New York State Dept of Health
Bacteriology Proficiency Testing Program
Instruction Sheet – Comprehensive
September 6, 2016

Samples for the Bacteriology – Comprehensive proficiency test were shipped on September 6, 2016. A minimum score of 80% is required to pass each test event. Failure to achieve a score of 80% on two of three consecutive testing events is considered unsuccessful performance.

Upon receipt and prior to testing, store the specimen in its original sealed pouch at 2°C to 8°C. Refer to the instruction sheet for processing the KWIK-STIK units.

These samples contain pathogenic microorganisms. Protect yourself by following your laboratory’s safety for proper handling. If any of the specimens are damaged in transit autoclave the entire set and inform our office at once. If there is any problem with the specimens, you MUST inform our office before close of business on September 13, 2016 by calling (518) 474-4177.

Proficiency testing specimens must be processed in the same manner as patient specimens.

1. Specimen content
   Below is a list of all specimens for this test event. Depending on your laboratory’s protocol, your shipment may or may not include a specimen for Group A Streptococcus or Chlamydia direct testing. Do not consider quantity or predominance of an organism when reporting. In order to accommodate as many laboratory protocols as possible, the sources listed below are very general. Please note: if, at any time (such as a physician's special request), you report a particular organism to the species or serogroup level, you must report this organism to the same level on the proficiency test. Likewise, if you ever process a particular specimen type, then you must process that specimen type on the proficiency test. Your questionnaire will be used to determine your laboratory’s level of identification.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Report</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Pathogens¹ only</td>
<td>Stool²</td>
</tr>
<tr>
<td>2</td>
<td>Pathogens only</td>
<td>Tracheal aspirate</td>
</tr>
<tr>
<td>3</td>
<td>All organisms</td>
<td>Intra-abdominal abscess</td>
</tr>
<tr>
<td></td>
<td>Aerobic / Anaerobic</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>All organisms &amp; Antibiotic susceptibility</td>
<td>Blood</td>
</tr>
<tr>
<td>5</td>
<td>Pathogens only</td>
<td>Wound</td>
</tr>
<tr>
<td>Chlamydia Direct Detection</td>
<td>Pos/Neg for Chlamydia</td>
<td>Cervix/Urine for C. trachomatis</td>
</tr>
<tr>
<td>Group A Streptococcus Direct Detection</td>
<td>Pos/Neg for Group A Streptococcus</td>
<td>Throat for Group A Streptococcus</td>
</tr>
</tbody>
</table>

¹ For proficiency testing purposes, an organism should be reported as a pathogen if it is generally considered by the medical community to cause disease when recovered from the specimen source indicated.

² Rule out ALL enteric pathogens according to your laboratory’s protocol including those organisms requiring a special request.

2. Antibiotic susceptibility –
   - Antibiotic susceptibility testing must be performed on the reportable isolate(s) in SPECIMEN NUMBER 4. Report your interpretations of susceptibility to penicillin and vancomycin. Zone sizes or MICs may be reported, but they must be accompanied by an interpretation. If your laboratory does not test these antibiotics, please indicate this. Do not report results for antibiotics other than those listed above.
3. Specimen preparation / reconstitution

Please note that your laboratory may not have received the *Chlamydia* &/or Group A Strep direct detection samples. Specimens are sent based on individual laboratory protocol. If you have any questions regarding the types of specimens your laboratory should be receiving, please contact us.

**Specimen reconstitution:**
- Refer to the instruction sheet for processing the KWIK-STIK units. These specimens are suitable for culture and for direct testing.
- After pinching the Kwik Stik tube to crush the ampoule of liquid you may need to wait a few minutes for the liquid to flow into the pellet area. This can be facilitated by tapping the tube on a hard surface. If a sufficient quantity of liquid does not flow into the pellet area you can add a few drops of sterile broth/saline to sufficiently re-hydrate the pellet.

**Antigen detection specimen (specimen A) – simulated throat swab:**
- Allow the specimen to reach room temperature before processing.
- Open and test for Group A *Streptococcus* using a direct detection (or rapid Strep) test system.
- Please note that some kits use very small amounts of liquid for the extraction process and the test swab may absorb all extraction liquid. If this occurs, add 3-5 drops sterile water to the extraction tube and continue with test procedure.

**Direct detection *Chlamydia* specimen – cervical/urine specimen**
- Process this sample following your kit's instructions. It can be processed as either a swab specimen in transport media or a urine specimen. We have increased the volume of this sample to 2 ml so that it can be processed as a urine sample.

**IMPORTANT:** Survey specimens, their progeny, unmodified derivatives or modifications thereof may not be transferred or incorporated into a product intended for Sale. Survey specimens, their progeny, unmodified derivatives or modifications thereof, reagents, and disposable equipment used in proficiency testing, when disposed of, should be autoclaved or incinerated and disposed of as hazardous waste.

4. Reporting of results

Report all results as you would for a patient specimen. Your report will be graded according to the Bacteriology Questionnaire which we have on file for your laboratory, including any changes or amendments submitted prior to the mailing date of this test. Your laboratory will only be graded on the test specimens which you evaluate and will not receive credit for those specimens which you do not process. All answers submitted will be graded. Credit will be deducted for conflicting reports such as, 'No pathogens reported/ found' along with a list of organisms identified. Conditional answers, such as 'Corynebacterium species, possibly xerosis' will be considered reported: in this instance, as if *Corynebacterium xerosis* was reported.

Credit will be deducted if additional, nonpathogenic organisms are reported in the specimens where only the pathogen is to be identified.

Please utilize the drop-down list of organisms if at all possible. If you need to enter a result that is not on the list, please select “Other” and type in your result. All fields must be completed. Do not enter an organism more than one time in order to enter multiple methods of identification. This causes the program to grade EACH additional entry as an extra INCORRECT organism, possibly resulting in a loss of credit. Please choose ONE system/kit as your primary method of identification and enter each organism only once.

5. Submission of results

- The deadline for submission of results is **September 27, 2016**.
- **Electronic reporting:** Do not submit paper results. Specific questions regarding electronic reporting should be addressed to the Clinical Laboratory Evaluation Program at (518) 485-5378 or by emailing clepeptrs@health.ny.gov

If you have any questions regarding the Bacteriology Proficiency Testing Program, please contact Dr. Wendy Archinal at (518) 474-4177 or BactiPTP@health.ny.gov.