TRAINING UPDATE

Lab Location: Department:

SGMC & WOMC Core Lab

Date Distributed:
Due Date:
Implementation:

11/3/2020 11/30/2020 **11/9/2020**

DESCRIPTION OF PROCEDURE REVISION

Name of procedure:

Specimen Processing for Microbiology SGAH.M04 v11

Plating Chart for Media by Source and Test Code AG.F191.5

Description of change(s):

SOP:

Section	Reason
5.3	Deleted section for environmental cultures (CHY no longer performs, order code XENVR has been inactivated in SQ)
	Note : the ordering process for water cultures has not changed, refer to Vista Sample Processing, Startup and Maintenance SOP

FORM: Deleted environmental culture details

The revised SOP and Form will be implemented on November 9, 2020

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

Non-Technical SOP

Title	Specimen Processing for Microbiology	
Prepared by	Ronald Master	Date: 4/14/2009
Owner	Ronald Master	Date: 4/14/2009

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

TABLE OF CONTENTS

1.	PURPOSE	2
2.	SCOPE	. 1
	RESPONSIBILITY	
4.	DEFINITIONS	. 1
5.	PROCEDURE	. 2
6.	RELATED DOCUMENTS	. 4
7.	REFERENCES	. 4
8.	REVISION HISTORY	. 4
9.	ADDENDA AND APPENDICES	. 4

1. PURPOSE

To describe the process for microbiology specimen setup, plating and management.

2. SCOPE

The scope of this SOP is to ensure the pre analytic processes for microbiology specimens are outlined. These procedures are imperative in determining what pathogenic organisms are present in specimens obtained from patients.

3. RESPONSIBILITY

It is the responsibility of all personnel assigned to Microbiology to read, understand and to perform all procedures as described in this SOP.

4. **DEFINITIONS**

Plating – inoculation of plated/tubes media with clinical specimen for microbiology culture.

Inoculation – to implant microorganisms or infectious material onto a culture medium.

Streaking – The use of a loop or other plating tool to inoculate a specimen in order to differentiate microorganisms by color or texture from its surroundings on a culture medium.

5. PROCEDURE

5.1 Routine Procedure for Plating Cultures:

All specimens are to be plated in a biosafety cabinet.

1. Media and its location:

All routine media will be stored in the refrigerator. Microbiology media should be kept in the refrigerator until needed. Media should be allowed to warm to room temperature before use. A working supply (minimum amount) is left at room temperature for use.

2. Loops, Swabs, and Pipettes:

- a. <u>Loops</u> A wire loop is used for streaking specimens, with the exception of urines. A 0.001 mL calibrated loop must be used to inoculate urine. For sterile urines (cystoscopy, suprapubic aspirate, etc.) use both 0.001 and 0.01 mL calibrated loops.
- b. <u>Swabs</u> Swabs are used in making the initial inoculation of plates, for preparing smears, and for inoculating specimens into broth media. If a specimen is submitted on a swab it must be submitted in a culturette containing holding medium to prevent drying out. A swab is convenient for inoculating certain specimens onto media, e.g., stool, sputum. Sterile swabs are available at the plating bench.
- c. <u>Pipettes</u> A sterile pipette may be used to inoculate liquid specimens into broth media, such as thioglycolate, and any plated media. A sterile pipette should be used to inoculate CSF and other body fluids and environmental cultures of liquids. To inoculate thioglycolate with a pipette, insert pipette to bottom of tube and slowly evacuate sample as you with draw the pipette.

3. Preparation of Smears and Gram Stains:

a. <u>Smears</u> - Write the accession number, specimen source, date and the patient's last name on the slide. Using a sterile loop or swab, make a smear about the size of a nickel near the center of the slide. Let the slide air dry, then heat fix. ALWAYS MAKE SMEAR AFTER INOCULATING MEDIA TO AVOID CONTAMINATING THE SPECIMEN.

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Page 2 of 7

4. Inoculation and Streaking of Media:

- a. The first process in the cultural examination of clinical specimens is the selection of appropriate isolation media. The plating table AG.F191 lists the media suitable for the isolation of microorganisms most commonly recovered from various clinical specimens. It is desirable to inoculate more than one kind of isolation medium unless a single organism is targeted.
- b. The purpose of isolation is to obtain bacterial colonies representing progeny of a single cell and thus provide the source of a pure culture. The streaking of materials onto the surface of the medium provides such results. The streaking method must be such that, (a) part of the medium is inoculated with a large amount of material, and (b) subsequent streaking will allow for growth of isolated colonies. When done properly, the completed streaking should cover essentially the entire surface of the medium. When streaking plates, flame the loop between the first and second streak area to avoid overly heavy growth or use a disposable loop. An illustration and explanation of an acceptable method is provided in Addendum A, Figure 1. Addendum A, Figure 2 illustrates the proper streaking procedure for a urine colony count.

5.2 Incubation of Plates:

Plates and broth media are incubated in a CO_2 incubator at 35 $\pm 2^{\circ}$ C except for chromogenic medium for MRSA which is incubated at 35-37°C in air (non- CO_2).

5.3 Specimens and Special Requirements:

5.3.1 Stool Cultures

- 1. If E. coli O157 is ordered, use test code XECOL.
- 2. Stool in transport media is to be sent to Chantilly. Stool specimens will be plated in Chantilly.

5.3.2 IV Catheter Tips

- 1. Perform all steps in a biological safety cabinet.
- 2. Using sterile forceps, remove catheter tip from transport tube.
- 3. Lay the catheter tip on a blood agar plate, and using sterile forceps, roll tip 4-5 times over entire plate. If the catheter tip is longer than 2 inches (5 cm), use sterile scissors or scalpel to cut the end closest to the top of the tube (proximal end) prior to rolling the distal end on the plate. The proximal end may be rolled on a second plate, if desired.
- 4. Leave the catheter tip on the plate, do not press it into the agar.

5.3.3 Cerebrospinal Fluid and Other Sterile Body Fluids

- 1. Sterile body fluids must be processed immediately upon receipt.
- 2. Include a chocolate agar plate for all sterile body fluids submitted for culture.
- 3. Include thioglycolate broth for CSF and synovial/joint fluids

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SOP version # 11 Page 3 of 7

6. RELATED DOCUMENTS

Plating Chart for Media by Source and Test Code (AG.F191) Sources for Anaerobic Culture Table (AG.F326)

7. REFERENCES

N/A

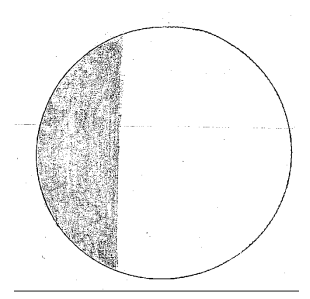
8. REVISION HISTORY

Version	Date	Reason for Revision	Revised By	Approved By
		Supersedes SOP M006.007		
000	5/26/09	Addenda D: media change for MRSA screen	L. Barrett	R. Master
001	3/10/10	Section 5: Change stool culture, add IV cath tip	R. Master	R. Master
002	5/17/10	Section 5.1: Delete requirement for date on opened	R. Master	R. Master
		media		
		5.2: Change temperature to $35 \pm 2^{\circ}$ C		
003	7/12/11	5.3.2 Specified catheter length	R. Master	R. Master
004	5/21/12	5.3.1 Deleted plated media for stool cultures	R. Master	R. Master
004	5/21/12	Figure 4: Updated stool cultures	R. Master	R. Master
005	4/9/13	5.2 Add exception for MRSA chromogenic medium	R. Master	R. Master
		5.3.3 Add centrifugation of sterile body fluids		
		5.3.4 Add environmental cultures		
006	4/27/15	5.3.3 Edited centrifugation of sterile body fluids	R. Master	R. Master
		Section 6: Moved plating chart from section 9,		
		added Anaerobic culture table		
		Footer: Version # leading zero's dropped due to		
		new EDCS in use as of 10/7/13		
7	3/21/17	Header: Add WAH	R. Master	R. Master
		5.1.4.a: Changed location of plating chart. Clarified		
		use of multiple media.		
8	3/14/18	Section 9: Added Figure 4: Z streak technique	R. Master	R. Master
9	12/4/19	Header: Changed WAH to WOMC	R. Master	R. Master
		5.3.4 Deleted dialysis and pharmacy cultures;		
		Added sending lab water to Chantilly		
		5.3.3 only need thio for CSF and synovial fluid		
10	11/2/20	5.3 Deleted environmental cultures	L Barrett	R Master

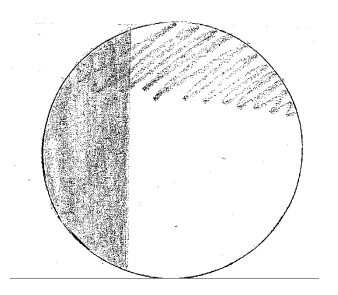
9. ADDENDA

- A. Figure 1 Acceptable Method of Plate Streaking
- B. Figure 2 Proper Streaking for a Urine Culture and Colony Count
- C. Figure 3 Proper Streaking for a Biplate
- D. Figure 4 Z-Streak Technique

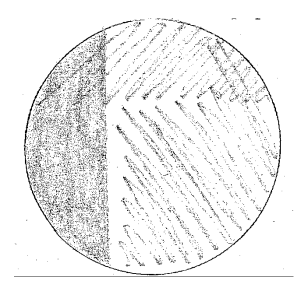
Figure 1 Acceptable Method of Plate Streaking



Step 1: Using a loop or a swab inoculate the specimen onto one edge of the plate, covering about one third of the plate.

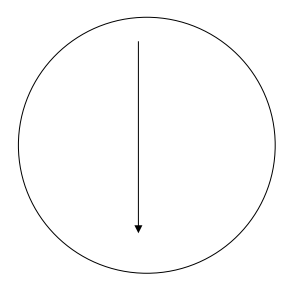


Step 2: Flame loop arid cool it by stabbing into the sterile agar. With cooled loop, streak at a right angle to the initial inoculum going back and forth

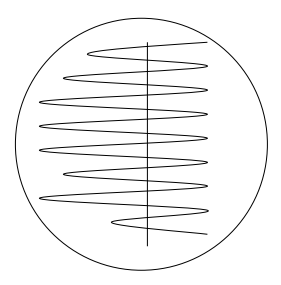


Step 3: Flame loop if specimen is likely to contain a lot of normal flora. Cool the loop. Rotate plate again, and entering only the isolation area, draw loop over the previously uninoculated portion of the plate. (Be careful to not streak over the initial inoculum.)

Figure 2 **Proper Streaking for a Urine Culture and Colony Count**

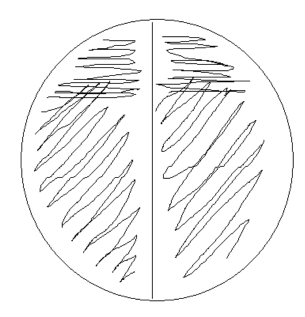


Step 1: Swirl urine to mix. Select sterile calibrated loop. Dip into the bottom of the urine sample and streak down the middle of the plate.



Step 2: Starting at the top, go back and forth numerous times over the initial streak line to facilitate the isolation of bacterial colonies.

Figure 3 **Proper Streaking of Bi-plates (non-urine specimens)**

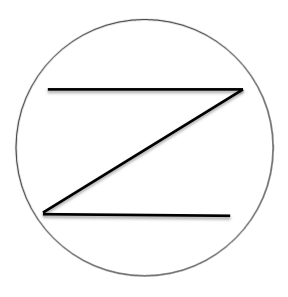


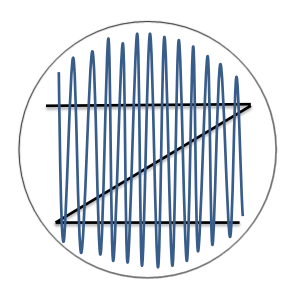
Biplates are inoculated by initially streaking about 15-20% of the plate, then flaming before performing the downward streak.

Figure 4 **Proper Z-streak Technique**

Step 1 Roll the swab over the plate in a "Z" pattern

Step2 Using a sterile loop, streak back and forth numerous times over the initial inoculum







PLATING CHART FOR MEDIA BY SOURCE AND TEST CODE

Culture	Test Code	Blood	MAC	СНОС	Martin/ Lewis	ParaPak C&S vial	CNA	Ana Blood	Thio	Gram Stain	V agar	Misc	Send specimen to Chantilly	Store On- Site
Stool Culture, Enteric Profile X														
Salmonella, Shigella,														
Campylobacter, Aeromonas,														
Plesiomonas,	15292					X						Send stool in transport to CHY	Yes	
Yersinia & Vibrio														
Stool, E. coli O157	4221					X						Send stool in transport to CHY	Yes	1 (1
Respiratory Culture	769	X	X	X			X						NO	1 month
Eye Culture	76950	X	X	X			X						NO	7 days
Ear Culture	76951	X	X	X			X						NO NO	7 days
Throat Culture	5870	X		X									NO NO	7 days
Group A Strep Only Culture	6470	X												7 days
Genital Culture	778		X	X			X				X		NO	7 days
GBS Screen	14537											Send swab to CHY	Yes	
GC Screen	657				X								NO	7 days
Urine Culture	775	X	X									0.001 mL loop	Yes	
Urine, sterile source	2983	X (0.01 +	X (0.01 +									0.001 mL and 0.01 mL loops	Yes	
		0.001)	0.001)											
CSF Culture	127350	X	X	X			X		X	X			NO	1 month
Sterile body Fluid Culture									Synovial					
	1273	X	X	X			X		or joint	X			NO	1 month
									fld only					
Wound Culture	783 / 78351	X	X	X			X						NO	7 days
Tissue Culture	78350	X	X	X			X		X	X			NO	1 month
Surgical Culture	78353	X	X	X			X		X				NO	1 month
Prosthetic Joint Culture	78355	X	X	X			X		X				NO	1 month
Anaerobic Culture	15871	X						X				ana BBE/LKV biplate	NO	7 days
VRE Screen	8557						X						NO	7 days
MDRO Screen	78354											Send swab to CHY	Yes	
Staph. aureus Nasal Screen	900942	X					X						NO	7 days
IV Catheter tip	78352	X											NO	7 days
Type 1 Lab water	4120											Send water to CHY	Yes	
Positive Blood Culture	18100 AER	X	X	X			X						NO	1 month
	18101ANA	X	X	X			X	X						