

# ***Gingeras Lab RNA-Seq Library Production Document***

## **ENCODE Transcriptome**

Sample Description: K562 Nucleoplasm RNA Biorep # 2

RNA ID: 026NP

Library ID: 9190

Protocol ID:

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### **Cold Spring Harbor Laboratory**

Genome Center

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### **LAB MEMBERS**

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**CELL CULTURE:** Cells are grown according to the ENCODE growth protocol and standards. Each bioreplicate grown and isolated independently.

**RNA ISOLATION:**

**Kits: miRNeasy Mini kit (QIAGEN cat#:217004)  
RNeasy MinElute cleanup kit (QIAGEN cat#:74204)**

1. Regular harvest and count cells and centrifuge for 10 min at 1900 rpm 4 °C.
2. Completely aspirate supernatant.
3. Resuspend all pellets in 10-30 mL of cold PBS by pipetting up and down.
4. Centrifuge for 5 min at 2000 rpm 4 °C.
5. Carefully aspirate the supernatant.
6. Add QIAzol Lysis Reagent to the pellet and vortex to mix.
7. If number of cells is  $\leq 3 \times 10^6$  then vortex for 1 min to homogenize the cells.  
If number of cells is  $>3 \times 10^6$  homogenize by placing 700  $\mu\text{L}$  of sample into a QIAshredder homogenizer and centrifuge for 2.5 min at maximum speed.
8. Place the tube(s) containing the homogenate on the benchtop at room temperature (15-25 °C) for 5 min.
9. Add 20% volumes of chloroform to the homogenate and cap it securely. Shake the tube vigorously for 15 s.
10. Place the tube containing the homogenate on the benchtop at room temperature for 2-3 minutes.
11. Centrifuge the homogenate for 15 min at 12,000 x g at 4 °C.
12. Transfer the upper, colorless, aqueous phase containing the RNA to a new collection tube.
13. Add and equal volume of 70% ethanol and mix thoroughly by vortexing. Do not centrifuge.
14. Pipet 700  $\mu\text{L}$  of the sample including any precipitate that may have formed into an RNeasy Mini spin column placed in a 2 mL collection tube. Close the lid gently and centrifuge at  $\geq 8000 \times g$  ( $\geq 10,000$  rpm) for 30 s at room temperature. Repeat this step until the whole sample has been pipetted into the spin column and discard the flow-through each time.
15. Pipet the flow-through (which contains miRNA) into a 2 mL RNase free tube. The spin column contains the large RNA.

**Large RNA Purification**

- L1. Pipet 700  $\mu\text{L}$  Buffer RWT into the RNeasy Mini spin column from step 15. Close the lid gently and centrifuge for 30 at  $\geq 8000 \times g$  ( $\geq 10,000$  rpm) to wash the spin column membrane. Discard the flow through.
- L2. Add 500  $\mu\text{L}$  Buffer RPE to the RNeasy Mini spin column. Close the lid gently and centrifuge for 30 s at  $\geq 8000 \times g$  ( $\geq 10,000$  rpm) to wash the spin column membrane. Discard the flow through.
- L3. Add another 500  $\mu\text{L}$  Buffer RPE to the RNeasy Mini spin column. Close the lid gently and centrifuge for 30 s at  $\geq 8000 \times g$  ( $\geq 10,000$  rpm) to wash the spin column membrane. Discard the flow through and the collection tube.
- L4. Place the RNeasy Mini spin column in a new 2 mL collection tube. Open the lid and centrifuge at full speed for 1 min.
- L5. Transfer the RNeasy Mini spin column to a new 1.5 ml collection tube. Pipet 30-50  $\mu\text{L}$  RNase-free water directly onto the RNeasy Mini spin column membrane. Close the lid gently. Wait one minute and then centrifuge for 1 min at  $\geq 8000 \times g$  ( $\geq 10,000$  rpm) to elute the total RNA (containing large RNA).
- L6. If the expected RNA yield is  $>30 \mu\text{g}$ , repeat step L5 with an additional volume of 30-50  $\mu\text{L}$  RNase-free water. Elute into the same collection tube.
- L7. Proceed to ethanol precipitation.

**Ethanol Precipitation**

1. Add 2.5 volumes of 100% ethanol and 1/10 volumes of NaOAc PH 5.5 to the eluted RNA.
2. Freeze in -80 °C for at least 30 min.
3. Centrifuge for 35 min at max speed at 4 °C.
4. Pipette and discard the supernatant making sure not to touch the pellet of RNA.
5. Wash with 1 mL of 70% ethanol and centrifuge at max speed for 5 min.
6. Pipette and discard the supernatant.
7. Open the cap and speed vacuum at low heat for 3-5 min making sure that the pellet is dry.
8. Resuspend the pellet with RNase-free water.

**DNase Digest (same for Small and Large RNA)**

<i>Reagents</i>	<i>100 <math>\mu\text{L}</math> Sample (100 <math>\mu\text{g}</math> RNA max)</i>	<i>50 <math>\mu\text{L}</math> Sample (50 <math>\mu\text{g}</math> RNA max)</i>
Total RNA (100 $\mu\text{g}$ max)	78 $\mu\text{L}$	39 $\mu\text{L}$
10X One-phor-all Buffer	10 $\mu\text{L}$	5 $\mu\text{L}$

10 U/ $\mu$ L DNase/RNase Free	8 $\mu$ L	4 $\mu$ L
20 U/ $\mu$ L RNasin/anti-RNase	4 $\mu$ L	2 $\mu$ L
<i>Total Volume</i>	<i>100 <math>\mu</math>L</i>	<i>50 <math>\mu</math>L</i>

1. Add all reagents to resuspended RNA and pipette to mix well.
2. Place in a 37°C waterbath for 30 min.
3. Proceed to RNA Cleanup, which is different for Small and Large RNA.

### Large RNA Cleanup

1. Add 350  $\mu$ L Buffer RLT to the 100  $\mu$ L (100  $\mu$ g) sample of RNA. Vortex to mix well.
2. Add 250  $\mu$ L of 100% ethanol to the reaction and mix by inverting.
3. Transfer the 700  $\mu$ L of sample to an RNeasy mini spin column placed in a 2 mL collection tube. Close the lid gently and centrifuge for 30 s at 8000 x g ( $\geq$ 10,000 rpm). Discard the flow through.
4. Add 700  $\mu$ L Buffer RW1 to the RNeasy mini spin column. Close the lid gently and centrifuge for 30 s at 8000 x g ( $\geq$ 10,000 rpm). Discard the flow through.
5. Repeat Step 4.
6. Add 500  $\mu$ L Buffer RPE to the RNeasy mini spin column. Close the lid gently and centrifuge for 30 s at 8000 x g ( $\geq$ 10,000 rpm). Discard the flow through.
7. Repeat Step 6.
8. Transfer the RNeasy spin column to a new collection tube. Centrifuge for 2 min at 10,000 x g to dry the RNeasy membrane.
9. Place the RNeasy spin column into a new 1.5 mL collection tube and discard the old tube. Add 30-50  $\mu$ L of RNase-free water directly on the spin column membrane. Close the lid gently and let stand for 1 min. Centrifuge for 1 min at 10,000 x g to elute the RNA.
10. Add another 30-50  $\mu$ L of RNase-free water onto the membrane using the same centrifuge tube. Wait 1 min and then centrifuge for 1 min at 10,000 x g to elute the RNA.
11. Proceed to ethanol precipitation.

### Ethanol Precipitation

1. Add 2.5 volumes of 100% ethanol and 1/10 volumes of NaOAc PH 5.5 to the eluted RNA.
2. Freeze in -80°C for at least 30 min.
3. Centrifuge for 30 min at max speed at 4°C.
4. Pipette and discard the supernatant making sure not to touch the pellet of RNA.
5. Wash with 1 mL of 70% ethanol and centrifuge at max speed for 5 min.
6. Pipette and discard the supernatant.
7. Open the cap and speed vacuum at low heat for 5 min making sure that the pellet is dry.
8. Resuspend the pellet with RNase-free water.

### RIBOMINUS TREATMENT:

#### Hybridization Step

Instructions are provided below to perform hybridization for 1–10  $\mu$ g of your total RNA sample with the RiboMinus™ Eukaryote Probe. To process >10  $\mu$ g total RNA sample, divide your sample into two samples, each containing <10  $\mu$ g total RNA.

1. Set a water bath or heat block to 70–75°C.
2. To a sterile, RNase-free 1.5 mL microcentrifuge tube, add the following:  
Total RNA (1–10  $\mu$ g): <10  $\mu$ L  
RiboMinus™ Probe (15 pmol/L): 10  $\mu$ L  
Hybridization Buffer: 100  $\mu$ L
3. Incubate the tube at 70–75°C for 5 minutes to denature RNA.
4. Allow the sample to cool to 37°C slowly over a period of 30 minutes by placing the tube in a 37°C water bath (a heat block works as well). To promote sequence-specific hybridization, it is important to allow slow cooling. **Do not** cool samples quickly by placing tubes in cold water.
5. While the sample is cooling down, proceed to **Preparing Beads**.

\*An earlier version of this protocol says to use RNA in less than 20 $\mu$ l, add 10 $\mu$ l of probe and 300 $\mu$ l hybridization buffer, this larger volume means you need to precipitate the ribominused RNA in a 2ml tube at the end. Either way works. It doesn't change anything else expect the supernatant volumes and the precipitation tube size.

### Preparing Beads

1. Resuspend RiboMinus™ Magnetic Beads in its bottle by thorough vortexing.
2. Pipet 750 uL of the bead suspension into a sterile, RNase-free, 1.5 mL microcentrifuge tube.
3. Place the tube with the bead suspension on a magnetic separator for 1 minute. The beads settle to the tube side that faces the magnet. Gently aspirate and discard the supernatant.
4. Add 750 uL sterile, DEPC Water to the beads and resuspend beads by pipetting
5. Place tube on a magnetic separator for 1 minute. Aspirate and discard the supernatant.
6. Repeat Steps 4–5 once.
7. Resuspend beads in 750 uL Hybridization Buffer and transfer 250 µL beads to a new tube and maintain the tube at 37°C for use at a later step.
8. Place the tube with 500 uL beads on a magnetic separator for 1 minute. Aspirate and discard the supernatant.
9. Resuspend beads in 200 uL Hybridization Buffer and keep the beads at 37°C until use.

### Removing rRNA

1. After the incubation at 37°C for 30 minutes of the hybridized sample (above), briefly centrifuge the tube to collect the sample to the bottom of the tube.
2. Transfer the sample (~120 uL- this will be ~330ul with the older protocol) to the prepared RiboMinus™ Magnetic beads from Step 9 (**Preparing Beads**, above). Mix well by pipetting up and down
3. Incubate the tube at 37°C for 15 minutes. During incubation, gently mix the contents occasionally. Briefly centrifuge the tube to collect the sample to the bottom of the tube.
4. Place the tube on a magnetic separator for 1 minute to pellet the rRNA-probe complex. **Do not discard the supernatant. The supernatant contains RiboMinus™ RNA.**
5. Place the tube with 250 µL beads from Step 7 (**Preparing Beads**, above) on a magnetic separator for 1 minute. Aspirate and discard the supernatant.
6. To this tube of beads, add ~320 µL (~500ul with older protocol) supernatant containing RiboMinus™ RNA from Step 4, above. Mix well by pipetting up and down or low speed vortexing.
7. Incubate the tube at 37°C for 15 minutes. During incubation, gently mix the contents occasionally. Briefly centrifuge the tube to collect the sample to the bottom of the tube.
8. Place the tube on a magnetic separator for 1 minute to pellet the rRNA-probe complex. **Do not discard the supernatant as the supernatant contains RiboMinus™ RNA.**
9. Transfer the supernatant (~ 320 uL- ~500ul with older protocol) containing **RiboMinus™ RNA** to a small filter column and spin at max speed for 2 minutes to remove any remaining magnetic particles.
10. Transfer flow through (ribominus RNA) to a new tube (1.5ml for small volume, 2ml for large volume)
11. EtOH precipitate with glycoblue

**SPIKE-INS:** NIST Spike-Ins beta set: Pool 14. 2 ng added to 100 ng of RNA. Use “corrected fasta” to map against.

**LIBRARY PROTOCOL:** Adapted from... *Transcriptome analysis by strand-specific sequencing of complementary DNA* Dmitri Parkhomchuk, Tatiana Borodina, Vyacheslav Amstislavskiy, Mariya Banaru, Linda Hallen, Sylvia Krobitsch, Hans Lehrach & Alexey Soldatov.

Use 100 ng of ribominused RNA (add spike ins, pool14 NIST beta) in 4.75ul or less of RNase free H2O for cDNA synthesis

### **cDNA- 1<sup>st</sup> strand:** Mix

4.75 ul sample r- RNA  
2ul 50ng/ul random primers  
2.5 50uM oligo-DT primer  
2ul NIST spike-ins  
1.25ul RNase inhibitor  
Up to 12.5ul with RNase free H2O if needed

98° 2 min  
70° 5 min  
.1°/s ramp to 15°  
15° 30 min  
.1°/s ramp to 25°  
25° 10 min  
.1°/s ramp to 42°  
42° 45 min

.1 °/s ramp to 50°  
50° 15 min  
75° 15 min  
4° hold

**As soon as** 15 degrees is reached (after ~15min), pause program and add:

5ul 5X First Strand Buffer  
1.25ul .1M MgCl<sub>2</sub>  
1.25ul 10mM dNTPs  
2.5ul .1M DTT

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22.5ul (total at this point)

After 30 minutes at 15 degrees, pause program and add (**before temp. ramp!**):

1.25ul Actinomycin-D (we have a 1mg/ml stock, dilute to 120ng/ul in 10mM Tris-Cp pH 7.6 before use)  
1.25ul Superscript III

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25ul final volume for 1<sup>st</sup> strand reaction

Rest of reaction takes about 1 hour 40 minutes  
Then, 4 degree hold

Bring reaction volume to 100ul (add 75ul Rnase free H<sub>2</sub>O)  
Add 5 volume PB (500ul) mix and apply to Minelute spin column  
Follow Qiagen Minelute cleanup protocol  
Elute 2 x 15ul EB

## **2<sup>nd</sup> Strand Synthesis**

Prepare 2<sup>nd</sup> strand mix:  
(22.5ul per sample)

1ul 5X 1<sup>st</sup> Strand Buffer  
15ul 5X 2<sup>nd</sup> Strand Buffer  
.5ul MgCl<sub>2</sub>  
1ul DTT  
2ul dUNTPs  
.5ul E. coli DNA ligase  
2ul E. coli DNA polymerase I  
.5ul RNase H

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22.5ul

Mix:  
30ul first strand reaction  
22.5ul second strand mix  
22.5ul RNase free H<sub>2</sub>O

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75ul final reaction volume

2 hours 16 degrees, 4 degrees hold in PCR machine  
Bringing volume to 100ul with H<sub>2</sub>O, then add 500ul PB, follow minelute cleanup protocol  
Elute 2 x 26ul (fragmentation takes place in 50ul).  
Bioanalyzer- high sensitivity DNA chip (to see if cDNA is full length, peak should be around 1000bp- if it is not, you need to lessen fragmentation time)

## **Fragment cDNA: Covaris**

If machine is not on:  
Fill appropriate chambers with autoclaved DI water  
Run degas program (~30 minutes)

Transfer your 50ul cDNA sample to the sonicator tube (using pipette)  
Place on machine (snaps in) and run program degas60snapcap100ul (60s sonication)  
Run Bioanalyzer- high sensitivity DNA chip to check fragment size (peak should be 200-300)

### **End-Repair cDNA**

48ul sample  
27ul H<sub>2</sub>O  
10ul T4 DNA ligase buffer with 10mM ATP ("10X ER")  
4ul dNTP mix 10mM  
5ul T4 DNA polymerase 3U/ul (NEB)  
1ul Klenow DNA polymerase 5U/ul (NEB)  
5ul T4 PNK 10U/ul (NEB)

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100ul final volume

Room temp. 30min.  
Add 500ul PB, follow Minelute cleanup, elute 2 x 16ul

### **Addition of single <A> Base**

32ul eluted cDNA  
5ul NEBuffer2  
10ul dATP (1mM)  
3ul Klenow fragment 3' to 5' exo- 5U/ul

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50ul final volume

37 degrees, 30 min.

Bring volume to 100ul (add 50ul H<sub>2</sub>O), then add 500ul PB  
Follow minelute cleanup, elute 1 x 19ul

### **Adapter Ligation**

19ul eluted cDNA  
25ul DNA ligase buffer  
1ul adapter oligo mix  
5ul DNA ligase 1U/ul (Enzymatics)

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50ul final volume

Room temp, 15 min.  
Bring volume to 100ul with H<sub>2</sub>O (add 50ul), then add 500ul PB  
Minelute cleanup, elute 1 x 15ul

### **UNG Treatment**

15ul eluted cDNA  
1.7ul 500 mM KCl  
1ul UNG

37 degrees, 15 min  
95 degrees, 10 min  
Hold on ice

Add 10ul loading buffer

Run on 2% Ultra-pure agarose gel for 2 hours, 70V (use 100bp ladder)  
Cut out 200bp band, and another band just slightly larger (freeze larger slice, -20)

If you do not see anything on the gel at this point, do not be alarmed, cut bands anyway

Use Qiaquick gel extraction kit, elute 2 x 15ul

**PCR Amplification:** Mix

15ul eluted cDNA from gel-extraction (freeze remaining cDNA)- If you suspect you need more or less for good amplification, use more or less

1ul PE primer 1

1ul PE primer 2

50ul HF Phusion Mix

33ul H2O (adjust this volume according to how much cDNA was used)

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100ul final PCR volume

98° 1 min

18 cycles of:

98° 10s

60° 30s

72° 30s

72° 5 min

4° hold

Add 500ul PB, Min-elute clean up, elute 1 x 15ul

Add 10ul loading dye, run on 2% gel at 70V for 2 hours

Cut distinct band (should be ~100bp larger than cDNA band)

You may have more than 1 band at this point, cut whichever band is ~100bp larger than your cDNA cut was

Gel purify as before

Elute 2 x 15ul

Precipitate

Resuspend 25ul H2O

Measure library with Nanodrop (1ul) –very inaccurate.

Run High sensitivity DNA chip (1ul)

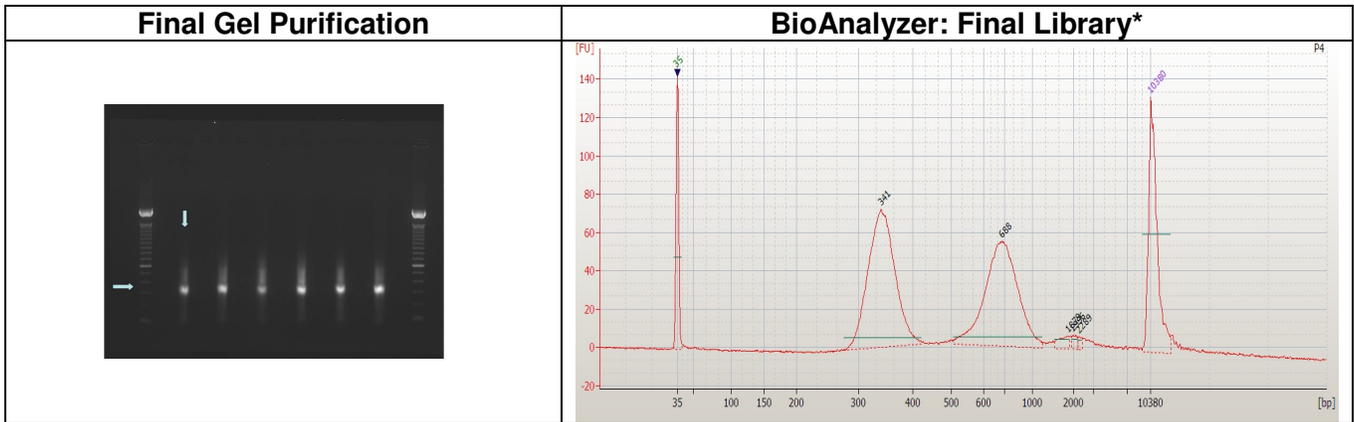
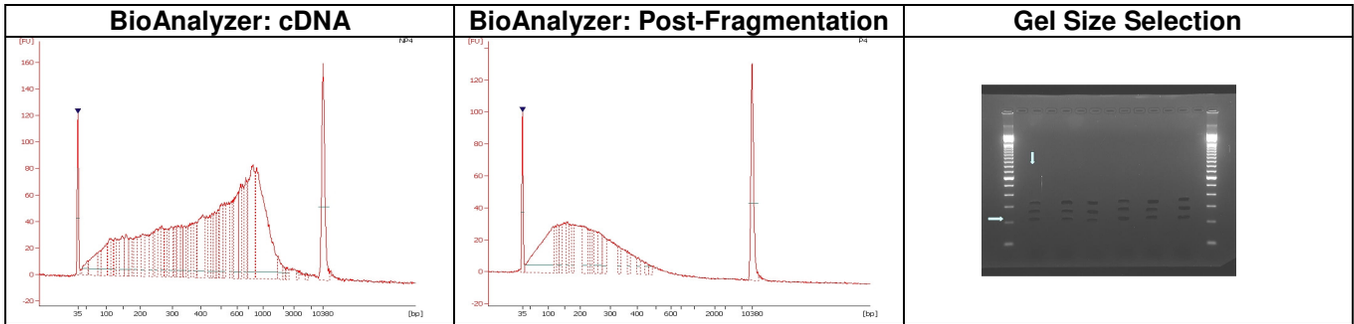
Measure concentration also with Tecan (pico green) (1ul per dilution)

KAPA Biosystems qPCR (1ul) per dilution

Dilute to 10nM (do not have to use whole library)

Prior to cluster generation we add PhiX at 1%.

<b>BioAnalyzer: Total RNA</b>	<b>BioAnalyzer: After oligo-dT</b>	<b>BioAnalyzer: After Ribominus</b>



\* Sometimes we see a doublet in the BioAnalyzer image of the final library. We take the height of the first peak to represent the library insert size when determining molarity. These doublets are not visible on gels, the libraries sequence fine and show inserts surrounding the first peak size.