Total RNA from Adult Male *Brugia pahangi*, Strain FR3

Catalog No. NR-42500

This reagent is the tangible property of the U.S. Government.

For research use only. Not for human use.

Contributor:
Steven A. Williams, Director of Filariasis Research Reagent Resource Center and Gates Professor, Department of Biological Science, Smith College, Northampton, Massachusetts, USA

Manufacturer:
Filariasis Research Reagent Resource Center supported by Contract HHSN272201000030I, NIH-NIAID Animal Models of Infectious Disease Program

Product Description:
NR-42500 is a preparation of total RNA extracted from adult male *Brugia pahangi* (B. pahangi), strain FR3. *B. pahangi*, strain FR3 was originally obtained from researchers in Malaysia by Dr. John Schacher.1,2

*B. pahangi* is a thread-like filarial nematode with a life cycle consisting of a mosquito intermediate host and a wide variety of carnivorous definitive hosts including human and felines.1,3 Mosquitos deposit infective third stage larvae (L3) on human skin. The larvae then penetrate and migrate to the lymphatic vessels where they develop into adult worms over several months. Development includes molting transitions into fourth stage larvae (L4) and juvenile adults to reach maturation. The matured female worms release large numbers of microfilariae into the host bloodstream. The microfilariae are ingested by a mosquito during a blood meal and penetrate the midgut and develop over a period of 10 to 14 days to L3.4,5 L3 are developmentally arrested in the mosquito. The process repeats when the mosquito's proboscis penetrates the definitive host's skin.4

Material Provided:
Each vial of NR-42500 contains 0.5 µg to 2.0 µg of DNase-treated RNA in TE buffer (1 mM Tris-HCl, 0.1 mM EDTA, pH ~ 8). The concentration is shown on the Certificate of Analysis. The vial should be centrifuged prior to opening.

Packaging/Storage:
NR-42500 was packaged in RNase/DNase-free plastic vials. The product is provided frozen and should be stored at -80°C or colder upon arrival. Freeze-thaw cycles should be minimized.

Citation:
Acknowledgment for publications should read "The following reagent was provided by the NIH/NIAID Filariasis Research Reagent Resource Center for distribution by BEI Resources, NIAID, NIH: Total RNA from Adult Male *Brugia pahangi*, Strain FR3, NR-42500."

Biosafety Level: 1


Disclaimers:
You are authorized to use this product for research use only. It is not intended for human use.

Use of this product is subject to the terms and conditions of the BEI Resources Material Transfer Agreement (MTA). The MTA is available on our Web site at www.beiresources.org.

While BEI Resources uses reasonable efforts to include accurate and up-to-date information on this product sheet, neither ATCC® nor the U.S. Government makes any warranties or representations as to its accuracy. Citations from scientific literature and patents are provided for informational purposes only. Neither ATCC® nor the U.S. Government warrants that such information has been confirmed to be accurate.

This product is sent with the condition that you are responsible for its safe storage, handling, use and disposal. ATCC® and the U.S. Government are not liable for any damages or injuries arising from the misidentification or misrepresentation of products.

Use Restrictions:
This material is distributed for internal research, non-commercial purposes only. This material, its product or its derivatives may not be distributed to third parties. Except as performed under a U.S. Government contract, individuals contemplating commercial use of the material, its products or its derivatives must contact the contributor to determine if a license is required. U.S. Government contractors may need a license before first commercial sale.

References:
Appendix I: Total RNA Isolation from Filarial Parasites

Before Starting: All reagents should be for RNA use only. Ethanol (EtOH) solutions should be made with nuclease free or DEPC treated water. Clean your entire workspace and pipettes with RNaseZap® (Ambion® catalog #9780). Wear a clean lab coat and be sure to change gloves frequently!

1. Defrost worms over ice and transfer to a 2 mL round bottom tube. Use of a round bottom tube is important to allow enough space for the BB to vibrate within the tube (step 4).

2. Add 750 μL TRIzol® LS (Invitrogen® 10296-010) for every 250 μL of worms in buffer (3:1).

Note: Be sure to measure the volume of worms because this ratio is very important.

*TRIzol® LS is a monophasic solution of phenol and guanidine isothiocyanate, specially formulated for use with tissues in buffer, therefore you are working with a volume of worms in buffer, not a dry weight of tissues.

3. Do 3 freeze/thaw cycles: 3 minutes in dry ice/EtOH bath followed by 3 min at 80°C.

4. Add one 3mm stainless steel BB to the 2 mL round bottom tube and attach to vortex with special adaptor [i.e., Vortex Genie® adapter (Mo Bio Laboratories Inc catalog #13000V1); or the tube can be taped on its side to the flat portion of a regular vortex mixer platform.]. Vortex on the highest speed for 45 minutes, stopping every 10 minutes to change the position of the tube to ensure all areas of the tube are equally mixed.

5. Spin tube briefly before opening and add 200 μL chloroform for every 250 μL of worms in buffer. Vortex briefly and incubate for 3 minutes at room temperature.

6. Transfer the entire sample (except BB) to a pre-spun 2 mL heavy Phase Lock Gel™ tube [Prior to use, pre-spin the tube at 12,000 to 15,000 x g for 30 seconds (5 prime catalog # 2302830)]. Mix by inversion. Do NOT vortex.

*Use of the Phase Lock Gel™ greatly decreases organic contamination from the aqueous phase.

7. Centrifuge at 4°C for 15 minutes at 11,900 x g (no more than 12,000 x g).

8. Transfer the aqueous phase to a new 1.5 mL tube being careful to avoid the gel interface.

9. To precipitate the RNA, add 500 μL cold isopropanol (per initial 250 μL of worms in buffer). Vortex briefly and incubate for 10 minutes at room temperature.

10. Centrifuge at 4°C for 30 minutes at 12,200 x g to precipitate the RNA.

Note: At this step you should be able to see a small white pellet.

11. Carefully remove supernatant without disturbing the pellet.

12. Wash the pellet with 1 mL cold 75% EtOH by gently rocking the tube back and forth. Centrifuge at 4°C for 5 minutes at 7,500 x g.

13. Carefully remove supernatant without disturbing the pellet. Spin briefly in nanofuge and remove any remaining supernatant.

14. Invert on kimwipe (or equivalent) and air dry for 5 to 10 minutes or until there is no visible liquid.

15. Resuspend in 50 μL nuclease free water. Flick tube gently to mix.

16. Incubate at 55°C for 10 minutes to ensure complete re-suspension of the pellet. Flick tube gently to mix and then spin briefly in nanofuge.

17. Measure the total RNA concentration using a NanoDrop™ spectrophotometer or Agilent bioanalyzer.