SARDRICS

Simplifying Progress

Label-free, real-time live cell assays for 3D Organoids embedded in Matrigel[®].

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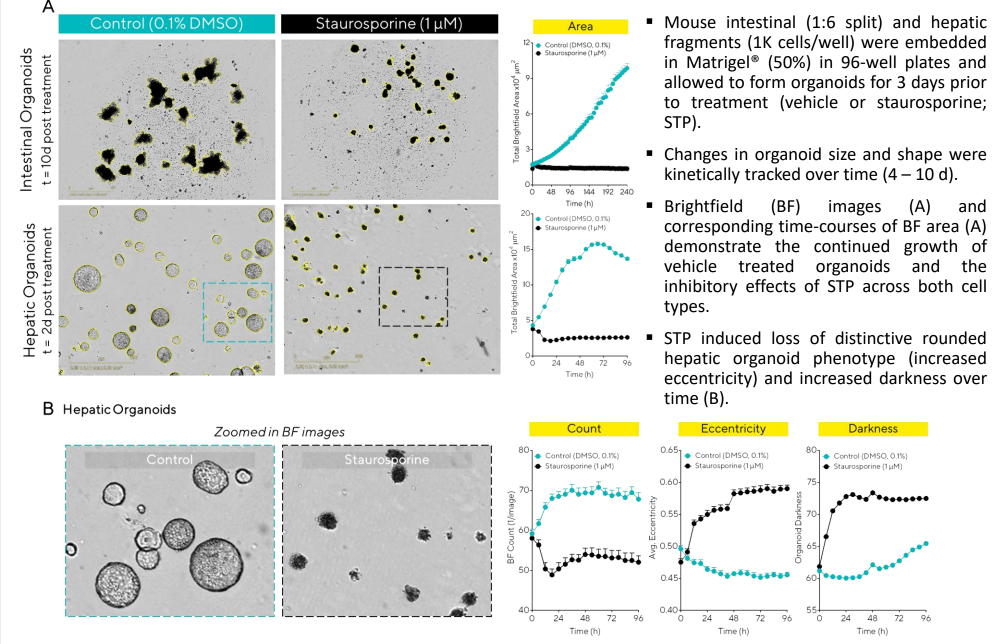
Summary & Impact

- As organoids exhibit structural, morphogenetic, and functional properties that recapitulate in vivo pathophysiology, they are increasingly being used *in vitro*.
- To successfully use these models across a variety of research disciplines and applications, technology pipelines to image & quantify these complex structures are key.
- Here, we demonstrate simple, robust workflows for monitoring and quantifying organoid growth, death and morphology.
- Incucyte[®]'s Organoid Analysis Software Module enables the ability to kinetically visualize and quantify distinct organoid

morphologies embedded in Matrigel[®].

- These validation data demonstrate the ability to characterize the differentiation and maturation of organoid cultures in 24-well plates and assess treatment effects on organoid growth in 96-well microplates.
- Integrated, label-free size and morphology metrics enabled real-time elucidation of compound mechanisms of action and assessment of CFTR function in vitro.
- These data exemplify the amenability of this approach for real-time compound profiling across a range of disease areas.

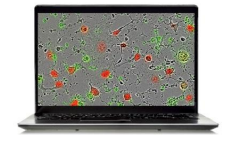
Label free quantification of organoid growth and death



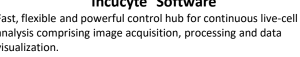
Incucyte[®] System for continuous live-cell analysis: Methodology



Incucyte[®] Live-Cell Analysis System A fully automated phase contrast and multi-color fluorescence system that resides within a standard cell incubator for optimal cell viability. Designed to scan plates and flasks repeatedly over time.









Sartorius Reagents and Consumables A suite of reagents, kits and protocols for cell health and function screening.

Assay Workflows

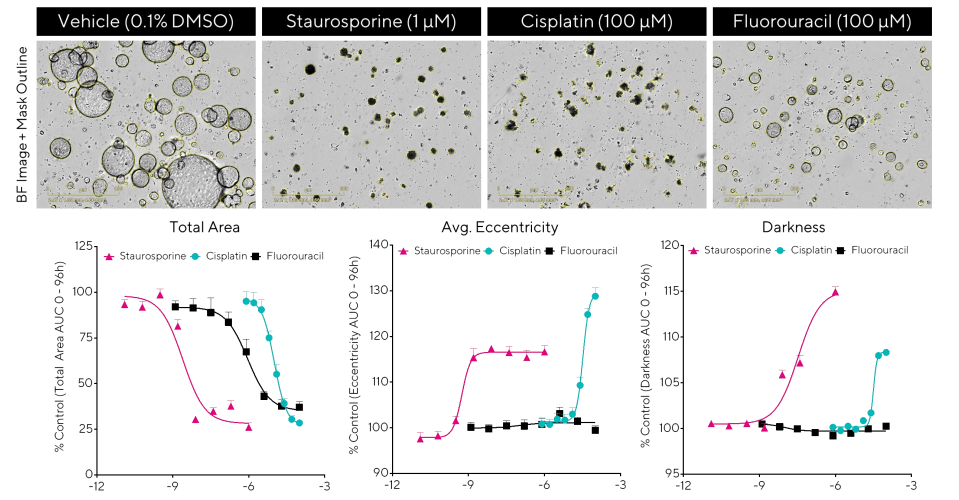
Resuspend

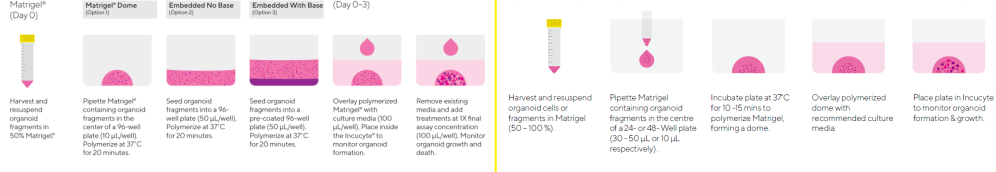
Organoid Assay Quick Guide Organoid QC Quick Guide 3 4 Polymerize matrigel Add media Add cells (Day 0) Harvest & resuspend Pipette matrigel Add media and Add treatments Monitor organoic monitor formation (Day 3) formation & grow

corresponding time-courses of BF area (A) demonstrate the continued growth of vehicle treated organoids and the inhibitory effects of STP across both cell

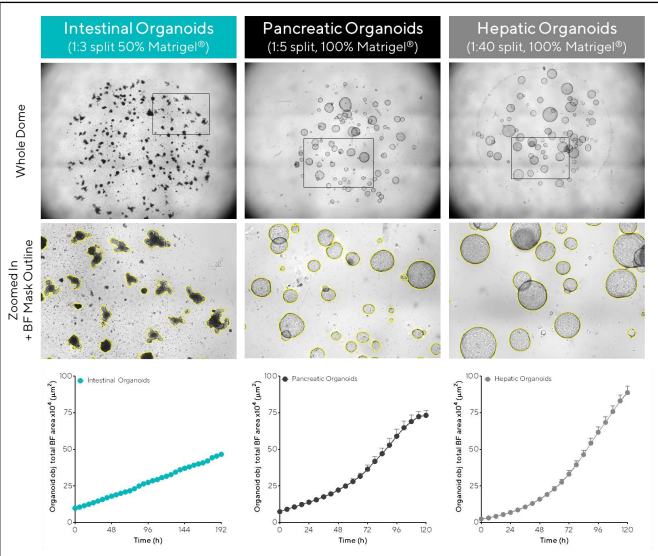
STP induced loss of distinctive rounded hepatic organoid phenotype (increased eccentricity) and increased darkness over

Probing mechanisms of action using morphology measurements





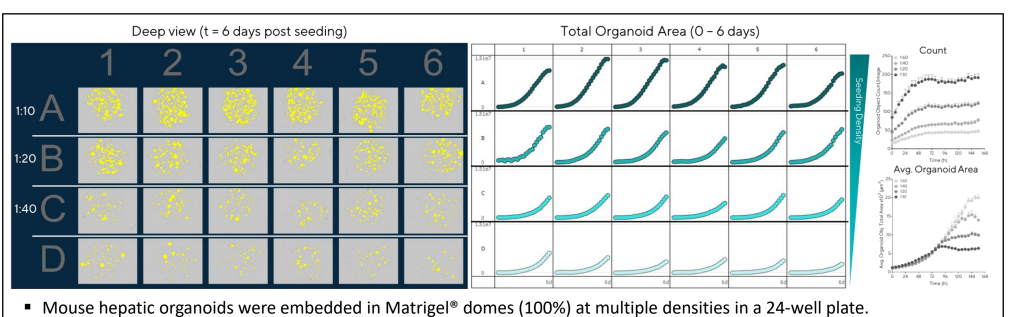
Monitoring and quantifying organoid growth in Matrigel[®] domes



Mouse intestinal, pancreatic and hepatic organoids were embedded in Matrigel® domes in 24-well plates and imaged every 6 hours.

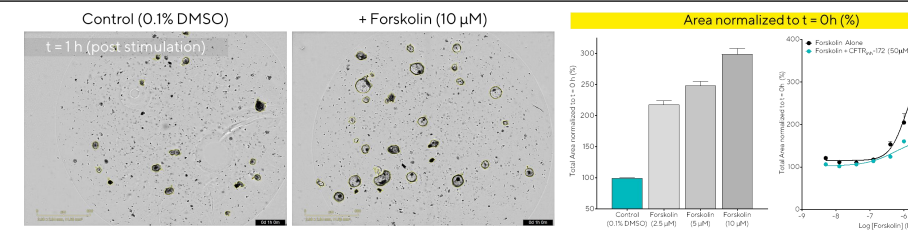
- Organoid growth, differentiation and maturation was measured using Incucyte[®]'s automated Organoid Analysis Software Module which tracks changes in organoid size (area) over time.
- Brightfield (BF) images of the entire Matrigel[®] dome (top) show organoid maturation 6 days post seeding.
- Note accurate segmentation (yellow) outline mask) and distinct phenotypes of mature organoids (bottom).
- BF area time-courses demonstrate cell type specific organoid growth.
- Comparable hepatic and pancreatic organoid growth was observed, while intestinal organoids appeared smaller and exhibited a distinct budding phenotype.

Real-time visualisation & quantification of culture conditions

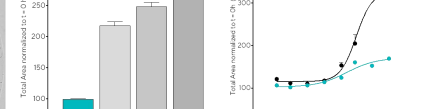


Log [Compound] (M) Log [C	Compound] (M) Log [Compound] (M)
 Hepatic organoids were treated with known chemotherapeutic agents & imaged every 6 h for 4 days. 	3 nM for staurosporine (STP), 9.7 μM for cisplatin (CIS) and 0.78 μM for fluorouracil (5-FU).
 Brightfield images (2d post treatment) show compound-specific effects on organoid size and morphology. Concentration response curves (CRCs) represent the area 	 Increases in eccentricity and darkness indicative of 3D structure disruption and cell death respectively were only observed in STP and CIS-treated organoids (cytotoxic MoA).
 All compounds exhibited concentration dependent inhibition of organoid growth (area), yielding IC₅₀ values of 	 Differences between the size and morphology readouts illustrated the cytostatic mechanism of 5-FU.

Organoid swelling in response to Forskolin stimulation



- Intestinal organoids formed for 3d were treated with increasing concentrations of forskolin and imaged in an Incucyte[®] every 15 – 20 mins for up to 7 hours.
- Brightfield (BF) images show effects of forskolin treatment on organoid size and phenotype.
- Following stimulation, organoids increased in size, exhibited a more rounded phenotype and a clear lumen.



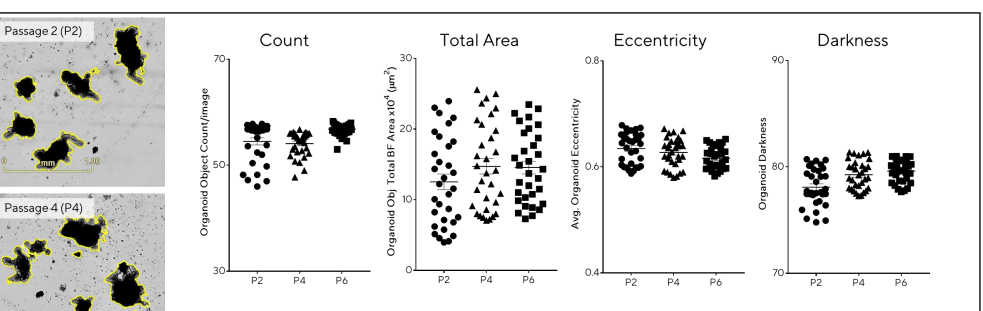
Bar chart of BF area normalized to t = 0 h demonstrates

Passage 6 (P6)

- that swelling is forskolin concentration-dependent.
- In the presence of CFTR inhibitor CFTR_{inh}-172 the maximal response was reduced by > 50% (~150% at 10 μ M) illustrating that swelling was cystic fibrosis transmembrane conductance regulator (CFTR)-dependent.

- Deep view images show brightfield images and segmentation mask overlay 6 days post seeding.
- Incucyte[®]'s real-time automated microplate vessel view and time-course plots demonstrate that organoid growth rate and size is proportional to cell number.

Growth & differentiation efficiency across multiple passages



- Intestinal organoids were embedded in 50% Matrigel[®] domes (1:3 split, 24-well plate) over multiple passages and evaluated for growth and differentiation consistency over time.
- When maintained at a consistent density, organoids exhibited comparable measurements across passages.
- Representative BF images (7 days post seeding) also demonstrated maintenance of distinct budding phenotype across multiple passages.
- Data shown exemplifies the amenability of this imaging and analysis approach to support robust and reproducible assessment of long-term organoid expansion.