PreAnalytiX Supplementary Protocol

Purification of total RNA from microdissected PAXgene® Tissue fixed, paraffin-embedded (PFPE) and PAXgene Tissue fixed, cryo-embedded (PFCE) tissues

This protocol is for using the PAXgene Tissue RNA Kit to purify total RNA from PAXgene Tissue fixed, paraffin-embedded (PFPE) and PAXgene Tissue fixed, cryo-embedded (PFCE) tissues that have been manually or laser microdissected (LMD) from a slide.

IMPORTANT: The tissue samples must be fixed and stabilized in PAXgene Tissue Containers. The PAXgene Tissue Container Product Circular includes information on tissue fixation and stabilization. For instructions on the preparation of PFCE tissue blocks, read the PreAnalytiX® Supplementary Protocol Cryo-embedding tissue specimens fixed and stabilized with the PAXgene Tissue System (PX14). To prepare sections from PFPE and PFCE tissue blocks for microdissection, follow instructions of the Supplementary Protocol Preparation of sections from PAXgene Tissue fixed, paraffin-embedded (PFPE) and PAXgene Tissue fixed, cryo-embedded (PFCE) tissue for manual or laser microdissection (PX20).

Also read the PAXgene Tissue DNA Kit Handbook, paying careful attention to the “Safety Information” and “Important Notes” sections, before beginning this procedure.

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Equipment and reagents

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, consult the appropriate safety data sheets (SDSs), available from the product supplier.

- RNase-free water
- Ethanol (96–100%, purity grade p.a.)
- Xylene
- 14.3 M β-mercaptoethanol, β-ME (commercially available solutions are usually 14.3 M)
- Variable-speed microcentrifuge* capable of attaining 1000–8000 x g, and equipped with a rotor for 2 ml microcentrifuge tubes
- Microtome or cryostat*

* Make sure that instruments have been checked and calibrated according to the manufacturer’s recommendations.
- Shaker–incubator* capable of incubating at 45–80°C and shaking at ≥400 rpm, not exceeding 1400 rpm (e.g., Eppendorf® Thermomixer Compact)†
- Vortex mixer*
- Adhesion slides for manual microdissection (e.g., SuperFrost® Plus Slides, VWR® Cat. no. 631-0108) or frame slides for laser microdissection, (e.g., PPS-membrane 25mm x 76 mm, Leica® Cat. no. 11505273)†
- Laser Microdissection System* (e.g., Leica LMD6500 & LMD7000, Leica Microsystems)†
- 1.5 ml safelock microcentrifuge tubes
- 0.5 ml PCR tubes suitable for the dedicated LMD system (e.g. CORNING® AXYGEN® 0.5 m Tube with Flat Cap, Cat. no. 10169-890† for Leica LMD6500 & LMD7000)

**Carrier RNA**

The PAXgene Tissue RNA Kit contains poly-A RNA for use as carrier RNA. When added to lysates from microdissected tissue, the carrier RNA may improve the recovery of total RNA. Carrier RNA is not required when processing more than 5000 cells. The small amounts of poly-A RNA used as carrier RNA do not interfere with subsequent RT-PCR, even when oligo-dT is used as a primer for reverse transcription. Reverse-transcription reactions typically contain an excess of oligo-dT primers, and the small amounts of poly-A used as carrier RNA are insignificant in comparison. Total RNA purified using poly-A RNA as carrier RNA can be amplified with the QIAGEN® QuantiTect® Whole Transcriptome Kit, which uses a mix of random and oligo-dT primers.

**Starting material**

Starting material for RNA purification is a PFPE or PFCE tissue section mounted on a glass adhesion slide for manual microdissection (procedure A) or on a frame slide with a membrane for laser microdissection (procedure B). Sections must be prepared according to the Supplementary Protocol Preparation of sections from PAXgene Tissue fixed, paraffin-embedded (PFPE) and PAXgene Tissue fixed, cryo-embedded (PFCE) tissue for manual or Laser microdissection (LMD).

Detach the microdissected specimen from the slide and transfer it to a dedicated collection device for direct purification of total RNA.

**IMPORTANT:** The tissue source, the preanalytical workflow and the choice of materials used to make a PFPE and PFCE specimen significantly affect quality and quantity of isolated RNA. Furthermore, variability in quality and quantity of isolated RNA may be observed across species and tissue type.

* Make sure that instruments have been checked and calibrated according to the manufacturer’s recommendations.
† This is not a complete list of suppliers and does not include many important vendors of biological supplies.
Things to do before starting

- If working with RNA for the first time, read “Appendix A: General Remarks on Handling RNA” in the PAXgene Tissue RNA Kit Handbook.
- Set the temperature of the shaker–incubator to 45°C for PFPE tissues or 56°C for PFCE tissues.
- Buffer TR1 may form a precipitate during storage. If necessary, warm to 37°C to dissolve.
- Add β-Mercaptoethanol (β-ME) to Buffer TR1 before use. Add 10 µl β-ME per 1 ml Buffer TR1. Dispense in a fume hood and wear appropriate protective clothing. Buffer TR1 containing β-ME can be stored at room temperature (15–25°C) for up to 1 month. Pre-cool the TR1 buffer on ice.
- Buffer TR3 is supplied as a concentrate. Before using the buffer for the first time, add the amount of ethanol (96–100%, purity grade p.a.) indicated on the bottle to obtain a working solution.
- If using the RNase-Free DNase Set for the first time, prepare a DNase I stock solution. Dissolve the solid DNase I in 550 µl DNase resuspension buffer (RNase-free water) provided with the set. Do not vortex the reconstituted DNase I. DNase I is especially sensitive to physical denaturation. Mixing should only be carried out by gently inverting the tube. Reconstituted DNase I can be stored at 2–8°C for up to 6 weeks and at −15°C to −30°C for up to 6 months.
- When processing <5000 cells, carrier RNA may be added to the lysate (see “Carrier RNA”, page 2). Before using carrier RNA for the first time, dissolve it (310 µg) in 1 ml RNase-free water. Store this stock solution at −20°C and use it to make fresh dilutions for each set of RNA preparations. The concentration of this stock solution is 310 ng/µl. Prepare a working solution of 4 ng/µl using Buffer TR1.

Procedure A: Purification of total RNA from manually microdissected PFPE and PFCE tissue

In this procedure, ● refers to PFPE tissue and ▲ refers to PFCE tissue.

1. Label the lid and the body of a 1.5 ml safelock microcentrifuge tube (not provided). Prepare a lysis mixture in the tube by mixing 60 µl Buffer TR1 with 290 µl RNase-free water. Mix by gently flicking the tube. Add 10 µl Proteinase K, mix again and centrifuge briefly to collect residual liquid from the sides of the tube.
   
   Note: Do not mix Buffer TR1 and Proteinase K before adding water.

2. Using a ● microtome or ▲ cryostat, make a tissue section of 6–12 µm thickness from a ● paraffin-embedded or ▲ cryo-embedded tissue block. Capture the tissue section on an RNase-free adhesion slide.

3. Remove embedding medium and stain (optional) according to the Supplementary Protocol Preparation of sections from PAXgene Tissue fixed, paraffin-embedded (PFPE) and PAXgene Tissue fixed, cryo-embedded (PFCE) tissue for manual or Laser microdissection (LMD). Keep freshly prepared slides in ice-cold ethanol until microdissection.
4. For manual microdissection, remove the slide from the ethanol. Using an absorbent
sheet, wipe away the liquid on the slide that surrounds the tissue section.

**Note:** Tissue areas of interest can be isolated by scratching away other tissue parts with a scalpel
blade or cell scraper. Remove tissue portions from the slide immediately after taking the section
out of the ethanol. Avoid completely drying the section because this makes it harder to dissolve
the tissue from the slide.

5. Place the slide on a horizontal working plate and overlay it with 100 µl ice-cold Buffer
TR1. Make sure that the whole section is covered.

**Note:** Work quickly. We recommend using a dark underlay as a work surface to make it easier to
see the tissue. The volume of Buffer TR1 needed depends on the tissue surface area.

6. Detach the tissue from the slide by pipetting the lysis mixture up and down. Transfer the
tissue and all liquid to the labeled 1.5 ml safelock microcentrifuge tube from step 1 and
mix by vortexing for 5 s.

**Note:** If the tissue does not dissolve easily, use the pipet tip to scrape it from the slide.

7. Incubate the tissue on a shaker–incubator for 15 min at 1400 rpm and • 45°C or ▲ 56°C.

8. Centrifuge for 3 min at maximum speed (but do not exceed 20,000 x g).

9. Carefully transfer the supernatant to a new 1.5 ml microcentrifuge tube without
disturbing the pellet.

10. Continue with the addition of ethanol as described in step 12 of the protocol “Purification
of Total RNA from Sections of PFPE Tissue” in the PAXgene Tissue RNA Kit Handbook
(page 17). Set the temperature of the shaker–incubator to 65°C for use in step 22.

**Note:** The elution volume can vary between 14–40 µl and strongly depends on the area and
number of microdissected cells. Low input requires lower elution volumes for high concentrations
of total RNA. Furthermore, repeated elution steps increase the overall total RNA yield of a sample.

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**Procedure B: Purification of total RNA from laser
microdissected (LMD) PFPE and PFCE tissue**

In this procedure, • refers to PFPE tissue and ▲ refers to PFCE tissue.

1. Using a • microtome or ▲ cryostat, make a tissue section of 6–12 µm thickness from a
• paraffin-embedded or ▲ cryo-embedded tissue block. Capture the tissue section on a
frame slide with membrane for laser microdissection.

2. Remove embedding medium and stain (optional) according to the Supplementary
Protocol Preparation of sections from PAXgene Tissue fixed, paraffin-embedded (PFPE)
and PAXgene Tissue fixed, cryo-embedded (PFCE) tissue for manual or Laser
microdissection (LMD). Keep freshly prepared slides in ice-cold ethanol until laser
microdissection.

3. Using a Laser Microdissection System, dissect a tissue area of ≥5000 µm² (≥50 cells) from
the PFPE or PFCE LMD slide.
Note: ≤50 cells may require pre-amplification of cDNA for downstream applications (use, for example, the QIAGEN REPLI-g® WTA Single Cell Kit).

4. Collect cells directly into a dedicated collection tube filled with 15 µl Buffer TR1 (e.g., use the cap of a 0.5ml PCR tube when using a Leica LMD system).

   Note: If not possible, add Buffer TR1 immediately after collecting the cells. Make sure to add β-ME to Buffer TR1 before use (see “Things to do before starting”).

5. Add 65 µl Buffer TR1. Mix by vortexing for 30 s and centrifuge briefly (1–2 s at 500–1000 x g) to remove drops from the tube lid.

   Note: The volume of Buffer TR1 depends on the collection vessel used for microdissection, but should not exceed 80 µl. Lysates in Buffer TR1 can be frozen at –80°C, however, freezing may impact RNA yield when a low number of cells is used. Thaw frozen lysates on ice and proceed with step 6.

6. If processing <5000 cells, 20 ng carrier RNA (5 µl of a 4 ng/µl solution) may be added to the lysate. Prepare the carrier RNA as described in “Things to do before starting”.

7. Add 145 µl RNase-free water and 5 µl Proteinase K and mix by vortexing for 5 s.

   Note: Do not mix Buffer TR1 and Proteinase K before adding them to the sample.

8. Incubate on a shaker-incubator for 15 min at 45°C or ▲ 56°C and 1400 rpm. After incubation, centrifuge briefly (1–2 s at 500–1000 x g) to remove drops from the tube lid.

9. Transfer the sample and Buffer TR1 to a 1.5 ml safelock microcentrifuge tube (not provided).

10. Continue with the addition of ethanol as described in step 12 of the protocol “Purification of Total RNA from Sections of PFPE Tissue” in the PAXgene Tissue RNA Kit Handbook (page 17). Set the temperature of the shaker-incubator to 65°C for use in step 22.

   Note: The elution volume can vary between 14–40 µl and strongly depends on the area and number of microdissected cells. Low input requires lower elution volumes for high concentrations of total RNA. Furthermore, repeated elution steps increase the overall total RNA yield of a sample.
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