Homogenization in the Bullet Blender[®] 50 Protocol for Mammalian Cell Culture

The protocol described in this document is for the use of the Bullet Blender[®] 50 for the homogenization of mammalian cell cultures. 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:

mammalian cells, Bullet Blender[®] 50, homogenization buffer, pipettor, 50mL skirted centrifuge tubes (Axygen[®] or Corning[®] brand), 0.15mm zirconium oxide beads (part number ZROB015) or 0.1mm glass beads (part number GB01).

Instructions

- 1. Detach cells from culture dish or flask by your chosen method (trypsinization, scraping, spontaneous detachment, etc.)
- 2. Wash cells from dish with PBS into 50ml tube
- **3.** Centrifuge cell suspension to yield a cell pellet (200-500g for five minutes at 0°C).
- **4.** Completely aspirate the supernatant liquid. Place tube on ice.
- **5.** Inspect the volume of the pellet. It should be 4mL or less to achieve efficient homogenization.
- **6.** Completely aspirate supernatant liquid. Place tube on ice.
- **7.** Add a volume of beads to the tube approximately equal to the volume of the pellet.
- **8.** Add 0.2 mL to 8mL buffer (2 volumes of buffer for every volume of sample).
- 9. Screw caps onto centrifuge tubes **TIGHTLY**.
- 10. Place tubes into the Bullet Blender[®] 50.
- 11. Set controls for SPEED 8 and TIME 12 minutes.
- **12.** Remove tubes from the instrument.
- **13.** Visually inspect samples, if homogenization is unsatisfactory, run for another six minutes at the **SPEED 9**.
- **14.** 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.

Date 05/06/2011



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