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③ Tissue Preparation for CLARITY

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Optical Clearing of Tissue

Human BioMolecular Atl...



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External link: http://wiki.claritytechniques.org/index.php/Solutions

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Protocol status: Working

We use this protocol and it's working

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Abstract

Tissue preparation for CLARITY includes fixation in 4% PFA, infusion with monomers in Hydrogel Solution, and clearing of lipids in Electrophoretic Tissue Clearing Solution.

Guidelines

The hydrogel solution components should be kept on ice during preparation to prevent polymerization. The thermal initiator is stable at low temperatures, but initiates polymerization at higher temperatures. To prevent polymerization, the hydrogel solution aliquots should be stored at -20°C until ready for use. However, the solution should be stable at 4°C for a few days and at room temperature for a couple hours.



Materials

Hydrogel Solution

(http://wiki.claritytechniques.org/index.php/Solutions)

Inq	gredient	Amou nt	Final Concentratio n	Purpose
)% ylamide	40 mL	4%	Hydrogel network monomer
	% Bis- ylamide	10 mL	0.05%	Small chemical crosslinker
	\-044 ator	1 g	0.25%	Polymerization thermal initiator
16 Par de	% aformaldehy	100 mL	4%	Biomacromolecule crosslinker
10	X PBS	40 mL	1X	Salt buffer
De wat	eionized er	210 mL	-	Aqueous solvent

Hydrogel Solution

Clearing Solution

Electrophoretic Tissue Clearing Solution C13001; logosbio.com, (https://logosbio.com/tissue-clearing_3d-imaging/tissue-clearing/x-clarity)

Equipment:

Platform rocker at room temperature (PFA Incubation) Rocking Stage at 4°C (hydrogel solution incubation) Incubator Shaker/ at 37°C (hydrogel polymerization) Incubator/Shaker at 45°C (tissue clearing)



Safety warnings



Paraformaldehyde and acrylamide are both toxic; therefore, preparation and use of the hydrogel solution should be done in a fume hood.



- 1 1. PFA Incubation: Trimmed tissues are placed fixed in 4% PFA for 20-24 hours at room temperature on a slow platform rocker.
- 2 2. Hydrogel Incubation: Tissues are transferred to Hydrogel Solution in for approximately 5 days at 4°C. Tubes are kept on a 2-axis rocker for the duration.
- 3. Hydrogel Polymerization: On day 5 the tissue is placed in pre-warmed 37°C PBS and a layer of mineral oil to deter oxygen exchange from air within tube. The hydrogel within the tissue is polymerized at 37°C for 4 hours with gentle rocking.
- 4. Passive Clearing: Following polymerization the tissue is placed in Clearing Solution and kept at 45°C with gentle rocking until optically clear.