

# Watershed Cells GUI

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## Table of contents

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- [Table of contents](#)
- [Installation](#)
  - [Option 1: MATLAB-Based GUI](#)
  - [Option 2: Standalone Application](#)
- [Usage](#)
  - [Start the GUI](#)
  - [Select Images](#)
  - [Segmentation](#)
  - [Classification](#)
  - [Save Data](#)
  - [Batch Process](#)
  - [Note](#)
- [More Information](#)
  - [Authors](#)
  - [Copyright](#)

## Installation

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The watershed\_cells\_gui can be implemented in two ways:

1. As a GUI run and accessed from MATLAB
2. As a standalone application installed on a Windows machine

### Option 1: MATLAB-Based GUI

#### Download the GUI files

Download the GUI (graphical user interface) files to a folder and save it somewhere. The folder should contain the following files:

```
watershed_cells_gui.m
watershed_cells_gui.fig
private\apply_function.m
private\apply_threshold.m
private\find_regions.m
```

You may also want to include the provided example image, `test.tiff`.

## Add the GUI folder to your MATLAB path

In MATLAB, go to the HOME tab and, in the ENVIRONMENT section, click "Set Path". In the window that opens, click "Add Folder..." and add the folder containing the GUI files to your path.

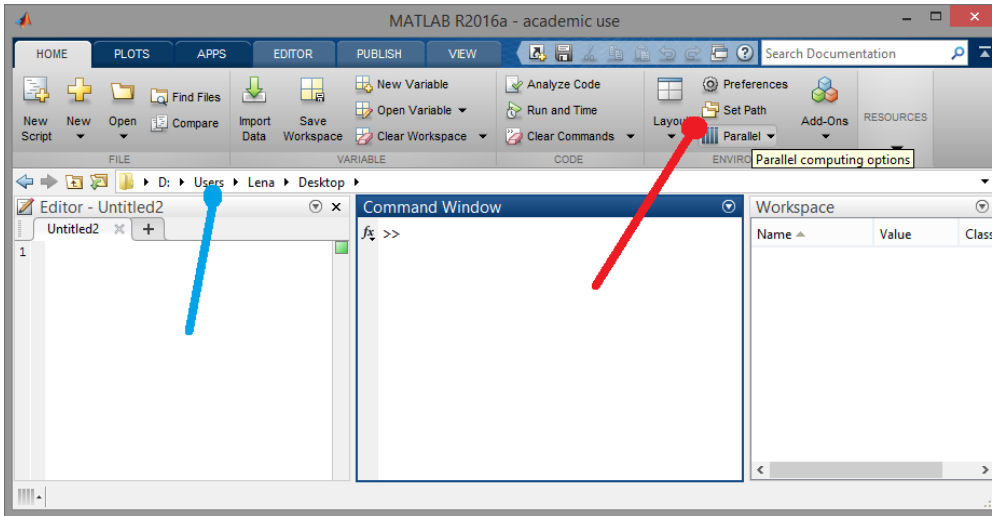


Figure 1. MATLAB window highlighting Set Path (red arrow) and current folder (blue arrow).

## Check your MATLAB version

This GUI will only work on MATLAB R2014b or later and requires the Image Processing Toolbox. To check your version of MATLAB, enter `ver` into the Command Window. You should get something that looks like this:

```
>> ver
-----
MATLAB Version: 9.0.0.341360 (R2016a)
...
-----
MATLAB                      Version 9.0      (R2016a)
Image Processing Toolbox     Version 9.4   (R2016a)
...
```

Note: This GUI has only been tested on Windows machines. I have no expectation that it will work on other operating systems.

## Start the GUI

To start the watershed cells GUI, run:

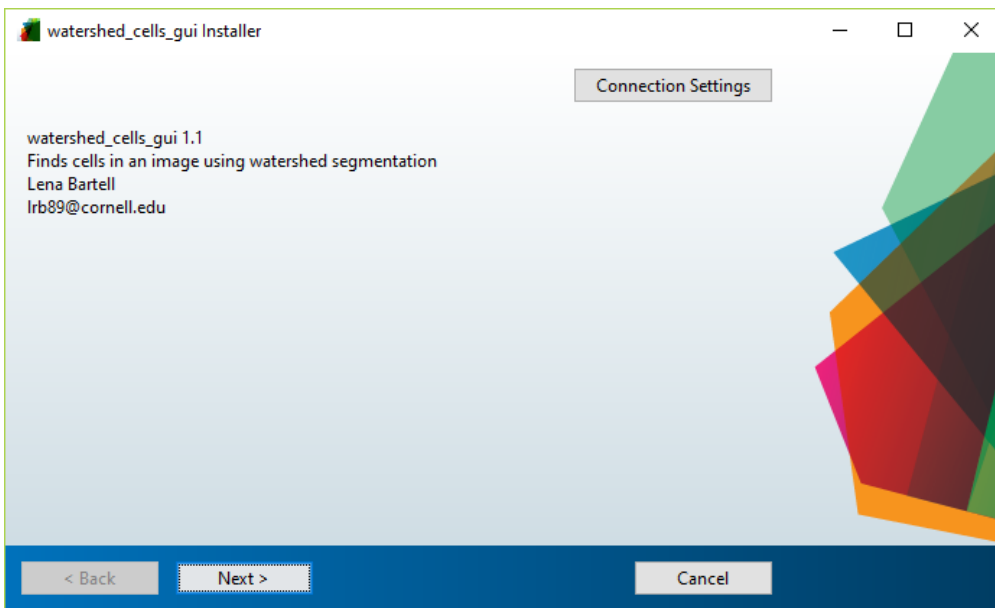
```
>> h = watershed_cells_gui
```

This will open the watershed cells GUI in a window with handle `h`. Within this handle, you can access the current state of the GUI data in the `UserData` field. In particular, `h.UserData` is a struct with the field `params`, which holds the current parameters from the GUI and `results`, which holds the current analysis results. For example, after the GUI opens, run `h.UserData.params.image.path` to return the path to the image being analyzed, or run `h.UserData.results.segmentation.number` to return the total number of regions found during segmentation. These fields are all populated with default values when the GUI is created, but will update as you use the GUI.

## Option 2: Standalone Application

*Note: The standalone application is not currently available.*

Download and run the application installer: `watershed_cells_gui_installer_mcr.exe` (be patient, it may take a few minutes to open). This will open a MATLAB-based installer to guide you through the installation process, as in Figure 2. Use this window to install the application and other related requirements, including the MATLAB Runtime.



*Figure 2. MATLAB Application installation window.*

After everything has finished installing, go to the installation folder. Now you can use the application by running the executable file `application\watershed_cell_gui.exe`. The installation also includes a test image, `test.tiff`, and a documentation PDF, `README.pdf`. You may wish to create a shortcut of `watershed_cell_gui.exe` somewhere else, like your Windows Desktop.

## Usage

When the GUI first opens, it should look like this:

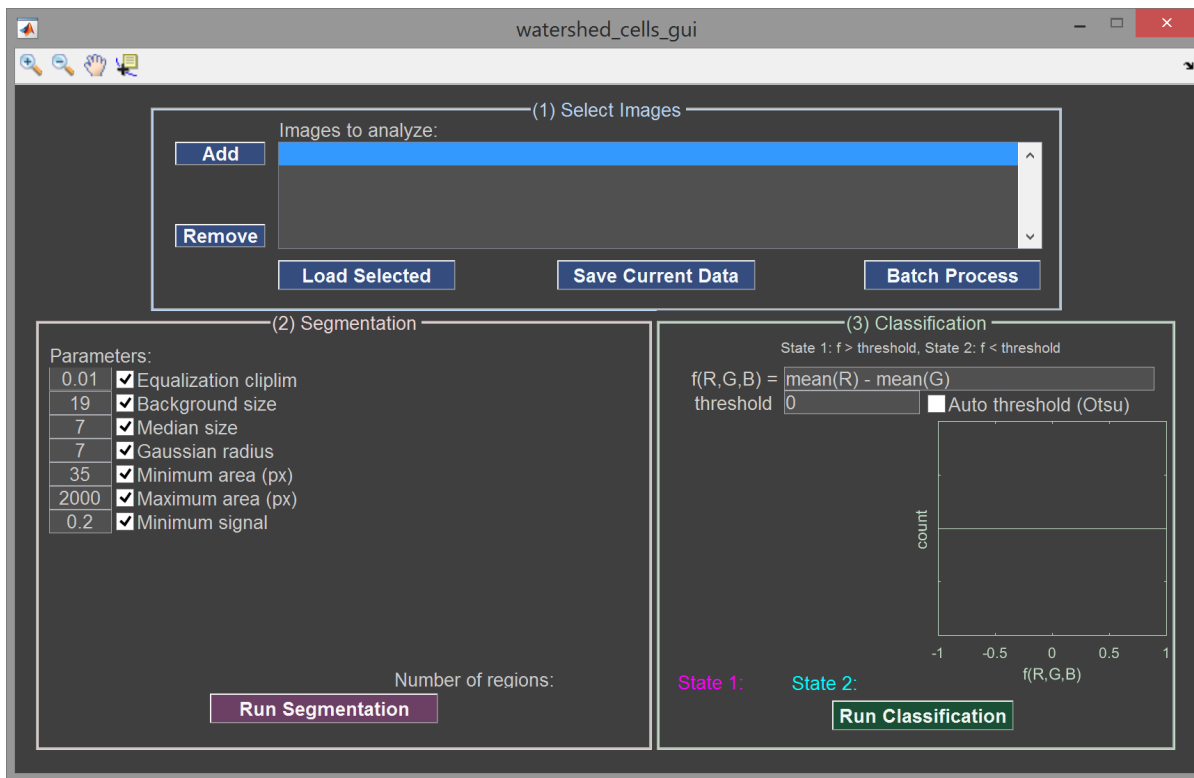


Figure 3. watershed\_cell\_gui initial appearance

## Select Images

In the **Select Images** panel, click the **Add** button to open a dialog box and select an image you want to process. To follow this example, use the included `test.tiff` file. Click **Load Selected** to import and display the selected image. The **Segmentation** panel shows a grayscale version of the image used for finding regions, while the **Classification** panel shows a full color version used for segmentation. At this point, the GUI should look like Figure 4. You can use the toolbar buttons to zoom/pan and inspect the images.

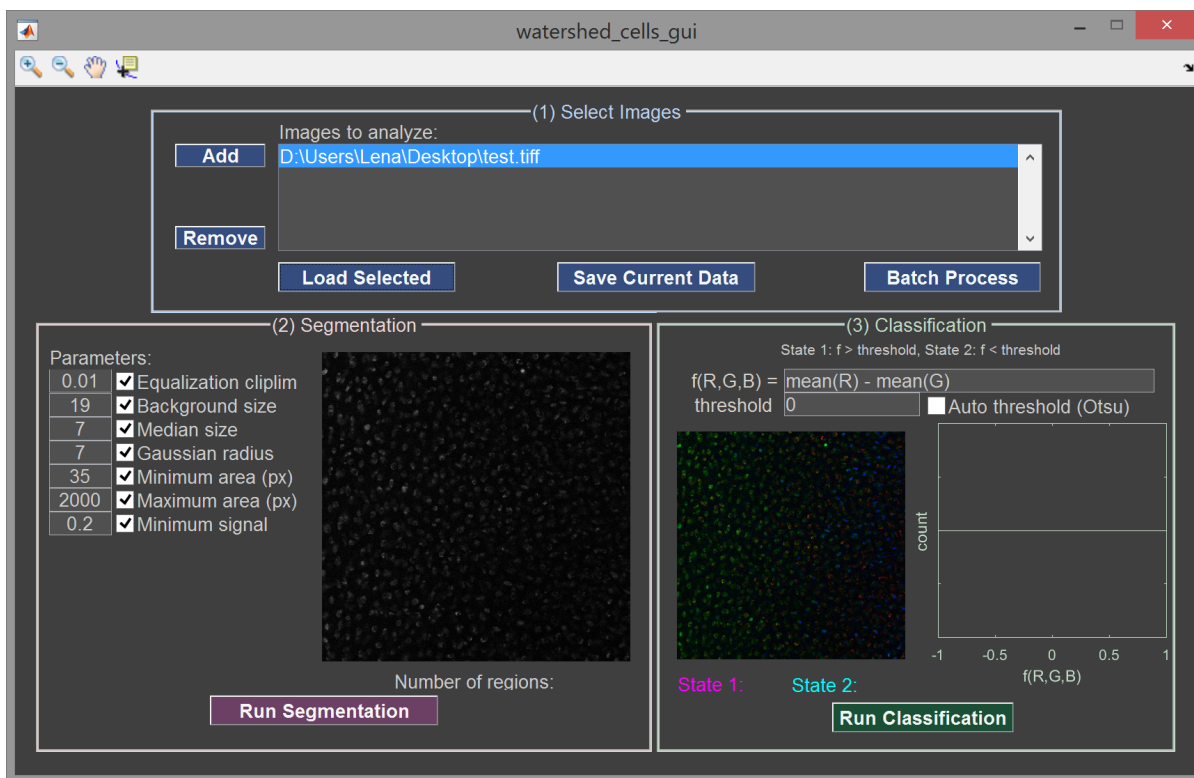


Figure 4. watershed\_cell\_gui after importing an image

## Segmentation

In the GUI, set the segmentation parameters by editing the text boxes in the Segmentation panel. To skip a particular step in the analysis, un-check the tick box next to the associated parameter. In general, you will need trial-and-error to pick the best parameters for your image. However, there are some rough guidelines below.

The parameters (and guidelines) are:

- *Equalization cliplim*: This is the "Clip Limit" parameter passed to the adaptive histogram equalization filter. This is a number between 0 and 1, but values around 0.01 are usually best. See MATLAB's function `adapthisteq` for more information.
- *Background size*: Background subtraction is performed using a median filter of this size (units: pixels). This should be an odd positive integer that is much larger than the diameter of your cells/objects. Background subtraction is usually the slowest step in the process, so skip it if you don't need it. Generally, background subtraction is important for epi-fluorescence images (due to unwanted out-of-focus/background signal), but optional for confocal images (because the confocal pinhole already blocks much of this background).
- *Median size*: The size of a separate median filter used for smoothing / reducing noise (units: pixels). This should be roughly the same size or smaller than the cells you are trying to find.
- *Gaussian radius*: The radius of a Gaussian filter, which is also used for smoothing / reducing noise (units: pixels). If the signal is discontinuous across an individual cell (e.g. if staining individual organelles rather than the cytoplasm), this radius should be approximately the radius of a cell. If the signal is relatively continuous across the cell (e.g. standard live/dead staining assay), this can be smaller. Values smaller than 0.5 px are not recommended.

- *Minimum area*: After finding objects/cells, discard any objects that have fewer than this many pixels in their area (units: pixels). This should be approximately  $r^2$ , where  $r$  is the average cell radius in pixels.
- *Maximum area*: Also discard any objects that have more than this many pixels in their area. This should be approximately  $(2r)^2$  to  $(5r)^2$ , where  $r$  is the average cell radius in pixels.
- *Minimum signal*: Also discard any objects that have an average signal-intensity smaller than this value (units: fraction of intensity range). This value is expressed as a fraction of the full range of possible intensities, so it should be a number between 0 and 1 (rather than 0 and 255, for example). This value will vary drastically based on imaging settings, image quality, other parameter values, and your desired output, so I won't provide any guidelines.

Once parameters are set, click **Run Segmentation** to run the image analysis process and display the result. Briefly, the image analysis procedure is:

1. Pre-processing
2. Import image
3. Adaptive histogram equalization
4. Background subtraction using a median filter (using "background size" parameter)
5. Median filter to smooth
6. Gaussian filter to remove noise
7. Watershed segmentation
8. Determine background using a conservative Otsu's threshold
9. Watershed segmentation, with background enforced
10. Post-processing
11. Remove cells that are too small
12. Remove cells that are too big
13. Remove cells that are too dim

Once the image segmentation is complete, regions that are found will be circled in yellow on the plot, as in Figure 5. You can use the toolbar buttons to zoom/pan and inspect the plots.

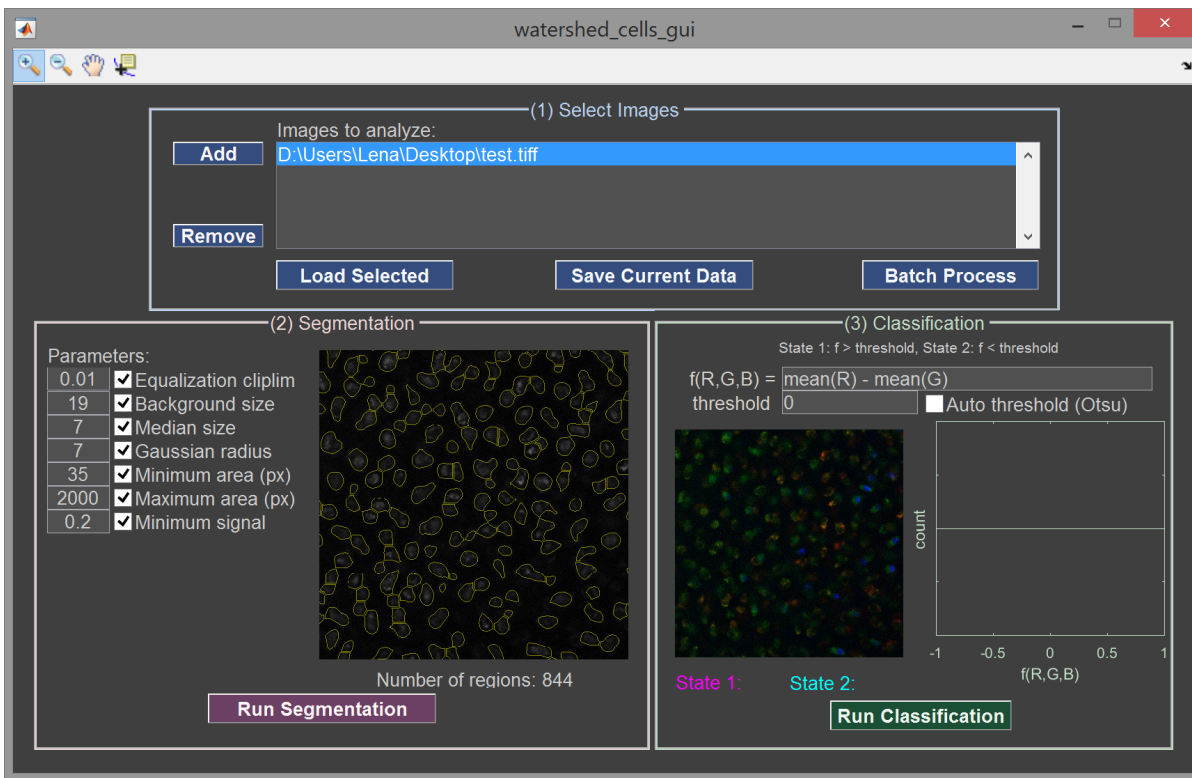


Figure 5. watershed\_cell\_gui segmentation result. Note that the number of regions found is also shown on the screen.

## Classification

To classify regions, the GUI applies a function,  $f(R,G,B)$  to each region, where R, G, and B, are, respectively, the list of red, green, and blue pixel values in the given region. This function should return one scalar number for each region. This number will then be thresholded according to the threshold parameter. If Auto threshold (Otsu) is selected, then the threshold value will be chosen automatically based on Otsu's thresholding scheme.

Enter the function definition and threshold (if manual) in the text boxes. The function may be a custom function and/or may call other Matlab and user-defined functions. Then, click Run Classification to run the classification process and display the results.

Once the classification is complete, the regions with f values above and below the threshold will be outlined in magenta and cyan, respectively. Also, the GUI shows a histogram of the f values from all the segmented regions. You can use the toolbar buttons to zoom/pan and inspect the plots. At this point, the GUI should look like Figure 6. You can use the toolbar buttons to zoom/pan and inspect the plots.

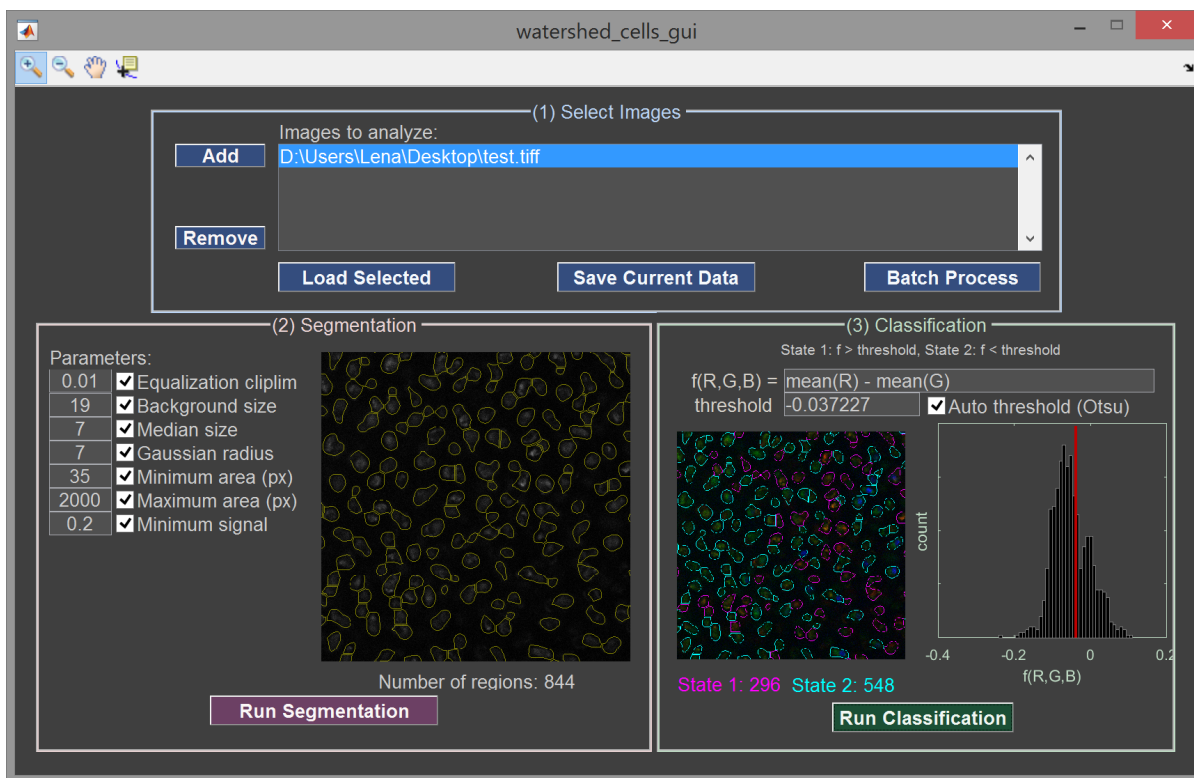


Figure 6. watershed\_cell\_gui classification result. Note that the number of regions above/below the threshold (state 1 / state 2) is also shown on the screen.

## Save Data

Click **Save Current Data** to save the results currently shown in the GUI. This will open a dialog box asking the user to choose a folder in which to save the results. In this folder, the GUI will save two files (note that both files use the base name from the image file):

1. `<image name>_data.mat`: A MATLAB data file containing the structure 'data'. This is the same structure output at the command line (i.e. data from `[h, data] = ...`). This structure contains the field `params`, which specifies the parameters from the GUI's current state, and `results`, which contains the results of the segmentation and classification.
2. `<image name>_display.tif`: A TIF image file showing the imported image with all regions outlined based on their state (above threshold = magenta, below threshold = cyan, equal to threshold = yellow). Open this image to visually check the results.

## Batch Process

You can also use the GUI to automatically process multiple images with the same parameters. To do this first use the **Add** button to select all the files you wish to process. Next, setup all the parameters in both the **Segmentation** and **Classification** panels. Finally, click **Batch Process** to analyze all the listed images. This will loop through each listed image file, load it, segment it, classify it, and save resulting data, before moving on to the next image on the list.

As the bath process continues, the GUI plots will dynamically update to show the current processing results. While the batch process is running click the **Cancel** button to stop the analysis. This will finish



completing the current calculation step and then cancel all remaining steps. Any unsaved data will be lost.

## Note

The “meat” of the segmentation algorithm is encoded in the private function `private\find_regions.m`. If you would like to alter the processing algorithm or use it externally or programmatically, start there. Similarly, the “meat” of the classification process is contained in `private\apply_function.m` and `private\apply_threshold.m`.

## More Information

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### Authors

This algorithm and GUI was written by Lena Bartell during the completion of her PhD thesis in Prof. Itai Cohen's group at Cornell University. Lena was generously supported by NSF GRFP DGE-1144153 and NIH 1F31-AR069977.

We welcome feedback. Please email Lena with any questions, comments, suggestions, or bugs.

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