



Pyxa[®]



Stellaromics



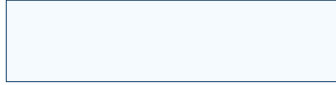
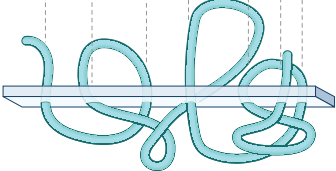
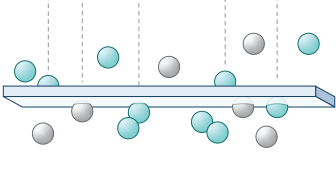
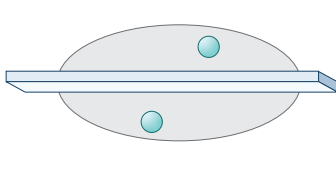
Discover the only technology that creates fully annotated, single-cell maps of 3D gene expression in thick tissue sections up to 100µm. Built on STARmap[™] and RIBOmap[™] technologies.



Stellaromics' Pyxa[®] platform delivers true 3D spatial multi-omics in thick tissue sections. Stellaromics[®] offers a streamlined end-to-end workflow from kit-based sample preparation to automated fluidics, confocal volumetric imaging and integrated GPU-enabled signal processing on Pyxa to intuitive 3D visualization and analysis with PyxaStudio[™] software.

[Stellaromics.com](https://www.stellaromics.com)

Pyxa is Spatial, Cubed

	Convolved Shapes	Complex Distributions	Sparse Objects
2D Slide View			
3D Reality			
Features	Vasculature, neurons & fibrosis	Spatial relationships between cell populations	Rare cell populations in large volumes of tissue
Applications	NASH, other fibrosis, neuroscience, ophthalmology, organoids	Immuno-oncology, immunology, metabolic disease	Gene therapy, CRISPR, CAR-T

Pyxa from Stellaromics is the first platform to deliver true 3D analysis of thick tissue slices. Based on Stellaromics' patented STARmap technology, the Pyxa system measures RNA and RNA translation within intact tissue at the subcellular level.

Biological reality is governed by 3D cellular organization and interactions within the tissue microenvironment. Therefore, to truly comprehend these processes and develop effective interventions, a 3D perspective is required. This represents a move from observing isolated cells and thin 2D tissue sections to understanding the systemic architecture of disease, profoundly changing the types of questions researchers can ask and answer.

3D spatial transcriptomics allows scientists to survey a larger volume of cells within the same spatial area, increasing the odds of capturing rare events such as gene edits, formation of neural connections, cellular differentiation, or immune cell activation. This technology has the power to rapidly advance numerous fields of research including neuroscience, oncology, developmental biology, and therapeutic development and drug delivery.

3D Spatial Transcriptomics

- **Enhanced Insights:** 3D spatial data with Pyxa provides a deeper understanding of cell-cell interactions and rare events missed by 2D assays.
- **Precision and Accuracy:** Whole cell analysis ensures precise localization of gene expression, critical for detailed molecular studies. Unlock new data modalities such as cell morphology through the analysis of intact cells.
- **Rare Event Detection:** Increased tissue volume imaged improves capture of rare cell populations and rare perturbations.
- **Versatility:** Curated and customizable gene panels for multiple assays make Pyxa™ adaptable to a wide range of research applications.
- **Ease of Use:** Onboard analysis and automated workflows simplify operations, making advanced spatial transcriptomics accessible to more researchers.

Analyze Tissue with Unprecedented Depth

The world's only confocal-imaging based spatial system

Volume Video™ enables 3D spatial analysis at unprecedented scale

Smart scheduling fluidics and incubation. Wells operate independently to minimize run time.

Thermal and vibration stabilized optical system

Real-time data processing with onboard GPUs



- Tissue Thickness: up to 100µm
- Resolution: Subcellular transcript localization at minimum resolution of 115nm lateral and 500nm axial
- Gene Panels: Hundreds of validated genes, with customization options available
- Sample Processing: 12-well plate format
- Workflow: Automated sample imaging and on-instrument data analysis
- Data analysis: PyxaStudio and compatibility with open-source tools for quantitative analysis

Key Features of Pyxa

- 3D Spatial Transcriptomics: Detailed visualization and analysis of gene expression within thick tissue sections, providing insights into cellular interactions and tissue architecture.
- STARmap Assay: Providing comprehensive spatial transcriptomics, along with nuclear detection.
- State of the Art: Custom optics and proprietary multi-channel confocal imaging system uniquely enable high plex, thick tissue analysis.
- Subcellular Resolution: Transcript localization at the subcellular level, allowing for precise identification and localization of gene expression patterns.
- 12-Well Plate Format: Efficient sample processing using a 12-well plate, compatible with high-throughput workflows.
- Curated and Custom Kits: Flexible panel selection for a variety of species and tissues.
- Automated 3D Imaging: Fully automated and integrated fluidics. Includes walkaway sample loading and processing.
- Click-button data analysis: Primary analysis performed on-instrument.
- Visualization Software: Custom-built 3D visualization software, PyxaStudio, included at no additional charge.

Offering a Streamlined End-to-end Workflow

Step 1

Tissue Sectioning

Tissue sectioning



Placement into 12 well plate



Tissue is fixed and attached to each well



Sample preparation in 12-well microplate format increases throughput (each well area is nearly equivalent to a standard 2D slide). Samples can be stored and shipped after preparation.

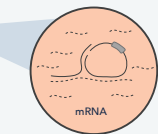
Step 2

STARmap Sample Prep

Thick tissue



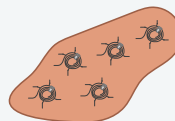
SNAIL™ probe hybridization and ligation



DNA amplicon generation



Tissue is embedded into a 3D hydrogel



Tissue is cleared for sequencing

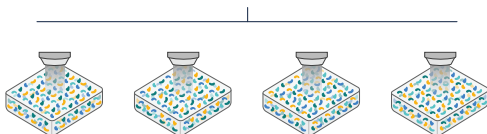


Pyxa chemistry leverages specific amplification of nucleic acids via intramolecular ligation via paired primer and padlock probes (SNAIL probes) to convert target RNA molecules into DNA amplicons with gene-unique codes. This enables highly multiplexed RNA detection in tissue hydrogels by multiple rounds of sequencing by ligation.

Step 3

Automated Imaging and Processing

3D sequencing using Pyxa system



Primary image processing on-instrument



Processed data for analysis

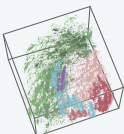
	Cell 1	Cell 2	...	Cell X
Gene A	3	0		11
Gene B	0	5		3
...
Gene X	1	7		0

SEDAL sequencing uses reading probes to decode bases, and fluorescence.

Step 4

Visualization and Data Analysis

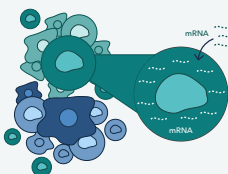
3D gene expression patterns in intact tissue



3D cell cluster and cell-cell Communication analysis



Identify and track cell manipulations



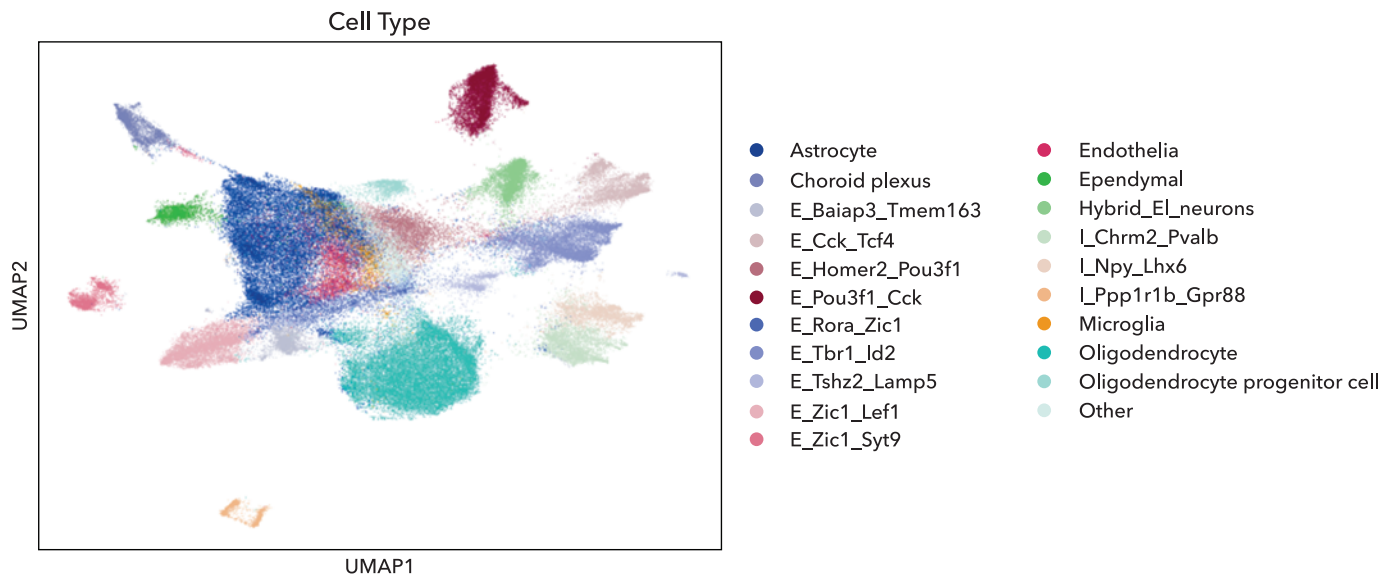
Rare cell communication events



Gene regulation and Translational analysis



Immediate Access to 3D Insights



When you purchase a Pyxa or partner with us for a services project, your data can be seamlessly explored in PyxaStudio, our dedicated visualization tool for navigating 3D gene expression maps, annotating spatial patterns, and quantifying transcript localization in volumetric context. Key features of PyxaStudio include:

- Visualize the spatial distribution of cell types across the tissue
- Explore multiple cell layers in 3D
- Explore cell-cell interactions through cell/transcript proximity in the 3D space
- Investigate subcellular transcript localization
- Filter by specific cell types or genes of interest
- Select individual cells or subclusters via the UMAP view
- Bookmark cells of interest
- Export a subset of the dataset for downstream analysis
- Share views directly with collaborators

Data files are compatible with numerous open-source tools for quantitative analysis

- Exploratory analysis of cellular spatial organization: Monkeybread (python), squidpy (python), Voyager (R and python), Giotto (R), StLearn (python)
- Exploratory analysis of single-cell gene expression: Seurat (R), scanpy (python)
- Analysis of spatially variable genes: Maxspin (python)

Technology Access Services

Be the first to explore spatial biology's next dimension.

TAS Offerings

- Cell typing panel analysis of 20µm-100µm thick, fresh frozen mouse or human tissue
- Predefined or custom gene panels available. Custom scan configurations available by request.
- Tissue sectioning, wetlab workflow, instrument processing, and primary data analysis performed by Stellaromics
- Minimum 4 samples processed

Reach out to discuss your project requirements.

*Working with a different tissue type?
Talk to us!*

Data Deliverables

- Comprehensive data report and technical presentation with Stellaromics Computational Biology team
- Cell by gene matrix including XYZ locations and subcellular localization.
- Customized data and QC reports
- Guidance on open-source visualization and analysis tools
- Access to Stellaromics 3D visualization software, PyxaStudio™
- 8 week sample-to-answer turnaround time for standard projects

