Nematode Homogenization Using the Bullet Blender

RS18-0238B.ANEM

Materials

- Bullet Blender® for 1.5/2 mL tubes using 5 mL adapters
- Homogenization Buffer
- FoamBlocker (Optional)
- Lysis Kit or Lysis Beads
 - PINK or RED Lysis Kit (from <u>PrecisionPak™</u> or purchased separately)
 - 0.5 mm Zirconium Oxide Beads in Eppendorf, GATOR, or RINO tubes
- Sample up to 300 mg

Table 1. Proper sample, bead and buffer volume ratios for 1.5/2 mL tubes using 5 mL adapters.

Lysis Kit and Bead Choices	Sample Volume	Bead Volume	Buffer Volume
PINK	Up to 100 mg	Pre-filled	200 - 300 μL
RED	100 - 300 mg	Pre-filled	300 - 600 μL
0.5 mm Zirconium Oxide Beads	Up to 300 mg	100 - 200 μL	200 - 600 μL

Procedure

- 1. Use the pre-filled bead lysis kit tubes OR prepare a tube with the recommended volume of bead choices from the table above.
- 2. Prepare the sample by pelleting the sample in a tube, resuspending it in appropriate amount of lysis buffer and then transfer it into the lysis tubes.
- 3. (Optional) To avoid excess foaming, add FoamBlocker up to 1-2% of the total volume of the homogenization buffer.
- 4. Close the tubes tightly and place into the Bullet Blender sample chamber. If using the Gold or Gold⁺ models, pre-cool the chamber before adding sample tubes.
- 5. Set the controls to speed 8, time 4 minutes then press Start.

 Note: Using single-size beads instead of pre-filled lysis kits may require additional time.
- 6. After the run, remove the tubes from the instrument and visually inspect the samples. If homogenization is incomplete, homogenize for an additional 30 seconds, or repeat the homogenization step with a higher speed.
- 7. Using a pipette, transfer the homogenized samples into new tubes.
- 8. Proceed with downstream application.

Notes

This protocol does not specify a particular buffer – choose a buffer that is most appropriate for the downstream application or use the lysis buffer provided in a <u>PrecisionPak™</u>, a simplified workflow solution which also includes a bead lysis kit, supplemental reagents for high quality nucleic acids isolations, and an optimized protocol for specific samples.

The provided homogenization conditions serve as a general guideline. Homogenization times, speeds, or beads may need to be optimized based on sample characteristics and desired outcomes.

