

Protocol for Brain Tissue Homogenization in the Bullet Blender®

The protocol described in this document is for the use of the Bullet Blender® for the homogenization of brain tissue (from a variety of animals). Note that the time and speed settings may differ due to the variation in consistency/texture of tissue from species to species. This protocol does not specify a particular buffer - you may choose which is most appropriate for your downstream application (nucleic acid isolation, protein extraction, etc.).

Materials Required: brain tissue, Bullet Blender[®], homogenization buffer, pipettor,

microcentrifuge tubes and Red bead lysis kit/Pink bead lysis kit/0.5 mm glass beads (part number GB05) or denser.

Instructions

1. Cut brain tissue into appropriately sized pieces for analysis (10mg-300mg).

- 2. **OPTIONAL:** Wash tissue 3x with ~1mL PBS. Aspirate. **NOTE:** This step removes external contaminants (blood, etc.).
- 3. a. Samples 100mg or greater

Place the sample in Red bead lysis kit tube.

- b. Samples less than 100mg
- Place the sample in Pink bead lysis kit tube.
- c. Alternate protocol step for bulk beads Place sample in microcentrifuge tube and add beads to the tube. Use a volume of beads equal to the mass of tissue. **NOTE:** $100 \text{mg} \cong 100 \mu\text{L}$.
- 4. Add 0.025mL to 0.6mL buffer (2 volumes of buffer for every mass of tissue).
- 5. Close the microcentrifuge tubes.
- 6. Place tubes into the Bullet Blender®.
- 7. Set controls for **SPEED 8** and **TIME 3** minutes. Press **Start**.
- 8. After the run, remove the tubes from the instrument.
- 9. Visually inspect samples. If homogenization is unsatisfactory, run for another two minutes at the **SPEED 9.**
- 10. Proceed with your downstream application.

SAFETY NOTE!!!

When using a centrifuge to separate your homogenate from the debris and beads, make sure your tubes are balanced!

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