

# *Supplemental Manual*

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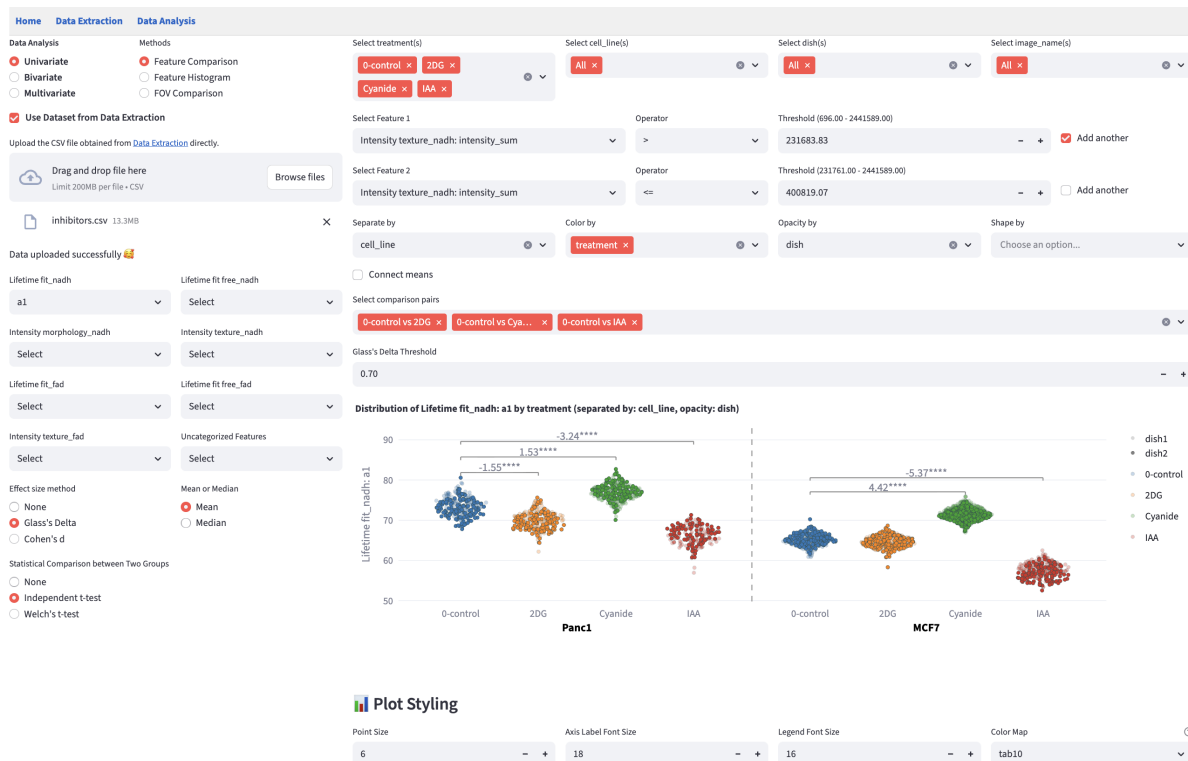
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# 1 Quick Start

Welcome to the FLIM Playground 😊🎉! This is an interactive graphical user interface (GUI) that allows you to extract single-cell features from fluorescence lifetime imaging microscopy (FLIM) raw data ([Data Extraction](#)) and analyze extracted features or datasets extracted via other methods using a built-in repertoire of methods ([Data Analysis](#)).

To quickly try out different analysis methods, download this [sample dataset](#) and try the [online demo](#). If you prefer to use your own data, read [this chapter on data analysis configuration](#) to learn how to configure the system.



Due to the limitation in local file system access, extracting features from raw data is not available in the online demo. Read on to discover how FLIM Playground transforms raw fluorescence decay data into biological insight through an integrated, interactive workflow.

The latest version of this FLIM Playground manual is available [online](#).

## 1.1 Installation

FLIM Playground is built entirely in Python and is open-source.

### 1.1.1 Download

Download the desktop app from [GitHub](#) and double-click it to run. Releases are currently available for Windows 11 and Mac OS 26. If your operating system is not either of these, you can build it yourself by following the instructions below.

### 1.1.2 Build it yourself

1. Clone the repo and navigate into the repository once cloned.
2. Install the Python environment
  - Install uv if not yet installed
  - run `uv sync`
3. Build the app
  - `run pyinstaller Flim-Playground.spec --clean`

## 1.2 Introduction

Fluorescence lifetime imaging microscopy (FLIM) measures the time it takes for a fluorescent molecule to emit light (return to the ground state) after being excited by a pulse of light (enter the excited state). It is sensitive to changes in fluorophore microenvironment including, pH, temperature, and conformational changes due to protein-binding and the presence of quenchers<sup>1</sup>. Coupled with modern automated cell-segmentation methods<sup>2</sup>, FLIM enables single-cell analyses that can reveal biological heterogeneity.

### Instrumentation

To acquire FLIM data, a light source—typically a pulsed laser for time-domain methods or a modulated continuous-wave source for frequency-domain methods—is used to excite the fluorophore of interest. The emission is detected using instrumentation capable of resolving fluorescence decay, such as time-correlated single-photon counting (TCSPC), time-gated, or phase/modulation-based detection. In time-domain FLIM, the delay between excitation and photon arrival is measured, and often a histogram is built, with the x-axis representing the delay time and the y-axis representing the number of photons falling into each time bin. Compared to intensity images, FLIM has an

additional dimension of time (e.g., 256 time bins per 12.5 nanoseconds).

In frequency-domain FLIM, the phase shift and modulation depth of the emission relative to the excitation are determined.

## 1.3 Challenges

A diverse set of [tools](#) — both open-source and commercial, ranging from libraries to code-free graphical user interfaces (GUIs) — is available to extract and analyze FLIM data, providing alternative methods and therefore flexibility to FLIM researchers. Examples include [PhasorPy](#)<sup>3</sup>, an open-source library for analyzing fluorescence lifetime using the phasor approach; [FLUTE](#)<sup>4</sup>, an open-source GUI for interactive phasor analysis; [FLIMPA](#)<sup>5</sup>, an open-source phasor analysis GUI enabling batch processing, ROI-based quantification, and experiment-level comparison through manual assignment; [FLIMLib](#), an open-source generic curve fitting library that can be used to fit fluorescence lifetime decay data; [SPCImage](#)<sup>6</sup>, a commercial software for fitting and phasor features.

However, while some tools offer code-free interfaces, users still need to write custom code—either to prepare data in the proper format as input, or to further process their outputs for downstream analysis. The fragmentation between tools arises because each focuses on only a subset of data levels: pixel, cell ROI, channel, field of view, and experiment.

### 1.3.1 Data Levels

- **Pixel:**
  - A single decay curve encoded in vendor-specific file formats (e.g., Becker & Hickl, Pico-Quant, etc.)
- **Region of Interest (ROI)**
  - Mask with cell labels
- **Channel**
  - Different fluorophores
  - Fluorophore-specific calibration files
  - Masks focusing on different parts of the cell (e.g., whole cell, cytoplasm, nucleus, stain, etc.)
  - Different feature extraction methods (fitting, phasor, morphology, texture)
- **Field of View (FOV)**
  - Input decay types (e.g., [2D](#), [3/4D](#), [pixel-level lifetime features already fitted](#))
- **Experiment**

- Different treatments, time points, cell lines, etc., and combinations thereof

An integrated framework should take into account all data levels ■ □ ▢ ▣ ▤ (•) while maintaining the flexibility to handle various input types (◦). **It should provide a level of abstraction to address fragmentation from data levels and input types.**

Additionally, the use of FLIM is rapidly evolving, and new methods are being developed all the time. An integrated framework should allow users to choose among alternative methods seamlessly, be the backbone of iterative explorations integral to research, and be ready to incorporate new methods: **The provided level of abstraction should address fragmentation from extraction and analysis methods.**

Finally, many of the existing tools, including GUI-based software, are not cross-platform, which limits their accessibility. The [Installation](#) addresses this last challenge.

A closer parallel to FLIM Playground’s integrated approach is the combination of [CellProfiler](#)<sup>7</sup>, which extracts per-object morphological and texture features, and [CellProfiler Analyst](#)<sup>8</sup>, which ingests these outputs or other feature tables for visualization and statistical analysis. However, FLIM Playground stands apart in that it can extract lifetime features from time-resolved data, alongside morphological and texture features from intensity images. Its general Data Analysis section incorporates FLIM-specific methods (e.g., [phasor analysis](#)) and provides statistical models that address additional analysis aspects such as data heterogeneity, in addition to training machine learning classifiers.

## 1.4 Method

### 1.4.1 Feature Classes

Tabular data columns can be categorized into three feature classes:

- **Identifiers**: unique row identifier and (optional) field of view identifier that allow us to investigate the biological heterogeneity at the single-cell level
- **Categorical features**: conceptually help us group the rows (e.g., `treatment` will group the rows into different treatment groups)
- **Numerical features**: quantify the differences/similarities between data groups

**Science, from the data perspective, is about closing the conceptual categorical gaps with quantitative measurements.**

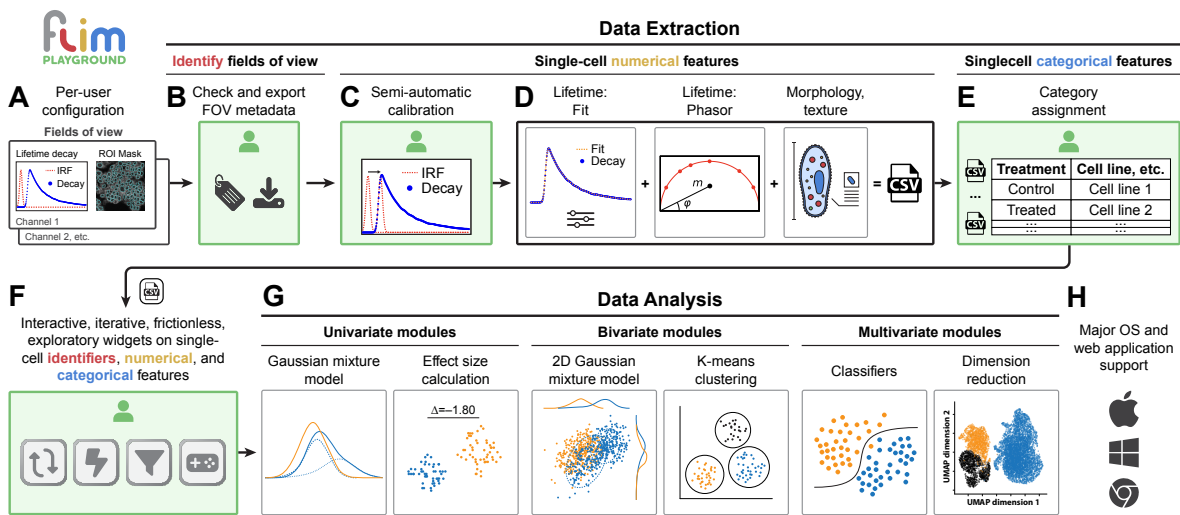
At [data levels](#), the data are processed to extract the feature classes in [Data Extraction](#), and the classes are used in the [Data Analysis](#) section.

Increasing complexity ↑

Data Levels	Data Processing	Feature Classes
Experiment	Assign category labels (e.g., treatment, cell line, etc.) to each cell; perform data analysis on labeled cell groups.	Categorical features
Field of View	Assign FOV identifier to each cell, obtain unique cell identifiers	Identifiers
Acquisition	Apply <b>per-channel</b> ROI mask, calibration and extraction methods to cell-level signals	Numerical features
Channel	Aggregate pixel-level decays to obtain cell-level signals	N/A
Cell ROI	Read pixel-level decays encoded in different formats	N/A
Pixel		

### 1.4.2 Design

FLIM Playground has two independent sections:



#### 1.4.2.1 Data Extraction

**Data Extraction** extracts single-cell features from the raw data. It adopts a framework that offers channel-level flexibility in input types and extraction methods without incurring too much overhead for users. Following the above **categorization**, it is divided into the following steps:

- **Data Extraction Configuration (A)**: allows users to choose among alternative input types and extraction methods (extractors).
- **Metadata organization (B)**: extracts the field of view **identifiers** and their configurations
- **Numerical Feature Extraction**: extracts single-cell **numerical features** based on user-selected extractors. More extractors can be integrated in the future.
  - Calibration (C): calibrate for IRF shift or use fluorescence lifetime standard
  - Alternative lifetime extractors (D):
    - \* **Fitting**
    - \* **Phasor**
  - Alternative intensity-based extractors (D):
    - \* **Morphology**
    - \* **Texture**
- **Categorical feature extraction (E)**: extracts single-cell **categorical features** and combines experiment-level datasets.

#### 1.4.2.2 Data Analysis

**Data Analysis** analyzes features—whether extracted through Data Extraction or by other methods—using visualizations and statistical modeling. It deploys a **shared framework (F)** built to handle the **feature classes** across all analysis methods, enabling the same interactive and frictionless exploration experience and allowing new methods to be integrated in the future easily.

- **Data Analysis (G)** goes in-depth into how FLIM Playground handles the feature classes and **Data Analysis Config** goes through how users can configure FLIM Playground to analyze datasets that are not extracted by **Data Extraction**.

Here is the list of analysis methods incorporated into FLIM Playground, grouped by the number of numerical features they take as inputs:

- **Univariate analysis**
  - **Feature Comparison**
  - **Feature Histogram**
  - **Field of View Comparison**
- **Bivariate analysis**
  - **Feature Distribution**
  - **Phasor Analysis**
- **Multivariate analysis**
  - **Dimension Reduction**

- Classification

Both sections are built in Python and as self-contained executables ready to run on major operating systems and in browsers (**H**).

### 1.4.3 Summary

**FLIM Playground** resolves these [challenges](#) with an interactive code-free graphical user interface (GUI) that spans the full pipeline. It integrates validation checks that guide users at every step, and has a built-in repertoire of analytical methods with interactive widgets that encourage hypothesis-driven, iterative exploration of large datasets. It is built on a modular architecture that enables incorporation of new algorithms in the future.

### 1.4.4 Citation

FLIM Playground is currently on [bioarchive](#). If it contributed to your research—whether through Data Extraction for single-cell feature generation or through Data Analysis for data exploration, visualization, selection of analysis methods, or hyperparameter tuning (UMAP, clustering, classification, etc.)—please cite this work in your publication. Your citation directly supports us in maintaining and improving it



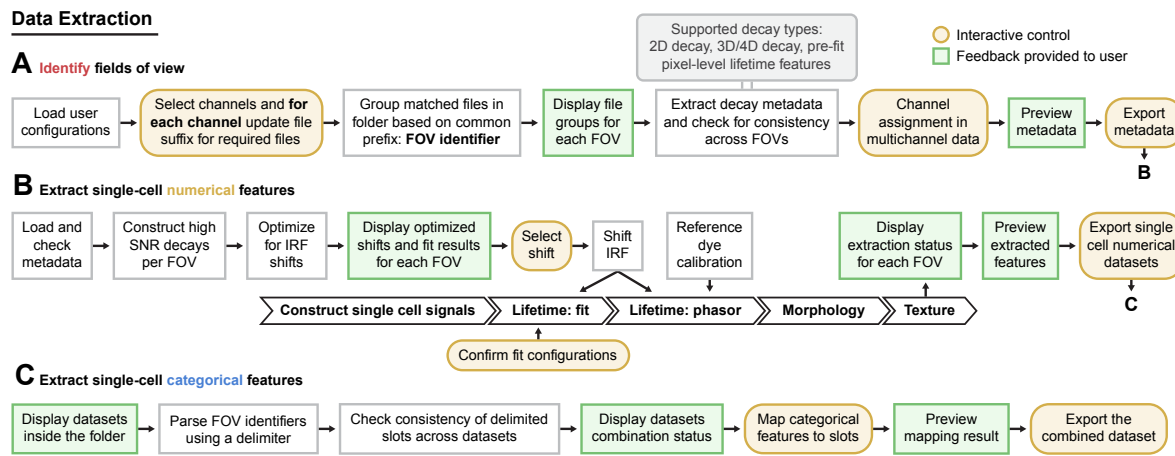
# **Part I**

# **Data Extraction**

## 2 Overview

FLIM Playground transforms raw FLIM data into single-cell numerical and categorical features, ensuring smooth integration with [downstream analysis](#). All steps are seamlessly connected through interactive widgets, with built-in error checking and reporting to ensure correctness.

A high-level overview of the data extraction process is shown below.



### 2.1 Stages

Users can [configure](#) the system once and apply it to future data. After configuration, users can extract features from raw data using the following steps, inspired by the [categorization](#):

- Select a step to perform [?](#)
- FOV Metadata Extraction
  - Numeric Feature Extraction
  - Categorical Feature Extraction

1. **FOV Metadata Organization (A)**: each experiment session consists of multiple fields of view (FOVs). This step helps users organize the metadata of the FOVs.

2. **Numerical feature extraction (B)**: it calibrates and then extracts single-cell numerical features.
  - Calibration
    - Fit Calibration
    - Fit Free Calibration
      - \* IRF Shift
      - \* Fluorescence lifetime standard
  - Per-channel feature extraction includes:
    - Lifetime fitting features: fit an exponential decay curve to the measured decay and extract lifetime features per cell ROI.
    - Phasor features: calculate the phasor features such as the phasor coordinates. It also provides phasor analysis.
    - Morphology features: calculate morphology features based on the ROI mask.
    - Texture features: calculate texture features based on the intensity image and the ROI mask.
3. **Categorical feature extraction (C)**: extract the categorical features such as the treatment, time point, etc. based on the FOV name.
  - Users can then upload the extracted datasets to perform [Data Analysis](#).

## 2.2 Input File Types

### 2.2.1 Decay

See [decay types](#) for more details.

- 2D decay:
  - .csv: a tabular data sheet with each row representing a cell and each column representing a time bin.
- 3D/4D decay: besides the spatial dimensions and the time dimension, the additional dimension in the 4D array represents acquisition channels.
  - .sdt: Becker & Hickl
  - .ptu: PicoQuant

## 2.2.2 Mask

Because of the single-cell focus of FLIM Playground, cell-level masks are needed for each channel. Channels can share the same mask, or use different masks (when they use different masks, the mask ID for each cell should match across masks).

The cell-level mask can focus on different parts of the cell region, such as the whole cell, the cytoplasm, the nucleus, or the stain part.

For an example, see [here](#).

- `.tiff/.tif`: a 2D array with background labeled as 0 and each cell ROI region labeled as a unique positive integer. That integer will be part of the [unique cell identifier](#).

## 2.2.3 IRF

IRF must be a 1D array. Currently, the supported formats are (extendable to more formats in the future):

- `.txt`: it uses `np.loadtxt` to parse a txt file. It returns a 1D array if the IRF numbers are in one row or in one column.

## 2.2.4 SPCImage t1

If the decay type is [3D/4D pixel-prefitted](#), users can provide pixel-prefitted lifetime feature files.

- `.asc`: a 2D array in spatial dimensions outputted by [SPCImage](#), with each (row, column) containing the value of the lifetime feature (e.g. `t1`: the first component's lifetime) of that pixel.
- Depending on the chosen number of lifetime components, the other pixel-prefitted feature files are included and their names are inferred from the `t1` file, by replacing `t1` in the file name with `t2, t3, a1[%]`, etc.

## 2.2.5 Fluorescence Lifetime Standard

If the fluorescence lifetime standard-based calibration is chosen, users are expected to provide the fluorescence lifetime standard file.

- `.tiff/.tif`: a 3D array, with one dimension representing the time axis.

### 2.2.6 Intensity (2D)

If the channel's [imaging modality](#) is **Intensity-only**, users are expected to provide a 2D intensity image in:

- `.tiff/.tif`: a 2D array.

## 2.3 Limitations

- To simplify the workflow and due to the framework chosen, FLIM Playground does not support pixel-level fitting and fit-free analysis, and there are many other open-source tools that can do so (for a comprehensive list, see [here](#)). Instead, it does cell-level fitting and fit-free (phasor) feature extraction by summing up all the decays belonging to the same cell ROI to get one cell-level decay as a preprocessing step<sup>9</sup>. It also accepts prefitted pixel-level lifetime features from [SPCIImage](#) and [aggregates pixel-level lifetime features to cell-level lifetime features](#).

# 3 Configuration

## Note

Users can configure the system once, and it will be applied to future data.

Constructing a framework that balances flexibility and usability is about choosing the right abstraction level. FLIM Playground chooses acquisition *channel* from [data levels](#) as the focus and adopts the **channel-centric framework**. Settings are divided into two categories: [cross-channel configurations](#) and [per-channel configurations](#).

## Configuration

The screenshot shows the FLIM Configuration interface with three main sections for different channels:

- Channel 1: nadhl**
  - FLIM Decay Input type: Decay (3/4D)
  - Laser rate (GHz) for Decay (3/4D): 0.08
  - Fit free calibration method: IRF (radio button selected)
  - Imaging modality: FLIM
  - Channel 1 name: nadhl
  - Extract feature types from nadhl: Intensity morph..., Intensity texture..., Lifetime fit free...
- Channel 2: fad**
  - FLIM Decay Input type: Decay (3/4D)
  - Laser rate (GHz) for Decay (3/4D): 0.08
  - Fit free calibration method: Reference Dye (radio button selected)
  - Imaging modality: FLIM
  - Channel 2 name: fad
  - Extract feature types from fad: Intensity morph..., Intensity texture..., Lifetime fit free...
- Channel 3: cd8**
  - FLIM Decay Input type: Decay (3/4D)
  - Laser rate (GHz) for Decay (3/4D): 0.08
  - Fit free calibration method: Reference Dye (radio button selected)
  - Imaging modality: Intensity-only
  - Channel 3 name: cd8
  - Extract feature types from cd8: Intensity morph..., Intensity texture...

Common settings across all channels include:

- Unique cell identifier column name: cell\_id
- FOV column name: image\_name
- Intensity (2D): r.ome.tif
- Mask: r.ome\_cellpose.tif

Other settings include:

- File suffixes: nadhl, fad, cd8
- Categorical columns (type to add more): cell\_line, treatment, condition, patient\_id, day

At the bottom is a "Update Configuration" button.

## 3.1 Cross-Channel Configurations

### 3.1.1 Categorical Features

Categorical columns (type to add more)

cell\_line ×treatment ×condition ×patient\_id ×day ×experiment

Add: experiment

Categorical features are useful to organize the data into groups of interest, and will be used extensively in the [Data Analysis](#) section. Users can specify the potential categorical feature names to be extracted in the [Categorical Feature Extraction](#) step.

### 3.1.2 Decay Types

FLIM Playground supports different decay types, including 2D decay, 3D/4D decay, and 3D/4D pixel-prefitted decay, through user specification.

#### 3.1.2.1 2D Decay

In the system developed by Samimi et al.<sup>10</sup>, single cells flow through and a decay curve is acquired for each cell by aggregating all the photon arrival time delays deemed to be from the same cell. In exchange for spatial distribution, the acquisition speed is increased enormously. The output of the system is a tabular data sheet with each row representing a cell and each column representing a time bin. **FLIM Playground currently supports 2D decay in the CSV format.**

#### 3.1.2.2 3D/4D Decay

In TCSPC-based FLIM, the data is stored in a 3D/4D array and in special format (e.g. .sdt, .ptu). In addition to the spatial dimensions and the time dimension (XYT), the channel dimension may also be present, making it 4-dimensional (CXYT). **FLIM Playground currently supports both .sdt and .ptu files up to those 4 dimensions.** It reads the duration and the number of time bins from the decay file metadata. Some systems can also record a time stack, which is not supported yet.

### 3.1.2.3 3D/4D pixel-prefitted

Since FLIM Playground currently does **not** support pixel-fitting, users have the option to provide pixel-prefitted outputs from [SPCImage](#) in the format of .asc. See [here](#) for how FLIM Playground processes SPCImage outputs to obtain cell-level lifetime fitting features.

### 3.1.3 Identifiers

Unique cell identifier column name	(?)	FOV column name
<code>cell_id</code>		<code>image_name</code>

#### 3.1.3.1 Cell Identifier

Users can specify the column name for the unique identifier of the cell to be stored in the output dataset. The cell identifier is constructed as {FOV Identifier}\_{cell\_label}, where the `cell_label` is the unique integer label in the mask or the row number if the decay type is [2D decay](#).

#### 3.1.3.2 FOV Identifier

Users can specify the column name for the field of view identifier. If the decay type is [2D decay](#), users may want to specify the column name as `exp_name`, because the FOV is not an image.

### 3.1.4 Decay Info

#### 3.1.4.1 Laser frequency

Users can specify the laser frequency in GHz. For example, 0.08 GHz equals 80 MHz. It will be used in the [phasor analysis](#). If you do not intend to use the phasor analysis, you can fill in any number.

#### 3.1.4.2 2D Decay-specific

Since the [2D decay](#) file does not provide the duration and the number of `time bins` per laser pulse interval, users need to specify them.

Decay (2D) duration (s)	Decay (2D) time bins
<code>20.00</code>	<code>1024</code>
-	+

### 3.1.5 Calibration Method

It supports two calibration methods for phasor analysis:

- IRF shift-based calibration
- [fluorescence lifetime standard-based calibration](#)

Since the instrumental response function (IRF) or the fluorescence lifetime standard is measured for each channel, they are set in the [per-channel configurations](#). The fluorescence lifetime standard's lifetime is channel-agnostic.

Fit free calibration method

IRF  
 Reference Dye

Reference dye lifetime (ns)

4.20

- +

Provide channel-specific Reference Dye file suffixes below in the File suffixes section.

### 3.1.6 Number of Channels

Each FOV is assumed to have the same number of channels. The configuration will dynamically adjust the [per-channel configurations](#) based on the specified number of channels. Currently, the maximum number of channels is 4, and it can be extended to house more channels.

## 3.2 Per-Channel Configurations

### 3.2.1 Channel Name

Users can customize the channel name based on, for example, the name of the fluorophore.

Channel 1 name

nadhl

Channel 2 name

fad

Channel 3 name

cd8

### 3.2.2 Imaging Modality

Users can specify the imaging modality for each channel. Currently, two modalities are supported:

- FLIM: The signal is time-resolved, i.e., has a time dimension, and is expected to be from one of the [decay types](#).
- Intensity-only: The signal is a spatial map of intensity values (2 spatial dimensions), i.e., has no time dimension. **Currently, the only supported signal type is a 2D intensity image in the format of .tiff/.tif.**

Imaging modality

Intensity-only |

FLIM

Intensity-only

💡 Tip

This setup allows users to image some channels in FLIM mode, and others in non-FLIM mode. It also allows users to extract and analyze their data if they image in non-FLIM microscopic systems such as epifluorescence, confocal, brightfield, etc. by selecting **Intensity-only** for all channels.

ℹ Note

If the decay type is **2D decay**, the **imaging modality** for all channels is restricted to **FLIM**.

### 3.2.3 Feature Extractor

Four types of feature extractors are available if the **imaging modality** is **FLIM**:

- Lifetime fit
- Lifetime fit free
- Intensity Morphology
- Intensity Texture

Extract feature types from nadh

Lifetime fit free ×

Intensity morph... ×

Lifetime fit ×

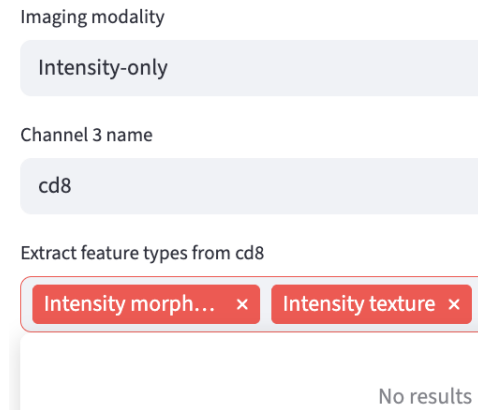
✖ ▾

Intensity texture

ℹ Note

If the decay type is **2D decay**, the **Intensity Morphology** and **Intensity Texture** feature extractors are **not** applicable.

The **Intensity Morphology** and **Intensity Texture** feature extractors are available if the **imaging modality** is **Intensity-only**.



Each channel can be assigned its own set of feature extractors. For example, if users do not intend to do lifetime extraction, they can de-select the `Lifetime fit` and `Lifetime fit free` options.

### 3.2.4 Number of Components

If users select the `Lifetime fit` under this channel, they can specify the number of components to fit.

### 3.2.5 File Suffix

The [FOV Metadata Organization](#) step looks for associated input files for all channels of a given FOV using file suffixes. Based on the [decay type](#), [calibration method](#), and the selected [feature extractors](#), the system automatically generates the file suffixes for required input files.

File suffixes: nadh	File suffixes: fad	File suffixes: cd8
Decay n.sdt	Decay f.sdt	Decay s.sdt
IRF NADH_IRF.txt	IRF FAD_IRF.txt	Mask stain_mask.tif
Mask n_photons_mask_cyto.tif	Mask n_photons_mask_cyto.tif	

In the example above, channels `nadh` and `fad` were asked to provide an IRF file, while channel `cd8` did not, because `Lifetime fit` was selected for the former two channels and not for the latter. The file suffixes of the IRF files were different for each of the two channels, but they shared the same ROI mask file suffix because they used the same ROI mask. The `cd8` channel used a different ROI mask than the other two channels, indicated by the mask file suffix.

For detailed information on what file formats are supported for each file type (Mask, IRF, etc.), see [input file types](#).

For detailed information on how the file suffixes are used, and what each file type (e.g. IRF, Mask, etc.) is, see the [FOV Metadata Organization](#) page.

### 3.3 Save

Finally, users can save the configuration for future use by clicking the `Update Configuration` button.

# 4 Field of View Metadata

It is the first step in the [Data Extraction](#) workflow. It organizes the metadata of each field of view (FOV)—including file paths for all required inputs and decay information—into a single metadata file. This file is then used for [numerical feature extraction](#), as well as for users' own bookkeeping and troubleshooting.

## Data Extraction

The screenshot shows the 'Data Extraction' interface. On the left, a configuration panel allows selecting steps like 'FOV Metadata Extraction' and 'Feature Extraction'. It includes dropdowns for 'Decay input type' (set to 'Decay (3/40)') and 'File suffixes' (set to 'nad' and 'fad'). A 'Feature extractors' section lists 'Intensity morphology', 'Intensity texture', and 'Lifetime fit'. A 'Laser rate (GHz)' slider is set to 0.80. Below these are sections for 'File suffixes: nadh' and 'File suffixes: fad', each with tabs for 'Mask', 'Decay', and 'IRF'. The 'nad' section shows 'n\_photons\_mask\_cyto.sdt' selected for Mask and 'NADH\_IRF.txt' for IRF. The 'fad' section shows 'f.sdt' selected for Mask and 'FAD\_IRF.txt' for IRF. A note at the bottom states 'Channels ['nadh', 'fad'] share the same mask suffix: n\_photons\_mask\_cyto.tif'. On the right, a large grid displays extraction status for 11 FOVs (H9\_DAY\_40\_5 to H9\_DAY\_40\_11). Most are marked as 'All files found.' Some show errors: 'H9\_DAY\_40\_11' has a red 'X' and 'Missing or duplicate files' with 'Missing nadh\_Decay: H9\_DAY\_40\_11.sdt' and 'Missing fad\_Decay: H9\_DAY\_40\_11.f.sdt'. A green bar at the bottom indicates successful loading of most views. A 'Copy the folder path here' field contains '/Users/allan/Downloads/40'. A 'Export FOV Metadata as CSV' button is at the bottom right.

It is divided into two sections:

- The left 1/3 of the screen is the metadata configuration panel, inherited from the [Data Extraction Configuration](#) step.
- The right 2/3 shows the extraction status for all FOVs.

## 4.1 Metadata

Metadata are configured earlier in the [Data Extraction Configuration](#) step and copied here. Some fields remain editable for flexibility, while others are fixed to ensure usability. Non-editable fields can be

modified in the Data Extraction Configuration step. After saving your modifications, return to this page and refresh it to see the changes.

#### 4.1.1 Decay Type

See [decay types](#) for more details. The decay type is not editable here.

Decay input type: Decay (3/4D)

#### 4.1.2 Channel Names

See [channel name config](#) for more details. The names are not editable here, though users can choose which channel to include.

has nadh

has fad

#### 4.1.3 Feature Extractors

See [feature extractor config](#) for more details. The feature extractors for each channel are shown but not editable here.

The image shows two separate configuration panels. The left panel is titled 'Feature extractors for nadh' and the right panel is titled 'Feature extractors for fad'. Both panels have a dropdown menu icon at the top right. The 'fad' panel is expanded, showing a list of three items: '0 : "Intensity morphology"', '1 : "Intensity texture"', and '2 : "Lifetime fit"'. The 'nad' panel is collapsed.

#### 4.1.4 Decay Info

See [decay info config](#) for more details. The decay information is editable here.

A horizontal input field with a light gray background. The text 'Laser rate (GHz)' is at the top left. In the center, the value '0.80' is displayed. At the far right are two small buttons: a minus sign '-' and a plus sign '+'.

#### 4.1.4.1 2D Decay-specific

Since the [2D decay](#) file does not provide the duration and the number of time bins per laser pulse interval, users need to specify them.

Duration (s)	Time bins	Laser rate (GHz)			
20.00	- +	1024	- +	0.50	- +

#### 4.1.5 File Suffix

Copied from the [file suffix config](#). Users can edit the file suffixes here.

##### File suffixes: nadh

Mask	Decay	IRF
n_photons_mask_cyto	n.sdt	NADH_IRF.txt

##### File suffixes: fad

Mask	Decay	IRF
n_photons_mask_cyto	f.sdt	FAD_IRF.txt

Channels ['nadh', 'fad'] share the same mask suffix: n\_photons\_mask\_cyto.tiff

#### 4.1.6 Fluorescence Lifetime Standard

If the selected fit free calibration method is [fluorescence lifetime standard](#), users are asked to specify the fluorescence lifetime standard file path for each channel and the fluorescence lifetime standard's lifetime, which are copied from the data extraction configuration.

#### 4.1.7 Folder Path

Finally, users can specify the folder path that contains all the required input files.

Copy the folder path here	?
/Users/allan/Downloads/40	Press Enter to apply

## 4.2 Metadata Extraction

### 4.2.1 FOV File Paths

The first step is to find the fields of view:

- It finds all the files recursively in the folder path that ends with the 1st file suffix of the 1st channel.
- The prefix of all the matched files is considered to be the field of view name.
  - FOV name = file name - 1st file suffix

Then, it uses the found FOV names to find all other files:

- The file name to search = FOV name + other file suffixes for each file suffix

The only exceptions are the calibration files, because they are not FOV-specific. They will be searched for based on their file suffixes only.

#### 4.2.1.1 Success

H9\_DAY\_40\_5

All files found.

#### 4.2.1.2 Missing

When it cannot find the file based on the file name (FOV name + file suffix for that file type) inside the folder path, it will complain.

H9\_DAY\_40\_11

Missing or duplicate files:

- Missing nadh\_Decay: H9\_DAY\_40\_11n.sdt

#### 4.2.1.3 Duplicate

Because the calibration file is not FOV-specific, it will be searched based on the file suffixes only. If there are multiple files that match the same file suffix, FLIM Playground will be confused about which one to use.

H9\_DAY\_40\_5

✖ Missing or duplicate files:

- Duplicate nadh\_IRF with suffix: \_IRF.txt

Fields of view with are loaded successfully . FOVs with (if any) will not be recorded.

Tip

If the issue persists after renaming the files to match the required prefix, force-quit and relaunch FLIM Playground to clear its internal cache.

#### 4.2.2 Decay Info

If the decay type is [3D/4D decay](#), FLIM Playground will try to infer the duration and the number of time bins per laser pulse interval from the decay file metadata. It will also check for inconsistencies across all decay files.

##### 4.2.2.1 Channel Assignment

If the decay file associated with the channel found by the previous step is a 4D array, then FLIM Playground will try to infer the channel number for that channel. For each channel name, it will list all non-empty channels as potential channels to be assigned. If there is only one, the assignment is automatic. If there are multiple, users need to select the channel intended. It will also check for inconsistencies across all decay files about their dimensions and complain if there are any.

##### 4.2.2.2 Fluorescence Lifetime Standard

If the file path to the [fluorescence lifetime standard](#) is specified, FLIM Playground will try to look for and read the file. It expects a `tiff` or `tif` file that contains a 3D array, with one of them as the time dimension. It will try to match the time dimension with the time dimension of the decay file. If it cannot find the matching time dimension, it will complain.

Error: Reference dye file 'test.tif' not found in folder. Please check the file name.

Error: Cannot find the time axis (256 time bins) in the reference dye file dimensions: (56, 512, 512)

Once the time axis is matched, the file path to the fluorescence lifetime standard, the fluorescence lifetime standard's lifetime, and the time axis will be recorded.

#### 4.2.3 Result Preview and Export

If there is a non-zero number of FOVs with and channel assignment and fluorescence lifetime standard checks (if applicable) are passed, a preview of the metadata data sheet will be shown for users to check.

	image_name	nadh_Mask	nadh_Decay	nadh_IRF	fad_Mask
0	H9_DAY_40_5	/Users/allan/Downloads/40/edited/H9_DAY_40_5n_photons_mask_cyto.tif	/Users/allan/Downloads/40/H9_DAY_40_5n.sdt	/Users/allan/Downloads/40/NADH_IRF.txt	/Users/allan/Downloads/40/edited/H
1	H9_DAY_40_1	/Users/allan/Downloads/40/edited/H9_DAY_40_1n_photons_mask_cyto.tif	/Users/allan/Downloads/40/H9_DAY_40_1n.sdt	/Users/allan/Downloads/40/NADH_IRF.txt	/Users/allan/Downloads/40/edited/H
2	<u>H9_DAY_40_2</u>	<u>/Users/allan/Downloads/40/edited/H9_DAY_40_2n_photons_mask_cyto.tif</u>	<u>/Users/allan/Downloads/40/H9_DAY_40_2n.sdt</u>	<u>/Users/allan/Downloads/40/NADH_IRF.txt</u>	<u>/Users/allan/Downloads/40/edited/H</u>

Export FOV Metadata as CSV

As an example, the following metadata are extracted:

- **image\_name**: The `image_name` column is the name of each FOV, which is specified in the [fov identifier config](#).
- **nadh\_Mask**, **nadh\_Decay**, **nadh\_IRF**, **fad\_Mask**, **fad\_Decay**, **fad\_IRF**: the [file paths](#) of the mask, decay, and IRF files for the nadh and fad channels.
- **nadh\_input\_type**, **nadh\_imaging\_modality**, **fad\_input\_type**, **fad\_imaging\_modality**: the [decay type](#) for each channel. If they share the same imaging modality, the input types should be the same. The only imaging modality supported by FLIM Playground is FLIM, therefore the `input_type` for all channels should be from one of the [decay types](#).
- **nadh\_Lifetime fit free**, **nadh\_Intensity morphology**, **nadh\_Lifetime fit**, **fad\_Intensity morphology**, **fad\_Intensity texture**, **fad\_Lifetime fit**: [feature extractors](#) for each channel.
- **nadh\_channel**, **duration**, **fad\_channel**, **time\_bins**, **laser\_rate**: the [decay info](#) for each channel and info shared by all channels.

The user can click the `Export FOV Metadata as CSV` button to export the metadata as a CSV file.

# 5 Numerical Feature Extraction

In this step, FLIM Playground extracts *single-cell* numerical features from the raw data in a folder. For each channel, the features to be extracted are specified in the [user-selected feature extractors](#). The available feature extractors are:

- [Lifetime fit](#): fit exponential models to decay curves to estimate lifetimes and their fractional contributions by minimizing an objective function.
- [Lifetime fit free](#): transform decay curves to phasor space using Fourier transform to obtain phasor coordinates.
- [Intensity Morphology](#): morphological features of single-cell ROI masks
- [Intensity Texture](#): texture features of single-cell ROI intensity images

## 5.1 Input

A CSV file extracted in the [fov metadata extraction](#) step that contains the metadata of the FOVs including:

- [required file paths](#)
- [decay type](#)
- [decay info](#)
- [selected feature extractors](#) for each channel
- [fit free calibration method](#) (if fit free calibration is required)

Users can either upload a previously extracted metadata file (left), or use the cached metadata file just extracted in the [fov metadata extraction](#) step (right).

Upload the field of view metadata csv

?

Browse files

Drag and drop file here  
Limit 200MB per file • CSV

Using the latest extracted metadata file:  
[/Users/allan/Downloads/40/fov\\_metadata\\_20250824\\_140117.csv](#). Refresh the page to use a different file.

For [Lifetime fit](#) and [Lifetime fit free](#) feature extractors, the first step is always [calibration](#).

## 5.2 Calibration

Calibration in FLIM is essential because raw decays are convolved with the instrument response function (IRF)—the timing profile of the detection system’s response to an ultrashort light pulse that broadens and shifts the measured decay—so without correcting for these, fitted lifetimes and phasor positions are biased and not comparable across days, samples, or instruments.

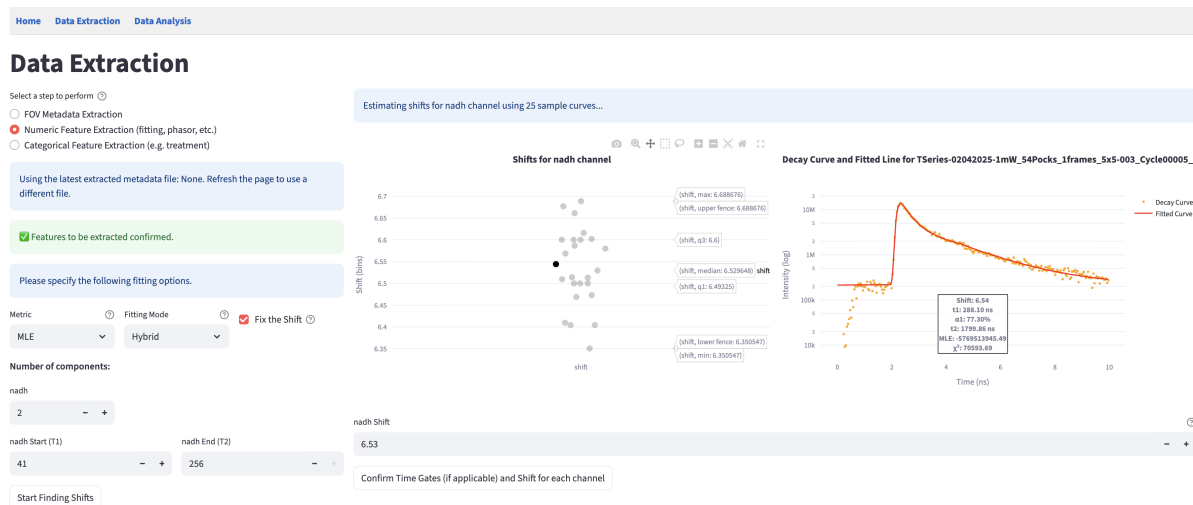
### 5.2.1 Fit Calibration

During different experiments, the IRF may shift differently with respect to the measured decay. Before applying deconvolution fitting, the IRF shift needs to be estimated for each channel if the Lifetime fit feature extractor is selected for the channel.

It is performed in two steps:

1. Gather the high signal-to-noise ratio (SNR) decay curves
2. For each curve, perform deconvolution fitting and set the shift value as the free parameter to be optimized/fitted.

The distribution of the shift values is displayed as an interactive scatter plot. When clicking on a point, the corresponding decay curve with the fitted curve and the key statistics are displayed for diagnostics. Based on the distribution or using certain prior knowledge, users can specify the shift value applied to all fields of view (if Fix the Shift is selected, see fitting options), or use the fov-specific shift value found by the fitting (if Fix the Shift is deselected).



### 5.2.1.1 Gather High SNR Decay Curves

The high SNR decay curves are constructed automatically based on the [decay type](#). If it is in [2D format](#), a total of the brightest 30 curves that are below 100000 photons are selected and evenly distributed across all fields of view (one CSV file is considered as one FOV). If it is in [3D/4D format](#), the fields of view are images, and one curve is constructed for each image that includes all the non-zero pixels within the [ROI mask](#).

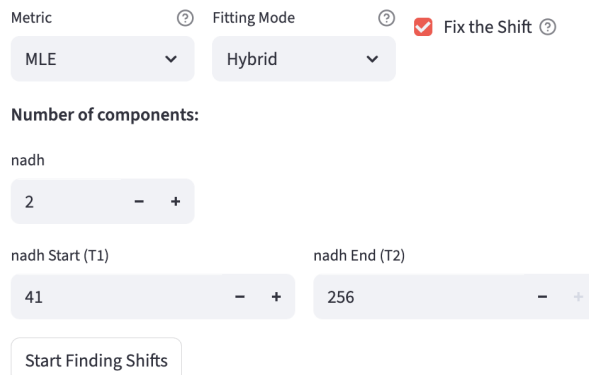
### 5.2.1.2 Reconvolution Fitting

Fitting is essentially an optimization problem: it minimizes the difference between the fitted curve and the measured curve. The fitted curve is modeled by an  $n^{th}$  component exponential function convolved with the shifted IRF, and the difference is modeled as an objective metric. Therefore, users are provided with controls over two parts of the fitting process through the fitting options panel:

1. How to construct the objective metric
2. How to perform the optimization

Implementation-wise, FLIM Playground uses the `lmfit` package that takes generic objectives to perform the optimization process that is flexible enough to handle the reconvolution fitting.

If at least one channel has `Lifetime fit` feature extractor selected, the fitting options panel is displayed.



Let's break down the fitting options one by one.

### 5.2.1.2.1 Number of Components

The number  $n$  in the  $n^{th}$  component exponential function:

$$I(\mathbf{t}) = \sum_{i=1}^n A_i e^{-\mathbf{t}/\tau_i}, \quad \tau_i > 0$$

Therefore,  $n$  determines the parameters to be fitted: the amplitudes  $A_i$  and the lifetimes  $\tau_i$ .  $\mathbf{t}$  is the time axis of the decay curve calculated by the duration and time bins from the [decay info](#):

$$\mathbf{t} = [0, \Delta t, 2\Delta t, \dots, (N-1)\Delta t], \quad \text{where } \Delta t = \frac{T}{N}.$$

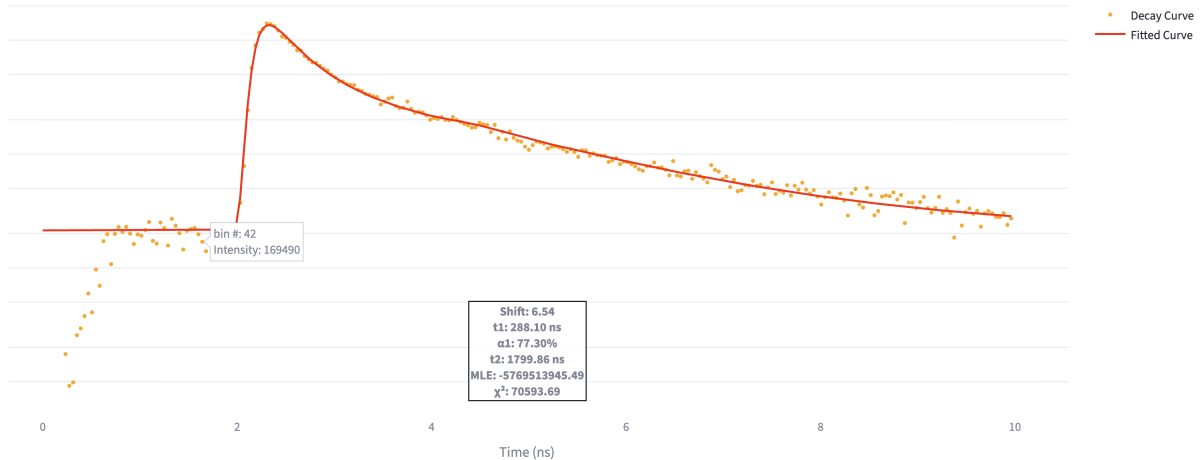
It supports  $n = 1, 2, 3$  components.

Additionally, the constant offset parameter ( $Z$ ) that represents a time-independent background (i.e., room light) is also optimized<sup>11</sup>.

$$\hat{y}(t) = [I(t) \otimes \text{IRF}(t)] + Z$$

### 5.2.1.2.2 Time Gates

Due to the dead time of the system certain time bins from the head and/or tail of the decay curve are not reliable. The T1 (head) and T2 (tail) gates are used to select the time range of the decay curve to be fitted. Users can inspect the decay curve by clicking the full-screen mode button of the plot on the right of the shift result. Hover-based interaction is implemented so users can see the time bin numbers to have a better sense of the time range.



nadh Start (T1)	nadh End (T2)				
41	-	+	256	-	+

Only the time bins within the T1 and T2 gates are used to calculate the cost metric.

#### 5.2.1.2.3 Metric



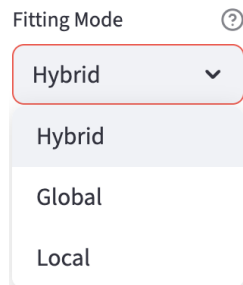
Maximum Likelihood Estimation (MLE) estimates the parameters by maximizing the likelihood function, which is the probability of the measured data given the model parameters. A mathematically convenient way to do this is to minimize the negative log-likelihood function:

$$\text{NLL}(\theta; t_s, t_e) = - \sum_{n=0}^{N-1} \mathbf{1}_{[t_s, t_e]}(t_n) [y_n \log \hat{y}(t_n) - \hat{y}(t_n)].$$

$\mathbf{1}_{[t_s, t_e]}$  is the indicator function that is 1 if  $t_n$  is within the time gates  $[t_s, t_e]$ , and 0 otherwise.  $y_n$  is the observed count at time  $t_n$ , and  $\hat{y}(t_n)$  is the model prediction at  $t_n$ .

Least Squares (LS) estimates the parameters by minimizing the sum of the squared differences between the measured curve and the model prediction.

#### 5.2.1.2.4 Fitting Mode



After constructing the objective metric, the fitting mode is used to determine the optimization algorithm. It is a trade-off between the speed and the effort to avoid local minima.

- Global: It uses the `differential evolution` algorithm, a derivative-free, population-based but slow global optimizer.
- Local: It uses the `leastsq` (least squares with Levenberg-Marquardt) algorithm if the chosen metric is LS, otherwise it uses the `nelder` (Nelder-Mead) algorithm.
- Hybrid: The most time-consuming option but combines the best of both worlds. It uses the `differential evolution` to find a good initial guess to all the parameters, and then uses the Local to drill in.

### 5.2.2 Fit Free Calibration

Users get to choose between the following two methods to calibrate the IRF shift in the [configuration step](#).

The first step of the calibration is to **subtract an offset value and clip each decay curve to 0 if the time bin is negative**. The offset  $C$  is estimated as the mean signal over the final 10% of time bins in each decay ('tail').

Then, depending on the chosen [calibration method](#), the following two methods are available:

#### 5.2.2.1 Shift IRF

It is performed similarly to the [fit calibration](#) steps, but in the second step, the shift is not optimized (fitted). It shares the same interface as the [fit calibration](#) step, where a scatter plot of the shift values is displayed for each channel, only that the plot is not interactive to show the fit. Instead, it outputs the shift that maximizes the cross-correlation between the IRF and each decay. If Fix the Shift is selected, users can specify the shift value that will be applied to all fields of view. Otherwise, the shift value is chosen to be the one that maximizes the cross-correlation between the IRF and the decay.

##### Note

If both the Lifetime fit and Lifetime fit free feature extractors are selected for this channel, the optimized IRF shift derived from the [fit calibration](#) step is reused for the Lifetime fit free feature extractor.

The signals of the shifted IRF are deconvolved from the offset-subtracted decay curves using `phasor.phasor_divide` from the phasorpy package.

### 5.2.2.2 Fluorescence lifetime standard

Because the fluorescence lifetime standard is measured in the same system as the decay curves and we know its lifetime, there is no need to account for the IRF shift and the offset. Therefore, after subtracting the offset, phasorpy's `lifetime.phasor_calibrate` function is used to calibrate all the decay curves behind the scenes when calculating the phasor coordinates.

### 5.2.3 Confirm Calibration

Once users are satisfied with the calibration settings (fitting options and shift values), they can click the `Confirm Calibration` button at the bottom of the page.

Confirm Time Gates (if applicable) and Shift for each channel

Once confirmed, the left panel prompts users to either apply the calibration settings to the downstream extractors or calibrate again. Users can also save the updated settings (fitting options and shift values) to the metadata file.

Confirm and Start Analysis

Go back and find shift

Download updated metadata

## 5.3 Feature Extraction

If the selected feature extractors do not require IRF shift calibration, or users have finished the calibration, they can proceed to the feature extraction step by clicking the `Confirm and Start Analysis` button.

Similar to the `fov metadata extraction` step, FLIM Playground extracts the features and displays the extraction status for each field of view.

The screenshot shows a user interface for feature extraction. At the top left, there's a section for selecting steps: 'FOV Metadata Extraction' (radio button), 'Numeric Feature Extraction (fitting, phasor, etc.)' (radio button, selected), and 'Categorical Feature Extraction (e.g. treatment)' (radio button). Below this is a file upload area for 'fov\_metadata.csv'. A file named 'fov\_metadata\_20250824\_162655.csv' (10.5KB) is shown with a 'Success!' message. There's also a note about 'Features to be extracted confirmed.' and buttons for 'Confirm and Start Analysis', 'Go back and find shift', and 'Download updated metadata'.

The main area consists of a grid of 10 boxes, each representing a different FOV. Each box contains the image name and a 'Success!' message. The last box on the right shows a progress bar and a note: 'Running fov\_extraction(...). g/fit free for nadh: for 23 cells...'.

A progress bar is rendered to show the feature extraction progress for each FOV (bottom right).

Warnings are displayed if cells have NaN values for some features. They are *not* excluded from the final CSV file.

## 5.4 Save Results

Once the feature extraction is finished, users can save the results to a CSV file by clicking the Download button at the bottom of the page.

Field of view features with  are extracted successfully ! FOVs with  (if any) are excluded. The first few rows of the features are shown below.							
cell_id	nadh_centroid_x	nadh_centroid_y	Intensity_morphology_nadh: area	Intensity_morphology_nadh: perimeter	Intensity_morphology_nadh: solidity	Intensity_morphology_nadh: eccentricity	Intensity_morphology_nadh: orientation
H9_DAY_40_5_3	214.1314	10.1134	388	116.5685	0.7854	0.7	
H9_DAY_40_5_4	242.4038	16.2271	634	118.4914	0.8476	0.6	
H9_DAY_40_5_6	155.2827	24.2655	2490	320.8356	0.8156	0.6	
H9_DAY_40_5_7	49.0648	25.8541	1172	178.9949	0.8919	0.7	
H9_DAY_40_5_8	77.3881	43.406	894	169.3381	0.8976	0	

Download single cell features as CSV

# 6 Lifetime Features (Fit)

Biological samples often contain mixtures of fluorescent species, each with its own characteristic lifetime. By modeling the decay as a sum of exponentials, the analysis can separate and quantify these different contributions. For example, NADH in cells exists in both free (short lifetime, ~0.4 ns) and protein-bound (longer lifetime, ~2–3 ns) states; by fitting the fluorescence decay with a bi-exponential model, one can estimate the fraction of each state, which provides direct insight into cellular metabolism and energy production pathways.

FLIM Playground extracts cell-level lifetime fitting features in this feature extractor, including the fraction of each lifetime component ( $\alpha_i$ ), their lifetimes ( $\tau_i$ ), and mean lifetime ( $\tau_{mean}$ ).

$$I(t) = \sum_{i=1}^n A_i e^{-t/\tau_i}$$

## 6.1 Fitting

If the decays have not been pre-fitted, FLIM Playground applies the confirmed fitting options ([number of components](#), [time gates](#), [metric](#), and [fitting mode](#)) to the same [reconvolution fitting process](#), as described in the [irf shift calibration step](#), with the only difference being that the IRF shift is no longer a free variable to be optimized. To recap, the reconvolution fitting minimizes the difference between the measured curve and the fitted curve modeled by a multi-exponential model convolved with the IRF, quantified by the cost metric.

Otherwise, FLIM Playground uses the [pre-fitted values](#) to calculate the fitting features.

In the final dataset, fitting features are prefixed by the combination of the feature extractor name (i.e. `Lifetime fit`) and the channel name, allowing [Data Analysis](#) to group the features. For example, `Lifetime fit_nadh`: `a1` means the fraction of the first lifetime component for the NAD(P)H channel.

### 6.1.1 Preprocessing

It sums up all the pixel decays belonging to the same cell ROI labeled by the ROI [mask](#) as one decay curve<sup>9</sup>. The ROI summing reduces variability and bias and allows for short integration times at acquisition in exchange for sub-cellular resolution (i.e., pixel-level). Users do not need to specify the

bin factor (each pixel sums up the surrounding pixels' decay curves) to account for insufficient photon counts. Also, this summing lets a single-threaded CPU application finish in a reasonable amount of time.

It also shifts the IRF using the shift values from the [IRF shift calibration](#) step. To shift an IRF, FLIM Playground upsamples the IRF 10 times using linear interpolation to fill the gaps. Then it shifts the IRF by the shift values  $\times 10$  and downsamples the IRF back to the original size.

### 6.1.2 Fraction of components

In addition to the absolute amplitudes of each component directly from the fitting result, FLIM Playground calculates the fraction of each component,  $\alpha_i$ , as the amplitude of the component divided by the sum of all amplitudes. This normalization allows lifetime to be independent of the absolute intensity of the signal.

### 6.1.3 Mean lifetime

The mean lifetime,  $\tau_{mean}$ , is calculated as the weighted average of the lifetimes of all components, where the weights are the fractions of each component.

$$\tau_{mean} = \sum_{i=1}^n \alpha_i \tau_i$$

## 6.2 Pixel-prefitted

Currently, FLIM Playground supports pixel-prefitted lifetime features from [SPCImage](#). The pixel-level lifetime fitting features are expected to be stored in [2D arrays](#) in spatial dimensions, with each row and column having the value of a lifetime feature outputted from SPCImage. SPCImage assigns 0 for pixels that are not fitted (e.g. thresholded out). To avoid them biasing the results, FLIM Playground uses `np.ma.masked_array` to create a masked array and disregards them when calculating the averages using `np.ma.average`.

Then it uses the ROI [mask](#) to calculate the cell-level lifetime features by averaging the pixel-level features within each ROI.

# 7 Lifetime Features (Fit-Free)

Phasor analysis provides a fast, model-free view of lifetimes by transforming fluorescence decays into phasor coordinates through a Fourier transform. This approach is considered “fit-free” because it does not require iterative curve fitting or predefined decay models—each decay is mapped directly to a point on the [phasor plot](#). It has the nice property that mono-exponential decays fall on the universal semicircle while mixtures lie inside as linear combinations, so clusters reveal distinct species and their fractional contributions. FLIM Playground currently extracts cell-level phasor features: `g`, `s`, `Tau_phase`, `Tau_m`, `g_2nd` (2nd harmonic), and `s_2nd`.

Fit-free features in the final dataset are prefixed by the combination of the feature extractor name (i.e. `Lifetime fit free`) and the channel name, allowing [Data Analysis](#) to group the features. For example, `Lifetime fit free_nadh`: `G` means the `g` coordinate for this cell in the NAD(P)H channel.

## 7.1 Preprocessing

It sums up all the pixel decays belonging to the same cell ROI labeled by the ROI [mask](#) as one decay curve<sup>9</sup>. The phasor features are calculated from the summed decay curve.

If the chosen [calibration method](#) is IRF shift calibration, FLIM Playground shifts the IRF using the shift values and subtracts the estimated offset as described in the IRF shift calibration step.

## 7.2 Phasor features

### 7.2.1 Step 1: raw phasor coordinates

The reference implementation from [phasorpy](#) assumes the decay curve comes from one period and time bins are evenly spaced (untruncated). So, their calculation is frequency ( $f$ ) and time ( $T$ ) independent. However, there are cases when the decay curve is truncated due to the deadtime of the system. For example, when the laser frequency is  $f = 0.08$  GHz, the theoretical period is  $T = 12.5$  ns, but in practice the time window is truncated to  $T = 10$  ns. FLIM Playground applies a custom implementation to the truncated decay curve. (The `sample_phase` option can be used to account for the truncation but it does not handle the second or higher harmonics)

### 7.2.1.1 Untruncated decay curve

It uses the `phasor_from_signal` function from the `phasorpy` package to calculate the uncalibrated phasor coordinates `g_raw` and `s_raw`, `g_raw_2nd` and `s_raw_2nd`, the first and second harmonics' real and imaginary parts, from the cell decay curve.

Either `g_irf`, `s_irf`, `g_irf_2nd`, and `s_irf_2nd` from the IRF, or `ref_mean`, `ref_real`, `ref_imag` from the fluorescence lifetime standard, depending on the chosen `calibration method`, are calculated using the same function.

Mathematically, the raw phasor coordinates are calculated as (copied from the `phasorpy` documentation):

$$F_{DC} = \frac{1}{K} \sum_{k=0}^{K-1} F_k, \quad g_{\text{raw}} = \frac{1}{F_{DC}} \frac{1}{K} \sum_{k=0}^{K-1} F_k \cos\left(2\pi h \frac{k}{K}\right), \quad s_{\text{raw}} = \frac{1}{F_{DC}} \frac{1}{K} \sum_{k=0}^{K-1} F_k \sin\left(2\pi h \frac{k}{K}\right)$$

where  $K$  is the number of time bins,  $F_k$  is the decay curve, and  $h$  is the harmonic number.

### 7.2.1.2 Truncated decay curve

The phasor coordinates of the IRF and the decay curves of both harmonics are calculated using the time axis  $t_k = k (T/K)$  ( $k = 0, \dots, K - 1$ ) and the angular frequency  $\omega = 2\pi h f$ :

$$g_{\text{raw}} = \frac{\sum_{k=0}^{K-1} F_k \cos(\omega t_k)}{\sum_{k=0}^{K-1} F_k}, \quad s_{\text{raw}} = \frac{\sum_{k=0}^{K-1} F_k \sin(\omega t_k)}{\sum_{k=0}^{K-1} F_k}$$

This formula is equivalent to the untruncated formula when  $fT = 1$ .

For the fluorescence lifetime standard, the phasor coordinates are calculated using `phasorpy`'s `phasor_from_signal` and the `sample_phase` option is set to  $\omega t_k$ .

## 7.2.2 Step 2: calibrated phasor coordinates

To get the calibrated phasor coordinates `g`, `s`, `g_2nd`, and `s_2nd`, FLIM Playground uses either

```
from phasorpy import phasor
g, s = phasor.phasor_divide(g_raw, s_raw, g_irf, s_irf)
g_2nd, s_2nd = phasor.phasor_divide(g_raw_2nd, s_raw_2nd, g_irf_2nd, s_irf_2nd)
```

or

```

from phasorpy import lifetime
g, s = lifetime.phasor_calibrate(g_raw, s_raw, ref_mean, ref_real, ref_imag, frequency=laser_
g_2nd, s_2nd = lifetime.phasor_calibrate(g_raw_2nd, s_raw_2nd, ref_mean, ref_real, ref_imag,

```

The `laser_rate` is specified during the [fov metadata extraction](#) and the [configuration interface](#).

### 7.2.3 Step 3: phasor-derived lifetime

`Tau_phase` and `Tau_m` are calculated using the first harmonic's phasor coordinates `g` and `s`.

```

import numpy as np
w = 2 * np.pi * laser_rate
phi = np.arctan2(s, g)
m = np.sqrt(g**2 + s**2)
tau_phase = 1/w * np.tan(phi)
tau_m = 1/w * np.sqrt(1/m**2 - 1)

```

$$\phi = \arctan2(s, g), m = \sqrt{g^2 + s^2}, \tau_\phi = \frac{\tan \phi}{\omega}, \tau_m = \frac{1}{\omega} \sqrt{\frac{1}{m^2} - 1}$$

## 8 Morphological Features

FLIM Playground extracts single-cell morphological features based on the cell region of interest (ROI) mask of each *channel*. By inputting the ROI mask to `skimage.measure.regionprops`, the following features are extracted:

- `area`: the number of pixels in the ROI.
- `perimeter`: length of the ROI boundary in pixels.
- `solidity`: Solidity =  $\frac{\# \text{ ROI pixels}}{\# \text{ convex hull pixels}}$ , where the convex hull is the smallest convex polygon that contains the ROI. A value close to 1 means the shape has few concavities (more solid), while lower values indicate more irregular boundaries.
- `eccentricity`: equals the eccentricity of the ellipse that has the same second moments as the region. It is given by the ratio of the focal distance (the distance between the foci) to the length of the major axis:  $e = \frac{c}{a}$ ,  $e \in [0, 1]$ , where  $a$  is the semi-major axis length,  $b$  is the semi-minor axis length, and  $c = \sqrt{a^2 - b^2}$  is the focal distance. When  $e = 0$ , the ellipse reduces to a circle. It quantifies how elongated the ROI is, with 0 being a perfect circle and 1 being a line.
- `major_axis_length`: the length of the major axis of the ellipse that has the same second moments as the region:  $2a$ . It represents the longest dimension of the ellipse that approximates the region's shape, essentially describing its maximum elongation.
- `minor_axis_length`: the length of the minor axis of the ellipse that has the same second moments as the region:  $2b$ . It represents the shortest dimension of that ellipse, capturing the region's minimum elongation.
- `circularity`: describes how close the ROI is to a perfect circle. It is defined as circularity =  $\frac{4\pi A_{\text{ROI}}}{P_{\text{ROI}}^2}$ ,  $\text{circularity} \in (0, 1]$ , where  $A_{\text{ROI}}$  is the area of the region and  $P_{\text{ROI}}$  is its perimeter. A value of 1 corresponds to a perfect circle.

Feature names in the final dataset are prefixed by the combination of the feature extractor name (i.e. `Intensity morphology`) and the channel name, allowing [Data Analysis](#) to group the features. For example, `Intensity morphology_nadh: area` means the area of each cell in the NAD(P)H channel.

# 9 Textural Features

FLIM Playground extracts single-cell texture features based on the intensity image and the ROI mask from each acquisition *channel*. For an example application, refer to this [paper](#) that used CellProfiler to extract the texture features.

The first step is to create a single-cell intensity image for each cell based on the ROI mask and the intensity image. Then a list of texture features (the list may be extended in the future) is calculated:

## 9.1 Granularity

$Granularity_n$  is the percentage of intensity contributed by bright objects of  $n$  pixels in radius. It is computed by performing a morphological opening with a disk of radius  $n$  on the cell intensity image and measuring the intensity lost as a percentage of the total intensity of the original cell ROI.

$n = 1, 3, 5, 7, 9$  are computed.

Semantically, higher granularity at smaller scales implies higher fragmentation.

## 9.2 Radial Distribution

To calculate the radial distribution, each cell intensity image is partitioned into 4 concentric rings, and the intensity fraction over the total intensity for each ring is calculated. For example, a higher fraction in ring 1 implies that signals are more concentrated at the center of the cell, whereas a higher fraction in ring 4 implies that signals are more concentrated at the cell edge.

## 9.3 Mass Displacement

Mass displacement is the Euclidean distance (in pixels) between the intensity-weighted centroid of the cell ROI and the geometric centroid of the same ROI.

## 9.4 Intensity Sum

Intensity sum is defined as the sum of pixel intensities within the cell intensity image.

In the final dataset, feature names are prefixed by the combination of the feature extractor name (i.e. `Intensity texture`) and the channel name, allowing [Data Analysis](#) to group the features. For example, `Intensity texture_nadh: granularity_1` means the percentage of intensity removed when bright objects of 1-pixel diameter are removed in the NAD(P)H channel.

 Visual explanation

[This post](#) provides a visual explanation of and code to calculate some of the texture features above.

# 10 Categorical Feature Extraction

Based on the datasets extracted in the [numerical feature extraction](#) step, FLIM Playground:

- merges the datasets into a single dataset
- assigns categorical features to each cell in the merged dataset

The screenshot shows the 'Data Extraction' step in the FLIM Playground interface. At the top, there are three tabs: Home, Data Extraction (which is selected), and Data Analysis. Below the tabs, a section titled 'Select a step to perform' has three options: FOV Metadata Extraction, Numeric Feature Extraction (fitting, phasor, etc.), and Categorical Feature Extraction (e.g. treatment). The third option is selected. A message box says 'Merging datasets...' and 'Finished combining all datasets and got 445 cells!'. Below this, a folder path is specified: '/Users/allan/Downloads/40'. A note says 'Now your task is to map the categories to (combination of) slots.' An example is given: 'Example fov\_name: H9\_DAY\_40\_5 has slots: [H9, 'DAY', '40', '5]'. A 'Choose Categorical features to populate' dropdown contains 'day' (selected) and '40'. A preview table shows the mapping between image names and categories:

image_name	day
0   H9_DAY_40_5	DAY_40
33   H9_DAY_40_1	DAY_40
83   H9_DAY_40_2	DAY_40
130   H9_DAY_40_6	DAY_40
180   H9_DAY_40_10	DAY_40

A 'Confirm category mapping & export the combined dataset' button is at the bottom.

## 10.1 Merging Datasets

It scans the specified folder path and finds all CSV files. It uses the [cell identifier column](#) to check for duplicates both within each dataset and across all datasets.

### ! Important

The FOV identifier uniquely identifies a field of view and is assumed to contain all information necessary to extract the categorical features (e.g., treatment, time point, etc.), by including segments separated by a certain delimiter (e.g., \_). Categorical features are assigned from individual segments or their combinations. Example: The FOV Panc1\_Cyanide\_dish1

denotes that it comes from the Panc1 cell line, under the Cyanide treatment, in the 1st dish.

Therefore, it checks if all field-of-view names include the same number of segments to determine the eligibility of [categorical feature assignment](#).

Field of View Name Delimiter ?

—

The image\_name column in /Users/allan/Downloads/DATA/donor6.csv has different number of parts. For example, check fov\_name: HC220113\_1n.

The image\_name column in /Users/allan/Downloads/DATA/donor5.csv has different number of parts. For example, check fov\_name: HC130319\_1n.

Found 7 available csv files ready to be assigned categories 😊:

```
▼ [ 0 : "/Users/allan/Downloads/DATA/donor8.csv" 1 : "/Users/allan/Downloads/DATA/donor9.csv" 2 : "/Users/allan/Downloads/DATA/donor1.csv" 3 : "/Users/allan/Downloads/DATA/donor2.csv" 4 : "/Users/allan/Downloads/DATA/donor3.csv" 5 : "/Users/allan/Downloads/DATA/donor7.csv" 6 : "/Users/allan/Downloads/DATA/donor4.csv" ]
```

It merges the datasets that pass the checks by finding common columns and vertically concatenating them.

Merging datasets...

Merging dataset 2 (/Users/allan/Downloads/DATA/donor9.csv) into the combined dataset.

Merging dataset 3 (/Users/allan/Downloads/DATA/donor1.csv) into the combined dataset.

Merging dataset 4 (/Users/allan/Downloads/DATA/donor2.csv) into the combined dataset.

Merging dataset 5 (/Users/allan/Downloads/DATA/donor3.csv) into the combined dataset.

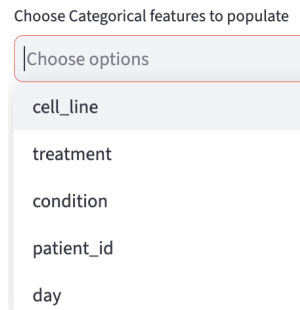
Merging dataset 6 (/Users/allan/Downloads/DATA/donor7.csv) into the combined dataset.

Merging dataset 7 (/Users/allan/Downloads/DATA/donor4.csv) into the combined dataset.

Finished combining all datasets and got 3037 cells!

## 10.2 Assigning Categorical Features

Available categorical features to be assigned are drawn from the [user-specified configuration](#).



Based on the assumptions, it breaks down the FOV identifier into individual fields and facilitates the assignment.

Now your task is to map the categories to (combination of) slots.

Example fov\_name: H9\_DAY\_40\_5 has slots: ['H9', 'DAY', '40', '5']

A live preview of the assignment is displayed.

A screenshot of a live preview interface. At the top, there's a header "Choose Categorical features to populate" with buttons for "cell\_line" and "day". Below this, two sections show assigned slots: "Slots for cell\_line" containing "H9" and "Slots for day" containing "DAY" and "40". Underneath, a table titled "Preview of category mapping:" shows a one-to-one mapping between image names and their assigned slots. The table has columns for "image\_name", "cell\_line", and "day". Rows include: H9\_DAY\_40\_5 (cell\_line: H9, day: DAY\_40), H9\_DAY\_40\_1 (cell\_line: H9, day: DAY\_40), H9\_DAY\_40\_2 (cell\_line: H9, day: DAY\_40), H9\_DAY\_40\_6 (cell\_line: H9, day: DAY\_40), and H9\_DAY\_40\_10 (cell\_line: H9, day: DAY\_40). At the bottom is a button "Confirm category mapping &amp; export the combined dataset".

And the merged dataset with categorical features is available for download.

## **Part II**

# **Data Analysis**

# 11 Overview

We are not done yet 😱! FLIM Playground also provides a suite of methods to analyze and visualize the extracted data (either using the [Data Extraction](#) section or in users' own way). They are designed to be interactive and frictionless ⚡ so that users can perform hassle-free exploration of their data with, hopefully, fun 😎. Inspired by the same data [categorization](#) in [Data Extraction](#), the Data Analysis section designs an [architectural blueprint](#) that all in-house analysis methods build on. It is modularized so that new methods and features can be added in the future easily.

## 11.1 General Workflow

- Select [numerical feature\(s\)](#) on the left.
- [Subset the data](#) to find data of interest on the top. *All the later interactions are based on this filtered dataset.*
- Use [visual channels widgets](#) to look at the data in different ways.
- Change the plot [style](#).
- [Hover](#) to find data points of interest.
- Interact with method-specific widgets that typically are associated with a modeling method.

The visualization and analysis results are updated in real time to reflect users' interactions with any of the above widgets.

## 11.2 Methods

- Depending on the number of numerical features in the analysis, the methods are categorized into 3 groups:
  - Univariate Analysis
    - \* [Feature Comparison](#)
    - \* [Feature Histogram](#)
    - \* [Field of View Comparison](#)
  - Bivariate Analysis
    - \* [2D Feature Distribution](#)
    - \* [Phasor Analysis](#)

- Multivariate Analysis
  - \* Dimension Reduction
  - \* Classification

All the methods [share a set of interactive widgets](#). They also have their own method-specific widgets, the descriptions of which are provided in the corresponding method pages.

## Data Analysis

**Univariate**

**Bivariate**

**Multivariate**

## Methods

**Feature Comparison**

**Feature Histogram**

**Image Comparison**

## Data Analysis

**Univariate**

**Bivariate**

**Multivariate**

## Methods

**2D Feature Distribution**

**Phasor Plot**

## Data Analysis

**Univariate**

**Bivariate**

**Multivariate**

## Methods

**Dimension Reduction**

**Classification**

## 11.3 Input

Users can upload the dataset output from the [Data Extraction](#) directly or *their own datasets* in **CSV** (Comma Separated Values) format after finishing the [interactive configuration](#) setup.



### Use Dataset from Data Extraction

Upload the CSV file obtained from [Data Extraction](#) directly.



### Use Dataset from Data Extraction

**Use the right panel to configure before loading your data ==>**

#### 11.3.1 Requirements

##### ! Important

In either case, following the [categorization](#), the dataset should have:

- a column that uniquely identifies each row
- an (optional) field of view identifier column
- a set of numerical features
- zero or more categorical features (e.g. treatment, day, patient id, etc.)

The unique row id and field of view id are used to help users identify the row (e.g. a single cell) and field of view (e.g. a single image) of data of interest through the built-in [hover-based interaction](#). Numerical and categorical features are used to render [shared widgets](#). Internally, FLIM Playground will check whether the dataset fulfills the requirements and output meaningful warning or error messages.

##### 💡 Tip

If you cannot find a certain **numerical feature** under any numerical widgets including **Uncategorized Features**, please inspect the dataset (e.g., using Excel filters) and look for non-numeric values in that column. One non-numeric value (e.g., “–”) will prevent it from being recognized.

FLIM Playground searches for **categorical features** in the uploaded dataset based on the [user-specified configuration](#) if the dataset is not extracted by [Data Extraction](#). Otherwise, it searches for categorical features specified in the [Data Extraction configuration](#).

### 11.3.2 Warning Messages

- `Empty columns`: will be dropped.
- `Duplicated columns`: Only the first occurrence of the duplicated column will be kept. Other occurrences will be dropped.
- `Duplicated rows based on the unique row id`: Only the first occurrence of the duplicated row will be kept. Other occurrences will be dropped.
- `Columns with NaN values`: won't be dropped, just a warning message.
  - The analysis will be performed on the rows that are not NaN in the [selected numerical features](#).

### 11.3.3 Error Messages

- Missing a column that uniquely identifies each row
- Cannot identify any numerical feature column
  - It uses `pd.api.types.is_numeric_dtype` to check if a column is numerical.

 Note

Warning messages will not prevent the analysis but error messages will.

## 11.4 Shared Interactive Widgets

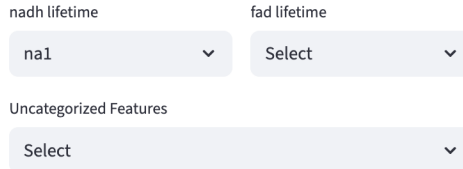
A list of shared widgets is provided to support the [general workflow](#) in analysis.

### 11.4.1 Numerical Selection Widgets

The Data Extraction recognizes numerical features using `pd.api.types.is_numeric_dtype` internally. Then it groups numerical features based on the [feature extractor type](#) and the channel it belongs to. For user-provided datasets, the numerical features are grouped based on the [user-specified configuration](#). Ungrouped numerical features are put into the [Uncategorized Features group](#). One selection widget is rendered for each numerical feature group.

#### 11.4.1.1 Univariate Analysis

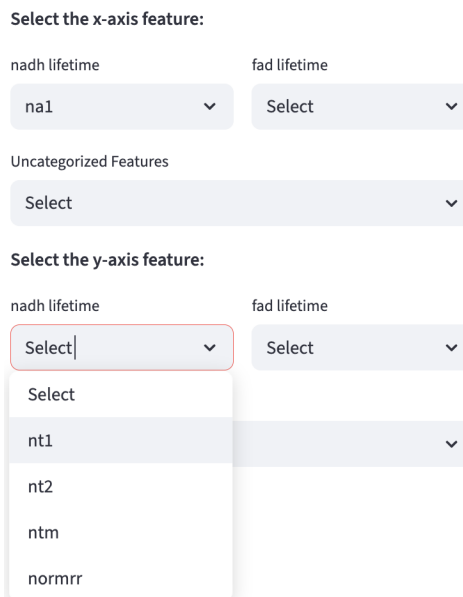
One single-select widget is rendered per group; choosing a feature in any group clears the selections and resets the others to Select.



The screenshot shows three selection widgets. The first two are grouped under 'nadhlifetime' and 'fad lifetime'. The first has an option 'na1' selected. The second has an option 'Select' selected. Below these are two more groups: 'Uncategorized Features' and another unnamed group. Both have a single 'Select' option.

#### 11.4.1.2 Bivariate Analysis

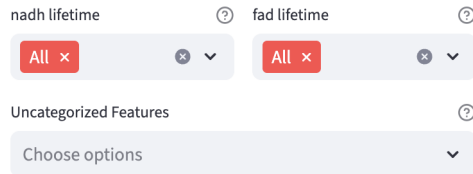
Two sets of selection widgets are rendered to allow maximum flexibility (the two features can be from the same or different feature groups). Each set of selection widgets behaves like the selection widgets in the univariate analysis. The first selected feature will be hidden in the second set of selection widgets.



The screenshot shows two sections. The top section is labeled 'Select the x-axis feature:' and contains the same two groups as the univariate analysis: 'nadhlifetime' (with 'na1' selected) and 'fad lifetime' (with 'Select' selected). The bottom section is labeled 'Select the y-axis feature:' and contains two groups. The left group under 'nadhlifetime' has a dropdown menu open with 'Select' at the top and other options 'nt1', 'nt2', 'ntm', and 'normrr' listed below. The right group under 'fad lifetime' also has a dropdown menu open with 'Select' at the top.

#### 11.4.1.3 Multivariate Analysis

One set of selection widgets, each can select multiple features from a feature group, is rendered. A special value All is introduced so that users can conveniently select all features under the feature group. If users select All, all the other options will be cleared, and vice versa.



## 11.4.2 Filter Widgets

### 11.4.2.1 Categorical Filters

Select experiment(s)	Select media(s)	Select cell_line(s)	Select treatment(s)
glytcafao <span style="color: red;">x</span>	All <span style="color: red;">x</span>	MCF7 <span style="color: red;">x</span> HeLa <span style="color: red;">x</span>	All <span style="color: red;">x</span>

For complex datasets that are collected over multiple days, experiments, treatments, etc., it is useful to filter the data to focus on a subset of the data (data of interest). One filter widget is rendered for each categorical feature so that users have the flexibility to filter the data based on combinations of categorical features. All is a special option that include all categories of the selected categorical feature. Once it is selected, all the other options are cleared, and vice versa.

### 11.4.2.2 Numerical Filters

Select Feature 1	Operator	Threshold (555.00 - 2508867.00)	<input checked="" type="checkbox"/> Add another
Intensity texture_nadh: intensity_sum	>	2000.00	- +
Select Feature 2	Operator	Threshold (2138.00 - 2508867.00)	<input checked="" type="checkbox"/> Add another
Intensity texture_nadh: intensity_sum	<=	2000000.00	- +
Select Feature 3	Operator	Threshold (5.00 - 2842.00)	<input checked="" type="checkbox"/> Add another
Intensity morphology_nadh: area	>	20.00	- +
Select Feature 4			
None			

Users can further filter the data based on numerical features. Each row specifies a condition to filter the data:

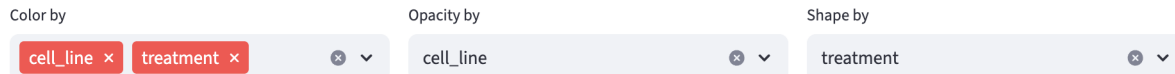
- **Feature:** the numerical feature to filter on
- **Operator:** > or <=
- **Threshold:** the range of values to filter on, dynamically updated based on previous filters, is displayed for reference. The entered value should be within the range.
- **Add Another:** add a new condition in the next row.

After the two sets of filters are applied, the final filtered data is used for visualization and analysis.

### ! Important

Make sure you clearly state all numerical filters used.

## 11.4.3 Visual Channels Widgets



Human vision is wired for rapid, parallel pattern and trend detection; good visual encodings (how to map data to visual elements such as color, shape, opacity, etc.) harness this to surface insights that raw numbers or text obscure<sup>12</sup>. FLIM Playground provides `color`, `opacity`, and `shape` channels for visualizations that support them:

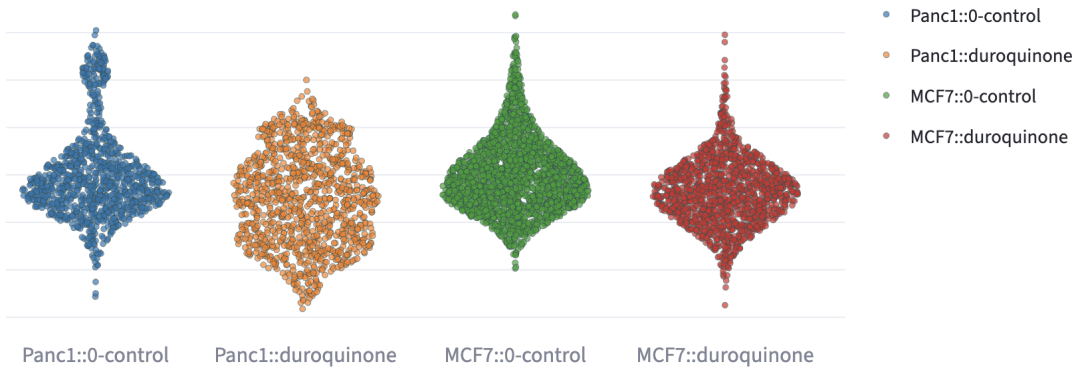
### 11.4.3.1 Color by

Color is supported in all methods except for [Classification](#). In `Color by`, users can select multiple categorical features, and each unique combination of available categories in selected features is assigned a distinct color. Groups created by `Color by` show up in the x-axis (if multiple features, categories from each feature are delimited by `::`).

#### i Note 1: How to order the groups

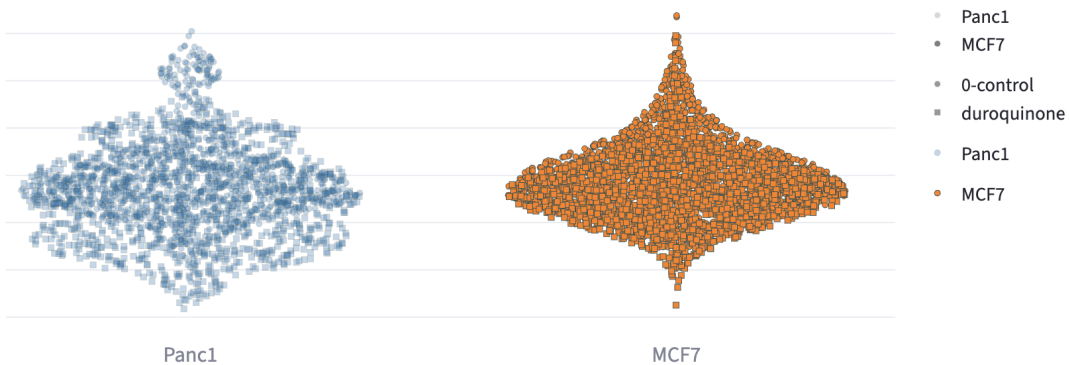
The order of the groups in x-axis and in the legend is determined by:

- the order of the selected features: categories of the first feature appear before those of the second, and so on and so forth. For example, the two treatments of Panc1 are together, because it is sorted first on the cell lines, then on treatments. If you want the treatments to be together, in `Color by` you can select `treatment` then `cell_line`.
- *Numeric-alphabetical sort*: within each feature, the category order is determined by the number inside (e.g. 1 in Panc1) first, then the alphabetical order of the category name. Therefore, although P is after M alphabetically, the 1 is before 7 numerically, making Panc1 appear before MCF7. This may be helpful when you have data from different days or hours. The default string sort will put Day 100 before Day 30, and this is not what we want.



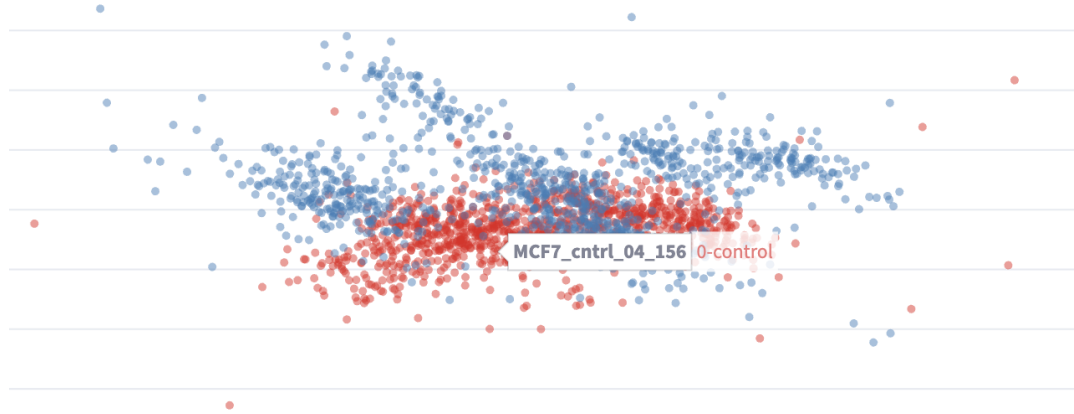
#### 11.4.3.2 Opacity and Shape by

Opacity and shape are supported in all point-based visualizations (e.g. [Feature Comparison](#), [2D Feature Distribution](#), [Phasor Analysis](#), [Dimension Reduction](#)). In **Opacity by** and **Shape by**, users can select one categorical feature, and each unique category is assigned a distinct opacity or shape. The order of the shape and opacity is also sorted [numeric-alphabetically](#).



#### 11.4.4 Unique ID Hover

- In all point-based visualizations, when hovering over a point, the unique id of the point will be shown.



#### 11.4.5 Plotting Configuration Widgets

##### Plot Styling ↻

Point Size	Axis Label Font Size	Legend Font Size	Color Map	?			
5	- +	18	- +	16	- +	tab10	▼

Users can interactively configure the following plot parameters:

- point size (if point-based)
- axis label size
- legend size
- the colormap used to color different groups
  - `colorblind`, `tab10`, `tab20`, `Set1`, `Set2`, `Set3`, `Pastel1`, `Pastel2`, `Accent`, `viridis`, `plasma`, `inferno`, `magma`, `cividis`

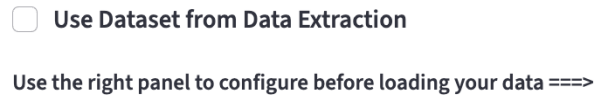
# 12 Configuration

FLIM Playground provides an interactive configuration workflow to help users prepare their datasets that are not extracted by [Data Extraction](#) so that they fulfill the [requirements](#).

The following sections use the [iris dataset](#) as an example: the `flower_id` is added to uniquely identify each row (i.e. a single flower). The other columns are: a categorical feature `species`, and a set of numerical features `sepal_length`, `sepal_width`, `petal_length`, and `petal_width`.

Specifically, the steps are:

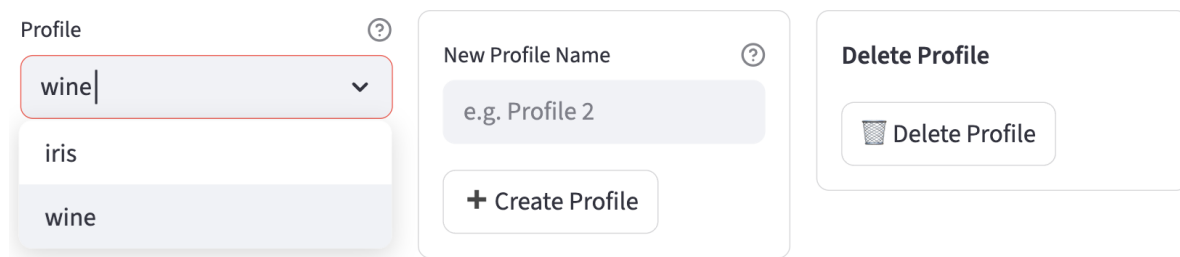
- On the left side of the screen, uncheck `Use Dataset from Data Extraction`.



Then the configuration interface will be displayed.

## 12.1 Analysis Profiles

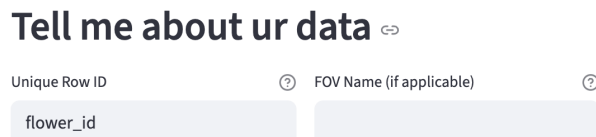
✨ NEW ✨: Users can now create, save, and switch between multiple analysis profiles—each a ready-to-go configuration tailored for different datasets.



- On the left you can select an existing profile to load.
- In the middle you can create a new profile.
- On the right you can delete an existing profile.

Everything described below is pertained to the current profile.

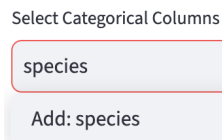
## 12.2 Identifiers Config



- The dataset needs to have a unique identifier column to identify each row. In this case, `flower_id` is the unique identifier column. This dataset does not have a field of view column, so we leave it blank.

## 12.3 Categorical Feature Config

- Add categorical columns you want FLIM Playground to recognize here. Those columns are converted to strings internally.



The default categorical feature list does not include `species`, so we add it manually.

## 12.4 Numerical Feature Config

Paste numerical features (comma, semicolon, or whitespace separated)

```
sepal_length,sepal_width,petal_length,petal_width
```

- Users can copy the numerical features from Excel or any text editor that opens their CSV files. The features can be separated by `,`, `;`, or whitespaces including newlines.
- After hitting the `ctrl/cmd + enter` key, the numerical features are parsed and added to Available Features. Users can add/delete groups and drag features from the Available Features to the groups they've just created. Any ungrouped features are added to a group called Uncategorized Features.

#### 12.4.1 Example

The screenshot shows the FLIM Playground interface with three main sections:

- Select Numerical Features:** A header "Select Numerical Features" with a bar chart icon. Below it is a list of selected features: "sepal\_length x", "sepal\_width x", "petal\_length x", and "petal\_width x".
- Feature Groups Management:** A header "Feature Groups Management". It contains two boxes:
  - Create New Feature Group:** A form with "Group Name" input (placeholder: "e.g. lifetime, morphology, texture") and a "Create Group" button.
  - Delete Feature Group:** A form with "Select group to delete" dropdown (selected: "sepal") and a "Delete Group" button.
- Drag & Drop Feature Assignment:** A header "Drag & Drop Feature Assignment". It shows feature assignments:
  - Available Features:** "petal\_length" (highlighted in red).
  - sepal:** "sepal\_length" (highlighted in red) and "sepal\_width" (highlighted in red).
  - petal:** "petal\_width" (highlighted in red).

Below these are "Save Configuration" and "Reset Configuration" buttons.

#### 12.5 Save

- After users finish the configuration and click the `Save Configuration` button, FLIM Playground saves configurations into the current profile and loads it for processing current and future datasets.

#### 12.6 Reset

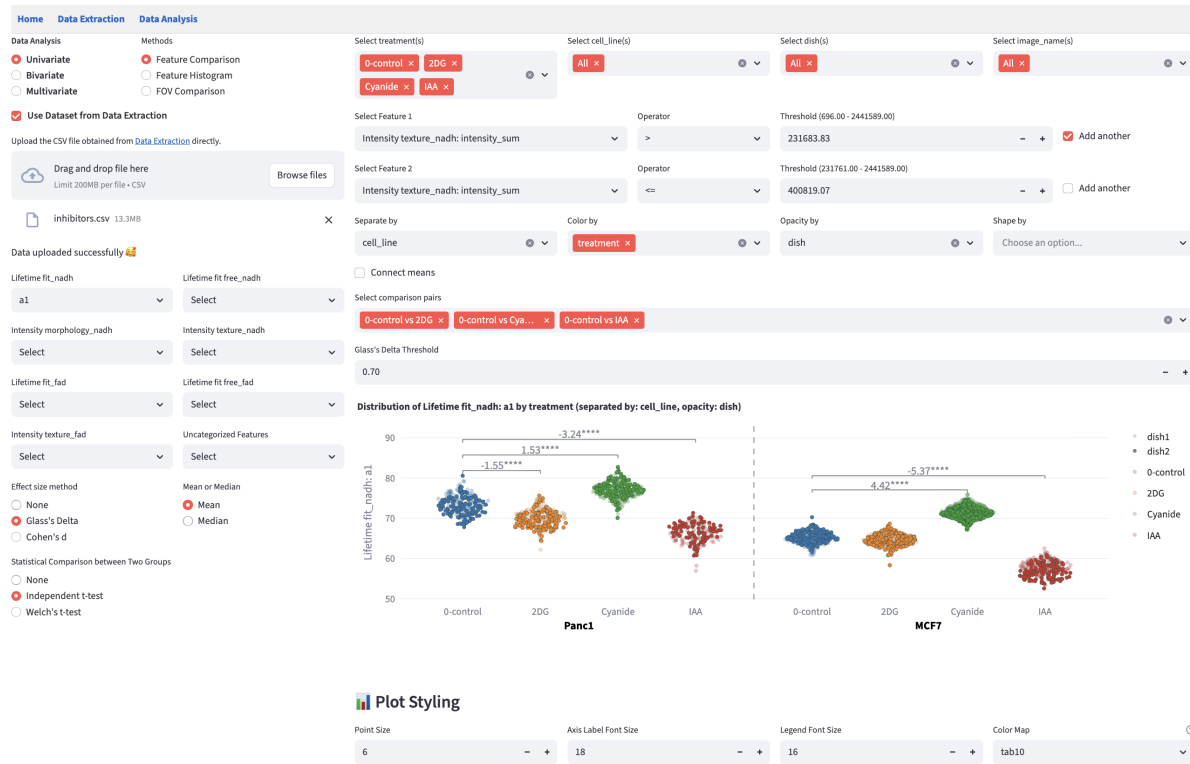
Users can reset the current profile to the default configurations.

## **Part III**

# **Univariate Analysis**

# 13 Feature Comparison

This method allows users to compare the distributions of a single numerical feature across groups. It can help users spot overarching trends between distributions, identify within-distribution shape-related characteristics such as skewness or bimodality that might signal sub-populations, and diagnose potential outliers.

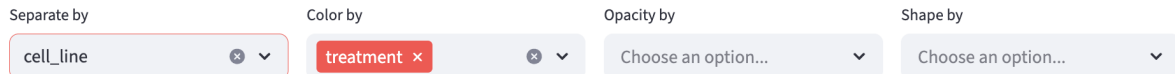


## 13.1 Shared Interface Components

- On the left, users can use the [selection widgets](#) to select the numerical feature to be compared. Exactly one feature can be selected from all feature groups.
- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Below the filters, users can apply the [visual channels widgets](#) that allow users to [Color by](#), [Opacity by](#), and [Shape by](#) categorical features.

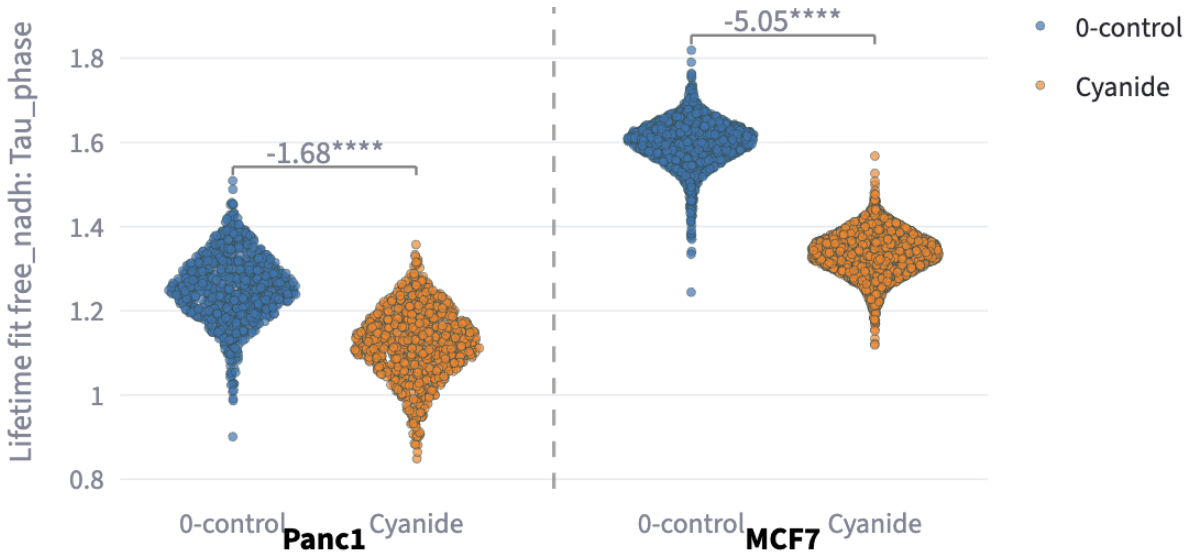
- On the bottom right, users can change the plot style using the [plot styling widgets](#).

## 13.2 Separate by

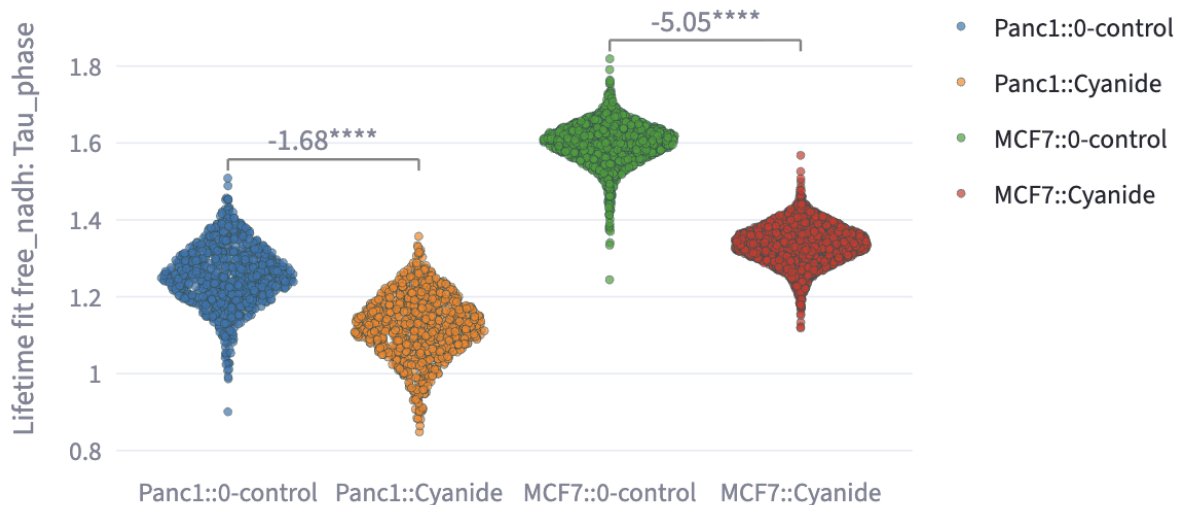


`Separate by` takes effect before `Color by`. `Separate by` divides the plot into sections separated by a dashed line, with each section labeled by one of the available categories of the selected categorical feature after [filtering](#). The selected feature of `Separate by` is automatically removed from available features in `Color by`. The section order is determined by the order of the categories which are sorted [numeric-alphabetically](#).

Then, visual channels (color, shape, opacity) and effect size annotation are applied within each section. For example, the visualization below was separated by `cell_line`, with each section labeled by the bold text of a cell line. Within each section, the x-axis rendered the `Color by` categories (treatments) so colors were consistent across sections. [Effect size](#) calculation and annotation (only effect size  $> 0.5$  is shown) were performed within each section.



If `Separate by` is left blank and both `cell_line` and `treatment` are selected in `Color by`, then colors are assigned to each cell line and treatment combination, and effect sizes are calculated across all combinations of group pairs (only pairs with effect size  $> 0.5$  are shown).



### 13.3 Comparison Widget

Statistical tests and effect size calculations are performed on the selected comparison pairs. FLIM Playground uses the groups on the x-axis populated by [Color by](#) to create comparison groups, starting from the leftmost group. For example, if there are 5 groups, then  $\binom{5}{2} = 10$  comparisons will be created. The number of groups can get combinatorially large and many of the comparisons are not meaningful. Therefore, two widgets are provided to filter the comparison groups:

- A selection widget that shows all possible comparisons by default and users can deselect groups they do not need.

Select comparison pairs

- 0-control vs 2DG ×
- 0-control vs Cya... ×
- 0-control vs IAA ×

- A threshold by effect size widget that filters out comparisons below the threshold.

Glass's Delta Threshold

## 13.4 Effect Size

Complementary to statistical tests, which address “is there a difference?”, effect size metrics answer “how large is the difference?”, enabling cross-study comparison. FLIM Playground offers two effect size calculation methods, Glass’s Delta and Cohen’s d, each can be in mean or median form.

Effect size method	Mean or Median
<input type="radio"/> None	<input checked="" type="radio"/> Mean
<input type="radio"/> Glass’s Delta	<input type="radio"/> Median
<input checked="" type="radio"/> Cohen’s d	

### 13.4.1 Glass’s Delta

Mean ( $\bar{X}$ ) form:

$$\Delta_G = \frac{\bar{X}_{\text{treatment}} - \bar{X}_{\text{control}}}{s_{\text{control}}}, \quad s_{\text{control}} = \sqrt{\frac{1}{n_c - 1} \sum_{i=1}^{n_c} (X_{\text{control}_i} - \bar{X}_{\text{control}})^2}$$

Median ( $\tilde{X}$ ) form:

$$\tilde{\Delta}_G = \frac{\tilde{X}_{\text{treatment}} - \tilde{X}_{\text{control}}}{\text{MAD}_{\text{control}}}, \quad \text{MAD}_{\text{control}} = 1.4826 \times \text{median} |X_{\text{control}_i} - \tilde{X}_{\text{control}}|, \quad i = 1, \dots, n_c$$

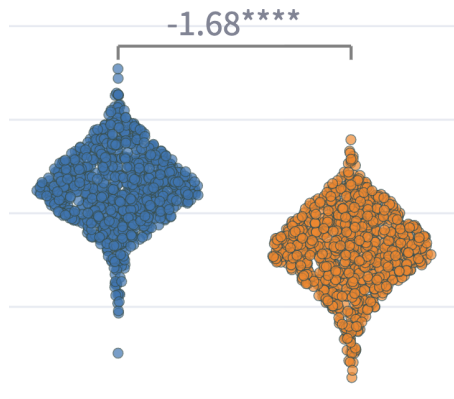
#### Note

MAD stands for [median absolute deviation](#). In order to make MAD asymptotically consistent with the standard deviation under normality, it is scaled by  $C = 1/\Phi^{-1}(0.75) \approx 1.4826$ . Implementation-wise, it uses `median_abs_deviation` from `scipy.stats`, and `scale` option is set to "normal" to account for the constant multiplier  $C$ . THE MEDIAN FORM of both Glass’s Delta and Cohen’s d IS NOT REPORTED IN THE LITERATURE.

#### Important

Which group is treatment and which is control matters! From the formula of  $\Delta_G$ , switching the order not only flips the sign of the result, but also changes the denominator, which is  $s_{\text{control}}$ , the standard deviation of the *control*. Currently, FLIM Playground treats the group on the left as control and the group on the right as treatment (the negative  $\Delta_G$  below implies this). In the future, it may support user-customizable x-axis order so that users can arrange the groups to make sure the control group is on the left. Or users can use [Cohen’s d](#) that does not assume the treatment-control

structure.



**i** Note

To keep the point-based visualization style that supports [interactive hover](#) while showing the violin-plot-like distribution, [Sina plot](#) is used: it uses `gaussian_kde` from `scikit-learn` to make sure the width of the point distribution is proportional to the kernel density.

### 13.4.2 Cohen's d

Mean ( $\bar{X}$ ) form:

$$d = \frac{\bar{X}_1 - \bar{X}_2}{s_p}, \quad s_p = \sqrt{\frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{n_1 + n_2 - 2}}$$

Median ( $\tilde{X}$ ) form:

$$\tilde{d} = \frac{\tilde{X}_1 - \tilde{X}_2}{s_{\tilde{p}}}, \quad s_{\tilde{p}} = \sqrt{\frac{(n_1 - 1) \text{MAD}_1^2 + (n_2 - 1) \text{MAD}_2^2}{n_1 + n_2 - 2}}$$

## 13.5 Statistical Test

Statistical tests are performed on the selected comparison pairs that exceed the effect size threshold (if users choose to calculate effect sizes). The statistical tests supported are:

- Independent t-test (student's t-test, for equal variances)

- Welch's t-test (for unequal variances)

Statistical Comparison between Two Groups

- None
- Independent t-test
- Welch's t-test

FLIM Playground uses `scipy.stats.ttest_ind` to perform the independent t-test and `scipy.stats.ttest_ind` with `equal_var=False` to perform the Welch's t-test.

## 13.6 Comparison Annotation

For each selected comparison pair that exceeds the effect size threshold (if any), the statistical significance, effect size, or both are annotated on the plot. It selects the lowest eligible position (over all points in both groups and over existing annotations) to annotate. If both statistical significance and effect size calculation are performed, the annotation is shown in the format of effect size followed by p-value. If only one is performed, then only the performed method's result is shown.

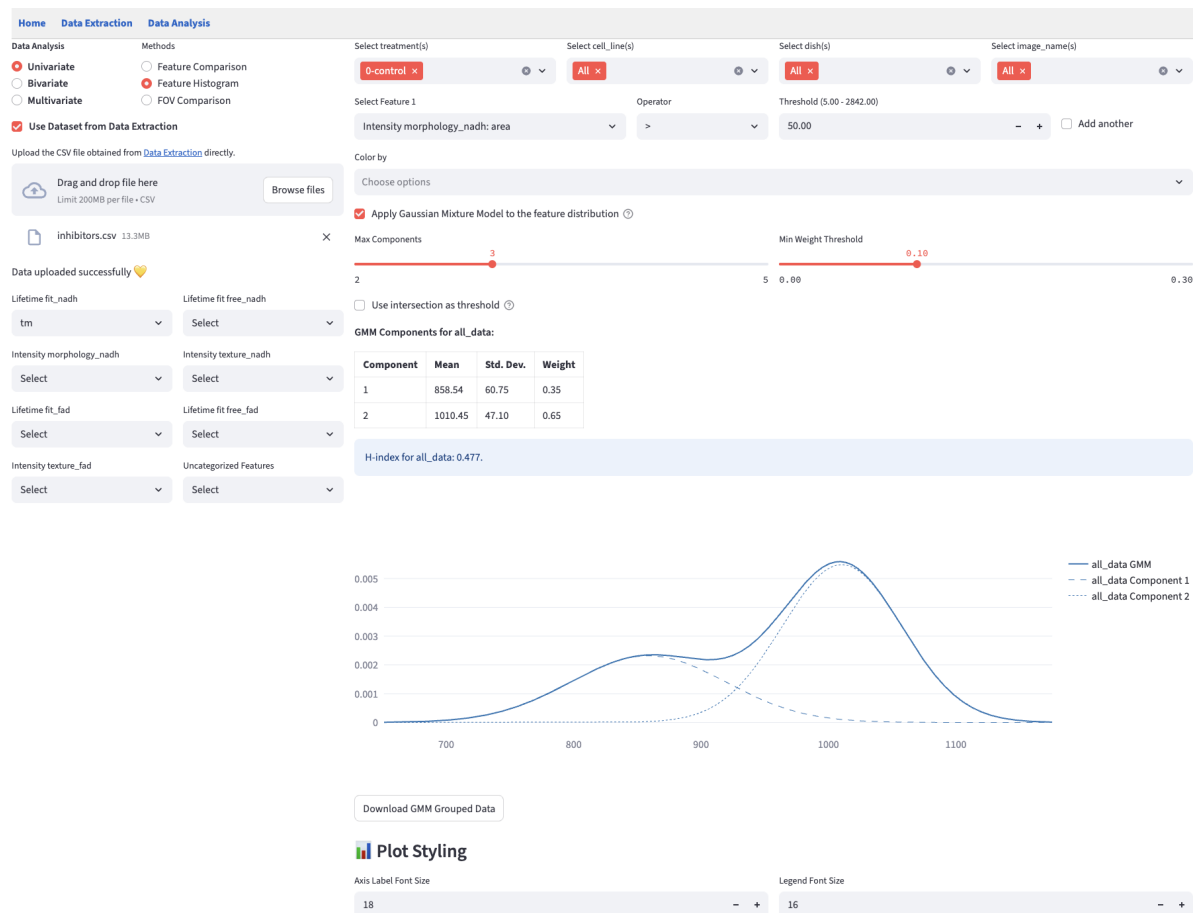
$\Delta=1.65$

\*\*\*\*\*

1.65\*\*\*\*\*

# 14 Feature Histogram

Feature histograms give an instant readout of each group's center, spread, skew and any multi-peaked shape that raw tables hide. FLIM Playground plots a histogram for each group based on the selected numerical feature. Additionally, the **Gaussian Mixture Model** mode can help find subcomponents of each histogram in an unsupervised way. To quantify the distribution heterogeneity, normalized **H-index** is calculated for each group.



## 14.1 Shared Interface Components

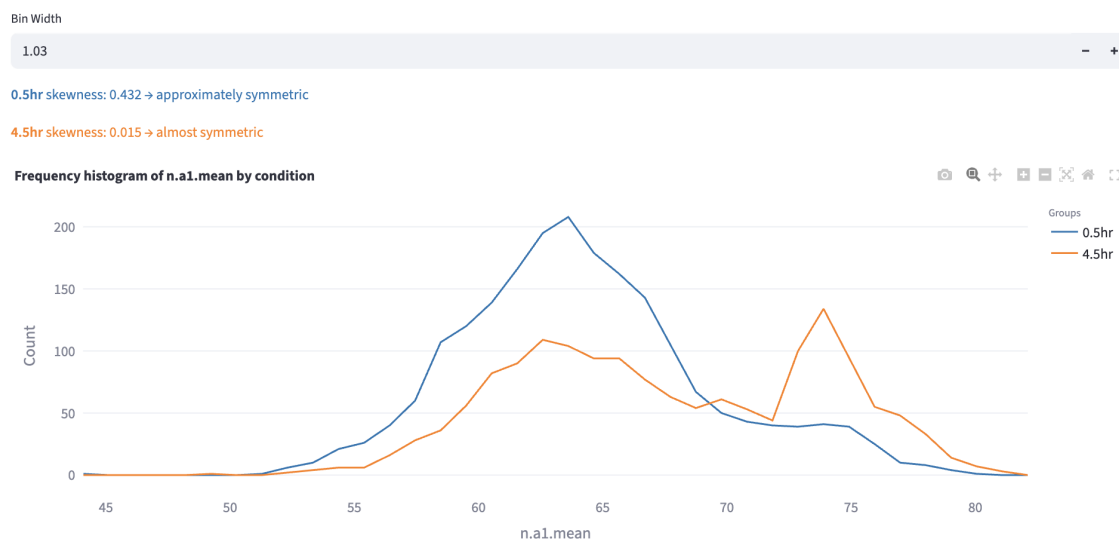
- On the left, users can use the [selection widgets](#) to select the numerical feature to see the histogram/GMM distribution. Exactly one feature can be selected from all feature groups.
- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Below the filters, users can apply the [visual channels widgets](#) that allow users to `Color by` categorical features. `Opacity by` and `Shape by` are *not* supported because there are no points in the histogram/GMM plot.
- On the bottom right, users can change the plot style using the [plot styling widgets](#).

## 14.2 Histogram Mode

With a single click, users can switch between the [histogram mode](#) and the [Gaussian Mixture Model mode](#). When the checkbox is unchecked, the histogram mode is selected.

Apply Gaussian Mixture Model to the feature distribution [?](#)

Histogram mode plots a histogram of the selected numerical feature for each group. The histogram bin width is user-controllable, with the default bin width determined automatically as the minimum bin width between the ‘sturges’ and ‘fd’ estimators (see [here](#) for more details). The maximum bin width is set to one-third of the range of the data. The step size is set to 1/50 of the range.



Additionally, the skewness of the distribution is computed and displayed<sup>13</sup>.

A rule of thumb categorization of skewness is as follows:

- Strongly left-skewed:  $< -1$
- Moderately left-skewed:  $-1$  to  $-0.5$
- Approximately symmetric:  $-0.5$  to  $-0.25$  and  $0.25$  to  $0.5$
- Almost symmetric:  $-0.25$  to  $0.25$
- Moderately right-skewed:  $0.5$  to  $1$
- Strongly right-skewed:  $> 1$

However, the skewness is just one way to quantify the distribution shape. In the example above, the orange distribution was clearly bimodal, but the skewness alone could not capture this. In applications where the distribution is multi-modal, and each mode can represent certain subpopulations (e.g. a cell type, a cell state, a cell cycle phase, etc.), users can use the [Gaussian Mixture Model mode](#) to find subcomponents of each distribution in an unsupervised way.

### 14.3 Gaussian Mixture Model Mode

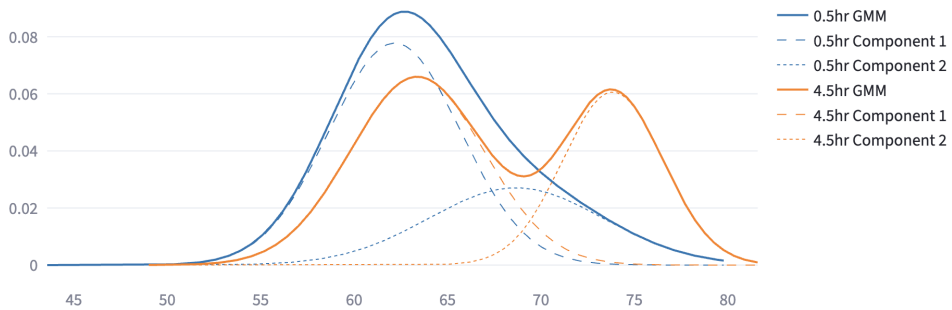
GMM Components for 0.5hr:

Component	Mean	Std. Dev.	Weight
1	62.18	3.51	0.68
2	68.67	4.66	0.32

GMM Components for 4.5hr:

Component	Mean	Std. Dev.	Weight
1	63.37	3.58	0.59
2	73.84	2.68	0.41

H-index for 0.5hr: 0.468. H-index for 4.5hr: 0.485.



A Gaussian Mixture Model (GMM) represents a dataset as a weighted sum of multiple Gaussian (normal) distributions, each defined by its own mean ( $\mu$ ) and (co)variance ( $\sigma^2$ ). By estimating the component

parameters and their weights (mixing coefficients) from the data—commonly using the Expectation-Maximization (EM) algorithm—GMMs can capture complex, multimodal distributions that a single Gaussian cannot.

### 14.3.1 Fit Gaussian Mixture Models



Figure 14.1: Gaussian Mixture Model Configuration

- Fits GMMs on the selected numerical feature up to Max Components components, each of which has its component weights ( $w_i$ ), means ( $\mu_i$ ) and variances ( $\sigma_i^2$ ).
- Retains only *valid* models in which every component has weight at least Min Weight Threshold (the weights of all components add to 1).
- FLIM Playground will choose the valid model with the lowest BIC score that penalizes the model complexity to avoid overfitting.

### 14.3.2 Heterogeneity index

To quantify the subpopulation structure, based on the GMM fit, FLIM Playground computes an “H $\square$ index” for each group as a weighted entropy $\square$ distance: each component contributes according to its weight’s uncertainty ( $-w_i \log w_i$ )<sup>14</sup> scaled by how far its mean  $\mu_i$  sits from the overall mixture mean  $\bar{\mu} = \sum_{i=1}^k w_i \mu_i$ , normalized by the standard deviation of  $\mu_i$ s.

$$H = \sum_{i=1}^k (-w_i \log w_i) d_i, \quad d_i = \frac{|\mu_i - \bar{\mu}|}{\sigma}$$

### 14.3.3 Classification

- Users can choose to perform **intersection thresholding** or **hard assignment** to determine which GMM component each data point belongs to. It appends a new categorical feature column called `GMM_Group` (recognizable by methods in Data Analysis as a categorical feature) to the filtered dataset, the values of which are `{color_group}_1`, `{color_group}_2`, etc. In the example above, the color groups were created by 0.5hr and 4.5hr because the user colored the data by hour. The classification result can be saved by clicking:

[Download GMM Grouped Data](#)

#### 14.3.3.1 Intersection thresholding

- Once the GMM is estimated, thresholds can be obtained by locating the intersection points of adjacent component densities:

$$w_1 \mathcal{N}(x; \mu_1, \sigma_1^2) = w_2 \mathcal{N}(x; \mu_2, \sigma_2^2)$$

where  $w_1, \mu_1, \sigma_1$  are the mixture weight, mean, and standard deviation of the first Gaussian component, respectively, and  $w_2, \mu_2, \sigma_2$  those of the second component.

- The above equation can be formed as a quadratic equation and solved for the unique root lying between  $\mu_1$  and  $\mu_2$ . In practice, we bracket this interval and apply a robust one-dimensional root-finding routine (e.g. Brent's method) to compute intersection points. Data points are then assigned to the class corresponding to the interval in which they fall.

! Important

In rare cases when there is more than one intersection point, the data point is assigned using hard assignment rather than intersection thresholding.

#### 14.3.3.2 Hard assignment

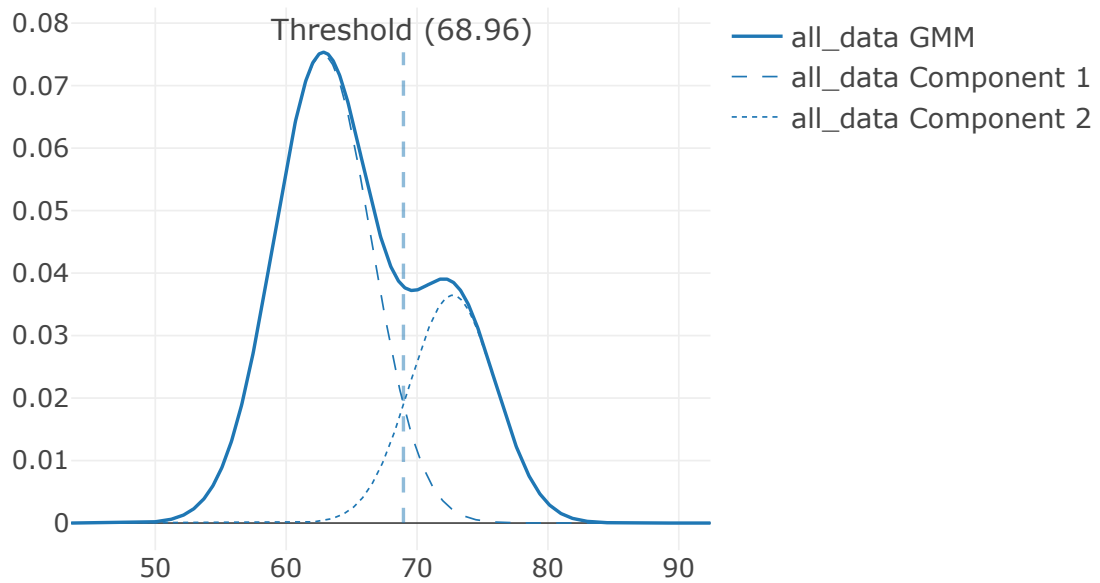
- For each data point, it computes the posterior probability (responsibility) of each component, and assigns the data point to the component with the highest responsibility.

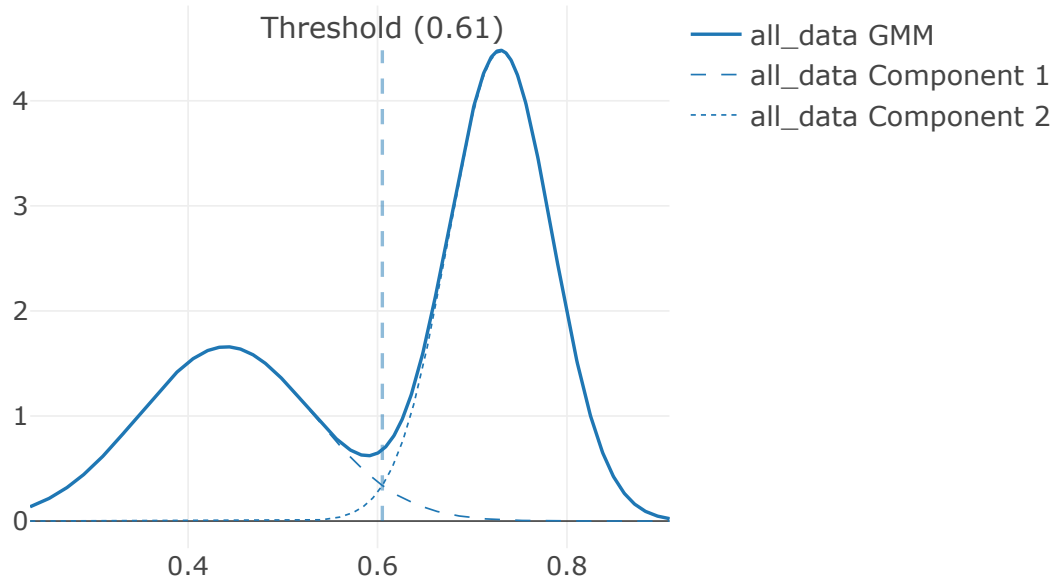
$$\text{label}(x) = \arg \max_i \gamma_i(x)$$

$$\text{where } \gamma_i(x) = \frac{w_i \mathcal{N}(x | \mu_i, \sigma_i^2)}{\sum_{j=1}^K w_j \mathcal{N}(x | \mu_j, \sigma_j^2)}$$

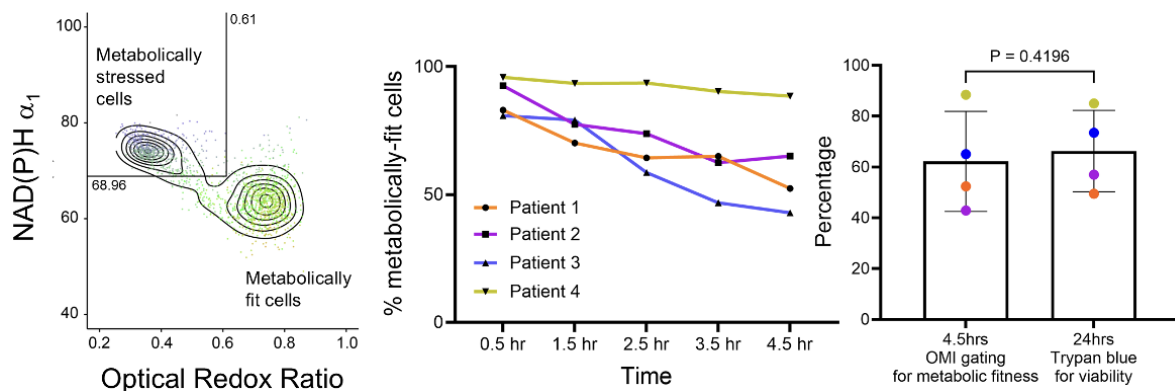
#### 14.3.4 Example

Pham et al.<sup>15</sup> used GMM with hard thresholding to create gates using NAD(P)H  $a_1$  and optical redox ratio. These gates were used to define metabolically stressed and fit cells in their study of the impact of cryopreservation on T cell metabolism.





They compared the GMM-derived gate with cell viability by Trypan blue staining:



It suggests that optical metabolic imaging (OMI) could serve as an early, non-destructive indicator of T cell viability post-thaw.

# 15 Field of View Comparison

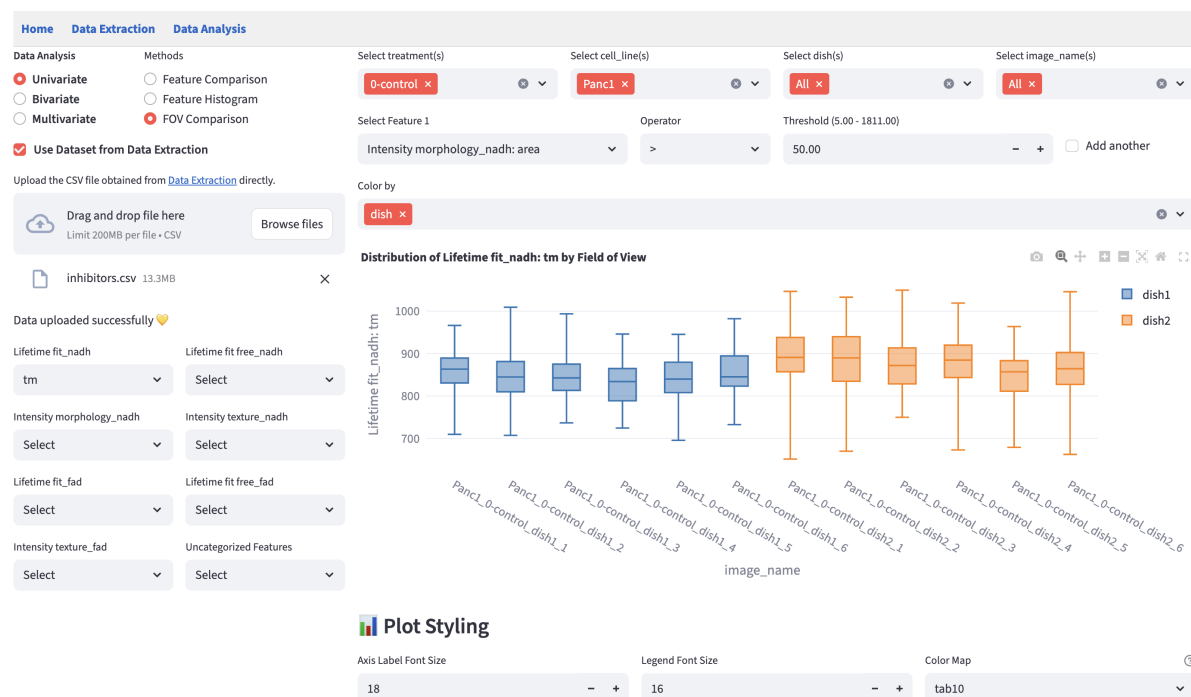
In imaging datasets like FLIM data, field of view (FOV) is an important categorical feature that documents where a certain set of rows (e.g. cells) originate. Visualizing each FOV side by side on a numerical feature can help identify potential FOV-level outliers.

FLIM Playground finds the FOV identifier column from the dataset using the `FOV column name` field in data analysis config (dataset not extracted by Data Extraction) or data extraction config (dataset extracted by [Data Extraction](#)).

If the FOV identifier column is not found in the dataset, then a warning message will be shown:

**Warning: We cannot find the fov column from your dataset.**

If found:



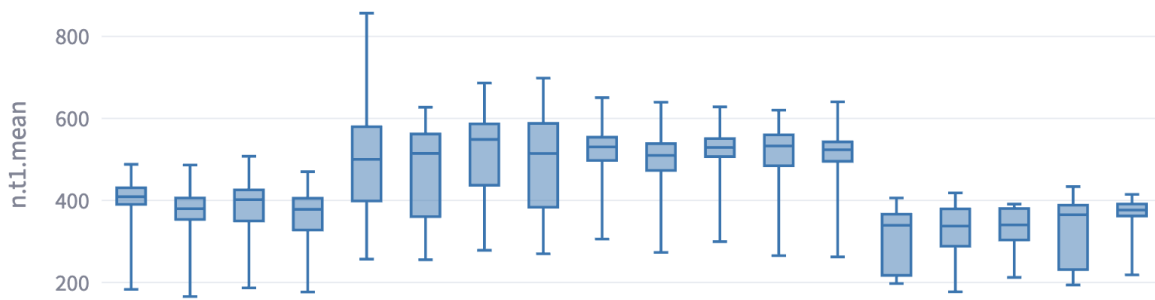
## 15.1 Shared Interface Components

- On the left, users can use the [selection widgets](#) to select the numerical feature to see the boxplot of each FOV. Exactly one feature can be selected from all feature groups.
- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Below the filters, users can apply the [visual channels widgets](#) that allow users to `Color by` categorical features. `Opacity by` and `Shape by` are *not* supported because there are no points in the boxplot.
- On the bottom right, users can change the plot style using the [plot styling widgets](#).

Besides the shared components, this module does not have any method-specific components.

## 15.2 Example

Sometimes, the FOV-level boxplots may show consistent divergence across different imaging days, which may suggest that there is experimental or data extraction inconsistency (e.g. shifts are not optimized correctly for one day versus the others). Below is an example that, despite the same conditions, systematically diverged across days, but was consistent within the same day.

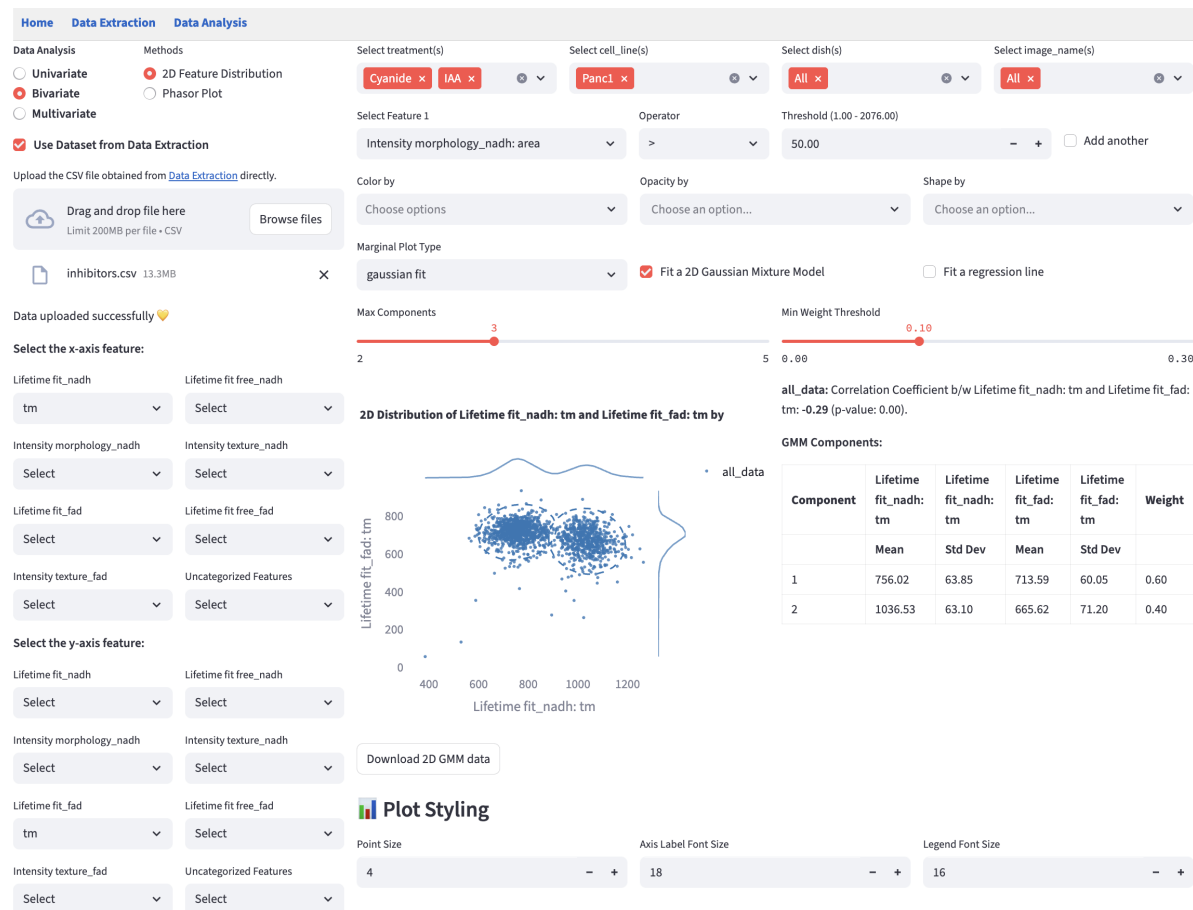


## **Part IV**

# **Bivariate Analysis**

# 16 Feature Distribution

This method extends the univariate [Feature Histogram](#) method to visualize the distribution of two numerical features on a scatter plot. It helps reveal patterns such as correlations, clusters, trends, or outliers.

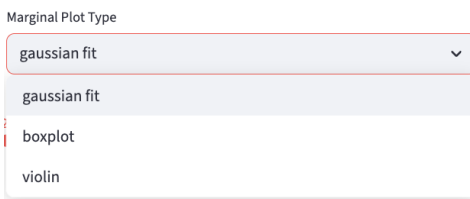


## 16.1 Shared Interface Components

- On the left, users can use the [selection widgets](#) to select the two numerical features on the x-axis and y-axis.
- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Below the filters, users can apply the [visual channels widgets](#) that allow users to `Color by`, `Opacity by`, and `Shape by` categorical features.
- On the bottom right, users can change the plot style using the [plot styling widgets](#).

## 16.2 Marginal Distribution

Marginal distributions (the distribution of a single feature) are plotted at the top and the right of the scatter plot. Users can choose the plot type from the dropdown menu.



`gaussian fit`: provides a kernel-density estimate using Gaussian kernels. It uses `gaussian_kde` from `scipy.stats` and sets all parameters to default values.

## 16.3 2D Gaussian Mixture Model

Using the same method as fitting a 1D [Gaussian Mixture Model](#), users can fit a 2D GMM by specifying `Max Components` and `Min Weight Threshold` for each color group.

Ellipses that capture approximately 95% of the points in each subcomponent are drawn. The color of the ellipses is determined by the color group that the subcomponent belongs to.

### **i** How ellipses are drawn

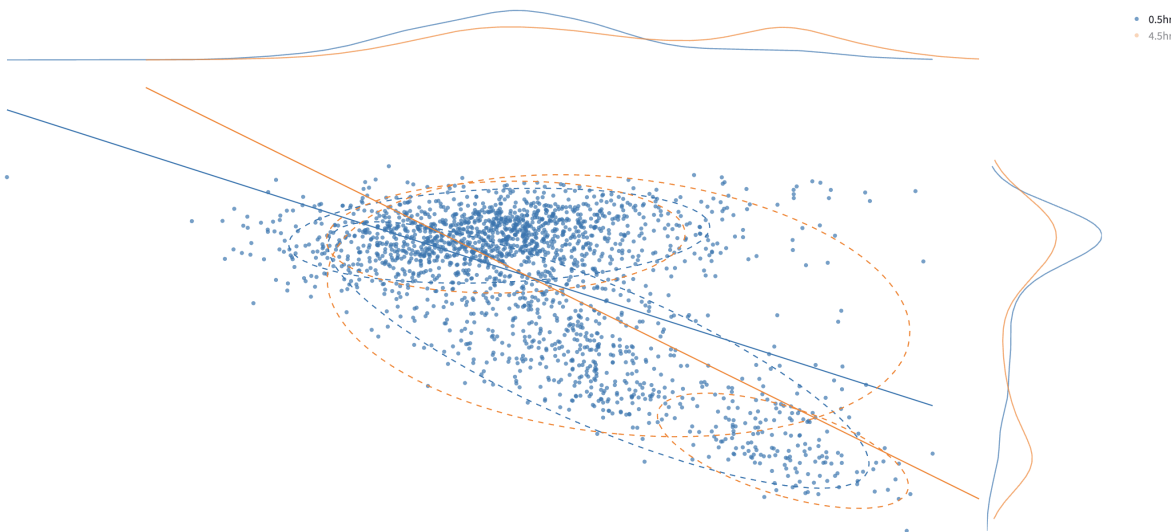
Each ellipse represents a subcomponent of the GMM.

- Center of the ellipse is the mean of the subcomponent  $(\mu_x, \mu_y)$ .
- The ellipse is rotated by  $\theta = \arctan 2(v_{1y}, v_{1x})$ , where  $\mathbf{v}_1 = \begin{bmatrix} v_{1x} \\ v_{1y} \end{bmatrix}$  is the first eigenvector of the covariance matrix, so that the major axis of the ellipse is aligned with the eigenvector

corresponding to the largest eigenvalue.

- The semi-axis lengths are  $r\sqrt{\lambda_1}$  and  $r\sqrt{\lambda_2}$ , where  $\lambda_1$  and  $\lambda_2$  are the covariance eigenvalues (variances along the principal axes) scaled by  $r = \sqrt{\chi^2_{df=2, p=0.95}}$  (Mahalanobis distance) that captures approximately 95% of the points in the subcomponent.

All the components below will be rendered for each color group, which is created by user-chosen categorical features in [Color by](#).



### 💡 Tip

Users can click on a legend group to hide/show the data points in that group. In the above example, to avoid cluttering the plot, the orange group is hidden.

Tables of subcomponents' means, standard deviations, and weights are reported on the side.

0.5hr: Correlation Coefficient b/w n.a1.mean and rr.mean: -0.56 (p-value: 0.00).

GMM Components:

Component	n.a1.mean	n.a1.mean	rr.mean	rr.mean	Weight
	Mean	Std Dev	Mean	Std Dev	
1	66.71	5.29	0.54	0.11	0.36
2	62.83	4.12	0.74	0.04	0.64

4.5hr: Correlation Coefficient b/w n.a1.mean and rr.mean: -0.76 (p-value: 0.00).

GMM Components:

Component	n.a1.mean	n.a1.mean	rr.mean	rr.mean	Weight
	Mean	Std Dev	Mean	Std Dev	
1	73.43	3.12	0.39	0.07	0.39
2	63.87	4.33	0.72	0.07	0.61

The classification result can be saved by clicking:

[Download 2D GMM data](#)

The classification is done by [hard assignment](#) to determine which GMM component each data point belongs to. It appends a new categorical feature column called 2D\_GMM\_group to the filtered dataset, the values of which are {color\_group}\_group1, {color\_group}\_group2, etc. The downloaded dataset keeps all the features plus the new categorical feature column, which is recognized by any method in Data Analysis as a categorical feature like others. The rows that pass the [filters](#) are included.

## 16.4 Regression line

A linear regression line is fitted to the data points in each color group. The key statistics are reported on the side.

**Regression R<sup>2</sup>:** 0.311 (Slope: -0.014, Intercept: 1.570)

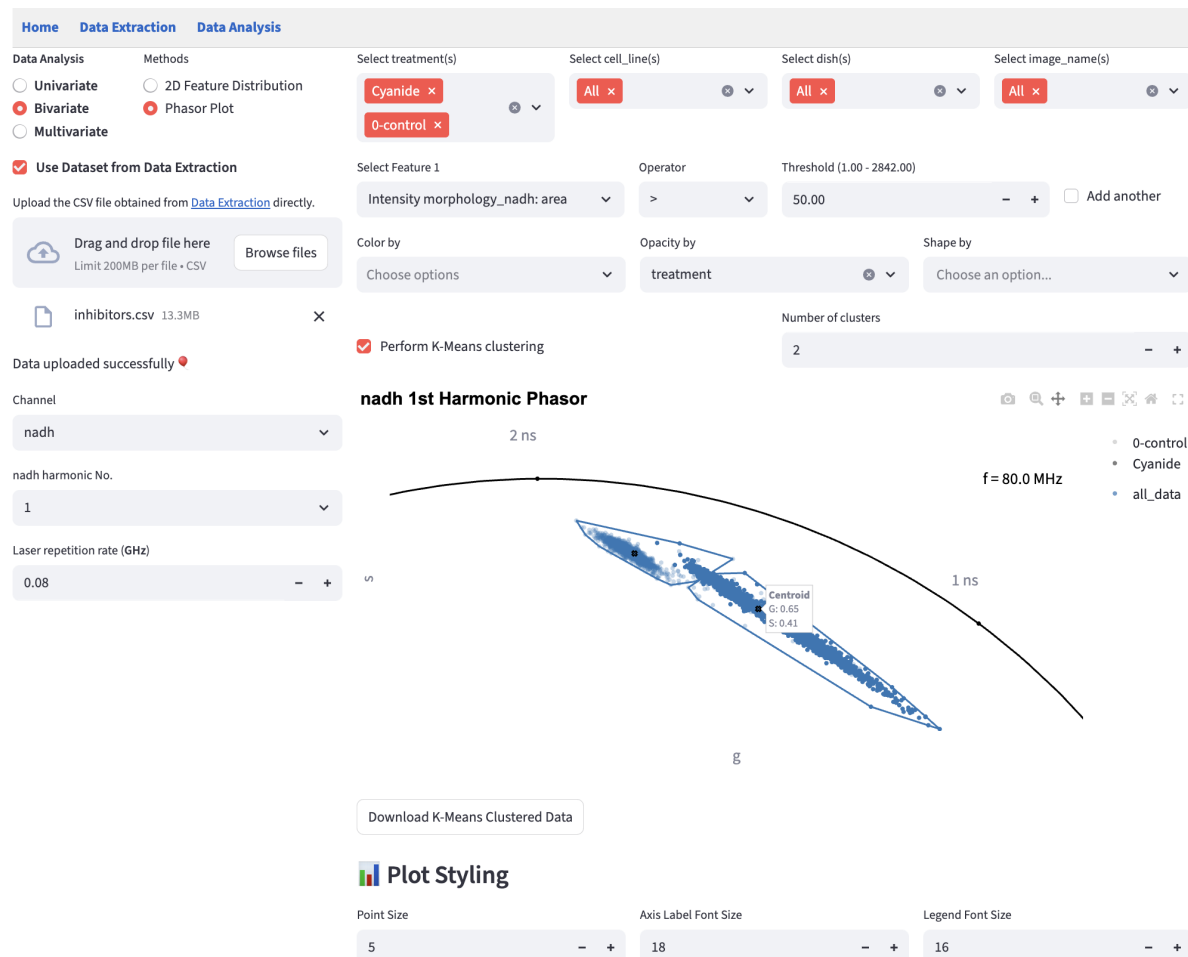
$R^2$  tells how much of the variability in  $y$  is captured by the model, compared to just predicting the  $\bar{y}$ .

  $R^2$

- $R^2 = 1 \rightarrow$  perfect prediction.
- $R^2 = 0 \rightarrow$  model predicts no better than the mean.

# 17 Phasor Analysis

This method focuses on visualizing the distribution of the 1st harmonic or 2nd harmonic phasor coordinates, extracted by the [Lifetime fit free](#) extractor, and performs optional clustering to highlight subpopulations or structural patterns.

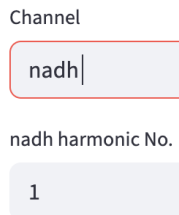


## 17.1 Shared Interface Components

- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Below the filters, users can apply the [visual channels widgets](#) that allow users to [Color by](#), [Opacity by](#), and [Shape by](#) categorical features.
- On the bottom right, users can change the plot style using the [plot styling widgets](#).

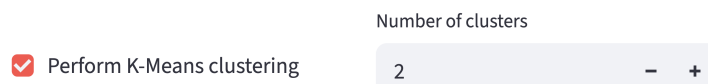
## 17.2 Select Phasor Coordinate

Instead of using the [bivariate selection widgets](#) that select two generic numerical features, this method renders a dropdown menu to select the channel and another to select the harmonic.



## 17.3 K-means Clustering

Phasor coordinates of single-cell ROIs from the selected channel and harmonic are plotted in the phasor space. K-means clustering can be performed for each color group (created by [Color by](#)). If [Perform K-means Clustering](#) is checked, users can specify the number of clusters to be created based on their prior knowledge. In the future, automatic tuning for this parameter, akin to selecting the number of components of [GMM](#), using silhouette analysis will be added.



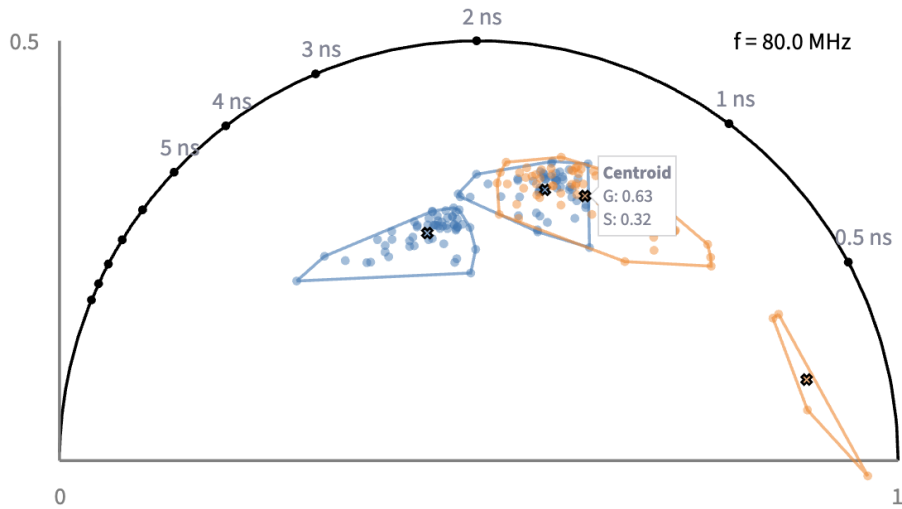
K-means clustering is performed given the number of clusters specified for each color group. Mathematically, it is an iterative algorithm that finds the best cluster assignment with regard to what cluster each point belongs to and what centroid the cluster has. The *best* is defined as the one that minimizes the sum of the squared distances between each point and the center of the cluster it belongs to.

$$\min_{\{C_k\}_{k=1}^K} \sum_{k=1}^K \sum_{x_i \in C_k} \|x_i - \mu(C_k)\|_2^2, \quad \mu(C_k) = \frac{1}{|C_k|} \sum_{x_i \in C_k} x_i.$$

First, FLIM Playground standardizes the phasor coordinates because  $g$  (real part) and  $s$  (imaginary) are not necessarily on the same scale.

Then, it performs k-means clustering using the `sklearn.cluster.KMeans` function. All the other hyperparameters control the optimization process (e.g. `init` controls the initialization of the centroids), and they are set to their [default values](#). 42 is used as the random seed to ensure reproducibility.

To visualize the result of clustering, FLIM Playground draws a polygon (convex hull) that contains all the points of each cluster, the color of which is determined by the color group the cluster belongs to. It uses the `scipy.spatial.ConvexHull` function to calculate the convex hull. The center of each cluster is marked as a black bordered cross and users can hover over it to see its coordinates.



### 17.3.1 Export Clustered Dataset

Users can download the clustered dataset by clicking:

[Download K-Means Clustered Data](#)

The downloaded dataset keeps all the features plus the new categorical feature column (`k_means_cluster`), which is recognized by any method in Data Analysis as a categorical feature like others. The rows that pass the [filters](#) are included.

# **Part V**

# **Multivariate Analysis**

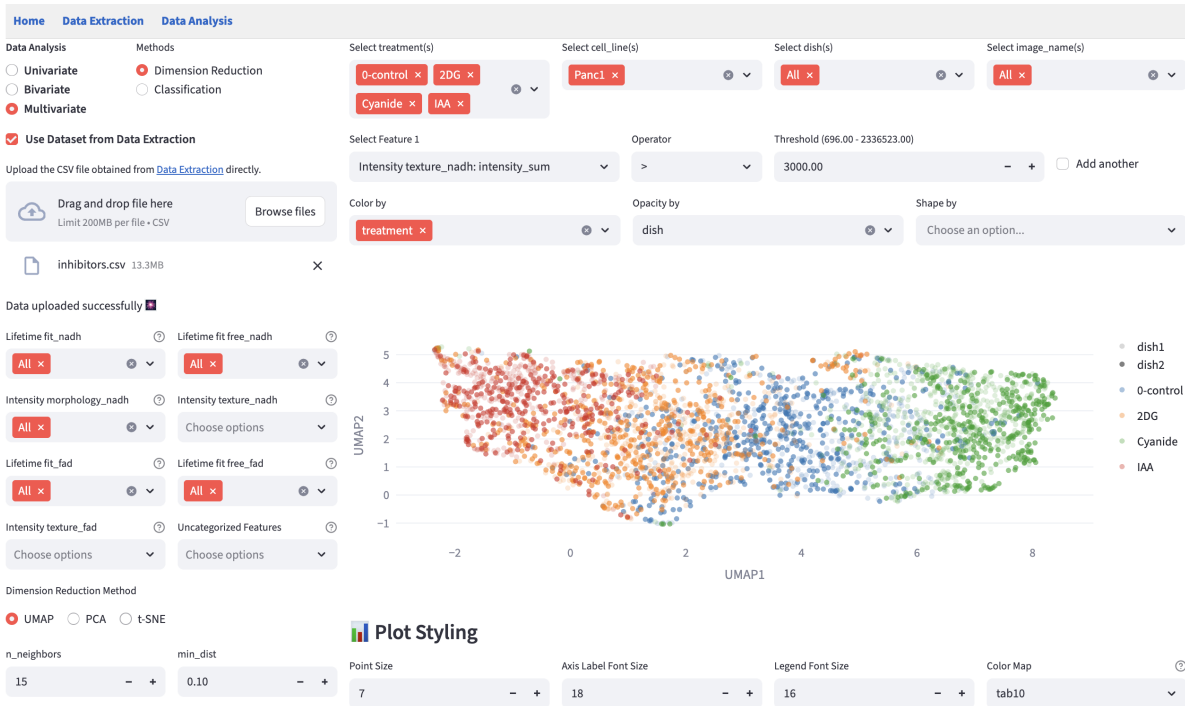
# 18 Dimension Reduction

To represent and visualize high-dimensional data in a meaningful way and help users interpret the data, dimension reduction methods project the data into a lower-dimensional space. Three popular dimension reduction methods are often used: UMAP, PCA, and t-SNE.

Principal Component Analysis (PCA) is a linear dimension reduction method that projects the data onto the directions of the largest variance. It is fast, deterministic, and easier to interpret (PCA loadings are provided), but it is limited by the linear nature of the projection. For a detailed discussion of PCA, please refer to [this post](#).

Uniform Manifold Approximation and Projection (UMAP) and t-Distributed Stochastic Neighbor Embedding (t-SNE) are non-linear dimension reduction methods that can capture more complex patterns in the data. They are more flexible and can handle non-linear relationships between features, but they are more computationally expensive, less interpretable, not deterministic (FLIM Playground uses the random seed 42 to ensure reproducibility), and dependent on the hyperparameters. Here is a [post](#) that explains how UMAP and t-SNE work at a high level and the meaning of some of the hyperparameters.

The numerical features are standardized before the dimension reduction.



## 18.1 Shared Interface Components

- On the left, users can use the [selection widgets](#) to select multiple numerical features from each feature group.
- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Below the filters, users can apply the [visual channels widgets](#) that allow users to [Color by](#), [Opacity by](#), and [Shape by](#) categorical features.
- On the bottom right, users can change the plot style using the [plot styling widgets](#).

## 18.2 Hyperparameter Widget

Hyperparameters really matter, so we provide a set of interactive widgets to help users change the hyperparameters and see their effects in real time.

### 18.2.1 UMAP

FLIM Playground uses the `umap-learn` implementation of UMAP. It chooses to expose `n_neighbors` and `min_dist` for users to tweak. Other hyperparameters are set to [default values](#).

### 18.2.2 t-SNE

FLIM Playground uses the `sklearn.manifold.TSNE` implementation of t-SNE. It chooses to expose `perplexity` and `early_exaggeration` for users to tweak. Other hyperparameters are set to [default values](#).

UMAP  PCA  t-SNE

n\_neighbors

15

- +

min\_dist

0.10

- +

UMAP  PCA  t-SNE

UMAP  PCA  t-SNE

perplexity

15

- +

early\_exaggeration

1

- +

# 19 Classification

A machine learning classifier learns patterns from categorized data and then predicts the category of new, unseen data. This module provides a set of machine learning classifiers and interactive widgets to help users perform classification.

All the classifiers are implemented using `scikit-learn` and the performance metrics are calculated using `sklearn.metrics`. A fixed random seed (42) is used for all the classifiers to ensure reproducibility.

The screenshot shows the Data Analysis pipeline interface. At the top, there are tabs for Home, Data Extraction, and Data Analysis. Under Data Analysis, there are sections for Data Analysis Methods (Univariate, Bivariate, Multivariate), Select treatment(s) (0-control, 2DG, Cyanide, Panc1), Select cell\_line(s) (All), Select dish(s) (All), and Select image\_name(s) (All). A checkbox 'Use Dataset from Data Extraction' is checked, and a file 'inhibitors.csv' (13.3MB) is uploaded. The 'Classify by' section shows 'treatment' selected for sampling method 'Undersampling'. The 'Performance Metrics' section displays overall accuracy of 0.8177 and per-class metrics for 2DG and the rest. The 'ROC Curve' and 'Confusion Matrix' plots are visible at the bottom.

Metric	Value
Accuracy	0.8177
N	1037

Class	N	Precision	Recall	Specificity	Youden's J	F1 Score
2DG	285	0.6159	0.8947	0.7886	0.6833	0.7296
the rest	752	0.9518	0.7886	0.8947	0.6833	0.8625
Average	518.50	0.7839	0.8417	0.8417	0.6833	0.7961

The performance metric plots are cut off in the screenshot. See the [performance metrics section](#) for the complete view.

An overview of the pipeline:

- Choose the numerical features on the left.
  - The z-score normalization is applied to the features before training if the chosen classifier is SVM or Logistic Regression, because they are sensitive to feature magnitudes, while Random Forest and Gradient Boosting are not.
- Choose the classifier and train-test split.
- Filter the dataset.
- Choose categorical features to form classes.
- Choose any of the methods to handle [class imbalance](#).
- Choose the classes to be classified.
- Look at the [performance metrics](#), [confusion matrix](#), and [ROC curve](#).
- Iterate over the previous options to improve the performance with regard to the metrics you care about.

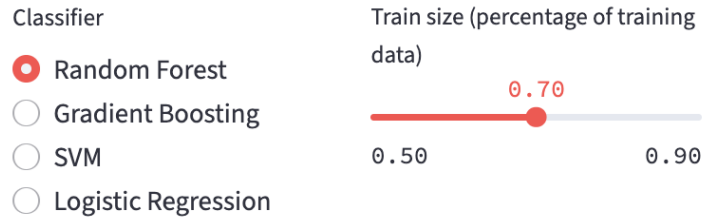
## 19.1 Shared Interface Components

- On the left, users can use the [selection widgets](#) to select numerical features for the classifier. Multiple features can be selected from each feature group.
- On the top right, users can use the [filters](#) to subset the data to find the groups of interest.
- Above the performance metrics plots, users can change the plot style using the [plot styling widgets](#).

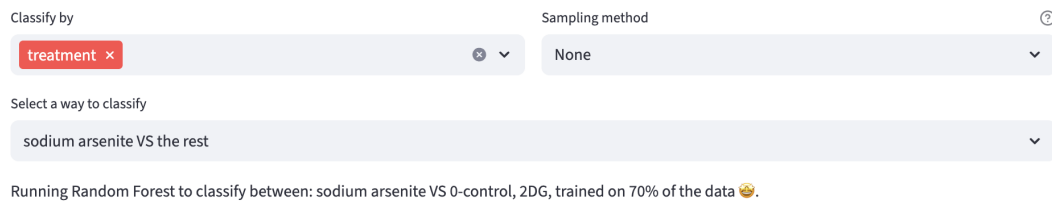
## 19.2 Method-specific Components

- Choose the classifier and train-test split: the slider controls the proportion of the data used to train the classifier and the rest is used as unseen data to test its generalizability. The following classifiers are available:
  - [Random Forest](#)
  - [Gradient Boosting](#)
  - Support Vector Machine (SVM)
  - [Logistic Regression](#)

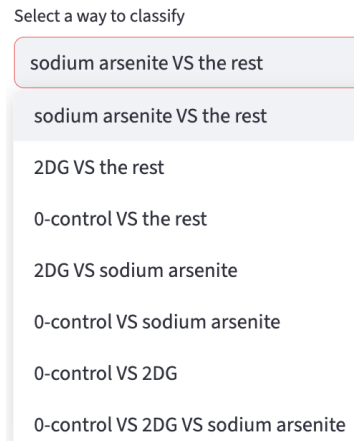
The [train-test split](#) is used to evaluate the generalizability of the classifier. The Validation set is used in hyperparameter tuning. Since the classifiers above do not have a lot of hyperparameters to tune, it does not partition a separate validation set. Though the k-fold cross-validation is used in tuning the threshold.



- **Classify by:** The `Color by` from the [visual channels widgets](#) is renamed to `Classify by` and it is used to form classes based on (a combination of) selected categorical features. Other visual channels are not supported in this method.



- Choose the classes to be classified: classes are formed based on the combinations of categories of the selected categorical features. In general, it supports binary classification and in general n-way classification where n is the number of classes. If one categorical feature is selected and after filtering has 3 categories, then all possible ways to classify are produced including the three-way classification. Additionally, the `class x vs the rest` option is provided for each class.



### 19.2.1 Class Imbalance Handling

We provide three options for handling class imbalance (i.e., one or more classes in a dataset have significantly fewer samples than others, which would otherwise cause a classifier to bias toward the

majority class). Note: to prevent data leakage, all the methods below are applied to the training set only.

- **Sampling methods:**

- None: stratified random sampling of the data on classification classes, regardless of the class size. It is implemented using `train_test_split` from `sklearn.model_selection`.
- Undersampling: All classes are downsampled to the size of the smallest class. It is implemented using `RandomUnderSampler` from `imbalanced-learn`.
- Oversampling: All classes are upsampled to the size of the largest class by randomly duplicating samples from minority classes. It is implemented using `RandomOverSampler` from `imbalanced-learn`.

Sampling method

None

None

Undersampling

Oversampling

- **Class weights:**

- None: no class weights.
- Balanced: the class weights are inversely proportional to the class size (support). Note: Gradient Boosting does not support class weights in the scikit-learn implementation, and thus this option is not available for it.

- **Threshold tuning:** Whereas the previous two options influence the probability estimation part of the classifier, this option adjusts only the decision boundary used to convert predicted probabilities into class labels.

- None: no threshold tuning. The default decision is always the class that has the highest predicted probability (in binary, anything above 0.5; in multi-class, the class with the highest predicted probability).
- Balanced Accuracy: the threshold is set to maximize the balanced accuracy score, which is the unweighted average of each class's recall (i.e., the minority class's recall is as important as the majority class's recall).
- F1 Score: the threshold is set to maximize the unweighted average of each class's F1 score.

Threshold tuning based on

None |

None

Balanced Accuracy

F1 Score

### i How does threshold tuning work?

It treats threshold selection as a separate optimization problem from model training, allowing thresholds to be tuned independently of the underlying classifier's probability estimates.

**K-fold cross-validation for the probability model:** The procedure begins by establishing a stratified [k-fold cross-validation](#) scheme (default k=5): the dataset is divided into k mutually exclusive subsets, and for each fold, one subset serves as the validation set while the remaining k-1 subsets form the training set.

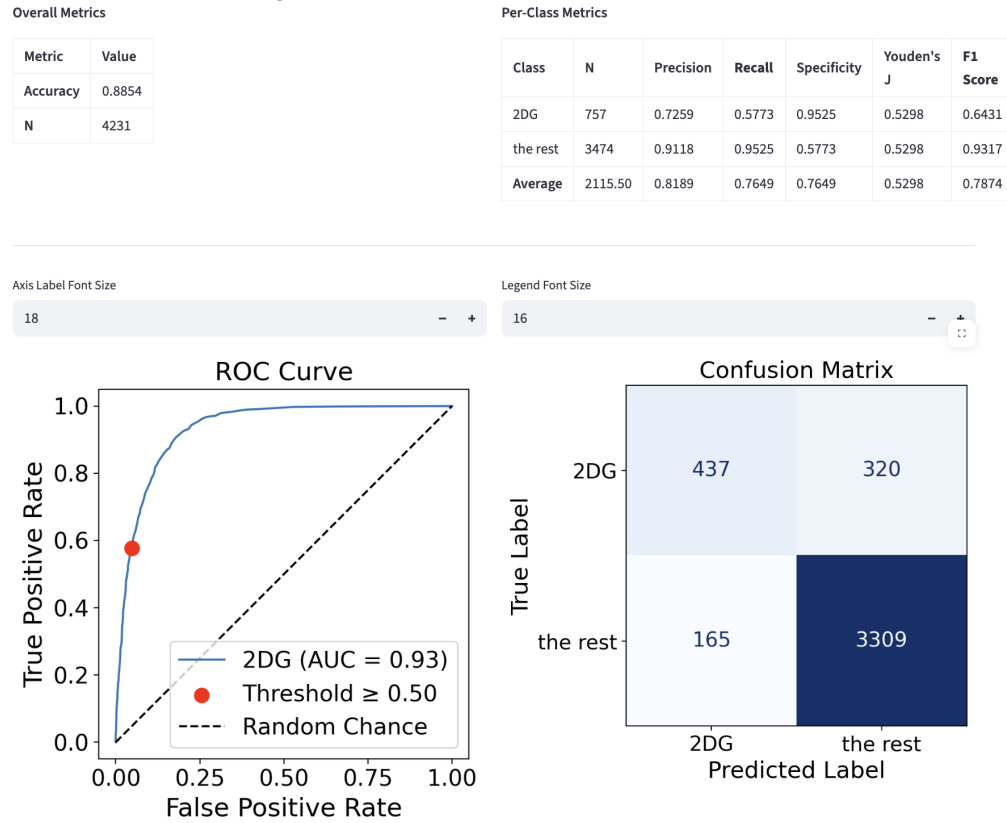
The probability models are trained separately for each cross-validation fold. This process yields k distinct probability models, each trained on a different subset of the data, along with their corresponding validation set probability predictions that collectively cover all samples exactly once.

**Threshold optimization:** A single threshold optimization procedure is performed once using all cross-validation folds. The threshold optimization process differs for binary and multi-class classification problems. For binary classification, a single threshold parameter  $\theta \in [0, 1]$  is optimized, where class 1 is predicted when  $P(\text{class } 1 \mid x) \geq \theta$ , and class 0 otherwise. For multi-class problems, a threshold vector  $\theta = [\theta_1, \theta_2, \dots, \theta_k]$  is optimized simultaneously for all k classes, where thresholds are applied through a normalized probability scheme: predictions are made by selecting the class with the highest normalized probability  $P(\text{class } i \mid x)/(\theta_i + \epsilon)$ , where  $\epsilon = 10^{-10}$  prevents division by zero.

The optimization objective function evaluates candidate thresholds by: (1) applying the same threshold(s) on top of the fold-specific probability models to generate prediction classes for the fold-specific validation set, (2) computing the specified evaluation metric (e.g., balanced accuracy, F1-score) between predicted and true labels for each fold, and (3) returning the negative mean cross-validated score across all folds (for minimization using the Nelder-Mead simplex algorithm). Finally, the optimized threshold(s) are applied on top of the probability model trained on the entire training set to the test set to generate the final predictions.

## *Threshold Tuning Example*

Before threshold tuning:



After threshold tuning based on the balanced accuracy metric:

## Performance Metrics

Overall Metrics

Metric	Value
Accuracy	0.8225
N	4231

Per-Class Metrics

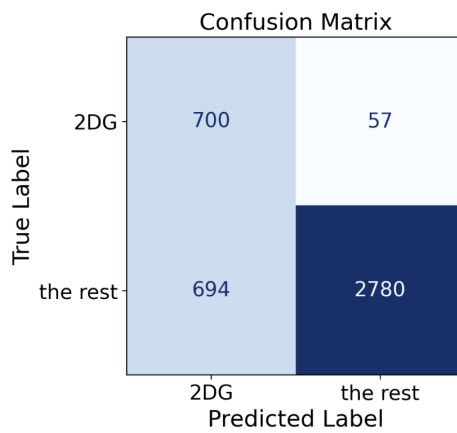
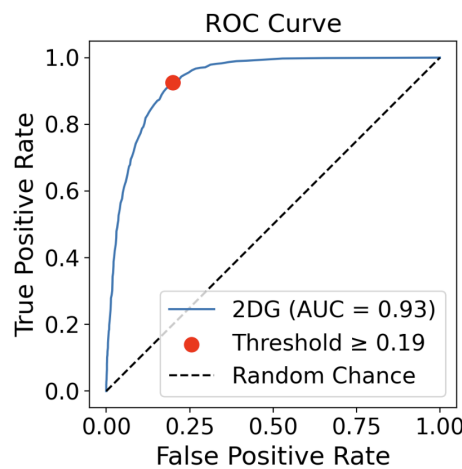
Class	N	Precision	Recall	Specificity	Youden's J	F1 Score
2DG	757	0.5022	0.9247	0.8002	0.7249	0.6509
the rest	3474	0.9799	0.8002	0.9247	0.7249	0.8810
Average	2115.50	0.7410	0.8625	0.8625	0.7249	0.7659

Axis Label Font Size

18

Legend Font Size

16

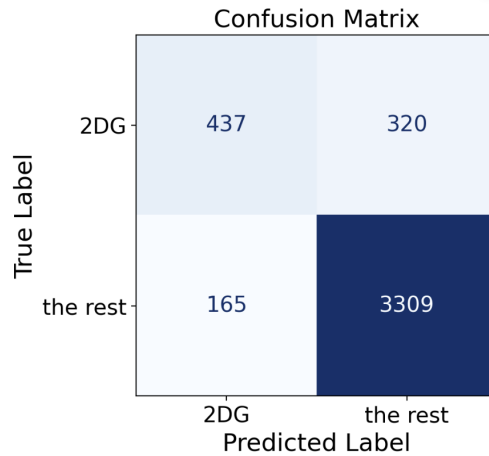
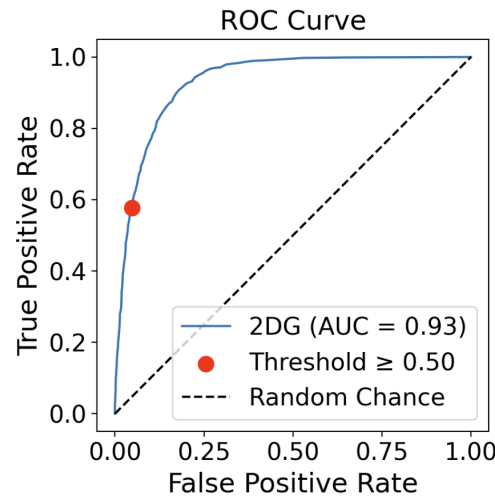


Things to **highlight**:

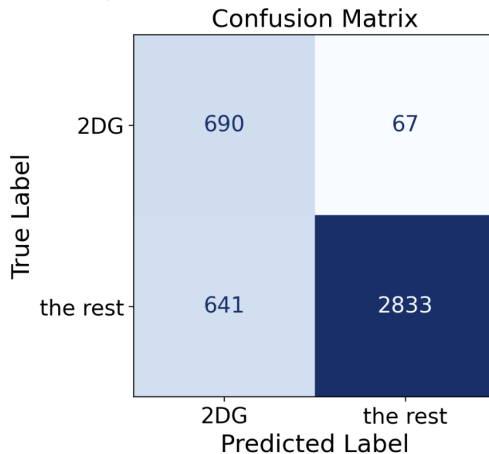
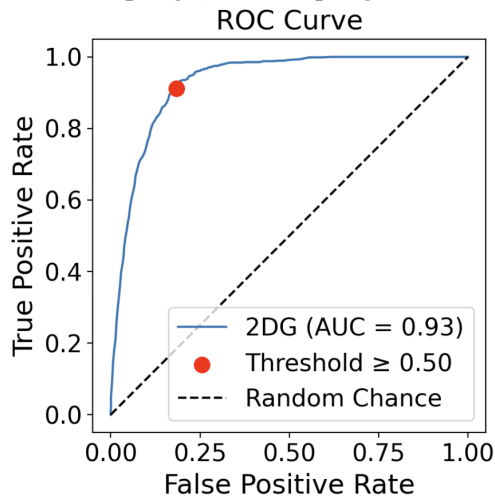
- The two ROC curves are the same, because the underlying probability models are the same.
- The annotated operating point in red is “better” *after* the threshold tuning because it achieves higher Recall (TRP) + Specificity (TNR), as can be seen in the metrics table (the optimized metrics are highlighted in red). The threshold before was 0.5, the default value, and after tuning it was optimized to 0.19. The reduction in thresholds means that the classifier is tuned to be more “confident” at predicting the positive class (2DG), which is the minority class. The effect of this threshold optimization is also visible in the confusion matrix.
- However, all of the above come at the cost of other metrics, such as the overall accuracy (0.82 to 0.82) and average precision (0.82 to 0.74). **Select the class imbalance methods/options that maximize the metrics most important to you.**

## Sampling Example

Before sampling:



After sampling (undersampling was used in this case):



Things to **highlight**:

- The ROC curve is different, because the underlying probability models are different.
- The thresholds before and after are the same (the default value of 0.5), but the operating points are at different locations and their associated confusion matrices are different.

## 19.2.2 Confusion Matrix

FLIM Playground evaluates the trained classifier on the held-out test set by converting the predicted probabilities into hard labels using the chosen threshold or by selecting the class with the highest predicted probability. The predictions are then summarized using `sklearn.metrics.confusion_matrix`. In the binary case, it is represented as:

	Predicted Positive	Predicted Negative
Actual Positive	<b>True Positive (TP)</b> – correct hits	<b>False Negative (FN)</b> – missed positives
Actual Negative	<b>False Positive (FP)</b> – false alarms	<b>True Negative (TN)</b> – correct rejections

## 19.2.3 Performance Metrics

The overall accuracy is calculated as the ratio of the sum of the true positives and true negatives to the sum of all the elements in the confusion matrix (the sum of the diagonal over the sum of all the elements).

- Accuracy:  $\frac{TP+TN}{TP+FP+TN+FN}$

Other metrics are calculated for each class based on the confusion matrix:

- Precision:  $\frac{TP}{TP+FP}$
- Recall (Sensitivity):  $\frac{TP}{TP+FN}$
- Specificity:  $\frac{TN}{TN+FP}$
- Youden's Index: Sensitivity + Specificity – 1
- F1 Score:  $\frac{2 \times \text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}}$

### 💡 Performance Metrics

Overall Metrics

Metric	Value
Accuracy	0.8854
N	4231

Per-Class Metrics

Class	N	Precision	Recall	Specificity	Youden's J	F1 Score
ZDG	757	0.7259	0.5773	0.9525	0.5298	0.6431
the rest	3474	0.9118	0.9525	0.5773	0.5298	0.9317
Average	2115.5	0.8189	0.7649	0.7649	0.5298	0.7874

### 💡 Performance Metrics Example

The reason that we need metrics in addition to accuracy is because *Accuracy is Not Enough*. For a visual explanation of metrics, see [here](#). In short, it is all about trade-off.

#### 19.2.4 ROC Curve

Another way to look at the performance of the classifier is to plot the Receiver Operating Characteristic (ROC) curve and its Area Under the Curve (AUC).

Confusion matrix looks at the performance of the classifier at a single threshold, while ROC curve looks at the performance of the classifier at all thresholds (i.e., from 0 to 1).

At the origin (0, 0), the threshold is 1, so the classifier predicts all samples as negative, making the false positive rate  $FPR = \frac{FP}{FP+TN} = 0$  and the true positive rate  $TPR = \frac{TP}{TP+FN} = 0$ , i.e. the origin. At the point (1, 1), the threshold is 0, so the classifier predicts all samples as positive, making the false positive rate 1 and the true positive rate 1, i.e. the point (1, 1).

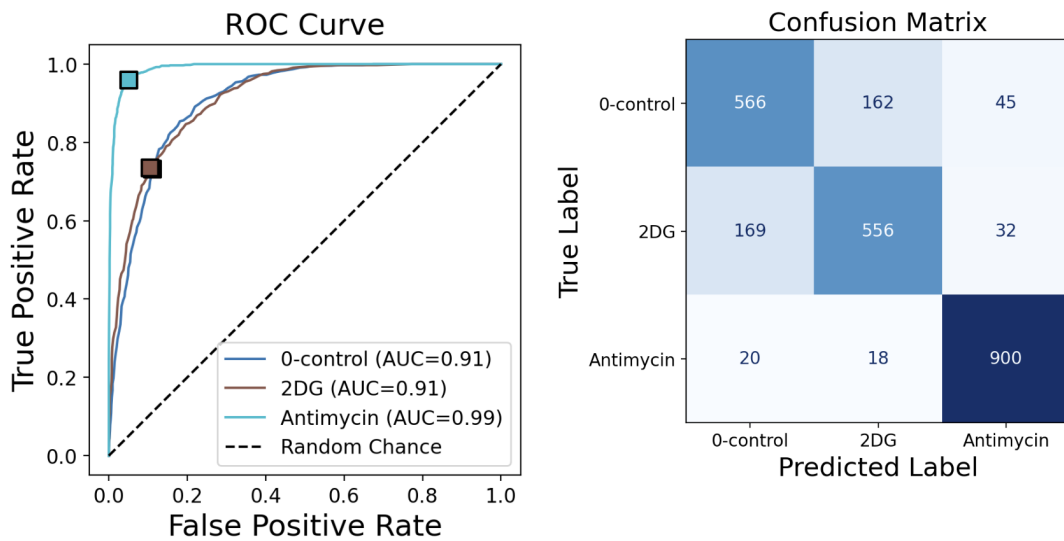
AUC summarizes the ROC curve by integrating it from 0 to 1, so it is threshold-free and has a nice interpretation: it is the probability that the classifier assigns a higher score to a randomly chosen positive sample  $X^+$  than to a randomly chosen negative sample  $X^-$ .

The ROC curve displays the operating point (FPR, TPR) associated with the confusion matrix on the right.

For a visual explanation, see [here](#).

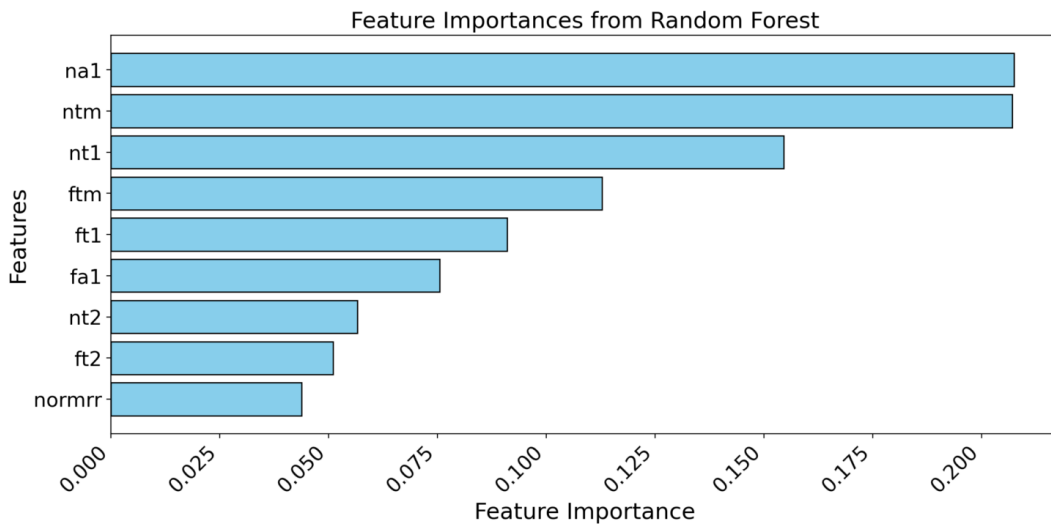
##### 19.2.4.1 Multi-class ROC Curve

In multi-class classification, each class's ROC curve is computed as a one-vs-rest binary classification problem. The operating point (FPR, TPR) associated with the confusion matrix is displayed for each class.



### 19.2.5 Feature Importance

Random Forest and Gradient Boosting classifiers provide feature importance scores.



# 20 Common Q & A

This chapter addresses frequently asked questions about FLIM Playground.

## 20.1 Feature Missing in Data Analysis

**Q:** Why I cannot find a certain **numerical feature** under numerical selection/filter widgets?

**A:** If you cannot find a certain **numerical feature**, try the following:

- look inside the **Uncategorized Features** feature group
- inspect the dataset (e.g., using Excel filters) and look for non-numeric values in that column. One non-numeric value (e.g., “–”) will prevent it from being recognized. After clearing those cells (Excel cell not real cells), saving the file, and uploading it again, the feature should be recognized.

**Q:** Why I cannot find a certain **categorical feature** under categorical filter/visual channel widgets?

**A:** - If the uploaded dataset was extracted by [Data Extraction](#), FLIM Playground will only match categorical features specified in the [Data Extraction configuration](#). Otherwise, FLIM Playground will only match categorical features specified in the [configuration](#). - If the categorical feature was in the corresponding configuration, and it is still not visible: - If you just added it, FLIM Playground may not have picked it up yet. Try refreshing the page. - If you have already added it, then there is only one unique value in that categorical feature column. FLIM Playground will not show it.

- It is ok if your dataset does not have all of the categorical features specified in either of the configurations.
1. Datta, R., Heaster, T. M., Sharick, J. T., Gillette, A. A. & Skala, M. C. [Fluorescence lifetime imaging microscopy: fundamentals and advances in instrumentation, analysis, and applications](#). *Journal of Biomedical Optics* **25**, 071203 (2020).
  2. Stringer, C., Wang, T., Michaelos, M. & Pachitariu, M. Cellpose: A generalist algorithm for cellular segmentation. *Nature methods* **18**, 100–106 (2021).
  3. Gohlke, C., Pannunzio, B., Schüty, B. & Blanco, R. Phasorpy/phasorpy: v0.7. Zenodo <https://doi.org/10.5281/zenodo.16923774> (2025).
  4. Gottlieb, D., Asadipour, B., Kostina, P., Ung, T. P. L. & Stringari, C. [FLUTE: A python GUI for interactive phasor analysis of FLIM data](#). *Biological Imaging* **3**, e21 (2023).

5. Kapsiani, S. *et al.* FLIMPA: A versatile software for fluorescence lifetime imaging microscopy phasor analysis. *Analytical Chemistry* (2025).
6. Becker, W. *The Bh TCSPC Handbook*. (Becker & Hickl GmbH, 2021).
7. Stirling, D. R. *et al.* CellProfiler 4: Improvements in speed, utility and usability. *BMC Bioinformatics* **22**, 433 (2021).
8. Stirling, D. R., Carpenter, A. E. & Cimini, B. A. CellProfiler analyst 3.0: Accessible data exploration and machine learning for image analysis. *Bioinformatics* **37**, 3992–3994 (2021).
9. Samimi, K. *et al.* Segmentation-guided photon pooling enables robust single cell analysis and fast fluorescence lifetime imaging microscopy. *bioRxiv* <https://doi.org/10.1101/2025.09.30.679660> (2025) doi:[10.1101/2025.09.30.679660](https://doi.org/10.1101/2025.09.30.679660).
10. Samimi, K. *et al.* Autofluorescence lifetime flow cytometry with time-correlated single photon counting. *Cytometry Part A* **105**, 607–620 (2024).
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14. Shannon, C. E. A mathematical theory of communication. *The Bell System Technical Journal* **27**, 379–423 (1948).
15. Pham, D. L. *et al.* Autofluorescence imaging reveals the impact of cryopreservation on t cell metabolism and activation response. *bioRxiv* <https://doi.org/10.1101/2025.11.14.688556> (2025) doi:[10.1101/2025.11.14.688556](https://doi.org/10.1101/2025.11.14.688556).