10x Genomics Scientific Symposium

Copenhagen, Borsen Stock Exchange | 2-3 June

https://10xgenomicsugm-copenhagen.eventbrite.com

- Day 1:
 - Presentations by industry leaders
 - Sample preparation workshop
 - Enjoy an evening social event
 - Book 1:1 meetings with our 10x-perts
 - · Visit our troubleshooting stations for expert advice
 - Four juniors researchers (PhD students/early PostDocs) will win a 500 euro travel award to cover their attendance!
 - Enter our Scientific Challenge to win free kits for your projects!
- Day 2 workshops:
 - Getting Started with Chromium Single Cell Gene Expression Data Analysis (by abstract submission)
 - Getting Started with Visium Spatial Gene Expression Data Analysis (by abstract submission)
 - Bioinformatics for Single Cell and Spatial Genomics (bioinformaticians)







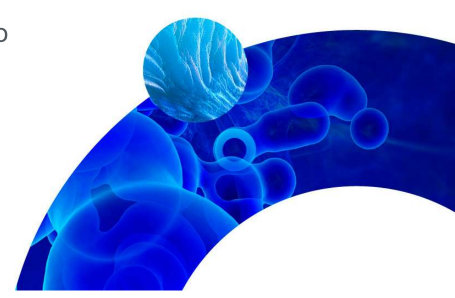


Visium Spatial transcriptomics

From single cell transcriptomics to multi-omics to spatial transcriptomics

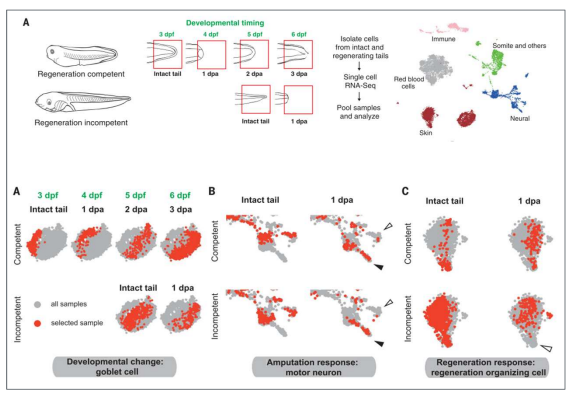
Christophe Fleury, PhD Science & Technology Advisor

User day Toulouse – 10/03/2020



Identification of a regeneration- organizing cell in the Xenopus tail

Aztekin et al., Science 2019



Single cell sequencing *X.laevis* tail (incompetent vs competent):

- Determine existence of amputation response in both lineage
- Regeneration response different: Identification of a new cell type

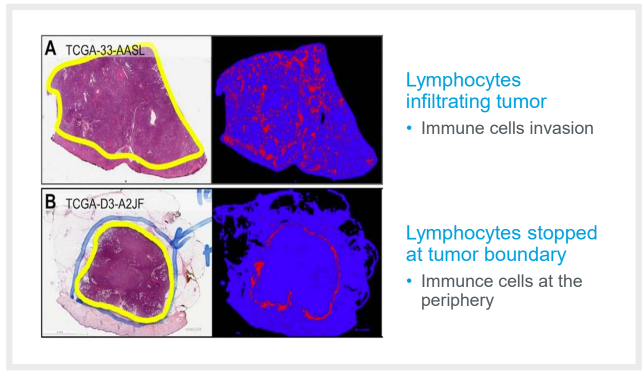
Regeneration organizing cells (ROC)

Where are these cells? Do they relocalise after amputation of the limb?



Why spatial analysis?

Gain deep biological insights with 10x Genomics Visium Spatial Gene Expression Platform





From Bulk to single cell to spatial transcriptomics





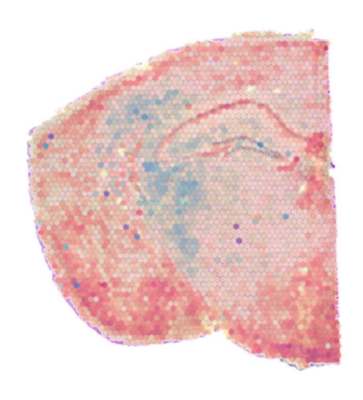
Visium Spatial Gene Expression

Applications



Visium Spatial Transcriptomics

Spatial transcriptomics: Ushering into the era of digital histology

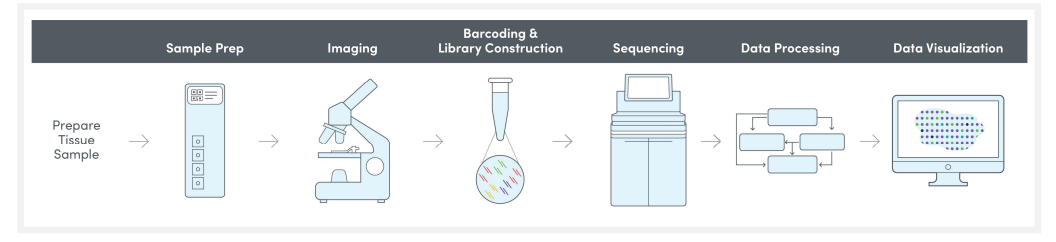


- Capture gene expression information in the context of a tissue section
- Unbiased capture of expressed transcripts (1000s->7000 genes/area)
- Superposition of H&E image and gene expression data
- Simple manipulation of datasets to analyse specific gene expression



The Spatial Gene Expression Solution

Streamlined Workflow from Sample Prep to Data Analysis



1-1.5 day(s)



Visium Spatial Gene Expression

Complete in 1 Working Day with Standard Tools for Tissue Studies

Staining & Imaging Permeabilization cDNA Synthesis CDNA Amplification Construction

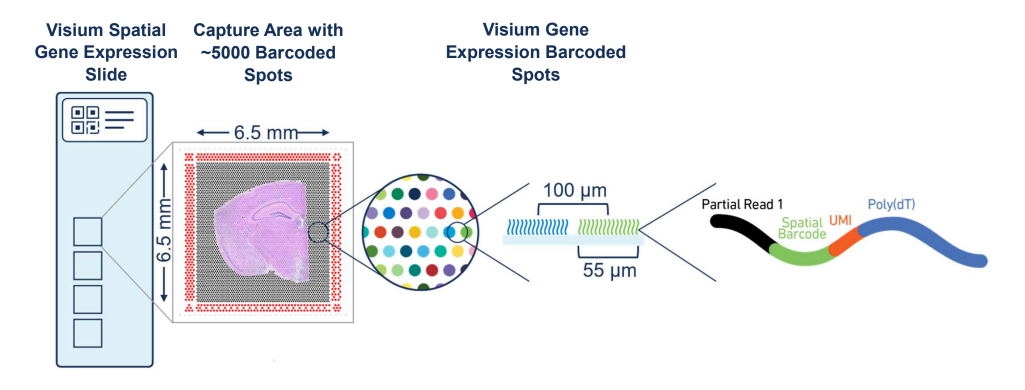
- Section tissue
- Place on slide
- Fix Tissue
- H&E Stain
- Brightfield Image

- Barcoded Spot Capture
- Generate cDNA
- Denature cDNA



Unbiased Gene Expression at High Spatial Resolution

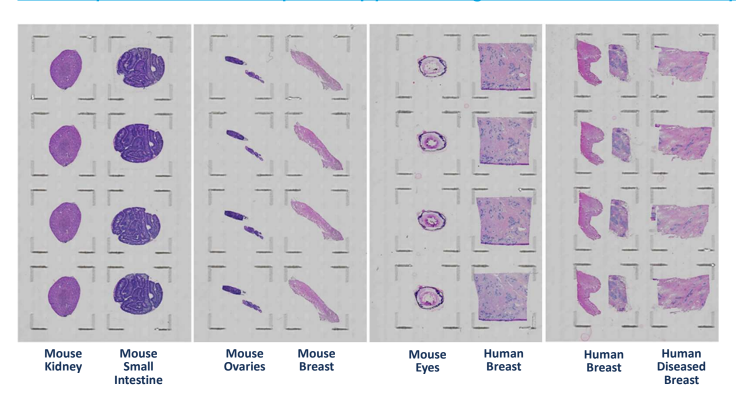
Utilizing Poly-A Capture and Unique Spatial Barcodes





Tissues Tested

Most up to date list: https://support.10xgenomics.com/docs/Spatial_GEX_tissues



HUMAN

Heart, Kidney (Normal & Nephritis),
Ovary, Breast (Normal, Invasive
Ductal Carcinoma, Invasive
Lobular Carcinoma), Lymph Node
(Normal
& Diseased)

MOUSE

Heart, Small & Large Intestine, Stomach, Liver, Kidney, Quadricep, Brain, Lung, Testes, Thyroid, Thymus, Eyes, Tongue, Spleen

RAT

Brain, Kidney, Heart

^{*}Solid tumors for a number of these tissues have also been tested. List will change over time as we are continually testing new tissues. This is not an all inclusive list, if your tissue of interest is not on here it is likely compatible, but we have not tested.

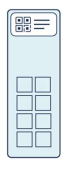




The Visium Spatial Gene Expression Solution

Tissue Optimization and Library Preparation Kits

- Visium Spatial Tissue Optimization Slide and Reagent Kit
 - 4 sample TO kit (4 slides)
 - Four new tissues can be optimized per TO kit (one new tissue per slide)
 - No spatial barcodes
- Visium Spatial Gene Expression Slide and Reagent Kit
 - -4 reactions LP kit (1 slide)
 - -16 reactions LP kit (4 slides)
 - Each slide has 4 capture areas containing spatial barcodes









CF2 Christophe Fleury, 09/03/2020

Visium sensitivity and reproducibility

	Reads per spots	Sequencing Saturation	Number of genes detected
Mouse brain	50 000	41%	4500
Mouse kidney	50 000	48%	3600
Human Lymph Node	50 000	60%	4100
Human Breast Cancer	50 000	63%	4000

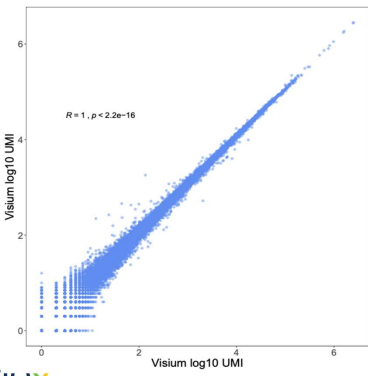


https://support.10xgenomics.com/spatial-gene-expression/datasets

Visium Reproducibility

Pearson Correlation Plot - Mouse Brain Data

4 consecutive slices per week for 6 weeks

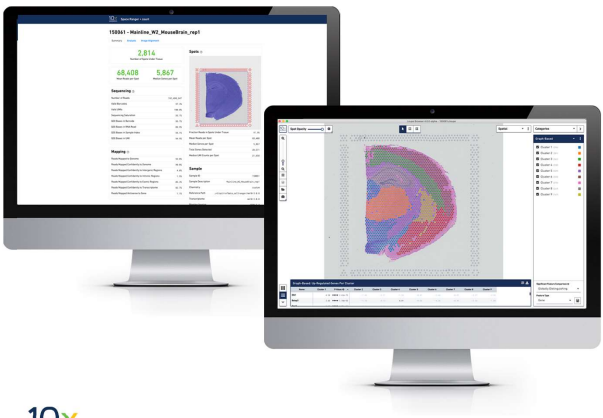


Visium Offers an Extremely Robust Workflow

- Repeatability amongst consecutive sections (replicates)
- Reproducible data at different times and days
- UMI and gene counts remain consistent based on Pearson correlation

Automated Analysis

Space Ranger and Loupe Browser



Freely available on support site

Space Ranger

- -Automated data and image processing
- Clustering analysis based on Gene Expression
- Actionable QC Statistics

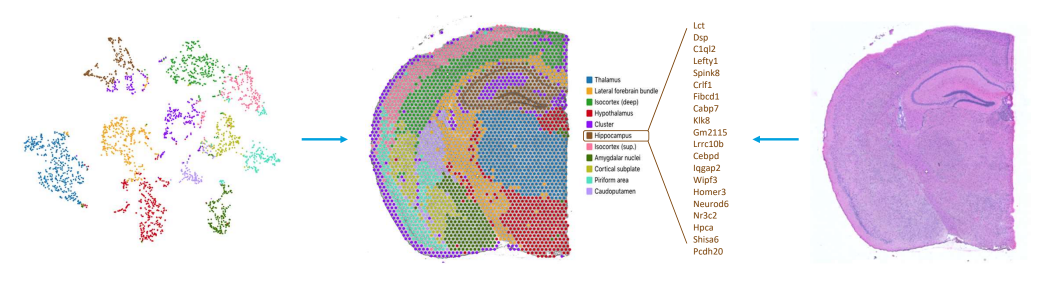
Loupe Browser

- View spot clusters and expressed genes in the context of the tissue image
- Manual alignment
- Interactive data visualization and exploration
- Easily export and share results



Cluster or Image Driven Analysis of Spatial Data

Start With the Gene Expression Data or microscopy images of the same section



Clustering/classification of glass slide spots based on their gene expression profile

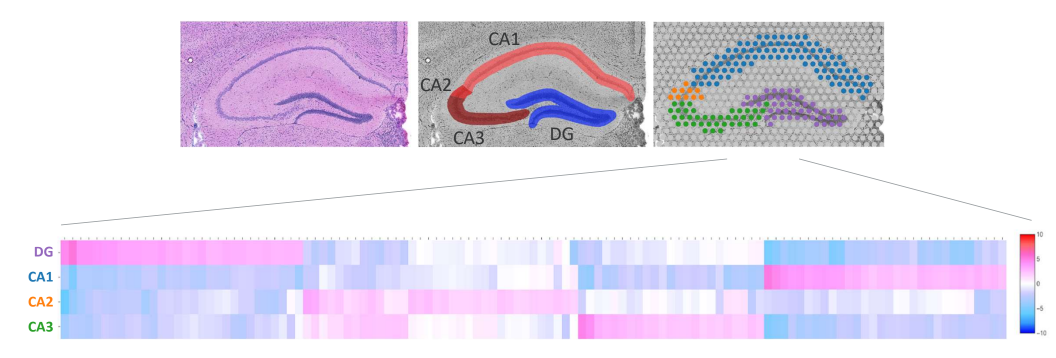
Annotation directly on H&E image

Captured H&E image from the microscope



Spatially Resolved Expression in the Mouse Brain

Dissecting Gene Expression within the Hippocampus





Applications

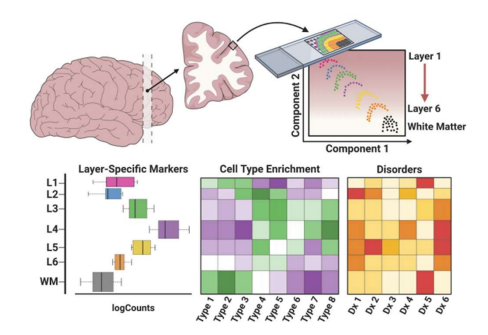


1st preprint in BiorXiv

Maynard et al. 2020 BiorXiv

Transcriptome-scale spatial gene expression in the human dorsolateral prefrontal cortex

Kristen R. Maynard^{1,*}, Leonardo Collado-Torres^{1,2,*}, Lukas M. Weber³, Cedric Uytingco⁴, Brianna K. Barry^{1,5}, Stephen R. Williams⁴, Joseph L. Catallini II³, Matthew N. Tran^{1,6}, Zachary Besich^{1,6}, Madhavi Tippani¹, Jennifer Chew⁴, Yifeng Yin⁴, Joel E. Kleinman^{1,7}, Thomas M. Hyde^{1,7,8}, Nikhil Rao⁴, Stephanie C. Hicks³, Keri Martinowich^{1,5,7*}, Andrew E. Jaffe^{1,2,3,5,6,7,9,*}



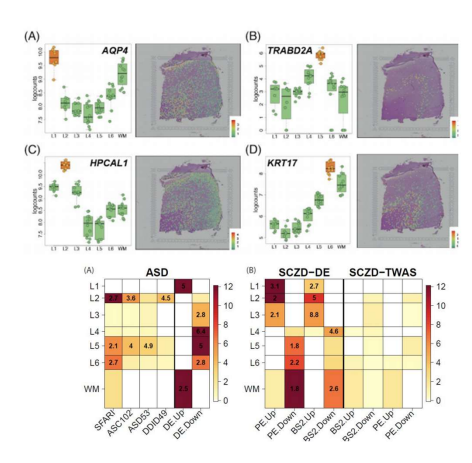


Why using spatial transcriptomics?

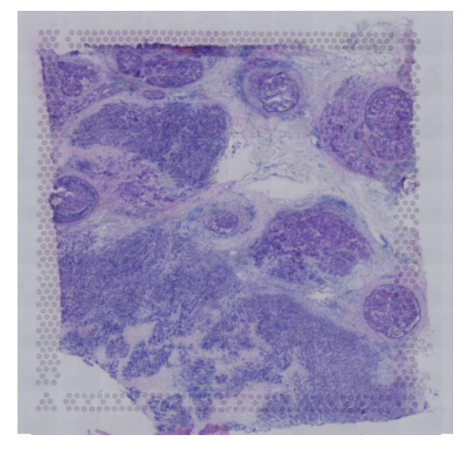
Maynard et al. 2020 BiorXiv

- Human neurons are large and fragile and may not be compatible with SnRNA-seq
- Post mortem tissues SnRNAseq rather than ScRNA-seq
- Differences in pathology and gene expression associated with neuropsychiatric disorders are localized to specific cortical layers
- Topography of human brain dorsolateral prefrontal cortex crucial in some neuropsychiatric disorders



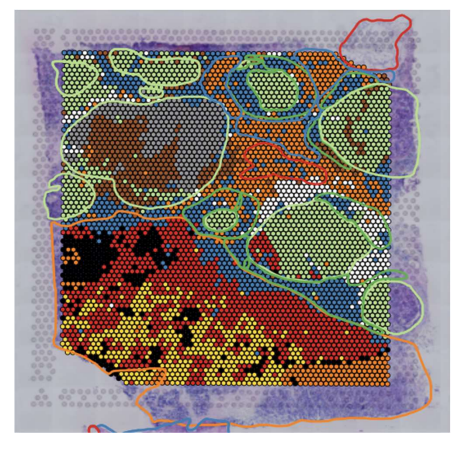


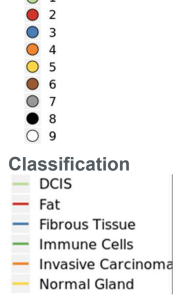
Pathologist Annotations Correlate with GeX Clusters





Pathologist Annotations Correlate with GeX Clusters



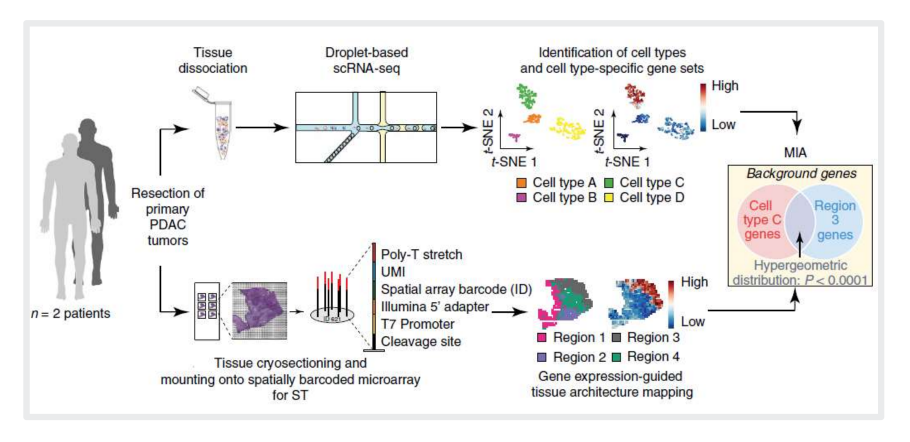


Cluster



Integrated approach reveals tissue architecture in pancreatic ductal adenocarcinomas

Understanding tissue architecture



The Visium Spatial Gene Expression Solution

Features



Sensitivity

Thousands of genes detected per spot



Efficient

1 day workflow time



Resolution

~5000 spots per capture area Spot size 55 μm diameter Close packing of spots; 100 μm c-t-c distance



All Inclusive

All slides and reagents included in kit



Unbiased

Assaying all mRNA with polyA capture technique



Cell Resolution

1-10 cells per spot; dependent on tissue type and thickness



But... This is just the beginning



Visium Spatial Portfolio



Targeted Gene Expression

Immunofluorescence Feature Barcode technology

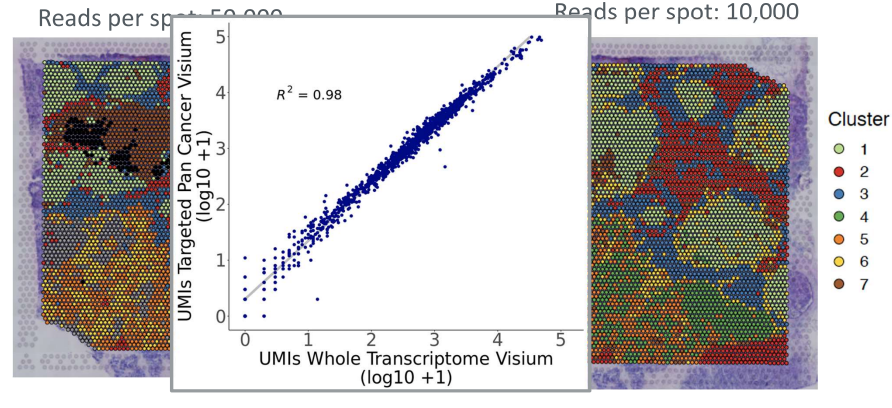
FFPE Solution



Targeted Gene Expression: Go Deep on What You Care About

Whole Transcriptome, Standard Visium

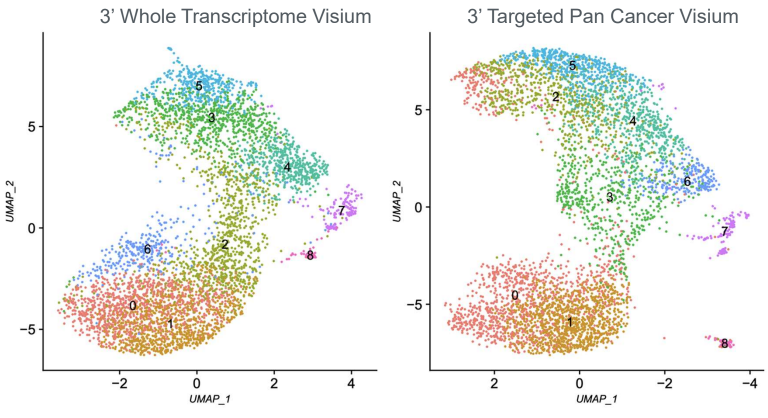






Targeted Gene Expression: Go Deep on What You Care About

Clustering on Pan-Cancer panel genes only



Visium Spatial Portfolio



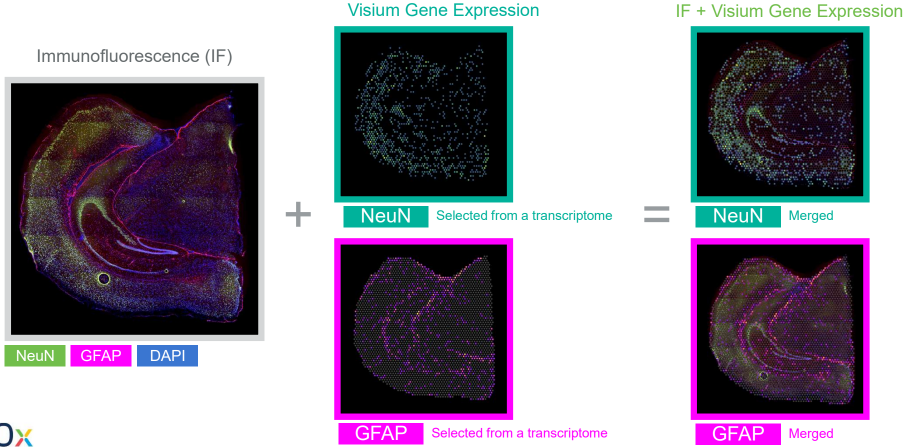
Targeted Gene Expression

Immunofluorescence Feature Barcode technology

FFPE Solution

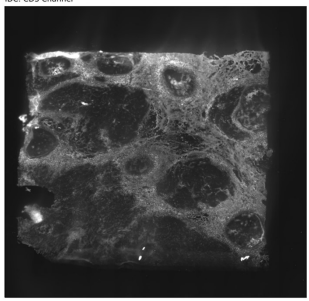


Visium: Immunofluorescence with Gene Expression



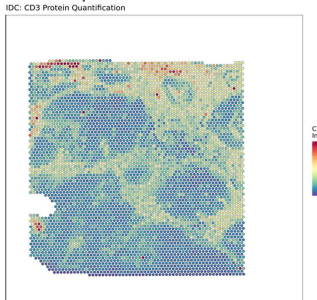
Merge Protein Data into "Spot Space" with mRNA

IDC: CD3 Channel

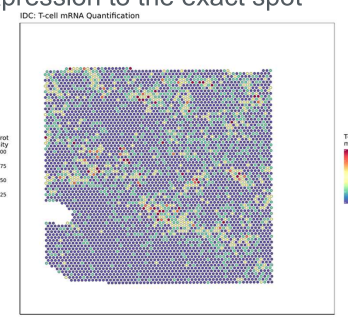


Protein image only

Match protein and mRNA expression to the exact spot



Protein Intensity per spot



T-cell mRNA expression per spot

Visium Spatial Portfolio



Targeted Gene Expression

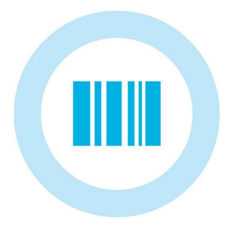
Immunofluorescence Feature Barcode Technology

FFPE Solution



Highly Multiplexed Protein Analysis of Tissues

Feature Barcode Technology

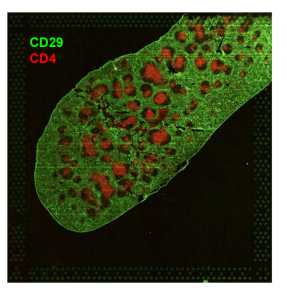


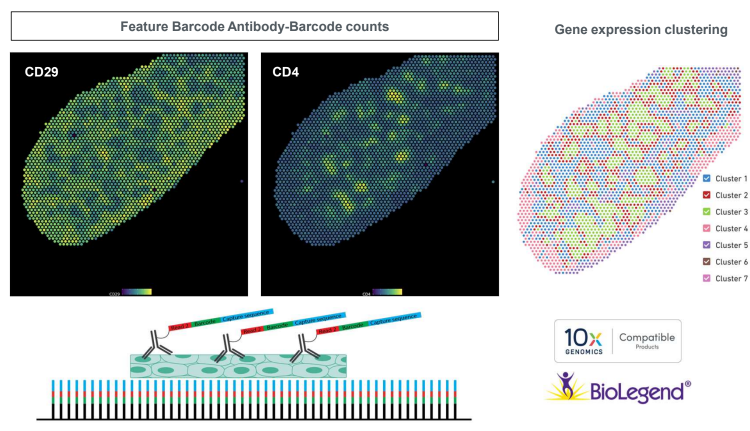
- Tens to hundreds of proteins analyzed at once
- Quantification of protein expression
- Simultaneous whole transcriptome and protein analysis
- No concern of spectral overlap or photobleaching



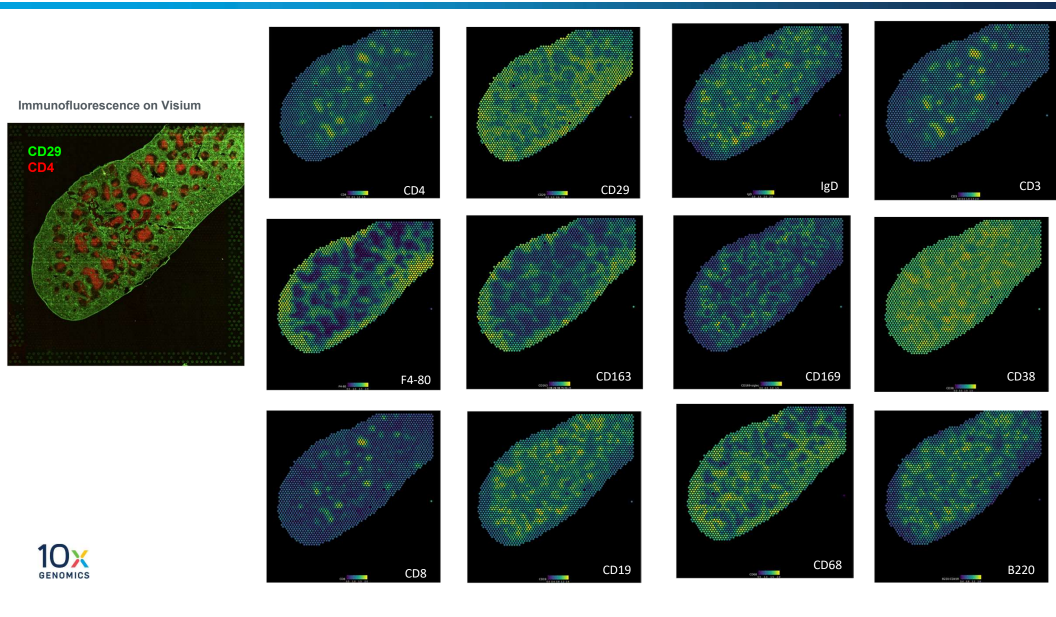
Feature Barcode Correlates with Immunofluorescence











Immunofluorescence on Visium **Feature Barcode** Protein **CD29** Protein Protein Gene expression clustering **mRNA** mRNA**mRNA** ☑ Cluster 1 Cluster 2 Cluster 3 Cluster 4 ☑ Cluster 5 Cluster 6

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Cluster 7

Visium Spatial Portfolio



Targeted Gene Expression

Immunofluorescence Feature Barcode technology

FFPE Solution



FFPE Enablement



- FFPE is the most commonly used preservation method
- RNA is difficult to recover due to fragmentation and crosslinking

10x FFPE protocol and reagent compatibility on current visium

A 10x solution dedicated for FFPE



FFPE Enablement



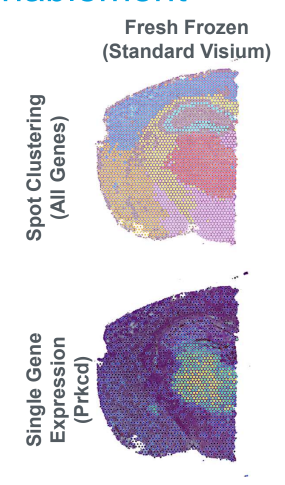
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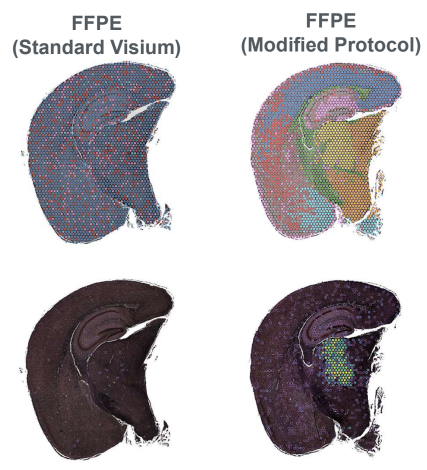
10x FFPE protocol and reagent compatibility on current visium





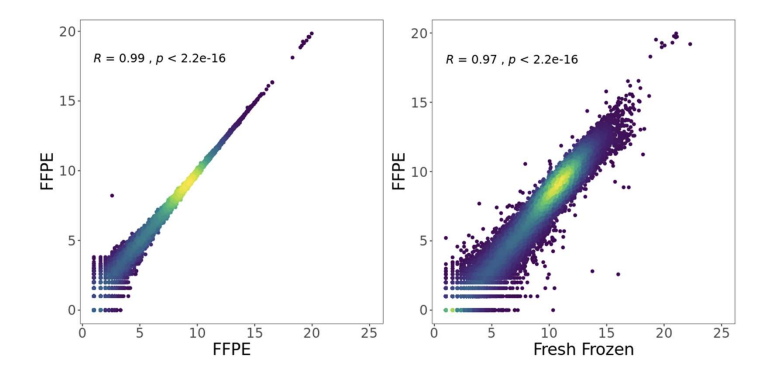
FFPE Enablement





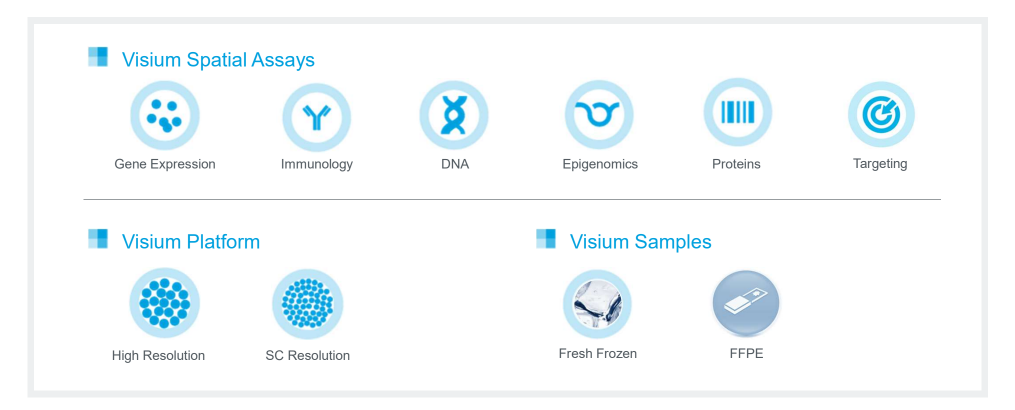


High Correlation Between FFPE and Fresh-Frozen





Towards a Comprehensive View of Biology





Thank You! Questions?

Christophe Fleury

GENOMICS

Science and technology advisor

10x Genomics Team

Jamal FAKIR

Sales Executive

Bashir SADET

Field Applications Scientist

