Protocol for Use of the Bullet Blender™ in Tandem with the QuickGene-Mini8O-Cultured Cells

The protocol described in this document is for the use of the Bullet Blender^m in tandem with the Fuji QuickGene-Mini80 RNA isolation system for cultured mammalian cells. Cells will first be homogenized using the Bullet Blender^m with the buffers provided in the QuickGene kit and biological sample provided by the researcher.

Materials Required:

Cultured Cells, Bullet Blender[™], QuickGene kit, pipetor, microcentrifuge tubes, and <u>0.15mm zirconium oxide beads (part</u> <u>number ZrOB015-RNA) or 0.1mm glass beads (GB01-RNA)</u>

Instructions

- 1. Detach cells from culture dish or flask by your chosen method (trypsinization, scraping, spontaneous detachment, etc.).
- **2.** Transport the appropriate number of cells according to the QuickGene protocol into a microcentrifuge tube.
- 3. Centrifuge cell suspension for 5 min at 300Xg to yield a cell pellet.
- **4.** Completely aspirate the supernatant liquid. Loosen pellet by flicking tube. Place tube on ice.
- 5. Inspect the volume of the pellet. It should be 300μ L or less.
- **6.** Add a volume of zirconium oxide beads (0.15mm) **OR** glass beads (0.1mm) to the tube equal to the volume of the pellet. One scoop of beads \approx 50µL.
- **7.** Add 2 volumes of LRP buffer (from QuickGene kit, with 2-mercaptoethanol) for every volume of cells.
- **8.** Close the microcentrifuge tubes.
- 9. Place tubes into the Bullet Blender[™].
- 10. Set controls for SPEED 5 and TIME 2 to 3 minutes. Press Start.
- **11.** After the run, remove tubes from the instrument. Visually inspect samples. If homogenization is unsatisfactory, run for another two minutes at **SPEED 5**.
- **12.** Prior to removing samples from the Bullet Blender, open tube cap, then add LRP to reach 350 μ l if you are using QuickGene Protocol A, 600 μ l if you are using QuickGene Protocol B, or 800 μ l if you are using QuickGene Protocol B'. Close cap. Run Bullet Blender at **SPEED 2** for **1 minute** to thoroughly mix.
- **13.** Proceed with QuickGene protocol (i.e. SRP addition and flash spin).

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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