



# PROSTATE CANCER **BIOREPOSITORY NETWORK**

SOP No: 006  
RNA Extraction from Frozen Tissues

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<b>STANDARD OPERATING PROCEDURE</b>	SOP No. 006 RNA Extraction from Frozen Tissue
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## 1. PURPOSE

To describe the procedure for the extraction of RNA from frozen tissue. NOTE: This SOP does not cover detailed procedures for handling Human Biological Materials or hazardous chemicals and it is recommended that personnel following this SOP refer to institutional safety guidelines.

**The following SOP for RNA extraction from frozen tissue is followed by the JHU and NYU Network Sites of the PCBN.**

## 2. RESPONSIBILITIES

**Authorized personnel** extracting RNA from frozen prostate tissue must ensure that:

- all procedures are followed correctly
- all samples are adequately coded during processing
- all documentation is completed, and accurate records maintained on all samples

## 3. HEALTH AND SAFETY

Personnel carrying out this procedure must maintain safe working practices and observe all relevant Environmental Health and Safety (EH&S) guidelines. This includes the appropriate use of Personal Protective Equipment (PPE), and the procedures for waste disposal, disinfection & spill clean-up and biosafety.

## 4. EQUIPMENT AND MATERIALS

Equipment	Materials
Cryostat, chuck, freezing bars	Chloroform
Centrifuge	Cryovials
gentleMACS Dissociator (JHU and NYU)	75% Ethanol in RNase free H <sub>2</sub> O
OCT	gentleMACS M Tubes
Refrigerated Microcentrifuge	Isopropyl alcohol
Vortex	Microcentrifuge tubes
	RNase Away
	RNase-free water
	Sterile blades for cryostat
	TRIzol® Reagent (Invitrogen Catalog # 15596-18)



## 5. PROCEDURES

### 5.1 Sample Preparation

- Collect tissue samples for RNA extraction from -80°C storage and place on dry ice.
- Cut 10 µm sections in a cryostat at -30°C.

NOTE: The tissue samples may require re-embedding in OCT, before 10 µm sections are cut.

*For each sample, a section is harvested at maximum diameter location at thickness of 5 µm both immediately before and immediately after the tissue sections collected in the tube for H&E staining and stored in the biorepository.*

- Collect sections into 3 ml cryovials (per sample). The starting sample weight should be no more than 50-100 mg for the TRIzol® Reagent.

NOTE: Use clean practice techniques for RNA handling (use of RNase Away or similar product on all surfaces, use RNase-free equipment, etc.). Change blade before cutting the next sample.

*We have tested the use of differing numbers of tissue sections for RNA extraction and have had success using up to 85, 10 µm sections of tissue, with greater yield from a greater number of sections. The only concern in maximizing the number of sections used is exhaustion of the block.*

### 5.2 RNA Extraction

- Add 1 ml of TRIzol® Reagent to tube with sectioned tissue. **NYU Networking site only:** no further homogenization step required.
- Incubate the homogenized sample for 5 min at 15-30°C to permit the complete dissociation of nucleoprotein complexes.
- **JHU Networking site only:** Transfer homogenous solution into a gentleMACS M Tube. Utilize the gentleMACS Dissociator on the RNA 02.01 pre-set program 3 times per sample to ensure complete homogenization of the tissue. Centrifuge M Tubes and transfer liquid into a fresh, sterile 1.5 ml microcentrifuge tube.
- Add 0.2 ml of chloroform per 1 ml of TRIzol® Reagent.
- Cap sample tubes securely. Shake tubes vigorously by hand for 15 sec and incubate at 15-30°C for 2-3 min.
- Centrifuge the samples at no more than 12,000 × g for 15 min at 2-8°C.

*Following centrifugation, the mixture separates into a lower red, phenol-chloroform phase, an interphase, and a colorless upper aqueous phase. RNA remains exclusively in the aqueous phase. The volume of the aqueous phase is about 60% of the volume of TRIzol® Reagent used for*



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*homogenization.*

- Transfer the aqueous phase to a fresh, sterile 1.5 ml microcentrifuge tube.
- Precipitate the RNA from the aqueous phase by mixing with isopropyl alcohol. Use 0.5 ml of isopropyl alcohol per 1 ml of TRIzol<sup>®</sup> Reagent used for the initial homogenization.
- Incubate samples at 15-30°C for 10 min.
- Centrifuge at no more than 12,000 × *g* for 10 min. at 2-8°C.

*The RNA precipitate, often invisible before centrifugation, forms a gel-like pellet on the side and bottom of the tube.*

- Remove the supernatant and discard, leaving the pellet untouched.
- Wash the RNA pellet once with 75% ethanol, adding at least 1 ml of 75% ethanol per 1 ml of TRIzol<sup>®</sup> Reagent used for the initial homogenization.
- Mix the sample by vortexing and centrifuge at no more than 7,500 × *g* for 5 min at 2-8°C.
- Remove the supernatant and discard, leaving the pellet untouched.
- Briefly dry the RNA pellet (air-dry or vacuum-dry for 5-10 min).

NOTE: Do not dry the RNA by centrifugation under vacuum. It is important not to let the RNA pellet dry completely as this will greatly decrease its solubility.

- Dissolve RNA in 200 µl (**JHU and NYU**) RNase-free water by passing the solution a few times through a pipette tip, and incubating for 10 min at 55-60°C.

This protocol is adapted from the Invitrogen handbook for TRIzol<sup>®</sup> Reagent (Catalog # 15596-018).