Amplicon Visualization and Quantification Protocols

Gel Electrophoresis

- 1. Prepare a 1% agarose gel by combining 1g of agarose with 100ml 1xTBE in an 250ml Erlenmeyer flask (add agarose first then liquid).
- 2. Microwave the mixture for ~2m (time is dependent on microwave) or until the mixture is clear. Note: During this step watch the mixture as it microwaves, when it begins to bubble pause the microwave, allow the bubbling to stop, and carefully mix by gently swirling the flask. Use hot hands or an autoclave mit and be very careful to ensure that the mixture does not boil over.
- 3. Allow mixture to cool for several minutes. It is cool enough when, while using a nitrile-gloved hand, you can hold your hand to the flask for 5s.
- 4. Pour gel into gel cassette. Be sure to add the comb to the gel.
- 5. Allow to cool at 4°C for ~15m (The gel should be firm at this point).
- 6. Combine 5ul of each PCR product with 5ul of ddH2O and 2ul of loading dye.
- 7. Prepare 100bp ladder by combining 1ul ladder, 2ul loading dye, 9ul ddH20
- 8. Place solidified gel into gel box in the correct orientation (DNA is negatively charged and will migrate toward the red positive post).
- 9. Fill gel box to fill line with 1xTBE.
- 10. Load gel starting with prepared 100bp ladder in the first lane.
- 11. Load PCR samples.
- 12. Connect the gel box to the power supply.
- 13. Run gel at 80v for ~45m.
- 14. Turn off power and disconnect the gel box.
- 15. Remove gel and place in ethidium bromide stain for 20m.
- 16. Remove gel (must be wearing gloves) and visualize.

 Note: Expected band size for 515f/806r is roughly 300 350 bp.

DNA Quantification with qubit (adapted from manufacturer's protocol*).

	Standard Assay tubes	User Sample Assay tubes
Volume of Working solution (see step 2 below)	190µL	199
Volume of DNA Standard	10μL	-
Volume user DNA	-	1µL
Total Volume	200μL	200μL

- 1. Set up two tubes for the DNA standards
- 2. Prepare the **Working Solution** by diluting the Qubit reagent 1:200 in Qubit buffer.
- 3. Prepare tubes as above for each pooled PCR reaction.
- 4. Vortex all tubes 2-3 seconds.
- 5. Incubate for 2m at room temp in the dark.
- 6. Insert tubes into Qubit and press DNA, then hsDNA
- 7. Read standards if appropriate (standards should be read at the very minimum weekly, but preferably daily)
- 8. Read sample
- 9. Calculate the ng/ul using the Qubit calculate function after reading each sample (note ng/ul must be selected from the drop down screen).

^{*}www.invitrogen.com/qubit