

# ATAC-seq on nuclei from frozen, sliced, *Drosophila melanogaster* embryo halves Version 2

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## Abstract

Protocol for performing ATAC-seq on nuclei isolated from *Drosophila melanogaster* stage 5 embryos that were flash frozen and then cut in half along the anterior-posterior midline. Data from this protocol are presented in the following paper:  
<https://www.biorxiv.org/content/early/2017/09/27/195073>.

This protocol is based adapted from the following sources:  
<https://www.nature.com/nmeth/journal/v10/n12/full/nmeth.2688.html> and  
<http://journals.plos.org/plosgenetics/article/>

**Citation:** Jenna Haines ATAC-seq on nuclei from frozen, sliced, *Drosophila melanogaster* embryo halves. **protocols.io**  
[dx.doi.org/10.17504/protocols.io.kj5cuq6](https://doi.org/10.17504/protocols.io.kj5cuq6)

**Published:** 02 Nov 2017

## Protocol

### Solutions

#### Step 1.

##### Lysis Buffer NO detergent

- 10mM Tris-HCl, pH 7.4
- 10mM NaCl
- 3mM MgCl<sub>2</sub>

**Store at 4°C- make fresh weekly**

##### Lysis Buffer Freezing Media

- 10ml Lysis Buffer No Detergent
- 1ml of 50% glycerol
- 1ul of bromoblue dye

##### Lysis Buffer + Spermine

- 10ml Lysis Buffer No Detergent
- 0.15mM spermine

### Step 2.

Put several hundred adult OregonR *Drosophila melanogaster* (<http://flybase.org/reports/FBsn0000276.html>) into a fly cage at 25°C for three days. Feed flies with standard molasses plates a spread of yeast paste. Collect embryos for 2 hours and then let them age at 25°C for an hour and thirty minutes. This should yield embryos between 2:00 and 3:30 hours old.

### Step 3.

Bleach embryos for 3 minutes in 30%-50% bleach to remove the chorion.

### Step 4.

Rinse well with water. Pat dry on towel.

### Step 5.

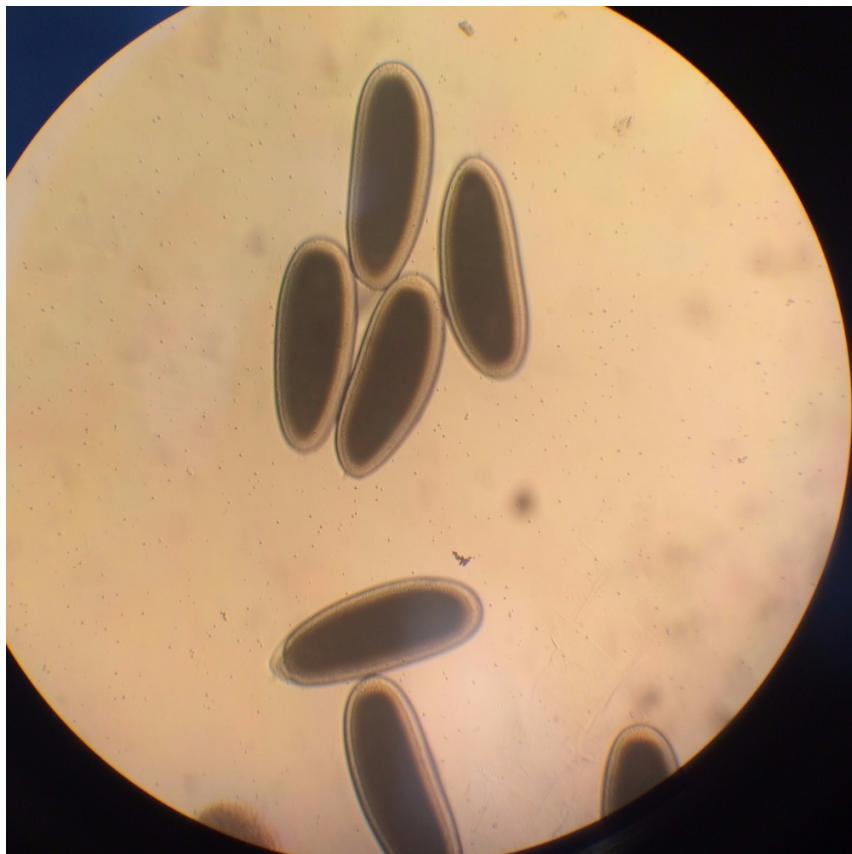
Rinse with 1xPBS + 0.5% Triton.

### Step 6.

Brush embryos into an 1.5ml tube filled with 1xPBS + 0.5% Triton. Let embryos settle. Then use a pipette tip to transfer a drop of embryos to a slide.

### Step 7.

Stage the embryos to the desired age. We usually go for Stage 5.



## Freezing and Slicing embryos

### Step 8.

Chill a cold block in the -80 for at least 30 minutes. Once it's cold, place in a box surrounded by dry ice to keep it cool while you slice



#### REAGENTS

Corning® CoolRack CF15, Holds 15 Cryovial or FACS Tubes (Product #432049) [432049](#) by [Corning](#)

## Freezing and Slicing Embryos

### Step 9.

Place staged embryos into ependorf tubes containing Lysis buffer freezing media.

### Step 10.

Keep the slides, poker tool, and scalpel on dry ice to keep from melting.



#### REAGENTS

Shandon™ Straight Point Teasing Needles, Straight Point, Wooden Handle, 5.5 in. (14.0cm) [19010](#) by [Thermo Scientific](#)

Fisherbrand™ High Precision #10 Style Scalpel Blade [12-000-162](#) by [Fisher Scientific](#)

### Step 11.

Underneath a dissection scope, space out staged embryos on the glass slide such that they are evenly separated and can be cut easily with the blade. I affix a grid pattern to the bottom side of the glass slide to offer contrast to the white embryos.

## Step 12.

Once the embryos are lined up on a slide, gently place the slide on dry ice for 2-5 minutes while you prepare the next slide. The embryos will turn from clear to completely white when frozen. I usually line up 10 embryos / slide and slice 5-6 slides at a time.

## Step 13.

Once embryos are frozen, immediately transfer the slide to a cold block that has been chilled. I surround the block with dry ice to keep it cool. Additionally, it is necessary to cool down the scalpel on dry ice before cutting. I leave the scalpel and 2 poker tools in dry ice while cutting to keep them cold. Be sure to wear gloves and keep the scalpel handle away from the dry ice to keep hands warm.

## Step 14.

Cut embryos along the midline with the dry ice chilled blade or scalpel. Move each half to a 1.5ml tube filled with 50ul of ATAC Lysis buffer + Spermine with the chilled poker tool.

- The embryos will thaw and dissolve as soon as they are transferred. To reduce contamination, we use 2 poker tools, one to handle anterior halves and one to handle posterior halves. Additionally, we wash pokers in 70% EtOH between slices

### AMOUNT

50 µl Additional info: ATAC Lysis buffer + Spermine

## Step 15.

Once your tube contains the desired number of embryo halves, move on to isolate the nuclei.

## Embryo Lysing

## Step 16.

Crush using a plastic pestle upwards of 20 times vigorously.



### REAGENTS

Pellet pestles [Z359947](#) by [Sigma](#)

## Step 17.

After homogenization, rinse the pestle with 50ul of lysis buffer to make sure not to lose any material. This will bring the total volume up to 100ul.



### AMOUNT

50 µl Additional info: Lysis buffer

## Step 18.

Vortex slightly at low speed and spin down on a table top centrifuge.

## Step 19.

Add 1ul of 10% IGEPAL CA-630 drop-wise to a final concentration of 0.1%. Mix well by flicking tube with a finger. **Let sit for about 10 minutes**

#### AMOUNT

1 µl Additional info: 10% IGEPAL CA-630

#### Step 20.

Spin in microcentrifuge at 800xg for 10 minutes at 4dC. Remove as much supernatant as possible to avoid nuclei loss.

#### Step 21.

Resuspend nuclei in enough DNase and RNase free water for the transposition reaction (below).

#### Tagmentation

#### Step 22.

Make sure the cell pellet is set on ice.

To make the transposition reaction mix, combine the following:

Sample	20 halves- 10 embryos
2x TD Buffer-	25
Tn5 Enzyme	7.5
H2O/ DNA	17.5
Total	50

#### REAGENTS

✓ Nextera DNA library preparation kit 24 samples [FC-121-1030](#) by Contributed by users

#### Step 23.

Gently pipette to resuspend nuclei in the transposition reaction mix.

#### Step 24.

Incubate the transposition reaction at **37°C for 30 min**. Immediately following transposition, purify using a **Qiagen MinElute Kit**.

#### REAGENTS

MinElute Reaction Cleanup Kit [28204](#) by [Qiagen](#)

#### Step 25.

Purified DNA can be stored at -20°C

I quantify DNA at this point with the qubit.

## PCR amplification

### Step 26.

To amplify transposed DNA fragments, combine the following in a PCR tube:

- 10  $\mu$ L Transposed DNA
- 5  $\mu$ L 25 $\mu$ M Primer 1
- 5  $\mu$ L 25 $\mu$ M Primer 2
- 25 $\mu$ L NebNext 2x Master Mix
- 5 $\mu$ L of PPC

50  $\mu$ L Total

Run PCR as follows:

- (1) 72°C, 5 min
- (2) 98°C, 30 sec
- (3) 98°C, 10 sec
- (4) 63°C, 30 sec
- (5) 72°C, 1 min
- (6) Repeat steps 3-5, **5x**
- (7) Hold at 4°C

## Optional - QPCR

### Step 27.

You can stop at this point and run a small qPCR reaction to calculate the number of additional cycles to use to avoid overamplification.

Take out 5  $\mu$ L of PCR reaction and run the qPCR reaction with the same conditions.

5  $\mu$ L Transposed DNA

0.25 ul 25uM Primer 1

0.25 ul 25uM Primer 2

5ul NebNext 2x Master Mix

3.9 ul of H<sub>2</sub>O

0.09 100x Sybr Green I Dye

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15 µL Total

Put in the QPCR machine and run:

1 cycle: 30 sec 98° C

20 cycles: 10 sec 98° C

30 sec 63° C

1 min 72° C.

To calculate the additional number of cycles needed, plot linear Rn (fluorescence) versus cycle and determine the cycle number that corresponds to one-third of the maximum fluorescent intensity.

### Step 28.

Once you decide how many additional cycles to run, run the rest of the PCR :

(2) 98°C, 30 sec

(3) 98°C, 10 sec

(4) **63°C**, 30 sec

(5) 72°C, 1 min **For X cycles**

(7) Hold at 4dC

Ampure (or Spri) bead Cleanup

### Step 29.

Bring Beads to RT.

#### Ampure (or Spri) bead Cleanup

##### Step 30.

Add 1.25x Beads, Mix well, let stand for 5 minutes.

#### Ampure (or Spri) bead Cleanup

##### Step 31.

Put on a magnet, let stand for 2 minutes.

#### Ampure (or Spri) bead Cleanup

##### Step 32.

Wash with 70% EtOH. (1/2)

#### Ampure (or Spri) bead Cleanup

##### Step 33.

Wash with 70% EtOH. (2/2)

#### Ampure (or Spri) bead Cleanup

##### Step 34.

Let dry for 10 minutes.

#### Ampure (or Spri) bead Cleanup

##### Step 35.

Resuspend in 20ul H<sub>2</sub>O.



20 µl Additional info: H<sub>2</sub>O

#### Ampure (or Spri) bead Cleanup

##### Step 36.

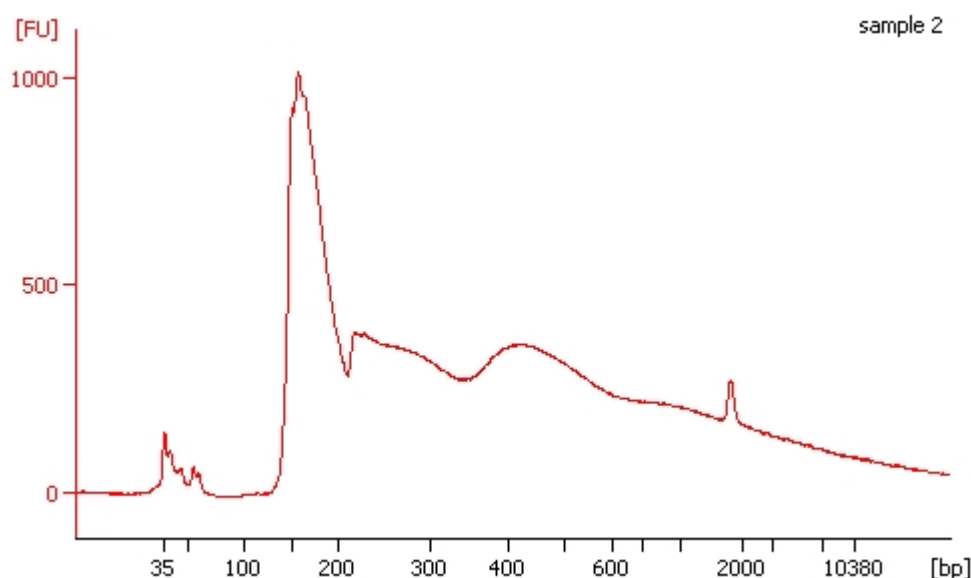
Let sit for one minute, put on magnet, take of super.



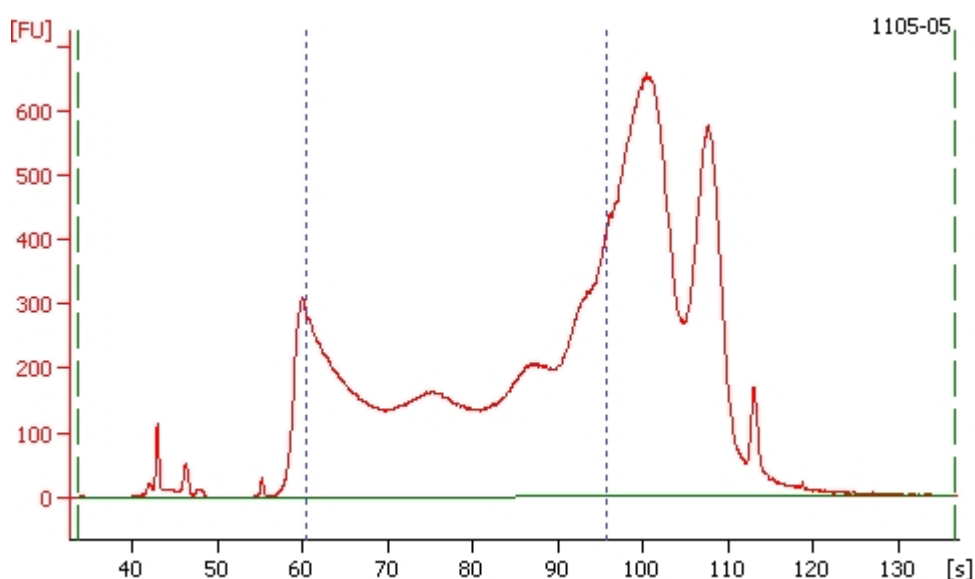
**Step 37.**

To validate your libraries, quantify them with qubit HS DNA and evaluate them with the Bioanalyzer High Sensitivity DNA assay.

Sometimes they look like this:



Sometimes they look like this: I usually see this when I am working with smaller amounts of embryos. Below is from one embryo half. I am not 100% sure what this means (whether it's actually undertagmented or what) but when I sequenced this sample the data was comparable to my other samples. If you have any ideas let me know! I usually try not to sequence these samples but they could be perfectly fine.



## Warnings

- Always wear proper PPE
- Be careful to not get frostbite when working with dry ice. Double glove when handling the chilled scalpel.