**The *Fluocell* Image Tool**

**Introduction**

***Fluocell*** is a software package for the analysis and visualization of fluorescence images for live cells, with applications in migration and cancer invasion. The main package contains a visualization and quantification tool for fluorescence intensity and FRET ratio images. Optional modules of diffusion analysis and polarity analysis can be obtained upon request. This package is mainly written in MATLAB. It is being developed by a group of researchers at the University of California, San Diego, and the University of Illinois, Urbana-Champaign. As developers, we are open for collaboration on using the software or developing additional features. If you find this software package of interest or useful for your work, please take the time to cite our work and write an email to tell us, since positive feedbacks are important to secure continuous support for this software package.

Citations for publication

1. Lu S et al. 2008 PLoS Computational Biology, The spatiotemporal pattern of Src activation at lipid rafts revealed by diffusion-corrected FRET imaging.
2. Lu S et al. 2011 PLoS ONE, Computational analysis of the spatiotemporal coordination of polarized PI3K and Rac1 activities in micro-patterned live cells.
3. Lu S et al. 2014 Scientific Reports, Decipher the dynamic coordination between enzymatic activity and structural modulation at focal adhesions in living cells.

Contact

Shaoying (Kathy) Lu: kalu@ucsd.edu

***Download the Source Code and Imaging Data***

1. Download the fluocell source code from here: <http://github/lu6007/p/fluocell>
2. Download the fluocell dataset from here: <http://wang.ucsd.edu/~kalu/fluocell_dataset/fluocell_sample.zip> , unzip, and the install it to a folder named “fluocell\_sample/” which is readable and writable by users. The relative location of the dataset README file from the fluocell root directory is: fluocell\_sample/README.txt

***Visualize the ECFP/FRET Ratio Images***

For instruction to install fluocell, refer to the “Installation and Usage” section for details.

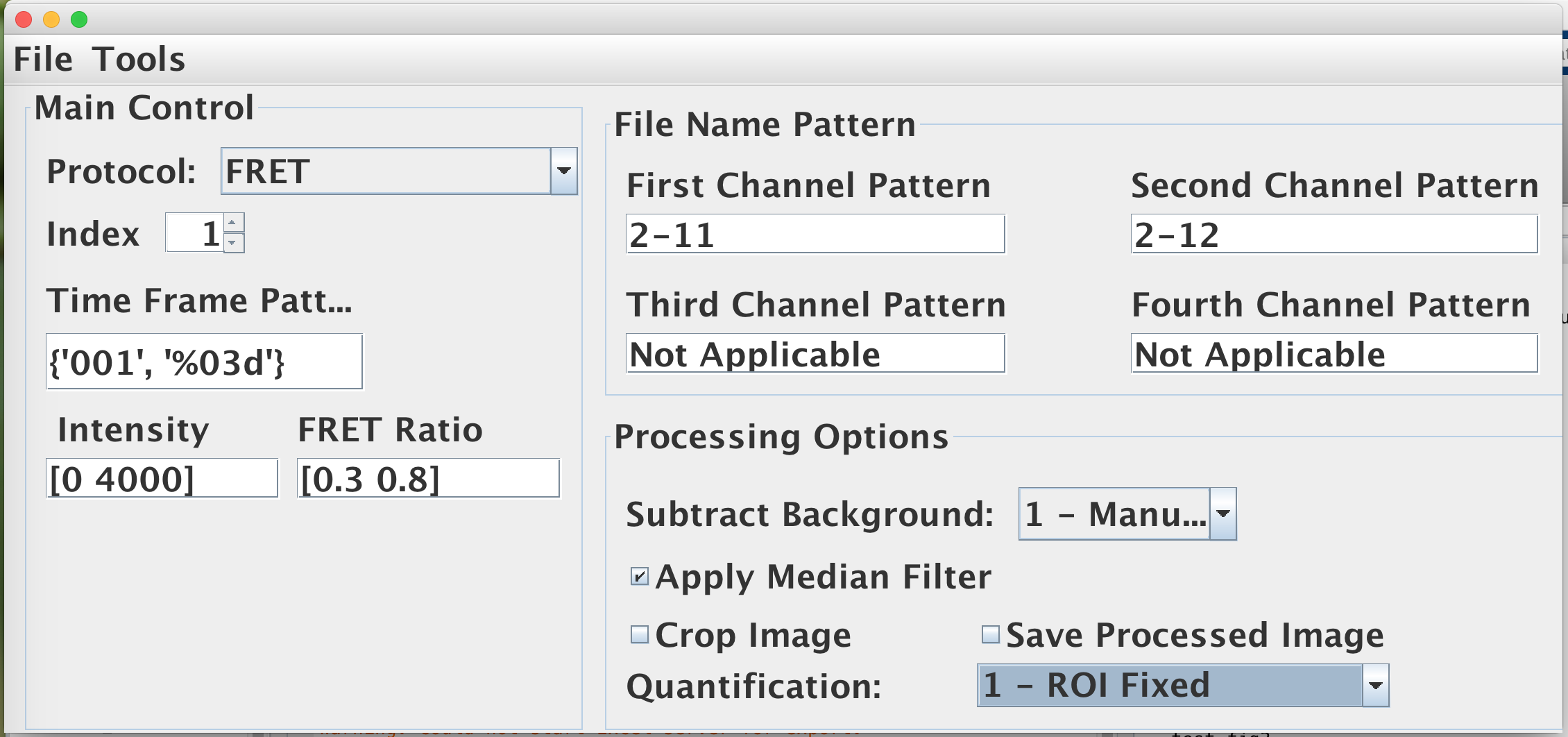
In the main menu of fluocell,

|  |  |
| --- | --- |
|  |  |
| ratio_image | intensity_image |

1. Select the **FRET** protocol. Also note the first channel pattern, second channel pattern, time frame pattern, intensity, FRET ratio and processing options.
2. Open the image files: File 🡪 Open Figure 🡪 Navigate to the image files 🡪 **Be sure to open the first channel file among the list of files.** Sample data image files can be found in the folder: fluocell\_sample\ 10\_24\_08\_Src\_fret\_pax\. The first channel image file is ‘2-11.001’. Note here how the file name patterns including “First Channel Pattern”, “Second Channel Pattern”, and “Time Frame Pattern” etc, are defined in the fluocell main window. As shown to the right, the CFP/FRET ratio image will be displayed in Figure 1 and the intensity image of FRET will be displayed in Figure 2. There are options to subtract background and apply median filter.

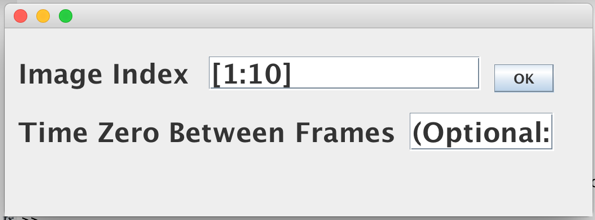
The intensity and FRET ratio information are needed for processing the ratio image. Independently, the intensity images can be adjusted using the colormap editor by clicking the Figure 2 menu: Edit 🡪 Colormap. The figures can be copied by clicking Alt+Print Screen, or Edit 🡪 Copy Figure, and pasted to a Microsoft PPT file.

1. Close files and quit the image tool: File 🡪 Close 🡪 Quit. Note: it is necessary to close the processed image using fluocell interface cleanly: File 🡪 Close Figure instead of closing it directly by clicking “x”.



***Quantification using regions of interest (ROIs).***

1. At the Fluocell Menu, there is a dropdown list that includes four different options. Select 1 – ROI Fixed.
2. Batch process: Tool 🡪 Batch Update Image 🡪 To process images from 1 to 10, set Image Index to [1:10] 🡪 click ‘OK’ **only once**.



1. To retrieve data, copy the output variables from the MATLAB Workspace to Excel: fluocell\_data 🡪 time, fluocell\_data 🡪 ratio, fluocell\_data 🡪 channel1 (average intensity of the first channel); fluocell\_data 🡪 channel2 (average intensity of the second channel). The fluocell\_data.time variable has two columns. The first column contains the index number of the image frames. The second column contains the time that the image frames were saved, in minutes starting from 12:00am of the day.

***Make Movies***

1. Choose settings and open a figure by going to File -> Open Figure
2. Save Ratio images by checking the box “Save Processed Image”, and in Tools -> Batch Update Image enter the range of files (e.g. [1:10]), click ok. The processed images have been saved as .tiff files in the data folder.

Note once the FRET images are saved, the options Intensity and FRET ratio bound will become ineffective. The background, ROI and processed image files are saved in a subdirectory output/ within the image path. When the user manually change the index, the ratio images will be saved automatically in the path: fluocell\_sample\10\_24\_08\_Src\_fret\_pax\output\0.3–0.8\ (the range is set in the interface of fluocell under ‘FRET Ratio’). The ratio images are convenient for visual inspection. To remove and re-define these files, the user needs to manually remove the saved files in output/ folder and repeat the image analysis steps.

1. Once the images have been saved, in the MATLAB command window run the make\_movie function. First, modify the function fluocell/app/sample\_init\_data.m. Replace the root variable in line 4 with the location of fluocell\_sample/ folder. Then run these commands:

>> cd ../../../app

>> data = sample\_init\_data('src\_pax','make\_movie');

>> make\_movie(data);

The function make\_movie() takes pre-existing images (.tiff) and assemble them into movies. The movie file “fret.avi” can be found in the “output\” folder within the data folder. Note: this feature currently does not work with Intensity and Intensity-DIC protocols.

***Visualize the Intensity Modified FLIM Images***

Load the images from this folder “data\fluocell\_sample\flim\_0505\_2014\”

|  |  |
| --- | --- |
|  | |
|  |  |

***Polarity Analysis***

Rotate the cells to horizontal position and quantify the polarity of signal. An example using the sample data:

1. Open the Image.

* Fluocell Menu 🡪 Protocol 🡪 Intensity, Time Frame Pattern:{‘001’,’%03d’}, select “Subtract Background: 1-manual”, and check the box “Apply Median Filter”. Leave the rest to default values.
* Open the image files by selecting File -> Open Figure. Navigate to the data folder fluocell\_sample\PH-Akt-GFP\_1\ -> double click the file “AKT-PH-YFP\_PDGF52.001”.

|  |  |
| --- | --- |
|  |  |

1. Cell boundary can be detected by selecting: Tools 🡪 Adjust Brightness Factor 🡪 Show Detected Boundary. Choose the Brightness Factor for cell detection.

Menu 🡪 Tool 🡪 Adjust Brightness Factor; Brightness Factor 1.0 (suggested value between 0.7 and 1.3), check the box “Show Detected Boundary”. Change the index value to show the boundary. Adjust the colormap to 0-5000 (for other data, adjust as needed). \*Note: The user may see different intensities of color depending on what their default colormap is in MATLAB.

1. In the file sample\_init\_data.m, confirm that the root variable in line 4 was updated to the location of fluocell\_sample/ folder.
2. Run the polarity quantification scripts

>> cd fluocell/app

>> cell\_name = 'akt\_1';

>> data = sample\_init\_data('akt\_1');

>> single\_cell\_analyzer('akt\_1',data);



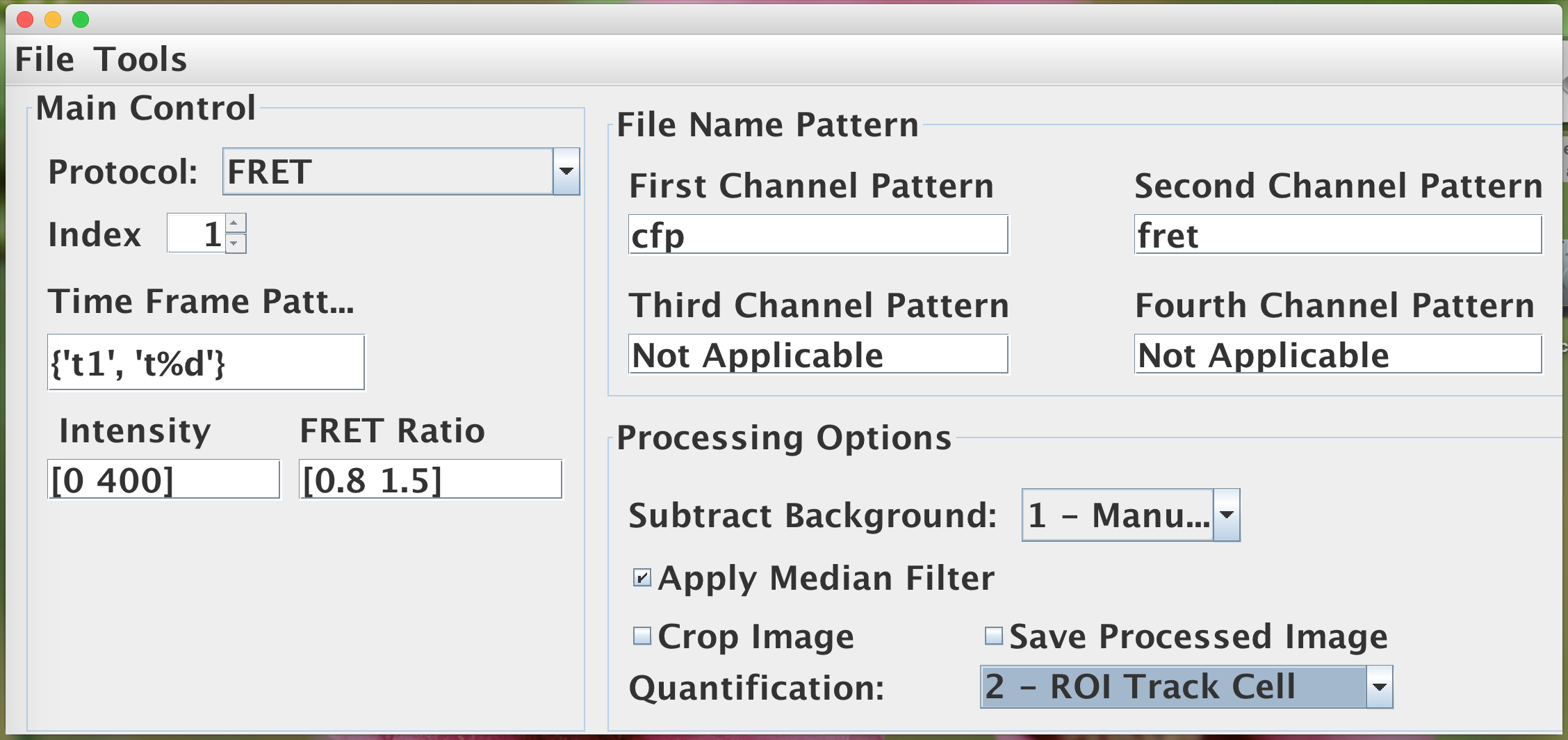
**Tracking Cells and Regions of Interest**

In the “Quantification” submenu, there are four options: (1) 0 – None: no quantification on region of interest; (2) 1 – ROI Fixed: quantify the region of interest without tracking cell; (3) 2 – ROI Track Cell: quantify the region of interest while tracking the cell.

**Note** that the option to “Apply Mask” may create a conflict as Fluocell detects another cell in the mask and move the ROIs by mistake. To avoid this conflict, we need to close the figures and re-open the figures each time when the “Apply Mask” is checked or unchecked.

>> fluocell\_data.num\_rois = 3;

Use the Fluocell GUI to navigate to another example data: tracking\_ex/ folder.



|  |
| --- |
|  |
|  |

To quantify subcellular layers, choose “3 – Quantify Subcell”. In this case, the default number of layers is 1 and user can change it by

>> fluocell\_data.num\_layers = 1;

These options also set the variable fluocell\_data.quantify\_roi to corresponding values.



>> fluocell\_data.num\_layers = 3;



The quantification output can be found in MATLAB workspace, under the variable “fluocell\_data”, such as fluocell\_data.time, fluocell\_data.ratio, fluocell\_data.donor, fluocell\_data.acceptor etc.

***FA Detection and Subcellular Quantification***

1. Detect cell mask. The following commands will detect a sequence of cell mask and output mask files to the subfolder output/

>> cd fluocell/app/fa\_analysis/

>> cell\_name = 'src\_pax';

>> data = sample\_init\_data(cell\_name, 'batch\_detect\_cell');

>> batch\_detect\_cell(cell\_name, data);



1. Detect Focal Adhesions

>> cell\_name = 'src\_pax';

>> data = sample\_init\_data(cell\_name, 'batch\_detect\_fa');

>> batch\_detect\_fa(cell\_name, data);



1. Programs to quantify local FRET ratio and FA intensity.

>> compute\_fa\_property(cell\_name, data);



***Useful Fluocell Functions:***

It is required that the image sequence of each cell is located in a separated folder. This function can be used to divide multiple position data into different folders: batch\_sort\_file\_multiple\_position(path, sub\_dir). Note: please don’t name a file with any space, which may confuse MATLAB programs.

1. Example:

>> path = 'E:\sof\fluocell\_2.1\data\migration\07\_01\_2010\';

>> sub\_dir = {'FN5\2\_2\', 'FN5\2\_3\', 'FN5\2\_FBS10\', 'FN10\1\', 'FN20\1\'};

>> batch\_sort\_file\_multiple\_position (path, sub\_dir)

1. Different types of regions of interest. By default, fluocell allows the user to choose a region of polygonal shape which is supplied by the user. There is an additional option to choose a circular region of a fixed diameter. This can be set up in MATLAB by

>> fluocell\_data.roi\_type = ‘circle with fixed diameter’;

>> fluocell\_data.roi\_diameter = 100;

If you want to switch back to a polygon region, simply remove the roi.mat file from the ourput/ folder and set:

>> fluocell\_data.roi\_type = ‘any’;

>> fluocell\_data.roi\_diameter = 0;

**Frequent Asked Questions**

1. *What is the channel patterns and time frame pattern for the images obtained using software other than metamorph.*

*Answer:* The channel patterns and time frame pattern are provided by the user to help Fluocell to recognize the donor and acceptor image files. The default values will work for the image files obtained from out Nikon microscope via metamorph. For example, the image files have the names such as fret11.001, fret11.002, …, fret12.001, fret12.002 etc.

The first channel pattern: fret11 ; the second channel pattern: fret12; the time frame pattern: {‘001’, ‘%03d’}, where the pattern string ‘%03d’ follows the convention in C++ or MATLAB, meaning an integer of 3 digits padded by 0. And the string ‘001’ should match the first donor image file that you click when opening the image. In another example, the image files have the names such as cfp\_t1.tiff, cfp\_t2.tiff, …, yfp\_t1, yfp\_t2 etc.. The first channel pattern: cfp; the second channel pattern: yfp; the time frame pattern:{‘t01’, ‘t%d’}, where ‘t%d’ means the letter ‘t’ followed by an integer.

1. *The first images opened correctly, but Fluocell cannot navigate through the time sequence.*

*Answer:* The Fluocell software use “time frame pattern” submitted by user to recognize file names and navigate through the time frame. For example, if the first image file has the pattern filename\_t1.TIF, the software will replace ‘t1’ with ‘t2’, ‘t3’, ‘t4’ etc. But, if fret18\_t1.TIF. In this case, Fluocell will think the next file in the time sequence as fret28\_t2.TIF by replacing ‘t1’ in two places the file name. In order to fix this problem, the file name pattern need to be defined as {‘\_t1’, ‘\_t%d’}, so that Fluocell can correctly replace the unique pattern ‘\_t1’ with ‘\_t2’, ‘\_t3’, ‘\_t4’ etc.

Note: Moving the Fluocell folder between operating systems may cause problems to arise. If Fluocell needs to be run on multiple systems, it is recommended that each system download its own copy of the Fluocell program.

***Installation and Usage***

Download and install fluocell either in the folder C:/Program Files/ or a local user folder. To Open Fluocell:

* On Windows/Mac
  + Double click “fluocellJava.jar” A window should appear and a new session of MATLAB should launch.
* On Linux
  + In the terminal, navigate to the directory containing “fluocellJava.jar.”
  + Enter “sudo java –jar fluocellJava.jar.” A window should appear and a new session of MATLAB should launch.

For future convenient usage, please make a shortcut of the executable file “fluocellJava.jar” at the desktop. If MATLAB does not launch when “fluocellJava.jar” is executed, the location of the MATLAB executable file needs to be specified. To do this, (1) In the fluocell/src/gui/java/ folder, make a copy from “win-default.property” (“mac-default.property” on Mac) to “default.property”; (2) Open “default.property” file with a text editor, and after “matlablocation=” enter the path to the MATLAB executable. In Windows, use double backslashes(\\) between directories, since a single backslash is the escape key (e.g.

“matlablocation=C\:\\Program Files\\MATLAB\\2013a\\bin\\matlab.exe”).

In MATLAB, add the fluocell programs to the searching path.

>> File 🡪 Set Path 🡪 Add with Subfolder 🡪 double click the folds to add 🡪 fluocell/app/ and fluocell/src/ 🡪 Save

The resulting pathdef.m file should be saved either to its default location (MATLAB/toolbox/local/) or the same directory as “fluocell.jar” (usually in fluocell/src/gui/java/).

**Updates**

1. Version fluocell-5.0