / pg		> 28		> 28
IPF / %		0 - 3		0 - 3

Sources:

- Sysmex Hematology Analyzer XN Series Instruction for use, revised 07/2015
- Roehrl, M.H., et al. Age-Dependent Reference Ranges for Automated Assessment of Immature Granulocytes and Clinical Significance in an Outpatient Setting. Arch Pathol Lab Med. 2011, 135(4):471-7.
- Kickle, T.S., et al. A Clinical Evaluation of High Florescent Platelet Fraction Percentage in Thrombocytopenia. Am J Clin Pathol 2006;125:282-287
- Reticulocyte hemoglobin content. Alan Mast, Morey Blinder, Dennis Dietzen. Blood: Vol 83, Issue 4, 307-310

Addendum 1 (continued):

Pediatric Reference Ranges

Pediatric CBC and Differential Reference Ranges											
Parameter/Units of Measurement	0d	2d	3d	2w	1m	2m	3m	6m	1y	2y	6y – 12y
WBC/ x10(3)/mcL	19.0-25.0	9.0-29.9	9.0-29.9	9.0-29.9	5.0-19.5	5.0-19.5	5.0-19.5	6.0-17.5	6.0-17.5	6.0-17.0	5.0-16.0
RBC/ 10⁶/μL	4.00-6.60	3.90-5.90	3.90-5.90	3.90-5.90	3.10-5.30	3.10-5.30	2.70-4.50	3.10-5.10	3.90-5.50	3.90-5.50	3.90-5.50
HGB/ g/dL	14.5-22.0	13.4-19.9	13.4-19.9	13.4-19.9	10.7-17.1	9.1-14.0	9.1-14.1	9.5-14.1	11.3-14.1	11.3-14.1	11.5-14.0
HCT/ %	45.0-65.0	42.0-65.0	42.0-65.0	42.0-65.0	33.0-55.0	28.0-42.0	29.0-41.0	29.0-41.0	31.0-41.0	31.0-41.0	34.0-42.0
MCV/ fL	95.0-121.0	88.0-123.0	88.0-123.0	88.0-123.0	88.0-123.0	91.0-112.0	74.0-108.0	74.0-108.0	70.0-86.0	70.0-86.0	73.0-87.0
MCH/ pg	31.0-37.0	31.0-37.0	31.0-37.0	31.0-37.0	27.0-36.0	27.0-36.0	25.0-35.0	25.0-35.0	23.0-31.0	23.0-31.0	24.0-30.0
MCHC/ g/dL	29.0-37.0	28.0-36.0	28.0-36.0	28.0-36.0	28.0-36.0	28.0-36.0	28.0-36.0	30.0-36.0	30.0-36.0	30.0-36.0	31.0-36.0
RDW/ %	11.5-14.0	13.0-18.0	13.0-18.0	11.5-16.0	11.5-16.0	11.5-16.0	11.5-16.0	11.5-16.0	11.0-15.0	11.0-15.0	11.0-15.0
PLT/ x10(3)/mcL	150-450	150-450	150-450	150-400	150-400	150-400	150-400	150-400	140-400	140-400	140-400
MPV/ fL	7.2-11.0	7.5-11.6	7.5-11.6	7.5-11.6	7.5-11.6	7.5-11.6	7.5-11.6	7.5-11.6	7.5-11.6	7.5-11.5	7.5-11.5
Absolute Neutrophils/ x10(3)/mcL	11.59-18.75	5.31-21.90	4.41-18.30	2.97-12.30	1.50-7.02	1.50-7.02	1.65-8.00	1.98-7.18	1.98-7.18	2.22-7.65	2.10-11.52
Absolute Lymphs/ x10(3)/mcL	3.61-5.75	2.43-11.40	2.61-12.00	4.68-20.70	2.75-14.04	2.75-14.04	2.80-12.29	3.30-12.60	2.88-11.03	2.88-10.71	1.75-7.68
Absolute Monocytes/ x10(3)/mcL	0.95-3.75	0.00-1.50	0.36-2.40	0.18-1.80	0.10-1.17	0.15-1.95	0.15-1.95	0.18-1.75	0.18-1.75	0.18-1.70	0.15-1.60
Absolute Eosinophils/ x10(3)/mcL	0.00-1.50	0.00-1.80	0.00-1.80	0.00-1.80	0.00-1.17	0.00-1.17	0.00-1.17	0.00-1.05	0.00-1.05	0.00-1.02	0.00-0.96
Absolute Basophils/ x10(3)/mcL	0.00-0.50	0.00-0.60	0.00-0.60	0.00-0.60	0.00-0.39	0.00-0.39	0.00-0.39	0.00-0.35	0.00-0.35	0.00-0.34	0.00-0.32
Nucleated RBC/ 100 WBC	0	0	0-8	0	0	0	0	0	0	0	0
IG / %	0.00-6.00	0.00-6.00	0.00-0.35	0.00-0.35	0.00-0.35	0.00-0.35	0.00-0.35	0.00-0.35	0.00-0.35	0.00-0.35	0.00-0.35
Retic / %	1.0-7.5	1.0-7.5	1.0-7.5	0.6-2.7	0.6-2.7	0.6-2.7	0.6-2.7	0.6-2.7	0.6-2.7	0.6-2.7	0.6-2.7

The reference ranges should be interpreted as from and including the age specified in the title of the column

Addendum 1 (continued):

Pediatric and Adult % Differential Reference Ranges

Parameter/Units of Measurement	0d	1d	2d	3d	4d	6d	8d	15d	1m	2m	3m	4m	7m	1y	13m	3y	4y	5y	6y	7y	12y+
Neutrophils/ %	61 - 75	60 -74	59-73	49-61	45-55	37-45	33-41	29-35	30-36	30-36	33-41	33-41	35-43	35-43	37-45	39-47	42-52	44-54	42-72	42-72	42-72
Bands / %	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15	1-15
Myelocyte/ %	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Metamyelocytes/ %	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1
Promyelocytes/ %	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Blast / %	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Lymphocytes/ %	19-23	22-28	27-33	29-35	32-40	40-48	46-56	52-64	55-67	55-67	56-68	55-67	52-64	48-58	48-58	43-53	42-52	39-47	35-43	34-42	17-45
Atypical Lymphocytes / %	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5	0-5
Monocyte / %	5-15	5-15	4-8	4-8	4-8	4-8	4-8	2-6	2-6	3-10	3-10	3-10	3-10	3-10	3-10	3-10	3-10	3-10	3-10	3-10	3-10
Basophil/ %	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2	0-2
Eosonophil/ %	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6	0-6
Plasma Cell/ %	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1	0-1

Reference: % Cell Differential were obtained from Sunquest Quality Assurance's Database

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Addendum 2:**Sysmex XN Maintenance**

1. Both Daily and Weekly shutdowns will take place between 3 and 4am.
2. To record the Pressure Vacuum:
Analyzer Mode – Maintenance – Pressure Adjustment
3. If for any reason daily shutdown cannot be done on a Sysmex XN, Daily Cleaning can be used as an alternative to the shutdown to keep one analyzer up and running at all times.

Daily Cleaning	
1.	Verify the analyzer is in the Ready state
2.	Click the Analyzer Menu button
3.	Select Maintenance
4.	Select Cleaning
5.	The tube holder will slide out
6.	Place a vial of CELLCLEAN AUTO in the sample tube holder
7.	Press the blue start switch.
Note: Cleaning will take approximately 20 minutes. The cleaning process will conclude with a background check and the analyzer will return to the Ready state in the Manual Mode.	

Instrument Startup after a Complete Shutdown	
1.	Verify the power switches on each device is turned on
2.	Press the Startup switch (green switch) on the samples
3.	Turn on the IPU and log in
4.	Once the power on the instrument turns on, a self-test automatically runs for about 10 minutes doing initialization of the mechanical parts, rinse & temperature stability

Sysmex XN Manual Shut down	
1.	Make sure the Status indicator LED on the analyzer is green
2.	Go to Menu
3.	Select Shut down and the tube holder (for manual/pediatric samples) slides out forward
4.	Place a CELLCLEAN AUTO in the tube holder
5.	Press the Start switch

Addendum 2:

Sysmex XN Maintenance (continued)

Sysmex XN Automatic Shut down	
1.	Make sure both the analyzer and the sampler are in the ready state
2.	Make sure that the tube holder (for manual/pediatric samples) is retracted into the analyzer
3.	<p>GEC: Place a CELLCLEAN AUTO in a regular patient rack in position 10; load the rack on the instrument.</p> <p>SG & WAH: Place a CELLCLEAN AUTO on a regular patient rack and load the rack on the instrument:</p> <ul style="list-style-type: none"> a. Position 9: Analyzer Left b. Position 10: Analyzer Right <p>You can shut down one particular analyzer by placing CELLCLEAN AUTO only in the position for that analyzer.</p>
4.	If the sampler auto – start function is ON, conveying automatically starts when the rack is loaded. If the sample auto – start function is OFF, click the sampler analyzer button in the control menu and then click start
5.	Shut down is performed automatically
6.	When the instrument is ready to be initialized the message Press Start SW will appear. Press the green button to initialize

Weekly IPU Shut down	
	SG & WAH, Note: This shut down does not affect the SP – 10 Stainer, so the unit could remain ON.
1.	Make sure XN analyzer(s) is/are in ready mode
2.	Open the front hood cover, use the black switch to turn the analyzer(s) OFF
3.	From Main Menu select Exit IPU
4.	From Windows screen restart the computer
5.	When the computer is back ON, log back in to IPU
6.	Now turn the analyzer(s) back ON

Addendum 3:

Sysmex Stainer SP-10 Maintenance and Quality Control

1. Stain quality and differential comparison for the SP-10 stainer is performed each day of use. Refer to the Automated Stainer Differential Comparison and Stain Quality Log for criteria.
2. Both Shutdown 1 and Shutdown 2 will be done by day shift.
3. Daily Shutdown Automatically:

Stainer SP – 10 Automatic Shutdown (Shutdown 1)	
1.	Before the daily shut down is initiated, verify the following: <ol style="list-style-type: none"> a. The instrument is in a ready mode and there are no racks/samples on board b. There is at least 12 single cassettes on the cassette supply table c. There is at least 450 mL of methanol in the methanol bottle
2.	Place a cell clean Auto in position 8 of a sample rack. Load the rack onto the right sampler pool (analyzer side)
3.	Shutdown is performed automatically

4. Daily Shutdown Manually:

Stainer SP – 10 Manual Shutdown (Shutdown 1)	
1.	Before the daily shut down is initiated, verify the following: <ol style="list-style-type: none"> a. The instrument is in a ready mode and there are no racks/samples on board b. There is at least 12 single cassettes on the cassette supply table c. There is at least 450 mL of methanol in the methanol bottle
2.	Select Conv.int. on the menu screen – Interrupt – Return – Select Shutdown on the menu screen – Select Shutdown 1 (daily)
3.	Place a cell clean Auto in position 10 of a sample rack
4.	Place the rack on the sampler (SP-10 side), lining the tube up with the gripper. Use the mark on the sampler as a guide for the left edge of the rack. Press OK .

5. Weekly Shutdown:

Stainer SP – 10 Shutdown (Shutdown 2)	
1.	Before the daily shut down is initiated verify the following: <ol style="list-style-type: none"> a. The instrument is in a ready mode and there are no racks/samples on board b. There is at least 9 single cassettes on the cassette supply table c. There is at least 800 mL of methanol in the methanol bottle
2.	Select Conv.int. on the menu screen – Interrupt – Return – Select Shutdown on the menu screen – Select Shutdown 2 (weekly)
3.	Place a tube of CELLCLEAN AUTO in the 10th position of a sample rack. Place the rack on the sampler (SP-10 side), lining the tube up with the gripper. Use the mark on the sampler as a guide for the left edge of the rack. Press OK .
4.	When shutdown ends, the device will automatically turn off. Slide the rack to the left on the analysis line, and then remove.

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Addendum 3:

Sysmex Stainer SP-10 Maintenance (continued)

6. To load a sample in a Manual Mode:

Note: This mode could be used for STAT specimens.

Manual Mode	
1.	Select Conv.int. in the SP-10's Main Menu screen
2.	Select Interrupt – Return - Manual in the main menu screen
3.	Select tube type and touch Closed. Enter Sample ID
4.	Mix the sample tube and place it in position 10 of a sample rack
5.	Set the rack so that its left end matches up with the label on the conveyor, select START
6.	Wait until the smear is prepared and the sample is returned to the rack then remove the rack
7.	Select Return in the manual screen - Select Conv.int. in the main menu screen – Select Stop int

7. Daily Maintenance:

Clean Spreader Glass	
1.	Select Maint. on the menu screen - Spreader glass – OK
2.	When prompted to replace the spreader glass, open the top cover - Clean the surface of the spreader glass with gauze moistened with reagent grade water
3.	Close the top cover, then select OK – Select Cancel
Note: Do NOT reset the counter after cleaning. If the counter is reset, the message to replace the glass will appear at the wrong time.	

Clean Cassettes	
1.	Remove the used cassettes from the instrument and place them in the assigned container
2.	Fill the container with Methanol until the tops of the cassettes are covered
3.	Take the cassettes out of the container immediately. Do NOT allow the cassettes to sit in Methanol for more than 5 minutes
4.	Allow the cassettes to air dry

8. Monthly Maintenance:

Monthly Maintenance	
1.	Use a damp gauze and wipe and clean the sample racks
2.	Use a damp gauze and wipe and clean the right and left sampler rack pools
3.	Use a damp gauze and clean the measurement line

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Addendum 4:**QC Instructions on the Sysmex XN****To document outliers of daily control runs:**

1. Click on **QC File**
2. Double click on the **QC level with the outlier** to view the Levy Jennings Chart
3. From the top menu select **Manage**
4. Under the Specify Excluded Section, select **Not Managed**
5. Under the Comments Settings, click the **drop down arrow** and choose a comment that qualifies with the outlier
6. Click on **OK** to save

To register a new lot:

1. From **Main Menu** Select **QC File**
2. High light one empty row then click **Register**
3. Update Control Level, Lot number and Expiration date
4. High light all methods from top to bottom then click on **Variable Target**
5. Click on **Restore**
6. Select the correct file for desired QC level, then select **OK**

Parallel test new controls by analyzing each level of control a minimum of twice a day for 5 days on each analyzer prior to expiration of previous lot. After a minimum of 10 data points are accumulated on each analyzer, auto set the targets using the steps below:

1. From **Main Menu**, select **QC File**
2. Select desired lot, then click **Modify**
3. Highlight all methods from top to bottom, then click on **Auto Setting**
4. Confirm that the check box for **Target** is set. Do **NOT** select the check box for Limit.

To adjust QC ranges:

1. From **Main Menu**, Select **QC File**
2. Double click on the desired lot
3. Click on **Range**
4. Drag the graph to cover the most recent shift/bias
5. Select **Modify**
6. Select the desired method, click on **Auto Setting**
7. Confirm that the check box for **Target** is set. Do **NOT** select the check box for Limit.

To be able to see two different lots of QC on the same page or to see the same QC of both instruments on the same page:

1. From **Main Menu** Select **QC File**
2. Open up desired QC Levy Jennings chart, on top click on **Reference**
3. Choose either **Compare QC Files** or **Compare Analyzers**

Addendum 5:

QC Instructions on Insight for Supervisor or Designee

Insight is the website used to submit and review QC data. All QC data is submitted to insight automatically.

To log onto Insight go to www.sysmex.com/Insight

Each lot of QC has two periods; QC needs to be reviewed and submitted by the Supervisor or Designee before the due date for each period, otherwise the lab's data will not be included in the peer group study.

Note: If for some reason the due date is missed, QC can still be submitted and a report will be generated, but the data will not be included in peer group.

Since QC is submitted automatically, Supervisor or Designee must log onto Insight to manage the data. If the outliers are not excluded before the due date of each Period, the QC report will include the data points that should be excluded such as QNS runs.

It is recommended that the supervisor or designee log onto Insight and check each lot to make sure data has been transmitting, before the due date for each Period.

To exclude/manage outliers:

- Log onto Insight
- On the Home Page, on the left side of the screen locate **QC data**
- Select **Review QC Data**
- Select the desire analyzer, lot and QC
- Click on **Review your data** – this page will show every run including date and time
- To exclude a run, click on **Manage Data**
- Select **Not Managed**
- Select a pre-typed comment from **Select Comment Type** or under **Enter Comment Description** free text a comment
- Save comment – back to review data

To pull a report or review weekly/monthly QC:

- Log onto Insight
- From Home Page, under **Report Center** – Select **Customer QC Report**
- When new page opens, click on **View All QC Reports (including Lot – to – Date Reports)**. This page will list all the lot numbers up to 2 years. Note that each lot appears three different times; Period 1, Period 2, and Cumulative.
- Cumulative has the data from Periods 1 and 2 included with peer group.

Reviewing the report:

The report includes every method's average run, Assay Mean and Group Mean. The following flags will appear under notes, depending on how the QC performs:

- W: Warning
- P: Positive Bias
- N: Negative Bias

If Warning appears for a method once, that means to check the instrument and do maintenance to make sure it is working properly.

If two consecutive Warnings, Positive Bias, or Negative Bias appear, call technical support and trouble shoot with them. They will examine the situation and might send service over to do a calibration and other trouble shooting.

To find the Insight calendar:

- Log onto Insight –
- From the Home Page, under **Reference/Documents** – Select **Lot Calendars**
- Select the **XN CHECK Calendar**. This provides information such as:
 - When each Period begins
 - When each Period ends
 - Due date to submit data

Addendum 6:**Smear Review and Manual Differential****A. General Instruction/Notes**

1. All smears for WBC differential must be made with the wedge pull film technique. A properly made Wright stain blood smear should have the following characteristics:
 - a. RBC - Pink with central pallor
 - b. NRBC - Dark purple nucleus.
 - c. WBC
 - Neutrophil - Dark purple nuclei with light pink cytoplasm dotted with lilac granules.
 - Lymphocyte - Dark purple nucleus. Cytoplasm with varying shades of blue (robin egg blue).
 - Monocyte - Cytoplasm of monocytes stains a faint bluish gray tinge.
 - Eosinophil - The eosinophilic granules, bright red to orange.
 - Basophil - The basophilic granules very dark bluish purple.
2. Prepare blood films within 4 hours of the blood collection in EDTA.
3. Stain the film within one hour of preparation with Wright stain.
4. The leukocytes must be well preserved, and anticoagulant effects such as excessive vacuolization or changes in nuclear shape must be minimal. Less than 2% of the leukocytes may be smudged, except in some lymphoproliferative disorders.
5. There should be sufficient working area with minimum 2.5 cm in length terminating at least 2 cm from the end of the slide.
6. Acceptable morphology within working area and no artifact introduced by the technique. Also there should be minimum distributional distortion.
7. A far end that becomes gradually thinner, without growing streaks, troughs, or ridges, all of which indicate an increased number of leukocytes carried into this area.

B. Prepare a blood smear and Scan:

1. Transfer a small drop of blood (2-3mm) to a pre-cleaned slide, frosted side up. The blood is placed in the centerline of the slide just past the frosting.
2. A second slide is used as a spreading slide. The "pusher" slide is placed at an angle of 30-45 degrees to the slide containing the blood drop. The slide is moved back to make contact with the drop, allowing the blood to spread the entire width of the slide.
3. The "pusher" slide is then quickly and smoothly pushed forward to the end of the smear, creating a wedge smear. It is important that the whole drop is picked up and spread quickly. Moving the pusher slide too slowly accentuates poor leukocyte distribution by pushing larger cells to the very end and the slides of the smear.
4. The drop of blood should be of such a size that the film is about 30mm in length. (If necessary repeat this process until an acceptable slide is obtained.)
5. Label the slide using a sharp pencil. On the frosted side, record the patient's last name, or as much as will fit on the slide and accession or barcode. Allow the slide to air dry. Load the slide on the slide stainer.

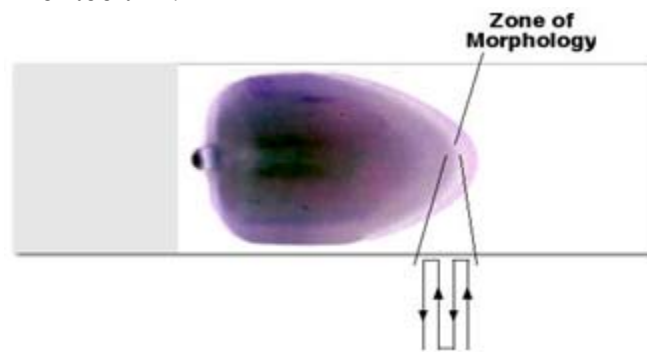
- Use the 50x oil immersion objective and examine the smear microscopically. Check the smear to see that it is well made, the distribution of the cells is uniform and the staining is satisfactory.

Criteria for a well-stained smear include:

- No precipitated stain should be seen.
 - The erythrocytes are orange or pink.
 - The nuclei of the leukocytes are purplish-blue.
 - Neutrophilic granules are reddish to pink-lilac.
 - Eosinophilic granules are red to orange.
 - Basophilic granules are very dark bluish-purple.
 - Platelets stain dark blue-purple.
- Scan the slide and look for abnormal or suspicious cells that may be in disproportionately low numbers. Look for nucleated red cells, immature cells, atypical lymphocytes and platelet clumps or large platelets. Estimate the white cell count to see if there is any gross error in the instrument count. This could also detect a clotted specimen or perhaps a mix-up in blood specimens.
 - If no significant abnormalities are noted the comment "*Smear review agrees with automated differential. No significant red cell or platelet abnormalities are noted*" should be entered.
 - If a smear review shows a discrepancy with the automated differential or reveals any significant abnormalities or problems for which a manual differential is deemed medically necessary, a manual differential will be performed.

C. Differential Counting Procedure

- Use the "battlement" tract for this examination. Each identified cell must be classified by cell type. Cells found in a Differential are: neutrophil, segmented; neutrophil, band; lymphocyte, normal; lymphocyte, variant or atypical; monocyte; eosinophil; basophile; other nucleated cells (except nucleated red cells). Include distorted cells that are clearly identifiable in the appropriate classification. While counting the cells, make a note of any abnormalities present in the cells. It is important to examine cellular morphology and to count leukocytes in areas that are neither too thick nor too thin.



2. On each slide, 100 leukocytes should be counted. If the blood is leukopenic, process additional slides in parallel. Exceptions to the 100 cell differential are as follows:

If	Then
WBC greater than 25,000/MM ³	Perform a 100 cell differential count and compare counts to the automated counts. Perform a 200 cells count if difference is greater than 10%.
WBC of greater than 2,000 and less than 25,000	A 100 cell differential count is to be performed
WBC less than 2,000	Perform a 100 cell count if possible; however, a 50 cell differential may be performed, insure correct percentage is entered in LIS.
WBC less than 500	Stain 2-3 slides and perform classifying as many cells as possible, ensure correct percentage is entered in LIS.

Note: When the standard 100 cell Differential is not performed, the number of cells counted must be noted in the LIS.

3. Express the results of the differential count as a percentage of all the leukocytes counted.
4. Count nucleated red blood cells present and report the result as the number per 100 leukocytes counted.
5. Examine the red cell morphology in a thin area of the slide where the red cells either do not overlap or lightly overlap. They should have a central pallor. In most cases an abnormality must be a consistent finding in order to be significant. Note any variations from normal and classify them according to section 13.7.
 - All clinically significant findings such as specific cell types, inclusions, polychromasia, etc., will be reported from the smear evaluation.

IF	Then
NO clinically significant findings to be added to a patient report.	Result as Normal.
ANY additions to the patient report, such as RBC morphology, cell differential, PLT morphology, etc.	Report all clinically significant findings using the Diff key board in the LIS or using the cell counter in DI

6. Examine the smear for platelets morphology and number. Find a thin area where red cells are not overlapping. Perform a PLT estimate and use the following if comparing to automated count.

IF	Then
In the presence of a platelet flag, a platelet estimate must be performed. Using the 100X objective	Count the PLT in each of 10 microscopic fields in areas of the slide where the RBCs are evenly dispersed. Divide the total # of platelets by 10 to establish the mean and multiply by 20,000.
The Sysmex platelet count and the platelet estimate do not agree within $\pm 20\%$	Repeat the platelet estimate and/or platelet count. If counts still do not agree, consult the supervisor or designee.

D. Germantown Emergency Center: differentials that are to be reviewed by the Pathologist will be sent via courier to Shady Grove Medical Center Hematology section with a Pathologist Slide Review Request. SGMC staff will take the slide and paperwork to the pathologist for review.

NOTE: If malaria is observed in the blood smear: Call the patient's physician and report your finding. The physician may request a malaria smear review and identification.

Addendum 7:

CBC DIFF/SCAN Action and Repeat Criteria

Criteria	Criteria Action Limit	Action 1	Action 2
WBC	< 0.5	Check for a clot, re-analyze sample in LW mode	SCAN
WBC	≤ 2.0	Check for a clot	DIFF
WBC	≥ 30.0	Check for Giant Platelets	SCAN
WBC	≥ 440.0	Re-analyze using dilution	SCAN
RBC	≥ 8.60	Re-analyze using dilution	SCAN
HGB	< 4.0	Re-analyze with the Aspiration Sensor OFF	None
HGB	≤ 6.0	Check for a clot	None
HGB	≥ 20.0	Re-analyze, check Coag sample if HCT ≥ 55	None
HGB	≥ 25.0	Re-analyze using dilution, check Coag sample if HCT ≥ 55	None
MCV	≤ 50.0	Check for a clot, re-analyze and verify morphology	SCAN
MCV	≥ 130.0	Warm sample for 30 min, re-analyze, verify morphology	SCAN
PLT	< 50	Check for a clot, re-analyze	None
PLT	> 5000	Re-analyze using dilution, perform PLT estimate	SCAN

Criteria	Criteria Action Limit	Action 1	Action 2
Lymph	≥ 70	Pathologist review for patient >17 y/old	DIFF
Neutro	≥ 90 no flags	Scan for bands, hyper-segmentation, toxic granulation	SCAN
Neutro	≥ 90 & flags	Rule out immature Neutrophils	DIFF
Mono	> 25	Rule out immature Monocytes	DIFF
Eos	≥ 35	Scan to verify Eosinophils. Check for parasites	SCAN
Baso	≥ 3.5	Mix for 5 min, re-run. If still elevated SCAN	SCAN
Retic	>30.0%	Check for a clot, Vortex 1 -2 min, re-analyze	Repeat

Criteria	Action
WBC Abn Scattergram	If dashes appear: re-analyze, if dashes still remain perform a DIFF If asterisk appears: scan, perform a DIFF if abnormal cells observed
NRBC Present	Release the result. No further correction of WBC is required
IG Present >5%	Perform a DIFF and check for the presence of: Promyelocytes, Metamyelocytes and Myelocytes
Blast/Abn Lymph	If dashes appear: re-analyze, if dashes still remain perform a DIFF If asterisk appears: perform a DIFF, check for blasts & immature granulocytes
Left Shift	If dashes appear: re-analyze, if dashes still remain perform a DIFF and check for bands, toxic granulation, hyper-segmentation, vacuolation of neutrophils
Atypical Lymph	Perform a DIFF
RBC Abn Distribution	SCAN, Perform a morphology (only note if findings are ≥2+)
Dimorphic Population	SCAN, Perform a morphology
RET Abn Scattergram	Vortex 1 -2 min, re-analyze. If flag remains, scan smear for abnormal cells

Criteria	Action
RBC Agglutination	Warm & re-analyze. If flag still remains perform plasma replacement
Turbidity/HGB interference	Re-analyze, if flag remains SCAN. Check for the presence of Spherocytes. If RBC agglutination is noted warm sample and re-analyze. If lipemic or Icteric, perform a plasma replacement.
Fragments	SCAN for fragmented RBCs & other poikilocytosis
PLT Abn Distribution & PLT Abn Scattergram	GEC: Vortex 1 – 2 min, then re-analyze WAH & SG: Vortex 1 – 2 min, re-analyze in PLT-F mode All sites: If flag remains, SCAN and perform a morphology
PLT Clumps	Check for clots. If present, cancel and request re-draw. If no clots, then GEC: Vortex 1 – 2 min, then re-analyze WAH & SG: Vortex 1 – 2 min, re-analyze in PLT-F mode All sites: If flag remains, SCAN and perform a morphology If fibrin or clumps are present, recollect a new sample in sodium citrate

Addendum 8:**Sysmex Rules Management**

	To add or modify a rule
1.	From the Main Menu select Rule
2.	For Sysmex Instrument rules – select the Rerun/Reflex/Comment Rule tab For the Stainer – select SP Rule tab
3.	High light the desired rule, then select Modify on top of the screen
4.	To set up a rule, for example RBC >= 8.6, a. Under Function , select item value b. Under Analysis items , select RBC c. Use the keyboard on the screen (not the PC keyboard) to add desired rule d. Click on Equation to check the formula e. Under Action/Comment , choose the repeat criteria and type the desired comment

	To import rules from a flash drive
1.	Insert the flash drive
2.	From the Main Menu, select Rule
3.	Select File on top of the screen
4.	Select Restore , locate the desired file on flash drive and accept to save.

	To export rules from a flash drive
1.	Insert the flash drive
2.	From the Main Menu, select Rule
3.	Select File on top of the screen
4.	Select Back Up , save the file on flash drive and accept to save.