STUDENT STERILE TECHNIQUE PRACTICE Module 3 Supplement

ONLY FOR PRACTICE you should

Re-use the loop and pipette and re-use their "sterile" wrappings Do not dispose of a culture dish even if you accidentally contaminate it.

If you were really doing a lab with bacteria you would

- Never reuse any loop or pipette.
- Always dispose of used loops and pipettes in the "biological trash beaker."
- Throw away any culture dish or other item that is accidentally contaminated, and ask for a replacement.
- Wash hands immediately if they contact any liquids containing bacteria.

Each member of the team should practice each of the following:

- 1. **Open, hold, and close** a sterile 15 ml culture tube.
- 2. **Open and close** a sterile 1.5 ml test tube.
- 3. **Open** a culture dish, observe it without breathing on it, and close it.

After each team member has practiced 1-3, use (practice) sterile technique as you

- 4. Transfer 250 I of liquid from the 1.5 ml test tube to the culture tube.
- 5. Pretend to **transfer** a colony from the surface of the culture dish to the liquid in the culture tube.
- 6. Use the pipette to **suspend** the imaginary bacteria in the culture tube liquid.

After each team member has practiced 4-6, use (practice) sterile technique as you

7. Transfer 100 I of culture tube contents to culture dish and spread it.

POSSIBLE SEQUENCE OF STEPS FOR STERILE TECHNIQUE DEMONSTRATION/PRACTICE

Delete parts as time requires

- 1. a. Teacher demonstrates opening/holding/closing of sterile 1.5 ml test tube, sterile 15 ml culture tube, sterile pipettes, measuring with pipettes.
 - b. Students practice with test tubes, caps, droppers. (opening, closing, transferring)
- 2. a. Teacher demonstrates cleaning lab bench, opening/closing of petri dish, how not-to-breathe-on an open plate.
 - b. Teacher passes around sample culture dish for touching and looking.
 - c. Teacher demonstrates labeling, opening inoculating loop, using loop, disposition of loop and other trash.
 - d. Students practice cleaning lab bench, label/open/close plate.
- 3. a. Teacher demonstrates sterilization of spreader with ethanol and flame, spreading sample on medium, closing, taping, waiting, then inverting.
 - b. Students practice spreading, waiting, taping, etc.

Transfer technique for new starter culture plates -Transformation Lab

Streaking is best done with a wire loop. Isolated colonies should occur in the section of the plate streaked last.

- Step (1)-Streak the inoculum over the surface of the plate. Start near the edge of the plate.
- Step (2) Flame the loop to get rid of the original bacteria. Cool the loop by gently touching a sterile portion of the agar.
- Step (3)-Rotate the agar plate 1/4 of a turn. Streak across the agar by transferring bacteria from the original streak(A).
- Step (4)-Rotate agar plate again 1/4 of a turn. Flame and cool the loop. Streak again by transferring bacteria from section "B".
- Step (5)-At this step do not flame the loop. Just rotate the plate and streak again as in step (4) this time by carrying bacteria from section C

