Issuing Authority: Director, Health Services

Next Review Date:

Type: Laboratory Services Program SOP

Policy Number: Date Approved:

PROGRAM Standard Operating Procedure – Laboratory Services			
Title: MIC20300 – Gram stain reporting	Policy Number:		
in LIS-Respiratory Specimens			
Program Name: Laboratory Services			
Applicable Domain: Lab, DI and Pharmacy Services			
Additional Domain(s):			
Effective Date:	Next Review Date:		
Issuing Authority:	Date Approved:		
Director, Health Services			
Accreditation Canada Applicable Standard: N/A			

GUIDING PRINCIPLE:

The culture of poorly collected respiratory specimens is a wasteful use of laboratory resources and can lead to erroneous reporting and treatment of patients. These specimens need to be scored for acceptability using the Q-score method.

PURPOSE/RATIONALE:

This standard operating procedure describes how to report the Gram stain results of respiratory specimens in the LIS in a consistent manner.

SCOPE/APPLICABILITY:

This procedure applies to Medical Laboratory Technologists (MLTs) reporting respiratory specimens in the LIS.

SAMPLE INFORMATION:

		Sputum, Endotracheal aspirates (ETT) and Auger Suction
		specimens are Q-scored for quality
Туре	•	Bronchial aspirates (washings), Bronchoalveolar lavage
Туре		(BAL) specimens and specimens from cystic fibrosis
		patients are NOT Q-scored for quality
	•	Refer to MIC10100-Microbiology Specimen Processing

REAGENTS and/or MEDIA:

- Methanol
- Gram Crystal Violet
- Gram Iodine (Stabilized)

- Gram Decolorizer
- Gram Safranin

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Policy Number: Date Approved: Page 1 of 8

Title: MIC20300-Gram stain reporting in LIS-Respiratory Specimens

Issuing Authority: Director, Health Services

Next Review Date:

Type: Laboratory Services Program SOP Policy Number:

Date Approved:

SUPPLIES:

Glass microscope slide

QC slide

Immersion oil

Slide storage tray

EQUIPMENT

- Hot plate
- Microscope

SPECIAL SAFETY PRECAUTIONS:

Containment Level 2 facilities, equipment, and operational practices for work involving infectious or potential infectious materials or cultures.

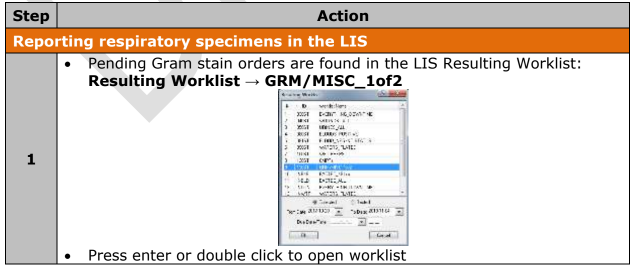
- Ensure that appropriate hand hygiene practices be used.
- Lab gown must be worn when performing activities with potential pathogens.
- Gloves must be worn when direct skin contact with infected materials is unavoidable.
- Eye protection must be used when there is a known or potential risk of exposure of splashes.
- All procedures that may produce aerosols or involve high concentrations or large volumes should be conducted in a biological safety cabinet (BSC).
- The use of needles, syringes and other sharp objects should be strictly limited.

All patient specimens are assumed to be potentially infectious. Routine Practices must be followed. Since viable micro-organisms are used, all cultures must be handled with appropriate precautions. All equipment in contact with cultures should be decontaminated by appropriate methods.

QUALITY CONTROL:

- Quality control is performed daily
- A TQC order is automatically generated daily to record the QC results
- Refer to MIC60060-Microbiology Stain Quality Control

PROCEDURE INSTRUCTIONS:



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Policy Number: Date Approved: Page 2 of 8

Issuing Authority: Director, Health Services Next Review Date:

Type: Laboratory Services Program SOP Policy Number: Date Approved:

2

3

- Enter the accession number on the slide and select enter to mark the order
- Select enter again to open Result Entry or double click on accession number to open

<u>Under low power (X10, LPF)</u>: screen slide to locate good specimen areas to obtain an overall impression of cell types present.

- Observe slide for stain crystals:
 - If an excess of precipitated stain is observed, prepare another smear
 - If precipitate continues, use freshly filtered crystal violet
- Determine if slide has been properly decolorized:
 - Depending on the source of the specimen, the background should be generally clear or Gram negative
 - If white blood cells are present, they should appear completely Gram negative
 - > If slide is over decolorized, prepare another smear
- Determine if thickness of smear is appropriate:
 - For proper interpretation, areas must be no more than one cell thick, with no overlapping of cells. Prepare a new slide if unreadable
- Examine for evidence of inflammation:
 - Determine areas representative of inflammation and areas of contamination with squamous epithelial cells

<u>Under low power (X10, LPF)</u>: average the number of epithelial cells and white blood cells:

4

None seen	No cells seen
1+	< 1 cell seen
2+	1 - 9 cells seen
3+	10 - 25 cells seen
4+	> 25 cells seen

Calculate the Q-score of the specimen. The Q-score is calculated by assessing the quantity of epithelial cells and neutrophils. Examine 20 to 40 fields and interpret as follows:

5

Q-score Table				
Eni celle/LDE	White blood cells /LPF			
Epi cells/LPF	0	1-9	10-25	>25
0	Q 0	Q 1	Q 2	Q 3
1-9	Q-1	Q 0	Q 1	Q 2
10-25	Q-2	Q 1	Q 0	Q 1
>25	Q-3	Q-2	Q-1	Q 0

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Policy Number: Page 3 of 8

Title: MIC20300-Gram stain reporting in LIS-Respiratory Specimens

Issuing Authority: Director, Health Services

Next Review Date:

Type: Laboratory Services Program SOP
Policy Number:
Date Approved:

6	Do not perform or report the Q-score on Bronchial aspirates (washings), Bronchoalyeolar layage (BAL) or specimens from cyclic fibrosis nationts				
Bronchoalveolar lavage (BAL) or specimens from cystic fibrosis patients. If the Q-score indicates the sample is of good quality (Q-score 1-3 or Q-score 0 or <0 if patient is immunocompromised), add one drop of immersion oil to the slide. In a representative area with predominance of inflammation or purulence using the oil immersion lens (100X), examine 20 to 40 fields to observe cell morphology and Gram reaction. Under oil immersion (X100, OIF): quantitate epithelial cells, white blood cells, red blood cells and bacteria as follows:					
		None seen	No cells seen		
		1+	< 1 cell seen		
8		2+	1 - 9 cells seen		
		3+	10 - 25 cells seen		
		4+	> 25 cells seen		
NOTE: Bacteria are not reported if the Q-score indicates specimen is unsatisfactory for culture					
Under the test code: STGM2 , use the STGM2 keypad to report the quantity of epithelial cells, white blood cells and bacteria if indicated by Qscore. Report cells in this order to maintain consistency with reporting.					
	Reporting Mixed oropharyngeal flora in respiratory gram stain: 1. If smear has ≥2 morphotypes and neither are predominant or				
10	 intracellular, mixed oropharyngeal flora can be reported If smear has ≥2 morphotypes and one or more are predominant or intracellular, the predominant or intracellular morphotypes are reported individually and other morphotypes are reported as mixed oropharyngeal flora 				

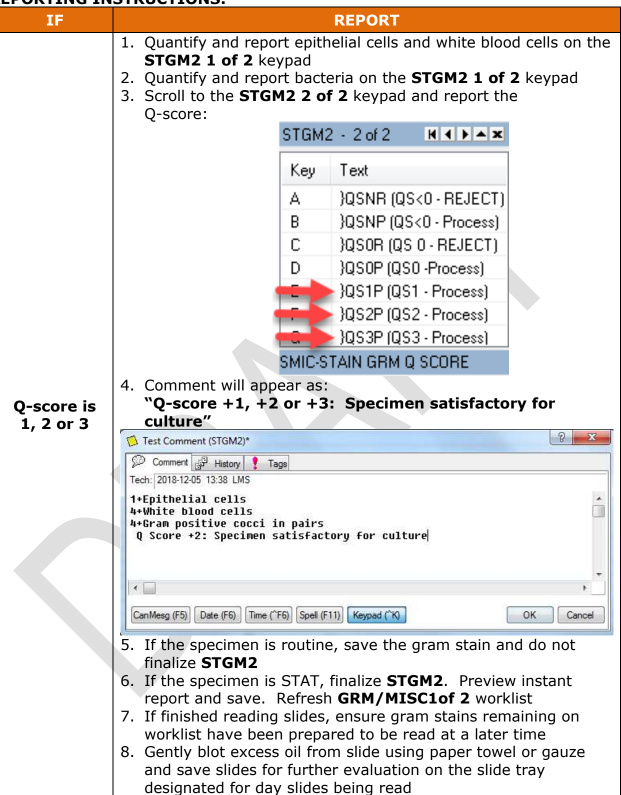
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Policy Number: Date Approved: Page 4 of 8

Issuing Authority: Director, Health Services Next Review Date:

Type: Laboratory Services Program SOP Policy Number: Date Approved:

REPORTING INSTRUCTIONS:



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Next Review Date:

Type: Laboratory Services Program SOP Policy Number:

Date Approved: IF REPORT 1. Quantify and report epithelial cells and white blood cells on the STGM2 1 of 2 keypad 2. Do not report bacteria 3. Scroll to the **STGM2 2 of 2** keypad and report the Q-score: H + F + XSTGM2 - 2 of 2 Text Kev)QSNR (QS<0 - REJECT) QSNP (QS<0 - Process) |}QSOR (QS 0 - REJECT)| (QSO -Process) D Ε QS1P (QS1 - Process) QS2P (QS2 - Process) G QS3P (QS3 - Process) SMIC-STAIN GRM Q SCORE 4. Comment will appear as: Q-score is "Q Score 0 or <0: Suggestive of poor quality; culture not 0 or <0 performed. Please recollect if clinically indicated" Test Comment (STGM2)* Patient is NOT Comment History ? Tags immune-Tech: 2018-12-05 13:40 LMS compromised 4+Epithelial cells No white blood cells seen Q Score 0: Suggestive of poor quality; culture not performed. Please recollect if clinically indicated. CanMesg (F5) Date (F6) Time (^F6) Spell (F11) Keypad (^K) Cancel 5. Select **OK**. Standard deviation rule violation box will pop up indicating that the culture will be cancelled. Select **OK** 6. Short cancellation reason box will pop up. Select **Key 0-**Report and select OK. This will cancel the culture 7. Finalize **STGM2**. Standard deviation rule violation box will pop up. Select **OK**. Preview instant report and save. 8. Refresh **GRM/MISC1of 2** worklist 9. If finished reading slides, ensure gram stains remaining on worklist have been prepared to be read at a later time 10.Gently blot excess oil from slide using paper towel or gauze and save slides for further evaluation on the slide tray

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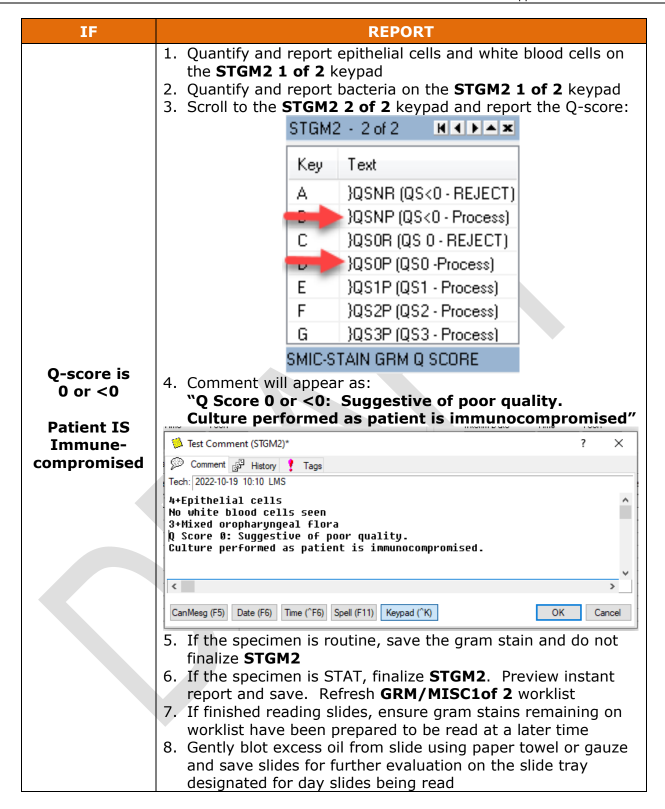
designated for day slides being read

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Title: MIC20300-Gram stain reporting in LIS-Respiratory Specimens

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Next Review Date:

Type: Laboratory Services Program SOP Policy Number:
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LIMITATIONS:

- 1. Use results of Gram stains in conjunction with other clinical and laboratory findings. Use additional procedures (e.g., inclusion of selective media, etc.) to confirm findings suggested by Gram-stained smears.
- Carefully adherence to procedure and interpretive criteria is required for accurate results. Accuracy is highly dependent on the training and skill of microscopists.
- 3. Gram stain-positive, culture negative specimens may be the result of contamination of reagents and other supplies, presence of antimicrobial agents, or failure of organisms to grow under usual culture conditions (medium, atmosphere, etc.).
- 4. False Gram stain results may be related to inadequately collected specimens or delays in transit.
- 5. Prior treatment with antimicrobial drugs may cause Gram-positive organisms to appear Gram-negative.

CROSS-REFERENCES:

- MIC10100-Microbiology Specimen Processing
- MIC60060-Microbiology Stain Quality Control

REFERENCES:

1. Leber, A. (2016). *Clinical microbiology procedures handbook.* (4thed.) Washington, D.C.: ASM Press

APPROVAL:	
Date	

REVISION HISTORY:

REVISION	DATE	Description of Change	REQUESTED BY
1.0	07 Feb 19	Initial Release	L. Steven
2.0	31 Mar 22	Procedure reviewed and added to NTHSSA policy template	L. Steven

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Policy Number: Date Approved: Page 8 of 8