pGLO Transformation Procedure

1. Label one closed micro test tube +pGLO and another -pGLO. Label both tubes with your group’s name. Place them in the foam tube rack.

2. Open the tubes and using a sterile transfer pipet, transfer 250 µl of transformation solution (CaCl₂). CaCl₂ punches holes in the cell membrane, allowing small DNA molecules like plasmids to slip through into the interior.

3. Place the tubes on ice.

4. Use a sterile loop to pick up a single colony of bacteria from your starter plate. Pick up the +pGLO tube and immerse the loop into the transformation solution at the bottom of the tube. Spin the loop between your index finger and thumb until the entire colony is dispersed in the transformation solution (with no floating chunks). Place the tube back in the tube rack in the ice. Using a new sterile loop, repeat for the -pGLO tube.

5. Examine the pGLO plasmid DNA solution with the UV lamp. Note your observations. Immerse a new sterile loop into the plasmid DNA stock tube. Withdraw a loopful. There should be a film of plasmid solution across the ring. This is similar to seeing a soapy film across a ring for blowing soap bubbles. Mix the loopful into the cell suspension of the +pGLO tube. Close the tube and return it to the rack on ice. Also close the -pGLO tube. Do not add plasmid DNA to the -pGLO tube. Why not?
6. Incubate the tubes on ice for 10 minutes. Make sure to push the tubes all the way down in the rack so the bottom of the tubes stick out and make contact with the ice.

7. While the tubes are sitting on ice, label your four agar plates on the bottom (not the lid) as follows:
   - Label one LB/amp plate: +pGLO;
   - Label the LB/amp/ara plate: +pGLO;
   - Label the other LB/amp plate: -pGLO;
   - Label the LB plate: -pGLO.

8. Heat shock. Heat shocking renders the cell membranes even more permeable, allowing the plasmid to penetrate the bacteria as thoroughly as possible. Using the foam rack as a holder, transfer both the (+) pGLO and (-) pGLO tubes into the water bath, set at 42 °C, for exactly 50 seconds. Make sure to push the tubes all the way down in the rack so the bottom of the tubes stick out and make contact with the warm water. When the 50 seconds are done, place both tubes back on ice. For the best transformation results, the change from the ice (0°C) to 42°C and then back to the ice must be rapid. Incubate tubes on ice for 2 minutes.

9. Remove the rack containing the tubes from the ice and place on the bench top. Open a tube and, using a new sterile pipet, add 250 µl of LB nutrient broth to the tube and reclose it. LB revitalizes the bacteria after the CaCl₂ and heat shocking with its influx of nutrients. Repeat with a new sterile pipet for the other tube. Incubate the tubes for 10 minutes at room temperature.
10. Tap the closed tubes with your finger to mix. Using a **new** sterile pipet for each tube, pipet 100 µl of the transformation and control suspensions onto the appropriate plates.

11. Use a new sterile loop for each plate. Spread the suspensions evenly around the surface of the agar by quickly skating the flat surface of a new sterile loop back and forth across the plate surface.

12. Stack up your plates and tape them together. Put your group name and class period on the bottom of the stack and place the stack upside down in the 37°C incubator until the next day.