# Linear Analysis of Midgut LAM v0.2.3

User Manual

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## 1 Description

Linear Analysis of Midgut (LAM) is a tool for reducing the dimensionality of microscopy image-obtained data, and for subsequent quantification of variables and feature counts while preserving regional context. LAM's intended use is to analyze whole *Drosophila melanogaster* midguts or their sub-regions for phenotypical variation due to differing nutrition, altered genetics, etc. Key functionality is to provide statistical and comparative analysis of variables along the whole length of the midgut for multiple sample groups. Additionally, LAM has K-D tree-based functions for the estimation of feature-to-feature nearest distances and for the detection of clusters, both of which also retain the regional context (4.6). LAM also approximates the widths of the samples along the antero-posterior axis and estimates border region locations based on multivariate scoring. The analysis is performed after image processing and feature detection. Consequently, LAM requires coordinate data of the features as input.

The analysis is based on approximation of the samples from end to end through the creation of vectors onto which features can be projected by minimal distance estimation (4.4). The vectors are divided into user-defined number of bins that can be readily compared between samples and sample groups. The feature-specific point of projection along the vector is linked to all relevant input data through feature identifiers, subsequently allowing quantification of cells and their differing characteristics along the length of the sample (4.5). All individual samples' quantification data is joined with their sample groups to find the groups' general characteristics, which can then be compared group-by-group in a statistical analysis. The joining of the data is based on anchoring of the vectors to each other via a user-provided input coordinate that corresponds to a distinguishable, real biological segment of the samples, e.g. R3-region (see 4.2.4). In practice, the anchoring means that all samples are inserted into a data matrix where all anchoring points are located at the same index position, with variable lengths of the vector at either side. This is necessary due to the inherent proportional variation within the midgut that leads to a compounding error when moving further along the bins. Consequently, insertion into the matrix guarantees that the samples are in 'focus' at the anchoring site and that the correct biological regions are compared against each other.

LAM has been tested on larval and adult midguts, but usage can most likely be extended to other tissues where linear quantification could be applicable. The analysis method is for the most part restricted to planar geometries, and consequently most functionalities of LAM do not offer resolution on the Z-axis. The limitation to XY-coordinates should be sufficient for most research questions regarding the midgut, as it is pseudostratified and has only minute layering of cells.

#### **Notification:**

While the positional input data that LAM accepts can indicate any sort of feature that is to be counted and analyzed, for the purpose of the user manual individual observations will be referred to as <u>cells</u>. LAM also accepts multiple data folders for each sample, and while each of these folders does not necessarily contain data from one microscopy channel, these separate data will be referred to as channels for simplicity. Microscopy channels will be referred to as such.

## 2 Installation of Dependencies

LAM is developed in Python 3.7 environment. The distribution includes requirements.txt and environment.yml that both contain names and version numbers of LAM dependencies. The environment.yml is used to create an **Anaconda virtual environment** in anaconda prompt:

- conda env create -n <yourenvname> -f <path\to\environment.yml>
  - E.g. conda env create LAM -f C:\LAM-master\LAMenv.yml
- conda activate <yourenvname>

while replacing all text with "<>" to desired environment name and the path to the file. Alternatively, the two required packages can be installed directly to Anaconda base environment (see box below).

If creating a **python virtual environment**, the needed packages can be directly installed by using *pip* with the requirements.txt in the terminal:

- 1. python -m venv <yourenvname>
  - Linux:
- source <yourenvname>/bin/activate
- Windows:
  - <yourenvname>\Scripts\activate.bat
- 2. pip install -r <path-to-requirements.txt>
  - On Windows you need to install Shapely separately (https://pypi.org/project/Shapely/). You can either remove shapely from the requirements.txt or add '#' in front of the line to pass it, in order to install all other necessary dependencies.

### Installation to Anaconda base environment

- 1. Install Anaconda3 distribution (<a href="https://www.anaconda.com/distribution">https://www.anaconda.com/distribution</a>/)
- 2. Add dependencies

Open Anaconda Prompt and write command:
 conda install –file <LAM-master\requirements.txt>
You may need to add conda-forge to channels:
 conda config –add channels conda-forge

<u>Dependencies</u>: matplotlib (3.1.3), numpy (1.18.1), pandas (1.0.1), pathlib2 (2.3.5), scipy (1.4.1), seaborn (0.10.0), shapely (1.7.0), scikit-image (0.16.2), scikit-learn (0.22.1), statsmodels (0.11.0)

## 3 Usage in brief

### Input

LAM accepts feature coordinates and feature-related data such as intensities in csv's and in wide-format as input. The data must be located so that each sample has a separate folder named as "<sample group>\_<sample name>", e.g. "holidic\_sample1". In the sample-folders each set of features are to be in their own folders named as "<sample group>\_<sample name>\_<feature set>\_<xyz>", e.g. "holidic\_sample1\_DAPI\_statistics". The coordinate data for each feature set must be in a file named "Position.csv" with columns "Position X", "Position Y", and "Position Z". All other data to be included to the analysis must be defined in *AddData*-variable in settings.py or in the GUI's *Other*-window. If using alternative anchoring of samples, i.e. the data of samples are to be adjoined at a specific point and not bin-to-bin, the user must also provide the anchoring point for each sample similarly to other feature sets, but with the folders named as the value in *MPname*-variable ("*MP label*" in GUI, default "MP").

Additionally, you can provide self-made vector coordinates for each sample to "./Analysis Data/Samples/<sample group>\_<sample name>/" named as "Vector.csv" or "Vector.txt".

### Analysis in GUI

With default settings, the GUI is launched by running "LAM-master/src/run.py". First, provide path to the directory with the sample data by clicking the *directory*-button at the top of the GUI. If the path is valid, LAM should update all sample groups and channels (i.e. feature sets) next to the *Detect*-button. In the upper part of GUI are the primary settings: **Process**, **Count**, **Distance**, **Plots**, and **Stats**. Ticking the boxes enables related options. The **Process**-setting creates vectors for the samples, **Count** projects features to vectors and counts them, **Distance** allows for feature-to-feature distances and clustering, and **Plots** and **Stats** enable plotting and statistical calculations, respectively. Some related settings are hidden behind the "Other/Plots/Stats"-buttons at the bottom of the GUI.

If the user has provided vector-files, the **Process**-setting can be left unticked. The **Count**-setting must be performed, but the rest are optional. Once **Count** has been performed once, the optional primary settings can be run at will unless changes have been made to samples or underlying data. All output by LAM will be created to "*Analysis Data*"-folder that is created to the root of the analysis directory. Most produced data can be found in "*Data Files*"-folder, but data of each sample's each feature set can be found in the Samples-folder.

### Vector creation in LAM

For the automatic vector creation, the samples must have been oriented along the x-axis of the image so that either end of the intestine has the lowest x-values (recommendation is anterior). As a rule, the *Median vector creation* functions best for samples that have no backwards movement along x-axis, while skeleton vector creation functions for curved intestines. Intestines that loop over itself should not be used for analysis. The user should check quality of created vectors from plots created to the Samples-folder. It is often appropriate to do multiple vector creations with different settings, at the same time collecting fitting vectors after each run. When proper vectors have been collected for all samples, the vector files can be moved to the Samples-folder overwriting the files from the last run of **Process**.

## 4 Usage in full

LAM is used by executing run.py in command line or in an integrated development environment (IDE). The execution by default opens the graphical user interface (GUI) which allows the use of all functionalities of LAM. Most analysis settings are handled through settings.py, but certain arguments can be parsed from command line: use 'python run.py -h' for help or refer to docs\CommandLine\_Args.txt. When executing from command line, the arguments described as toggles will switch the default value in settings.py to the opposite Boolean value, e.g. argument -C switches clustering from default value False to True.

The workflow is as follows:

O. (Feature detection from microscopy images) (user)

1. Organizing files for input (user)

2. Creation of sample-specific vectors (automated or user)

3. Gathering of data and projection onto vector

4. Determining of sample width

5. Calculation of cell numbers and additional data

6. Finding nearest cells and clusters

7. Border region detection

8. Calculation of statistics

9. Plotting

The analysis requires a specific folder and file hierarchy, all of which need to be named respecting a convention in order to provide information for the program (see 4.2.1). The sample vectors that are used to reduce the dimensionality of data can be either made by the user or alternatively by LAM (see 4.4). The rest of LAM's functionalities are performed automatically in accordance to user-defined settings.

## 4.1. Image pre-processing

To assure the best functioning of LAM, certain pre-processing steps are required before feature detection and subsequent data feeding to LAM. Of these, the masking and orienting of the microscopy images is paramount. The vector creation also functions best with straightened midguts; inordinate curvature may cause premature end of vectorization or other artifacts. The samples are to be masked so that no excess features are present, i.e. the image contains only intensities from the midgut. When using automatic creation of vectors, the samples should also be oriented so that either end of the midgut has the smallest X-coordinate value. This orientation must be the same between samples of an experiment, e.g. all samples have the anterior end at the lowest X-axis values. The orientation is necessary in order to create proper vectors and to provide comparable quantification. LAM also includes a script to rotate feature coordinate csv's around the origin, if necessary (comp/rotator.py). When providing user-made vectors, the coordinates must be provided in the same orientation, e.g. from anterior end to posterior

### 4.2. Input

Originally, LAM was designed to be an extension to Imaris-analyses and consequently exported data files from Imaris's feature detection and other tools are in a format that is immediately usable by LAM. However, LAM only requires that the data is numerical, in csv-file format, and with certain column labels.

The main functionality of LAM, i.e. the locational quantification of cells, requires only X- and Y-coordinates of cells within a csv-file with all variables on columns and each cell on separate row, i.e. in wide-format. For finding nearest cells and clusters, the Z-coordinate of the cells is also required. Any additional data that is to be analyzed must be defined in the settings ('AddData') and can either be located within the same csv-file with the positional data or separately. All samples are not expected to have data from all microscopy channels; LAM only adds the data to analysis if it is found.

The vectors are created based on the positional data of the 'vector channel', defined by the given value in 'Channel'-setting ('vectChannel' in settings.py). The creation begins from cells with the lowest X-coordinates, and consequently all samples are expected to be oriented the same in the coordinate system. In case of wrongly oriented samples, a coordinate system-rotation script (rotator.py) can be found in the 'comp'-folder of LAM.

#### **Notification:**

Required **sample size** for each sample group depends on multiple factors including the strength of the effect caused by the experiment, but also how variable the response is. Exact sample size is difficult to determine beforehand, but **we recommend at least 10 samples per group**.

### 4.2.1. File organization and naming

LAM expects to find all input data in a certain hierarchy of directories and with specific filenames that contain necessary information (Fig. 1). As LAM scans the directory for files and folders without user-defined samples, sample groups, or channels, the recommendation is not to have any unrelated files or folders within the input data-directory, as this can break the analysis. LAM automatically gathers the names of the samples, groups, and channel-names from the paths of the files.

The full naming convention for the paths in the analysis is as follows:

```
<group>_<<descriptor>_<sample>>_<channel>_<xyz>,
```

where group denotes the sample group, descriptor and sample are used to identify individual samples (e.g. date of imaging and sample name), and channel is an identification for a data folder. The descriptor and sample can also be replaced with just one identifier. The "xyz" at the end can be any string of text and is not used by LAM; it is present for convenience, as e.g. Imaris-exported data includes "\_Statistics" at the end.

Within the analysis directory, the files and folders for the analysis must be organized in the manner shown by Fig. 1. All samples should be located at the root of the analysis directory in separate folders named as <group>\_<descriptor>\_<sample>, e.g. Control\_09-06-

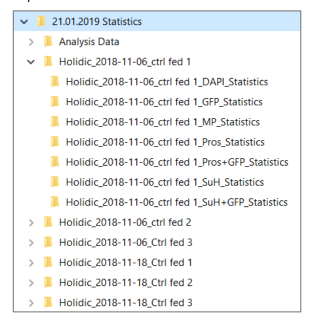


Fig. 1. Analysis directory hierarchy. All sample folders are to be located in the root of the analysis directory. Within the sample folders, each channel must have its own folder that contains all channel-specific data files. Additionally, the folders must be named consistently and according to a convention.

2019\_sample1. Within these sample folders, each sample should have a separate channel folder for each channel it has data for, named with the before mentioned name extended with <channel>\_xyz, e.g. Control\_09-06-2019\_sample1\_DAPI\_Stats. Each data file that relates to the specific channel and sample must be found within these channel folders.

Underscore '\_' is used as a delimiter for information, and is consequently reserved for LAM. Any use of underscore by the user will likely interfere with the analysis. Naming should be restricted to letters [A-Z], numbers [0-9], and plus and minus sign [+ -].

### 4.2.2. Data file column labels

Within the data files, specific column headers are required. In the Position.csv, the coordinates of cells should be marked with column labels 'Position X', 'Position Y', and 'Position Z'. Additionally, each cell should have its own unique identification number in a column labeled as 'ID'. This cell ID should be unique within the sample and shared between all data files to correctly associate data. If the data files contain for example metadata-headers, the 'header\_row'-setting can be changed to point at the correct row with column labels, with row numbering starting from zero. LAM expects all user input data files to have the same header row index number.

All data specific to a sample must be located in its sample-specific folder at the root of the analysis directory. Similarly, all data related to one channel must be in the specific channel-folder of the respective sample folder.

Position X,Position Y,Position Z,Unit,Catego	ry,Collection,Time,ID,
4735.07,390.268,0.7,um,Spot,Position,1,0,	
4741.71,397.542,0.7,um,Spot,Position,1,1,	
4726.84,399.752,0.7,um,Spot,Position,1,2,	
4705.06,405.75,0.7,um,Spot,Position,1,3,	
4698.85,411.068,0.7,um,Spot,Position,1,4,	

Fig. 2. Example of Positions.csv column labels and data in Imaris–format. LAM requires the positional coordinates in columns labeled with 'Position X', 'Position Y', and 'Position Z'. Additionally, the file must contain 'ID'–column where each cell of the channel has a unique identifier that is used for merging data. Of these column labels, the unit, category, and collection are not used by LAM.

### 4.2.3. Additional data

LAM considers all data in addition to the positional coordinates and ID as 'additional data', e.g. area, volume, and intensities. These data files must be defined within the 'AddData'-dictionary variable in settings.py, or alternatively in the 'Other'-window if using the GUI. Additional data can be in the Position.csv, but its location must be given in the variable.

In the case of additional data, its key (in bold below) in the 'AddData' dictionary defines the required column label for the data:

AddData = {"**Area**": ["Area.csv", "Area, \$\u03BCm^2\$"]}.

The values given to the key, in order, are the name of the data-containing file, and the unit of the variable. Any special characters should be given in Unicode and between '\$'-signs. The unit is only used for plots.

When inputting a data type that has multiple values, e.g. intensities from each microscopy channel, LAM expects the column labels to have an identifier separated by underscore, e.g. "Intensity Median\_Ch=1" or "Intensity Median\_ch2". The identifiers do not need to be defined for LAM as they are only used to separate the data from each other, but they can be defined in order to change them to a form that is more informative, e.g. from Ch=4 to DAPI. The replacement can be done with 'replace file ID'-option in GUI's Other-window (replaceID and channeIID in settings.py).

It should be noted that internally LAM uses '-' as separator instead of '\_' when handling identifiers of additional data with multiple column labels, such as intensities from various microscopy channels.

### 4.2.4. Sample anchoring (MP)

On some experiments the size proportions of different regions may alter, e.g. when comparing starved and fully fed midguts. In these cases, results that are more accurate can be obtained by dividing the image/data into multiple analyses. A typical way to do this is to run separate analyses for R1-2, R3, and R4-5. Alternatively, a user-defined coordinate (MP = measurement point) at a distinguishable point can be used to anchor the individual samples for comparison. For example, MP at R2/3-border of each sample causes them to be lined at a specific bin, with each sample having variable numbers of bins on either side (Fig. 3). The proportional variation however likely leads to a compounding error as distance from the MP grows. When MP is not used, the samples are lined at bin 0 and compared bin-by-bin. The MP-input is given similarly to channel data, i.e. as a separate directory that contains position.csv for a single coordinate, the MP. The anchoring is controlled by the 'useMP'-setting.

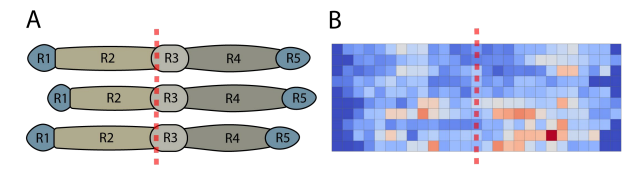


Fig. 3. Sample anchoring using Measurement Points. Due to possible proportional variation between sample groups, LAM has the option to anchor samples based on user-given coordinates. The anchoring coordinate is called a measurement point, or MP. When using MPs, the samples are lined so that every MP is located at same bin index. As a result, the samples are in 'focus' at the given positions and comparison between samples and sample groups gives biologically relevant data. a) MP marked by the red dashed line. Each sample has the same number of bins, i.e. 100% of user-defined bins, but after sample anchoring the total length of the dataset can be greater than 100% due to proportional variation. b) Heat maps of real data, with each row representing one whole-midgut sample. MP marked by the black dashed line. Shared characteristics between the samples are most clearly seen near the MP-bin.

### 4.3. Primary functionalities

LAM has several primary functionalities:

- Process
- Count
- Distance
- Plots
- Stats

The *Process*-setting creates the vectors, while *Count* projects cells from all found channels and counts their numbers. The output files from *Count* are required for the rest of the functionalities. Making of plots and statistical calculations are controlled via the *Plots* and *Stats* settings. The *Distance* setting is used for the calculation of distances to nearest cells and for finding clusters of cells. In the GUI, clicking any of these primary settings will enable or disable input for the related settings.

**NOTE:** using *Count* clears the 'Analysis Data'-folder that holds many LAM-created data files. Move any earlier data you may want to keep.

### 4.3.1. Default values

The default values for the GUI (and the analysis) are stored in settings.py and can be altered at will. Any changes to the settings within the GUI will revert to the default values when the program is run again.

### 4.4. Vector creation

The vector creation of LAM is limited to planar geometries by the *Shapely* package, i.e. only XY-coordinates are used. The limitation should not cause variation in the counting of cells as the midgut is pseudostratified and there is little Z-coordinate difference between cells, assuming the sample is flat when imaged.

The vectors for the samples can be created automatically in two ways: by a running median or by skeletonization. Both methods have their own benefits and drawbacks (4.4.1). The whole analysis can be performed in one run; however, the best and most accurate results can typically be obtained by subjecting the samples to multiple rounds of vector creation and then passing only the well-fitting vectors for further analysis.

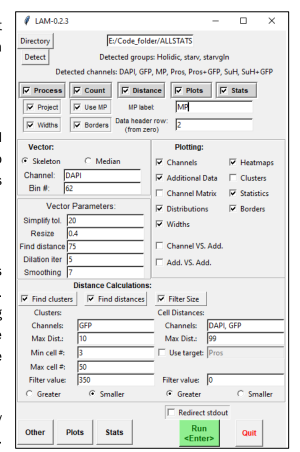


Fig. 4. LAM graphical user interface. The five primary functionalities of LAM (dark boxes) control which settings are enabled. Use of 'Count' requires that either 'Process' has been run or user has provided self-made vector csv's. The three other functionalities require that 'Count' has been performed.

A typical workflow starts by selecting only the *Process*-setting to create vectors for all of the samples. After each round of vectorization, the quality of each sample's vector can be verified from the vector plots

located at './Analysis Data/Samples', and when deemed fit, the respective LAM-created sample folders containing the vector.csv-files can be collected to be used later. Next, vector creation can be performed with other settings, again collecting acceptable vectors. Any unfit vectors can also be directly modified by changing any offending coordinates or by adding new ones to the vector.csv. Alternatively, the user can generate the vector themselves and place it in vector.csv (or .txt) (4.4.2).

Once all vectors are created, the collected sample folders can be transferred back to './Analysis Data/Samples', overwriting any non-usable data.

### 4.4.1. Vector types

Both vector creation types share two settings: the vector channel and simplify tolerance (Fig. 5). The vector channel designates the data channel that is used for creating the vector. Best approximation of the sample is achieved by using a channel that is densely populated by cells. Consequently, the recommendation is to use data files containing nuclear label data or similarly dense features.

Different methods of vector creation can lead to different kinds of artifacts in the approximation. The artifacts can often be removed with vector simplification, i.e. the straightening of the vector by moving its points. The 'Simplify tol.'—setting gives the maximal distance of the movement.

### Median vector

The median vector-creation performs ideally with fully straightened samples. However, steep curvature in the sample or any backwards movement on X-axis will cause problems for this vector type. The vector creation is simple; it is based on calculating the middle Y-axis coordinate between the top-most and bottom-most cells in each 'median bin'. The number of the bins is defined by the 'Median bins'-setting in vector parameters (Fig. 5a). Equidistant points between the lowest and highest X-coordinates of the data set's cells define the edges of the median bins. The left-most bin is assumed to be the first bin, meaning that the vectors

Vector: a) Skeleton Median DAPI Channel: Bin # 100 Vector Parameters: Simplify tol. 40 Median bins 70 b) Vector Parameters: Simplify tol. Resize 0.4 Find distance 30 Dilation iter 0 Smoothing 1

Fig. 5. Vector creation settings. a) In the upper part are the general settings for vector creation and in the lower part the median vector-related settings. b) All settings related to skeleton vector creation.

are created from left to right (smallest X-value to largest), and the samples should be oriented to account for this.

To assure the best quality of sample approximation for the vector, the number of median bins should be set high. The median bin number is ultimately limited by the density of cells on the channel that is used for vector creation; too many bins might lead to bins with no cells, in which case the previous bin's value is copied. The empty bins can consequently lead to stair-like increases in the median, but this can be remedied with vector simplification, set by 'Simplify tol.'.

### Skeleton vector

The skeleton-method offers better approximation for highly curved samples, especially if the sample has any back and forth movement on the X-axis. The downside is that some settings may sample-specifically cause branching in the skeleton, resulting in a broken vector. Recommendation is to ascertain the quality from the vector plots; there is no guaranteed warning when a vector is faulty. When using skeleton vectors, LAM produces an additional plot of the original binary image and the resulting skeleton to each sample's folder. This skeleton plot can be useful in determining how the creation settings should be changed to get a fit vector.

The creation settings for the skeleton vector are 'resize', 'find distance', 'dilation iter.', and 'smoothing' (Fig. 5b). Modifying these settings is encouraged; optimal settings are highly dependent on the shape of each sample. Most important of these are 'resize' and 'smoothing' (Fig. 6). Both of these settings remove roughness from the edges of the sample and consequently reduce branching of the skeleton. However, in samples where loops in the midgut bring the intestinal walls close together, these settings can cause the different segments to merge. In these cases, it is necessary to have a smaller size decrease and less smoothing. When any resizing would lead to merging of midgut segments, increasing the iteration number of binary dilation ('dilation iter.') may help in making the midgut uniform in the transformed binary image and consequently reduce skeleton branching.

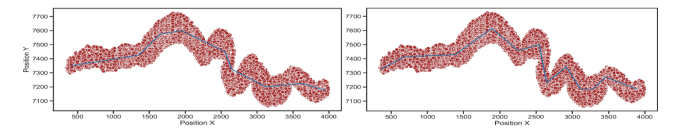


Fig. 6. Effects of resizing and smoothing on skeleton vector creation. On the left values of resizing and smoothing are 0.4 and 1, respectively. The settings cause segments of the latter half of midgut to merge, subsequently leading to a mistaken approximation and false peaks in cell counts. On the right, using a lesser reduction in size (0.7) and only a little smoothing (0.3) leads to a much-improved vector.

The transformation of the skeleton into a vector is performed by following pixels beginning from the smallest X-coordinate. From the start point, the creation algorithm adds new points to the vector by scoring pixels based on distance and the direction of the existing vector. Due to the scoring method, the vector creation can at times follow the wrong pixels if there is any branching in a steep curve. In these cases, it is necessary to reduce the branching with for example increased smoothing in order to get a fit vector.

### 4.4.2. User-generated vectors

When opting to use self-created vectors, it should be noted that the first coordinate in the file is the beginning of the first bin (i.e. bin zero) of the vector. Consequently, all vectors must be given in the same orientation, e.g. from the anterior to the posterior end of the sample. The number of the coordinates that define the vector is arbitrary, and the binning of the vector for projection of cells is done relative to the geometrical length of the vector. A convenient method for self-drawn vectors is to use one of Fiji ImageJ's line tools on the microscopy image and then saving as XY coordinates.

The user-generated vectors have to be placed similarly to LAM-created vectors, in the vector.csv's (or txt) located at respective sample-folder in './Analysis Data/Samples'. An exception to the required csv-format are the XY-coordinate files from ImageJ line tools, which can be directly used as vector files when named as 'Vector.txt'. These files contain only the X and Y coordinate values on each row, separated by tabulator ('\t').

## 4.5. Cell projection and counting

After vector creation, all channels found in the sample folders located at the root of the analysis directory are projected onto the vector if the *Count* setting is selected. For counting, the most critical value is '*Bin* #' located within the *vector*-frame in the GUI (*projBins* in settings.py). The bin number determines the length of LAM-created data arrays and consequently affects plotting and statistical analysis. All vectors are divided into the same number of bins. The length of the bins is uniform within a sample but can be variable between samples due to length differences of the vectors. Consequently, the binning functions well between sample groups that have similar proportions of different parts of the midgut. If proportional variance exists, the samples should be cropped so that the binning corresponds to the biological proportions.

When anchoring the samples at some specific point along the midgut, e.g. R3-region, the 'use MP'-option can be selected and input given to 'MP label' to collect the coordinates of the point for each sample. The coordinate must be located in each sample's folder in a channel folder named after the MP label. When 'use MP' is not selected, the samples are anchored bin to bin.

The projection is done by minimal distance estimation and consequently the cells are counted as belonging to the bin sector that is nearest to them. Data on the projection can be found within csv's named by respective channels at './Analysis Data/Samples/'. The new data columns created by the projection are 'VectPoint', 'NormDist', and 'DistBin', which are the XY-coordinates of the point of projection along the vector, the normalized distance (0–1) along the vector, and the projection's bin number, respectively.

### Cropping samples and combining subsets (split & combine)

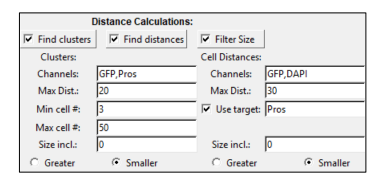
The simplest way to divide samples into comparable subsets of data is to provide cut-off coordinates for each sample in a similar manner to a MP and then use the script split\_dataset.py in 'comp'-folder of LAM. Alternatively, the script can be provided with a csv-file that contains bin-numbers of cut-off points for each sample. One way to do this is to run border detection for the whole midgut samples and determining the cut-off points for each sample from the output files. In practice, if proportional variance exists at R4-5 region between the sample groups, each sample's data could include as a channel an individual coordinate point that corresponds to R3/4 border that can be used to split all of the sample's data at the point of projection using the split\_dataset-script. The script also provides bin length suggestions for each subset of the split data in order to preserve a more standardized biological length of each bin. Afterwards, each subset can be analyzed separately. As a note, when analyzing the subsets, the 'header\_row'-setting must be set to zero.

Another way of doing the following analysis is to only perform *Count* for each of the data subsets, and later combining the sets with the companion script CombineSets.py. Combining the datasets in effect rejoins all data while retaining the refined binning of the subsets for further analysis, e.g. statistics and plotting of the full-length midgut. Note that after using *Count* on the subsets, all cells have already been projected before combining, and consequently the subsequent *Count* for the whole data set should be performed with 'project' set to False in order to preserve projection data of each subset. The combining of the data breaks the equality of bin lengths within a sample but will align the sample groups in a way that is more biologically relevant if done properly.

### 4.6. Distance calculations

The two functionalities controlled by the *Distance* -setting, i.e. the computation of feature-to-feature nearest distances and the detection of clusters, function based on K-Dimensional tree calculations. Only the smallest distance is used when computing nearest distances, but when finding clusters, neighborhoods of cells are grouped to form 'seeds' that are later merged based on shared cell ID's in order to form the final clusters.

Both functionalities are controlled individually, but they share similar settings (Fig. 7). Both can be performed on multiple channels within the same analysis, set by 'Channels'. The 'Max Dist.' limits how distant cells are considered neighbors. The cells can also be filtered by a variable of choice such as volume ('Other'-window in GUI or 'incl\_col' in settings), including cells with either smaller or larger value than the value set in 'Size incl'. In case of feature-to-feature distance calculation, the distance can be set to be calculated against a specific target channel, defined by 'Use target'. With this setting, the user can for example find nearest Pros-positive cells for each GFP-positive cell. When finding clusters, the user can define cell number limits for what is considered a cluster by modifying the 'Min cell #' and 'Max cell #'-settings.



*Fig. 7. The settings for distance calculations.* The distance calculation performs two separate functionalities, i.e. the detection of clusters and finding of nearest cells.

Both functionalities create new data columns to the channel csv's in the './Analysis Data/Samples/'-folders. The 'Find distances' creates two columns, labeled 'Nearest\_ID' and 'Nearest\_Dist', which are the ID of the nearest feature and the distance to it, respectively. The 'Find clusters' creates only one column labeled 'ClusterID', which is a unique identifier that is shared between members of same cluster.

### 4.7. Width estimation

The estimation of sample width is performed on the vector channel ('vectChannel') and is controlled by the 'Widths'-checkbox in the GUI or the 'measure\_width'-setting in settings.py. Additionally, LAM can create plots of the widths by checking the 'Widths'-box in the Plotting-frame of GUI or by setting 'plot\_width' to True in the settings. The width is calculated by following the vector bin-by-bin and summing the average distances to the most distant decile of cells on both hand sides of the vector. The width estimation uses twice the size of bins than the other functionalities to provide increased resolution for border detection (4.8). The estimation creates two output files: the "Sample\_widths.csv" that contains non-anchored values, and "Sample\_widths\_norm.csv" that contains sample values anchored to the MP.

### 4.8. Border detection

For the most part border detection is controlled from settings.py, but there is an ON/OFF checkbox at the upper frame of GUI and the 'Borders' checkbox found in the 'Plotting'-frame which controls the creation of border plots. The settings.py has a border detection subsection that has settings for what variables are used for scoring and what are the scoring weights of the variables: 'border\_vars' and 'scoring\_vars', respectively. In the 'scoring\_vars'-setting, adding extensions "\_diff" or "\_std" to the ends of the variable names causes LAM to compute either differentials or standard deviations of the variable for every bin along each sample's vector. On default settings, the border detection requires LAM-created width data (4.7) in addition to feature-to-feature distance data on the channel provided to the 'border\_channel'-setting (default channel is the vector creation channel) (4.6). The default values are intended for border detection on whole-midgut DAPI-data. Any setting changes regarding border detection should be made in settings.py before launching the GUI.

The found border regions of each group can also be added to other plots via the Boolean 'add\_peaks'-setting. Additionally, if 'select\_peaks'-setting is True the user will be asked to define which peaks will be plotted, otherwise all found peaks are added. The border detection algorithm will as a default produce a "All-Border\_Scores"-plot into "Analysis Data\Plots\Borders" that can be used in determining each sample groups' valid peaks.

The border detection outputs two data files: "Borders\_peaks.csv" that contains the peaks of the border scores and their prominences for each sample group, and "Borders\_scores.csv" that contains the border score of each bin of each sample.

### 4.9. Statistics

With the selection of **Stats**-setting, LAM can compute two different kinds of statistics, i.e. total statistics and versus statistics. The total statistics create comparisons of sample group data regarding total cell numbers on each channel and averages of additional data in each bin of sample groups. The versus statistics are bin-to-bin comparisons of data between the control group and a test group and are channel-specifically calculated for cell counts and additional data, e.g. for GFP cell counts or for the areas of the GFP cells. Additionally, the statistical analyses are performed for any LAM-calculated data such as clusters and feature-to-feature distances. For statistical plots to be created, both *Stats* and *Plots*-primary settings must be selected. The calculated statistical data can be found in '. /Analysis Data/Stats'-folder.

The versus statistics are calculated with Mann-Whitney-Wilcoxon U-test with corrections for continuity and multiple testing. The total statistics are done similarly, but without multiple test correction. The projection bin comparing versus statistics can be performed either bin-by-bin, or with a sliding window with adjustable leading and trailing edge sizes. On channels with low count numbers, increasing the size of the window leads to greater number of non-zero observations per test, consequently increasing its power at the cost of resolution.

### 4.10. Plotting

Plots are drawn by selecting the *Plots*–setting and all wanted plots from '*Plotting*'. All plotting excluding statistical plots can be performed in a separate run from the other functionalities of LAM; the plots are created from the data files that are found in the analysis directory. To obtain statistical plots, the *Stats*–setting must also be selected. Similarly, the 'borders'–plot option also requires 'border detection' to be in use.

The LAM-included plotting options are:

**Additional Data** – Bin-by-bin line plots of additional data, including LAM-created cell-to-cell distances. The line is formed from each sample's averages at the respective bin marked by X-axis. The band around the line indicates the standard deviation of the averages. Output files named e.g. "Additional Data – GFP"

Borders – Creates a file that contains plots regarding scores and found borders for each sample group. The 'Raw group score"-plot shows average scores of samples in the groups. To obtain the 'Smoothed mean scores'-plot, the raw scores are fitted with a curve that is promptly subtracted from the raw values to simulate more local changes in the score, which are then normalized between one and zero. As additional plots, by setting 'plot\_samples' to True in settings.py, LAM creates individual sample score plots. The "Normalized variables" contains normalized values of each used variable, from which a fitted curve is subtracted and resulting deviances are multiplied by the scoring weights to provide the 'Raw scores'-plot. The 'Smooth score differential'-plot contains the smoothed differential of the sum of raw scores. The differential of sum score corresponds better to borders marked by an expert than only sum score. Output to subfolder 'Borders'.

**Channels** – Bin-by-bin line plots of counts for each channel, with bins of the total, anchored matrix on X-axis and cell counts on Y-axis. Output file is named "Channels – All".

**Channel Matrix** – A grid of channel count data against each other, with channel count distribution on the diagonal axis. Each scatter marker on the grid represents the average cell counts of the samples in one bin on the respective channels. The bands around the regression lines indicate standard deviation. Output file: "Channels – Matrix".

**Clusters** – Creates three kinds of plots based on computed clusters: heatmaps and line plots of counts of clustered cells, and positional plots of clusters within samples. On the positional scatter plots, the tancolored markers are all cells on the channel, and each found cluster has an individual randomized color. Requires data from *Distances / Find clusters*. Output files created to "Clusters"-subfolder.

**Distributions** – Probability density distributions of bin values of channel counts and additional data, including LAM-created cell-to-cell distances. Output files named e.g. "Distributions – Additional GFP Data".

**Heatmaps** – Heat maps of bin-by-bin cell count averages of sample groups ("Heatmaps – Groups") and cell counts of samples on all channels ("Heatmaps – Samples").

**Statistics** – Two types of plots: total and versus plots. Total statistics creates violin plots with each sample group for every channel and additional data type. The versus statistics are plotted as bin-by-bin box plots of two sample groups, with P-value significance marked either with stars or negative log2 line plot, and with a possibility for coloring of significant bins. Additionally, the 'observations'-setting in the Plots-window can be selected to plot individual observations on top of the box plot. The Y-limit of the negative log2 line can be changed in the GUI's Plots-window or by changing 'ylim' in settings.py. The stars from one to three indicate significances of 0.05, 0.01, and 0.001, respectively. Color fill of significant regions is based on the same significances but begins from the P-value defined by 'alpha' (GUI's Stats window). The whiskers of the boxplots extend to 1.5 times interquartile range and scatters mark data points beyond this. Output to "Plots/Stats".

**Widths** – Creates a line plot that contains the average width of each sample group along the anteroposterior axis. The band around the average line is the standard deviation. Output file is named "Widths – All".

Channel VS. Add. – A bivariate density estimate of the channel counts and additional data, using averages of each bin of a sample group. Plots created for all possible combinations and may consequently create many plots depending on settings and data. The plotting combinations can be restricted by inputting only wanted data variables in GUI's Plots-window under the 'versus plots'-section ('vs\_chans' and 'vs\_adds' in settings.py), e.g. to DAPI and Intensity Mean-Ch=3. The plots represent density estimation of one sample groups values of two variables. The marginal plots are distributions of the variable on the opposing axis line. Output to "Versus"-subfolder.

Add. VS. Add. - As in 'Channel VS. Add.' but with additional data type against another type.

## 5 Output Files

LAM creates myriad of output files depending on used settings. All created files can be found in '. /Analysis Data' and its subfolders; the root of the analysis directory is read-only, except for log files. Analysis-related data is always stored as csv's, but the type of plot-files can be changed within the 'Plots'-window in the GUI or by 'saveformat' in settings.py.

Collections of the used channels, samples, and sample groups can be found at the root of the 'Analysis Data'-folder. The folder also contains the following subfolders for storing various data:

**Data Files** – For storing collections of data from samples. As a baseline, the files have either samples or sample groups in the columns and the projection bins in order as rows. The exception to this are 'Total Counts.csv' with the total cell numbers for each channel on each row, the 'MPs.csv' with the bin number of each sample's MP on the row, and 'Length.csv' with the length of each samples vector on the row.

The rows of files with the 'All\_'-prefix contain cell counts for a bin of each sample. The data with 'Norm\_'-prefix is the same data as with 'All\_', but with MP anchoring. The rows of 'Avg\_'-prefixed files contain the average values of the variable for all cells that fall in to the bin. Similarly, the rows of files with 'ChanAvg\_'-prefix contain the average channel cell counts of the sample groups for each bin. The 'Clusters' and 'ClNorm' contain the numbers of clustered cells without and with anchoring, respectively.

**Plots** – Contains all plots created by LAM, with the exception of vector-related plots that are in 'Samples'-subfolder.

**Samples –** Stores LAM-created and -edited sample-specific data. Within each sample's folder can be found the vector.csv and channel-specific csv's. The channel csv's contain all collected input data for the sample, in addition to LAM-created columns. The created columns are:

- Projection 'VectPoint', 'NormDist', and 'DistBin' are the XY-coordinates of the point of projection along the vector, the normalized distance (0-1) along the vector, and the projection's bin number, respectively.
- <u>Clustering</u> 'ClusterID' indicates the given identification number to the cluster that the respective cell belongs to.
- <u>Distance</u> 'Nearest\_XYZ', 'Nearest\_Dist', and 'Nearest\_ID' are the coordinates of the nearest cell, the distance to it, and its ID, respectively.

The root of the 'Samples'-folder also contains the plots of each sample's vector, and within the subfolders can be found the skeleton plots if using the respective vector creation.

**Statistics** – Contains all files with statistical data. Each file contains the data of bin-to-bin or windowed tests of the control group against another group. The test variable is indicated after the '='-sign in the file name, i.e. "Stats\_<test groups> = <channel> (<additional data>)".

### Column labels:

- U Score The MWW U-score of the compared populations
- Corr. <Greater/Lesser> Corrected P-value that control group is <Greater/Lesser>
- P <Greater/Lesser> Non-corrected P-value that control group is <Greater/Lesser>
- Labels with 'Two-sided' indicate the probability that there is a significance in either way

Additionally, the Statistics-folder contains the 'Total Count Stats.csv' with the two-way statistics of each channel for the test groups.

## 6 Definitions

The following definitions are named after the settings in the GUI. The corresponding variables in settings.py are marked with brackets.

## 6.1. Top Frame

GUI	Settings.py	Description
Borders	Border_detection	Calculate border region scores for samples and groups
Count	process_counts	Projection, anchoring, and counting of data on all channels
Data file	header_row	The expected row of column labels in user given data files
header row		
Distance	process_dists	Calculation of cell-to-cell distances and detection of clusters
Directory	workdir	Path to analysis directory where sample data is located
MP label	MPname	The name of the csv data files containing anchoring point locations
Plots	Create_Plots	Creation of any/all plots
Project	project	Whether to project features onto vector during <b>Count</b>
Process	process_samples	Creation of vectors for samples
Stats	statistics	Calculation of statistics
Use MP	useMP	Whether to use user given anchoring points
Widths	Measure_width	Estimate widths of each sample

### 6.2. Vector Frame

GUI	Settings.py	Description
Bin #	projBins	Number of bins onto which data is projected on all vectors
Channel	vectChannel	The channel on which vector creation is based on
Skeleton /	SkeletonVector	The type of vector to be created
Median		

## 6.3. Vector Parameters Frame

GUI	Settings.py	Description
Dilation iter.	BDiter	Iterations of binary dilation (2x2) on resized and image-transformed
		position coordinates
Find	find_dist	The maximum coordinate distance of next pixel (coordinate) of
distance		vector when creating from skeleton

Median bins	medianBins	The number of medians that are calculated for the sample when
		creating a median vector
Resize	SkeletonResize	Binary image resizing factor when creating skeleton vector
Simplify tol.	simplifyTol	Tolerance of coordinate adjustment for vector simplification
Smoothing	SigmaGauss	Sigma for Gaussian smoothing of binary image

## 6.4. Plotting Frame

GUI	Settings.py	Description
Add. VS. Add.	Create_AddVSAdd_Plots	Plot additional data against additional data
Additional Data	Create_AddData_Plots	Plot additional data averages per bin
Borders	Create_Border_Plots	Plot border scores of sample groups
Channel pair	Create_Channel_PairPlots	Create plot matrix of all channels against all channels
plots		
Channel VS.	Create_ChanVSAdd_Plots	Plot channels versus additional data
Add		
Channels	Create_Channel_Plots	Plot box plots of channel data
Clusters	Create_Cluster_Plots	Create box plots and heat maps of clustered cells
Distributions	Create_Distribution_Plots	Plot distribution densities for all channel and
		additional data
Heat maps	Create_Heatmaps	Plot sample and sample group heatmaps of channel
		counts
Statistics	Create_Statistics_Plots	Plot statistical box plots of all data

## 6.5. Distance Calculations Frame

Compared to cell-to-cell distances, the variables specific to clustering algorithm have the prefix 'Cl\_' in settings.py.

GUI	Settings.py	Description
Channels	Distance_Channels,	The channels to which distances are calculated
	Cluster_Channels	
Filter size	N/A	Whether to filter cells based on volume
Find clusters	Find_Clusters	Whether to perform clustering of cells
Find	Find_Distances	Whether to find cell-to-cell distances
distances		

Greater /	incl_type,	The direction of size filtering, includes cells of either smaller or
Smaller	Cl_incl_type	greater size
Max cell #	CI_max	Maximum cell number that is considered to be a cluster
Max Dist.	maxDist, Cl_maxDist	Maximum distance between cells to be considered neighbors.
		In clustering Max Dist is a hard distance limit for inclusion into
		a cluster.
Min cell #	CI_min	Minimum cell number of a cluster
Size incl.	Vol_inclusion,	The volume limit of filtering
	CI_Vol_inclusion	
Use target	use_target +	Calculate cell-to-cell distances from one channel to the target
	target_chan	channel

## 6.6. Other Window

GUI	Settings.py	Description
Column label /	AddData	Gathering of additional data. The label of the column where data is
csv-file /		located / the name of the file where data is / unit of the data for
Unit		plotting
Distances i. col	incl_col	Define variable for filtering during <b>Distance</b> functionalities
File descriptor /	channelID	Replace these ID's found in file names / the proper names of the ID's
Change to		
Replace file ID	replaceID	Whether to replace channel ID's found in file names

## 6.7. Plots Window

## **General settings:**

GUI	Settings.py	Description
Drop outliers	Drop_Outliers	Whether to drop data point outliers based on Std. dev
Pairplot jitter	plot_jitter	Create jitter for pair plot scatters for easier visualization of
		discretized data
Plotted add. data	vs_adds	Define additional data to be plotted in versus plots
Plotted channels	vs_channels	Define channel data to be plotted in versus plots
Save format	saveformat	The saving format of all plots
Std dev.	dropSTD	The standard deviation limit for <b>Drop outliers</b>

## Statistical Plotting:

GUI	Settings.py	Description
Sign. color	fill	Do color fill for statistically significant regions
Sign. stars	stars	Include significance stars to statistical plots
Neg. log2	negLog2	Create negative log2 significance line for statistical plots
y-limit	ylim	Y-axis value limit for <b>Neg. log2</b> in plot
Observations	observations	Plot individual observations (DEPRECATED)

## 6.8. Stats Window

GUI	Settings.py	Description
Alpha	alpha	The P-value limit for the rejection of null hypothesis
Control Group	cntrlGroup	Name of the control group
Group vs. Group	stat_versus	Create control group versus sample group statistics
Total Statistics	stat_total	Create statistics of total cell counts
Trailing /	trail / lead	Number of included bins in front and behind current index
Leading window		position when using windowed statistics
Windowed	windowed	Statistics done in a sliding window defined by <b>trail</b> and <b>lead</b>
statistics		

## 6.9. Redirect stdout

GUI	Settings.py	Description
Redirect stdout	non_stdout	Redirects output to a separate window

## 6.10. Non-GUI settings

Settings.py	Description
border_channel	Define channel name for border detection
plot_samples	Create border plots for individual samples
border_vars	Define variables that are collected for border detection
scoring_vars	Define scoring weights for border detection variables
add_peaks	Add detected border regions to other plots
select_peaks	Only subset of peaks will be plotted by 'add_peaks'
seaborn_style	Change plot style. Can break plots.
seaborn_context	Change plot element sizes. Can break plots.

Palette colors	Change sample group plot colors. Groups given color in alphabetical order.
, 4,0110_00,070	Change cample group plot colorer all cape given color in alphabetical craci-

## 7 Command line arguments

usage: run.py [-h] [-p PATH] [-o OPTIONS] [-b BINS] [-v CHANNEL]

[-g CONTROL\_GROUP] [-H HEADER] [-M] [-m MP\_NAME] [-G] [-F]

[-f DISTANCE\_CHANNELS] [-C] [-c CLUSTER\_CHANNELS]

[-d CLUSTER\_DISTANCE] [-B] [-W] [-r] [-D]

Perform LAM analysis from command line. Args described as toggle alter the default value in settings.py to the opposite Boolean. All settings that are not altered through parsed arguments will use default values from settings.py.

optional arguments:

optional arguments.	
-h,help	show this help message and exit
-p PATH,path PATH	Analysis directory path
-o OPTIONS,options OPTIONS	option string: r (process), c (count), d (distance), l
	(plots), s (stats)
-b BINS,bins BINS	Sample bin number
-v CHANNEL,channel CHANNEL	Vector channel name
-g CONTROL_GROUP,	Name of control group
control_group CONTROL_GROUP	
-H HEADER,header HEADER	Header row number
-M,measurement_point	Toggle useMP
-m MP_NAME,mp_name MP_NAME	Name of MP
-G,GUI	Toggle GUI
-F,feature_distances	Perform feature-to-feature distances
-f DISTANCE_CHANNELS,	f-to-f distance channels
Distance_channels DISTANCE_CHANNELS	
-C,clusters	Perform feature clustering
-c CLUSTER_CHANNELS,	Clustering channels
cluster_channels CLUSTER_CHANNELS	
-d CLUSTER_DISTANCE,	Clustering max distance
cluster_distance CLUSTER_DISTANCE	
-B,borders	Toggle border detection
-W,widths	Toggle width calculation
-r,no_projection	Projection to false
-D,force_dialog	Force no user input

### **Examples:**

python src\run.py -p C:\experiment\DSS -o cls -b 62 -MGD

python src\run.py --options rcd --control\_group ctrl -F -f GFP -f DAPI -C -c GFP --borders

## 8 Companion Scripts

The LAM-master folder includes several companion scripts in the 'comp'-folder that perform functionalities related to LAM. These are:

• ChannelPositionPlots.py Create plots of cell locations on different channels.

### combineSets.py

Used to combine LAM-created data sets from cropped samples.

#### rotator.py

Rotate coords around origin to properly orientate samples for LAM vector creation.

### ImarisFileConverter.ijm

Fix broken Aurox confocal images for the Imaris file converter & stitcher

ManualVectorPlots.py

Create vector plots from vector files

FileMove.py

Move and rename image files after splitting channels and focal planes (ImarisFileConverter.ijm)

split dataset.pv

Split all data and vectors in a dataset based on cut point channels

stitcher-grandma-pro.py

Stitches tiff-images in a level-plane, i.e. without jumps in z-stack.

Refer to files of individual scripts for more information.

### 9 Test Data

The distribution includes a small test data set that can be used to try LAM. The data consists of three sample groups with four samples each. The sample groups are Holidic, Starved, and StarvGln, i.e. fully-fed, fed with starvation media, and fed with starvation media with glutamine supplementation. The samples contain data from DAPI, Prospero, and Su(H) stainings, and additionally GFP lineage labelling via Esg-Flip out –system.

It should be noted that four samples per sample group is not enough for a proper analysis, and any results from these samples are only directional.

### 10 Troubleshoot

As a first rule, when you get errors and/or modify files you should always restart LAM between runs. You should also remove all folders from "Analysis Data" except for the Samples-directory that holds the vector files. Critical errors are often caused by missing data due to faulty vectors and projections. You should always check the LAM-created vector-plots in the Samples-folder, or alternatively when giving user-created vectors, use the companion-script ManualVectorPlots.py to look for any offending vectors. You can also look at files in the "Data Files"-folder to determine samples with faulty data. If you can't figure how any faulty data comes to be, easiest way to handle it is by removing any offending samples. When removing samples, you should remove the sample from both the root of the analysis directory and from the "Samples"-folder of "Analysis Data", and then restart LAM.

### The resulting cell counts have peaks at the ends of the bin range

Either the vectors are not optimally created, or the dataset has cell detection artifacts at the ends of the midgut. Make sure that both ends of the vector extend approximately to the ends of the sample. For example, when creating vectors through skeletonization, resizing of the transformed binary image may lead to incorrect approximation of the sample. As a result, the skeletonization produces a truncated vector and causes overrepresentation of cells in the last bins. Alternatively, check that cell detection was done properly.

### > The vector ends abruptly when created through skeletonization

Certain patterns in the sample may cause branching in the skeletonization. As the algorithm follows the skeletonized pixels to create the vector, sometimes these branches can confuse the algorithm into a dead-end. Increasing 'find distance' can allow the vectorization to jump back to the correct skeleton. Alternatively, trying different resizing or greater smoothing can also solve the problem. The best solution can be determined by looking at the skeleton plot in './Analysis Data/Samples/'.

### The vector jumps in an unexpected manner when created through skeletonization

Sometimes the algorithm that finds the next pixels of the binary image is fooled by branching of the skeleton. If the vector jumps into a branch that has already been passed, reducing 'Find distance' can solve the problem.

### LAM will not run because system cannot find path at start up

Modify 'workdir' variable in settings.py to point at the analysis directory. Double check that the path is written correctly and is surrounded by r"", for example r"C:\experimentDirectory".

### LAM outputs a ValueError when normalizing data

This happens when data is missing for some reason, e.g. a faulty vector for a sample or missing MP projection when using measurement points. Look at "./Analysis Data/Data Files/" for the error-producing channel's file with prefix "All\_", e.g. "All\_DAPI.csv" and find samples with abnormal values. Easiest way to handle the error is to remove the offending samples from the analysis by removing the sample from the root of the analysis directory **and** from "./Analysis Data/Samples/". If data of all samples seems fine,

there is likely an error with MP-projection. Check "MPs.csv" and look for samples with abnormal or missing values.