

ABSTRACT

Automated Measurement of a Cell's Morphological Evolution via Image Analysis

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Area and eccentricity of mouse-embryo fibroblasts are measured as these cells release the surface of a collagen gel. Image sets of the cells are collected at regular time-intervals during and after detachment. Software was written to segment selected cells and measure both the area and eccentricity of each segmented cell. As a main product of this research, the Matlab code for segmenting selected objects is listed in the appendix.

LAB-METHODS

Mouse embryo fibroblasts exhibiting green fluorescent protein (cell line NIH3T3/GFP) are placed atop a collagen-gel and suspended in growth medium(see the diagram in figure 1). The fibroblasts attach to the surface of the collagen at 37°C for 15 to 30 hours in a CO₂-regulated incubator. The cells were not seen to undergo mitosis.

Cells are imaged every regularly beginning 40 minutes before trypsinization via TrypLE Select. The images are collected with a confocal microscope using a 70/30 transmission/reflection beam-splitter. The images are collected in image sets called z-stacks.

A z-stack is a sequence of images taken in quick succession. After the first, lowest image, each image is focused a few microns higher than the preceding image. Imaged in this way, z-stacks essentially provide a "snapshot" of the state of specific cells within the sample.

The stage is adjusted before imaging a sample so that the z-stacks image from ten microns below the surface of the collagen to ten microns above the surface of the collagen. Ten microns is approximately the radius of these cells when not attached to the collagen. Z-stacks are captured every five to fifteen minutes.

Imaging the cells occurs by irradiating the sample with a 488nm laser. The green fluorescent proteins fluoresce at a different wavelength than the 488nm excitation light, thereby distinguishing fluorescence from reflection. Light from 500nm to 550nm is observed for emissions of the GFP.

The fibers were also observed for alignment phenomena which were not seen. To image the fibers, the sample is excited by a 532nm laser and wavelengths in the spectrum 531nm to 536nm are observed for reflection and transmission of the incident light off of the fibers. The observation spectrum imaged for the fibers was found by visually determining the spectrum in which the fibers appear most distinct.

Light detected from the fibers is maximized by observing a spectrum symmetric about the wavelength of the incident light, say 526nm to 536nm. However, such a spectrum has a higher signal to noise ratio because more of the light emitted by the GFP is detected below 531nm.

Both sets of images are acquired with a 30% reflection, 70% transmission beamsplitter. Using the same beamsplitter for both images means time is not spent changing the beamsplitter between images. This time savings between frames allows a z-stack to be captured closer to instantaneously. An instantaneous "snapshot" of the z-stack is ideal.

During imaging, the growth-medium surrounding the cells is rinsed with 1XPBS twice. The PBS is next replaced with TrypLE Select. The TrypLE Select breaks the proteins binding the cells to the collagen-matrix.

The proteins that bind the cells to the collagen are on bulges in the cell-membrane. These protrusions are called filopodia. After the bonds break, these filopodia flatten back into the cells.

This flattening of the filopodia leaves the cells somewhat spherical. The spherical fibroblasts appear as circles when seen through the microscope. While the cells return to a spherical shape, their lack of sphericity can be measured as eccentricity, as in the eccentricity of an ellipse.

FIG. 1. A diagram of orange cells submerged in violet growth medium atop green collagen

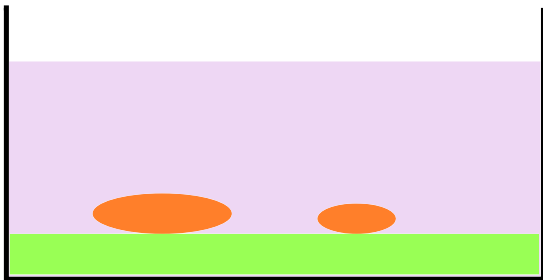


IMAGE PROCESSING

A single experiment yields a folder containing all z-stacks acquired during that experiment. Each z-stack is comprised of several images, as described before. These images are processed to yield numerical data.

The first step in processing the images from experiments is to transform each z-stack into single image. There is certainly data lost in the transformation of the

z-stacks from several images to a single image. However, this experiment is only concerned with whether a cell is present at each pixel's location.

For this reason, a maximal intensity projection (MIP) is used to transform all images comprising a z-stack into a single image. An MIP creates a maximal intensity projection for each z-stack. This maximal intensity projection is a single image. The maximal intensity projection is made by using the value of the brightest pixel at position (x_1, y_1) chosen from among the values of each image's (x_1, y_1) pixel. An example of the MIP function is shown in figure 2.

$$\text{mip}(\begin{array}{|c|c|} \hline \text{■} & \text{■} \\ \hline \end{array}, \begin{array}{|c|c|} \hline \text{■} & \text{■} \\ \hline \end{array}) = \begin{array}{|c|c|} \hline \text{■} & \text{■} \\ \hline \end{array}$$

FIG. 2. An illustration of the MIP function creating a projection from two simple example images

The values of all pixels in the MIP-images is multiplied by 50, effectively "brightening" the images. This is done to facilitate masking and subsequently segmenting entire cells. Seemingly, reducing the brightening factor would reduce the noise at the edges of the segmented cells. If so, this would improve precision accuracy of the numerical data.

The Matlab Image Processing Toolbox function `im2bw` uses a brightness threshold to create a binary (black and white) image from a grayscale image. Pixels brighter than the threshold value are converted to white while other pixels are turned to black. The threshold used here to create binary images is 0.15.

The function `maskmaker` is used to draw a region loosely enclosing the cell at its most stretched out. `Maskmaker` sets pixels outside this region to black. This creates a mask loosely surrounding the cell of interest. Multiplying this mask leaves the user-selected cell as the largest object in the resulting masked binary image.

In this way, the function `stackmasker` applies this mask to all images in the stack. All masked images contain the entire cell of interest. The mask is drawn around the cell when the cells are stretched out to ensure the entire cell is visible in later images.

Masking works by ensuring that the cell of interest is the largest object shown. The reason a mask is made enclosing a cell is so that the user can specify that cell for software to segment precisely. Cells are precisely outlined by software to save time and to ensure uniformity in tracing the cell.

Many experiments show multiple cells in each image. For example, consider a fictitious image that contains cells A and B. Each cell's shape and eccentricity is measured as follows.

Cell A is loosely masked by the user so that it's the largest object visible. Cell A is segmented in the images. These images showing only cell A are saved. The area and eccentricity of cell A is measured from cell A's segmented images.

A cell is measured by segmenting a set of images for that cell only. The cell is segmented by ensuring it is the only object visible in its image-set. This is why masking needs to specify a single cell as it does.

After cell A is segmented, the user segments cell B with the same process as used for cell A. The process begins by loosely enclosing cell B in a binarized MIP containing both cells. In this way, information about each cell in an image can separately gathered from a single experiment's images.

In figure 3, the two images shown are the initial image and the final image from the same experiment. figure 4 shows the region of interest outlined in figure 3. The image-processing procedure is summarized in figure 4.

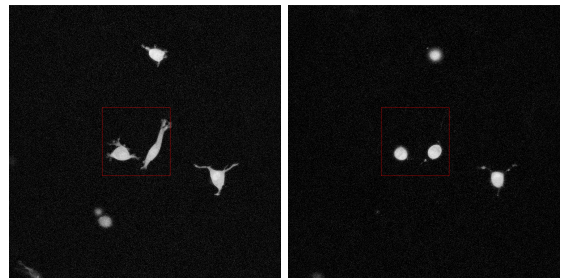


FIG. 3. MIP of first and last z-stacks from an experiment with the region of interest outlined in red

CONCLUSIONS

The plotted numerical data in figure 5 appears to show that area and eccentricity both decrease as time progresses. Analysis of data from further experiments might allow one to model size and eccentricity of cells as they release a substrate. One could use this model to predict size and area of fibroblasts when trypsinized.

To improve the results of this data, I would take a taller z-stack, take images more frequently, and adjust the brightening factor to increase smoothness in images. Using taller z-stacks should better capture entire cells as they move away from the surface of the collagen. More frequent images would increase data. Adjusting the brightening factor could reduce noise which is amplified when using the `im2bw` function.

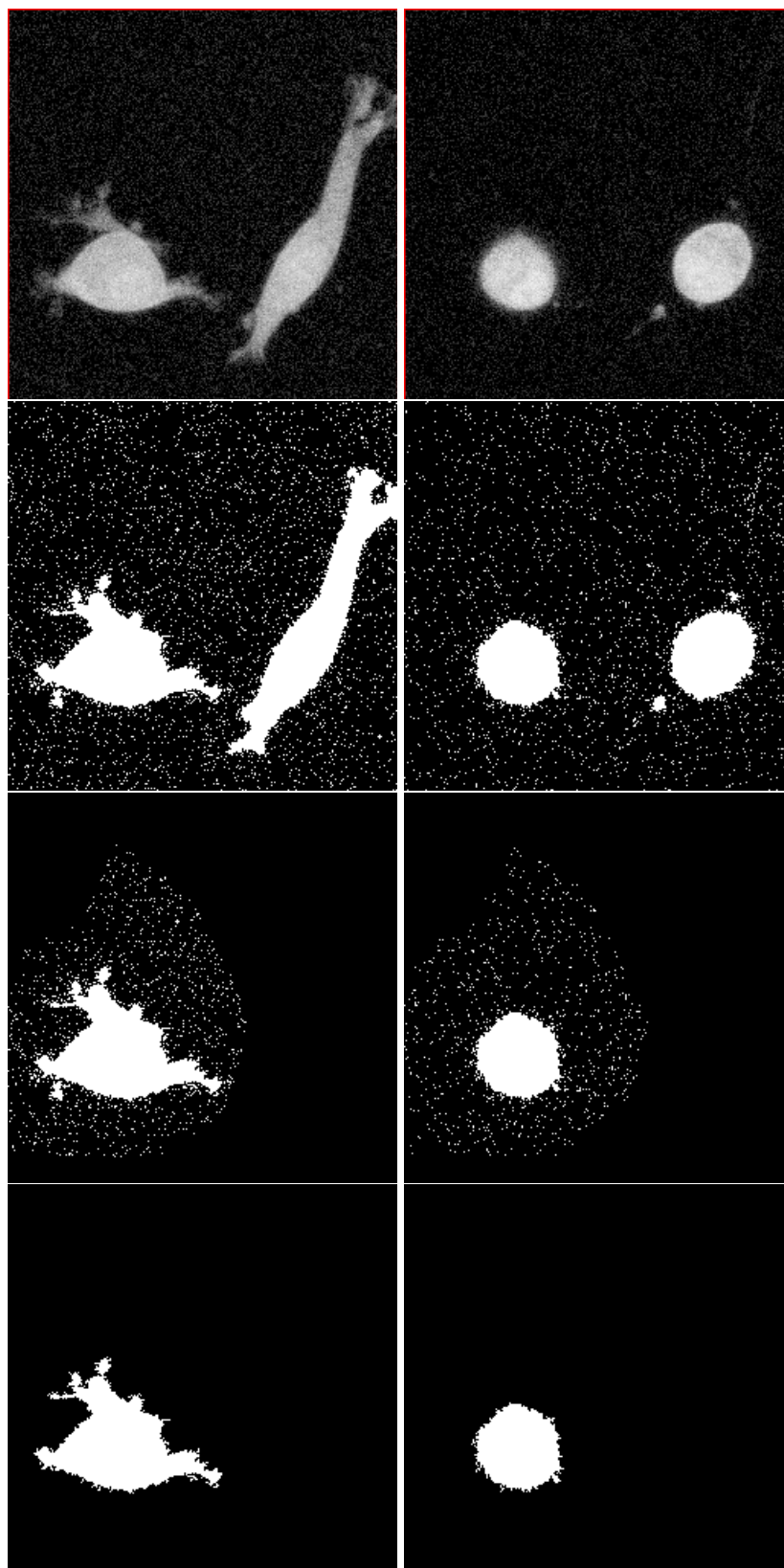


FIG. 4. Region of interest from FIG. 3 at several stages throughout image processing; from top to bottom, the rows show the original images, the result of 'binarization', the result of masking, and the segmented cell from the images.

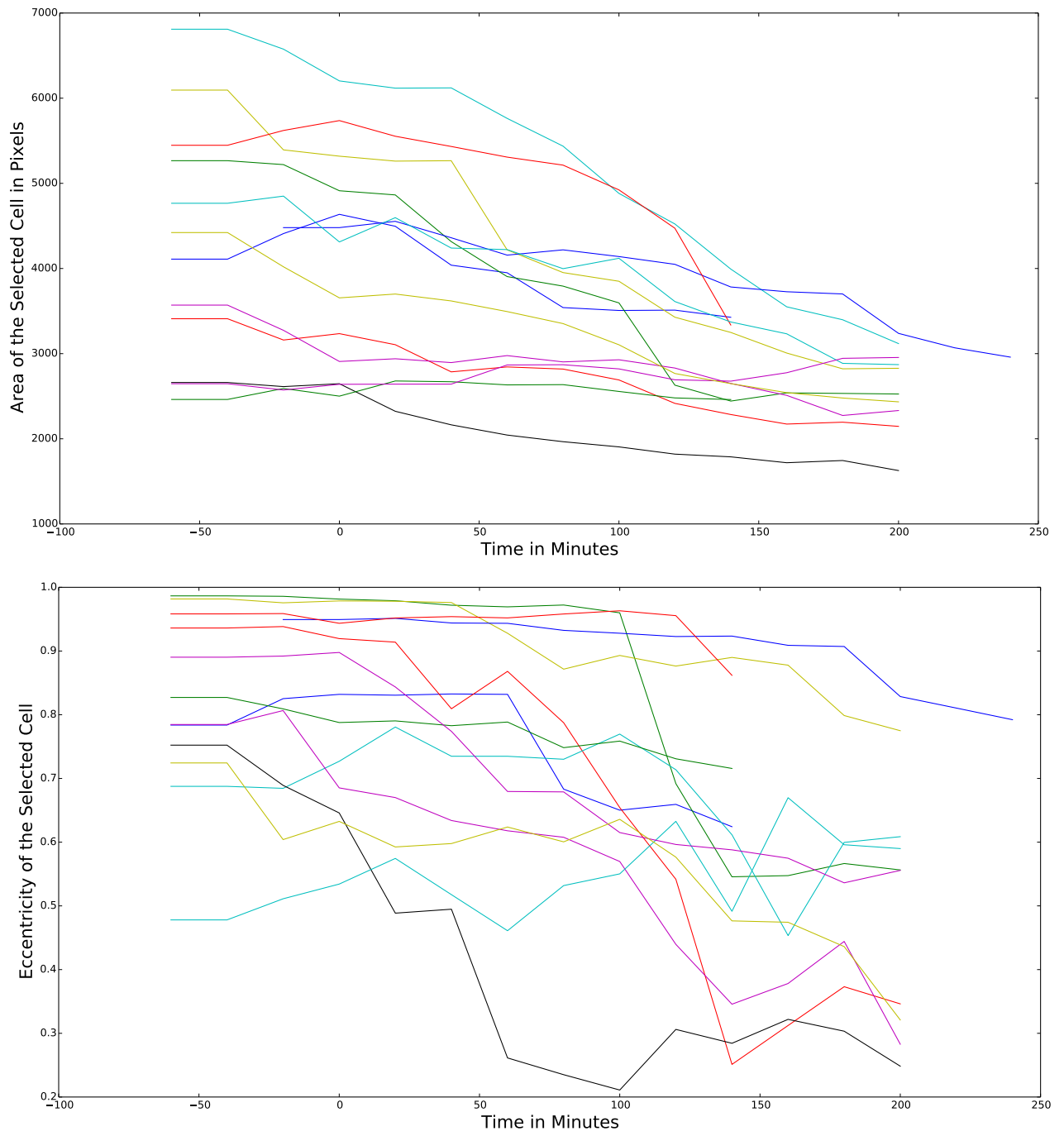


FIG. 5. Plots of experimental data: Area Vs. Time and Eccentricity Vs. Time

Appendix: Code (to be used and distributed freely)

```

1 % this script segments each object of a tiff-stack into its own stack
2 % by asking the user to mask a region constaining only one object.
3 % user decides how much to brighten and threshold the image for
4 % segmentation. Put all files for each sequence of mip images into
5 % a separate folder within "directory"

7 % directory contains folders which each contain an image sequence:
directory = '../..//data/0-good-images-no-beads';
9 % where to put output files:
outputPath = fullfile(directory, '..', '3-segmented');
11 % senesitivity for segmentation:
threshold = 0.15;
13 % brighten by this factor by default:
defaultBrighteningFactor = 15;
15 % get folders or file-sets containing stacks:
folders = ls(directory); folders = folders(3:end,:);
17 % save all intermediate stacks? (true or false)
saveIntermediateStacks = true;
19 % attempt to resume from the folder that the program left off at
if exist('k','var')
21     resume = input(sprintf(['Enter the folder number to start'...
22                             ' with or press enter to attempt\nresuming where the'...
23                             ' program left off.\n']));
25     if isempty(resume)
        startingfolder = k;
27     else
        startingfolder = resume;
29     end
31 else
    startingfolder = 1;
33 end

35 for k=startingfolder:length(folders(:,1))
    % read all folders in directory and skip folders with no tiffs
    stackDirectory = fullfile(directory, strtrim(folders(k,:)));
    if size(dir(fullfile(stackDirectory, '*.tif')),1)==0 &&...
37         isdir(stackDirectory)
        continue
39    end

41    % read stack in current folder
    disp(['FOLDER #' num2str(k) ': ' folders(k,:)]);
    stack = stackreader(stackDirectory, '*.tif');

45    % loop through binarizing stack until ok
    if exist('brightenedStack','var')
47        clear('brightenedStack');
    end
    ok = 0;
    while ~isempty(ok)
51
        % get brightening factor from user or use default
53        if ok ~= 0
            brighteningFactor = ok;
55        end
        if ~exist('brighteningFactor','var')
57            brighteningFactor = defaultBrighteningFactor;
        end
59
        % brighten, binarize, and fill the stack
61        bwStack = zeros(size(stack));
        for page=1:size(stack,3)
63            bwStack(:,:,page) =...
                imfill(im2bw(brighteningFactor*stack(:,:,page)),...
65                'holes');
        end
67

```

```

69     % show stack and ask if ok, not ok => redo w/ new parameters
    implay(bwStack);
    ok = input(['PRESS ENTER IF B&W STACK IS OK OR '...
71              'ENTER NEW BRIGHTENING FACTOR: ']);
    end
73
74 % write the binary stack if saveIntermediateStacks is true
75 if saveIntermediateStacks
76     sprintf(['SAVING BINARIZED STACK IN\n',...
77            fullfile(outputPath,'..','1-binarized')])
78     stackwriter(bwStack,fullfile(directory,'..','1-binarized'),...
79             ['binarized-' folders(k,:),'],'','');
    end
81
82 % let user mask region containing mth object and decide whether to
83 % save the selected object's stack
84
85 % ask how many objects in the current stack will be masked
    numobjects = input(sprintf(['How many objects will be'...
87                               'segmented in this stack?\n']));
88
89 % for each object, let user create mask
    for m = 1:numobjects
90         disp(['MASKING OBJECT #' num2str(m) ...
91              ' IN STACK ' folders(k,:) ]);
92         mask = maskmaker(bwStack);
93         while 1==1
94             appendObjectToFilename = ['object-' num2str(m)];
95             maskedstack = stackmaker(bwStack,mask,threshold);
96             % write the masked stack if saveIntermediateStacks is true
97             if saveIntermediateStacks
98                 sprintf(['SAVING MASKED STACK IN\n',...
99                        fullfile(directory,'..','2-masked')])
100                 stackwriter(maskedstack,fullfile(directory,...
101                        '..', '2-masked'), ['masked-' folders(k,:),'],'...
102                        ',appendObjectToFilename);
103             end
104             segmentedstack = maskedstack;
105             for page=1:size(maskedstack,3)
106                 segmentedstack(:,:,page) = ...
107                     ExtractNLargestBlobs(maskedstack(:,:,page),1);
108             end
109
110 % show segmented stack. ask if ok...
111 % if ok => save single-object-stack
112 implay(segmentedstack);
113 ok = input(sprintf(['Either enter 1 to save'...
114                    ' the segmented stack, press enter to skip,'...
115                    '\nor enter anything else to tinker.\n']));
116 if ok == 1
117     sprintf(['WRITING SEGMENTED STACK\n:' folders(k,:) '.tif']);
118     stackwriter(segmentedstack,outputPath,...
119             ['segmented-' folders(k,:),'],'',...
120             appendObjectToFilename);
121     sprintf(['SUCCESSFULLY WROTE THE STACK NAMED\n'...
122            'segmented-', folders(k,:),appendObjectToFilename,...
123            '.tif' ])
124     break
125 elseif isempty(ok)
126     break
127 else
128     tinker
129 end
130 end
131     continue
132 end
133 end

```

Listing 1. segmenter.m

```

% reads images in the directory or multi-page image matching
2 % the given pattern into a stack in memory
% stack = stackreader(stackDirectory, filenamematchingstring)
4 function stack = stackreader(stackDirectory, filenamematchingstring)

6 % reads cellarray of fileinfo for images in directory matching pattern
files = dir(fullfile(stackDirectory, filenamematchingstring));

8
% if only one matching image file then assume a multipage image
10 % (e.g., reads a multipage .tif)
if length(files) == 1
12     fname = files.name;
    filenameWithPath = fullfile(stackDirectory, fname);
14     info = imfinfo(filenameWithPath);
    numImages = numel(info);
16     stack = zeros([1024 1024 numImages]);
    for page = 1:numImages
18         stack(:,:,page) = imread(filenameWithPath, page, 'Info', info);
    end

20 % WARN: didn't find any matching files
22 elseif isempty(files)
    disp('ERROR: No matching image files');
24     return

26 % else assumes each file is a single page tif and reads files as 2d
% images into an array named stack
28 % (reads filename-specified tifs in specified directory into stack)
else
30     numImages = length(files);
    stack = zeros([1024 1024 numImages], 'uint8');
32     for page = 1:numImages
        filenameWithPath = fullfile(stackDirectory, files(page).name);
34         stack(:,:,page) = imread(filenameWithPath);
    end
36 end

```

Listing 2. stackreader.m

```

% stackwriter writes a stack to a multipage image
2 % function stackwriter(stack,outputpath,baseFilename,...
%     prependToFilename,appendToFilename)
4 function stackwriter(stack,outputpath,baseFilename,...
    prependToFilename,appendToFilename)
6 filename = [prependToFilename, baseFilename, appendToFilename, '.tif'];

8 % Write first stack-slice to new image file
imwrite(stack(:,:,1), fullfile(outputpath, [prependToFilename...
10     baseFilename...
    appendToFilename...
12     '.tif']))

14 % Append rest of stack-slices to image file
for k = 1:size(stack,3)
16     imwrite(stack(:,:,k), fullfile(outputpath, filename),...
        'writemode', 'append');
18 end

```

Listing 3. stackwriter.m

```

function mask = maskmaker(brightenedstack)
2 imshow(brightenedstack(:,:,1));
[r,c] = ginput();
4 mask = roipoly(brightenedstack(:,:,1),r,c);
close all;
6 imshow(mask);

```

Listing 4. maskmaker.m


```

% mask the stack to observe only the user-specified region
2 function maskedstack = stackmasker(brightenedstack,mask,threshold)

4 % imshow(stack(:,:,1));
%
6 % [r,c] = ginput();
%
8 % mask = roipoly(stack(:,:,1),r,c);

10 % figure, imshow(mask)

12 maskedstack = zeros(size(brightenedstack));

14 for k=1:length(maskedstack(1,1,:))
    bw = im2bw(brightenedstack(:,:,k),threshold);
16     maskedstack(:,:,k) = immultiply(mask,bw);
18 end
close all

```

Listing 5. stackmasker.m

```

function binaryImage = ExtractNLargestBlobs(binaryImage, numberToExtract)
2 %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% commented try/catch, investigate this if broken
4 %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% try

6
% Get all the blob properties. Can only pass in originalImage in version R2008a and later.
8 [labeledImage, numberOfBlobs] = bwlabel(binaryImage);
blobMeasurements = regionprops(labeledImage, 'area');
10 % Get all the areas
allAreas = [blobMeasurements.Area];
12 if numberToExtract > 0
    % For positive numbers, sort in order of largest to smallest.
14     % Sort them.
    [sortedAreas, sortIndexes] = sort(allAreas, 'descend');
16 elseif numberToExtract < 0
    % For negative numbers, sort in order of smallest to largest.
18     % Sort them.
    [sortedAreas, sortIndexes] = sort(allAreas, 'ascend');
20     % Need to negate numberToExtract so we can use it in sortIndexes later.
    numberToExtract = -numberToExtract;
22 else
    % numberToExtract = 0. Shouldn't happen. Return no blobs.
24     binaryImage = false(size(binaryImage));
    return;
26 end
% Extract the "numberToExtract" largest blob(a)s using ismember().
28 biggestBlob = ismember(labeledImage, sortIndexes(1:numberToExtract));
% Convert from integer labeled image into binary (logical) image.
30 binaryImage = biggestBlob > 0;

32 %%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%%
% catch ME
34 %     errorMessage = sprintf('Error in function ExtractNLargestBlobs().\n\nError Message:\n%s', ME.
    message);
%     fprintf(1, '%s\n', errorMessage);
36 %     uiwait(warndlg(errorMessage));
end

```

Listing 6. ExtractNLargestBlobs.m


```

1 one = 1;
  eval(tinkerings);
3 while one ~= 0
    tinkering = input(sprintf('(Enter tinkerings and remember one = 0 breaks things (like this loop
        :)\n'),'s');
5     eval(tinkering)
end

```

Listing 7. tinker.m

```

folder = '../.. / data/3-segmented';
2 files = ls(fullfile(folder, '*.tif'));
for file = 1:size(files,1)
4
    % ask for time interval between z-stack acquisition
6    stack = stackreader(folder, files(file, :));

8    % get time, area, and eccentricity values of object for each page
    disp('#####');
10   disp(['PROCESSING SEGMENTED OBJECT #' num2str(file) ':']);
    disp( files(file, :) );
12   disp('#####');
    timeInterval = input('\nTIME BETWEEN Z-STACKS: ');
14   timeOffset = input(['NUMBER OF Z-STACKS BEFORE '...
        'TRYPLE SELECT WAS ADDED: ']);

16   time = zeros(size(stack,3),1);
    area = zeros(size(stack,3),1);
18   eccentricity = zeros(size(stack,3),1);
    for page = 1:size(stack,3)
20       time(page) = (page - timeOffset - 1)*timeInterval
        info = regionprops(stack(:, :, page), 'Area', 'Eccentricity');
22       for region = 1:size(info,1)
            if info(region).Area ~= 0
24                 area(page) = info(region).Area
            end
            if info(region).Eccentricity ~= 0
26                 eccentricity(page) = info(region).Eccentricity
            end
28       end
30   end
    % store time, area, and eccentricity values of object for each page
32   data = cat(2,time,cat(2,area,eccentricity));

34   % write time, area, and eccentricity to csv
    csvFilename = fullfile(folder,[ files(file, :) '.csv' ]);
36   dlmwrite(csvFilename,data,'newline','pc')
    disp(sprintf(['WROTE DATA TO FILE ' ]));
38
end

```

Listing 8. regionPropsScript.m