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## seqFISH Tissue Preservation V.1

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ABSTRACT

Tissue Preparation for seqFISH

MATERIALS TEXT

## Fixation

- 1 Clean surface with RNAzap. Make the following solutions.
  - 1L PBS 1x, RNAse-free
  - 1. take 1 L RNAse-free water bottle
  - 2. use serologic pipet and draw up 100 mL water into another container
  - 3. add 100 mL of 10x RNase-free PBS
  - 4. mix thoroughly

## 50% sucrose

- 1. tare 1 L RNase-free disposable container
- 2. take 500 g RNAse-free sucrose container
- 3. add RNAse-free water to sucrose and transfer to 1 L container
- 4. continue to add RNAse-free water until mass to 1  $\mbox{kg}$
- 5. mix thoroughly, may need agitation on platform at RT for some time
- 6. store at 4°C

NOTE: for fixation, minimum PFA to tissue volume is 20:1, ideal closer to 50:1

2 Prepare 4% PFA in 1x PBS, 120 mL total for 4 tissues 30 mL each in 50 mL conicals.

prepare 3x conicals for 4 tissues	
16% PFA	10 ml
10x PBS	4 ml
RNAse free water	26 ml
total	40 ml

- 2 Cut freshly dissected tissue to small pieces for heart, try cutting slab, 2 cm x 1 cm x 0.4 cm thickness: 2 cm dimension from epicardium to endocardium.
- Place each piece in a 50 ml tube (or smaller if you can cut to smaller pieces) in 4% PFA leave at RT for 3 hrs.

  \*\* the sample should be completely covered.
- 5 Wash three times with equal volume (30mL) of 1x PBS to remove PFA leave a bit of extra liquid in the bottom between each wash.

## Sucrose gradient

- 6 Place the tissue in 10% Sucrose at RT and wait for it to sink in 50 mL conical, add 10 mL of sucrose wait for sample to sink to bottom minimum incubation time of 30 min remove supernatant for next bath.
- 7 Place the tissue in 20% Sucrose at RT and wait for it to sink.
- Place the tissue in 30% sucrose in 4°C ready for shipping. Make sure there is enough 30% sucrose solution in container e.g. if in 50 mL conicals, have 50 mL of 30% sucrose.

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