

User Guide | CG000685 | Rev D

Visium HD Spatial Gene Expression Reagent Kits

For use with:

Visium HD Reagent Kit, Small, PN-1000668

Visium Human Transcriptome Probe Kit v2 - Small, PN-1000466

Visium Mouse Transcriptome Probe Kit v2 - Small, PN-1000667

Visium HD Cassettes, 6.5 mm, 4 rxns, PN-1000669

Visium HD Slide, 6.5 mm, 2 rxns, PN-1000670

Visium CytAssist Reagent Accessory Kit, PN-1000499

Visium CytAssist Alignment Aid Kit, 6.5 mm, PN-1000886

Dual Index Kit TS Set A, 96 rxns PN-1000251



View videos for key techniques on slide preparation and use, slide drying, and more. To view, click here.



Take 1 minute to evaluate this protocol. Scan this code or click here.



Notices

Document Number

CG000685 | Rev D

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Document Revision Summary

Document Number

CG000685

Title

Visium HD Spatial Gene Expression Reagent Kits User Guide

Revision

Rev D

Revision Date

August 25, 2025

Description of Changes

- Updated Visium S3 Cassette part numbers in Reagent Kits on page 9.
- · Added information regarding Visium HD best practice videos to cover page and Videos on page 23.
- Added information on varying Visium HD Slide appearance to Visium HD Slides on page 16.
- Added information on the Visium CytAssist Alignment Aid to Visium CytAssist Alignment Aid on page 33.
- Added information on lint-free laboratory wipes to Lint-free Laboratory Wipes on page 24.
- Added information on Visium S3 Cassette appearance to Reagent Addition to & Removal from Wells on page 26.
- Adjusted Perm Enzyme B preparation instructions in 4.0 Get Started on page 67.
- Updated Visium HD Slide drying guidance in 3.1 Visium HD Slide Wash on page 64.
- Added information on using the Visium CytAssist Alignment Aid to 4.1 CytAssist-Enabled Probe Release & Capture on page 75.
- Updated tissue slide drying and Visium HD Slide drying language in 4.1 CytAssist-Enabled Probe Release & Capture on page 75.
- Updated Sequencing Depth example in Sequencing Depth on page 97.
- Added information on Space Ranger segmentation pipeline to 3. Tissue Segmentation Failure due to High Connective Tissue on page 116.

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Reagent Kits

Reagent Kits	Part Number	Components	Component Part Number
Visium HD, Human Transcriptome, 6.5 mm*	1000675	Visium HD Slide, 6.5 mm, 2 rxns	1000670
		Visium Human Transcriptome Probes v2 - small	1000466
		Visium HD Reagents, small	1000668
		Visium HD Cassettes, 6.5 mm, 4 rxns	1000669
Visium HD Mouse, Transcriptome, 6.5 mm*	1000676	Visium HD Slide, 6.5 mm, 2 rxns	1000670
		Visium Mouse Transcriptome Probes v2 - small	1000667
		Visium HD Reagents, small	1000668
		Visium HD Cassettes, 6.5 mm, 4 rxns	1000669

^{*}Also available in a pack of 4 as a 16 rxn kit.

Visium HD Slide, 6.5 mm, 2 rxns PN-1000670



Visium HD Cassettes*, 6.5 mm, 4 rxns PN-1000669

Visium HD Cassettes, 6.5 mm 4 rxns PN-1000669 (store at ambient temperature)		
	#	PN
Visium 2-port Cassette S3, 6.5 mm		
Visium Cassette 2-port gasket, 6.5 mm	2	3001831 or 3002329
Visium Cassette Bottom	2	3001830 or 3002328
Visium Tissue Slide Cassette S3, 6.5 mm		
Tissue Slide Cassette Top	4	3001826 or 3002327
Movable Tissue Gasket 6.5 mm (preassembled with translator)	4	3001828
Movable Tissue Gasket Translator (preassembled with gasket)	4	3001927 or 2002330
Tissue Slide Cassette Bottom	4	3001825 or 3002326
Visium Slide Seals, 12 pack	2	2000283
		10x GENOMICS

*The Visium HD Workflow is run with Visium Tissue Slide Cassettes S3 and Visium Cassettes S3. These are referred to as Tissue Slide Cassettes and Visium Cassettes in this document, respectively. Consult the Visium Cassette S3 Quick Reference Card (CG000730) for assembly and disassembly information.

Visium HD Reagent Kit - Small, PN-1000668



Visium Human Transcriptome Probe Kit v2 - Small, PN-1000466

Visium Human Transcriptome Probe Kit v2 - Small PN-1000466 (store at -20°C)			
		#	PN
0	FFPE Hyb Buffer	1	2000423
0	FFPE Post-Hyb Wash Buffer	1	2000424
	Human WT Probes v2 – RHS	1	2000657
	Human WT Probes v2 – LHS	1	2000658
0	Probe Ligation Enzyme	1	2000425
0	2X Probe Ligation Buffer	1	2000445
0	Post Ligation Wash Buffer	1	2000419
			10x genomics

Visium Mouse Transcriptome Probe Kit v2- Small, PN-1000667



Dual Index Kit TS Set A, 96 rxns PN-1000251

Dual Index Kit TS Set A 96 rxns PN-1000251 (store at -20°C)		
	#	PN
Dual Index Plate TS Set A	1	3000511

10x Genomics Accessories

Visium CytAssist Alignment Aid Kit, 6.5 mm PN-1000886



Visium CytAssist Reagent Accessory Kit PN-1000499



Third-Party Items

Successful execution of the Visium HD workflow requires third-party reagents, kits, and equipment in addition to those provided by 10x Genomics. All third-party reagents and consumables should be obtained prior to starting this workflow.

Consult the Visium HD Spatial Gene Expression Protocol Planner (CG000698) for a list of the following third-party items:

- · Additional reagents, kits, and equipment
- Tested pipette tips
- Tested thermal cyclers
- · Tested blank slides



10x Genomics has tested all items listed in the Protocol Planner. Unless otherwise noted, 10x Genomics recommends using these items to ensure optimal assay performance. Substituting materials may adversely affect assay performance.

Workflow Overview

Sample Preparation

Before starting this User Guide, consult these documents to prepare samples. Choose **one** handbook for guidance on tissue preparation and staining.

Visium HD FFPE Tissue Preparation Handbook

Prepare formalin fixed & paraffin embedded tissue blocks, section tissue onto slides, stain, and image.

Demonstrated Protocol CG000684

Visium HD FxF Tissue Preparation Handbook

Prepare fixed frozen tissue blocks, section tissue onto slides, stain, and image.

Demonstrated Protocol CG000764

Visium HD Spatial Gene Visium HD Spatial Expression Protocol Planner

Information on third-party items.

Planner CG000698

Applications Imaging Guidelines

Visium HD FF Tissue

Preparation Handbook

slides, stain, and image.

Demonstrated Protocol

CG000763

Prepare fresh frozen tissue

blocks, section tissue onto

Optimize imaging settings.

Technical Note CG000688

Visium Cassette S3 Quick Reference Card

Practice cassette assembly and disassembly.

Quick Reference Card CG000730

Visium CytAssist **Accessory Kit Quick Reference Card**

Determine slide allowable

Quick Reference Card CG000548

Library Construction

Visium HD Spatial Gene **Expression Reagent Kits User Guide**

Construct Visium HD Spatial Gene Expression - Probebased Libraries.

User Guide CG000685

Consult the 10x Genomics support website for additional documents

Protocol Steps & Timing

Steps	Timing	Stop & Store
Day 1		
Step 1: Probe Hybridization (page 48)		
1.1 Probe Hybridization (page 50)	Overnight	
Day 2		
Step 2: Probe Ligation (page 53)		
2.1 Post-Hybridization Wash (page 55)	18 min	
2.2 Probe Ligation (page 57)	60 min	
2.3 Post-Ligation Wash (page 58)	12 min	stop 4°C ≤24 h
Step 3: Visium HD Slide Preparation (page 60)		
3.1 Visium HD Slide Wash (page 62)	20 min	
4.1 CytAssist-Enabled Probe Release & Capture (page 69)		
4.1 CytAssist-Enabled Probe Release & Capture (page 69)	60 min	
4.2 Probe Extension (page 81)	60 min	STOP 4°C ≤24 h
4.3 Probe Elution (page 82)	15 min	
Step 5: Pre-Amplification and SPRIselect (page 83)		
5.1 Pre-Amplification (page 85)	40 min	
5.2 Pre-Amplification Cleanup - SPRIselect (page 86)	30 min	4°C ≤72 h or -20°C ≤4 weeks
Day 3		
Step 6: Visium HD Spatial Gene Expression – Probe-based Library Construction (p	page 87)	
6.1 Cycle Number Determination – qPCR (page 89)	45 min	
6.2 Sample Index PCR (page 91)	40 min	
6.3 Post-Sample Index PCR Cleanup – SPRIselect (page 93)	30 min	-20°C long-term
6.4 Post-Library Construction QC (page 94)	50 min	

Stepwise Objectives

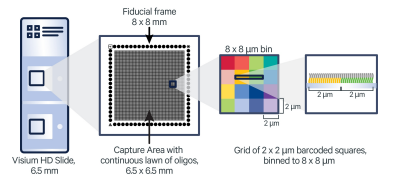
The Visium HD Spatial Gene Expression assay is designed to analyze mRNA in tissue sections derived from formalin fixed & paraffin embedded (FFPE), fresh frozen (FF), or fixed frozen (FxF) tissue samples. It uses probes targeting the whole transcriptome. Each Visium HD Slide contains Capture Areas with barcoded squares that include oligonucleotides required to capture gene expression probes. Before the assay, tissue sections are processed as described in their relevant Demonstrated Protocol. See Workflow Overview on page 13 for documentation references.

Human or mouse whole transcriptome probe panels, consisting of ~3 pairs of specific probes for each targeted gene are added to the tissue, enabling hybridization and ligation of each probe pair. Tissue slides and Visium HD Slides are loaded into the Visium CytAssist instrument, where they are brought into proximity with one another. Gene expression probes are released from the tissue upon CytAssist-Enabled Probe Release & Capture, enabling capture by the spatially-barcoded oligonucleotides present in a hydrogel on the Visium slide surface. The Visium HD Slide is removed from the Visium CytAssist for downstream library preparation. Gene expression libraries are generated from each tissue section and sequenced. Spatial Barcodes are used to associate the reads back to the tissue section images for spatial mapping of gene expression.

Visium HD Slides

The Visium HD Slide, 6.5 mm has 2 Capture Areas. Each Capture Area is 6.5 x 6.5 mm and is surrounded by a fiducial frame that is used for image alignment. The fiducial frame + Capture Area is 8 x 8 mm. The Capture Area is a continuous lawn of oligos comprised of 2 μ m barcoded squares. Each barcoded square has oligos with an Illumina TruSeq partial read 1 sequencing primer, unique molecular identifier (UMI), Spatial Barcode, and 30 nt poly(dT) sequence (captures ligation product). Each Capture Area is surrounded by a spacer. This spacer creates a reaction chamber that facilitates proper reagent addition.

The active surface of the slide has an etched label that includes the serial number. The label should be legible. If the label appears reversed, the active surface is facing down. The slide should be loaded onto the instrument with the active surface facing up.

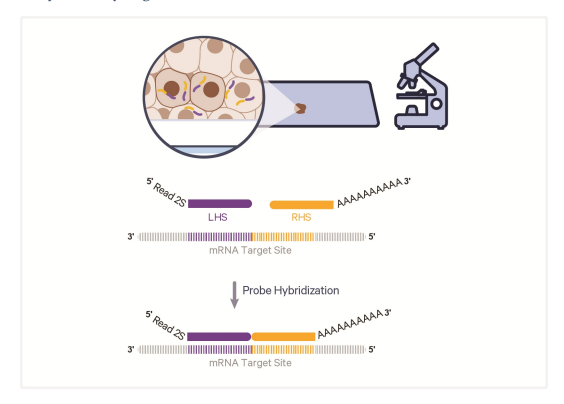


Visium HD Slide appearance may vary, but all Visium HD Slides have the same functionality and performance.



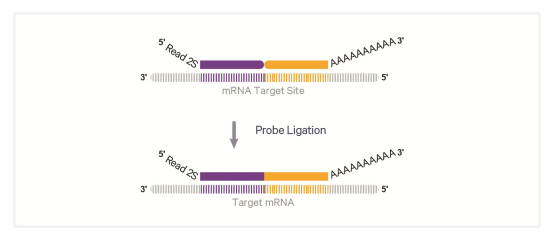
Step 1: Probe Hybridization

The human or mouse whole transcriptome probe panel, consisting of ~3 specific probes for each targeted gene, is added to the deparaffinized, destained, and decrosslinked tissues. Together, probe pairs hybridize to their complimentary target RNA.



Step 2: Probe Ligation

After hybridization, a ligase is added to bridge the junction between the probe pairs that have hybridized to RNA, forming a ligation product.



Step 3: Visium HD Slide Preparation

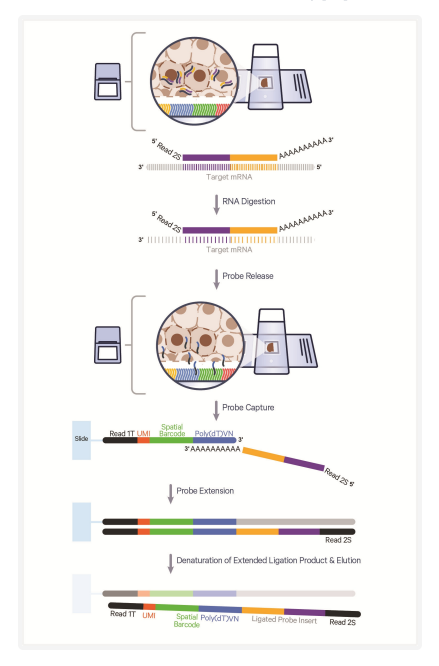
Visium HD Slides are thawed, washed, and equilibrated prior to placement on the Visium CytAssist instrument.



Step 4: Probe Release & Extension

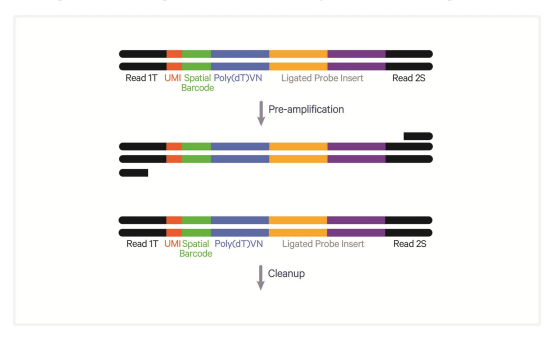
Probe release and capture occurs in the Visium CytAssist instrument. The single stranded ligation products are released from the tissue upon RNase treatment and captured on the Visium slide. Once ligation products are captured, the slides can be removed from the instrument.

Ligation products are extended by the addition of the Spatial Barcode, UMI, and partial Read 1 primer. This generates spatially-barcoded ligation products, which can then be carried forward for library preparation.



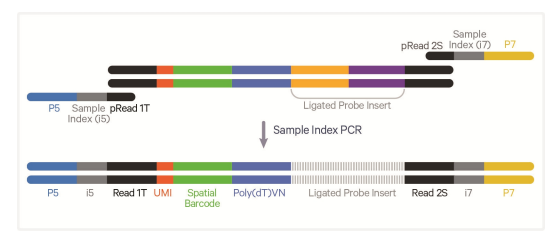
Step 5: Pre-amplification and SPRIselect

To generate ample material for library construction, barcoded ligation products are amplified. This amplification is followed by SPRIselect cleanup.



Step 6: Visium HD Spatial Gene Expression - Probe-based Library Construction

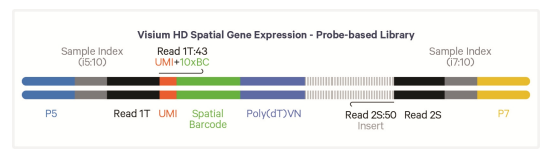
Pre-amplification material is collected for qPCR to determine sample index PCR cycle number for gene expression libraries. The amplified material then undergoes indexing via sample index PCR to generate final library molecules. The final libraries are cleaned up by SPRIselect, assessed on a Bioanalyzer or a similar instrument, quantified by qPCR, and then sequenced.



Sequencing

A Visium HD Spatial Gene Expression - Probe-based library comprises standard Illumina paired-end constructs, which begin and end with P5 and P7 adaptors. The 43 bp UMI and Spatial Barcode are encoded in TruSeq Read 1, while Small RNA Read 2 (Read 2S) is used to sequence the ligated probe product.

Illumina sequencer compatibility, sample indices, and library loading information are summarized in the Sequencing section.



See Oligonucleotide Sequences on page 129



Tips & Best Practices

Icons



Tips & Best Practices section includes additional guidance



Signifies critical step requiring accurate execution



Troubleshooting section includes additional guidance



Indicates a video is available for the specified technique

Videos

- Videos describing key steps in the protocol can be found here.
- When a relevant video is available for a protocol step, it will be marked with a video icon.

General Reagent Handling

- Fully thaw and thoroughly mix reagents before use.
- When pipette mixing, unless otherwise specified, set pipette to 75% of total volume.

Visium HD Slide Storage

- Keep Visium HD Slide at -80°C until ready to use.
- Do not open the mylar bag containing the Visium HD slides until slides are ready to be used.

Pipette Calibration

- Follow manufacturer's calibration and maintenance schedules.
- Pipette accuracy is particularly important when using SPRIselect reagents.

Lint-free Laboratory Wipes

- All laboratory wipes used in the protocol must be 100% polyester and lintfree.
- Lens paper or non-lint free laboratory wipes are not suitable alternatives. See Visium HD Spatial Gene Expression Protocol Planner (CG000698) for

tested part numbers.



Visium HD Slide Handling

- Wipe Visium HD Slide Mailer with a lint-free laboratory wipe to ease in handling.
- Prior to beginning an instrument run, Visium HD Slides are thawed, washed, equilibrated, and dried. Using an external light source (like a flashlight) can help confirm that the Visium HD Slide is fully dried.
- Occasionally, small chips can arise on the edge of the Visium HD Slide.
 These are mainly cosmetic in nature and do not typically impact assay performance.



Sharp edges on the slide's corners may cause injury or damage if mishandled. Handle with care. Wear proper PPE when handling and avoid contact with the chipped corner to prevent cuts or scratches.



• Visium HD Slide preparation steps should be done gently to avoid damaging the slide.

- Visium HD Slides should only be washed if proceeding with a CytAssist instrument run.
- · Always wear gloves when handling slides.
- Ensure that the active surface of a slide faces up and is never touched. The active surface is defined by a readable label. The image of the chipped Visium HD slide demonstrates the active surface facing up.
- Minimize exposure of the slides to sources of particles and fibers.
- When pipetting reagent onto a slide, avoid generating bubbles.
- If debris appears on the Visium HD Slide after performing slide washes, slide may be re-immersed in 0.1X SSC in the slide mailer to remove debris before proceeding. If performing an additional wash, ensure the slide is dried as described in the protocol.
- If necessary, unwashed, room temperature Visium HD Slides may be returned to -80°C. Protect from light. DO NOT exceed 3 h at room temperature. DO NOT exceed three freeze/thaw cycles.
- DO NOT allow the Visium HD Slide to dry, except where indicated in the protocol steps.
- If using a barcode scanner to read the serial number, ensure that the label is not wet.
- When disassembling the Visium HD Slide from the Visium Cassette, ensure that the cassette gaskets DO NOT touch any surfaces to avoid debris accumulation.

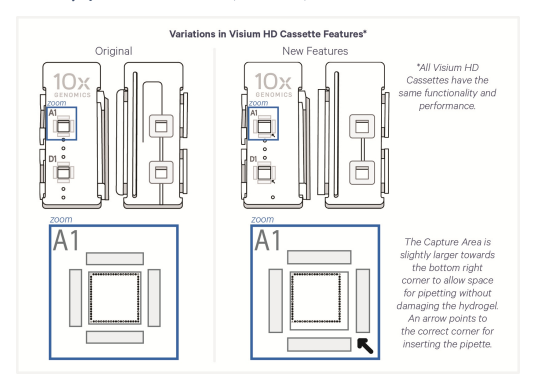
Tissue Slide Handling

- To ensure compatibility with the Visium CytAssist, tissue sections must be placed in specific areas on a blank slide. Tested slides, as well as appropriate tissue placement areas, are listed in the Visium HD Spatial Gene Expression Protocol Planner (CG000698).
- Always wear gloves when handling slides.

Reagent Addition to & Removal from Wells

Reagent Addition

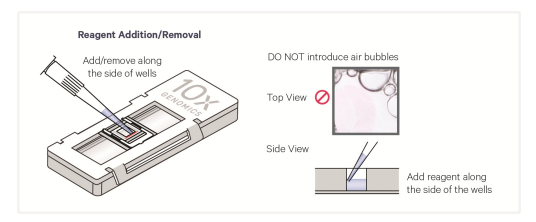
• Assemble slide into the cassette flat on a clean work surface. See Cassette Assembly Quick Reference Card (CG000730).



- Dispense and remove reagents along the side of the wells without touching the slide surface, tissue sections (when applicable), and without introducing bubbles.
- DO NOT forcefully insert pipette tip into well corners to avoid damaging the Visium HD Slide.
- If applicable, perform washes next to the thermal cycler to avoid significant changes in temperature during washes.
- When processing two or more Visium Tissue Slide Cassettes, remove and add reagent from the first cassette before proceeding to the next cassette. Ensure tissue sections are always covered with reagent in between removal and addition steps to avoid drying out of tissue samples.
- Unless indicated otherwise, when adding and removing reagent from a cassette, do so one well at a time. For example, remove reagent from A1 and add the next reagent to A1, before moving into onto D1, to avoid drying the well.

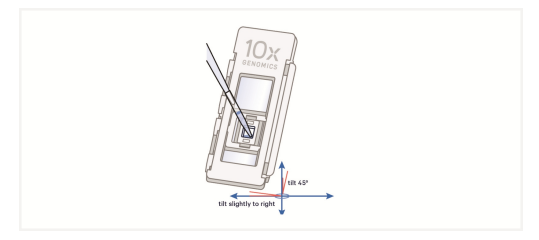


• Always cover the Capture Area or tissue completely when adding reagents to the well. A gentle tap may help spread the reagent more evenly.



Reagent Removal from Wells

- Slightly tilt the cassette while removing the reagent. DO NOT completely tilt the cassette at a 90 degree angle to avoid liquid from entering other grooves within the gasket. See image below for proper technique.
- Place the pipette tip on the bottom edge of the wells.
- Remove reagents along the side of the wells without touching the tissue sections (when applicable) 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.



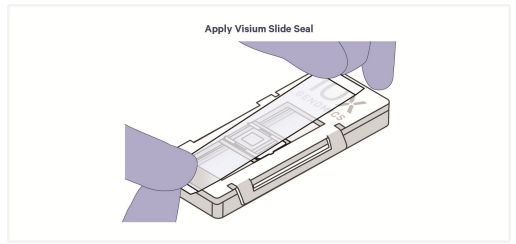
Visium Slide Seal Application & Removal

Application

• If applying a Visium Slide Seal to a Tissue Slide Cassette, the seal must be cut in half lengthwise. Cut the seal as shown in the image below. Six precut seals per tissue slide are necessary for this assay.

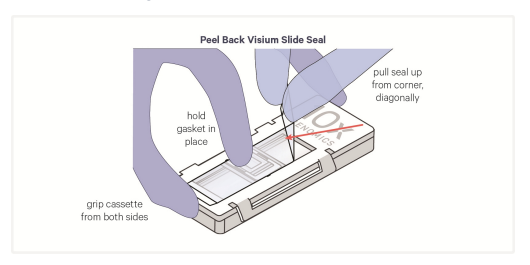


- Place the cassette flat on a clean work surface. Ensure surface of cassette is dry.
- Remove the back of the adhesive Visium Slide Seal.
- Align the Visium Slide Seal with the surface of the cassette and apply while firmly holding the cassette with one hand.
- Press on the Visium Slide Seal to ensure uniform adhesion.
- Use a fresh Visium Slide Seal when prompted during the protocol. Steps that do not require a new slide seal will specify that the slide seal should be pulled back and reapplied instead.



Removal

- Place the cassette flat on a clean work surface.
- Grasp a Visium Slide Seal corner. Carefully pull on the Visium Slide Seal up and across the cassette while firmly holding the cassette. Ensure that no liquid splashes out of the wells. Movement of the gasket during slide seal removal could damage the tissue section.



Slide Incubation Guidance

Incubation at a Specified Temperature

- Position a Low Profile Thermocycler Adapter on a thermal cycler that is set at the incubation temperature.
- Ensure that the Low Profile Thermocycler Adapter is in contact with the thermal cycler block surface uniformly. Pre-equilibrate adapters at the HOLD temperature described in the protocol steps for at least five minutes.
- Open and close the thermal cycler lid gently. If the lid is adjustable, tighten lid only as much as necessary. Avoid overtightening.

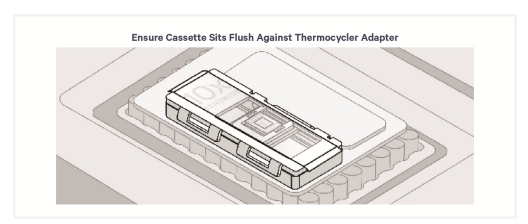
Incubation at Room Temperature

• Place the slide/cassette on a flat, clean, nonabsorbent work surface.

Cassette Incubation

Incubation using a Thermal Cycler:

- Position a Low Profile Thermocycler Adapter on a thermal cycler that is set at the incubation temperature. Low Profile Thermocycler Adapter must come to temperature before cassette placement.
- Ensure that the Low Profile Thermocycler Adapter is in contact with the thermal cycler surface uniformly. Consult the Visium HD Spatial Gene Expression Protocol Planner (CG000698) for information on thermal cycler compatibility.
- Ensure that the cassette sits flush with the raised portion of the Low Profile Thermocycler Adapter.
- Allow Low Profile Thermocycler Adapter to cool before removing from thermal cycler.



Tissue Detachment on Tissue Slides

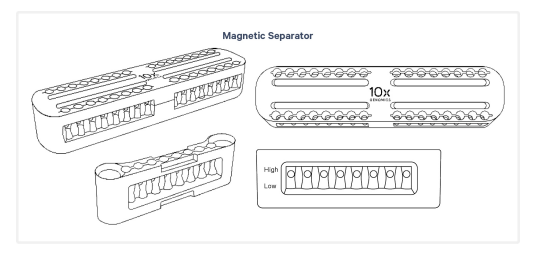


- Monitor section adhesion on tissue slides throughout the workflow.
- Ensure that tissue sections are on Superfrost Plus or positively charged slides. A list of tested slides can be found in the Visium HD Protocol Planner (CG000698).
- 10x Genomics recommends placing tissues with large amounts of connective tissue (like breast or colon) on Schott Nexterion Slide H - 3D Hydrogel Coated Slides (Schott North America, PN-1800434) to minimize risk of tissue detachment. Schott H slides may also be used if prior detachment has been observed with tissues of interest.

- Follow manufacturer instructions for Schott Nexterion Slide H slides, which have specific requirements for storage and equilibration.
- Tissue detachment before the completion of Probe Release during the workflow can negatively impact performance. Ensure that all guidance in tissue preparation handbook is followed.
- For more information, see Troubleshooting on page 107.

10x Magnetic Separator

• Images below are illustrative - actual appearance of magnetic separator may vary. Guidance applies to all 10x Magnetic Separators.



- Offers two positions of the magnets (high and low) relative to a tube, depending on its orientation.
- Flip the magnetic separator over to switch between high (magnet-High) or low (magnet-Low) positions.



• If using MicroAmp 8-Tube Strips, use the high position (magnet•**High**) only throughout the protocol.

Magnetic Bead Cleanup Steps

- During magnetic bead-based cleanup steps that specify waiting "until the solution clears", visually confirm clearing of solution before proceeding to the next step. See panel below for an example. If solution is not clear, leave on magnet until separation is complete.
- The time needed for the solution to clear may vary based on specific step, reagents, volume of reagents, etc.
- Images below are illustrative actual appearance of magnetic separator may

1. Magnetic beads mixed with reagent

2. Separation in progress

3. Separation continuing

4. Separation complete; solution clear

vary. Guidance applies to all 10x Magnetic Separators.

SPRIselect Cleanup & Size Selection

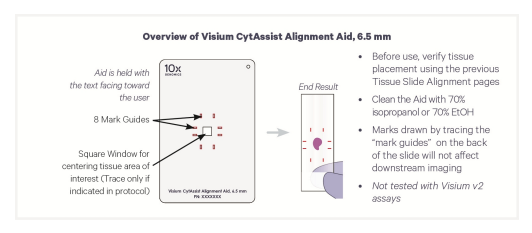
- Use fresh preparations of 80% Ethanol.
- Ensure SPRI beads are not expired and are thoroughly mixed before use.
- After aspirating the desired volume of SPRIselect reagent, examine the pipette tips before dispensing to ensure the correct volume is transferred.
- Pipette mix thoroughly as insufficient mixing of sample and SPRIselect reagent will lead to inconsistent results.
- Ensure solution is completely clear before removing supernatant.

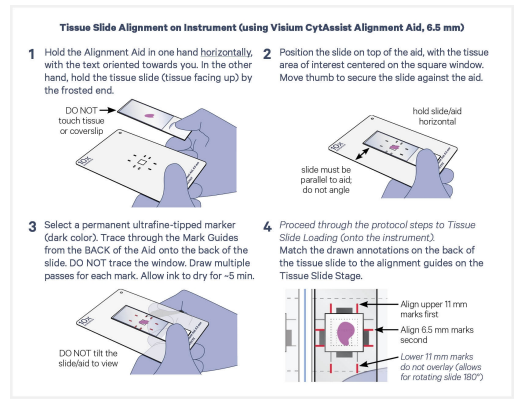
Sample Indices (i5/i7) in Sample Index PCR

- Choose the appropriate sample index sets to ensure that no sample indices overlap in a multiplexed sequencing run.
- Each well in the Dual Index Plate TS Set A contains a unique i7 and a unique i5 oligonucleotide.
- To avoid the risk of cross-contamination, use each plate well only once.
- Record the index plate well location for each sample.

Visium CytAssist Alignment Aid

 The Visium CytAssist Alignment Aid (6.5 mm, PN-1000886; available for purchase separately) is an optional accessory used to draw annotations onto the back of the tissue slide to assist with alignment onto the Visium CytAssist instrument.





- Use aid when indicated in the protocol.
- Marks drawn by tracing the "mark guides" shown in the image above on the back of the slide using the aid will not affect downstream CytAssist

imaging.

- Before use:
 - Ensure back of tissue slide is dry.
 - If a coverslip is present, remove excess mounting medium by gently touching the slide to a lint-free laboratory wipe.
 - ° Clean aid with 70% isopropanol or 70% ethanol.
 - Obtain a permanent, ultrafine-tipped, dark-colored marker.
- During use:
 - If a coverslip is present, do not move coverslip.
 - o Draw multiple passes for each mark.
 - Allow ink to dry for ~5 min.
- Once marks are drawn, avoid wiping the back of the slide vigorously to prevent mark removal. Instead, tap the back of the slide onto a lint-free laboratory wipe to remove excess moisture.



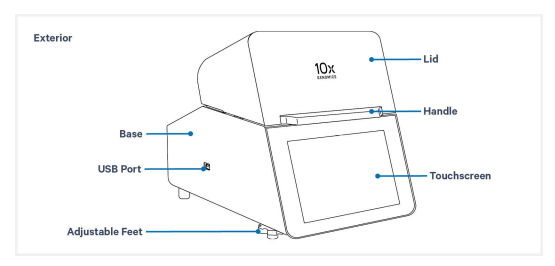
Visium CytAssist

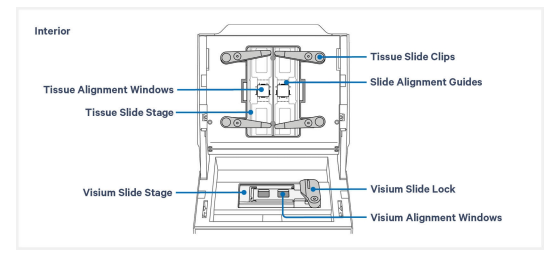
Instrument Orientation	36
Instrument Loading Guidelines	37

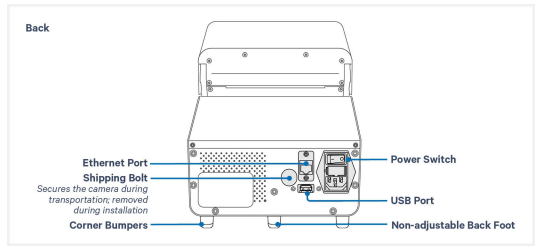
Instrument Orientation



Firmware version 2.0.0 or higher is required in the Visium CytAssist used for this protocol.







Visium CytAssist 10xgenomics.com 36

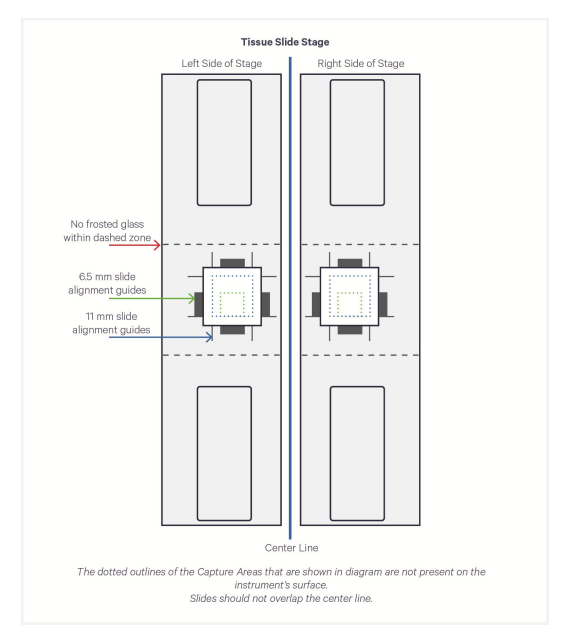
Instrument Loading Guidelines

- Ensure tissue section placement is compatible with Visium CytAssist by using the Visium CytAssist Accessory Kit Quick Reference Card (CG000548). A list of tested slide types can be found in the Visium HD Spatial Gene Expression Protocol Planner (CG000698).
- Consult the Visium CytAssist Training Kit UG (CG000549) for information on practicing tissue slide loading.
- Each tissue slide may be used for one Capture Area on a Visium HD Slide.

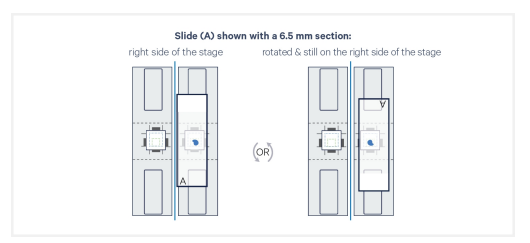
Determine Slide Placement

a. On the diagram below, align tissue to the center of the alignment guides for 6.5 mm (rectangles) or 11 mm (lines) Capture Areas on either the left or right side of the stage. Align the center of the tissue region of interest to the center of the Capture Area, instead of aligning the edge of the region to the edge of the Capture Area. If the Alignment Aid was used, use the annotations to assist with aligning. See Visium CytAssist Alignment Aid on page 33 for instructions.

If printing the image, select "actual size" or "100%" to print to scale. The blue Capture Area should measure 11 mm and the green capture area should measure 6.5 mm.



b. If necessary, rotate the slide 180° as shown. Place off-center tissues closer to the center line (shown as a blue line in the image below). Slides should not overlap the center line. Frosted glass should not be within the dashed zone of the tissue slide stage.

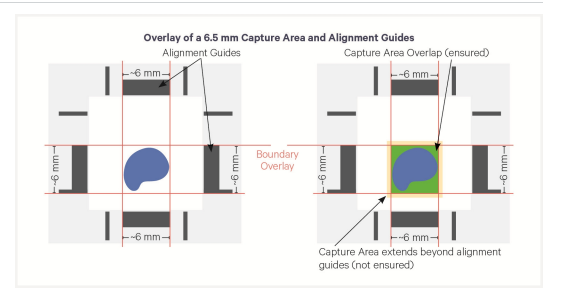


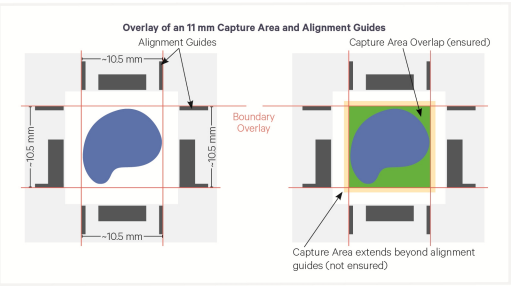
Fine Adjustments

The **outside** of the alignment guides, shown in red, mark the inner boundaries of the fiducial frame on a Capture Area. They mark a distance **approximately** 6 mm (6.5 mm slides) or 10.5 mm (11 mm slides) wide on each side and surround the target region ensured to align within the fiducial frame. Though the Visium CytAssist instrument displays guides for 11 mm Capture Areas, the Visium HD Spatial Gene Expression Assay is only compatible with 6.5 mm Capture Areas.

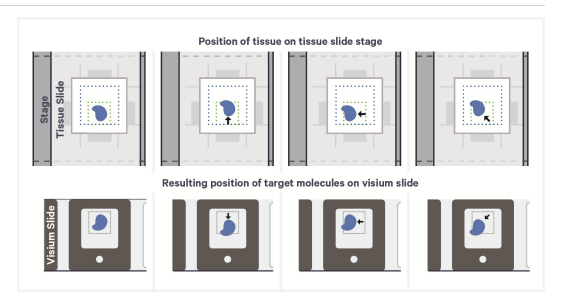
When placing a tissue slide on the Tissue Slide Stage, center the tissue or area of interest within the area inside the outer edges of the alignment guides (area in green in the images below).

Alignment guide dimensions and locations are designed with a high tolerance to minimize data loss. Tissue within the alignment guides is ensured to overlap with the Capture Area during a successful run. Tissue outside of these guides may be captured (area in yellow in the images below), but it is not ensured. 10x Genomics recommends placing high-priority tissue areas within the alignment guides.





The image below demonstrates how movement of the Tissue Slides affects where target molecules will end up on the Visium HD Slide.



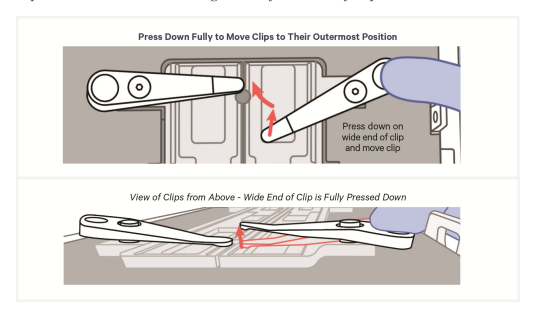
Tissue Slide Loading



Tissue slide loading instructions are updated in this document revision (Rev C and later).

a. Press down fully on the wide end of each clip to lift and rotate the narrow end to its outermost position.

Cradle the instrument lid for increased stability and fully press down on the clips with the thumb. See image below for extent of clip rotation.



b. Lay the tissue slide flat on the stage surface and against the center line. If necessary, rotate the slide 180° to better place off-center tissues closer to the center line. Slides should not cross/overlap the center line.



DO NOT touch the tissue section.

- **c.** Use one hand to hold slide in place with two fingers, one on either side of the tissue. Use the other to both cradle the instrument lid and fully press down on the first clip. Rotate the clip over the slide.
- **d.** While clip is hovering over the slide, make fine adjustments to align the tissue or area of interest to the center of the alignment guides. Gently release when adjustments are complete.

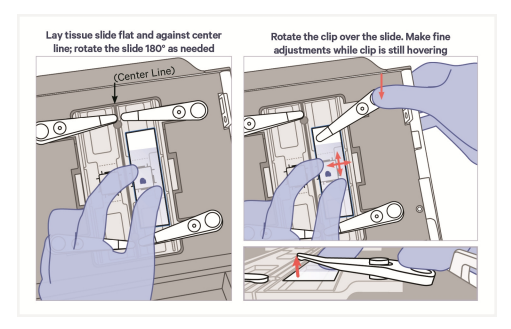


DO NOT adjust slide while tissue clip is touching the slide, which may result in slide damage. DO NOT touch the tissue section.



DO NOT place clips on the edge of the slide, which may result in slide damage.

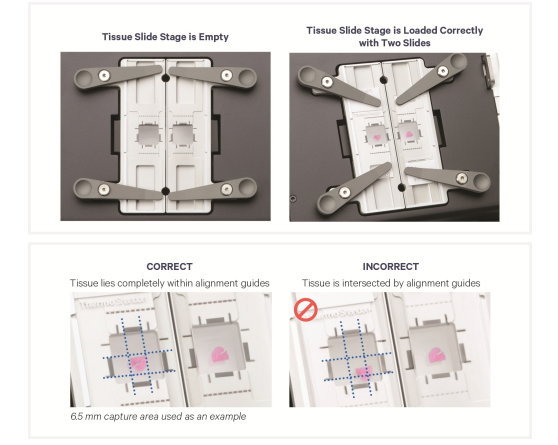
Reference images below for steps b-d.



e. Continue to hold the slide in place while pressing down fully on the wide end of the second clip. Rotate the second clip onto the slide and gently release.

Use both clips whenever possible. The second clip may be omitted if the tissue slide is beyond its reach.

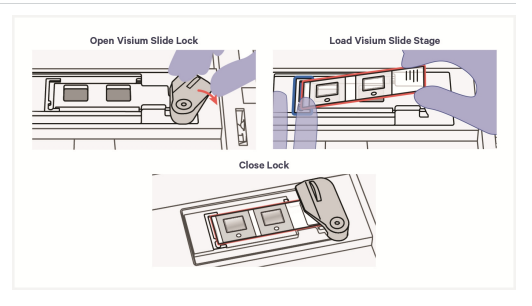
f. If only one tissue on one slide will be analyzed, use a blank slide for the second position on the Tissue Slide Stage. If using only one tissue slide, the unused Capture Area on the Visium HD Slide cannot be used in another instrument run.



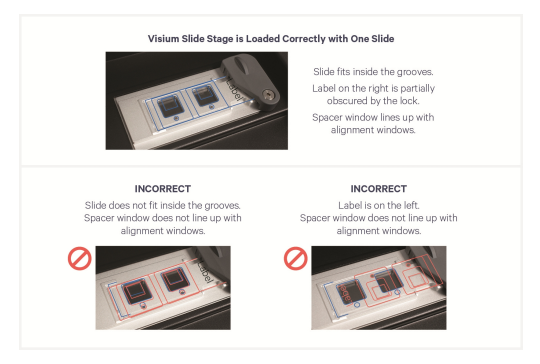
Visium HD Slide Loading

Before loading, record the slide serial number and note which tissue is placed in which Capture Area. Note which Capture Area will be placed on which side of the Visium Slide Stage (Left = D1, Right = A1).

- a. Open the Visium Slide Lock by using one finger to pivot.
- **b.** Line up the slide with the label face up and oriented to the right.
- **c.** Fit the slide within the raised grooves on the left, top, and bottom.



d. Hold slide in place with one hand while slowly closing the Visium Slide Lock. Lock will partially obscure the slide label when correctly secured. Close lock gently to avoid slide damage. In the images below, the slide (red outline) does not line up with the stage (blue outline).





Sample Preparation & Staining Guidelines

Sample Preparation

Proper tissue handling and preparation techniques preserve the morphological quality of the tissue sections and the integrity of RNA transcripts. Maintaining high-quality RNA is critical to assay performance.



Before performing the protocol steps in this User Guide, ensure that all samples have followed the appropriate documentation as described in the Workflow Overview on page 13. Listed below are key considerations described in the tissue preparation documentation.

Key Con	siderations
FFPE Ti	ssue Sectioning & Section Placement
	Assess RNA quality of the FFPE tissue block or from archived sections.
	Practice sectioning and section placement with a nonexperimental block before placing sections on slides.
	Section the tissue block and place sections on compatible blank slides using a water bath.
	Place tissue sections on the allowable area on the slide. Allowable area may vary depending on slide choice.
Tissue S	Slide Handling
	Ensure slides are completely dried after tissue placement, using a fan if necessary.
	Store the slides containing FFPE sections for up to six months in a desiccator at room temperature.
	After preparing tissue slides, stain sections according to the desired staining protocol in the Visium HD FFPE Tissue Preparation Handbook (CG000684).
Tissue S	Slide Processing
	After preparing tissue slides, process sections according to the desired staining protocols in the Visium HD FFPE Tissue Preparation Handbook (CG000684). The handbook contains information on tissue slide staining and imaging.
Vov Con	oidevetions for FF Comples
	siderations for FF Samples
Freezing	g and Embedding Perform either separate or simultaneous tissue freezing & embedding.
	Store frozen samples in a sealed container at -80°C for long-term storage.
Slide Ha	ndling
	Equilibrate slides to cryostat temperature before cryosectioning.
Cryosec	tioning
	Practice sectioning and section placement with a nonexperimental block before placing sections on slides.
	Equilibrate OCT tissue block to the cryostat chamber temperature for 30 min.
	Section the tissue block and place sections on compatible blank slides.
	Assess RNA quality of the tissue block.

Key Consi	derations for FF Samples
	Optional - assess tissue morphology via DAPI and H&E staining.
Tissue Sli	de Handling
	Maintain slides containing sections in a low moisture environment.
	Keep slides cold and transport slides on dry ice.
	Store tissue slides in a slide mailer at -80°C for up to two months.
Tissue Sli	de Processing
	After preparing tissue slides, follow the remaining steps in the Visium HD FF Tissue Preparation Handbook (CG000763). The handbook contains information on tissue slide staining and imaging.
Key Consi	derations for FxF Samples
Fixation,	Embedding, and Freezing
	Fix samples in 4% PFA or formaldehyde
	Cryopreserve samples in a sucrose gradient.
	Embed fixed samples in OCT and freeze.
Sample B	ock Storage
	Store frozen samples in a sealed container at -80°C for long-term storage.
Slide Han	dling
	Equilibrate slides to cryostat temperature before cryosectioning.
Cryosecti	oning
	Assess RNA quality of the tissue block.
	Optional - assess tissue morphology via DAPI and H&E staining.
	Practice sectioning and section placement with a nonexperimental block before placing sections on slides.
	Equilibrate OCT tissue block to the cryostat chamber temperature for 30 min.
	Section the tissue block and place sections on compatible blank slides.
Tissue Sli	de Handling
	Maintain slides containing sections in a low moisture environment.
	Keep slides cold and transport slides on dry ice.
	Store tissue slides in a slide mailer at -80°C for up to two months.
Tissue Sli	de Processing
	After preparing tissue slides, process sections according to the desired staining protocols in the Visium HD FxF Tissue Preparation Handbook (CG000764). The handbook contains information on tissue slide staining and imaging.



Step 1:

Probe Hybridization

1.0 Get Started	49
1.1 Probe Hybridization	50

1.0 Get Started

Each 10x Genomics reagent tube is enough for two 6.5 mm Visium HD Slides (four Tissue Slides). Number of reactions in reagent preparation tables refers to the number of Tissue Slides.

Items	s		10x PN	Preparation & Handling	Storage
Equil	librate to	room tempera	ture		
	0	FFPE Hyb Buffer	Tube: 2000423 Kit: 1000466 or 1000667	Thaw at room temperature. If precipitate persists, heat at 37°C until dissolved. Avoid vortexing to prevent bubble formation. Pipette mix 10x. Keep the buffer at room temperature after thawing and while performing the workflow.	-20°C
Place	e on ice				
		Human WT Probes v2 - RHS	Tube: 2000657 Kit: 1000466	Thaw at room temperature. Vortex and centrifuge briefly. Maintain on ice.	-20°C
		Human WT Probes v2 - LHS	Tube: 2000658 Kit: 1000466	Thaw at room temperature. Vortex and centrifuge briefly. Maintain on ice.	-20°C
	•	Mouse WT Probes v2 - RHS	Tube: 2000913 Kit: 1000667	Thaw at room temperature. Vortex and centrifuge briefly. Maintain on ice.	-20°C
	•	Mouse WT Probes v2 - LHS	Tube: 2000912 Kit: 1000667	Thaw at room temperature. Vortex and centrifuge briefly. Maintain on ice.	-20°C
Obta	in				
		Nuclease- free Water	-	-	Ambient
		10X PBS, pH 7.4	-	-	Ambient
		Visium Slide Seals	Component: 2000284 Kit: 1000669	See Tips & Best Practices. 3 pre-cut slide seals are necessary for the complete workflow.	Ambient
		10% Tween- 20	-	-	Ambient

Step 1: Probe Hybridization

1.1 Probe Hybridization

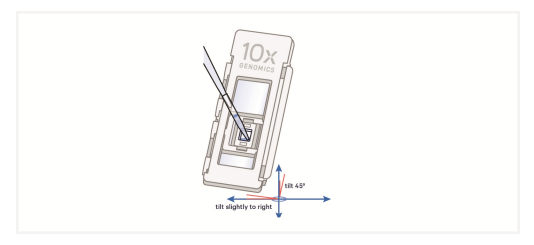


During reagent removal steps, ensure that **ALL liquid is removed** from wells. Use a P20 pipette to remove any remaining liquid. See Tips & Best Practices for guidance on Reagent Removal.

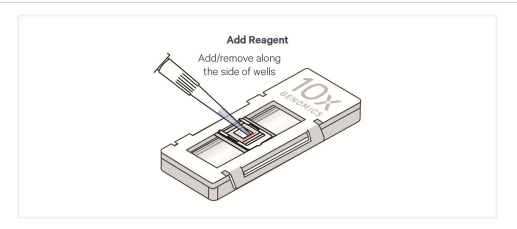
a. Prepare Pre-Hybridization Mix according to the table shortly before use. Add reagents in the order listed. Vortex and centrifuge briefly. Maintain at room temperature.

Pre-Hybridization Mix	10x PN	1Χ (μl)	2X +10% (μl)	4X +10% (μl)
Nuclease-free Water	-	134.2	295.3	590.6
10X PBS, pH 7.4	-	15.0	33.0	66.0
10% Tween-20	-	0.8	1.7	3.4
Total	-	150.0	330.0	660.0

- b. Retrieve Tissue Slide Cassettes containing H&E stained or IF stained sections.
- c. Peel back Visium Slide Seals.
- d. Using a pipette, remove all buffer from each well at well corners. Use a P20 pipette to remove any remaining liquid. See image below for proper liquid removal technique.



- e. Add 150 μl Pre-Hybridization Mix along the side of each well to uniformly cover tissue sections, without introducing bubbles.
- f. Re-apply Visium Slide Seal on each Tissue Slide Cassette.
- **g.** Incubate for **15 min** at **room temperature**.





h. Place a Low Profile Thermocycler Adapter in the thermal cycler. Prepare thermal cycler with the following incubation protocol and start the program. See Tips & Best Practices for more information on cassette incubation in a thermal cycler.

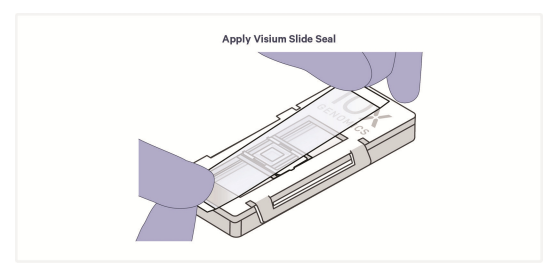
Lid Temperature	Reaction Volume	Run Time
50°C	100 μΙ	Overnight (16 - 24 h)
Step	Temperature	Time hh:mm:ss
Pre-equilibrate	50°C	Hold
Hybridization	50°C	Overnight (16 - 24 h)
Post Hybridization Wash	50°C	Hold

i. Prepare Probe Hybridization Mix according to the table shortly before use. Add reagents in the order listed. Keep at **room temperature**. Pipette mix 10x and centrifuge briefly.

	Probe Hybridization Mix	10x PN	1X (µl)	2X +10% (μl)	4X +10% (μl)
\bigcirc	FFPE Hyb Buffer	2000423	70.0	154.0	308.0
	Nuclease-free Water	-	10.0	22.0	44.0
	Human WT Probes v2 - RHS or Mouse WT Probes v2 - RHS	2000657 or 2000913	10.0	22.0	44.0
	Human WT Probes v2 - LHS or Mouse WT Probes v2 - LHS	2000658 or 2000912	10.0	22.0	44.0
	Total	-	100.0	220.0	440.0

- j. Remove Visium Slide Seals from Tissue Slide Cassettes.
- k. Remove all Pre-Hybridization Mix from each well.
- l. Add $100 \mu l$ room temperature Probe Hybridization Mix to each well.

m. Apply a new pre-cut Visium Slide Seal to each Tissue Slide Cassette and place on the Low Profile Thermocycler Adapter on the thermal cycler. Close the thermal cycler lid.



n. Skip Pre-equilibrate step to initiate Hybridization.



Step 2:

Probe Ligation

2.0 Get Started	54
2.1 Post-Hybridization Wash	55
2.2 Probe Ligation	57
2.3 Post-Ligation Wash	58

2.0 Get Started

Each 10x Genomics reagent tube is enough for two 6.5 mm Visium HD Slides (four Tissue Slides). Number of reactions in reagent preparation tables refers to the number of Tissue Slides.

Items	5		10x PN	Preparation & Handling	Storage
Equil	ibrate to	room tempera	ture		
		FFPE Post- Hyb Wash Buffer	Tube: 2000424 Kit: 1000466 or 1000667	Thaw at room temperature. If precipitate persists, heat at 37°C until dissolved. Vortex briefly. Flick tube until liquid settles at the bottom of the tube.	-20°C
	\bigcirc	2X Probe Ligation Buffer	Tube: 2000445 Kit: 1000466 or 1000667	Thaw at room temperature until no precipitate remains. Vortex and centrifuge briefly.	-20°C
	0	Post-Ligation Wash Buffer	Tube: 2000419 Kit: 1000466 or 1000667	Thaw at room temperature. If precipitate persists, heat at 37°C until dissolved. Vortex.	-20°C
Place	on ice				
	\circ	Probe Ligation Enzyme	Tube: 2000425 Kit: 1000466 or 1000667	Centrifuge briefly. Maintain on ice.	-20°C
Obta	in				
		Nuclease- free Water	-	-	Ambient
		20X SSC Buffer	-	-	Ambient

2.1 Post-Hybridization Wash

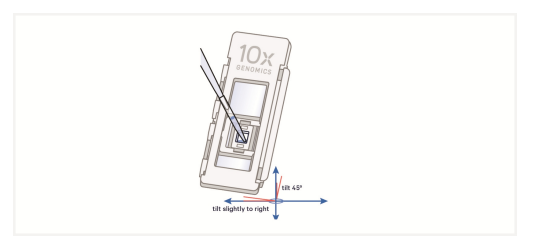
- a. Aliquot FFPE Post-Hyb Wash Buffer (495 μl/per 6.5 mm sample) and preheat to 50°C in a water bath or thermomixer. Maintain pre-heated wash buffer at 50°C throughout washes.
- **b.** Prepare 2X SSC Buffer according to the table. Add reagents in the order listed. Maintain at **room temperature**. DO NOT discard excess buffer, as it will be used for subsequent washes in the protocol.

SSC Buffer	Stock	Final	1X (µl)	2X +10% (μl)	4X +10% (μl)
SSC	20X	2X	150	315	615
Nuclease-free Water	-	-	1,350	2,835	5,535
Total	-		1,500	3,150	6,150

c. Remove Visium Tissue Slide Cassettes from Low Profile Thermocycler Adapter and place on a flat, clean work surface.



d. Peel back Visium Slide Seals and using a pipette, remove all Probe Hybridization Mix from each well. See Tips and Best Practices for slide seal removal guidance.





e. Immediately add 150 μ l pre-heated FFPE Post-Hyb Wash Buffer to each well. Avoid well drying or cooling to room temperature. Removal and addition of buffers should be done quickly.



f. Re-apply Visium Slide Seal on each Tissue Slide Cassette and place on Low Profile Thermocycler Adapter on pre-heated thermal cycler. Close thermal cycler lid. See Tips & Best Practices for more information on cassette incubation on a thermal cycler.



Ensure that Tissue Slide Cassette sits snugly on the Low Profile Thermocycler Adapter for all wash steps.

- **g.** Skip the Hybridization step on thermal cycler and initiate Post-Hybridization Wash.
- h. Incubate Visium Tissue Slide Cassettes in thermal cycler at 50°C for 5 min.



- i. Remove Tissue Slide Cassettes from Low Profile Thermocycler Adapter and place on a flat, clean work surface. Exercise caution, as slide is hot.
- -<u>Ö</u>-
- **j.** Peel back Visium Slide Seals and using a pipette, remove all FFPE Post-Hyb Wash Buffer from each well.



- **k. Immediately** add **150** μ l pre-heated FFPE Post-Hyb Wash Buffer to each well. Avoid well drying or cooling to room temperature. Removal and addition of buffers should be done quickly.
- Re-apply Visium Slide Seal on each Tissue Slide Cassette and place on Low Profile Thermocycler Adapter on pre-heated thermal cycler. Close thermal cycler lid.
- m. Incubate in thermal cycler at 50°C for 5 min.



- **n. Repeat** steps i-m one more time for a total of three washes.
- **o.** Peel back Visium Slide Seals and remove FFPE Post-Hyb Wash Buffer from each well.
- p. Add 150 μl 2X SSC Buffer to each well and re-apply Visium Slide Seal.
- **q.** Let the Tissue Slide Cassettes cool to **room temperature** (**~3 min**) before proceeding to the next step.

2.2 Probe Ligation

a. Place a Low Profile Thermocycler Adapter onto a thermal cycler (if not already placed), prepare with the following incubation protocol, and start the program.

Lid Temperature	Reaction Volume	Run Time
37°C (lid may be turned off if the instrument does not enable 37°C)	100 μΙ	1 h
Step	Temperature	Time hh:mm:ss
Pre-equilibrate	37°C	Hold
Ligation	37°C	01:00:00
Hold	4°C	Hold

b. Prepare Probe Ligation Mix according to the table shortly before use. Add reagents in the order listed. Pipette mix 10x and centrifuge briefly. DO NOT vortex. Maintain on ice.

	Probe Ligation Mix	10x PN	1X (µl)	2X +10% (μl)	4X +10% (μl)
	Nuclease-free Water	-	24.0	52.8	105.6
\circ	2X Probe Ligation Buffer	2000445	30.0	66.0	132.0
\bigcirc	Probe Ligation Enzyme	2000425	6.0	13.2	26.4
	Total	-	60.0	132.0	264.0

- **c.** Remove Visium Slide Seals from each Tissue Slide Cassette and remove all 2X SSC Buffer from each well.
- **d.** Add **60** μ **l** Probe Ligation Mix to each well, without introducing bubbles. Tap Tissue Slide Cassettes gently to ensure uniform coverage.
- **e.** Apply a new pre-cut Visium Slide Seal on each Tissue Slide Cassette and place on the Low Profile Thermocycler Adapter on the pre-heated thermal cycler. Close the thermal cycler lid.
- **f.** Skip Pre-equilibrate step to initiate Ligation.

2.3 Post-Ligation Wash



Use room temperature Post-Ligation Wash Buffer at first wash step (step 2.3f). Use pre-heated Post-Ligation Wash Buffer at second wash step (step 2.3k).

- a. If the stopping point after the Post-Ligation Wash will not be used, remove Visium HD slide mailer from -80°C. Remove slide mailer from mylar bag. Keep slide mailer upright, capped, and thaw at room temperature for 30 min 3 h. Two Visium HD slides may be prepared at the same time.
- **b.** Pre-heat Post-Ligation Wash Buffer (110 μ l/sample) to 57°C. Only 100 μ l per 6.5 mm sample is needed.
- **c.** Remove the Tissue Slide Cassettes from the Low Profile Thermocycler Adapter and place on a flat, clean work surface.
- **d.** Immediately prepare thermal cycler with the following incubation protocol and start the program.

Lid Temperature	Reaction Volume	Run Time
57°C	100 μΙ	-
Step	Temperature	Time
Incubate	57°C	Hold

e. Remove Visium Slide Seals from each Tissue Slide Cassette and using a pipette, remove all Probe Ligation Mix from each well.



f. Immediately add **100** μ **l room temperature** Post-Ligation Wash Buffer to each well. Removal and addition of buffers should be done quickly.



g. Apply a new pre-cut Visium Slide Seal on each Tissue Slide Cassette and place on Low Profile Thermocycler Adapter on pre-heated thermal cycler. Close thermal cycler lid. See Tips & Best Practices for more information on cassette incubation in a thermal cycler.



Ensure that Tissue Slide Cassette sits snugly on the Low Profile Thermocycler Adapter for all wash steps.

- h. Incubate at 57°C for 5 min.
- **i.** Remove Tissue Slide Cassettes from the Low Profile Thermocycler Adapter and place on a flat, clean work surface.
- **j.** Peel back Visium Slide Seals and using a pipette, remove all Post-Ligation Wash Buffer from each well.



k. Add **100** μ**l pre-heated** Post-Ligation Wash Buffer to each well.

- **1.** Re-apply Visium Slide Seal on Tissue Slide Cassette and place on Low Profile Thermocycler Adapter on pre-heated thermal cycler. Close thermal cycler lid.
- m. Incubate at 57°C for 5 min.
- **n.** Remove Tissue Slide Cassettes from Low Profile Thermocycler Adapter and place on a flat, clean work surface.
- **o.** Peel back Visium Slide Seals and using a pipette, remove all Post Ligation Wash Buffer from each well.
- **p.** Add **150** μ **l** 2X SSC Buffer prepared in 2.1 Post-Hybridization Wash on page 55 to each well.
- q. Remove all 2X SSC Buffer from each well.
- r. Add 150 μl 2X SSC Buffer to each well.
- s. Re-apply Visium Slide Seals on each Tissue Slide Cassette.
- t. Allow Tissue Slide Cassettes to come to room temperature for 5 min.



u. Store at 4°C for up to 24 h or proceed to next step.



Step 3:

Visium HD Slide Preparation

3.0 Get Started	61
3.1 Visium HD Slide Wash	62

3.0 Get Started

Each 10x Genomics reagent tube is enough for two 6.5 mm Visium HD Slides. Number of reactions in reagent preparation tables refers to the number of Visium HD slides.



The Visium CytAssist instrument requires firmware version 2.0.0 or higher for this protocol.

Items		10x PN	Preparation & Handling	Storage
Obtain				
	Nuclease- free Water	-	-	Ambient
	20X SSC	-	-	Ambient
	Visium Cassette	Component: 3001830/ 3001831 Kit: 1000669	See Cassette Assembly Quick Reference Card (CG000730)	Ambient

3.1 Visium HD Slide Wash

a. If Visium HD Slide has not been thawed, remove Visium HD Slide mailer from -80°C. Remove slide mailer from mylar bag. DO NOT uncap slide mailer. Keep slide mailer upright and thaw at room temperature for 30 min - 3 h. Two Visium HD Slides may be prepared at the same time, though only two tissue slides should be processed at a time.

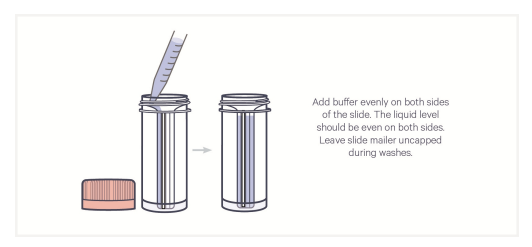
DO NOT touch Visium HD Slide spacer during slide washes.

b. Prepare 0.1X SSC Buffer according to the table below in **two** 50-ml centrifuge tubes (e.g. two tubes for one Visium HD slide). Vortex. Maintain at room temperature. 0.1X SSC prepared at this step is sufficient for remaining steps in the protocol.

0.1X SSC Buffer	Stock	Final	1 Visium HD Slide (ml)*	2 Visium HD Slides (ml)*
Nuclease-free Water	-	-	59.7	119.4
SSC	20X	0.1X	0.3	0.6
Total	-		60.0	120.0

^{*}Volumes are in ml instead of µl

- c. Open slide mailer.
- d. Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on one side of the slide.
- e. Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on the other side of the slide, so that both sides of the slide are immersed in 0.1X SSC Buffer.



f. Incubate at **room temperature** for **1 min**. DO NOT close the mailer.

g. Remove 0.1X SSC Buffer by securing slide in place with one finger and pouring SSC Buffer from mailer.



Three 0.1X SSC 5 min washes:

h. 5 Min Wash 1:

- Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on one side of the slide.
- Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on the other side of the slide, so that both sides of the slide are immersed in 0.1X SSC Buffer.
- Incubate at room temperature for 5 min.
- Remove 0.1X SSC Buffer by securing slide in place with one finger and pouring SSC Buffer from mailer.

i. 5 Min Wash 2:

- Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on one side of the slide.
- Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on the other side of the slide, so that both sides of the slide are immersed in 0.1X SSC Buffer.
- Incubate at **room temperature** for **5 min**.
- Remove 0.1X SSC Buffer by securing slide in place with one finger and pouring SSC Buffer from mailer.

i. 5 Min Wash 3:

• Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on one side of the slide.

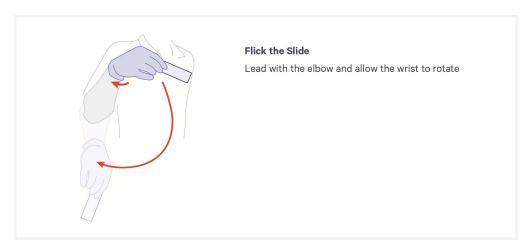
- Add 7 ml 0.1X SSC Buffer slowly along the side of the mailer on the other side of the slide, so that both sides of the slide are immersed in 0.1X SSC Buffer.
- Incubate at room temperature for 5 min.



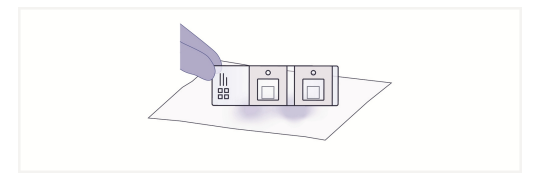
• Remove Visium HD Slide from mailer.

During the last wash, save SSC Buffer in the mailer in case Visium HD Slide needs additional immersion.

- k. Inspect the Visium HD Slide for debris. If debris is visible, immerse the slide back in the mailer containing SSC and quickly remove.
- 1. Flick the Visium HD Slide.



- m. Ensure back of Visium HD Slide (side without spacers) is dry. If necessary, remove excess moisture from the back of the slide with a lint-free laboratory wipe.
- **n.** Place a fresh, lint-free laboratory wipe on the work surface.
- o. Gently touch long edge of Visium HD Slide on a lint-free laboratory wipe 5x to remove excess SSC buffer.



p. Inspect the Visium HD Slide and ensure it is free of particulate matter. If debris is visible, immerse the slide back in the mailer containing SSC and

- quickly remove. Repeat flicking and wiping as previously described.
- q. Record Visium HD Slide serial number.
- r. Place Visium HD Slide in a new 6.5 mm Visium Cassette. See Visium Cassette S3 Quick Reference Card (CG000730) for assembly instructions.
- **s.** Add $100 \mu l$ 0.1X SSC to each well in the cassette.
- t. Apply a new Visium Slide Seal on the Visium Cassette.
- u. Leave 0.1X SSC Buffer in the Visium Cassette at room temperature until Visium HD Slide equilibration. DO NOT exceed 2 h before proceeding with a CytAssist run. Keep Visium Cassette and Visium HD slide free from debris.



Step 4:

Probe Release & Extension

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4.3 Probe Elution	82

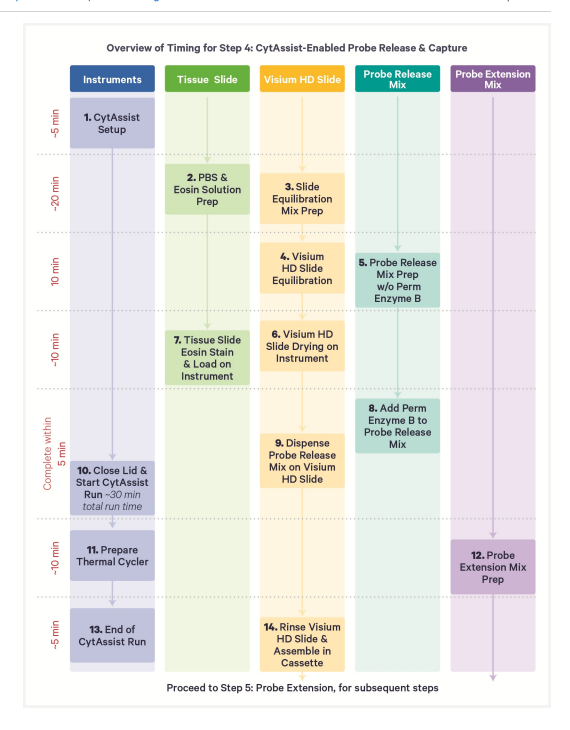
4.0 Get Started

Each 10x Genomics reagent tube is enough for two 6.5 mm Visium HD Slides (four tissue slides). Number of reactions in reagent preparation tables refers to the number of Tissue Slides. After instrument run, number of reactions refers to the number of wells in a Visium Cassette.



Ensure that the Visium CytAssist is powered on, running firmware version 2.0.0 or higher, is clean, and ready to perform an experimental run.

Item	s		10x PN	Preparation & Handling	Storage
Equi	librate to	o room temperature			
		2X RNase Buffer	Tube: 2000411 Kit: 1000668	Thaw at room temperature, vortex, centrifuge briefly.	-20°C
		Extension Buffer	Tube: 2000409 Kit: 1000668	Thaw at room temperature, vortex, centrifuge briefly.	-20°C
Plac	e on ice				
	•	RNase Enzyme	Tube: 3000605 Kit: 1000668	Pipette mix, centrifuge briefly. Maintain on ice until ready to use.	-20°C
		Extension Enzyme	Tube: 2000389 Kit: 1000668	Pipette mix, centrifuge briefly. Maintain on ice until ready to use.	-20°C
	•	Perm Enzyme B	Tube: 3000553 Kit: 1000668	Pipette mix, centrifuge briefly. Perm Enzyme B is added to the Probe Release Mix immediately before running the CytAssist instrument.	-20°C
Obta	in				
		Visium HD Slide	-	Obtain washed Visium HD Slide.	Ambient
		Nuclease-free Water	-	-	Ambient
		Tris 1 M, pH 8.0 (Tris-HCl)	-	Manufacturer's recommendations.	Ambient
		Alcoholic Eosin	-	Manufacturer's recommendations.	Ambient
		10X PBS	-	Use 10X PBS stock to prepare 1X PBS.	Ambient
		8 M KOH Solution	-	Manufacturer's recommendations.	Ambient
		Qiagen Buffer EB	-	Manufacturer's recommendations.	Ambient
		Visium Slide Seals	Component: 2000283 Kit: 1000669	See Tips & Best Practices.	Ambient



4.1 CytAssist-Enabled Probe Release & Capture

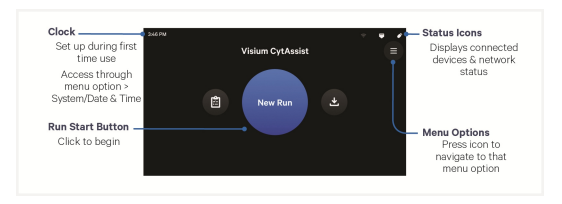
Videos demonstrating key techniques in this section are available on the 10x Genomics website.

If processing more than two tissue slides, keep remaining tissue slides at 4°C with 2X SSC buffer. For more guidance, see Appendix on page 120.



a. Ensure that Visium CytAssist is powered on, clean, and ready for an experimental run. Visium CytAssist firmware version 2.0.0 or higher is required for this protocol.

The home screen is the most common state of the instrument. There are several key functions accessible directly from the home screen.



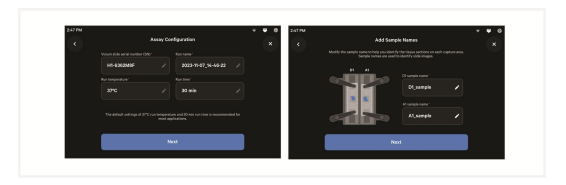
- **b.** Press blue New Run button on the touchscreen to initiate run.
- **c.** Enter the following assay configuration information:



 Visium Slide serial number. Ensure serial number is accurate. If serial number is entered in the wrong format, the check mark button to proceed will be grayed out. Record serial number for future use.

A slide scanner may be used to scan the Visium HD slide for automatic serial number input prior to cassette assembly. See Visium HD Spatial Gene Expression Protocol Planner (CG000698) for compatible part numbers. The barcode scanner cannot scan the Visium HD Slide after cassette assembly, as the QR code will not be visible.

- Custom run name, temperature, and time (37°C for 30 min is recommended for most applications)
- **d.** Enter the following sample information:
 - Sample names and the locations of each sample on the instrument (A1 for right side, D1 for left side)



e. Prepare 1X PBS. Maintain at room temperature.

1X PBS	10x PN	2 Tissue Slides (μl) (includes overage)	4 Tissue Slides (μΙ) (includes overage)
Nuclease-free Water	-	5,940	11,880
10X PBS	-	660	1,320
Total	-	6,600	13,200

f. Prepare 10% Eosin. Vortex and centrifuge briefly. Eosin should be prepared fresh for each CytAssist instrument run.

10% Eosin	10x PN	1Χ (μl)	2X +10% (μl)	4X +10% (μl)
Alcoholic Eosin	-	15	33	66
1X PBS	-	135	297	594
Total	-	150	330	660

g. Prepare Visium HD Slide Equilibration Mix according to the table below. Pipette mix and centrifuge briefly. Maintain on ice.

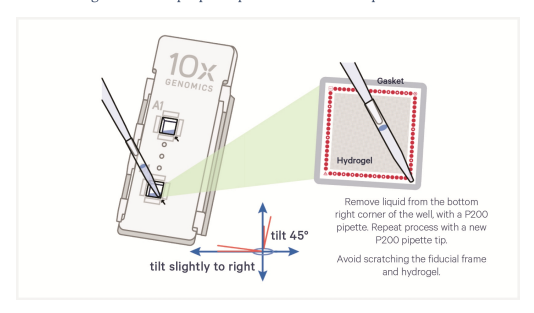
Visium HD Slide Equilibration Mix	10x PN	1X (µl)	2X +10% (µl)	4X +10% (μl)
Nuclease-free Water		44.0	96.8	193.6
2X RNase Buffer	2000411	50.0	110.0	220.0
RNase Enzyme	3000605	6.0	13.2	26.4
Total	-	100.0	220.0	440.0

h. Retrieve Visium Cassette with Visium HD Slide.



- i. Remove Visium Slide Seal and perform the following:
 - Hold the cassette at a 45° angle slightly tilted to the right so that the buffer pools to the lower right hand corner of the well.
 - Set a P200 pipette to 200 μl.

- Place the pipette tip at the bottom right corner of the well without scratching the fiducial frame or hydrogel.
- Remove 0.1X SSC from the well.
- Repeat this process using a new P200 tip.
- Once all the liquid has been removed, repeat for the remaining well.
- See image below for proper liquid removal technique.



- j. Add 100 µl Visium HD Slide Equilibration Mix to each well in the cassette. Gently tap to ensure uniform coverage of the Capture Area.
- k. Apply a new uncut Visium Slide Seal on the Visium Cassette and incubate at room temperature for 10 min.
- 1. Prepare Probe Release Mix during Visium HD Slide equilibration. DO NOT prepare Probe Release Mix for more than two tissue slides at a time. Pipette mix thoroughly until solution is homogenous. Avoid generating bubbles. Maintain on ice.



Probe Release Mix will also require Perm Enzyme B, which will be added at step af.

Probe Release Mix	10x PN	2 Tissue Slides (µl) (includes overage)
2X RNase Buffer	2000411	20.0
RNase Enzyme	3000605	17.5
Total	-	37.5

m. Remove Visium Slide Seal and using a P200 pipette, remove Visium HD Slide Equilibration Mix from the right corner of each well in the cassette without scratching the fiducial frame or hydrogel. Use a P200 pipette with a fresh tip to remove any remaining liquid. See step i for proper liquid removal technique.



Failure to remove Slide Equilibration Mix completely may result in reduced assay performance.



n. Remove top half of Visium Cassette, leaving Visium HD Slide resting in bottom half of Visium Cassette. Rest top half of Visium Cassette such that the gaskets face up, as shown in the image below. This minimizes the risk of introducing debris on the top half of the cassette after disassembly. Avoid introducing debris to the cassette after disassembly. See Visium Cassette S3 Quick Reference Card (CG000730) for more information on cassette disassembly.



o. Remove Visium HD Slide from Visium Cassette and inspect the slide. (avoid touching active surface). If necessary, remove excess moisture from the back of the slide with a lint-free laboratory wipe. DO NOT flick the Visium HD slide.

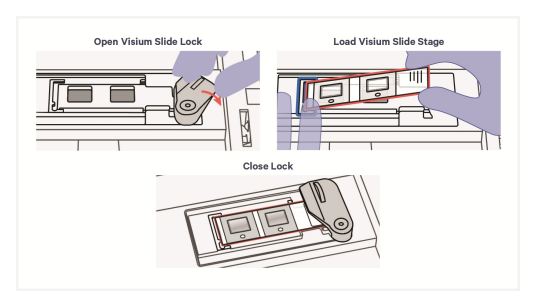
The time between completing step o and starting step q should not exceed 5 min.



Place a cover over the Visium Cassette and save for use after the instrument run, keeping all parts free from debris. Wiping the front of the Visium HD Slide may damage the slide.



p. Load Visium HD Slide against the grooves of Visium Slide Stage, using one hand to load the slide and the other to close the Visium Slide Lock.



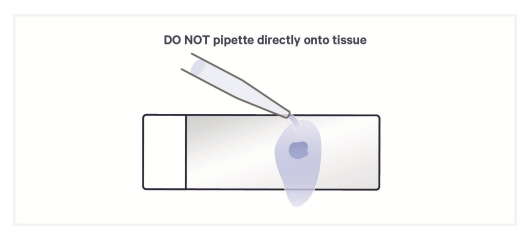


q. Allow Visium HD Slide to dry on the Visium Slide Stage for 10 min. Inspect entire spacer chamber. Look at the slide at different angles to confirm the presence of liquid. If liquid remains on the slide or anywhere in the spacer chamber, continue drying and proceed immediately when no liquid remains within the spacer chamber.

Ensure area around the Visium CytAssist instrument is free from debris. While the Visium HD Slide is drying, prepare Tissue Slides as described in steps r-ae.



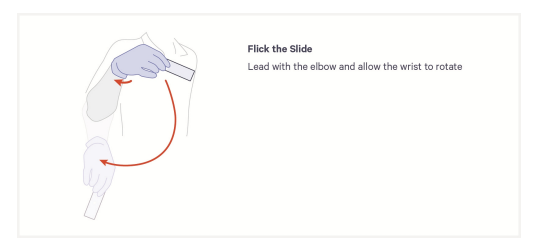
- r. Retrieve Tissue Slide Cassettes.
- s. Using a pipette, remove all 2X SSC Buffer from each well of the Tissue Slide Cassettes.
- t. Remove tissue slides from Tissue Slide Cassettes.
- **u.** Add **100** μ**l** 10% Eosin to uniformly cover each tissue section per slide.
- v. Incubate 1 min at room temperature.
- w. Remove 10% Eosin by holding slide at an angle over a liquid waste container.
- x. While holding the slide over the liquid waste container, rinse with 1 ml 1X PBS. DO NOT pipette directly onto tissue.



y. Repeat step x two more times for a total of three washes.



z. Flick the tissue slide to remove excess PBS.



- aa. Inspect the slide for the presence of any large droplets. If present, repeat flicking.
- ab. Once the slide has no liquid droplets on the tissue surface, remove any excess PBS with a lint-free laboratory wipe in areas outside of the tissue, without touching the tissue sections.

Lint-free laboratory wipes are strongly recommended to ensure no debris is introduced. See the Visium HD Spatial Gene Expression Protocol Planner (CG000698) for a list of tested part numbers.

- ac. Gently remove any remaining liquid on the back of the slides with a lintfree laboratory wipe.
- ad. If not using the Visium CytAssist Alignment aid, proceed directly to step af.
- ae. If using the Visium CytAssist Alignment Aid, use the aid to draw alignment marks by tracing the mark guides onto the tissue slide. See Visium CytAssist Alignment Aid on page 33 for more information.



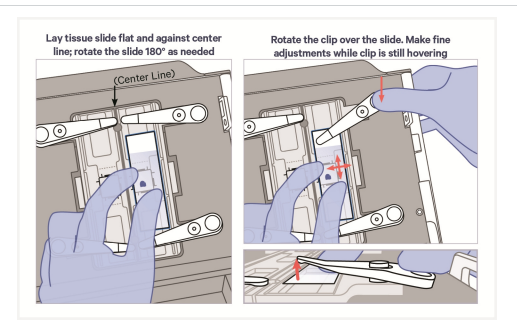
DO NOT trace the square window.



af. Load tissue slides into Visium CytAssist. Ensure tissue section is completely dry prior to instrument run. Examine the aligned tissue slide from multiple angles to ensure that tissue is completely dry. See Instrument Loading Guidelines on page 37 for more information.



Complete alignment aid use and tissue slide alignment before Visium HD slide is done drying.





Before proceeding to next step, ensure that Visium HD slide is completely dry. See step q.

ag. Pipette mix Perm Enzyme B (PN-3000553) and centrifuge briefly. Remove Probe Release Mix from ice. Add 2.5 μ l of Perm Enzyme B to 37.5 μ l of Probe Release Mix (prepared at step 4.11) to complete Probe Release Mix. Pipette mix 15x with pipette set to 30 μl. Avoid generating bubbles. Centrifuge for **5 sec**.



The time between adding Perm Enzyme B to Probe Release Mix and starting the Visium CytAssist instrument run should be less than 5 min.

Probe Release Mix	10x PN	2 Tissue Slides (µl) (includes overage)
2X RNase Buffer	2000411	Already added in step I.
RNase Enzyme	3000605	Already added in step I.
Perm Enzyme B	3000553	2.5
Total	-	40

ah. Using a P20 pipette, slowly aspirate 17 μl of Probe Release Mix and inspect the pipette tip. Ensure that no bubbles were drawn up into the pipette tip.

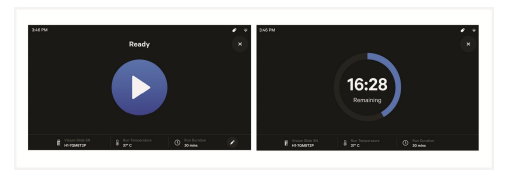
ai. Slowly dispense $17 \mu l$ of Probe Release Mix into the center of each spacer well on the Visium HD Slide, using a fresh pipette tip for each dispense. Do not push plunger beyond first stop to avoid generating bubbles.



aj. Close lid and press Next.

The home screen will now display a play symbol and run information along the bottom of the screen.

- ak. Press play button to start the run. 37°C for 30 min is recommended for most applications.
 - Midrun progress bar will show the time remaining in the run.



• Yellow before a run begins indicates a user-recoverable error. If an error occurs, follow on-screen prompts.

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

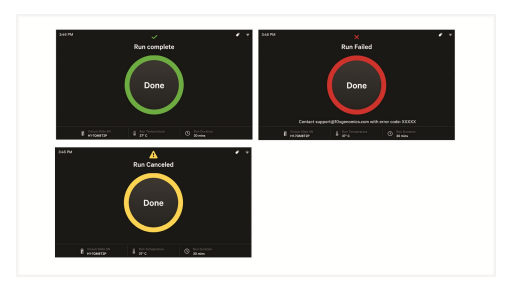
Lid Temperature	Reaction Volume	Run Time
53°C (lid may be turned off if the instrument does not enable 53°C)	100 μΙ	60 min
Step	Temperature	Time
Pre-equilibrate	53°C	Hold
Probe Extension 1	53°C	00:30:00
Cool Down	4°C	00:03:00
Hold	4°C	Hold
Probe Extension 2	53°C	00:30:00
Cool Down	4°C	00:03:00
Hold	4°C	Hold

am. Prepare Probe Extension Mix. Pipette mix. Centrifuge briefly. Maintain on ice.

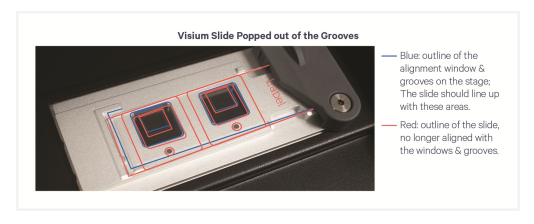
Probe Extension Mix	10x PN	1X (µl)	2X +10% (μl)	4X +10% (μl)
Extension Buffer	2000409	147.0	323.4	646.8
Extension Enzyme	2000389	3.0	6.6	13.2
Total	-	150.0	330.0	660.0



- an. At the end of a run, the button will display "Done" and a "Run Info" tab at the bottom of the screen. DO NOT allow sample to sit in the Visium CytAssist after run completion. Immediately move to next step.
 - Green indicates a successfully completed run.
 - Red indicates a failed run/error
 - Yellow at the end of a run indicates an incomplete run.
 - For more information on errors, consult the Visium CytAssist Instrument User Guide (CG000542).



ao. Click "Done" button and open lid. DO NOT power off the instrument at this time, as it needs to process support data. It is normal for the Visium slide to move after opening instrument.

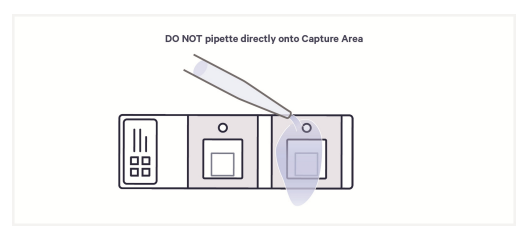


ap. Immediately remove Visium HD Slide. It is normal if tissue remains of tissue slides after run completion.

Leave the CytAssist lid open to facilitate cleaning of the instrument at a safe stopping point.

aq. While holding Visium HD Slide over liquid waste container, rinse each Capture Area with 1 ml Buffer EB. DO NOT pipette directly onto Capture Areas, which are surrounded by the fiducial frames.

Rinse slide near Visium CytAssist instrument to ensure prompt washing of Capture Areas.



- ar. Repeat step al two more times for a total of three washes per Capture Area.
- as. Place Visium HD Slide in the same Visium Cassette from earlier in this step.
 - Some moisture remaining on the Visium HD Slide is normal.
- at. Proceed immediately to Probe Extension. CytAssist instrument should only be cleaned and data transferred at a safe stopping point.

4.2 Probe Extension

Probe Extension occurs over two rounds, Probe Extension 1 and Probe Extension 2.



a. Add 75 μl Probe Extension Mix to each well. Gently tap Visium Cassette to ensure uniform coverage of Capture Area. Keep remaining Probe Extension Mix on ice.



- **b.** Apply new uncut Visium Slide Seal on Visium Cassette and place on the Low Profile Thermocycler Adapter on pre-heated thermal cycler. Close thermal cycler lid.
- **c.** Skip Pre-equilibrate step to initiate Probe Extension 1.
- d. After the first Cool Down, open thermal cycler lid and remove cassette from Low Profile Thermocycler Adapter,
- e. Peel back Visium Slide Seal and remove Probe Extension Mix from each well.
- f. Add 75 μl Probe Extension Mix to each well.



Gently tap Visium Cassette to ensure uniform coverage of Capture Area.

- g. Re-apply Visium Slide Seal and place cassette on Low Profile Thermocycler Adapter. Close thermal cycler lid.
- **h.** Skip Hold step to initiate Probe Extension 2.



- i. Sample may remain at 4°C in the thermal cycler for up to 24 h.
- j. Clean CytAssist instrument. Consult Visium CytAssist Instrument User Guide (CG000542) for more information.
- k. Export run data and review CytAssist images. Consult Visium CytAssist Instrument User Guide (CG000542) for more information on exporting run data. If images look abnormal, contact support@10xgenomics.com.

4.3 Probe Elution

a. Prepare fresh 0.08 M KOH Mix according to the appropriate table shortly before use, adding reagents in the order listed. Vortex and centrifuge briefly. Maintain at room temperature. Discard unused KOH Mix after use. Volumes provided are sufficient for use with up to 4 reactions.

6.5 mm Slides				
КОН Mix	Stock	Final	1Χ (μΙ)	4Χ +10% (μΙ)
Nuclease-free Water	-	-	49.5	217.8
КОН	8 M	0.08 M	0.5	2.2
Total	-	-	50.0	220.0

- b. Remove Visium Cassette from Low Profile Thermocycler Adapter and place on a flat, clean work surface after Probe Extension is complete.
- c. Remove Visium Slide Seal and using a pipette, remove all Probe Extension Mix from wells.
- **d.** Add **150** μ**l** Buffer EB to each well.
- e. Remove all Buffer EB from wells.
- f. Add 50 µl 0.08 M KOH Mix to each well. Gently tap Visium Cassette to ensure uniform coverage of Capture Area.
- **g.** Incubate at **room temperature** for **10 min**.
- **h.** Add $3 \mu l 1 M$ Tris-HCl pH 8.0 to a tube in an 8-tube strip for each sample.



i. Transfer all solution for each sample containing the probes to a tube in an 8-tube strip containing 1 M Tris-HCl. Vortex, centrifuge briefly, and place on ice. DO NOT leave behind any solution in the wells. See Tips & Best Practices for reagent removal instructions.



Step 5:

Pre-Amplification and SPRIselect

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5.0 Get Started

Number of reactions in reagent preparation tables refers to the number of samples from Probe Elution.

Item		10x PN	Preparation & Handling	Stora	ge	
Equilibrate t	to room tempe	rature				
	TS Primer Mix B	Tube: 2000537 Kit: 1000688	Thaw at room temperature, vortex, and centrifuge briefly.	-20°C	;	
	Beckman Coulter SPRIselect Reagent	-	Manufacturer's recommendations.	-		
Place on ice						
	Amp Mix B	Tube: 2000567 Kit: 1000668	Vortex, centrifuge briefly.	-20°C	;	
Obtain						
	Qiagen Buffer EB	-		Ambie	ent	
	10x Magnetic Separator	Component: 2001212 Kit: 1000499	See Tips & Best Practices.	Ambie	ent	
	80% Ethanol	-	Prepare fresh. Prepare 1 room temperature.	ml per re	action. S	tore at
			80% Ethanol Store at room temperature S	itock	Final	1000 μΙ 1Χ
			100% Ethanol 1	100%	80%	800 µl
			Nuclease-free Water	-	-	200 µl

5.1 Pre-Amplification

a. Prepare Pre-Amplification Mix on ice. Add reagents in the order listed. Pipette mix and centrifuge briefly. Maintain on ice.

	6.5 mm Slides						
	Pre-Amplification Mix	PN	1Χ (μl)	2X + 10% (µl)	4X + 10% (μl)		
	Nuclease-free Water		19.5	42.9	85.8		
\bigcirc	Amp Mix B	2000567	25.0	55.0	110.0		
	TS Primer Mix B	2000537	2.5	5.5	11.0		
	Total	-	47.0	103.4	206.8		

- b. Add 47 µl Pre-Amplification Mix to each tube from 4.3 Probe Elution on page 82 (regardless of slide type). Pipette mix and centrifuge briefly.
- **c.** Incubate in a thermal cycler with the following protocol.

Lid Temperature	Reaction Volume	Run Time
105°C	100 μΙ	~25 min
Step	Temperature	Time hh:mm:ss
1	98°C	00:03:00
2	98°C	00:00:15
3	63°C	00:00:20
4	72°C	00:00:30
5	Go to Step 2, repeat 9X	for a total of 10 cycles
6	72°C	00:01:00
7	4°C	00:03:00
8	4°C	Hold

5.2 Pre-Amplification Cleanup - SPRIselect

If using MicroAmp 8-Tube Strips, use the high position (magnet•High) only throughout the protocol.

- a. Vortex to resuspend SPRIselect reagent. Add 120 µl SPRIselect reagent (1.2X) to each pre-amplification reaction in an 8-tube strip (100 µl) and pipette mix 15x (pipette set to 130 µl).
- b. Incubate 5 min at room temperature.
- **c.** Place on the magnet•**High** until solution clears .
- **d.** Remove supernatant.
- e. Add 300 µl 80% ethanol to pellet. Wait 30 sec. Pipette carefully as 300 µl is at tube limit.
- f. Remove ethanol.
- g. Add 200 µl 80% ethanol to pellet. Wait 30 sec.
- h. Remove ethanol.
- i. Centrifuge briefly and place on magnet•Low.
- j. Remove any remaining ethanol without disturbing the beads. Air dry for 2 min. DO NOT exceed 2 min as this will decrease elution efficiency.
- **k.** Remove from magnet. Add 105 μl Buffer EB. Pipette mix 15x (pipette set to 100 μl).
- 1. Incubate 2 min at room temperature.
- m. Place tube strip on magnet •High until solution clears
- **n.** Transfer **100** μ **l** sample to a new tube strip.



o. Store at 4°C for up to 72 h, -20°C for up to 4 weeks, or proceed to next step.



Step 6:

Visium HD Spatial Gene Expression – Probebased Library Construction

6.0 Get Started	88
6.1 Cycle Number Determination – qPCR	89
6.2 Sample Index PCR	91
6.3 Post-Sample Index PCR Cleanup – SPRIselect	93
6 4 Post-Library Construction QC	94

6.0 Get Started

Item		10x PN	Preparation & Handling	Storage
Equilibrate t	o room tempera	ature		
	TS Primer Mix B	Tube: 2000537 Kit: 1000688	Thaw at room temperature, vortex, and centrifuge briefly.	-20°C
	Dual Index Plate TS Set A	Tube: 3000511 Kit: 1000251	Thaw at room temperature, vortex, and centrifuge briefly.	-20°C
	Beckman Coulter SPRIselect Reagent	-	Manufacturer's recommendations.	-
	Library QC Reagents		Use Agilent Bioanalyzer, Perkin Elme or Agilent TapeStation for QC. Obtain appropriate reagents based on methology.	า
Place on ice				
	Amp Mix B	Tube: 2000567 Kit: 1000668	Vortex, centrifuge briefly.	-20°C
	KAPA SYBR Fast qPCR Master Mix	-	Manufacturer's recommendations.	-
Obtain				
	Nuclease- free Water	-	-	Ambient
	Qiagen Buffer EB	-		Ambient
	10x Magnetic Separator	Component: 2001212 Kit: 1000499	See Tips & Best Practices.	Ambient
	80% Ethanol	-	Prepare fresh. Prepare 400 µI per reaction.	Ambient

6.1 Cycle Number Determination - qPCR

- a. Dilute TS Primer Mix B 1:10 in nuclease-free water. The amount of diluted primer mix required to generate the qPCR mixes is described in the table below.
- **b.** Prepare qPCR Mix on ice according to the table below. Add reagents in the order listed. Vortex and centrifuge briefly. Maintain on ice.

A passive reference dye, such as ROX, may be required. Consult the Roche KAPA SYBR FAST qPCR Kit website or the appropriate qPCR system manufacturer for guidance.

Gene Expression qPCR Mix	Stock	Final	1X (µl)	3X* + 10% (µl)	5X* + 10% (μl)
KAPA SYBR FAST qPCR Master Mix Minimize light exposure	2X	1X	5.0	16.5	27.5
Diluted TS Primer Mix B	-	-	1.0	3.3	5.5
Nuclease-free Water	-	-	3.0	9.9	16.5
Total			9.0	29.7	49.5
*Includes one negative cont	trol				

- c. Add 9 µl qPCR Mix to one well per sample in a qPCR plate (a well for negative control may be included).
- d. Dilute 2 μl sample from Pre-Amplification Cleanup SPRIselect in 8 μl nuclease-free water. Pipette mix, centrifuge briefly.
- e. Transfer $1 \mu l$ diluted sample from Pre-Amplification Cleanup SPRIselect to each qPCR plate well containing qPCR Mix. For the negative control, add 1 μl nuclease-free water to the corresponding well. Pipette mix.
- f. Apply seal and centrifuge briefly. Record which sample is in which well of the qPCR plate.

g. Prepare a qPCR system with the following protocol, place the plate in the thermal cycler, and start the program.

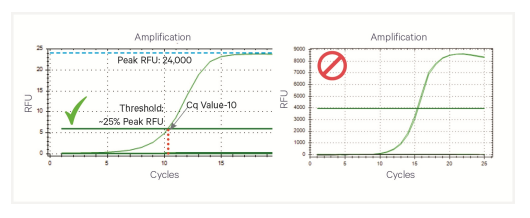
Lid Temperature	Reaction Volume	Run Time
105°C	10 μΙ	35 min
Step	Temperature	Time hh:mm:ss
1	98°C	00:03:00
2	98°C	00:00:05
3	63°C	00:00:30
	Read signal	
4	Go to step 2, 24x (total of 25 cycles) -	

h. Record the Cq value for each sample.

Set the y-axis to a linear scale. Plot RFU on the y-axis if not using a reference dye, or ΔRn if using a reference dye. The threshold for determining the Cq value should be set along the exponential phase of the amplification plot, at ~25% of the peak fluorescence value.

If the Cq value is > 15 cycles, refer to 3. High Cq Value on page 112 or contact support@10xgenomics.com before proceeding.

Representative qPCR Amplification Plots



6.2 Sample Index PCR



- **a.** Choose the appropriate sample index sets to ensure that no sample indices overlap in a multiplexed sequencing run. Record the 10x Sample Index name (PN-3000511 Plate TS Set A well ID) used.
- **b.** Prepare Amplification Master Mix and pipette mix 10x shortly before use.

	Amplification Master Mix	10x PN	1Χ (μl)	2X +10% (μl)	4X +10% (μl)
	Nuclease-free Water	-	45	99	198
\bigcirc	Amp Mix B	2000567	25	55	110
	Total	-	70	154	308

- **c.** Add **70 μl** Amplification Master Mix to a tube in an 8-tube strip for each sample.
- **d.** Add 25 μ l of each sample from pre-amplification to a separate tube previously aliquoted with Amplification Master Mix.
- e. Add 5 µl of an individual Dual Index Plate TS Set A to each tube and record the well ID used. Pipette mix and centrifuge briefly.
- **f.** Incubate in a thermal cycler with the following protocol.

Lid Temperature	Reaction Volume	Run Time		
105°C	100 μΙ	Variable		
Step	Temperature	Time hh:mm:ss		
1	98°C	00:03:00		
2	98°C	00:00:15		
3	63°C	00:00:20		
4	72°C	00:00:30		
5	Go to step 2, use the Cq Value +1 as the total # of cycles. See table below for total # of cycle examples.			
6	72°C	00:01:00		
7	4°C	Hold		

Round Cq values up to the nearest whole number and add one cycle (examples below). Samples within ±1 cycle numbers can be combined in a single SI-PCR run at the higher cycle number.



Do not combine samples if the cycle number difference is greater than 1 to avoid library overamplification.

Example Cycle Numbers

Cq Value from qPCR	+1	Total Cycles
7.2	+1	9
8.5	+1	10
13.7	+1	15

Example Batched Cycles

Cq Value from qPCR	+1	Total Cycles	Batched Cycles
7.2	+1	9	10
9.5	+1	11	10
13.7	+1	15	16
14.6	+1	16	10



Any remaining pre-amplification material can be stored at $4^{\circ}C$ for up to 72~hor at -20°C for up to 4 weeks for generating additional libraries.

6.3 Post-Sample Index PCR Cleanup - SPRIselect

If using MicroAmp 8-Tube Strips, use the high position (magnet•High) only throughout the protocol

- a. Vortex to resuspend SPRIselect Reagent. Add 85 µl SPRIselect Reagent (0.85X) to each sample. Pipette mix 15x (pipette set to 175 μ l).
- **b.** Incubate **5 min** at **room temperature**.
- c. Place on magnet•High for 3 min. Verify solution is clear.
- d. Remove supernatant.
- e. Add 200 μl 80% ethanol to pellet. Wait 30 sec.
- **f.** Remove ethanol.
- g. Add 200 µl 80% ethanol to pellet. Wait 30 sec.
- h. Remove ethanol.
- i. Centrifuge briefly. Place on the magnet•Low.
- j. Remove any remaining ethanol. Air dry for 2 min. DO NOT exceed 2 min as this will decrease elution efficiency.
- k. Remove from magnet. Add 27 µl Buffer EB. Pipette mix 15x.
- 1. Incubate 2 min at room temperature.
- m. Place on the magnet•Low for 3 min. Verify solution is clear.
- **n.** Transfer **25** μ **l** sample to a new tube strip on ice.

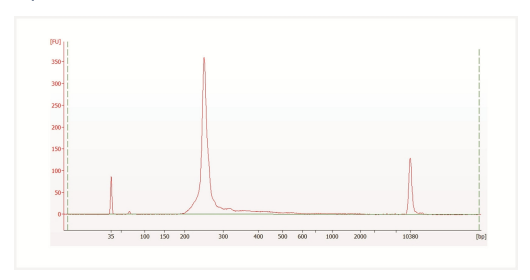


o. Store at **-20°C** for **long-term** storage.

6.4 Post-Library Construction QC

- a. Dilute sample (1:50 dilution, i.e 1 µl sample in 49 µl of EB buffer) until it is at an appropriate concentration for the Bioanalyzer.
- **b.** Run **1 μl** of diluted sample on an Agilent Bioanalyzer High Sensitivity chip. If peak is too small or flat, retry with a lower dilution. See 5. Flat Line in Bioanalyzer Library Trace on page 113 for more information.

Representative Trace



Determine the average fragment size from the Bioanalyzer trace. The expected average fragment size is 250 bp. This will be used as the insert size for library quantification.

Alternate QC Methods:

- Agilent TapeStation
- Perkin Elmer LabChip

See Appendix on page 120 for representative trace

See Post Library Construction Quantification on page 126

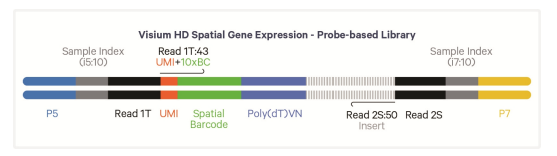


Sequencing

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Sequencing Libraries

Libraries contain standard Illumina paired-end constructs, which begin with P5 and end with P7. Spatial Barcodes are encoded at the start of TruSeq Read 1 (Read 1T), while i7 and i5 sample index sequences are incorporated as the index read. Read 1T, and Small RNA Read 2 (Read 2S) are standard Illumina sequencing primer sites used in paired-end sequencing. Read 1T are used to sequence the 43 bp Spatial Barcode and UMI. Read 2S is used to sequence the Ligated Probe Insert. Sequencing these libraries produces a standard Illumina BCL data output folder.



Sequencing Depth

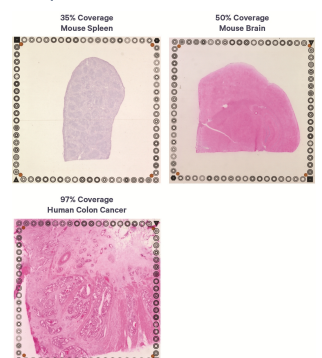
The minimum sequencing depth for Visium HD is 275 million read pairs per fully-covered Capture Area. The recommended minimum sequencing depth was chosen because it achieved >50% sequencing saturation for >50% formalin fixed, paraffin embedded (FFPE) tissues tested at 10x Genomics. To achieve >50% saturation for >50% of fresh frozen (FF) tissues tested by 10x Genomics, 700 million read pairs per fully-covered Capture Area were required. To achieve >50% saturation for >50% of the Fixed Frozen (FxF) tissues tested by 10x Genomics, 500 million read pairs per fully-covered Capture Area were required.

10x Genomics recommends calculating the minimum sequencing depth for libraries by multiplying the percentage of the Capture Area covered by tissue by 275 million read pairs. For example, a Capture Area that is 50% covered tissue would require at least 137 million read pairs (0.50 X 275,000,000).

Calculating sequencing depth requires estimating the approximate Capture Area (%) covered by tissue. This may be performed visually or by using the Visium Manual Alignment Wizard in Loupe Browser for a more accurate measurement. See examples below for estimating coverage area visually. If using Loupe Browser, the number of bins associated with tissue can be calculated during the "Identify Tissue" step. For more information, consult the 10x Genomics Support website.

Example: Sequencing Depth for a Sample

- Estimate the approximate Capture Area (%) covered by the tissue section.
- Calculate total sequencing depth= (Coverage Area x 275,000,000 read pairs
- Example calculation for 50% coverage:
 0.50 x 275,000,000 read pairs = 137,500,000 total read pairs for that sample



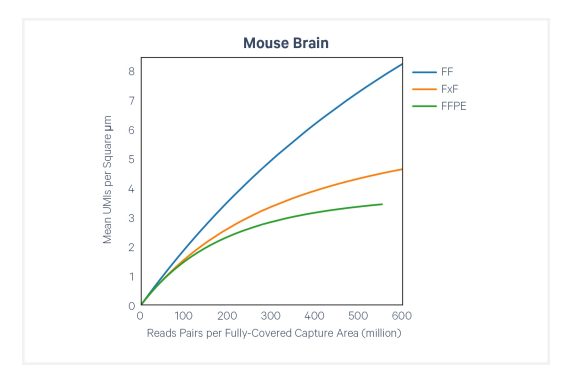
Sequencing saturation, at a particular sequencing depth, is driven by the complexity of the library and the percent usable read pairs. To achieve saturation with high complexity libraries (unique UMIs), more sequencing is required to read all unique UMIs present in the library. Saturation is achieved more efficiently (with fewer read pairs) for libraries with a high percentage of usable read pairs. The main driver of saturation for Visium HD libraries is library complexity.

Visium HD library complexity, or the UMIs recovered from a tissue sample, is dependent on fixation method, tissue composition, tissue type, and RNA quality. Visium HD FF libraries will typically be more complex compared to equivalent FFPE tissue libraries. At a particular sequencing depth, a FF tissue run with Visium HD will typically result in a higher number of UMIs per square μm and a lower sequencing saturation compared to an equivalent FFPE sample of the same tissue type.

The figures below show the relationship between sequencing depth, sequencing saturation, and mean UMIs per square μm across different tissue preservation methods. The first graph shows the relationship between mean UMIs per square μm and sequencing depth. At the same sequencing depth, FF with good RNA quality will typically have higher UMIs per square μm compared to FFPE. Around the recommended sequencing depth, the slope of the line representing the FFPE samples becomes smaller, indicating faster saturation and smaller increases in UMIs per square μm with additional sequencing.

The second graph shows the relationship between sequencing depth and sequencing saturation for the same samples shown in the first graph. At the same sequencing depth, FF with good RNA quality will typically have lower saturation compared to FFPE due to higher complexity. In these cases, additional sequencing is often desirable.

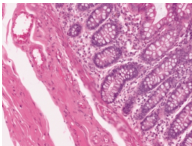


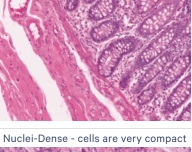


Tissue composition can also influence the ideal sequencing depth for a sample. The figures below demonstrate the relationship between sequencing depth and mean UMIs per square µm for a variety of sample types to help guide selection of sequencing depth. All H&E images below were taken at 20X magnification.

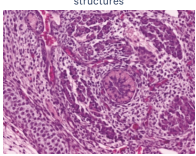
Tissue Compositions Tested

Stroma-Rich - extracellular matrix, connective tissue, or adipose tissue

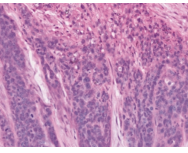


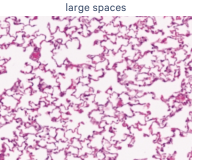


Heterogenous - variety of cell types and structures

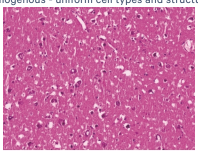


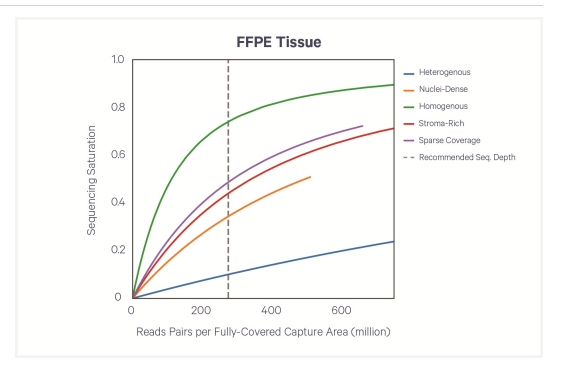
Sparse Coverage - cells are separated by



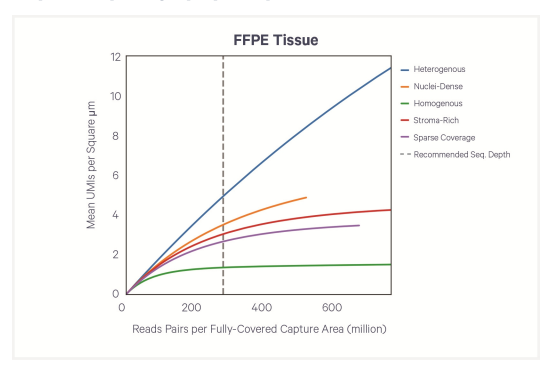


Homogenous - uniform cell types and structures

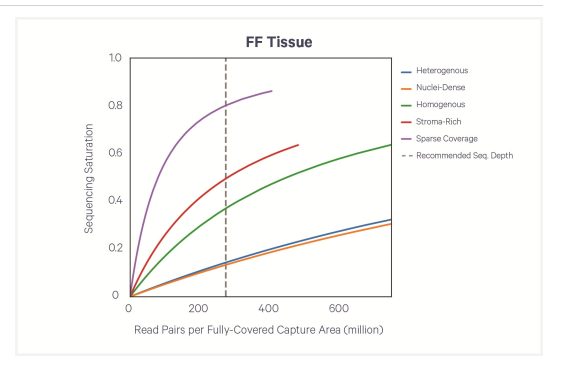




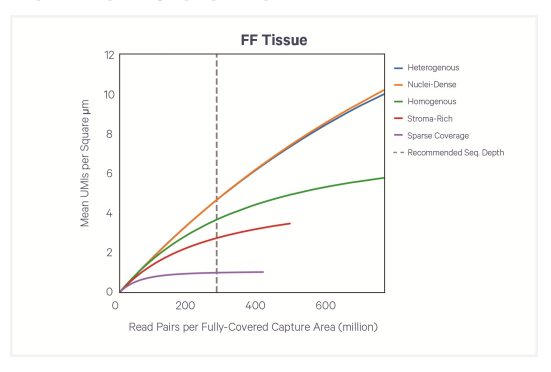
The above graph shows the relationship between sequencing saturation per sample and sequencing depth per sample.



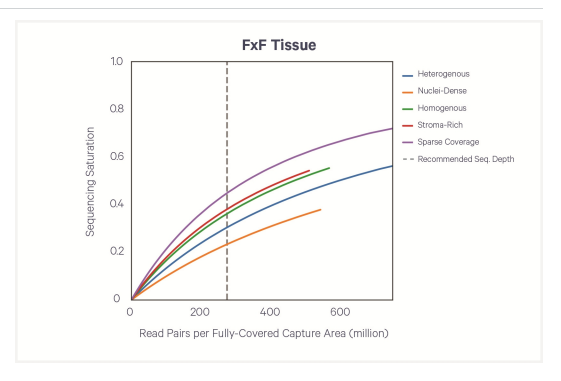
The above graph shows the relationship between mean UMIs per square μm of the Capture Area covered by tissue and sequencing depth per sample.



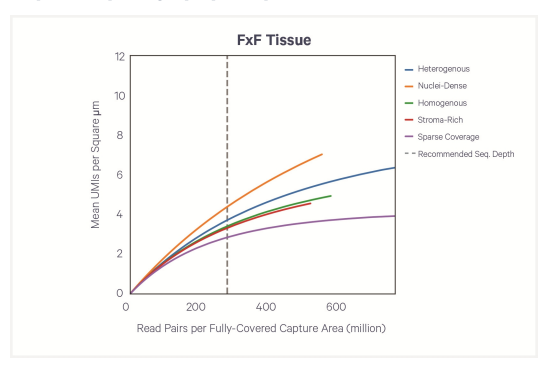
The above graph shows the relationship between sequencing saturation per sample and sequencing depth per sample.



The above graph shows the relationship between mean UMIs per square μm of the Capture Area covered by tissue and sequencing depth per sample.



The above graph shows the relationship between sequencing saturation per sample and sequencing depth per sample.



The above graph shows the relationship between mean UMIs per square μm of the Capture Area covered by tissue and sequencing depth per sample.

Sequencing Type & Run Parameters

Use the sequencing run type and parameters indicated.

Visium HD Spatial Gene Expression - Probe-based Library

Paired-end, dual indexed sequencing

Read 1T: 43 cycles i7 Index: 10 cycles i5 Index: 10 cycles Read 2S: 50 cycles

Illumina Sequencer Compatibility

The compatibility of the listed sequencers has been verified by 10x Genomics. Some variation in assay performance is expected based on sequencer choice.

- NextSeq 2000
- NovaSeq 6000
- NovaSeq X Plus

Sample Indices

Each well of the Dual Index Kit TS Set A (PN-1000251) contains a mix of one unique i7 and one unique i5 sample index. If multiple samples are pooled in a sequence lane, the sample index name (i.e. the Dual Index Plate TS Set A plate well ID, SI-TS-) is needed in the sample sheet used for generating FASTQs with "spaceranger mkfastq". Samples utilizing the same sample index should not be pooled together or run on the same flow cell lane, as this would not enable correct sample demultiplexing.

Color balance in the index read is important for optimal quality and demultiplexing results. For example, when pooling samples on the NovaSeq X Series instruments, Illumina recommends avoiding index combinations which only have signals from A bases, G bases, or G+A bases in any given cycle. Refer to Illumina's guidance on index color balancing.

Library Pooling

Visium HD libraries derived from samples prepared using different methods (such as FFPE, FF, FxF) may be pooled for sequencing, taking into account differences in tissue-covered spot on a Capture Area and per-spot read depth requirements.

Pooling Visium HD libraries with other 10x Genomics libraries is not recommended due to their unique sequencing configuration: R1: 43, i7:10,

i5:10, R2: 50. Additionally, on NovaSeq X Plus, Visium libraries cluster optimally at a higher loading concentration (300-400 pM) than most Single Cell Gene Expression libraries (150-200 pM). Limited testing of Visium HD and Flex libraries pooled at a 1:1 molar ratio at range of concentrations on NovaSeq X Plus did not yield any equal sequencing representation of Visium HD and Flex libraries, likely due to the differences in optimal loading concentrations for each library type.

Library Loading

Library quantification should be done with the KAPA DNA Quantification Kit using the average insert size determined by Agilent Bioanalyzer, Perkin Elmer LabChip, or Agilent TapeStation QC. Alternate methods to KAPA qPCR for final library concentration may result in underquantification and consequently overloading.

Once quantified and normalized, libraries should be denatured and diluted as recommended for Illumina sequencing platforms. Consult Illumina documentation for denaturing and diluting libraries.

Instrument	Loading Concentration (pM)	PhiX (%)
NextSeq 2000	650	1
NovaSeq 6000 Standard	100-150	1
NovaSeq 6000 XP Workflow	150-200	1
NovaSeq X Plus	300-400	1

Sequencing Metrics

FFPE samples were processed according to documentation listed in the Workflow Overview on page 13. Libraries were pooled and sequenced at the indicated run parameters to generate the % Base and % ≥Q30 plots shown in the following figures. These same libraries were used to illustrate sequence compatibility. All libraries were quantified with the KAPA DNA Quantification Kit and sequenced with 1% PhiX. Though only FFPE libraries are shown in this section, libraries from fresh frozen (FF) and fixed frozen (FxF) tissue sections are expected to perform similarly. All libraries followed the following sequencing configuration and run parameters:

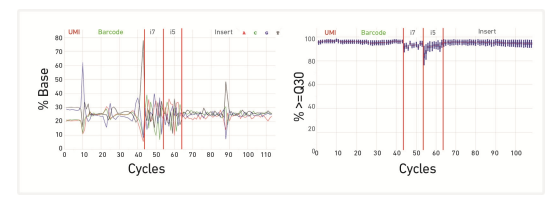
- Targeted sequencing depth: Minimum of 275 million read pairs multiplied by fraction Visium HD slide tissue coverage.
- Paired-end, dual indexing: Read 1: 43 cycles; i7 Index: 10 cycles; i5 Index: 10 cycles; Read 2: 50 cycles

% Base by cycle and % ≥Q30 Quality Score distribution showed highly consistent profiles for all sequencing platforms tested. The data serve as guidelines for assessing the quality of library sequencing. Additional factors that may contribute to overall success of a sequencing run and impact downstream application performance metrics include:

- Starting with a high-quality tissue block.
- Generating sequencing data from sections that remain adhered to the slide.
- Final libraries with fragment lengths within the expected size range for each library type.
- Reliable and accurate library quantification using the KAPA qPCR DNA
 Quantification Kit for Illumina Platforms (Roche, KK4824) and the average
 insert size determined by Agilent Bioanalyzer, Perkin Elmer LabChip, or
 Agilent TapeStation QC. Alternative methods to KAPA qPCR, such as Qubit,
 for final library quantification may result in underquantification and result
 in overloading flow cells.
- Sequencing platform loading concentrations follow recommendations
 described in Library Pooling on page 103, which are based on KAPA qPCR
 quantification. Overloading/over-clustering may result in poor run
 performance, decrease sequencing quality, and lower total data output as
 compared to optimally loaded runs. The loading recommendations for an
 individual sequencer are listed as general guidance and additional
 optimization may be required.

Probe-based Libraries

Representative % Base and % ≥Q30 plots are from a pool of sixteen Probe-based libraries sequenced on a NovaSeq 6000 SP flow cell.



Libraries were sequenced on the NextSeq 2000, NovaSeq 6000 and NovaSeq X Plus. "Data by Cycle" plots from the Illumina SAV software displaying the percentage of base calls and Q30 quality scores are shown. Mapped Reads (%) refers to Reads Mapped Confidently to Probe Set for Probe-based (GEX).

Instr.	Load Conc. (pM)	% Occupancy	%PF	R1	R2	R1	i7	i5	R2	GEX
					er Lane 6b)		% ≥(230		Mapped Reads (%)
NextSeq 2000	650	81.2	73.1	51.4	59.9	95.6	96.6	95.1	94.4	97.2
NovaSeq 6000 SP	300	93.2	82.8	135	160	95	92	89	93	92.1
NovaSeq X Plus	400	84.7	71.79	41.4	48.3	95.9	94.6	94.7	95.3	97.7



Troubleshooting



Before CytAssist Instrument Run

1. 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 targeted reads usable (see table below). A similar effect is observed when washing for less than the recommended 5 min, 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.

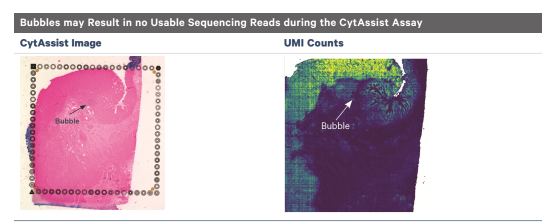
Wash	Number of Washes	Fraction Reads Usable (Mean)		
Post Hybridization Wash	1	0.29		
	2	0.41		
	3	0.79		
Post Ligation Washes	1	0.69		
	2	0.75		

2. Inadequate Visium HD Slide Preparation

- Leaving Visium HD Slide in 0.1X SSC inside slide mailer longer than 4 h after Visium HD Slide washing may result in spacer detachment.
- Failure to remove excess liquid from the back of the Visium HD Slide before loading the slide onto the Visium CytAssist may result in distorted fiducial imaging, which can impact fiducial registration.
- Insufficient drying of Visium HD Slide prior to instrument run may result in bubble entrapment and uneven flow of reagents.

During CytAssist Instrument Run

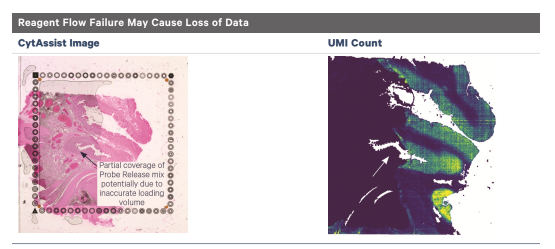
1. Bubbles Trapped During Visium CytAssist Run



Bubbles during a sample run are rare, but may result in a lack of useable reads in the tissue area where the bubbled occurred. The most common cause of bubbles is incomplete drying of the tissue slide and/or the Visium HD slide. Bubbles can also result from inaccurate dispensing volume of reagents onto the slide. Ensure that the entire spacer chamber is inspected for bubbles, not just the Capture Area. Ensure that the entire spacer chamber is also dry during the drying step, not just the Capture Area.

Avoid generating bubbles during reagent dispensing by pipetting slowly and avoiding expelling air from the pipette tip. Additionally, ensure that the frosted area of the tissue slides are outside of the alignment guide lines on the Tissue Slide Stage and that tissue slides are compatible with the CytAssist instrument. Consult the Visium HD Spatial Gene Expression Protocol Planner (CG000698) for information on tested slides.

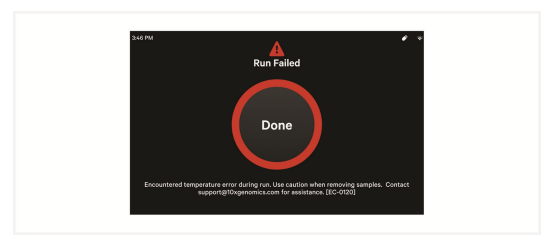
2. Reagent Flow Failure



Inappropriate flow of reagents during a Visium CytAssist experiment run may result in transcript mislocation. This may be caused by improper loading of Probe Release Mix onto the Visium HD Slide or debris on the Visium HD Slide. If the UMI map appears abnormal (complete loss of UMIs from a significant portion of the tissue), contact support@10xgenomics.com.

Ensure that frosted areas of the tissue slide are not within the dashed zone on the Visium Tissue Slide Stage. Keep Visium slides and tissue slides free from physical damage, dust, and debris. After Eosin wash, remove as much PBS as possible by flicking the tissue slide and wiping any remaining droplets around tissue with a lint-free laboratory wipe. Both tissues slides and Visium HD slides should be completely dry. See Lint-free Laboratory Wipes on page 23 for more information.

3. Visium CytAssist Overheating

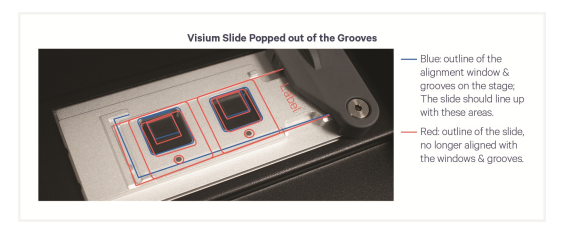


If the Visium CytAssist instrument overheats during an experiment run, the run will fail. A warning message will appear instructing users to carefully remove slides and contact support@10xgenomics.com.

After CytAssist Instrument Run

1. Slide Popped Off of Visium Slide Stage

Upon opening the lid after a successful run, the Visium slide may pop out of the grooves on the Visium Slide Stage. This does not impact assay performance or slide quality.



2. Visium HD Slide Removal Delayed

A delay in removing and processing the Visium HD Slide after run completion may impact data quality. Immediately proceed with slide washes while adjacent to the instrument.

3. High Cq Value

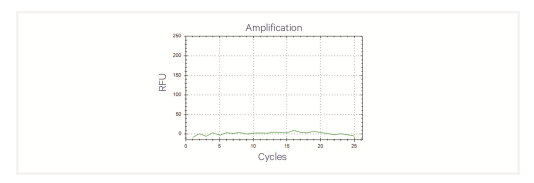
Cq values higher than 15 may indicate a workflow error. Tissues that are small, not complex (e.g. high amounts of connective tissue), or are poor quality may result in higher Cq values (15-20) than expected. 10x Genomics does not recommend proceeding with the assay if Cq values are higher than 20. Contact support@10xgenomics.com for further assistance.

4. No qPCR Amplification

No amplification during Cycle Number Determination may be due to the following:

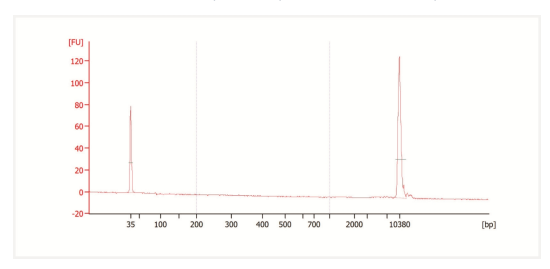
- Incorrect qPCR machine programming (i.e. detecting a fluorophore other than SYBR Green)
- Issues with pipetting small volumes

- Failure to neutralize KOH
- Failure to add primers to pre-amp or qPCR mix
- Unusually low recovery from tissue due to high percentage of connective tissue in Capture Area
- Incorrect preparation of Probe Release Mix
- Mistake in Probe Release Mix addition timing
- · Leakage from the cassette during workflow



5. Flat Line in Bioanalyzer Library Trace

A normal qPCR output but no peak visible in the Bioanalyzer trace may be due to a mistake in the SI-PCR step, errors during SPRIselect cleanup, an overdilution of the final library, Bioanalyzer issue, or thermal cycler failure.

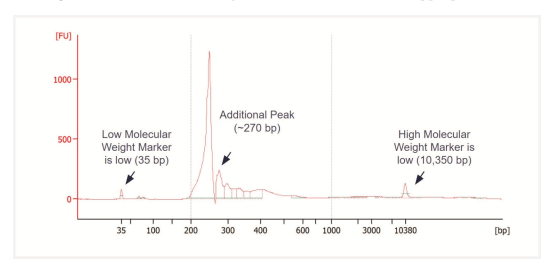


6. Traces with Additional Peaks

Overloaded Trace

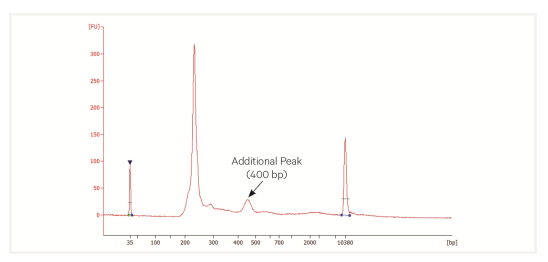
The image below is an example of an overloaded trace. Note the double peak at around 270 bp. The low and high molecular weight markers are in low

abundance compared to the sample. Ensure the library was diluted prior to loading per protocol recommendations. If the library was diluted, ensure the library concentration is within the specification range of the automated electrophoresis kit being used. The library can be quantified using an Invitrogen Qubit dsDNA HS Assay Kit and diluted further if appropriate.



Overamplified Trace

The image below is an example of an overamplified trace. Note peak at 400 bp.



Ensure interpretation of qPCR plot is correct. If necessary, batch samples into separate SI-PCR runs if Cq is > 1 cycle. If batching samples, the cycle number determined should be ±1. If needed, select the value in between.

For example, a sample with Cq value of 7.5 should go through 9 cycles during SI-PCR (round up to 8 and add 1) and a sample with a Cq value of 9.2 should go through 11 cycles during SI-PCR (round up to 10 and add 1). If running together, samples should undergo 10 cycles during SI-PCR.

Issues Impacting Tissue Analysis

1. Tissue Detachment and Folding

Tissue detachment may result in a lack of Fraction Reads Usable in the region where detachment occurred. If the tissue has folded on itself, this may also cause elevated UMI counts in the overlapping areas or a loss of spatiality due to ineffective permeabilization. Tissue folding may also cause lower UMI capture. Inspect images carefully to identify these areas.

Ensure that slides tested by 10x Genomics were used for tissue placement. For a list of tested slides, refer to the Visium HD Spatial Gene Expression Application Protocol Planner (CG000698). For more information, consult the sample preparation documentation described in Workflow Overview on page 13.

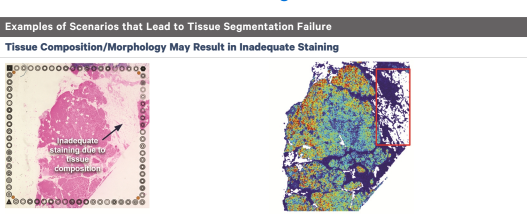


2. Area of Interest Not Within Allowable Area

Area of Interest Outside of Allowable Area is Not Analyzed **Tissue Larger than Capture Area Tissue Not Properly Aligned** AOI Outside Allowable Area 0000000000000000000000

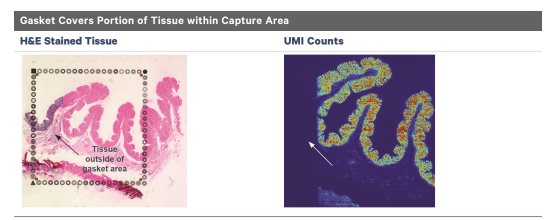
Areas of interest that are not placed within the allowable area on compatible glass slides will not be analyzed. This may occur if the tissue is larger than the Capture Area or if the tissue slide is not properly aligned when loading onto the Tissue Slide Stage. Tissue outside of the allowable area is fine if the tissue outside the allowable area is not an area of interest. Consult the Visium HD Protocol Planner (CG000698) for information on tested glass slides.

3. Tissue Segmentation Failure due to High Connective Tissue



Space Ranger may fail to detect tissue for a variety of reasons, which may necessitate manual fiducial alignment and tissue detection via Loupe Browser. If the contrast between background and tissue is poor (due to inadequate staining, technical error, or tissue composition) tissue segmentation failure may occur. Consider processing H&E images through Space Ranger's segmentation pipeline (spaceranger segment) prior to proceeding with the assay to gauge expected performance. For troubleshooting tips related to image quality, refer to the Visium HD Spatial Applications Imaging Guidelines Technical Note (Document CG000688).

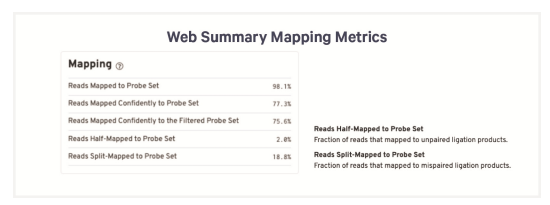
4. Tissue within Capture Area Not Analyzed



If the gasket covers a portion of the tissue within the Capture Area, that area will not be exposed to the appropriate reagents; thus, probes will not be captured. In the example above, a lack of destaining on the left side of the tissue implies that this area was obscured by the gasket. Though the area is within the Capture Area, probe capture will not occur due to this obstruction. Gasket obstruction may be caused by improper gasket assembly, resulting in the gasket being placed directly onto areas of interest. In Loupe Browser, manual alignment must be performed to exclude the tissue area obscured by the gasket. Ensure that the area of interest is centered within the gasket.

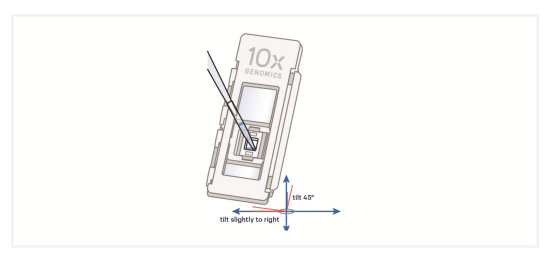
Additionally, tissue placed within the Capture Area may be excluded from the assay workflow due to improper tissue slide alignment when loading the tissue slide into the Visium CytAssist instrument.

5. High Split-Mapped or Half-Mapped Reads



High split-mapped or half-mapped reads can result from insufficient and/or inefficient washing of the tissue (especially during the post-hybridization and post-ligation wash steps), poor tissue quality, or poor sample preparation. Follow all best practices, including:

- Following all sample preparation guidelines outlined in the appropriate tissue handbook (see Workflow Overview on page 13 for documentation).
- Completely removing residual liquid prior to proceeding to subsequent steps (e.g. using a P20 pipette to remove residual volume, tilting the cassette to collect the liquid at a corner, see illustration below)
- Performing the wash steps close to the thermal cycler
- Ensuring that the tissue section(s) do not dry out during the washes.
- Checking that the cassette is fully seated on the Low Profile Thermocycler Adapter between performing each wash
- Cassette is fully sealed when necessary to prevent evaporation
- Ensuring that the thermal cycler lid is set to the appropriate temperature to prevent condensation during the incubation steps.

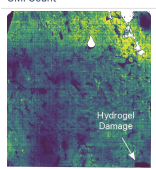


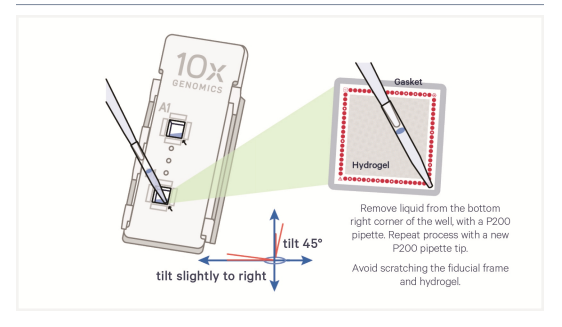
6. Data Loss due to Hydrogel Damage

Damage to the hydrogel may result in data loss. Avoid damaging the hydrogel by adhering to best practices when removing reagent from the well as described in Reagent Addition to & Removal from Wells on page 26. In the example below, a scratch on the hydrogel noted by the arrow has resulted in a loss of UMI count.

Hydrogel Damage May Result in Data Loss

UMI Count





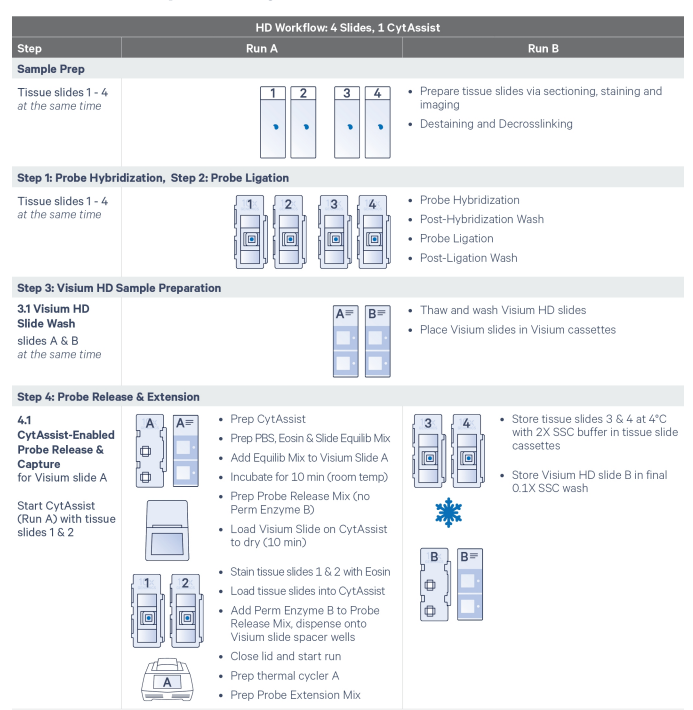


Appendix

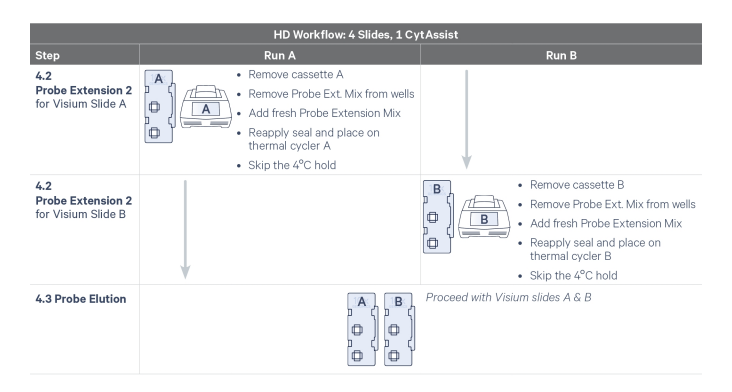
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HD Workflow Overview for Processing 4 Tissue Slides with 1 Visium CytAssist Instrument

To execute this workflow, two thermal cyclers are required. This table is meant as a reference only. Consult the detailed steps in this user guide for execution.



HD Workflow: 4 Slides, 1 CytAssist		
Step	Run A	Run B
Equilibration for tissue slides 3 & 4 during Run A	Once CytAssist has completed the run and is cooling down, begin equilibration for slides 3 & 4.	 Remove slides 3 & 4 from storage Equilibrate for 5 min (room temp.)
CytAssist (Run A) complete	 Open CytAssist Wash Visium slide A Place in Visium Cassette 	
4.2 Probe Extension 1 for Visium Slide A	 Add Probe Extension Mix Apply seal and place on thermal cycler A 	
Ensure 20 min cooldown period after completion of Run A Start CytAssist (Run B) with tissue slides 3 & 4		Prep CytAssist Prep PBS, Eosin & Slide Equilib Mix Add Equilib Mix to Visium Slide B Incubate for 10 min (room temp) Prep Probe Release Mix (no Perm Enzyme B) Load Visium Slide on CytAssist to dry (10 min) Stain tissue slides 3 & 4 with Eosin Load tissue Slides into CytAssist Add Perm Enzyme B to Probe Release Mix, dispense onto Visium slide spacer wells Close lid and start run Prep thermal cycler B Prep Probe Extension Mix
CytAssist (Run B) complete		 Open CytAssist Wash Visium slide B Place in Visium Cassette
4.2 Probe Extension 1 for Visium Slide B		 Add Probe Extension Mix Apply seal and place on thermal cycler B



HD Workflow Overview for Processing 4 Tissue Slides with 2 Visium CytAssist Instruments

To execute this workflow, two thermal cyclers are required. This table is meant as a reference only. Consult the detailed steps in this user guide for execution.

HD Workflow: 4 Slides, 2 CytAssists			
Step	Run A, CytAssist A	Run B, CytAssist B	
Sample Prep			
Tissue slides 1 - 4 at the same time		 Prepare tissue slides via sectioning, staining and imaging Destaining and Decrosslinking 	
Step 1: Probe Hybridization, Step 2: Probe Ligation			
Tissue slides 1 - 4 at the same time		Probe HybridizationPost-Hybridization WashProbe LigationPost-Ligation Wash	
Step 3: Visium HD Sample Preparation			
3.1 Visium HD Slide Wash slides A & B at the same time	A=	Thaw and wash Visium HD slidesPlace Visium slides in Visium cassettes	

HD Workflow: 4 Slides, 2 Cyt Assists			
Step	Run A, CytAssist A	Run B, CytAssist B	
Step 4: Probe Release & Extension			
4.1 CytAssist- Enabled Probe Release & Capture for Runs A & B	A	Prep CytAssist A Prep PBS, Eosin & Slide Equilibration Mix for Runs A & B	
4.1 CytAssist- Enabled Probe Release & Capture for Visium slide A Start CytAssist A with tissue slides 1 & 2	 Add Equilib Mix to Visium Slide A Incubate for 10 min (room temp) Prep Probe Release Mix (no Perm Enzyme B) Load Visium Slide on CytAssist to dry (10 min) Stain Tissue slides 1 & 2 with Eosin 	• Store tissue slides 3 & 4 at 4°C with 2X SSC buffer in tissue slide cassettes	
	 Load Tissue Slides into CytAssist Add Perm Enzyme B to Probe Release Mix, dispense onto Visium slide spacer wells Close lid and start run Prep thermal cycler A Prep Probe Extension Mix 	• Store Visium HD slide B in final 0.1X SSC wash with Visium Slide Seal	
Equilibration for tissue slides 3 & 4		 Remove slides 3 & 4 from storage Equilibrate for 5 min (room temp.) 	
Start CytAssist B with tissue slides 3 & 4		Prep CytAssist B Add Equilib Mix to Visium Slide B Incubate for 10 min (room temp) Prep Probe Release Mix (no Perm Enzyme B) Load Visium Slide on CytAssist to dry (10 min) Stain tissue slides 3 & 4 with Eosin Load tissue Slides into CytAssist Add Perm Enzyme B to Probe Release Mix, dispense onto Visium slide spacer wells Close lid and start run Prep thermal cycler B Prep Probe Extension Mix	

HD Workflow: 4 Slides, 2 CytAssists			
Step	Run A, CytAssist A	Run B, CytAssist B	
CytAssist (Run A) complete	• Open CytAssist • Wash Visium slide A • Place in Visium Cassette		
4.2 Probe Extension 1 for Visium Slide A	 Add Probe Extension Mix Apply seal and place on thermal cycler A 		
CytAssist (Run B) complete		 Open CytAssist B Wash Visium slide B Place in Visium Cassette 	
4.2 Probe Extension 1 for Visium Slide B		 Add Probe Extension Mix Apply seal and place on thermal cycler B 	
4.2 Probe Extension 2 for Visium Slide A	 Remove cassette A Remove Probe Extension Mix from wells Add fresh Probe Extension Mix Reapply seal and place on thermal cycler A Skip the 4°C hold 		
4.2 Probe Extension 2 for Visium Slide B		Remove cassette B Remove Probe Extension Mix from wells Add fresh Probe Extension Mix Reapply seal and place on thermal cycler B Skip the 4°C hold	
4.3 Probe Elution		Proceed with Visium slides A & B	

Post Library Construction Quantification

- a. Thaw KAPA Library Quantification Kit for Illumina Platforms.
- **b.** Dilute $2 \mu l$ sample with deionized water to appropriate dilutions that fall within the linear detection range of the KAPA Library Quantification Kit for Illumina Platforms. (For more accurate quantification, make the dilution(s) in duplicate).
- **c.** Make enough Quantification Master Mix for the DNA dilutions per sample and the DNA Standards (plus 10% excess) using the guidance for 1 reaction volume below.

Quantification Master Mix	1X (µl)
SYBR Fast Master Mix + Primer	12
Water	4
Total	16

- **d.** Dispense **16** μ **l** Quantification Master Mix for sample dilutions and DNA Standards into a 96 well PCR plate.
- e. Add $4 \mu l$ sample dilutions and $4 \mu l$ DNA Standards to appropriate wells. Centrifuge briefly.
- **f.** Prepare a qPCR system with the following protocol. Insert the plate and start the program.

Lid Temperature	Reaction Volume	Run Time
-	20 μΙ	35 min
Step	Temperature	Time
1	95°C	00:03:00
2	95°C	00:00:05
3	67°C Read signal	00:00:30
4	Go to Step 2, 29X (Total 30 cycles)	

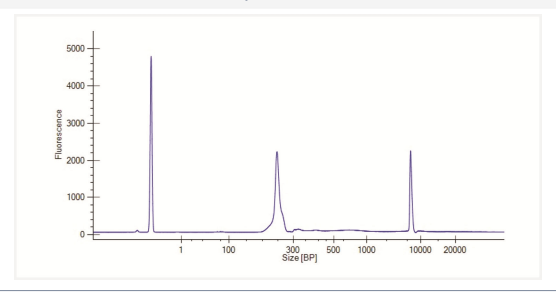
g. Follow the manufacturer's recommendations for qPCR-based quantification. For library quantification for sequencer clustering, determine the concentration based on insert size derived from the Bioanalyzer/TapeStation trace.

Perkin Elmer LabChip Traces

Protocol Step 6.3 - GEX Post Library Construction QC

Representative Trace

Run manufacturer's recommended volume of diluted sample (1:5 dilution)



DNA High Sensitivity Reagent Kit was used. Protocol steps correspond to the steps in this User Guide.

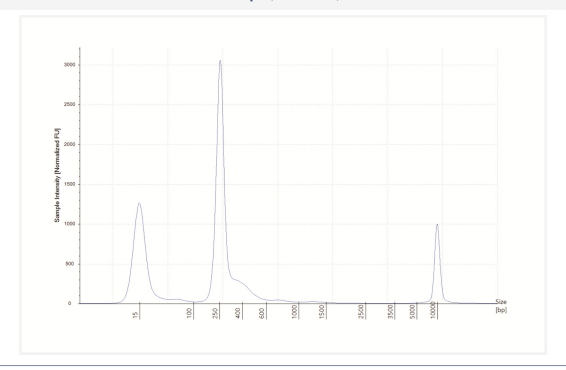
Agilent TapeStation Traces

Agilent TapeStation High Sensitivity D5000 ScreenTape was used. Protocol steps correspond to the steps in this User Guide.

Protocol Step 6.4 - GEX Post Library Construction QC

Representative Trace

Run manufacturer's recommended volume of diluted sample (1:20 dilution)



Oligonucleotide Sequences

ongonacieotide Sequences



5'-AATGATACGGCGACCACCGAGATCTACAC-N10-ACACTCTTTCCCTACACGACGCTCTTCCGATCT-UMI-SpatialBarcode-GVV-T30-VN-Ligated_Probe_Insert-TGGAATTCTCGGGTGCCAAGGAACTCCAGTCAC-N10-ATCTCGTATGCCGTCTTCTGCTTG-3'

3'-TTACTATGCCGCTGGTGGCTCTAGATGTG-N10-TGTGAGAAAGGGATGTGCTGCGAGAAGGCTAGA-UMI-SpatialBarcode-GVV-T30-BN-Ligated_Probe_Insert-ACCTTAAGAGCCCACGGTTCCTTGAGGTCAGTG-N10-TAGAGCATACGGCAGAAGACGAAC-5'