

User Guide | CG000749 | Rev B

Xenium In Situ Gene Expression

with Cell Segmentation Staining

For use with:

Xenium Slides & Sample Prep Reagents (2 slides, 2 rxns) PN-1000460

Xenium Decoding Consumables (1 run, 2 slides) PN-1000487

Xenium Pre-Designed Gene Expression Panel - (2 rxns)*

Xenium Add-on Custom Gene Panel - (4 & 16 rxns)*

Xenium Standalone Custom Gene Panel - (4 & 16 rxns)*

Xenium Cell Segmentation Add-on Kit - (2 rxns) PN-1000662

Xenium Instrument Accessory Kit Module A PN-1000530

Xenium Slide, Kit, 4 Slides PN-1000659

Xenium Slide Kit, 16 Slides PN-1000660

*Refer to the 10x Genomics Support Website for the most current list of available panels and part numbers.

Notices

Document Number

CG000749 | Rev B

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Document Revision Summary

Document Number

CG000749

Title

Xenium In Situ Gene Expression with Cell Segmentation Staining User Guide

Revision

Rev B

Revision Date

September 13, 2024

Description of Changes

- Added storage temperature to Reagent Kits (pages 7-11).
- Added Xenium Cell Segmentation Add-on Kit (2 rxns), PN-1000662 to Reagent Kits list on cover and page 10.
- Updated Tips & Best Practices for Xenium Cassette wash and storage conditions (pages 30 and 31).
- Added guidance about cracked or broken slides on page 51.
- Updated thawing guidance for Xenium Amplification mix on page 59.
- Updated preparation guidance for Xenium Multi-Tissue Stain Mix on page 66.
- Updated Cassette Lid guidance during Cell Segmentation Staining (pages 66 and 69).
- Added guidance about keeping slides assay-specific on page 75.
- Updated PBS-T wash step formatting throughout.
- Updated for general minor consistency of language, format, and terms throughout.

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Introduction

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Reagent Kits

Xenium In Situ Gene Expression Reagent Kits

Refer to SDS for handling and disposal information.

Xenium Slides & Sample Prep Reagents - (2 Slides, 2 rxns) PN-1000460

Xenium Slides & Sample Prep Reagents (2 slides, 2 rxns), PN-1000460 Shipped in dry ice Store at -20°C				
		#	PN	
	Xenium Probe Hybridization Buffer	1	2000390	
0	Xenium Post Hybridization Wash Buffer	1	2000395	
	Xenium Ligation Buffer	1	2000391	
	Xenium Ligation Enzyme A	1	2000397	
	Xenium Ligation Enzyme B	1	2000398	
	Xenium Amplification Mix	1	2000392	
	Xenium Amplification Enzyme	1	2000399	
0	Reducing Agent B	1	2000087	
•	Xenium Autofluorescence Mix	1	2000753	
•	Xenium FFPE Tissue Enhancer*	1	2000798	
•	Xenium Nuclei Staining Buffer	1	2000762	
	Perm Enzyme B	1	3000553	
	Xenium Slides (2 pack)	1	3000941	
			10x	

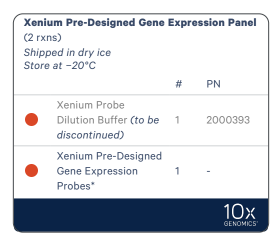
All items, except Xenium FFPE Tissue Enhancer (PN-2000798) and Perm Enzyme B (PN-3000553), are needed for this workflow.

Xenium Decoding Consumables - (1 Run, 2 Slides) PN-1000487

Xenium Decoding Consumal (1 run, 2 slides), PN-1000487 Shipped at ambient temp Store at ambient temp	bles	
	#	PN
Xenium Cassette Kit (2 cassettes + 16 lids)	1	1000566
Extraction Tip	1	2000757
Pipette Tips	1	3000866
Xenium Buffer Cap	4	3000949
Xenium Objective Wetting Consumable	1	2000749
Deionized Water (bottle)	1	3001198
Xenium Sample Wash Buffer A (bottle)	1	3001199
Xenium Sample Wash Buffer B (bottle)	1	3001200
Xenium Probe Removal Buffer (bottle)	1	3001201
		10x

Only the Xenium Cassette Kit (2 cassettes + 16 lids) is needed for this workflow.

Xenium Pre-Designed Gene Expression Panel - (2 rxns)



Example Xenium Pre-Designed Gene Expression Panel kit. Pre-designed panel kits contain one or two tubes - Xenium Probes with or without Xenium Probe Dilution Buffer.

*Refer to the 10x Genomics Support Website for the most updated list of available panels and part numbers.



Use TE Buffer instead of Xenium Probe Dilution Buffer (to be discontinued) without any impact on assay performance. Relevant guidance is provided in the workflow steps.

Xenium Add-on Custom Gene Panel - (4 & 16 rxns)



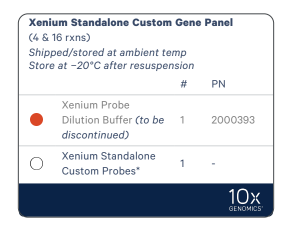
Example Xenium Add-on Custom Gene Panel kit. Add-on custom panel kits contain one or two tubes - Xenium Probes with or without Xenium Probe Dilution Buffer.

*Refer to the 10x Genomics Support Website for the most updated list of available panels and part numbers.



Use TE Buffer instead of Xenium Probe Dilution Buffer (to be discontinued) without any impact on assay performance. Relevant guidance is provided in the workflow steps.

Xenium Standalone Custom Gene Panel - (4 & 16 rxns)



Example Xenium Standalone Custom Gene Panel kit. Standalone custom panel kits contain one or two tubes - Xenium Probes with or without Xenium Probe Dilution Buffer.

*Refer to the 10x Genomics Support Website for the most updated list of available panels and part numbers.



Use TE Buffer instead of Xenium Probe Dilution Buffer (to be discontinued) without any impact on assay performance. Relevant guidance is provided in the workflow steps.

Xenium In Situ Gene Expression with Cell Segmentation Staining Reagent Kits

Reagent Kits	Part Number	Components	Component Part Number
Xenium Cell Segmentation Add-on Kit (2 rxns)	1000662	Xenium Cell Segmentation Staining Reagents (2 rxns)	PN-1000661
		Xenium Cell Segmentation Detection Reagents (2 rxns)*	PN-1000639
		*For use with the Xenium In Situ Gene Expression with Cell Segmentation Staining workflow on the Xenium Analyzer Instrument. Consult the Xenium Analyzer User Guide (CG000584) for complete information.	

The Xenium Cell Segmentation Add-on Kit (2 rxns) is only compatible with this workflow.

Xenium Cell Segmentation Staining Reagents - (2 rxns) PN-1000661

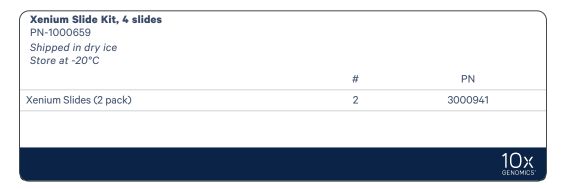
(2 rx	jents ns), PN-1000661		
Ship	ped in dry ice		
Store	e at -20°C		
		#	PN
•	Xenium Block and Stain Buffer	1	2001083
	Xenium Multi-Tissue Stain Mix	1	2000991
•	Xenium Stain Enhancer	1	2000992
	Xenium Cassette Insert	2	3001885
	Slide Seals (2 pack)	1	2001124

Xenium Instrument Accessory Kit, Module A PN-1000530

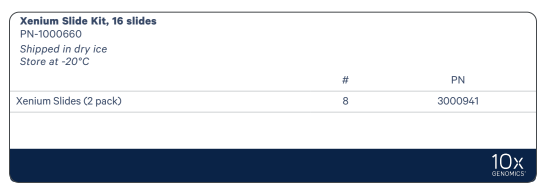
Xenium Instrument Accessory Kit Module A PN-1000530		
Shipped at ambient temp Store at ambient temp		
	#	PN
Waste Bottle	1	3000955
Xenium Waste Tip Tray	1	3000957
Xenium Thermocycler Adaptor	1	3000954
		10x GENOMICS

Only the Xenium Thermocycler Adaptor (PN-3000954) is needed for this workflow.

Xenium Slide Kit, 4 Slides PN-1000659



Xenium Slide Kit, 16 Slides PN-1000660



Recommended Thermal Cyclers

Supplier	Description	Part Number
Bio-Rad	C1000 Touch Thermal Cycler with 96-Deep Well Reaction Module (discontinued)	1851197
Analytik Jena	Biometra TAdvanced 96 SG with 96-well block (silver, 0.2 mL) and gradient function	846-x-070-241 (where x=2 for 230 V; 4 for 115 V; 5 for 100 V, 50-60 Hz)
VWR	Gradient thermal cycler, XT ⁹⁶ Gradient, with 96-well gradient block and standard lid	76452-153
Marshall Scientific	MJ Research PTC-200 Thermal Cycler (discontinued)	05434-05

Additional Kits, Reagents & Equipment

10x Genomics has tested all items listed below. These items perform optimally with the assay. **Substituting materials may adversely affect system performance.** For items with multiple options listed, choose option based on availability and preference. Refer to the manufacturer's website for regional part numbers.

Item	Description	Supplier	Part Number (US)
Plastics			
1.5 ml tubes	DNA LoBind Tubes, 1.5 ml	Eppendorf	022431021
	Low DNA Binding Tubes, 1.5 ml	Sarstedt	72.706.700
15 ml tubes	15 ml PP Centrifuge Tubes (or equivalent)	Corning	430791
50 ml tubes	Self-Standing Polypropylene Centrifuge Tubes (50 ml), sterile (or equivalent)	Corning	430921
Pipette tips	Tips LTS 20UL Filter RT-L20FLR (or equivalent)	Rainin	30389226
	Tips LTS 200UL Filter RT-L200FLR (or equivalent)	Rainin	30389240
	Tips LTS 1ML Filter RT-L1000FLR (or equivalent)	Rainin	30389213
Kits & Reagents			
Nuclease-free Water	Nuclease-free Water (not DEPC treated)	Thermo Fisher Scientific	AM9937
TE Buffer	TE Buffer, TRIS-EDTA, 1X Solution, pH 8.0 (nuclease-free)	Fisher Scientific	BP24731
PBS	PBS - Phosphate Buffered Saline (10X) pH 7.4, RNase-free	Thermo Fisher Scientific	AM9624
10% Tween-20	Tween 20 Surfact-Amps Detergent Solution (10% solution) (not 100% Tween diluted to 10%)	Thermo Fisher Scientific	28320
	10% Tween 20	Bio-Rad	1662404
Ethanol	Ethyl alcohol, Pure (200 Proof, anhydrous)	Millipore Sigma	E7023-500ML
Glycerol	Glycerol, 99.5%, for molecular biology, DNAse, RNAse and Protease free (optional, if storing slides long-term)	Acros Organics	327255000
Equipment			
Pipettes	Pipet-Lite LTS Pipette L-20XLS+	Rainin	17014392
	Pipet-Lite LTS Pipette L-200XLS+	Rainin	17014391
	Pipet-Lite LTS Pipette L-1000XLS+	Rainin	17014382
Forceps	Fisherbrand Curved Medium Point General Purpose Forceps (or any equivalent forceps)	Fisher Scientific	16-100-110

Item	Description	Supplier	Part Number (US)	
	Cassette Insert assembly)			
Mini centrifuge	VWR Mini Centrifuge (or any equivalent mini centrifuge)	VWR	76269-064	
Refrigerated Microcentrifuge	Eppendorf Microcentrifuge (or any equivalent microcentrifuge)	Eppendorf	5425 R	
Thermomixer	Eppendorf ThermoMixer C (or any equivalent Thermomixer)	Eppendorf	5382000023	
Thermoblock	Eppendorf SmartBlock 2.0 mL (or any equivalent Thermoblock)	Eppendorf	5362000035	
Additional Materials				
Waterbath (bead bath untested)				
Thermal Cycler (see Recommended Ther	rmal Cyclers)			
Ice bucket				
Vortex				
Lens-cleaning Paper or Lint-free Laborat	ory Wipes (for wiping thermal cycler adaptor)			
PCR Sealing Film (optional, if storing long	g-term)			
Ultrapure/Milli-Q Water for Water Bath from Milli-Q Integral Ultrapure Water S				

This list may not include some standard laboratory equipment.

Protocol Steps & Timing

Steps	Timing	Stop & Store
Day 1		
Step 1: Probe Hybridization		
1.1 Buffer Preparation 1.2 Custom Probe Preparation (optional) 1.3 Probe Hybridization	20 min 10 min 16-24 h (overnight)	
Day 2		
Step 2: Post Hybridization Wash		
2.1 Post Hybridization Wash	35 min	
Step 3: Ligation		
3.1 Ligation	~2 h	
Step 4: Amplification		
4.1 Amplification	~2 h	
4.2 Post Amplification Wash	15 min	
Step 5: Cell Segmentation Staining		
5.1 Buffer Preparation	20 min	
5.2 Block and Stain	~1 h + 16-24 h (overnight)	
Day 3		
5.3 Stain Enhancement	~1 h	
Step 6: Autofluorescence Quenching		
6.1 Autofluorescence Quenching 6.2 Nuclei Staining	30 min	4°C overnight (in the dark)
0.2 Nuclei Stailing	10 min	4°C overnight or 1 week (in the dark)*

^{*}After Nuclei Staining, the slides can also be stored long term in a cryoprotectant at -20°C as described in Xenium Cassette Storage on page 31.



Storing slides for more than one month risks ~5% decrease in sensitivity. These risks are dependent on many factors, including input tissue quality and how nuclease-free and microbe-free the workflow is.

Stepwise Objectives

Xenium In Situ Gene Expression assays RNA at the subcellular level by using targeted probes in formalin fixed & paraffin embedded (FFPE) or fresh frozen (FF) tissue sections. FFPE tissue sections placed on Xenium Slides are deparaffinized and decrosslinked as described in Xenium In Situ for FFPE - Deparaffinization & Decrosslinking (Demonstrated Protocol – CG000580). FF tissue sections placed on Xenium slides are fixed and permeabilized as described in Xenium In Situ for Fresh Frozen - Fixation & Permeabilization (Demonstrated Protocol – CG000581).

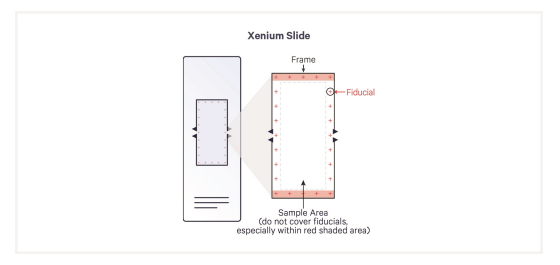
Pre-designed, add-on custom, or standalone custom probe panels are then added to the tissue. Each circularizable DNA probe contains two regions that hybridize to the target RNA and a third region that encodes a gene-specific barcode. The two ends of the probes bind the target RNA and are ligated to generate a circular DNA probe. Following ligation, the circularized probe is enzymatically amplified, generating multiple copies of the gene-specific barcode for each RNA target. Cell segmentation reagents added during an optional staining workflow allow for labeling of cell nuclei, membranes, and interiors that are inputs for automated morphology-based cell segmentation analysis.

Xenium slides containing FFPE or FF tissue sections are then loaded for imaging and analysis on the Xenium Analyzer instrument for high-throughput, automated in situ analysis. Fluorescently labeled oligos bind to the amplified DNA probes. Cyclical rounds of fluorescent probe hybridization, imaging, and removal generate optical signatures specific for each barcode, which are converted into a gene identity. Identified transcripts can be visualized using Xenium Explorer software.

This document outlines the protocol for generating Xenium In Situ Gene Expression data from FFPE and FF tissue sections placed on Sample Areas of a Xenium slide.

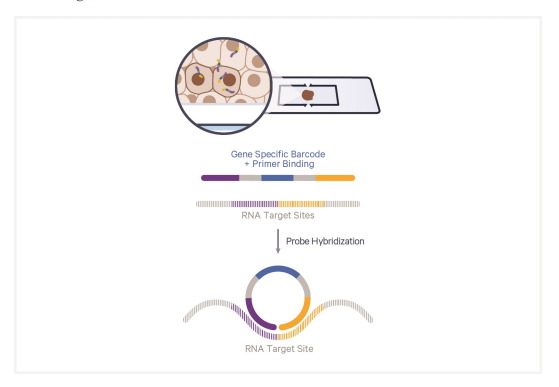
Xenium Slide

The Xenium slide has one Sample Area measuring $10.45 \times 22.45 \text{ mm}$ and is defined by a fiducial frame. The imageable area, measuring $12 \text{ mm} \times 24 \text{ mm}$, includes the area within the Sample Area + fiducial frame. FFPE or FF tissue sections are placed within the Sample Area for downstream processing and analysis.



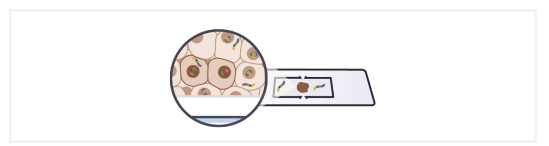
Step 1: Probe Hybridization

Pre-designed, add-on custom, or standalone custom probe panels are added to the FFPE or FF tissue sections. The DNA probes are flanked by two regions that independently hybridize to the target RNA and also contain a gene-specific barcode sequence. The probes hybridize to their complementary target RNA in an overnight incubation.



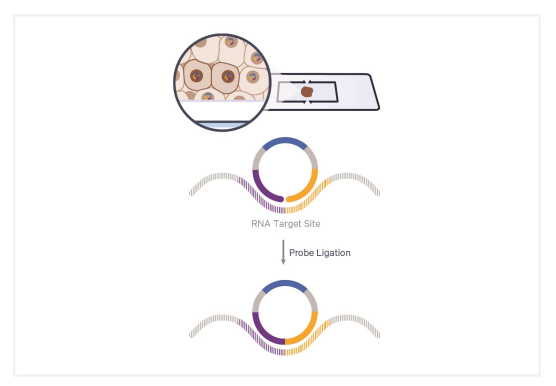
Step 2: Post Hybridization Wash

Excess, unbound probes are washed away in the post hybridization wash step.



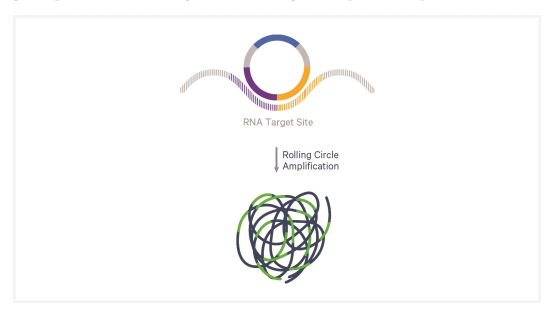
Step 3: Ligation

After removal of unbound probes, a ligase is added to seal the junction between the probe regions that have hybridized to RNA. Ligation of the probe ends on the targeted RNA region generates a circular DNA probe. This ligation ensures a unique level of probe specificity to the target region.



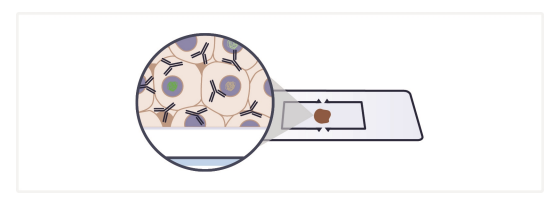
Step 4: Amplification

The ligation products are enzymatically amplified. Hundreds of copies of the gene-specific barcode are generated during the amplification process.



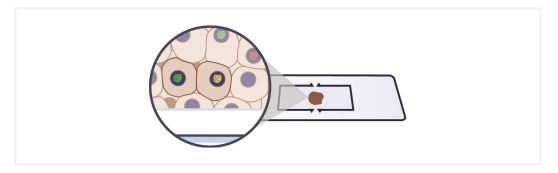
Step 5: Cell Segmentation Staining

Following a series of ethanol washes, blocking reagent is added to the tissue sections to reduce non-specific binding during staining. Cell segmentation antibodies bind their antigens in an overnight incubation, while excess antibodies are washed away in a post-incubation wash. Stain is then enhanced by the addition of Xenium Staining Enhancer reagent.



Step 6: Autofluorescence Quenching

Autofluorescence Quenching diminishes unwanted autofluorescence and enhances signal-to-noise ratio in the treated FFPE and FF tissue sections. Next, nuclei are stained with DAPI (derived from Xenium Nuclei Staining Buffer) to assist in identification of tissue or regions of interest during an instrument overview scan. Finally, tissue sections on Xenium slides assembled into Xenium Cassettes are loaded into the Xenium Analyzer for imaging and decoding.





Tips & Best Practices



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Tissue Detachment on Xenium Slides

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Icons



Tips & Best Practices section includes additional guidance



Signifies critical step requiring accurate execution



Troubleshooting section includes additional guidance

General Reagent Handling

- Fully thaw reagents at indicated temperatures. Thoroughly mix reagents before use.
- When pipette mixing reagents, unless otherwise specified, set pipette to 75% of total volume.
- Keep all enzymes and Master Mixes on ice during setup and use, unless otherwise stated.
- Promptly move reagents back to the recommended storage.

Pipette Calibration

• Follow manufacturer's calibration and maintenance schedules.

Probe Panel Handling

- 10x Genomics provides the following types of probe panels: pre-designed add-on custom, and standalone custom. Add-on custom panels are used to supplement pre-designed panels. Standalone custom probe panels are used alone and do not require pre-designed panels.
- Pre-designed probes are good for two Xenium slides. Add-on and Standalone custom probes are good for either four or sixteen Xenium slides.
- Add-on and Standalone custom probes are delivered lyophilized at room temperature and should be stored at -20°C upon resuspension.
- Custom probes must be resuspended prior to use. See Custom Probe Preparation (optional) on page 46 for more details.
- Record the Custom Panel Design ID and Slide Number before starting the workflow. This information is critical for identifying the correct electronic decode file when setting up the Xenium Analyzer in downstream steps.

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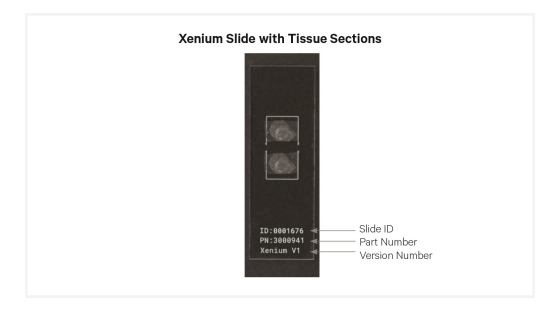
Probe Panel Storage & Shipping

- Pre-designed probe panels are ready to use and are shipped on dry ice. Pre-designed probes should be stored at -20°C upon receipt. The expiration date is listed on the kit label.
- Pre-designed probes are stable short-term at room temperature (<8 hours). Pre-designed probes should be discarded if left at room temperature for more than 8 hours.
- Add-on and Standalone custom probe panels are lyophilized and are shipped at room temperature. Store at -20°C upon receipt.
- Add-on and Standalone custom probes expire three years from the manufacture date in lyophilized form when stored at -20°C. Alternatively, custom probes expire one year from time of resuspension if stored in TE Buffer at -20°C or at kit expiration date (whichever comes first).
- Custom probes must be resuspended prior to use in TE Buffer, pH 8.0. After resuspension, they are stable for short-term at room temperature (<8 hours). Resuspended custom probes should be discarded if left at room temperature for more than 8 hours.
- Custom probes (in lyophilized form) accidentally left at room temperature for an extended period of time are likely stable. Contact support@10xgenomics.com if further assistance is needed.

Xenium Slide Handling

- Always wear gloves when handling slides.
- The bottom of the slide is indicated by the etched label, which should be readable when in the proper position.
- The tissue sections should always be placed within the Sample Area on etched label side of the slide.
- Hold the slide on the label. DO NOT touch the tissue sections or near fiducials.
- Minimize exposure of the slides to sources of particles and fibers.
- Keep the slide cassette flat on the bench when adding reagents to the Sample Area.
- Ensure that no absorbent surface is in contact with the reagents on the slide during incubation.
- When pipetting reagent onto a slide, avoid generating bubbles. Avoid pipetting directly onto the tissue.
- After aspirating reagent from a slide, pipette new reagent onto same slide before moving onto aspiration of second slide.

The Xenium Slide (PN-3000941) may not always include the suffix v1 in the etched label.



Processing a Single Xenium Slide

- Xenium reagent kits are sufficient for two reactions, and for optimal Xenium Analyzer throughput, two slides should be run at the same time.
- It is possible to perform the Xenium In Situ Gene Expression workflow with a single slide. To do this, ensure the following best practices are followed for optimal assay performance:
 - Assemble a mock Xenium Cassette using a blank slide and a cassette from the Xenium Cassette Kit (2 cassettes), PN-1000566.
 - Insert the blank slide into the Xenium Cassette. Cassettes should be assembled following the instructions in Troubleshooting for Cassette Assembly Failure on page 84.
 - Attach a Xenium Cassette Lid from the Xenium Cassette Kit (2 cassettes), PN-1000566 to the cassette containing the blank slide.

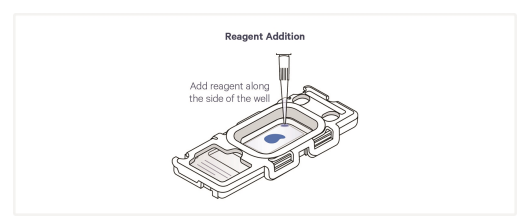


When handling an assembled cassette with the lid applied, always hold from the bottom of the cassette and not the lid.

 For all incubation steps with the thermal cycler lid closed, ensure the mock slide cassette is placed alongside the Xenium slide cassette containing tissue on the Thermocycler Adaptor.

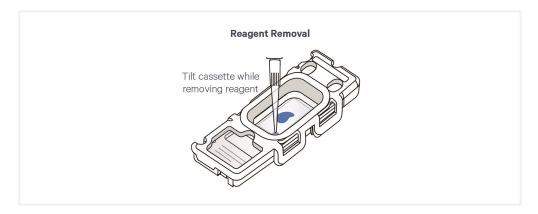
Reagent Addition to Wells

- Place assembled cassette flat on a clean work surface.
- Dispense and remove reagents along the side of the well without touching the tissue sections and without introducing bubbles.
- Always cover the Sample Area completely when adding reagents to the well. A gentle tap may help spread the reagent more evenly.



Reagent Removal from Wells

- Place assembled cassette flat on a clean work surface.
- Slightly tilt the cassette while removing the reagent.
- Place the pipette tip on the bottom edge of the well.
- Remove reagents along the side of the well without touching the tissue sections.
- Remove all liquid from the well in each step.

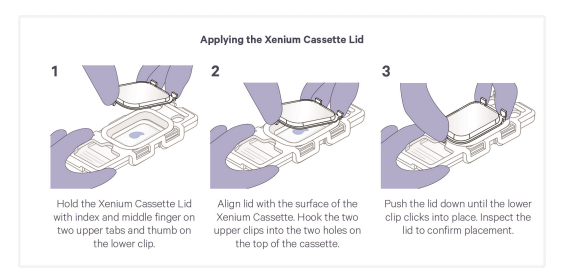


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Xenium Cassette Lid Application & Removal

• Work on a clean surface.

Application



Removal



Xenium Cassette Lids are a single use item and should be discarded after each use unless otherwise indicated. PBS-T washes DO NOT require sealing of the cassette with a lid.

Xenium Cassette Storage

- Store an assembled Xenium cassette with slide by applying a Xenium Cassette Lid or slide seal at the indicated stopping points listed throughout the protocol and as outlined in the Protocol Steps & Timing on page 16.
- Cassettes should always be stored hydrated with recommended reagent and stored at the recommended temperature.



- Short-term Storage (≤ 1 week):
 - ° Store in **1,000 \mul** PBS-T at **4**°**C** in the dark. Ensure that the slide is stored in microbe-free and nuclease-free conditions, with a Xenium Cassette Lid or slide seal applied to prevent evaporation.
- Long-term Storage (1 week 2 months):
 - Remove all PBS-T from the cassette well.
 - Add 1,000 μl 70% ethanol, incubate for 2 min at room temperature, remove the ethanol.
 - Add 1,000 μl 100% ethanol, incubate for 2 min at room temperature, remove the ethanol.
 - Add 1,000 μl 100% ethanol, incubate for 2 min at room temperature, remove the ethanol.
 - Remove slide from the cassette and place in a slide mailer containing
 10 ml or more cryoprotectant (30% Glycerol prepared in PBS) to fully submerge the slide. Clean the cassette as described in Xenium Cassette, Xenium Cassette Insert, and Lid Cleaning on page 87. Save for a subsequent instrument run.
 - Store at -20°C for up to 2 month.

When ready to use:

- Equilibrate the mailer with the slide to room temperature (takes ~30 min)
- Once completely thawed, rinse the mailer 3X with 10 ml PBS-T
- Remove the slide from the mailer, assemble in the cassette (as described on page 84)
- Add **1,000 µl** PBS-T to the cassette well.
- The above storage guidelines can also be used for storing Xenium cassettes with slides after a Xenium Analyzer run (or in the event of a run failure). Contact support@10xgenomics.com for more information.

Slide Incubation Guidance

Incubation at a specified temperature

• Position a Xenium Thermocycler Adaptor on a thermal cycler that is set at the incubation temperature. Ensure thermal cycler has reached appropriate temperature prior to starting incubation.



- Ensure that the Thermocycler Adaptor is fully inserted into the thermal cycler and is in contact with the thermal cycler block surface uniformly.
- When incubating a slide, position the slide on the Thermocycler Adaptor with the tissue side facing up.





- Ensure the Sample Area is aligned with the corresponding area on the Thermocycler Adaptor. DO NOT close the lid.
- When incubating a slide encased in a cassette, place the assembled unit on the Thermocycler Adaptor with the well facing up. Ensure the cassette is in complete contact with the Thermocycler Adaptor. The cassette should always be sealed with a Xenium Cassette Lid when on the Thermocycler

Adaptor unless indicated otherwise.



- For steps not using a Xenium Cassette Insert, where indicated, after thermal cycler incubation, at least 400 μl should be aspirated from cassette well. If less than 400 μl is recovered, contact support@10xgenomics.com.
- Before each incubation step, always inspect the adaptor and wipe it using a lint-free laboratory wipe. If debris or liquid is noticed, wipe the adaptor using 70% isopropanol, followed by wiping with a lint-free laboratory wipe. Failing to do so could result in slide cracking.

Tightening the thermal cycler lid

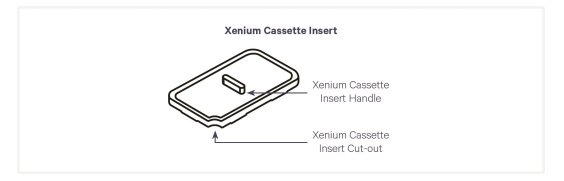
- Thermal cycler lid contact with the Xenium Cassette Lid is critical for assay performance.
- For thermal cyclers with adjustable lids, tighten the lid until an audible click is heard. Tightening past the click risks breaking the slide.

Incubation at room temperature

- Place the assembled cassette on a flat, clean, non-absorbent work surface.
- Ensure that no absorbent surface is in contact with the reagents on the slide/cassette during incubation.



Xenium Cassette Insert

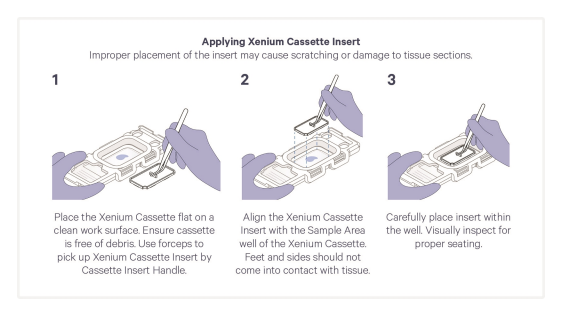


- The Xenium Cassette Insert sits in the Sample Area of the assembled Xenium Cassette and creates a flow cell in the well for adding reagents.
- Feet on the bottom of the insert help to prevent damage to the tissue sample by creating a gap between the sample and cassette.
- Place the insert in the cassette only when specified.
- Pick up the insert by using forceps on the Xenium Cassette Insert Handle. Forceps are needed for all handling steps. See Additional Kits, Reagents & Equipment on page 14 for recommended forceps.
- Reagents are loaded onto the sample by pipetting into the Xenium Cassette Insert Cut-out. The well is sealed with the application of Slide Seal.
- The insert is assembled manually into the cassette. See Xenium Cassette Insert Application & Removal on page 37 for more details.

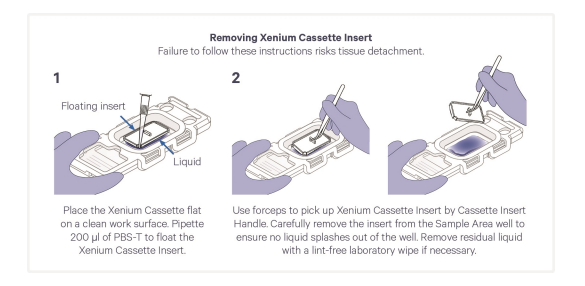
Xenium Cassette Insert Application & Removal

• Work quickly during application and removal steps to avoid drying out of tissue sections.

Application

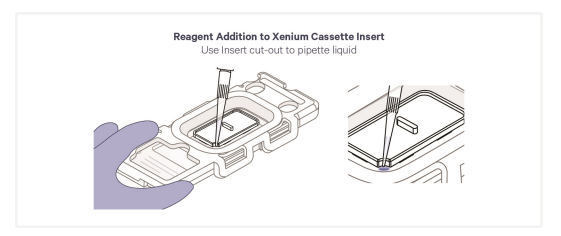


Removal



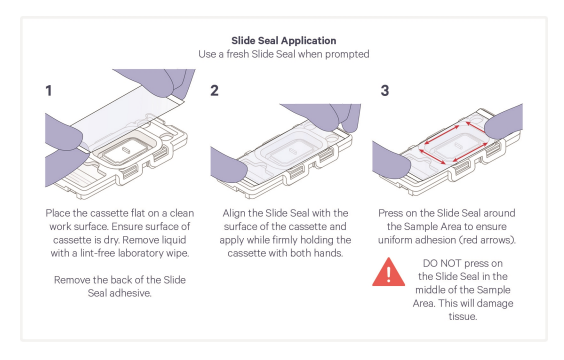
Reagent Addition to Xenium Cassette Insert

- Place the assembled cassette with insert flat on a clean work surface.
- Before executing the steps requiring an insert, retrieve all items and reagents needed for the steps to minimize chances of tissue drying.
- Slowly dispense reagents under insert by pipetting into the Xenium Cassette Insert Cut-out along the side of the well.
- Avoid dispensing reagent too quickly as this may lead to bubble formation.
- Gently support the pipette tip while dispensing reagent to provide more stability during this step.
- Allow reagent to fill the bottom of the insert.
- DO NOT dispense past first stop of the pipette to avoid introducing bubbles.

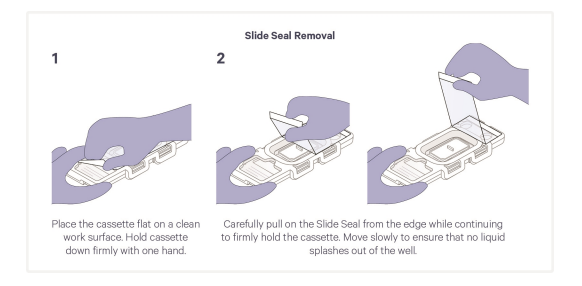


Slide Seal Application & Removal

Application



Removal



Xenium Cassette Insert Practice

Listed below are the practice steps that may be performed prior to using the Xenium Cassette Insert for Cell Segmentation Staining on page 63.

Obtain the following items:

- Xenium Cassette
- Superfrost Plus Slide
- Xenium Cassette Insert
- Slide Seal
- PBS-T



Cassettes may be obtained from previous runs and washed according to Xenium Cassette, Xenium Cassette Insert, and Lid Cleaning on page 87 instructions. Xenium Cassette Inserts can also be cleaned as per these instructions and reused. If an unused Xenium Cassette Insert is available, that can also be used for the practice.

Practice Steps	Guidance
1. Assemble Xenium Cassette	Cassette Assembly Failure on page 84
Assemble a Xenium Cassette and blank slide.	
2. Place Insert	Xenium Cassette Insert Application & Removal on
Place the Xenium Cassette Insert gently onto the	page 37
Xenium Cassette using forceps.	
3. Add Reagent	Reagent Addition to Xenium Cassette Insert on
Add 100 μl PBS-T through the Xenium Cassette	page 38
Insert Cut-out to uniformly cover the Sample	
Area, without introducing bubbles.	
4. Apply Slide Seal	Slide Seal Application & Removal on the previous
Apply PCR film to seal the well. Press down	page
around the Sample Area for uniform adhesion.	
5. Remove Slide Seal	Slide Seal Application & Removal on the previous
Remove the PCR film carefully	page
6. Remove Insert	Xenium Cassette Insert Application & Removal on
Add 200 µI PBS-T into Xenium Cassette Insert	page 37
Cut-out to float the insert. Remove the insert	
using forceps.	

Tissue Detachment on Xenium Slides



- Monitor section adhesion on Xenium slides throughout the workflow.
- Tissue detachment during the workflow can negatively impact performance. If observed, contact support@10xgenomics.com.
- For more information, refer to Troubleshooting.



Step 1:

Probe Hybridization

1.0 Get Started	43
1.1 Buffer Preparation	45
1.2 Custom Probe Preparation (optional)	46
13 Probe Hybridization	47

1.0 Get Started

Items			10x PN	Preparation & Handling	Storage
Equilibr	ate to ro	om temperature			
	•	Xenium Probe Hybridization Buffer	2000390	Thaw at room temperature for 15 min or until completely thawed. Check for precipitate and invert until clear. Maintain at room temperature after thawing.	-20°C
	•	Xenium Probe Dilution Buffer (to be discontinued)	2000393	Thaw at room temperature. Vortex and centrifuge briefly. Maintain at room temperature after thawing. Use TE Buffer instead of Xenium Probe Dilution Buffer without any impact on assay performance. See details below.	-20°C
	•	Xenium Pre- Designed Gene Expression Probes*	-	Thaw at room temperature. See Probe Hybridization for additional handling instructions.	-20°C
	0	Xenium Add-on Custom Probes*	-	Resuspend add-on custom probes according to Custom Probe Preparation. For additional handling instructions, see Probe Hybridization. Xenium v1 assay is not compatible with Xenium Prime addon custom priming oligos.	-20°C
	0	Xenium Standalone Custom Probes*	-	Resuspend standalone custom probes according to Custom Probe Preparation. For additional handling instructions, see Probe Hybridization. Xenium v1 assay is not compatible with Xenium Prime standalone custom priming oligos.	-20°C
Obtain					
		Assembled cassettes containing FFPE or FF tissue samples	-	Consult Xenium in Situ for FFPE - Deparaffinization & Decrosslinking (Demonstrated Protocol CG000580) or Xenium in Situ for Fresh Frozen - Fixation & Permeabilization (Demonstrated Protocol CG000581), respectively.	-
		Nuclease-free Water	-	-	Ambient
		10X PBS, pH 7.4	-	-	Ambient
		10% Tween-20	-	-	Ambient
		Heatblock or waterbath	-	Preheat to 95°C.	Ambient
		Xenium Cassette Lids (16 ct)	3001046	See Tips & Best Practices.	Ambient
		Xenium	3000954	See Tips & Best Practices.	Ambient

Items		10x PN	Preparation & Handling	Storage
	Thermocycler Adaptor			
	TE Buffer, TRIS- EDTA, 1X Solution, pH 8.0 (nuclease-free)	-	The pH of the stock solution should be 8.0. Readjusting the pH is NOT recommended.	Ambient

^{*}Thaw appropriate probe panels based on experimental needs. Refer to the 10x Genomics Support Website for the most updated list of available panels and part numbers.



Pre-heat appropriate equipment to: 37°C & 95°C.

Program a thermal cycler with the Probe Hybridization incubation protocol.

1.1 Buffer Preparation

Prepare the following buffers fresh before starting the Xenium In Situ Gene Expression workflow. The volumes of each buffer are sufficient for washes in all subsequent steps.

a. Prepare 1X PBS according to the table below before use and maintain at **room temperature.** Add reagents in the order listed and mix.

1X PBS	Stock	Final	1X+10% (ml)	2X+10% (ml)
Nuclease-free Water	-	-	18.0	36.0
10X PBS, pH 7.4	10X	1X	2.0	4.0
Total	-	-	20.0	40.0

b. Using 1X PBS from step 1.1a, prepare PBS-Tween Buffer (PBS-T) according to the table below before use and maintain at **room temperature.** Add reagents in the order listed. Invert gently to mix.

PBS-T	Stock	Final	1X+10% (ml)	2X+10% (ml)
1X PBS (prepared at Step 1.1a)	-	-	14.93	29.85
10% Tween-20	10%	0.05%	0.07	0.15
Total	-	•	15.0	30.0

1.2 Custom Probe Preparation (optional)

Proceed to Probe Hybridization, step 1.3, directly if using pre-designed probes only. Add-on and standalone custom probes are delivered lyophilized and must be resuspended before use. Resuspend add-on or standalone custom probes according to the instructions below before proceeding with Probe Hybridization.



Confirm the number of reactions provided for the add-on or standalone custom probes prior to resuspension.

- **a.** Centrifuge custom probe panel tube briefly.
- **b.** Resuspend lyophilized custom probes in TE Buffer according to the following table.

Custom Probe Resuspension	10x PN	TE Buffer (μl)
4 reactions/kit		
Xenium Add-on	varies	140
Xenium Standalone	varies	140
16 reactions/kit		
Xenium Add-on	varies	700
Xenium Standalone	varies	700

Xenium Probe Dilution Buffer may be used interchangeably with TE Buffer if sufficient volume is available.

- **c.** Replace the cap firmly. Vortex twice for **15 sec** each. Maintain at **room temperature** for **5 min**.
- **d.** Centrifuge custom probe panel tube briefly and maintain at **room temperature**.
- **e.** If custom probes are already resuspended, thaw at **room temperature** prior to starting Probe Hybridization.



Custom probes resuspended in Xenium Probe Dilution Buffer or TE Buffer can be stored at -20°C until the expiration date specified on the kit.

1.3 Probe Hybridization

Before starting this protocol, ensure that tissue sections have been appropriately deparaffinized and decrosslinked if working with FFPE tissues. Ensure that tissue sections have been appropriately fixed and permeabilized if working with fresh frozen tissues. Consult Xenium In Situ for FFPE - Deparaffinization & Decrosslinking Demonstrated Protocol (CG000580) or Xenium In Situ for Fresh Frozen - Fixation & Permeabilization Demonstrated Protocol (CG000581), respectively, for more information.



During reagent removal steps, ensure that **ALL the liquid is removed** from the wells. See Reagent Addition to Wells on page 29 for guidance.



Fluid on the Thermocycler Adaptor may indicate a reagent leak from the cassette. See Troubleshooting for more details.

- **a.** Obtain probes that have been thawed or equilibrated to **room temperature**. Briefly centrifuge the probe tube and remove an aliquot appropriate for the number of desired Xenium slides (34 μl/ slide).
- **b.** Immediately before use, preheat probes by incubating at **95°C** for **2 min** in a heatblock or thermal cycler, followed by **1 min** on **ice**. Maintain on ice.
- **c.** Prepare Probe Hybridization Mix according to the options below. Preparation instructions for each option can be found on the following page.
 - Pre-designed probe panels only
 - Add-on custom probe panels used with pre-designed probe panels
 - Standalone custom probe panels only

Prepare Probe Hybridization Mix shortly before use and maintain at **room temperature**. Add reagents in the order listed. Pipette mix and centrifuge briefly.

Option 1: Probe Hybridization Mix (pre-designed probe panels only)

Probe Hybridization Mix (pre-designed probe panels only)	10x PN	1X+5% (µl)	2X+5% (μl)
Xenium Probe Hybridization Buffer	2000390	315.0	630.0
TE Buffer	-	177.0	354.0
Xenium Pre-Designed Gene Expression Probes*	-	33.0	66.0
Total	-	525.0	1,050.0

Option 2: Probe Hybridization Mix (add-on custom probe panels used with pre-designed probe panels)

	Probe Hybridization Mix (add-on custom probe panels used with pre-designed probe panels)	10x PN	1X+5% (μl)	2X+5% (μl)
	Xenium Probe Hybridization Buffer	2000390	315.0	630.0
	TE Buffer	-	144.0	288.0
	Xenium Pre-Designed Gene Expression Probes*	varies	33.0	66.0
\bigcirc	Xenium Add-on Custom Probes*†	varies	33.0	66.0
	Total	-	525.0	1,050.0

Option 3: Probe Hybridization Mix (standalone custom probe panels only)

	Probe Hybridization Mix (standalone custom probe panels only)	10x PN	1X+5% (μl)	2X+5% (μl)
	Xenium Probe Hybridization Buffer	2000390	315.0	630.0
	TE Buffer	-	177.0	354.0
0	Xenium Standalone Custom Probes*†	varies	33.0	66.0
	Total	-	525.0	1,050.0

^{*}Refer to the 10x Genomics Support Website for the most updated list of available panels and part numbers.

[†]Custom probes resuspended in Xenium Probe Dilution Buffer or TE Buffer can be stored at -20°C until the expiration date specified on the kit. Record the Custom Panel Design ID and Slide Number before starting workflow. This information is critical for identifying the correct electronic decode file when setting up the Xenium Analyzer in downstream steps.

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

Lid Temperature	Reaction Volume	Run Time
50°C	100 μΙ	-
Step	Temperature	Time hh:mm:ss
Pre-equilibrate	50°C	Hold
Probe Hybridization	50°C	Overnight (16 - 24 h)
Hold	50°C	Hold

e. Retrieve the assembled Xenium Cassette and remove all the buffer.

- **f.** Add **500 \mu l** room-temperature Probe Hybridization Mix along the side of the well to uniformly cover the tissue sections, without introducing bubbles.
- **g.** Apply Xenium Cassette Lid on the cassette and place on the Thermocycler Adaptor on the pre-heated thermal cycler. Tightly close the thermal cycler lid until an audible click is heard.



Audible click will be heard in thermal cyclers with an adjustable lid (i.e. Bio-Rad C1000 Touch Thermal Cycler).

- **h.** Skip Pre-equilibrate step to initiate Probe Hybridization.
- i. After Probe Hybridization is complete, proceed to the next step.



Step 2:

Post Hybridization Wash

2.0 Get Started	51
2.1 Post Hybridization Wash	52

2.0 Get Started

Items			10x PN	Preparation & Handling	Storage
Equilibrate to room temperature					
	0	Xenium Post Hybridization Wash Buffer	2000395	Thaw at room temperature for 30 min or until thawed completely. Vortex and centrifuge briefly. Keep the buffer at room temperature after thawing.	-20°C
Obtain					
		PBS-T	-	Prepared at Step 1.1.	Ambient



Program a thermal cycler with Post Hybridization Wash incubation protocol.

2.1 Post Hybridization Wash

a. Remove the Xenium Cassette from the Thermocycler Adaptor and place on a flat, clean work surface.



DO NOT let the cassette cool down before proceeding to PBS-T washes.

DO NOT proceed with assay if slide is cracked or broken. Cracked or broken slides will result in assay failure.



Fluid on the Thermocycler Adaptor may indicate a reagent leak from the cassette. See Troubleshooting for more details.

- **b.** Remove the Xenium Cassette Lid. Using a pipette, remove all Probe Hybridization Mix from well corners. Discard old Cassette Lids.
- **c.** Using a pipette, remove all the buffer from well corners.

If processing two slides at a time, remove and add reagent from first slide before proceeding to second slide. Ensure tissue sections are always covered with reagent in between removal and addition steps to avoid drying out of tissue samples. At least 400 µl of liquid should be aspirated from each slide.

Two PBS-T washes (PBS-T prepared at step 1.1):

- d. Wash 1: Immediately add 500 μ l. Incubate for 1 min at room temperature. Remove all PBS-T.
- e. Wash 2: Add 500 µl PBS-T. Incubate for 1 min at room temperature.
- **f.** Prepare a thermal cycler with the following incubation protocol and start the protocol.

Lid Temperature	Reaction Volume	Run Time
37°C (lid may be set to the lowest temperature if the instrument does not enable 37°C)	100 μΙ	-
Step	Temperature	Time hh:mm:ss
Pre-equilibrate	37°C	Hold
Post Hybridization Wash	37°C	00:30:00
Hold	37°C	Hold

- g. Remove all PBS-T to complete wash 2.
- h. Add 500 µl Xenium Post Hybridization Wash Buffer to the well.
- i. Apply Xenium Cassette Lid on the cassette and place on the Thermocycler Adaptor on the pre-heated thermal cycler. Close the thermal cycler lid.
- j. Skip Pre-equilibrate step to initiate Post Hybridization Wash.



Start thawing Ligation reagents during Post Hybridization Wash incubation as outlined in Get Started on page 55.

k. After the Post Hybridization Wash is complete, immediately proceed to the next step.



Step 3:

Ligation

3.0 Get Started	55
3.1 Ligation	56

3.0 Get Started

Items			10x PN	Preparation & Handling	Storage
Equilibrate	to room	temperature	9		
		Xenium Ligation Buffer	2000391	Thaw at room temperature for 15 min or until completely thawed. Pipette MIX and centrifuge briefly. Maintain at room temperature after thawing.	-20°C
Place on ic	е				
	•	Xenium Ligation Enzyme A	2000397	Pipette mix and centrifuge briefly. Maintain on ice until ready to use.	-20°C
		Xenium Ligation Enzyme B	2000398	Pipette mix and centrifuge briefly. Maintain on ice until ready to use.	-20°C
Obtain					
		PBS-T	-	Prepared at Step 1.1.	Ambient



Program a thermal cycler with Ligation incubation protocol.

Step 3: Ligation 10xgenomics.com 55

3.1 Ligation

a. Prepare Ligation Mix shortly before using. Add reagents in the order listed. Pipette mix 10X and centrifuge briefly. Maintain on ice.

Ligation Mix	10x PN	1X+10% (μl)	2X+10% (µl)
Xenium Ligation Buffer	2000391	481.2	962.5
Xenium Ligation Enzyme A	2000397	13.8	27.5
Xenium Ligation Enzyme B	2000398	55.0	110.0
Total	-	550.0	1,100.0

- **b.** Remove the Xenium Cassette from the Thermocycler Adaptor and place on a flat, clean work surface.
- **c.** Remove the Xenium Cassette Lid. Using a pipette, remove all Xenium Post Hybridization Wash Buffer from the well. Discard used cassette lids.



If processing two slides at a time, remove and add reagent from first slide before proceeding to second slide. Ensure tissue sections are always covered with reagent in between removal and addition steps to avoid drying out of tissue samples. At least 400 µl of liquid should be aspirated from each slide.

Three PBS-T washes (PBS-T prepared at step 1.1):

- **d. Wash 1: Immediately** add **500 μl** PBS-T. Incubate for **1 min** at **room temperature**. Remove all PBS-T.
- **e. Wash 2:** Add **500 μl** PBS-T. Incubate for **1 min** at **room temperature**. Remove all PBS-T.
- f. Wash 3: Add 500 μl PBS-T. Incubate for 1 min at room temperature.
- **g.** Prepare a thermal cycler with the following incubation protocol. Place a Thermocycler Adaptor on the thermal cycler and start the program.

Lid Temperature	Reaction Volume	Run Tie
37°C (lid may be set to the lowest temperature if the instrument does not enable 37°C)	100 μΙ	-
Step	Temperature	Time hh:mm:ss
Pre-equilibrate	37°C	Hold
Ligation	37°C	02:00:00
Hold	37°C	Hold

Step 3: Ligation 10xgenomics.com 56

- **h.** Remove all PBS-T to complete wash 3.
- i. Add 500 μl Ligation Mix to the well.
- **j.** Apply a new Xenium Cassette Lid on the cassette and place on the Thermocycler Adaptor on the pre-heated thermal cycler. Close the thermal cycler lid.
- k. Skip Pre-equilibrate step to initiate Ligation.



Start thawing Amplification reagents (except enzymes) during Ligation incubation as outlined in Get Started on page 59.

1. After Ligation is complete, immediately proceed to next step.

Step 3: Ligation 10xgenomics.com 57



Step 4:

Amplification

4.0 Get Started	59
4.1 Amplification	60
4.2 Post Amplification Wash	62

4.0 Get Started

Item		10x PN	Preparation & Handling	Storage
Place on ice				
	Xenium Amplification Mix	2000392	Immediately after initiating Ligation step, thaw on ice. Vortex and centrifuge briefly. Ensure completely thawed before use, with no precipitate remaining.	-20°C
	Xenium Amplification Enzyme	2000399	Transfer to ice before use. Pipette mix and centrifuge briefly. Maintain on ice until ready to use.	-20°C
Obtain				
	PBS-T	-	Prepared at Step 1.1.	Ambient
	TE Buffer, TRIS- EDTA, 1X Solution, pH 8.0 (nuclease- free)	-	-	Ambient

4.1 Amplification

a. Prepare Amplification Master Mix on ice shortly before use. Add reagents in the order listed. Pipette mix 10X and centrifuge briefly. Maintain on ice.

Amplification Master Mix	10x PN	1X +10% (μl)	2X +10% (μl)
Xenium Amplification Mix (Thaw, vortex, centrifuge briefly before use)	2000392	495.0	990.0
Xenium Amplification Enzyme	2000399	55.0	110.0
Total	-	550.0	1,100.0

- **b.** Remove the Xenium Cassette from the Thermocycler Adaptor and place on a flat, clean work surface.
- **c.** Remove the Xenium Cassette Lid. Using a pipette, remove all the buffer from the well. Discard used cassette lids.

If processing two slides at a time, remove and add reagent from first slide before proceeding to second slide. Ensure tissue sections are always covered with reagent in between removal and addition steps to avoid drying out of tissue samples. At least 400 µl of liquid should be aspirated from each slide.

Three PBS-T washes (PBS-T prepared in step 1.1):

- **d. Wash 1: Immediately** add **500 μl** PBS-T. Incubate for **1 min** at **room temperature**. Remove all PBS-T.
- **e. Wash 2: Immediately** add **500 μl** PBS-T. Incubate for **1 min** at **room temperature**. Remove all PBS-T.
- f. Wash 3: Immediately add 500 μl PBS-T. Incubate for 1 min at room temperature.
- **g.** Prepare a thermal cycler with the following incubation protocol. Place a Thermocycler Adaptor on the thermal cycler and start the program.

Lid Temperature	Reaction Volume	Run Time
30°C (lid may be set to the lowest temperature if the instrument does not enable 30°C)	100 μΙ	-
Step	Temperature	Time hh:mm:ss
Pre-equilibrate	30°C	Hold
Amplification	30°C	02:00:00

h. Remove all PBS-T to complete wash 3.

- i. Immediately add $500 \mu l$ Amplification Master Mix to the well.
- **j.** Apply a new Xenium Cassette Lid on the Xenium Cassette and place on the Thermocycler Adaptor on the thermal cycler. Close the thermal cycler lid.
- k. Skip pre-equilibrate step to initiate Amplification.



Start thawing Block and Stain Buffer during Amplification incubation as outlined in Get Started on page 64.

1. After Amplification is complete, **immediately** proceed to the next step.

4.2 Post Amplification Wash

- **a.** Remove the Xenium Cassette from the Thermocycler Adaptor and place on a flat, clean work surface.
- **b.** Remove the Xenium Cassette Lid. Using a pipette, remove all Amplification Mix from the well.

Three TE Washes:

- c. Wash 1: Add 500 μ l TE Buffer to the well. Incubate 1 min at room temperature. Remove all TE buffer.
- **d.** Wash 2: Add 500 μl TE Buffer to the well. Incubate 1 min at room temperature. Remove all TE buffer.
- e. Wash 3: Add 500 μ l TE Buffer to the well (buffer removal will be in next step to complete wash 3).
- **f.** Apply a new Xenium Cassette Lid on the Xenium Cassette for Staining Buffer Preparation step. Maintain cassette at **room temperature** during buffer preparation.



Step 5:

Cell Segmentation Staining

5.0 Get Started	64
5.1 Buffer Preparation	65
5.2 Block and Stain	66
5.3 Stain Enhancement	69

5.0 Get Started

Item		10x PN	Preparation & Handling	Storage
Maintain on le	се			
	Xenium Block and Stain Buffer	2001083	Thaw at room temperature for 30 min. Vortex and centrifuge briefly. Maintain on ice.	-20°C
	Xenium Multi- Tissue Stain Mix	2000991	Prepare according to Block and Stain on page 66. Maintain on ice.	-20°C
	Xenium Stain Enhancer	2000992	To be prepared next day according to Stain Enhancement on page 69. Maintain on ice.	-20°C
Equilibrate to	room temperature			
	Xenium Cassette Insert	3001885	Remove from -20°C and equilibrate to room temperature.	-20°C
	Slide Seals	2001124	Remove from -20°C and equilibrate to room temperature.	-20°C
Obtain				
	Nuclease-free Water	-	-	Ambient
	Ethanol	-	-	Ambient
	1X PBS	-	Prepared at Step 1.1. To be obtained next day for Stain Enhancement.	Ambient
	PBS-T	-	Prepared at Step 1.1.	Ambient
	Forceps	-	-	Ambient
	Ice bucket	-	-	Ambient

5.1 Buffer Preparation

- a. Retrieve PBS-Tween (PBS-T) prepared in step 1.1.
- **b.** Prepare ethanol dilutions in 15-ml conical tubes using Nuclease-free Water. Vortex and centrifuge briefly. Maintain at room temperature.

	For Ethanol Washes				
Items		Preparation & Handling			
	100% Ethanol	Label one 15-ml conical tube as 100% ethanol. Add 15 ml 100% ethanol.			
	70% Ethanol	Label one 15-ml conical tube as 70% ethanol. Add 10.5 ml 100% ethanol and 4.5 ml Nuclease-free Water.			



Less 70% ethanol will be needed if not performing Cell Segmentation Staining. Adjust volumes accordingly, if desired.

c. Prepare 1X Diluted Xenium Block and Stain Buffer (for two slides) in a centrifuge tube for both blocking and staining steps. Add reagents in the order listed. Vortex and centrifuge briefly. Maintain on ice.

	1X Diluted Xenium Block and Stain Buffer (for 2 slides)				
Items		Stock	Final	Total Amount (µl)	
	Nuclease-free Water	-	-	990.0	
	Xenium Block and Stain Buffer	4X	1X	330.0	
	Total	-	-	1,320.0	

5.2 Block and Stain

- **a.** Retrieve the Xenium Cassette from the previous step and place on a flat, clean work surface.
- **b.** Remove the Xenium Cassette Lid. Using a pipette, remove all TE Buffer from the well. **Save the lid** for the next indicated steps.

If processing two slides at a time, remove and add reagent from first slide before proceeding to second slide.



DO NOT let tissue dry out during the ethanol wash steps and between removal of ethanol and addition of PBS-T. Left over ethanol volumes may be used for ethanol wash steps in Autofluorescence Quenching on page 73.

Four ethanol washes:

- **c. Wash 1:** Add **1,000 μl** <u>70%</u> ethanol. Incubate for **2 min** at **room temperature**. Remove the ethanol.
- d. Wash 2: Add 1,000 μ l 100% ethanol. Incubate for 2 min at room temperature. Remove the ethanol.
- e. Wash 3: Add 1,000 μl 100% ethanol. Incubate for 2 min at room temperature. Remove the ethanol.
- f. Wash 4: Add 1,000 μ l 70% ethanol. Incubate for 2 min at room temperature. Remove the ethanol.
- **g.** Immediately add 1,000 μl PBS-T. Incubate for 1 min at room temperature. Remove all PBS-T.
- **h.** Add **500 μl** 1X Diluted Xenium Block and Stain Buffer (from 5.1 Buffer Preparation on the previous page) to cassette for blocking.



Keep remaining volume for later indicated steps.

- **i.** Apply the previously used Xenium Cassette Lid on the Xenium Cassette and incubate for **1 hour** at **room temperature**.
- **j.** During incubation, prepare Xenium Multi-Tissue Stain Mix. Centrifuge tube for **5 sec**.
- **k.** Add **220 µl** 1X Diluted Xenium Block and Stain Buffer (from **5.1** Buffer Preparation on the previous page) to the Xenium Multi-Tissue Stain Mix tube.
- **l.** Pipette mix 15X with (pipette set to 175 μ l) and centrifuge briefly. Avoid bubbles if possible.

	Xenium Multi-Tissue Stain Mix	10x PN	1X Diluted Xenium Block and Stain Buffer (μΙ)
	Xenium Multi-Tissue Stain Mix	2000991	220.0

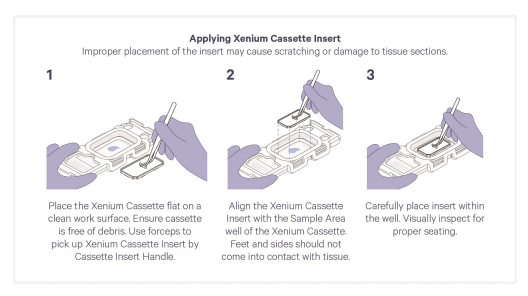


DO NOT add any additional antibodies to the Xenium Multi-Tissue Stain Mix as that can compromise assay performance.

- m. Incubate resuspended Xenium Multi-Tissue Stain Mix for 30 min at room temperature.
- n. Centrifuge Xenium Multi-Tissue Stain Mix for 10 min at 14,000 rcf at 4°C. Maintain on ice.
- o. Obtain forceps and Xenium Cassette Insert to prepare for application of the insert onto the cassette
- p. Remove the Xenium Cassette Lid and using a pipette, remove 1X Diluted Xenium Block and Stain Buffer from well corners. Save the lid for the next indicated steps. DO NOT let the tissue sections dry out.
- **q.** Gently place Xenium Cassette Insert onto the Xenium Cassette using forceps.



Leave second slide with 1X Diluted Xenium Block and Stain Buffer until the first cassette has been assembled with insert and Xenium Multi-Tissue Stain Mix has been added.

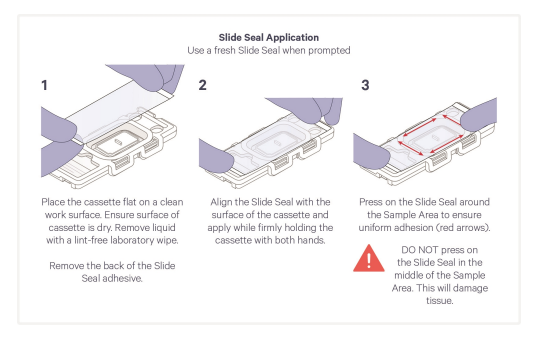


r. Using a pipette along the side of the tube (avoid touching pellet), withdraw 100 µl Xenium Multi-Tissue Stain Mix and add to the Xenium Cassette Insert using the cut-out to pipette solution under the Xenium Cassette Insert.



Pipette slowly into cut-out to avoid introducing bubbles as bubbles may impact staining. See Bubbles under Xenium Cassette Insert on page 82 for guidance on resetting the cassette insert if bubbles are present.

s. Carefully seal the cassette well using Slide Seal. Press down around the cassette well edges to ensure complete sealing between cassette and Slide Seal.



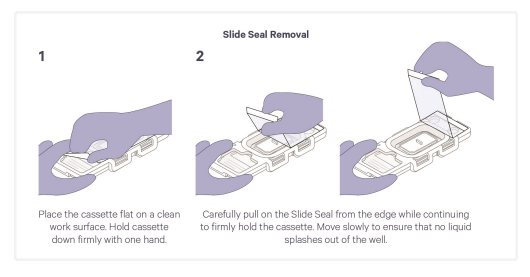
t. Incubate the Xenium Cassette overnight (16-24 h) at 4°C (in a refrigerator or incubator).

5.3 Stain Enhancement

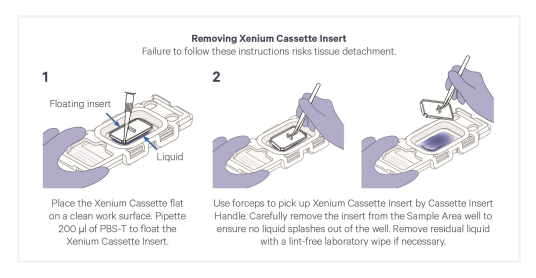
a. Following overnight incubation, prepare Xenium Stain Enhancer. Thaw at **room temperature** for **10 min.** Centrifuge briefly. Visually inspect the tube to ensure white powder is at the bottom of the tube before opening. Add **1,100 µl** 1X PBS. Pipette mix 5X and centrifuge for **5 sec**.

	Xenium Stain Enhancer	10x PN	1X PBS (μl)
	Xenium Stain Enhancer	2000992	1,100

b. Remove Slide Seal from Xenium Cassette. Carefully pipette **200** μ l PBS-T into Xenium Cassette Insert Cut-out to float the insert.



c. Carefully remove Xenium Cassette Insert with forceps. **Discard** used Xenium Cassette Inserts.



d. Using a pipette, remove all Xenium Multi-Tissue Stain Mix from well corners.

Three PBS-T Washes (PBS-T prepared in step 1.1):

- e. Wash 1: Add 500 µl PBS-T. Incubate for 1 min at room temperature. Remove all PBS-T.
- f. Wash 2: Add 500 μl PBS-T. Incubate for 1 min at room temperature. Remove all PBS-T.
- g. Wash 3: Add 500 µl PBS-T. Incubate for 1 min at room temperature. Remove all PBS-T.
- h. Add 500 μl resuspended Xenium Stain Enhancer to the well.
- i. Apply the previously used Xenium Cassette Lid on the cassette. Incubate at room temperature for 20 min.



Start thawing Autofluorescence Quenching reagents during Stain Enhancement as outlined in Get Started on page 72.

j. Remove the Xenium Cassete Lid. Using a pipette, remove all Xenium Stain Enhancer from well corners. Discard old Cassette Lids.

Two PBS-T Washes (PBS-T prepared in step 1.1):

- k. Wash 1: Add 500 µl PBS-T. Incubate for 1 min at room temperature. Remove all PBS-T.
- 1. Wash 2: Add 500 μl PBS-T. Incubate for 1 min at room temperature. Remove all PBS-T.
- m. Add **500 μl** PBS-T.
- **n.** After Cell Segmentation Staining is complete, proceed **immediately** to the next step.



Step 6:

Autofluorescence Quenching

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6.0 Get Started

Items			10x PN	Preparation & Handling	Storage		
Equilibrate to room temperature							
	•	Xenium Autofluorescence Mix	2000753	Thaw in a thermomixer (with 2.0-ml thermoblock) for 15 min at 37°C, 300 rpm with shaking. Cool to room temperature for 5 min. Vortex for 30 sec and centrifuge briefly. Alternatively, thaw in a waterbath	−20°C		
				for 15 min at 37°C. Cool to room temperature for 5 min. Vortex for 30 sec and centrifuge briefly.*			
	\bigcirc	Reducing Agent B	2000087	Thaw at room temperature. Vortex and centrifuge briefly.	-20°C		
	•	Xenium Nuclei Staining Buffer	2000762	Thaw at room temperature. Vortex and centrifuge briefly. Keep in the dark until ready to use.	-20°C		
Obtain							
		Nuclease-free Water	-	-	Ambient		
		1X PBS	-	Prepared at Step 1.1.	Ambient		
		PBS-T	-	Prepared at Step 1.1.	Ambient		
		100% Ethanol	-	-	Ambient		

^{*}If processing only one slide, remove the volume required (specified in the following section), and refreeze the remaining reagent in the tube at -20 °C.



Pre-heat appropriate equipment to: 37°C.

100% ethanol is needed for washes in addition to making 70% ethanol dilutions.

6.1 Autofluorescence Quenching

- a. Prepare the following for Autofluorescence Quenching:
 - **i. Prepare diluted Reducing Agent B.** Add reagents in the order listed and vortex to mix. Maintain at room temperature.

	Diluted Reducing Agent B	10x PN	Stock	Final	1X+10% (μl)	2X+10% (μl)
	1X PBS (prepared at Step 1.1)	-	-	-	544.5	1,089.0
\circ	Reducing Agent B	2000087	-	-	5.5	11.0
	Total	-	-	-	550.0	1,100.0

ii. Prepare 70% ethanol. Add reagents in the order listed and vortex to mix. Maintain at room temperature.

70% Ethanol	10x PN	Stock	Final	1X+10% (μl)	2X+10% (μl)
Nuclease-free Water	-	-	-	330.0	660.0
100% Ethanol	-	100%	70%	770.0	1,540.0
Total	-	-	-	1,100.0	2,200.0

iii. Prepare Autofluorescence Solution using thawed Xenium Autofluorescence Mix prepared according to step 6.0. Add reagents in the order listed and vortex to mix. Maintain at room temperature in the dark until ready to use.

If processing only one slide, remove the Xenium Autofluorescence Mix volume required for 1X+10% (specified in the table below), and refreeze the remaining reagent at -20° C. Once diluted with ethanol, the Xenium Autofluorescence Mix should not be refrozen.

Autofluorescence Solution	10x PN	Stock	Final	1X+10% (μl)	2X+10% (μl)
100% Ethanol	-	100%	-	544.5	1,089.0
Xenium Autofluorescence Mix	2000753	-	-	5.5	11.0
Total	-	-	-	550.0	1,100.0

- **b.** Retrieve the Xenium Cassette from step 5.3 and place on a flat, clean work surface.
- **c.** Using a pipette, remove all PBS-T from the well.
- **d.** Add **500 μl** Diluted Reducing Agent B prepared at step 6.1ai to the well.

- **e.** Apply a new Xenium Cassette Lid on the cassette, and incubate for **10 min** at **room temperature**.
- **f.** Remove the Xenium Cassette Lid. Using a pipette, remove all Diluted Reducing Agent B from the well. **Save the lid** for use in following indicated steps.

Three ethanol washes:

- g. Wash 1: Add 1,000 μ l 70% ethanol. Incubate for 1 min at room temperature. Remove the ethanol.
- h. Wash 2: Add 1,000 μl 100% ethanol. Incubate for 1 min at room temperature. Remove the ethanol.
- i. Wash 3: Add 1,000 μl 100% ethanol. Incubate for 1 min at room temperature. Remove the ethanol.
- j. Pipette mix Autofluorescence Solution prepared at step 6.1aiii thoroughly before dispensing onto sample to prevent settling of reagent. Add **500 μl** Autofluorescence Solution to the well.
- **k.** Reapply Xenium Cassette Lid on the cassette, and incubate for **10 min** at **room temperature in the dark**.
- **1.** Prepare a thermal cycler with the following incubation protocol. Place a Thermocycler Adaptor on the thermal cycler and start the program.

Lid Temperature	Reaction Volume	Run Time
37°C (lid may be set to the lowest temperature if the instrument does not enable 37°C)	100 μΙ	-
Step	Temperature	Time hh:mm:ss
Step Pre-equilibrate	Temperature 37°C	

m. Remove the Xenium Cassette Lid. Using a pipette, remove all the buffer. Discard old Cassette Lids.

Three ethanol washes:



Ethanol washes DO NOT need to be performed in the dark.

- n. Wash 1: Add 1,000 μ l 100% ethanol. Incubate for 2 min at room temperature. Remove the ethanol.
- o. Wash 2: Add 1,000 μl 100% ethanol. Incubate for 2 min at room temperature. Remove the ethanol.

- p. Wash 3: Add 1,000 μ l 100% ethanol. Incubate for 2 min at room temperature. Remove the ethanol.
- **q.** Place Xenium Cassette **without lid** on the Thermocycler Adaptor on the thermal cycler to dry. DO NOT close the thermal cycler lid.
- **r.** Skip pre-equilibrate step to initiate Drying.
- **s. Immediately** remove the cassette from the Thermocycler Adaptor and place on a flat, clean work surface.
- t. Add 1,000 μl 1X PBS prepared at step 1.1 to rehydrate the tissue and incubate for 1 min at room temperature in the dark.
- u. Remove all 1X PBS.
- v. Add 1,000 μl PBS-T and incubate for 2 min at room temperature in the dark.
- O

Optional: photograph the slide against a white background. This image can be used for comparison purposes to identify tissue detachment downstream in the workflow. See Troubleshooting for more details.



w. Store slides **16-24 h** (overnight) at **4°C in the dark** with a Xenium Cassette Lid applied on the Xenium Cassette. If storing slides, DO NOT discard the lid; instead save for Step 6.2 Nuclei Staining. Alternatively, proceed to the next step.

6.2 Nuclei Staining

- a. Retrieve thawed Xenium Nuclei Staining Buffer prepared as outlined in 6.0 Get Started on page 72.
- **b.** Retrieve the Xenium Cassette from the previous step 6.1y and place on a flat, clean work surface.
- c. If stored, remove the Xenium Cassette Lid . Using a pipette, remove all PBS-T from the well. Save the lid for use in following indicated steps.
- d. Add 500 µl Xenium Nuclei Staining Buffer and incubate 1 min at room temperature in the dark.
- e. Remove all Nuclei Staining Buffer.

Three PBS-T washes (PBS-T prepared at step 1.1):

- f. Wash 1: Add 1,000 µl PBS-T. Incubate for 1 min at room temperature in the dark. Remove all PBS-T.
- g. Wash 2: Add 1,000 µl PBS-T. Incubate for 1 min at room temperature in the dark. Remove all PBS-T.
- h. Wash 3: Add 1,000 µl PBS-T. Incubate for 1 min at room temperature in the dark. Remove all PBS-T
- i. Add 1,000 ul PBS-T.



j. Store slides (as specified below) or alternatively, proceed directly to the Xenium Analyzer User Guide (CG000584).



Slides from different Xenium assay workflows cannot be run together on the same instrument run.

Short-term storage for ≤1 week at 4°C in the dark with a Xenium Cassette Lid or slide seal applied on the cassette.



Ensure that the slide is stored in microbe-free and nuclease-free conditions, with a Xenium Cassette Lid or slide seal applied to prevent evaporation. Storing slides for more than recommended time risks decrease in sensitivity. These risks are dependent on many factors, including input tissue quality and how nuclease-free and microbe-free the workflow is.

Long-term storage for (1 week - 2 months) at -20°C. Perform serial ethanol washes to dehydrate, transfer the slide to a mailer, and store in 10 ml cryoprotectant (30% Glycerol prepared in PBS), as described in Xenium Cassette Storage on page 31



Storing slides for more than one month risks ~5% decrease in sensitivity. These risks are dependent on many factors, including input tissue quality and how nuclease-free and microbe-free the workflow is.

The above storage guidelines can also be used for storing Xenium cassettes with slides after Xenium Analyzer run (or in the event of a run failure). Contact support@10xgenomics.com for more information.



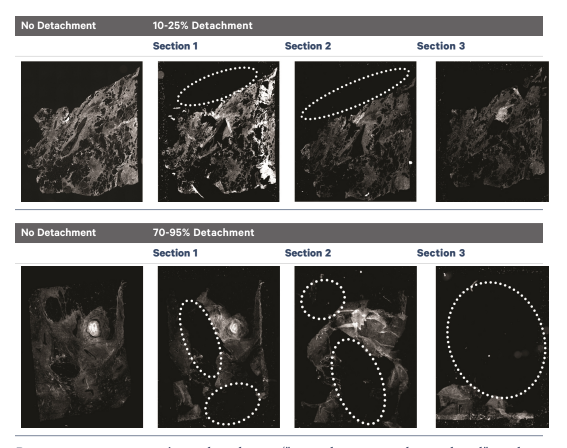
Troubleshooting



Tissue Detachment and Folding

Tissue detachment may result in a lack of decodable data in the region where detachment occurred. If the tissue has folded on itself, this may also cause elevated signal in the overlapping areas. Inspect images carefully to identify these areas. If tissue detachment is observed during this workflow, contact support@10xgenomics.com

Tissue Detachment in Human Breast as viewed on Xenium Analyzer Overview Scan



Percentages represent tissue detachment/"area that cannot be analyzed" at the end of the Xenium Analyzer workflow. White circles indicate areas of tissue detachment.

Tissue Detachment on Xenium Slides

Tissue sections may detach from Xenium slides during on-slide workflows. Tissue adhesion to the slide is impacted both by tissue processing and tissue quality.

Careful tissue preparation is critical for adhesion in on-slide workflows.

Consult Xenium In Situ for Fresh Frozen - Tissue Preparation Guide (Document CG000578) and Xenium In Situ for FFPE - Tissue Preparation Guide (Document CG000579) for tissue QC and sectioning best practices. Below are some additional best practices for minimizing detachment during on-slide workflows:

- Do not pipette directly onto the tissue.
- Gently add and remove reagents from the well. Forceful addition or removal of reagents can agitate tissue and lead to detachment.
- Avoid touching in and around the Sample Area of the Xenium slide.
- Work quickly and carefully during reagent addition and removal.

In addition to following best practices, it is possible to monitor section adhesion on Xenium slides throughout the workflow. Taking a photograph of the slide at the beginning of the on-slide workflow and comparing with post-assay workflow images can help identify whether tissue shape has changed significantly, an indication of detachment. Steps when slide photos can be taken are noted in the protocol. These QC images can be compared with the DAPI overview scan as part of the Web Summary file to see whether tissue morphology has changed in the workflow.

If tissue detachment occurs, send pictures to support@10xgenomics.com for further assistance.

Bubbles during Workflow

Bubbles may occur throughout the workflow, including during PBS-T washes. Bubbles floating on the surface of the slide are unlikely to compromise assay performance. However, bubbles that are in contact with the tissue during a Xenium Analyzer run may result in a lack of decodable data in the tissue area where the bubbles occurred.

Avoid generating bubbles during reagent dispensing by pipetting slowly and avoiding expelling air from the pipette tip. Gently tap or rock the cassette after reagent dispension and inspect the cassette carefully to ensure liquid is fully covering the tissue. DO NOT aspirate or pop the bubbles as this could lead to tissue detachment or scratching of the tissue. Ensure there are no bubbles on the assembled cassette before loading it into the Xenium Analyzer.

Bubbles under Xenium Cassette Insert



Bubbles when dispensing reagents



Bubbles may occur when dispensing Xenium Multi-Tissue Stain Mix under the Xenium Cassette Insert.

If bubbles are observed, DO NOT aspirate or pop the bubbles as this could lead to tissue detachment or scratching of the tissue. Instead, remove the insert and reapply the insert to expel these bubbles.



These bubbles may impact staining quality and can potentially impact assay performance.

- When dispensing a reagent: pipette slowly and avoid expelling any air from the pipette tip.
- After dispensing the reagent: visually inspect the cassette to ensure that the reagent is fully covering the tissue.
- • If bubbles are observed, remove the insert and reapply the insert to expel these bubbles.
 - Gently lift the insert peeling it off from one corner until the bubbles are released.
 - Place the insert back the way it was removed.

Bubbles after overnight incubation

Bubbles can also appear after overnight incubation of the slide (Probe Hybridization or Xenium Multi-Tissue staining) due to bubble nucleation and are unlikely to impact staining or assay performance.

Number of Washes

Post Hybridization and post-Ligation washes are critical for assay performance. Failure to perform the correct number of washes can reduce the fraction of usable decodable data. A similar effect is observed when washing for less than the recommended time, or when reagent is carried over during the washes. Remove all liquid from the well when washing, and refer to User Guide for correct number of washes and incubation times.

Samples Dry Out

Drying of tissue samples may lead to decreased decoding efficiency and unusable data and will impact overall assay performance and sensitivity. Work quickly and ensure reagents are dispensed evenly across tissues during incubation and wash steps throughout the workflow to prevent drying out of tissues. If processing two slides at a time, remove and add reagent from first slide before proceeding to second slide. Ensure tissue sections are always covered with reagent in between removal and addition steps. Note that there are no safe stopping points except for those described in the protocol and outlined specifically in the Protocol Steps & Timing on page 16.

Cassette Assembly Failure

Incorrect assembly of the Xenium cassettes with a Xenium slide can negatively impact assay performance. Always dry the front and the back of the slide completely using a lint-free laboratory wipe while avoiding touching or damaging of the tissue sections. Inspect the slide carefully to ensure it is seated fully within the cassette before assembly.

Example scenarios that may indicate incorrect Xenium Cassette assembly are described below:

- If a gap appears between the two halves of the cassette after assembly.
- If the cassette does not click shut or appears domed after assembly.
- For Xenium Cassette v2, slide is not placed underneath the slide clip.
- If the Xenium Thermocycler Adaptor is wet following removal from the thermal cycler, indicating reagent leakage from the cassette.

If the cassette is incorrectly assembled, disassemble and reassemble the cassette as instructed in the following pages.



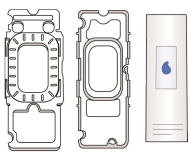
Inspect gasket during cassette assembly. Damaged gaskets can lead to leaks in the cassette.

Xenium Cassette Assembly

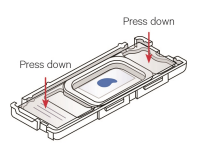


Exercise caution when handling slide edges to prevent injury.

1 Place top and bottom halves of cassette on bench with the top cassette facing down and bottom cassette facing up.



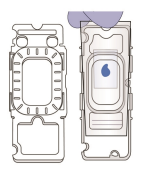
3 Press slide down into grooves of the bottom half of the cassette until it sits firmly in place.



Apply even pressure on top of cassette until all clips click shut. Verify that clips are completely secured over tabs.

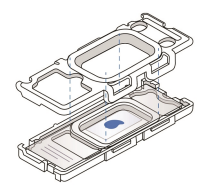


2 Place Xenium slide with tissue side facing upwards into bottom half of cassette; ensure label is toward bottom of cassette.



Slide facing up

4 Secure clips of top half with tabs of bottom half (on both sides).



TIPS

Once cassette is assembled, DO NOT remove slide until after Xenium Analyzer imaging and decoding for optional H&E staining step.

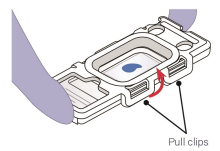
Xenium Cassette Removal

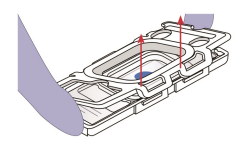


Exercise caution when handling slide edges to prevent injury.

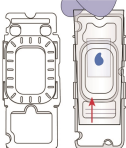
1 Pull inner clips from inner tabs to detach top and bottom halves of cassette.







3 Hold slide by the label and lift slide out from bottom half.



Xenium Cassette, Xenium Cassette Insert, and Lid Cleaning

Xenium Cassettes, Inserts, and Lids are single use items and are to be discarded after use (unless specified in the protocol step that the lid should be saved and reused). Cassettes (prior to cassette assembly) or lids that are accidentally dropped may be reused after thorough cleaning. Note that PBS-T washes DO NOT require sealing of the cassette.

Cleaning Procedure:

- Rinse the lid under running Milli-Q Water
- Spray with 70% isopropanol
- Rinse under running Milli-Q Water
- Spray with 70% isopropanol a second time
- Rinse under running Milli-Q Water
- Air dry

Incorrect Autofluorescence Quenching

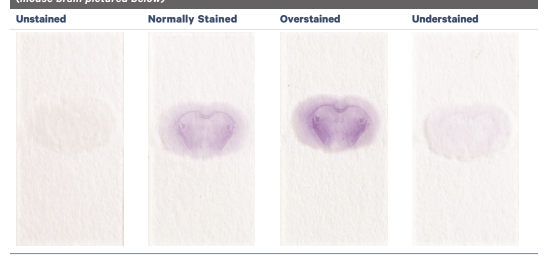
Variation in stain color is normal and tissue-type dependent in tissue sections correctly stained with Autofluorescence Solution. Incorrect staining scenarios are listed below:

- Uneven staining with Autofluorescence Solution may be visible as a non-uniform stain across a tissue section.
- Overquenching can cause tissue to overheat on the Xenium Analyzer, and data generated in the overheated spots may be compromised or missing.

Improper Autofluorescence Quenching risks lower transcript quality scores and reduced median transcripts per cell, but depends on the sample type. Underquenching is lower risk than overquenching.

Ensure Autofluorescence Solution is well mixed and dispensed uniformly across the tissue sections to avoid uneven staining. Autofluorescence Quenching has been optimized for a large number of tissue types and modifying the dilution listed in the User Guide is not recommended. Cassette should be sealed properly and firmly during incubation to prevent reagent evaporation.

Incorrect or insufficient Autofluorescence Quenching may cause variation in staining of tissues (mouse brain pictured below)

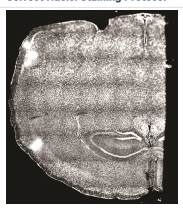


Incorrect Nuclei Staining

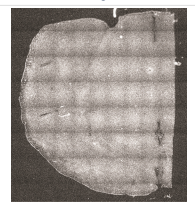
Incorrect staining of nuclei may lead to poor image quality and an inability to easily identify tissue or regions of interest when selecting areas to image during a Xenium Analyzer overview scan. Follow the Nuclei Staining protocol as instructed using the Xenium Nuclei Staining Buffer provided in the Xenium Slides & Sample Prep Reagents Kit - (2 slides, 2 rxns), PN-1000460. Confirm Xenium Nuclei Staining Buffer is well mixed and applied uniformly across tissue sections. All incubations with Xenium Nuclei Staining Buffer should be performed in the dark. If an alternate staining protocol or buffer is used, lower quality images may be obtained.

Incorrect or insufficient nuclei staining may impact image quality and region of interest selection (as viewed on a Xenium Analyzer overview scan)

Correct Nuclei Staining Protocol



No Nuclei Staining Performed



Improper Placement of Xenium Cassette Insert

Improper placement of the Xenium Cassette Insert in the Xenium Cassette can negatively impact assay performance and may cause damage to tissue sections. Inspect the insert carefully to ensure it is seated fully within the cassette. Practice placement of insert with a blank slide if necessary to avoid damage to experimental samples. Refer to Xenium Cassette Insert Practice on page 40.

Place the assembled cassette on a white surface to check insert placement. If the insert is incorrectly placed (see image below), remove and reapply the insert into the cassette. Gently lift the cassette insert by the handle using forceps. DO NOT lift the insert straight up as this may cause tissue detachment. Instead, gently peel back one corner and lift the insert. Finally, place insert back the way it was removed.





Appendix

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Sample Shipping	92

Probe Panel Selection

Ensure that a compatible gene panel has been selected prior to executing the Xenium In Situ Gene Expression workflow. 10x Genomics provides the option of using pre-designed gene panels, pre-designed panels that are customized by adding genes of interest, and standalone custom gene panels.

Pre-designed Gene Panels

Refer to the 10x Genomics Support Website for the most current list of available panels and part numbers.

Custom Gene Panels

Contact your 10x Genomics Sales Executive for information about designing a standalone custom gene panel or add-on custom gene panels that are compatible with pre-designed panels. If you do not know your Sales Executive, please contact customerservice@10xgenomics.com.

If utilizing a custom panel, the Design ID on the label of the tube containing the custom panel should match with the first portion of the custom gene panel electronic file name.

Sample Shipping

Processed Xenium slides may be shipped within 1 week of completing the Xenium In Situ assay workflow. After Nuclei Staining, remove all PBS-T, disassemble the Xenium Cassette, and place no more than two slides in a mailer. Ship using one of the two options:

- Fill the mailer (containing the slides) to capacity with PBS-T, place the mailer in a container with ice packs, and ship (overnight to 2 days).
- Dehydrate the slides by serial ethanol washes as described on Xenium Cassette Storage on page 31. Fill the mailer (containing the slides) to capacity with a cryoprotectant (30% Glycerol), place the mailer in a container with dry ice, and ship (overnight to 2 days).

Note that assay performance may be compromised post-shipping and handling.

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