Prenatal Screening for Group B *Streptococcus* (Non-pigmented broth, serologic or molecular testing for identification)

Annual Review:

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Revision History:

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Approval Signatures:

Approved by: _______________________________  Date: _________________

Author

____________________________________________________________

Print Name and Title

Approved by: _______________________________  Date: _________________

Supervisor/Technical Specialist-Director

____________________________________________________________

Print Name and Title

Approved by: _______________________________  Date: _________________

Quality Manager

____________________________________________________________

Print Name and Title
Purpose/Principle:

The purpose of this document is to provide instructions for performing prenatal screening for Group B *Streptococcus* from appropriately collected vaginal/rectal specimens using non-pigmented broth enrichment with serologic or molecular testing for identification.

Scope:

This document applies to laboratory assistants, technicians and technologists within the Division of Microbiology at XXXX.

Related documents:

1. Provider procedure for collection of specimens for Prenatal Screening for Group B *Streptococcus* - # xxxx. Note: Procedure should specify the acceptable specimen is a vaginal-rectal swab and that swabs from other sites (e.g. cervical or perianal swabs only) are not acceptable. Procedure should list acceptable type of swab and transport medium.
2. Quality Control procedure for tube and plated media - # xxxx.
3. Procedure for Group B *Streptococcus* serologic testing from broth culture - # xxxx.
4. Procedure for Group B *Streptococcus* molecular testing from broth culture - # xxxx.
5. Procedure for the identification of streptococci - # xxxx.
6. Notification procedure for positive Prenatal Screening for Group B *Streptococcus* - # xxxx.
7. Procedure for susceptibility testing of Group B *Streptococcus* - # xxxx.
Responsibility:

1. It is the responsibility of the technical staff of the microbiology laboratory to review and assure they are familiar with the processes outlined in this document which are within their job duties.

2. It is the responsibility of the technical specialist to keep this procedure updated and current with national standards/guidelines/recommendations.

3. It is the responsibility of the microbiology supervisor to accurately train and assess the competency of all microbiology staff involved in performing this procedure.

4. It is the responsibility of the microbiology data technologist to assure that reporting of all results is appropriate in all computer systems.

5. It is the responsibility of the technical director to approve this procedure on an annual basis.

Definitions:

N/A

Equipment/Materials:

1. Trans-Vag broth supplemented with 5% defibrinated sheep blood, LIM broth (or equivalent)

2. Agar plates for subculture for susceptibility testing (e.g., tryptic soy agar with 5% defibrinated sheep blood, or equivalent)

3. Sterile loops

4. Incinerator
5. Incubator

6. Serologic instrumentation and/or reagents

7. Molecular instrumentation and/or reagents

Quality Control:

Follow Quality Control procedures for tube and plated media - # xxxx.

Follow Quality Control procedures for serologic testing for Group B *Streptococcus* - # xxxx.

Follow Quality Control procedures for molecular testing for Group B *Streptococcus* - # xxxx.

Safety Precautions:

Use standard precautions when performing this procedure.

Procedure: (Appendix includes one-page flowchart)

1. Remove patient specimen (swab; swabs) from transport medium. Specimens should be processed within 4 days of collection. Specimens requiring more than 24 hr of transport time should be refrigerated until received and processed by the laboratory.

2. Inoculate swab(s) into a recommended selective enrichment broth medium, such as Trans-Vag broth supplemented with 5% defibrinated sheep blood, LIM broth or equivalent. [Before inoculating into broth, laboratories may choose to direct plate by rolling the swab(s) on a sheep blood agar plate, NNA agar, CNA agar, other selective *Streptococcus* agar. If Group B *Streptococcus* is identified from the primary plate, report Group B streptococcus positive on the culture report. If Group B *Streptococcus* is not identified from the primary plate, plate should be discarded, and]
enriched broth processed as per protocol below and results reported out from enriched broth].

3. Incubate the inoculated broth for a minimum of 18-24 hours at 35-37°C in ambient air or 5% CO₂ or for the length of time and conditions recommended by the manufacturer. The broth should display visible turbidity as an indicator of adequate specimen collection. A clear broth after 24 hours of incubation indicates failure to obtain a proper sample and a new specimen should be requested.

4. Perform serologic testing or molecular testing on the incubated broth for Group B Streptococcus.

5. Report as negative for Group B Streptococcus when no Group B Streptococcus has been identified by serologic testing or molecular testing.

6. If Group B Streptococcus is identified by serologic testing or molecular testing, report Group B Streptococcus positive on the report.

7. If the positive specimen is from a penicillin-allergic patient at high risk for anaphylaxis, subculture the incubated broth to a sheep blood agar plate (e.g., tryptic soy agar with 5% defibrinated sheep blood, or equivalent) and incubate for a minimum of 18-24 hour at 35-37°C in 5% CO₂.
   a. Inspect and identify colonies characteristic of Group B Streptococcus (i.e., gram-positive cocci, catalase negative, narrow zone of β-hemolysis or no hemolysis) using routine laboratory identification protocols. (Refer to procedure # xxxx – Procedure for the identification of streptococci).
   b. Perform susceptibilities for clindamycin and erythromycin, including inducible clindamycin resistance, using appropriate laboratory susceptibility
Experts from the American Society for Microbiology (ASM) and the Centers for Disease Control and Prevention’s (CDC) *Streptococcus* Laboratory co-authored this document.*  

For more GBS laboratory resources, visit [http://www.cdc.gov/groupbstrep](http://www.cdc.gov/groupbstrep)

*Collaboration does not imply HHS/CDC endorsement of ASM*
Flowchart #2: Using non-pigmented broth and serologic or molecular testing for identification of GBS

1. **Vaginal-rectal swab**
   - Place in non-pigmented enrichment broth
   - Incubate in enrichment broth x 18-24 hrs at 35-37°C

2. Latex agglutination, DNA probe, Nucleic Acid Amplification Test (NAAT), or other molecular testing

3. **Identify GBS**
   - GBS+
   - GBS−

4. **Report as GBS+**
5. **Report as GBS−**

*Without enrichment, the false negative rate can be as high as 50%*

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**Remember**
- Giving PCN-allergic women the most effective drug depends on YOU!
- Test appropriate samples for susceptibility to clindamycin and erythromycin, INCLUDING inducible clindamycin resistance.

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If PCN-allergic & at high risk for anaphylaxis, subculture to plate and incubate 18-24 hrs at 35-37°C to conduct susceptibility testing.

For more information, see: [http://www.cdc.gov/groupbstrep/lab](http://www.cdc.gov/groupbstrep/lab)