Title: MIC32600-Eye Culture-Deep Issuing Authority: Director of Health Services Next Review Date:

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

PROGRAM Standard Operating Procedure – Laboratory Services			
Title: MIC32600 - Eye Culture-Deep	Policy Number:		
Program Name: Laboratory Services			
Applicable Domain: Lab, DI and Pharmacy Services			
Additional Domain(s):			
Effective Date:	Next Review Date:		
Issuing Authority:	Date Approved:		
Director of Health Services			
Accreditation Canada Applicable Standard: N/A			

GUIDING PRINCIPLE:

The inner eye structure consists of sclera, cornea, iris, lens, vitreous, retina, uvea, macula, and optic nerve. Early clinical and laboratory diagnosis of inner eye infections is paramount to the patient having a good outcome. Inner eye infections should be urgently identified and treated to prevent loss of visual acuity.

PURPOSE/RATIONALE:

This standard operating procedure describes how to determine the significance of growth in deep eye specimens.

SCOPE/APPLICABILITY:

This procedure applies to Medical Laboratory Technologists (MLTs) processing specimens for deep eye culture.

SAMPLE INFORMATION:

T	Swab	
Туре	Amie's with or with charcoal	
Source	 Canaliculitis: inflammation of the canaliculus Wound on external lacrimal duct or pus Dacryoadenitis/Dacryocystitis: infection of lacrimal glands External lacrimal duct or pus Bacterial keratitis: acute and chronic inflammation of the cornea Corneal scrapings collected by ophthalmologist Bacterial endophthalmitis: inflammation of the ocular cavities and intraocular tissue (uvea and retina) Aqueous and vitreous fluid collected by aspiration 	

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Stability	 If the sample is received in the laboratory and processed greater than 48 hours from collection: Add specimen quality comment: "Delayed transport may adversely affect pathogen recovery" 	
Storage Requirements	Room temperature	
Criteria for rejection	 Unlabeled/mislabelled swabs Specimen container label does not match patient identification on requisition Improperly collected, labeled, transported, or handled specimens should be processed. Waiver of responsibility form SCM40110 needs to be filled out by the responsible nurse 	

NOTE:

- Refer to MIC34100-Body Fluid Culture for intraocular fluid
- Refer tissue or biopsy specimens for culture to DynaLIFE

REAGENTS and/or MEDIA:

- Blood agar (BA), Chocolate agar (CHO), MacConkey agar (MAC), Brucella agar (BRU) and Thioglycollate broth (THIO)
- Identification reagents: catalase, oxidase, Staph latex test, Strep latex test, etc.

SUPPLIES:

- Disposable inoculation needles
- Microscope slides

- Anaerobic jar and pouch
- Wooden sticks

EQUIPMENT:

- Biosafety cabinet
- 35° ambient air and 35° CO₂ incubators
- Vitek 2 and supplies

SPECIAL SAFETY PRECAUTIONS:

Containment Level 2 facilities, equipment, and operational practices for work involving infectious or potentially 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

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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:

· Refer to Test Manual for reagent quality control procedures

PROCEDURE INSTRUCTIONS:

Step	Action
Proce	essing specimens for deep eye culture
1	 In the biosafety cabinet: Inoculate BA, CHO, MAC, BRU and THIO with the swab or specimen Ensure all surfaces of swab make contact with the agar Streak for isolated growth using a disposable inoculation needle Prepare smear by rolling the swab gently across the slide to avoid destruction of cellular elements and disruption of bacterial arrangements
2	 Incubate the media: Place BA and CHO in the CO₂ incubator Place MAC in the O₂ incubator Label THIO with day 2 date and day 5 date and place in the THIO rack in the O₂ incubator NOTE: If the clinical information provided indicates canaliculitis, dacryoadenitis/dacryocystitis or endophthalmitis, label broth with Day 10 date Place BRU in anaerobic jar with anaerobic pouch and indicator as soon as possible after inoculation. Label jar with day 2 date and place in the O₂ incubator NOTE: Anaerobes should not be exposed to air for 42 to 48 hours after inoculation
3	Allow smear to dry and perform gram stain. Gram stain must be read before culture plates. Refer to MIC20115-Gram stain procedure.
4	Interpret deep eye smears immediately. During the regular Microbiology lab hours of 08:00 to 20:00, turnaround time for these gram stains is <1 hour. Outside the regular Microbiology lab hours, Microbiology Technologist may be called in if ordering physician determines the stain must be read immediately.
5	Immediately phone positive deep eye gram stain results to ordering location and document in the LIS.

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Canaliculitis and Dacrocystitis/Dacroadenitis:

Probable pathogens	Comments
 Actinomyces spp. Pseudomonas aeruginosa Propionibacterium spp. Staphylococcus aureus Streptococcus pneumoniae 	Gram-stained smear can help determine the presence of <i>Actinomyces</i>

Bacterial Keratitis:

	2000114111014411101		
Probable pathogens	Comments		
Cornea trauma/ulcer: Candida albicans Haemophilus influenzae Moraxella spp. Neisseria gonorrhoeae Neisseria meningitidis *+ Nocardia spp. Pseudomonas aeruginosa Staphylococcus aureus Streptococcus pneumoniae Viridans group Streptococci Contact lens associated: Bacillus spp. *+ Enterobacteriaceae Pseudomonas aeruginosa	 Other primary pathogens include: Acanthamoeba, Fusarium spp., Mycobacterium Haemophilus parainfluenzae can cause conjunctivitis, corneal ulcers, and bacterial keratitis. Report if no other pathogens isolated Identify yeasts to the species level 		

Bacterial Endophthalmitis:

Probable pathogens	Comments
 Anaerobes Bacillus spp. Coagulase-negative staphylococci Gram-negative organisms Haemophilus influenzae Neisseria gonorrhoeae Neisseria meningitidis *+ Staphylococcus aureus Streptococcus pneumoniae Viridans group streptococci 	 Fungi, AFB and Nocardia species should be ruled out in chronic postsurgical and traumatic infection Viral cultures should be done, particularly for patients with trigeminal herpes zoster infection Blood cultures should be obtained Post-cataract surgery can result in chronic infection occurring months to years after surgery

^{*} Risk group 3 organism. If suspected, refer to Policy B-0160: "Specimens Containing Suspected Risk Group 3 Pathogens" for Primary Specimen Handling Flow Chart

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⁺ All work-up should be performed in the BSC

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INTERPRETATION OF RESULTS:

Step	Action
Inter	pretation of aerobic growth in deep eye specimens
1	Ensure growth on culture media correlates with gram stain results. If discordant results are found between the gram stain and growth: Re-examine smear and culture plates Check for anaerobic growth Re-incubate media to resolve Consider re-smearing or re-planting specimen
2	 Observe BA and CHO plates at 24 hours, 48 hours, and 72 hours Observe MAC plate at 24 hours and 48 hours
3	 If organism is a probable pathogen: Perform and report full identification Perform and report susceptibility testing as per ASTM
4	Contact Dyna <i>LIFE</i> microbiologist regarding any growth other than probable pathogens to determine suitable workup.

Step	Action	
Interpretation of anaerobic growth for deep eye specimens		
1	 Observe BRU and THIO after 48 hours Re-incubate BRU and THIO for an additional 72 hours If anaerobic growth is suspected, perform gram stain. If gram stain resembles growth on aerobic plates, further workup is not indicated. If growth does not resemble growth on aerobic plates, perform aerotolerance test. Refer to MIC53700-Aerotolerance Test NOTE: If specimen is from suspected canaliculitis, endophthalmitis or dacryoadenitis/dacryocystitis, re-incubate BRU and THIO for a total of 10 days 	
2	 If single morphology growing on anaerobic plates: If growth is same as aerobic growth: Re-incubate BRU and THIO for anaerobic growth If growth does not resemble growth on aerobic plates: Perform identification If organism is a probable pathogen: Report full identification Refer to DynaLIFE for susceptibility testing Contact DynaLIFE microbiologist regarding any growth other than probable pathogens to determine suitable workup. 	
3	 If multiple morphologies growing on anaerobic plates: Contact DynaLIFE microbiologist regarding any growth other than probable pathogens to determine suitable workup. 	

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REPORTING INSTRUCTIONS:

IF	REPORT	
No growth after 1 day	PRELIM: • Report: "No Growth After 1 Day" • Report: "Further report to follow"	
No aerobic growth at 3 days and no anaerobic growth	 INTERIM: Report: "No growth aerobically after 3 days" Report: "@Anaerobic Culture to follow" 	
Aerobic growth at 2 or 3 days and no anaerobic growth	 INTERIM: Report aerobic growth Report: "@Anaerobic culture to follow" 	
No anaerobic growth after 5 days and specimen source indicates 10-day incubation	FINAL: • Report: "No anaerobes isolated after 5 days" • Add test comment }AC10	
No growth on anaerobic media after 5 days	FINAL: • Report: "No anaerobes isolated after 5 days"	
Growth of pathogen	Report organism identificationList quantitationReport susceptibility results as per ASTM	
Neisseria gonorrhoeae isolated and gonorrhea culture was not ordered	 Add organism: "Neisseria gonorrhoeae" List quantification as: "Isolated" Report susceptibility results as per ASTM Add isolate comment &REF6 Refer isolate to APL for susceptibility testing Freeze isolate and log into stored isolates log 	

NOTE:

- Refer to Reportable Diseases Public Health Act as of September 2009 for reporting to OCPHO (HPU1)
- Refer to LOM70620-Laboratory Critical Results List-Microbiology for results that need to be phoned to ordering location
- Refer to MIC36100-Nosocomial Infection Notification Job Aid to determine if organism needs to be copied to Infection Prevention and Control
- Refer to MIC36200-Referral of Category A Specimens to APL for sending category A isolates to APL
- Refer to MIC36300-Referral of Category B Specimens to APL for sending isolates to APL
- Refer to MIC36400-Referral of Category B Specimens to DL for sending isolates to *Dyna*LIFE

LIMITATIONS:

- 1. False positive cultures can result from contamination of the specimen or plates with skin flora.
- 2. False negative results can occur if antimicrobial agents are given prior to collection of the specimen.

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CROSS-REFERENCES:

- LQM70620-Laboratory Critical Results List-Microbiology
- MIC20115-Gram stain procedure
- MIC33500-Gonorrhoeae Culture
- MIC34100-Body Fluid Culture for intraocular fluid
- MIC36100-Nosocomial Infection Notification Job Aid
- MIC36200-Referral of Category A Specimens to APL
- MIC36300-Referral of Category B Specimens to APL
- MIC36400-Referral of Category B Specimens to DL
- MIC53700-Aerotolerance Test

REFERENCES:

- 1. Leber, A. (2016). *Clinical microbiology procedures handbook.* (4thed.) Washington, D.C.: ASM Press
- Jorgensen J.H., Pfaller M.A., Carroll K.C., Funke G., Landry M.L., Richter S.S., Warnock D.W. (2015). *Manual of Clinical Microbiology*, 11th edition. Washington, D.C: ASM Press

APPROVAL:	
Date	

REVISION HISTORY:

REVISION	DATE	Description of Change	REQUESTED BY
1.0	28 May 18	Initial Release	L. Steven
2.0	26 Feb 21	Procedure reviewed and added to NTHSSA policy template	L. Steven
3.0	27 Feb 23	Procedure reviewed	L. Steven

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