Imaging based barcode readout in tissue sections

This protocol is built upon Askary, et al. 2020. *Nature Biotechnology* 38 (1): 66–75. We have mostly used this procedure on cryosections of mouse retinal tissue. It has also been used successfully in other contexts such as chick neural tube, mouse brain, bone marrow, and skin.

For this protocol, it is important to not fix samples with formaldehyde prior to the in vitro transcription step. So the tissues are freshly frozen in OCT and sectioned at 15-20µm thickness. It is also important to keep RNases away from your sample and reagents to the extent possible. Use RNase free water and buffers, spray the surfaces and your gloves with RNase Away, do not touch anything without gloves, and use filtered RNase free tips. The most common source of RNase contamination in the labs is from miniprep buffers that contain high concentration of RNase (e.g. Buffer P1).

To image the samples multiple rounds with an inverted microscope, you need to section them on a coverslip with surface coating that promotes adhesion of the tissue. You can find the protocol for preparation of such coverslips here:

https://www.protocols.io/view/coverslips-preparation-for-seqfish-36wgq5rpogk5/v1

The readout procedure here is based on seqFISH+ (Eng, et al. 2019. *Nature* 568 (7751): 235–39). So for more information check out the paper and this related protocol:

https://protocolexchange.researchsquare.com/article/nprot-7333/v1

Here we use two types of probes. Readout probes are 15nt ssDNA oligos that are coupled with a fluorescent dye. Primary probes are longer ssDNA oligos (usually around 150nt) that bind to transcribed barcode RNAs and also include sequences complementary to readout probes. Primary probes are not fluorescent themselves. This scheme provides flexibility in assigning fluorescence channels to barcodes and is cost effective. In most cases the region of primary probes that anneals to transcribed barcodes (i.e. footprint) is 20nt long. Short footprint of the primary probes makes it necessary to circularize them to reduce the chance of losing them during multiple wash and rehybridization rounds. If primary probes with longer footprint (e.g. 35nt) are used for an experiment circularization and post-fix steps in Day 3 can be skipped.

**Day 1**

Air dry the slides by leaving them at room temperature (RT) for ~ 5-10 min.

Fix and permeabilize in a jar of MAA (37.5 ml of Methanol + 12.5 ml of Acetic Acid) for 3 hours at RT.

Wash with PBS (in jars) two times, ~5 min each.
Start preparing the transcription mix at this point by mixing nuclease free water (NFW), NTPs, and 10X buffer at RT. However, keep the enzyme at -20°C and don’t add it until the slides are ready. We usually use MEGAscript T3 Transcription Kit (ThermoFisher; Catalog number: AM1338).

The amount of the transcription mix that is needed for each slide depends on the type of chamber being used. We usually use hybridization chambers from Grace Biolabs (for example: https://gracebio.com/product/secureseal-hybridization-chambers-621505/). To save transcription mix, you can also put as low as 10µl of the mix on the center of your sections and spread it by placing a clean piece of parafilm on top.

To make a transcription mix of 10µl with MEGAscript T3 enzyme: Mix 1µl of each NTP with 1µl of 10X transcription buffer, and 4µl of NFW. 1µl of enzyme is added when you are ready to start the reaction. It is very important to use the cleanest water possible here, otherwise you not only don’t get a barcode signal but also destroy the endogenous mRNAs.

Transfer slides to a jar with NFW. After a few minutes, take the slides out one by one, wait until they are mostly dry but there is still water on the sections, put the hybridization chamber on, and then add NFW on the sections. It is important that the sections don’t over dry at this step, otherwise solutions may not easily cover the sections in the following steps.

When all the slides are ready, add the enzyme to the transcription mix, replace water on the slides with the mix, seal the holes on the chambers, and keep the slides at 37°C overnight in a humid box.

**Day 2**

Replace the transcription mix with 4% PFA and fix the sections for 10 min at RT.

Permeabilize in 8% SDS in PBS for 10 minutes.

Prepare 8% SDS fresh every time: 400µl of 20% SDS + 100µl of 10X PBS + 500µl of NFW

Wash with PBS, three times ~5 min each.

Wash with 5X SSC, three times ~5 min each.

Pre-hyb with hybridization buffer for at least 30min at 37°C in a humid box.

The choice of hybridization buffer depends on the application. If the goal is to detect single base pair variations, we use 30% hyb buffer from Molecular Instruments. If we only want to check the presence or absence of a barcode sequence, we usually use a 10% hyb buffer which is: 10% formamide, 10% dextran sulfate (Sigma D8906), 2X SSC in NFW.

Hybridize the primary probes at 4nM/probe final concentration in the hybridization buffer overnight at 37°C in a humid box. Seal the holes of the chamber for this step to avoid excessive evaporation.
Day 3
Wash 4 times at 37°C over a one hour period with a wash buffer containing the same percentage formamide as the hybridization buffer.

Wash buffer ingredients are 10 or 30% formamide, 0.1% Triton X-100, 4X SSC in NFW. Stringency of wash should be comparable to that of hybridization. So use the same percentage of formamide as you used for hybridization, depending on the application.

Wash 3 times with 5x SSCT at RT.
You can pause at this point and leave the samples in 5x SSC 4°C.

If primary probes with 20nt footprint are used and sequential rounds of hybridization/imaging are needed, proceed to circularization and postfix as follows. Otherwise skip the bullet points below and continue with hybridization of readout probes.

- Hybridize samples with 50nM ligation bridge in 20% hyb buffer 4xSSC at 37°C for at least 4 hours.
  - Ligation bridge is a ssDNA oligo that bridges the two arms of the primary probes. It has a 15nt complementary sequence on each arm.
  - The hyb buffer here is: 4X SSC, 20% Formamide, 10% Dextran Sulfate (Sigma D8906) in NFW.
- Wash with 10% wash buffer (10% formamide, 0.1% Triton X-100, 4X SSC in NFW) for 5 minutes at room temperature.
- Wash 3 times with 4xSSC at room temperature.
- Wash each slide with 1:1 mix of NFW and NEB Quick Ligase Reaction Buffer (2X) for 10 min at room temperature.
- Replace wash with ligation mix (125µl Quick ligase buffer, 87.5 ul NFW, 25µl 10mM ATP, 12.5µl Quick ligase) and incubate at RT overnight.

Day 4
- Wash with 10% wash buffer (10% formamide, 0.1% Triton X-100, 4X SSC in NFW) for 5 minutes at RT.
- Wash 3 times with 4X SSC.
- Wash once with 1x Label-IT buffer (Mirus MIR3900; stock 10x, diluted with NFW).
- Incubate with 1x Label-IT (stock 10x) in 1x Label-IT buffer at RT for 45 minutes (or 37°C for 30 min might be better).
- Wash 3 times with 1xPBS.
- Cross-linking with 1.5 mM BS(PEG)5 in 1xPBS at room temperature for 1 hr.
  - Prepare a 15mM solution by dissolving a 1mg vial of BS(PEG)5 (ThermoFisher Cat# A35396) in 125μL of dry DMSO. Mix by repeated pipetting or by replacing the cap and vortexing. The maximum usable volume of the vial is 800μL. BS(PEG)5 is a translucent solid and will most likely not be visible at the bottom of the vial.
- Wash 3 times with 1M Tris-HCl pH7.4.
- Wash with 55% wash buffer (55% formamide, 0.1% Triton-X 100, 4× SSC in NFW) for 5 minutes.

**Continue here if circularization was not needed:**

At this point the samples are ready for cycles of readout probe hybridization and imaging. The fluorescent readout probes are hybridized at 50nM each in 10% EC buffer for 20 minutes at room temperature. See below for instructions on how to make the EC buffer.

Wash for readout probes is done in 10% Wash Buffer (10% formamide, 0.1% Triton-X 100, 4× SSC in NFW) for 5 minute at room temperature followed by 1 minute wash in 4× SSC.

Stain with DAPI diluted in 4xSSC to 1µg/ml for 10 min.

Wash with 4x SSC for three times, ~5 min each.

Image in anti-bleaching buffer.

Anti-bleaching buffer is made up of 1% (w/v) glucose, 1:100 diluted catalase (Sigma C3155), 1.0 mg/ml Glucose oxidase (Sigma G2133), and 50mM pH8 Tris HCl in 4x SSC.

First, make a 1.43x mix of Ab base:

1M Tris HCl pH8 - 3.575mL
20x SSC - 14.3mL
0.715g of glucose
0.0375g trolox
Water to 50mL

Make sure to leave the Ab base overnight at 4°C after everything (except enzymes) is mixed and dissolved, to let Trolox convert into Trolox quinone which helps for anti-bleaching more. Add Catalase and Glucose oxidase before use.
If imaging many samples, it is possible to make a batch of anti-bleaching buffer. However, make sure to add the anti-bleaching buffer to the sample right before imaging and replace it with 4x SSC when the imaging is done. Also, use mineral oil to cover the top of the buffer to limit the oxidation reaction - solution should be stable for days.

After each round of image acquisition the samples are washed 3 times using 55% wash buffer at room temperature for 5 minutes to strip off readout probes.

Washing multiple times is crucial for completely stripping off readout probes. This is then followed by 2x SSC wash for 1 minute each round.

Then, the fluorophore-coupled readout probes are incubated at 50nM each concentration at room temperature for 20 minutes in 10% EC buffer followed by 5 minute wash in 10% wash buffer and then 1 minute wash in 4X SSC and DAPI staining.

This procedure can be repeated with the next round of readout hybridization until the completion of all rounds.
**Reagents**

**Primary probes**

We usually order oligoPools from IDT as 50pmol per probe. Dissolve probes in one tube in 50µl of IDTE pH8.0 to get 1µM stock (250x).

**10% wash buffer stock (1x) / 55% wash buffer stock**

<table>
<thead>
<tr>
<th>Source</th>
<th>Stock concentration</th>
<th>Final concentration</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>UltraPure SSC</td>
<td>Gibco LifeTech</td>
<td>20x</td>
<td>2x</td>
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<tr>
<td></td>
<td>Cat# 15557-036</td>
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<tr>
<td>Formamide</td>
<td>Ambion</td>
<td>-</td>
<td>10% / 55%</td>
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<tr>
<td></td>
<td>Cat# AM9342</td>
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<td></td>
</tr>
<tr>
<td>TritonX-100</td>
<td>Sigma-Aldrich</td>
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<td></td>
<td>Cat# 93443</td>
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</tr>
<tr>
<td>UltraPure water</td>
<td>100% EC</td>
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<td>7.9 ml / 3.4 ml</td>
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<tr>
<td></td>
<td></td>
<td>TOTAL</td>
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**100% EC**

EC chunks from the original stock (Sigma-Aldrich E26258)

Put in 50ml tubes and warm up in the 42°C water bath for 30-60 min until it melts.

**50% (v/v) EC solution**

<table>
<thead>
<tr>
<th>100% EC solution</th>
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<tbody>
<tr>
<td>Ultrapure water</td>
<td>6 mL</td>
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</table>

Note: should NOT vortex
### 10% (v/v) EC solution (1.67x)

<table>
<thead>
<tr>
<th>Source</th>
<th>Stock concentration</th>
<th>Final concentration</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>UltraPure SSC</td>
<td>Gibco LifeTech</td>
<td>20x</td>
<td>3.34x</td>
</tr>
<tr>
<td>Cat# 15557-036</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>50% EC solution</td>
<td>above</td>
<td>50% = 5x</td>
<td>16.7% = 1.67x</td>
</tr>
<tr>
<td>Dextran sulfate LW</td>
<td>Sigma-Aldrich</td>
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<td>16.7% (w/v)</td>
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<tr>
<td>UltraPure water</td>
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<td></td>
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<td>TOTAL</td>
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