MTQUANT++ DOCUMENTATION

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1 MTQuant++ Overview

MTQuant++ builds on the original MTQuant by providing functions for the following:

- 1. Calculating statistical significance of different organization parameters
- 2. Splitting the axon and calculating organization parameters for each split
- 3. Microtubule Simulations

2 Setup & System Requirements

Download both MTQuant and MTQuant++ from their respective websites:

- MTQuant: http://roscoope.github.io/MTQuant
- MTQuant++: http://roscoope.github.io/MTQuant++

The code can be downloaded as a ZIP file or checked out as a Github repository. Download all of the files and keep them together. The run the following code to add all of the subdirectories to the path:

```
>> addpath('C:/path/to/MTQuant')
>> addpath('C:/path/to/MTQuant/em')
>> addpath('C:/path/to/MTQuant/misc')
>> addpath('C:/path/to/MTQuant/neuronTracing')
>> addpath('C:/path/to/MTQuant/redScan')
>> addpath('C:/path/to/MTQuant/singleMT')
>> addpath('C:/path/to/MTQuant++')
>> addpath('C:/path/to/MTQuant/misc')
>> addpath('C:/path/to/MTQuant/simulations')
>> addpath('C:/path/to/MTQuant/splitNeuron')
>> addpath('C:/path/to/MTQuant/stats')
```

The setup.m file contains these commands assuming that you are in the MTQuant++ directory, and that the MTQuant and MTQuant++ directories are in the same parent directory. Modify setup.m as necessary with the correct path names for easy startup.

If you copy and paste commands from this document into MATLAB directly, some characters, such as curly braces ({}) might not copy correctly; be sure to review copied commands before running them.

Thus far, MTQuant has been tested on Windows 7 and 8, using MATLAB versions R2014b and R2013b.

3 Calculate Statistics

There are two functions that take a CSV written by MTQuant and calculate the statistical significance of the organization parameter relationships. One is used for rank correlation (e.g., how a parameter changes as a function of age), and the other is used to compare various groups to wild-type.

3.1 compareStatsAge

This function calculates the correlation and p-value using the Spearman Rank Correlation.

3.1.1 Input & Output Arguments

| Argument | Description | | | |
|-------------------|------------------------------------------------------------------------|--|--|--|
| INPUT ARGUMENT | Input Arguments | | | |
| statsFile | the name of an output file generated by MTQuant. | | | |
| groups (optional) | a list of group numbers to compare. These are the values of the DirNum | | | |
| | column of the output file (see Section 4 of the MTQuant documentation | | | |
| | for more information). If groups is not inputted, or it is empty (i.e. | | | |
| | groups=[];), the function compares all groups. | | | |
| stats (optional) | list of columns in statsFile for which to calculate the cor- | | | |
| | relation. This can be a list of column numbers or a list of | | | |
| | strings, where each string is the column name (See Section 5.2 | | | |
| | of MTQuant documentation for complete list). By default, stats = | | | |
| | {'Avg_Spacing','Std_Dev_Spacing','Avg_Coverage','Avg_Length'} | | | |
| OUTPUT ARGUMENTS | | | | |
| allRhos | cell array with Spearman Rank Correlation for parameters in stats | | | |
| allPs | cell array containing the p-values associated with allRhos | | | |

3.1.2 Example Usage

```
>> statsFile = 'C:/Users/Roshni/Desktop/MTQuantTest/axonTest_2.csv'
>> groups = [1;2;5;6]
>> [allRhos,allPs] = compareStatsAge(statsFile,groups,stats)
allRhos =
    'Avg_Spacing''Std_Dev_Spacing''Avg_Coverage''Avg_Length'
    [ -0.1156] [ -0.1657] [ 0.7217] [ 0.7556]
allPs =
    'Avg_Spacing''Std_Dev_Spacing''Avg_Coverage''Avg_Length'
    [ 0.0709] [ 0.0094] [ 1.0424e-40] [ 1.5150e-46]
```

3.2 compareStatsToWT

This function calculates the two-sample t-test between every group of animals and wild-type.

3.2.1 Input & Output Arguments

| Argument | Description | | |
|--------------------|--------------------------------------------------------------------------|--|--|
| INPUT ARGUMENTS | PUT ARGUMENTS | | |
| statsFile | the name of an output file generated by MTQuant | | |
| groups (optional) | a list of group numbers to compare. These are the values of the DirNum | | |
| | column of the output file (see Section 4 of the MTQuant documentation | | |
| | for more information). If groups is not inputted, or it is empty (i.e. | | |
| | groups=[];), the function compares all groups. | | |
| wtGroup (optional) | The group number for the wild-type animals. If this input is not in- | | |
| | cluded, wtGroup is set to be the smallest value of DirNum in statsFile | | |
| stats (optional) | list of columns in statsFile for which to calculate the correlation. | | |
| | This can be a list of column numbers or a list of strings, where each | | |
| | string is the column name (See Section 5.2 of MTQuant documentation | | |
| | for complete list) | | |
| OUTPUT ARGUMEN | Output Arguments | | |
| allPs | cell array containing the p-values of two-way t-tests for each parameter | | |
| | in stats. N is the number of animals in each group | | |
| allHs | cell array containing the decision to reject the null hypothesis (1 for | | |
| | p-value less than 0.05) | | |
| allMeans | cell array containing the mean values of each parameter for each group | | |
| allStds | cell array containing the standard deviation of each parameter for each | | |
| | group | | |

3.2.2 Example Usage

```
>> statsFile = 'C:/Users/Roshni/Desktop/MTQuantTest/axonTest_2.csv';
>> groups = [3;4];
>> wtGroup = 2;
>> stats = {'Avg_Spacing'; 'Avg_Coverage'; 'Avg_Length'};
>> [allPs,allHs,allMeans,allStds] = compareStatsToWT(statsFile,groups,wtGroup,stats)

allPs =
    [] 'N''Avg_Spacing''Avg_Coverage''Avg_Length'
    'L4 17-4'[45] [ 0.7808] [ 0.2911] [ 0.8126]
    'L4 arl-8'[41] [ 0.2968] [ 0.1671] [ 0.0194]

allHs =
    [] 'N''Avg_Spacing''Avg_Coverage''Avg_Length'
```

```
'L4 17-4'[45] [ 0] [ 0] [ 0]
'L4 arl-8'[41] [ 0] [ 0] [ 1]

allMeans =
    [] 'N''Avg_Spacing''Avg_Coverage''Avg_Length'
    'L4 17-4'[45] [ 12.3305] [ 1.7442] [ 21.7853]
    'L4 arl-8'[41] [ 11.4005] [ 1.7129] [ 19.9497]

allStds =
    [] 'N''Avg_Spacing''Avg_Coverage''Avg_Length'
    'L4 17-4'[45] [ 1.7165] [ 0.3266] [ 2.6545]
    'L4 arl-8'[41] [ 1.4945] [ 0.3592] [ 3.2746]
```

Split the Neuron 4

4.1 splitNeuron

This function takes every line scan file catalogued in an output file of MTQuant and calculates the organization parameters for each segment of the neuron.

Input & Output Arguments 4.1.1

| Argument | Description | |
|----------------------------|-------------------------------------------------------------|--|
| NPUT ARGUMENTS | | |
| statsFile | the name of an output file generated by MTQuant | |
| splitSegLen | length of each segment in the split in pixels (not microns) | |
| groups (optional) | a list of group numbers to compare. These are the values | |
| | of the DirNum column of the output file (see Section 4 | |
| | of the MTQuant documentation for more information). If | |
| | groups is not inputted, or it is empty (i.e. groups=[];), | |
| | the function compares all groups. | |
| rPeakTol (optional) | See MTQuant documentation | |
| toUseRandRPeaks (optional) | See MTQuant documentation | |
| rPeakCorr (optional) | See MTQuant documentation | |
| toUseHalfMTs (optional) | See MTQuant documentation | |
| OUTPUT ARGUMENTS | | |
| fileOut | name of the output file containing the split information. | |
| | It is the same as statsFile, but with the string '_split' | |
| | appended to it. The first three columns of the output file | |
| | are the same as the first three columns of statsFile. | |
| | The remaining columns are the average spacing, std dev | |
| | of spacing, single MT brightness, average coverage, std | |
| | dev of coverage, and average length for each segment. | |
| | The column names are the same as in statsFile, but | |
| | with $'S*'$ prepended to each, where the askterisk (*) is | |
| | the segment number. The segment number 1 refers to the | |
| | segment with the green star in Figure 1 in the MTQuant | |
| | documentation. | |

4.1.2 Example Usage

```
>> statsFile = 'C:/Users/Roshni/Desktop/MTQuantTest/axonTest_2.csv';
```

>> splitSegLen = 15/0.170;

>> fileOut = splitNeuron(statsFile,splitSegLen);

4.2 splitAxonComm

This function is almost identical to splitNeuron, but it also creates segments for the axon close to the cell body and the commissure. It expects the line scan files to be '*_acbLineScans.csv', '*_commLineScans.csv', and '*_axonLineScans.csv' for the line scans around the cell body, of the commissure, and the remainder of the axon, respectively.

4.2.1 Input & Output Arguments

| Argument | Description |
|----------------------------|--------------------------------------------------------------|
| Input Arguments | |
| statsFile | the name of an output file generated by MTQuant |
| splitSegLen | length of each segment in the split, in pixels (not microns) |
| groups (optional) | a list of group numbers to compare. These are the values |
| | of the DirNum column of the output file (see Section 4 |
| | of the MTQuant documentation for more information). If |
| | groups is not inputted, or it is empty (i.e. groups=[];), |
| | the function compares all groups. |
| rPeakTol (optional) | See MTQuant documentation |
| toUseRandRPeaks (optional) | See MTQuant documentation |
| rPeakCorr (optional) | See MTQuant documentation |
| toUseHalfMTs (optional) | See MTQuant documentation |
| OUTPUT ARGUMENTS | |
| fileOut | name of the output file containing the split informa- |
| | tion. It is the same as statsFile, but with the string |
| | '_splitComm' appended to it. The first three columns |
| | of the output file are the same as the first three columns |
| | of statsFile. The remaining columns are the average |
| | spacing, std dev of spacing, single MT brightness, av- |
| | erage coverage, std dev of coverage, and average length |
| | for each segment. The column names are the same as |
| | in statsFile, but with 'S*_' prepended to each, where |
| | the askterisk (*) is the segment name or number. The |
| | segment number 1 refers to the segment with the green |
| | star in Figure 1 in the MTQuant documentation. |

4.2.2 Example Usage

```
>> statsFile = 'C:/Users/Roshni/Desktop/MTQuantTest/axonTest_2.csv';
```

>> splitSegLen = 15/0.170;

>> fileOut = splitAxonComm(statsFile,splitSegLen);

4.3 splitNeuronEM

This function takes every segment from a '*_split.csv' file and applies the distribution refinement to every organization parameter, DirNum, and segment combination.

4.3.1 Input & Output Arguments

| Argument | Description | | |
|-----------|--------------------------------------------------------------------------|--|--|
| INPUT ARG | NPUT ARGUMENTS | | |
| statsFile | the name of a '*_split.csv' of a '*_splitComm.csv' output file gen- | | |
| | erated by splitNeuron or splitAxonComm | | |
| OUTPUT A | RGUMENTS | | |
| fileOut | name of the output file containing the split information. It is the same | | |
| | as statsFile, but with the string '_em' appended to it. The first three | | |
| | columns of the output file are the same as the first three columns of | | |
| | statsFile. The remaining columns are the average spacing, std dev of | | |
| | spacing, single MT brightness, average coverage, std dev of coverage, | | |
| | and average length for each segment. The column names are the same | | |
| | as in statsFile, but with 'S*_' prepended to each, where the askterisk | | |
| | (*) is the segment name or number. The segment number 1 refers to the | | |
| | segment with the green star in Figure 1 in the MTQuant documentation. | | |

4.3.2 Example Usage

```
>> statsFile = 'C:/Users/Roshni/Desktop/MTQuantTest/axonTest_2_split.csv';
>> fileOut = splitNeuronEM(statsFile);
```

4.4 splitNeuronPlots

This function takes a '*_split.csv', '*_splitComm.csv', '*_split_em.csv', or '*_splitComm_em.csv' file and generates multple plots of the organization parameters over the segments in the file. For every parameter specified by the input stats, and for every animal group specified by the input groups, the function generates two plots. One is the average value of the parameter over each segment for that group with error bars. The second plot is the parameter for each animal over each segment. The plots are saved with different suffixes added to the string in fileOut.

4.4.1 Input & Output Arguments

| Argument | Description | | |
|-------------------|------------------------------------------------------------------------|--|--|
| INPUT ARGUMENT | PUT ARGUMENTS | | |
| statsFile | the name of an output file generated by MTQuant | | |
| fileOut | the base name of the output plots. The names of the parameters and/or | | |
| | the numbers of the groups are added to fileOut and the plots are saved | | |
| | as .BMP files. | | |
| groups (optional) | a list of group numbers to compare. These are the values of the DirNum | | |
| | column of the output file (see Section 4 of the MTQuant documentation | | |
| | for more information). If groups is not inputted, or it is empty (i.e. | | |
| | groups=[];), the function compares all groups. | | |
| stats (optional) | list of columns in statsFile for which to calculate the correlation. | | |
| | This can be a list of column numbers or a list of strings, where each | | |
| | string is the column name (See Section 5.2 of MTQuant documentation | | |
| | for complete list) | | |

4.4.2 Example Usage

>> splitNeuronPlots('C:/Users/Roshni/Desktop/MTQuantTest/axonTest_2_split.csv')

5 Simulations (runModel)

To run the model, use the function runModel. This function accepts as input the number of times to run the model, i.e. re-simulate the MTs and analyze them, and the scaling factors by which to change the spacing and the length. It also accepts the same name-value parameters that would be sent to MTQuant in order to see the results of the model with various settings in the algorithm. The function generates numRuns MTs and corresponding blurred, noisy line scans. It saves those line scans into 'LineScans.csv' files in a temporary directory created in the location from which runModel is being called. Then MTQuant is called, on the line scans in that temporary directory. That temporary directory is deleted at the end of the function.

5.1 Input & Output Arguments

| Argument | Description | | |
|-------------|----------------------------------------------------------------------------|--|--|
| INPUT ARGUM | INPUT ARGUMENTS | | |
| numRuns | The number of times to run the model | | |
| spcFactor | The factor by which to multiply the randomly drawn spacing of each | | |
| | MT. If in doubt, set this input to 1. To increase coverage, set this input | | |
| | to something less than 1 | | |
| lenFactor | The factor by which to multiply the randomly drawn spacing of each | | |
| | MT. If in doubt, set this input to 1. To increase coverage, set this | | |
| | number of something greater than 1 | | |
| MTQuantArgs | Cell array of strings containing a list of name-value pairs to send to | | |
| (optional) | MTQuant | | |
| OUTPUT ARGU | UMENTS | | |
| modelB | Randomly selected single MT brightness for each run of the simulation | | |
| modelS | Actual spacing for each model run, extracted from generated MTs | | |
| modelC | Actual coverage for each model run, extracted from generated MTs | | |
| modelL | Actual length for each each model run, extracted from generated MTs | | |
| calcB | Single MT brightness for each model run, calculated with MTQuant | | |
| calcS | Spacing for each run of the simulation, calculated using MTQuant | | |
| calcC | Coverage for each run of the simulation, calculated using MTQuant | | |
| calcL | Length for each run of the simulation, calculated using MTQuant | | |

5.2 Example Usage

```
>> MTQuantArgs = {'toEM',false,'rPeakTol',0.01};
>> [modelB,modelS,modelC,modelL,calcB,calcS,calcC,calcL] = ...
    runModel(100,1,1,MTQuantArgs);
```