# Visium CytAssist Spatial Gene Expression for Fixed Frozen – Rehydration, H&E Staining, Imaging & Decrosslinking

#### Introduction

The Visium CytAssist Spatial Gene Expression for Fixed Frozen assay is designed to analyze mRNA in tissue sections derived from fixed frozen tissue samples. The Visium CytAssist instrument requires a glass slide with intact tissue sections as input. This protocol outlines rehydration, Hematoxylin & Eosin (H&E) staining, imaging, and decrosslinking of tissue for use with the 10x Genomics Visium CytAssist Spatial Gene Expression for Fixed Frozen assay. Rehydrated, stained, and decrosslinked tissue sections are inputs for the downstream Visium CytAssist Spatial Gene Expression workflow.

## **Additional Guidance**

Consult the Visium CytAssist Spatial Gene Expression for Fixed Frozen - Tissue Preparation Guide (CG000663) for complete information on fixing, cryopreserving, embedding, and sectioning of tissue blocks and placing sections on slides. Consult the Visium CytAssist Spatial Gene Expression Imaging Guidelines (CG000521) to verify imaging settings prior to starting this Demonstrated Protocol. After completing this Demonstrated Protocol (CG000662), proceed with the Visium CytAssist Spatial Gene Expression - User Guide (CG000495), Rev D or later.

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#### **Reagent Kits**

#### **Visium Spatial Gene Expression for FFPE Reagent Kits**

Refer to SDS for handling and disposal information

## Visium CytAssist Slide and Cassettes, 6.5 mm, 2 rxns PN-1000519

Visium CytAssist Slide and Cassettes, 6.5 mm 2 rxns PN-1000519		
Store at ambient temperature	#	PN
Visium Cassette, 8 port	1	3000811
Visium Tissue Slide Cassette*		
Visium CytAssist moveable gasket small (pre-assembled with translator)	2	3000814
Visium CytAssist moveable translator (pre-assembled with gasket)	2	3000816
Visium CytAssist moveable Cassette, frame	2	3000813
Visium CytAssist Slide Seals, 40 pack*	1	2000284
Visium CytAssist Spatial Gene Expression Slide v2, 6.5 mm	1	2000549

\*Only these items are used in this protocol.

Visium CytAssist Slide and Cassettes, 11 mm, 2 rxns PN-1000518

#### Visium CytAssist Slide and Cassettes, 11 mm 2 rxns PN-1000518 Store at ambient temperature # ΡN 3000812 Visium Cassette, 2 port 1 Visium Tissue Slide Cassette\* Visium CytAssist moveable gasket 2 3000815 large Visium CytAssist moveable Cassette, 2 3000813 frame Visium CytAssist Slide Seals, 40 pack\* 1 2000284 Visium CytAssist Spatial Gene Expression 1 2000701 Slide v2, 11 mm 10x

\*Only these items are used in this protocol.

10x

#### **Reagent Kits**

#### Visium Spatial Gene Expression for FFPE Reagent Kits

Refer to SDS for handling and disposal information

## Visium FFPE Reagent Kit v2 - Small PN-1000436

Enough reagent is provided for processing two 6.5 mm slides or one 11 mm slide.

PN-	<b>Visium FFPE Reagent Kit – Small</b> PN-1000436				
Stor	e at -20°C	#	PN		
$\bigcirc$	Amp Mix B	1	2000567		
	Extension Enzyme	1	2000389		
	Extension Buffer	1	2000409		
	RNase Enzyme	1	3000593		
	RNase Buffer B	1	2000551		
	Tissue Removal Enzyme	1	3000387		
	Tissue Removal Buffer B**	1	2000543		
	Tissue Removal Buffer Enhancer**	1	2000557		
	Decrosslinking Buffer*	1	2000566		
	TS Primer Mix B	1	2000537		
	Block and Stain Buffer	2	2000554		
			10)		

\*Only this reagent is used in this protocol.

\*\*These tubes may not be included in the kit. They are not used in this assay.

#### **10x Genomics Accessories**

Product	#	Kit and Part Number	Part Number (Item)
Low Profile Plate Insert	2	Visium CytAssist	3000823
10x Magnetic Separator	1	Reagent Accessory Kit 1000499	120250

## **Recommended Thermal Cyclers**

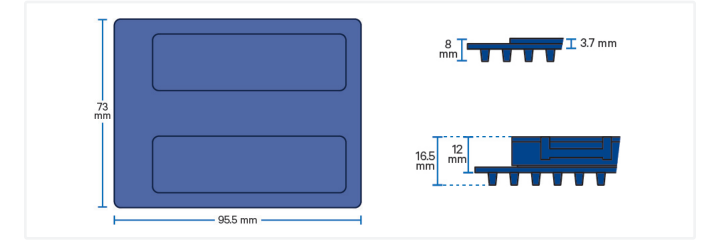
Supplier	Description	Part Number
Bio-Rad	C1000 Touch Thermal Cycler with 96-Deep Well Reaction Module	1851197
Eppendorf	MasterCycler Pro (discontinued)	North America 950030010 International 6321 000.019
	MasterCycler X50s	North America 6311000010
Thermo Fisher Scientific	Veriti 96-Well Thermal Cycler (discontinued)	4375786
Analytik Jena	Biometra TAdvanced 96 SG with 96-well block (silver, 0.2 mL) and gradient function	846-x-070-241

If using thermal cyclers other than the BioRad C1000, ramp rates should be adjusted for all the steps as described below:

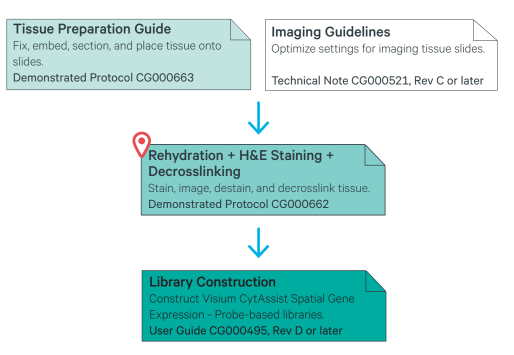
- Eppendorf MasterCycler X50s: 3°C/sec heating and 2°C/sec cooling
- Analytik Jena Biometra TAdvanced: 2°C/sec heating and cooling

Thermal cycler must be able to accommodate the Low Profile Plate Insert (also referred to as the Low Profile Thermocycler Adapter):

- Well depth: 4.5 mm
- Distance between block and heated lid: 12 mm
- Reaction block dimensions 95.5 x 73 mm



#### **Workflow Overview**



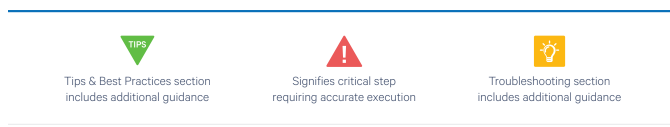
Visit the 10x Genomics Support website for the most current documentation.

### **Specific Reagents & Consumables**

For each item, a number of vendor options are listed. Choose item based on availability and preference. **Substituting materials may adversely affect system performance**.

ltem	Alternatives/Options	Vendor	Part Number
Ethanol	Ethyl Alcohol, 200 Proof	Millipore Sigma	E7023
	Ethanol absolute ≥99.5%, TechniSolv, pure (Europe Only)	VWR	83813.360DP
Eosin	Eosin Y-solution, Alcoholic	Millipore Sigma	HT110116
Hematoxylin	Hematoxylin Solution, Mayer's	Millipore Sigma	MHS16
Bluing Reagent	Bluing Reagent, Dako	Agilent	CS70230-2
1X PBS	Phosphate-Buffered Saline, 1X without calcium and magnesium, PH 7.4	Corning	21-040-CV
Glycerol	Glycerol Solution	Millipore Sigma	49781
	Glycerol	Acros Organics	327255000
0.1 N HCI	Hydrochloric Acid Solution, 0.1 N Or any equivalent HCl	Fisher Chemical	SA54-1
Coplin Jar/	Coplin Jar	VWR	100500-232
Staining Dishes	Staining Dishes	VWR	25608-906
<b>Green Marker,</b> Optional, if annotating slide	Sharpie Argyle Green Permanent Marker	Sharpie	1785396
Slide Holders	Slide Holders, 24-place	VWR	25608-868
Coverslips	Fisherbrand Cover Glasses: Rectangles	Fisher Scientific	12-544-EP
	Cover Glasses, Rectangular	VWR	16004-322
Pipettes	Pipet-Lite Multi Pipette L8-200XLS+	Rainin	17013805
	Pipet-Lite LTS Pipette L-2XLS+	Rainin	17014393
	Pipet-Lite LTS Pipette L-10XLS+	Rainin	17014388
	Pipet-Lite LTS Pipette L-20XLS+	Rainin	17014392
	Pipet-Lite LTS Pipette L-100XLS+	Rainin	17014384
	Pipet-Lite LTS Pipette L-200XLS+	Rainin	17014391
	Pipet-Lite LTS Pipette L-1000XLS+	Rainin	17014382
Wide Bore	Tips RT LTS 200UL FLW	Rainin	30389241
Pipette Tips	Tips RT LTS 1000UL FLW	Rainin	30389218
Pipette Tips	Tips LTS 200UL Filter RT-L200 FLR	Rainin	30389240
	Tips LTS 1ML Filter RT-L1000 FLR	Rainin	30389213
	Tips LTS 20UL Filter RT-L20 FLR	Rainin	30389226
Additional Mate	erials		
1000-ml Beakers	s (6)	-	-
Ultrapure/Milli-C	<b>Q Water,</b> gral Ultrapure Water System or equivalent	-	-

## **Tips & Best Practices**



## **General Reagent Handling**

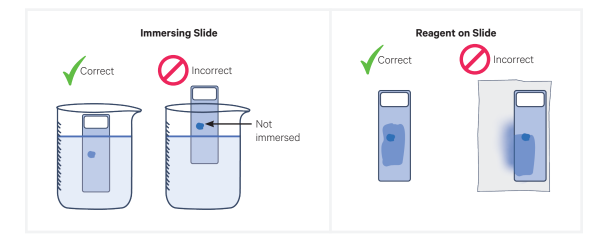
• Thoroughly mix reagents before use.

#### **Pipette Calibration**

• Follow manufacturer's calibration and maintenance schedules.

### **Slide Handling**

- · Always wear gloves when handling slides.
- DO NOT touch the tissue sections on slides.
- Keep tissue slides in a container on dry ice.
- Minimize exposure of slides to sources of particles and fibers.
- When immersing slides in prepared solutions and water, ensure that the tissue sections are completely submerged.
- Keep slides flat on the bench when adding reagents to the tissue.
- Ensure that no absorbent surface is in contact with the reagents on slides during incubation.



#### **Slide Incubation Guidance**

#### Incubation at a specified temperature

#### Incubation using a Thermal Cycler:

- Position a Low Profile Plate Insert (also referred to as Low Profile Thermocycler Adapter) on a thermal cycler that is set at the incubation temperature. Move Low Profile Thermocycler Adapter back and forth to ensure that it is seated properly.
- Ensure that the Low Profile Thermocycler Adapter is in contact with the thermal cycler surface uniformly.
- When incubating a slide, position the slide on the Low Profile Thermocycler Adapter with the tissue surface facing up.
- Ensure that the entire bottom surface of the slide is in contact with Low Profile Thermocycler Adapter.
- When incubating a slide encased in a cassette, place the assembled unit on the Low Profile Thermocycler Adapter with the wells facing up. Cassettes should always be sealed when on the Low Profile Thermocycler Adapter.
- Allow Low Profile Thermocycler Adapter to cool before removing it from the thermal cycler.



#### Incubation at room temperature

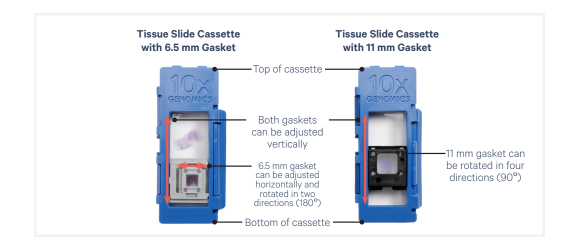
• Place the Visium CytAssist Tissue Slide Cassette on a flat, clean, non-absorbent work surface.

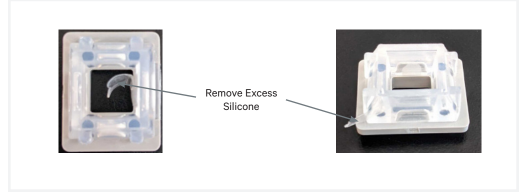
#### Visium CytAssist Tissue Slide Cassette

- The Visium CytAssist Tissue Slide Cassette encases the slide and creates a leakproof well for adding reagents on tissue slides.
- The cassette is a single use item.
- Gaskets are adjusted by the user to ensure that the tissue section or area of interest is encased in a well.
- Refer to Visium CytAssist Tissue Slide Cassette Assembly & Removal instructions for details.



- Prior to use, inspect the moveable gasket to ensure that the gasket perimeter and corners are free of excess silicone prior to assembly.
- Excess silicone should be safely removed with forceps or a pipette tip prior to assembly.
- Assembly should occur against a white background for easy tissue visualization during alignment.
- Practice assembly with a blank slide (75 x 25 x 1 mm).
- Place slides in the cassette only when specified.





## Visium CytAssist Tissue Slide Cassette Assembly





Exercise caution when handling slide edges to prevent injury.

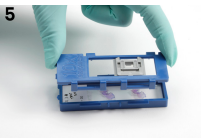
Break cassette into two halves by bending each half at the hinge until they snap apart



The 6.5 mm gasket can be adjusted horizontally and rotated in two directions (180°) while 11 mm gasket can be rotated in four directions (90°). Determine the appropriate configuration that allows the gasket to encompass the tissue area of interest.



Gently place top half of cassette over bottom half. DO NOT assemble together until Step 7.



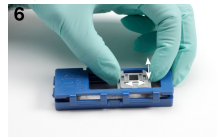
Place tissue slide into lower half of cassette with tissue facing up



Securely combine gasket with top half of cassette until the gasket snaps into place.



Adjust gasket such that gasket is over the tissue area of interest. The 6.5 mm gasket can be adjusted horizontally as well as vertically.



Apply even pressure on top of cassette until it clicks shut. Verify that clip is completely secured over hinges.

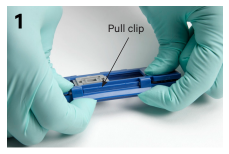


Turn cassette over and verify tissue area of interest is within gasket. DO NOT move gasket once cassette is closed. If necessary, open cassette and recenter gasket.



## Visium CytAssist Tissue Slide Cassette Removal

Pull clip up to detach upper and lower halves of cassette



Hold slide by the label and lift slide out from lower half

Open cassette by continuing to lift clip upward. If slide sticks to gasket, continue to apply even upward pressure to separate slide from gasket



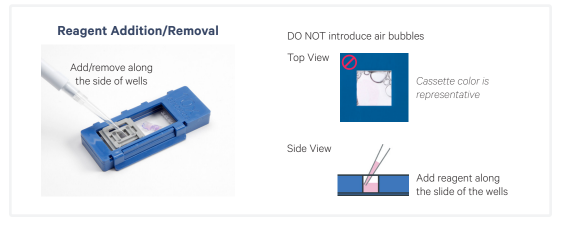


#### **Reagent Addition to Wells**

- Place the assembled slide in the Visium CytAssist Tissue Slide Cassette flat on a clean work surface.
- Dispense reagents along the side of the wells without touching the tissue sections and without introducing bubbles.



• Always cover the tissue section completely when adding reagents to the well. A gentle tap may help spread the reagent more evenly.



#### **Reagent Removal from Wells**

- Slightly tilt the cassette while removing the reagent.
- Place the pipette tip to the bottom edge of the wells.
- Remove reagents along the side of the wells without touching the tissue sections and without introducing bubbles.
- Remove all liquid from the wells in each step. To ensure complete removal, check the bottom of the well by tilting the cassette slightly. A meniscus at the bottom of the well will indicate the presence of liquid in the well. Repeat removal steps until no reagent remains.



#### Visium CytAssist Slide Seal Application & Removal

#### Application

- If applying a Visium Slide Seal to a Tissue Slide Cassette, the seal must be cut in half lengthwise.
- Four pre-cut seals per tissue section are needed for the entire Visium CytAssist Spatial Gene Expression assay.
- Cut the seal as shown in the image below. Ensure scissors are cleaned with a RNase decontamination solution.
- Place the CytAssist Tissue Slide Cassette flat on a clean work surface.
- Remove the back of the adhesive Visium Slide Seal.
- Align the Visium Slide Seal with the surface of the cassette and apply while firmly holding the cassette with one hand.
- Press on the Visium Slide Seal to ensure uniform adhesion.

#### Removal

- Place the CytAssist Tissue Slide Cassette flat on a clean work surface.
- Carefully pull Visium Slide Seal up and over from the edge while firmly holding the cassette.
- Ensure that no liquid splashes out of the wells.



#### Visium CytAssist Tested Slides

The following slides have been tested for use with the Visium CytAssist Tissue Slide Cassette and instrument.

Item	Length (mm)	Width (mm)	Thickness (mm)
Epredia Shandon ColorFrost Plus	75.0	25.0	1.0
Fisherbrand SuperFrost Plus	75.0	25.0	1.0
Sigma-Aldrich Poly Prep Slides	75.0	25.0	1.0
VWR SuperFrost Plus Micro Slide, Premium	75.0	25.0	1.0

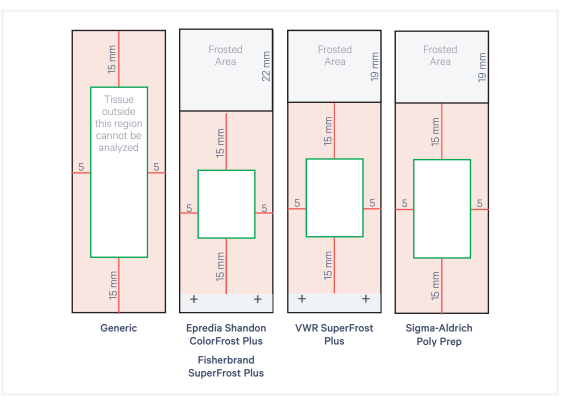
If unsure of slide part number, refer to the "generic" slide diagram below for general guidance (images not to scale). Diagrams for verifying that tissue sections are placed in the allowable area can also be found in the Visium CytAssist Quick Reference Cards - Accessory Kit (CG000548). The diagrams demonstrate allowable areas that are far enough away from frosted sections to not interfere with gasket closure during the CytAssist assay. Frosted sections include the opaque area of the slide as well as any etching or writing on the slide.



While slides are specified as being 25 mm x 75 mm, manufacturing tolerances may lead to dimensions that are too small or large to be compatible with 10x Genomics products. Tissue slide dimensions must be within 24.8 mm - 25.3 mm in width and 74.4 mm - 76.2 mm in length to fit the Visium CytAssist Tissue Slide Cassette.

Minimum slide dimensions: 24.8 x 74.4 mm

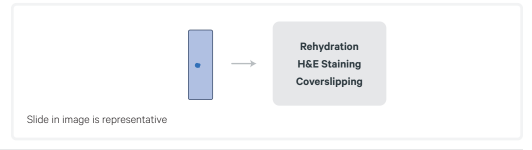
Maximum slide dimensions: 25.3 x 76.2 mm



# 1. Rehydration & H&E Staining

### **1.0 Overview**

This chapter provides guidance on rehydration and H&E staining of tissue slides containing fixed frozen tissue sections. Ensure that microscope settings have been verified and imaging programs have been created prior to starting this protocol. Consult the Visium Spatial Gene Expression Imaging Guidelines Technical Note (CG000521) for more information.



## **1.1 Preparation - Buffers**

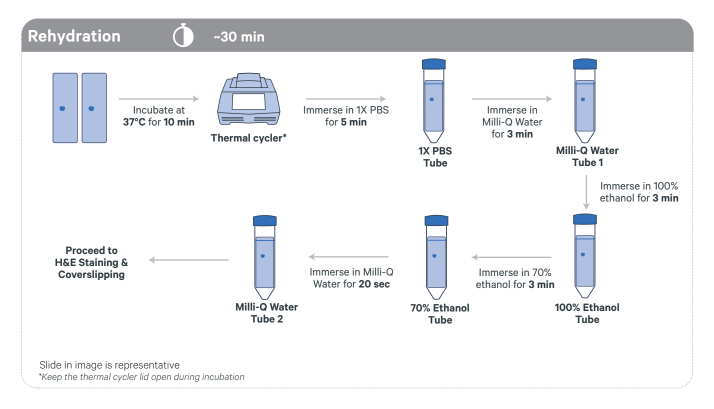
For	For Rehydration					
	Prepare fresh. If using 50-ml centrifuge tubes, two slides can be faced back to back (tissue facing out) per tube.					
Ite	ms	Preparation & Handling				
	100% Ethanol	Label one 50-ml centrifuge tube as 100% Ethanol Tube. Dispense 30 ml 100% ethanol. <i>Alternatively, use a coplin jar.</i>				
	70% Ethanol	Label one 50-ml centrifuge tube as 70% Ethanol Tube. Dispense 30 ml 70% ethanol. Alternatively, use a coplin jar.				
	Milli-Q or UltraPure Water	Label two 50-ml centrifuge tubes as Water Tube 1 and 2. Dispense 30 ml water in each. Alternatively, use a coplin jar.				
	1X PBS	Label one 50-ml centrifuge tube as 1X PBS Tube. Dispense 30 ml 1X PBS. Alternatively, use a coplin jar.				

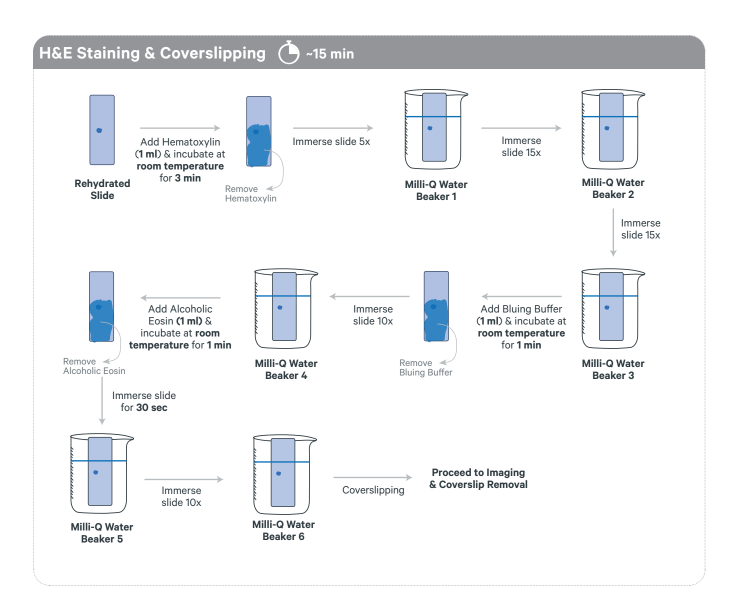


Reagent volume of 30 ml is sufficient for both 50-ml centrifuge tubes and coplin jars.

For	For H&E Staining					
Iter	ms	<b>Preparation &amp; Handling</b>				
	Milli-Q or UltraPure Water		Label six 1000-ml beakers as Water Beakers 1 – 6. Dispense 800 ml of water into each beaker. Dispensed volumes in each beaker can be used for two slides.			
For	Coverslipping					
Iter	ms	<b>Preparation &amp; Handling</b>				
	Mounting Medium	The dilution below is not necessary if stock glycerol is already at 85%. Invert or pipette gently with a wide-bore pipette tip to mix. Briefly centrifuge to remove bubbles.				
		Mounting Medium	Stock	Final	1X (µl)	2X+ 15% (µl)
		Glycerol	100%	85%	85.0	195.5
		Nuclease-free Water	100%	15%	15.0	34.5
		Total	-	-	100.0	230.0
	DO NOT let the attached coverslip dry. DO NOT use Cytoseal or nail polish for securing the coverslip.					

## **Protocol Overview**





#### **1.2 Rehydration**

- **a.** Place a Low Profile Thermocycler Adapter on a thermal cycler and preheat thermal cycler to **37°C.**
- **b.** Retrieve slide containing tissue from **-80°C** and place on dry ice.
- **c.** Place slide on the Low Profile Thermocycler Adapter with the tissue side facing up and incubate **10 min** at **37°C.** DO NOT close the thermal cycler lid.



Ensure tissue is dry before proceeding to next step. Damp tissue may lead to detachment. DO NOT exceed 20 min incubation time.



**d.** Remove slide from thermal cycler. Gently immerse slide in the 1X PBS Tube and incubate for **5 min**.

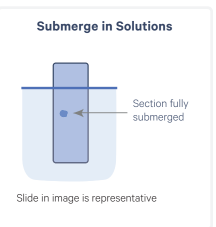


When immersing slides in solutions, ensure that the tissue sections are completely submerged.

- e. Gently immerse slide in the Milli-Q Water Tube 1 and incubate for **3 min**.
- **f.** Gently immerse slide in the 100% Ethanol Tube for **3 min**.
- **g.** Gently immerse slide in the 70% Ethanol Tube for **3 min**.
- **h.** Gently immerse slide in the Milli-Q Water Tube 2 for **20 sec**.
- i. Proceed **immediately** to H&E Staining & Coverslipping.



To ensure even staining, DO NOT let the slides dry.



#### 1.3 H&E Staining

- **a.** Place slide on a flat, clean, non-absorbent work surface. Some residual droplets may remain.
- **b.** Add **1 ml** Hematoxylin per slide to uniformly cover the entire tissue section.
- c. Incubate 3 min at room temperature.
- **d.** Discard reagent by draining and/or holding slide at an angle with the bottom edge in contact with a laboratory wipe.



- e. Immerse slide 5x in Water Beaker 1.
- f. Immerse slide 15x in Water Beaker 2.
- g. Immerse slide 15x in Water Beaker 3.
- **h.** Wipe excess liquid from the back of the slide without touching the tissue section.
- i. Place slide on a flat, clean, non-absorbent work surface. Some droplets may remain.
- **j.** Add **1 ml** Bluing Buffer per slide to uniformly cover the entire tissue section. Incubate **1 min** at **room temperature**.
- **k.** Discard reagent by draining and/or holding slide at an angle with the bottom edge in contact with a laboratory wipe.
- **1.** Immerse slide 10x in Water Beaker 4.
- **m.** Wipe excess liquid from the back of the slide without touching the tissue section. Place slide on a flat, clean, non-absorbent work surface. Some droplets may remain.
- n. Add 1 ml Eosin per slide to uniformly cover the entire tissue section. Incubate
   1 min at room temperature. DO NOT use diluted Eosin.
- **o.** Discard reagent by draining and/or holding slide at an angle with the bottom edge in contact with a laboratory wipe.
- p. Immerse slide for 30 sec in Water Beaker 5.
- **q.** Immerse slide 10x in Water Beaker 6.



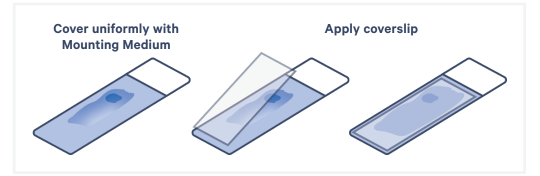
DO NOT air dry the slides.

#### **1.4 Coverslipping**

**a.** Place slide on a flat, clean, non-absorbent work surface. Some residual droplets may remain.

Wipe excess liquid from the back of the slide without touching the tissue section.

- **b.** Using a **wide-bore** pipette tip, add **100 µl** Mounting Medium to uniformly cover the entire tissue section.
- **c.** Apply the coverslip at an angle on one end of the slide. Slowly lower the coverslip, without introducing bubbles. Allow Mounting Medium to spread and settle.
- **d.** If needed, remove any large excess of Mounting Medium by carefully wicking away from the edge of the coverslip with a laboratory wipe. Be careful not to move the coverslip and disturb the tissue.



STOP

e. Once coverslipping is complete, **immediately** proceed with imaging or store slide laying flat in a squared slide mailer or tissue slide book at **4°C** in the dark for up to **two weeks**. Ensure multiple slides do not come in contact with one another. Storage can be done before or after imaging. DO NOT exceed two weeks of storage time.



# 2. Tissue Imaging

## 2.0 Overview

This chapter provides guidance on imaging tissue slides containing H&E stained fixed frozen tissue sections and coverslip removal.



## 2.1 Imaging System Recommendations

The following table shows imaging systems used by 10x Genomics in the development of this protocol. Any equivalent imaging system can be used as an alternative.

Supplier	Model	Configuration				
Thermo Fisher Scientific	EVOS M7000	Inverted				
Leica	Aperio Versa 8	Upright				
	Leica DMi8	Inverted				
MetaSystems	Metafer	Upright				
Nikon	Nikon Eclipse Ti2	Inverted				
BioTek	Cytation 7	Inverted or Upright				
Keyence	Keyence BZX800	Inverted				
Olympus	VS200	Upright				
Zeiss	Imager.Z2	Upright				
Brightfield Recommended Conf	Brightfield Recommended Configuration					
Color camera (3 x 8 bit, 2,424 x 2,424 pixel resolution)						
White balancing functionality						
2.18 µm/pixel minimum capture resolution						
Exposure times 2-10 milli sec						

## 2.2 Imaging

- **a.** Image each tissue section individually at the desired magnification using brightfield imaging settings.
- **b.** Consult the Visium CytAssist Spatial Gene Expression Imaging Guidelines Technical Note (CG000521) for additional information.
- c. After imaging, proceed immediately to Coverslip Removal or store slide laying flat in a squared slide mailer or tissue slide book at 4°C in the dark for up to two weeks. Storage can be done before or after imaging. DO NOT exceed two weeks of storage time.

#### 2.3 Coverslip Removal

STOP

- **a.** Dispense **800 ml** Milli-Q Water in a beaker. Up to 10 slides may be processed using this beaker.
- **b.** Immerse slide sideways/horizontal in the beaker containing **800 ml** water with the coverslipped surface fully sideways.
  - ImmerseHoldCoverslip detachesImmerse</td





To avoid damaging the tissue sections or causing tissue detachment, DO NOT move the slide up and down, shake forcibly, or manually move the coverslip.

- d. Gently immerse slide 30x in water to ensure all Mounting Medium is removed.
- e. Wipe the back of the slide with a laboratory wipe. Place on a flat, clean, nonabsorbent work surface and air dry for a minimum of **5 min** until the tissue is mostly dry. DO NOT exceed **20 min**.

f. Incubate slide on the Low Profile Thermocycler Adapter with the thermal cycler lid open for 3 min at 37°C to dry the slide.



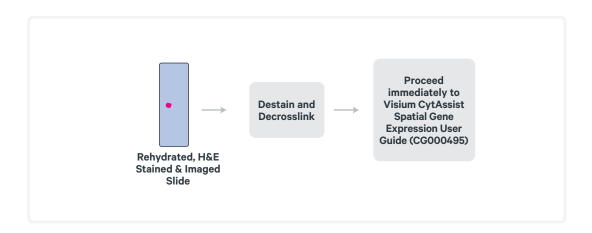
g. Proceed immediately to Decrosslinking.

## 3. Decrosslinking

### 3.0 Overview

This chapter provides guidance on performing decrosslinking to release RNA that was sequestered by formalin fixation from rehydrated and H&E stained tissues. Ensure that the coverslip is removed and tissue section is dry before destaining and decrosslinking.

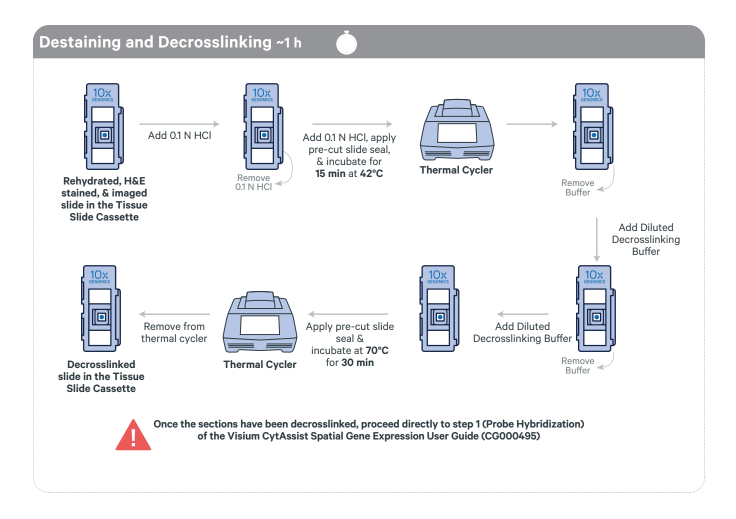
Once sections have been decrosslinked, step 1 (Probe Hybridization) of the Visium CytAssist Spatial Gene Expression User Guide (CG000495) should be immediately performed.



## **3.1 Preparation - Buffers**

Item	ns						
		Items Preparation & Handling					
	0.1 N HCl	If necessary, prepare 0.1 N HC	l using n	uclease-f	ree wate	er.	
	Decrosslinking	Thaw Decrosslinking Buffer a briefly after preparing Diluted Store excess stock buffer at 4	Decross			x and centri	fuge
		Diluted Decrosslinking	Stock	Final	1X (µl)	2X+ 10% (µl)	4X+ 10% (μl)
	ł	Decrosslinking Buffer	10X	1X	25	55	110
		Nuclease-free Water	-		225	495	990
		Total	-	-	250	550	1,100
		11 mm Gaskets Diluted Decrosslinking Buffer	Stock	Final	1X (µl)	2X+ 10% (µl)	4X+ 10% (µl)
		Decrosslinking Buffer	10X	1X	50	110	220
		Nuclease-free Water	-		450	990	1,980
		Total	-	-	500	1,100	2,200

#### **Protocol Overview**



### **3.2 Destaining**

■ denotes volumes for 6.5 mm gaskets and ▲ denotes volumes for 11 mm gaskets

**a.** Place a Low Profile Thermocycler Adapter in the thermal cycler. Prepare the thermal cycler with the following incubation protocol and start the program.

Lid Temperature	Reaction Volume	Run Time
42°C (lid may be set to lowest setting if instrument does not enable 42°C)	100 µl	15 min
Step	Temperature	Time
Pre-equilibrate	42°C	Hold
Destaining	42°C	00:15:00
Hold	22°C	Hold

b. Place the slide in the Visium CytAssist Tissue Slide Cassette.



See Tips & Best Practices for assembly instructions. Practice assembly with a blank slide.

- **c.** Add ■150 µl or ▲300 µl 0.1 N HCl along the side of the wells to uniformly cover the tissue sections, without introducing bubbles. Tap cassette gently to ensure uniform coverage.
- d. Remove HCl from the wells.



- e. Add ■100 µl or ▲200 µl 0.1 N HCl along the side of the wells to uniformly cover the tissue sections, without introducing bubbles. Tap cassette gently to ensure uniform coverage.
- **f.** Apply pre-cut slide seal on cassette and place the cassette on the Low Profile Thermocycler Adapter at **42°C**.



g. Close the thermal cycler lid. Skip Pre-equilibrate step to initiate Destaining.

**h.** Remove the cassette from the Low Profile Thermocycler Adapter and place on a flat, clean work surface. Some color remaining in the tissue after Destaining is normal.



### **3.3 Decrosslinking**

**a.** Prepare the thermal cycler with the following incubation protocol and start the program.

Lid Temperature	Reaction Volume	Run Time
70°C	100 µl	30 min
Step	Temperature	Time
Pre-equilibrate	70°C	Hold
Decrosslinking	70°C	00:30:00
Cooling	22°C	00:10:00
Hold	22°C	Hold

**b.** Peel back slide seal and using a pipette, remove all the HCl from the well corners.



- c. Add ■150 µl or ▲300 µl Diluted Decrosslinking Buffer along the side of the wells.
- **d.** Remove Diluted Decrosslinking Buffer from the wells.
- e. Add ■100 µl or ▲200 µl Diluted Decrosslinking Buffer along the side of the wells.
- **f.** Re-apply slide seal on the cassette and place the cassette on the Low Profile Thermocycler Adapter at **70°C**.
- g. Close the thermal cycler lid. Skip Pre-equilibrate step and initiate Decrosslinking.



**h.** Proceed **immediately** to Visium CytAssist Spatial Gene Expression User Guide (CG000495).

## Troubleshooting

detachment. Refer to Visium CytAssist Tested Slides.         Bubbles <ul> <li>Avoid bubble formation during coverslipping. Introduction of bubbles can be mitigated by applying the coverslip at an angle and slowly lowering it on to the slide, allowing air to escape. Briefly centrifuge Mounting Medium to remove bubbles before use. Avoid introducing bubbles when pipetting Mounting Medium onto slide.           Bubbles              <ul> <li>Bubbles may cause blackening of tissue</li> <li>Bubbles may cause blackening of tissue</li> <li>Bide in image is representative</li> </ul>            Uneven Staining              <ul> <li>Ensure fresh staining reagents are applied to the tissue uniformly and adequate washes are performed. A gentle tap may help reagents spread uniformly. Consider filtering hematoxylin.</li> </ul>            Slide in image is representative    Slide in image is representative            Incorrect Staining              <ul> <li>Ensure that the correct staining protocol with fresh reagents was followed.</li> <li>Ensure that the correct staining protocol with fresh reagents was followed.</li> </ul></li></ul>		Notes
bubbles can be mitigated by applying the coversilp at an angle and slowly lowering it onto the slide, allowing air to escape. Briefly centritique Mounting Wedium to remove bubbles before use. Avoid introducing bubbles when pipetting Mounting Medium onto slide. <b>Subbles may cause blackening of tissue</b> Slide in image is representative Uneven Staining Uneven Staining Slide in image is representative Slide in image is representative	Tissue Detachment	
Slide in image is representative         Uneven Staining         • Ensure fresh staining reagents are applied to the tissue uniformly and adequate washes are performed. A gentle tap may help reagents spread uniformly. Consider filtering hematoxylin.         • Ensure fresh staining reagents are applied to the tissue uniformly and adequate washes are performed. A gentle tap may help reagents spread uniformly. Consider filtering hematoxylin.         • Ensure that he correct staining protocol with fresh reagents was followed.         • Ensure that the correct staining protocol with fresh reagents was followed.	Bubbles	bubbles can be mitigated by applying the coverslip at an angle and slowly lowering it onto the slide, allowing air to escape. Briefly centrifuge Mounting Medium to remove bubbles before use. Avoid introducing bubbles when pipetting Mounting
Slide in image is representative         Uneven Staining         • Ensure fresh staining reagents are applied to the tissue uniformly and adequate washes are performed. A gentle tap may help reagents spread uniformly. Consider filtering hematoxylin.         • Ensure fresh staining reagents are applied to the tissue uniformly and adequate washes are performed. A gentle tap may help reagents spread uniformly. Consider filtering hematoxylin.         • Ensure that he correct staining protocol with fresh reagents was followed.         • Ensure that the correct staining protocol with fresh reagents was followed.		Bubbles may cause blackening of tissue
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• Ensure that the correct staining protocol with fresh reagents was followed.	Uneven Staining	uniformly and adequate washes are performed. A gentle tap may help reagents spread uniformly. Consider filtering
was followed.		Slide in image is representative
Incorrect staining protocol (right image) may result in poor staining	Incorrect Staining Protocol	was followed.
		Incorrect staining protocol (right image) may result in poor stainin

Slide in image is representative

#### **Document Revision Summary**

Document Number	CG000662
Title	Visium CytAssist Spatial Gene Expression for Fixed Frozen – Rehydration, H&E Staining, Imaging & Decrosslinking Demonstrated Protocol
Revision	Rev A
<b>Revision Date</b>	March 2023

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