User manual

MSSegregation

Macro package for Fiji-ImageJ

ver 1.51

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Software and hardware requirements:

All MSSegregation macros run in Fiji-ImageJ application. Installation of Fiji-Image application before using MSSegregation macros is required (for details see the step-by-step guide of individual macros). The macros have been tested in Windows7/10, 64-bit, Fiji-ImageJ 1.53c

The MSSegregation package was run on two different notebook computers and three desktop computers (8Gb RAM, 64-bit Intel/AMD processors). We believe typical modern desktop computer can successfully run our macros. However, handling large image sets requires more powerful computer.

Availability

The ImageJ distribution package FIJI is freely available online: https://imagej.net/Fiji

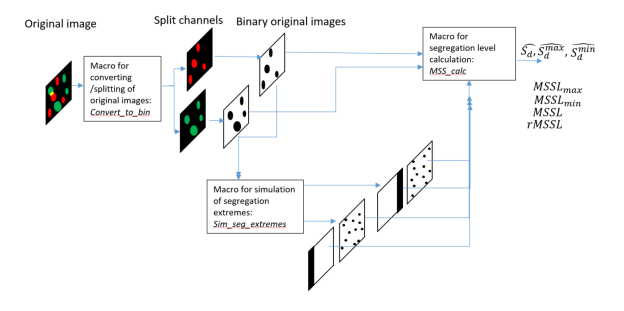
The MSSegregation package is freely available at https://github.com/IztokD/MSSegregation-package

Package: MSSegregation

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A brief overview

The purpose of MSSegregation package is to perform segregation analysis in digital images of two component mixtures (mixtures of two types of particles, particles and background...). MSSegregation package consists of three scripts that can be run as macros in Fiji (ImageJ) environment. The macro <code>Convert_to_bin_ver1.3.ijm</code> is used to split input images depicting two components into two separate images, where each image represent only one component. The macro <code>Sim_seg_extremes_ver1.3.ijm</code> simulates and depicts maximal and minimal segregation of two components system depicted in input digital images. Based on digital images of individual components, their maximal segregation and minimal segregation images, the Macro <code>MSS_calc_ver1.34.ijm</code> can perform spatial scale segregation analysis. The workflow of multiscale spatial segregation analysis of digital images is shown in the next page. For more details about usage, please refer to the individual descriptions of macros.



Workflow of multiscale spatial segregation analysis of digital images. The original image containing two types of particles (for example, microscopy image displaying red and green bacteria) is split into single type particle images, following conversion into binary images (i.e. black & white). Binary original images serve as input for macro that generates maximum and minimum segregation images for each type of the particle. In such images the number of displayed particles is typically proportional to the area occupied by the pixels representing them. From binary original images the segregation level (Sd^{Λ}) as a function of dimension (d) of field of view (FOV) is calculated; maximum and minimum segregation images are used for calculation of maximum and minimum segregation level ($Sdmax^{\Lambda}$, $Sdmin^{\Lambda}$). Finally, multiscale spatial segregation levels (MSSL), relative multiscale spatial segregation level (rMSSL) are calculated from segregation levels.

Macro 1: Convert_to_bin_ver1.3.ijm

Description

This script runs as a macro in Fiji (ImageJ) environment. Its purpose is to split and convert a digital image that depicts a mixture of two components, (two types of particles, particles and background..., called *ch1*, *ch2* particles) into two images in a 8-bit binary format (black and white), where each image represent one type of the particle. The separation is done via user assisted thresholding of pixel values. These images can serve as input for Multiscale spatial segregation analysis implemented in script *MSS_calc_ver1.3* and in *Sim_seg_extremes_ver1.3* for more details refer to part of this manual corresponding to these macros). The stacks of images are supported by the macro.

Step by step guide

- 1. If you do not already have Fiji, download and install it from: https://imagej.net/software/fiji/downloads
- Run Fiji (ImageJ-win64.exe file, if you are running 64-bit version).
 If asked to make an update, you can choose »Remind me later«, the update is usually not needed to successfully run Convert_to_bin_ver1.3.ijm

A. Installation

3. In Windows explorer (or equivalent File browser) go to the folder where you have installed Fiji and find plugins folder within Fiji folder. In our case:

C:\Users\iztokd\Downloads\fiji-win64\Fiji.app\plugins

Create folder \MSSegregation within plugins folder

In running Fiji app press Ctrl+shift+M (=Install Macro) and choose a file Convert_to_bin_ver1.3.ijm .

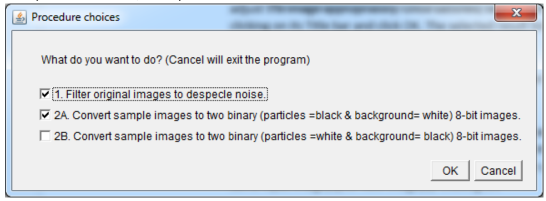
Save it to directory \MSSegregation, created above.

Close and restart Fiji app. If asked to save LUTMenuTool.ijm, just say "No".

- 4. Now you can run the macro from the Menu Bar: Plugins> MSSegregation>Convert to bin ver1.3.
- B. Direct run of the Macro
- 3. Open Windows explorer and "drag and drop" the file *Convert_to_bin_ver1.3.ijm* to the running Fiji application.

4. New window with displaying the code of *Convert_to_bin_ver1.ijm* will open (script editor) and you can run it by pressing Ctrl+R or by choosing Run>Run from menu of the script editor.

5. First you will be asked, what you want to do?



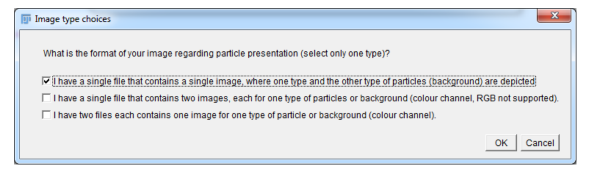
Option 1:

By default the filtering of original (input images) of pixel noise is selected. Usually this improves the output.

Option 2:

Also, because the main purpose of this macro is to obtain two binary images (8-bit), where on first image the particles of one type are shown (with bit value of 255) and on the second image the particles of other type are shown (with bit value of 255) the option 2 is selected by default. The background or anything which is not a particle of the particular type shown in the image will have bit value of 0. By (ImageJ) default the particles in these 8-bit images are displayed in black colour (option 2A), but you can also choose to display them in white (option 2B).

- 6. After you decided what you want to do, press "OK". Now you will be asked to choose a directory to save the output binary images. Select the directory and press "Select".
- 7. Now the menu "Image type choice" should appear:



Your input image(s) representing the mixture of two components (particles) can be of various formats:

Option 1:

user can have one file in which one digital image is decoded where both type of the particles are depicted (i.e. single file, single image). As the result of the opening procedure, two identical images will be displayed (duplicates), if the image is not RGB. In case of an RGB image, the original image will be split to three grayscale images (red, green, blue).

Option 2:

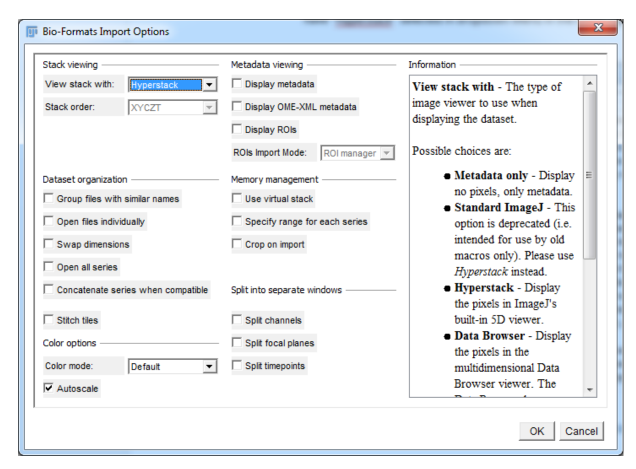
user can have one file in which two digital images are decoded, each for one type of the particles (i.e. singe file, two images). As the result of the opening procedure, two images will be displayed.

Option 3:

as last, you can have two files that each decode one image representing one type of the particle. As the result of the opening procedure, two images will be displayed.

In short, the user needs to know, which of the above holds for his input image and make the corresponding choice in the displayed menu and then click "OK".

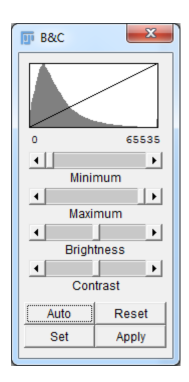
8. In next step select the corresponding image file and click "Open". Note that varies file formats are supported by ImageJ (tif, jpg...) and many more in Fiji through the plug-in Bio-formats that automatically starts, when file of particular format is to be opened (like in the case of Zeiss .czi format). If the Bio-formats menu pops up, you should not thick anything (you can leave Autoscale), but you need to have "Hyperstack" selected in dropdown Menu in the upper left corner. Once you click "OK" the next Bio-Formats window may appear, offering you a selection of the size of the images that are going to be opened. By default, choose ONLY the biggest resolution series.



9. In the case the user has opened an RGB image (*Option 1 in step 7*) three images will be displayed, representing red, green, blue channels. (S)he will be asked to close the image with the worst contrast of the particles of one type against the other type. So, the remaining two images, should show, each separately, as good as possible one type of the particles.

There are now two images opened (we will call them <code>image_ch1 & image_ch2</code>), each representing one type of the particle and to observe them in best possible contrast the auto-contrast option is automatically applied by the macro. Each of the two images will be now sequentially converted to two binary images.

10. Let us start with the *image_ch1*. The user can manually improve the contrast through B&C menu that appears simultaneously on the right hand side next to the image in focus, if this is necessary.

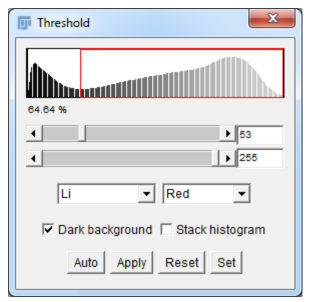


Unless the user is an expert, we suggest, only the sliders are adjusted in this menu if this is necessary at all. Once satisfied, the user should click "OK" in the simultaneously displayed window:



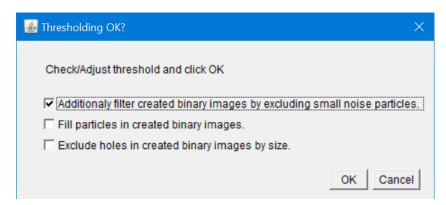
Note that contrasting is applied on selected window (*image_ch1*), which can be accidentally changed by clicking the other window (*image_ch2*). In the case you did accidentally click on the *image_ch2*, just click again on the *image_ch1* and then adjust contrast. Also, in the case you have two stacks of images (like in the case of CLSM images of biofilms composed from two bacterial strains, labelled with different fluorochromes), you can slide through the stack prior clicking "OK", just to make sure you adjusted the contrast satisfactorily through all slices in stack.

11. After clicking OK in previous step, the Threshold menu will open. The red area in the <code>image_ch1</code> denotes the particles that are supposed to be extracted from the <code>image_ch1</code>. By default it is assumed that the background in <code>image_ch1</code> is darker than particles. If this is not the case the user can uncheck the "Dark background" choice. By using the sliders in the Threshold menu the user can adjust the lower and upper boundary of pixel values to be included in the selection to select the particles of interest in the best possible way.



Unless the user is an expert, we suggest, only the sliders (and choice of Dark background) are adjusted in this menu if this is necessary.

Sometimes the thresholding cannot be set ideally- there are some parts of the particles that are not coloured red and some pixels or group if pixels that are not part of the particles of interest, but are coloured red. In this case it is a good idea to select Option 1 and 2 in the simultaneously displayed window:



Option 1:

The obtained binary image may display particles (noise, like some shadows...) that are not part of the "true" particles that you want to obtain. Often they are small in size, and can be filtered out. Choose this option if needed.

Option 2:

The obtained binary image may display particles that are the "true" particles that you want to display. However, some pixels that are part of them are missing (displayed as background, due to some

shadows etc.). If the extend of missing pixels is not too high, you can fill the missing parts by choosing this option. However, all holes in the particles will be filled regardless of their size.

Option 3:

This option differs from the above option. Here, you can specify the maximum size of the hole that will be filled.

Once satisfied with set thresholds, you can click "OK" to continue.

- 12. If you checked *Option 1* in previous step you will be asked to encircle the biggest particle (that are not of your interest) you want to remove. You can use the Oval or Rectangle tool, which you can select from ImageJ-Fiji toolbar. All particles of the encircled and smaller size will be removed after clicking "OK".
- 13. Now the result binary image is displayed. You will be asked if you are satisfied. If Yes, select the result binary image by clicking on its Title bar (the top of the image with title) and click OK. If Not, you can manually use additional built-in tools in ImageJ-Fiji to adjust the image appropriately. Once satisfied, select the result binary image by clicking on its Title bar and click OK. The selected result binary image will be saved to the save directory you have set in step 6 and closed.
- 14. The steps from 10 to 13 are now repeated, but this time the *image_ch2* is in the focus.
- 15. Once *image_ch1* and *image_ch2* had been converted to two binary images (*ch1_bin* and *ch2_bin*), where each image separately presents one type of the particle (or component, background) the user can choose either to process next image set (user can select again the same input image(s), or new image(s). Clicking OK brings the user again to step 5, while No or Cancel terminates the macro.

The saved output images from step 13 can serve as input images in macro *sim_seg_extremes* and *MSS_calc*.

Macro 2: Sim_seg_extremes_ver1.3.ijm

Description

This script runs as a macro in Fiji (ImageJ) environment. Its purpose is to simulate and depict maximal and minimal segregation of two components system (or particles of two types, called *ch1* and *ch2* particles) depicted in input digital images. Two input images (8-bit binary format, black and white) are required, each representing one type of the particle (*ch1_bin_, ch2_bin_*) in the same space. The two images can be obtained by the macro *Convert_to_bin_ver1.3.ijm*, but the user can obtain them also in other ways (e.g. manually by splitting via ImageJ tools). The output of the simulation are images (8-bit) representing minimal and maximal segregation case for each particle type (component type), together four images (*segmin_ch1_bin_, segmax_ch1_bin_* and *segmin_ch2_bin_, segmax_ch2_bin_*) stored to the disk as corresponding files. The particles are simulated as circles of user defined diameter; their number is obtained by dividing particle area in an input image by the sphere area. In addition, the macro can also simulate segregation for only part of the image (ROI) i.e. by creating the user defined mask. The masked original images are also outputted (*M_orig_ch1_bin_, M_orig_ch2_bin_*), together with the image representing mask itself (*Mask_*) and stored to the disk as corresponding files. Also, the stacks of images are supported by the macro. *segmin* and *segmax* files together *ch1_bin, ch2_bin* serve as input for *MSS_calc_ver1.3*, where Multiscale spatial segregation analysis is performed.

Step by step guide

- 1. If you do not already have Fiji, download and install it from: https://imagej.net/software/fiji/downloads
- 2. Run Fiji (ImageJ-win64.exe file, if you are running 64-bit version). If asked to make an update, you can choose »Remind me later«, the update is usually not needed to successfully run Sim seq extremes ver1.3, unless you are running really old version of Fiji-ImageJ.

A. Installation

3. In Windows explorer (or equivalent File browser) go to the folder where you have installed Fiji and find \plugins folder within Fiji folder. In our case:

C:\Users\iztokd\Downloads\fiji-win64\Fiji.app\plugins

Create folder (if it is not there already) \MSSegregation within plugins folder

In running Fiji app press Ctrl+shift+M (=Install Macro) and choose a file Sim seg extremes ver1.3.ijm

Save it to directory \MSSegregation, created above.

Close and restart Fiji app. If asked to save LUTMenuTool.ijm, just say "No".

- 4. Now you can run the macro from the Menu Bar: Plugins>MSSegregation> Sim seg extremes ver1.3
- B. Direct run of the Macro
- 3. Windows explorer (or equivalent File browser) and "drag and drop" the Sim seg extremes ver1.3.ijm to the running Fiji application.
- 4. New window with displaying the code of *Sim_seg_extremes_ver1.3.ijm* will open (script editor) and you can run it by pressing Ctrl+R or by choosing Run>Run from menu of the script editor.
- 5. Now you will be asked to choose a directory to save the output binary images, which are the result of the simulation. Select directory and press "Select".
- 6. Next you will be asked to select an image file that represent *ch1* particles. The image needs to be in 8-bit format. The particles must have pixel value of 255 the rest of pixels in the image must be 0. If you had used the macro *Convert_to_bin_ver1.3* than this is the file that ends with *_ch1_bin.tif* Click "OK" and select the file. Repeat the same for the image file that represent *ch2* particles.

7. Next the following question appears:

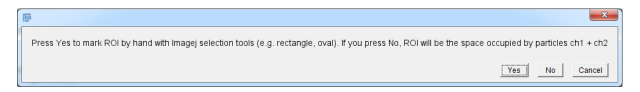


If you press "Yes" the macro will simulate segregation extremes only on the specified region (ROI = region of interest) of the input images from step 6. This can be handy, if the image has some background that you do not want to include into analysis and/or particles can occupy only one particular region in the image.

If you press "No" the macro will simulate segregation extremes on the entire area of input images from step 6. This is the most common situation. In this case you can go directly to step 9.

If you press "Cancel" the macro will terminate.

8. If you pressed "Yes" in previous step the following will appear on your screen:

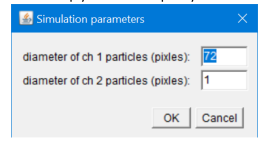


If you press "Yes" you will be asked to mark ROI on ch1 image (_ch1_bin). Use ImageJ-Fiji selection tools from the toolbar (e.g. Oval, Rectangle...) to encircle the ROI. The same ROI will be automatically copied and applied to ch2 image (_ch2_bin).

If you press "No", the ROI will become the space that is occupied by the particle *ch1* plus the space occupied by the particle *ch2*. This is useful, when two component or particles are restricted to certain area in the image. For example, oats and raisins can be restricted by the plate that occupies only part of the entire image. Oats or raisins cannot "move" out of the plate. Further example: you want to preserve the shape of two species biofilm in microscopy image and just simulate segregation within biofilm. You will be also asked to fill the holes in the common space. If they are not any holes present or the displayed holes are true empty space, just press No, otherwise press OK. If you are satisfied press OK, when asked about the result, No will undo fill.

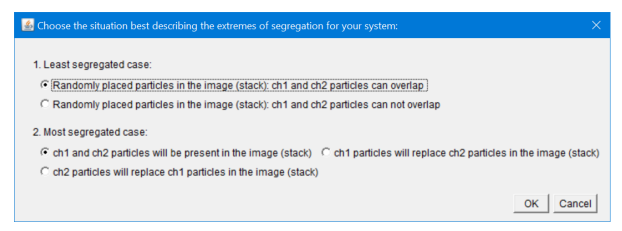
If you press "Cancel" the macro will terminate.

9. In this step you need to specify the diameter of the particles (component).



The particles can be only 1 pix in size, but can be also bigger. Only circular particles are supported at present time. It is advisable that user opens the image of particular particles in a separate instance of Fiji app and measures the particle size by, for example, by Straight tool from ImageJ-Fiji toolbar. However, if particles can be regarded as a continuous matrix than their size is the image resolution, i.e. 1 pixel. For example, in the case of mixture of oats and raisins, the oats can be considered as continuous matrix (diameter 1 pix) and raisins as circular particles(diameter 72 pix).

10. After you entered the diameter of respective particles and have pressed "OK" the following menu will appear:



1. Least segregated case:

Here you decide, if the randomly placed particles of one type (*ch1*) can overlap (i.e. share the same location in an image) with particles of other type (*ch2*) in the image. By default particles of the same type cannot overlap.

2. Most segregated case:

Here you decide about the type of maximal segregation:

The two type of particles (*ch1* and *ch2*) are restricted to the space of image (or ROI, step 8) and cannot "escape" from it. However, they can segregate to the sides of the image in order to maximize the distance from not alike particles. On the other hand, the alike particles will be placed maximally to each other, their individual shape will be ignored (the area of the particles is held constant).

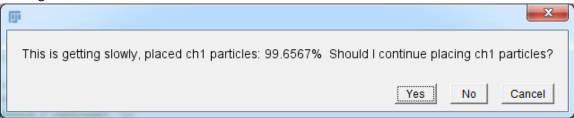
If the most segregated case is such that particles *ch1* can completely exclude particles *ch2* then choose the second option.

If the most segregated case is such that particles *ch2* can completely exclude particles *ch1* then choose the third option.

11. After clicking OK in the previous step the simulation of placing the particles in the image will begin, without displaying anything, except in Log window "start of the simulation:" together with time will be printed.

However, you may get warnings that "Total area of simulated spherical particles outlined by square exceeds the available image space". This can happen in the case you did not allow particle overlap in step 10 and the area of square outlined spherical particles is higher than total area of image or ROI. Outlining spheres by squares is very conservative approach in calculating total area, so there is a good chance that the simulation will be able to place all the particles, especially if at least one of the

particles is very small. Therefore, you should try and click "OK" and soon you will likely encounter a message like this:



This occurred because the space available to particles become very scarce during simulation and not all of the particles have been placed. If the total area of simulated particles exceeds the available image space than the placing all of the particles without overlapping will never happen. If the percentage is small, you should press Yes and see in the next message if the percentage improved significantly. If it did, than you should press Yes again and continue pressing Yes as long as you see significant improvements. Once you have reached high enough number of the % of placed (typically >95 %) you can press No to finish simulation and check the result images. On the other hand, if you did not observe significant steps in increase in % of placed particles after clicking Yes, you can either abort the macro by clicking Cancel and rerun the simulation with different parameters (allowing overlap, or different size of particle) or press No to finish simulation and check the result images.

- 12. Once all the particles (ch1 and ch2) have been placed or user agrees not to continue placing the particles (read step 11) the simulation finishes, prints to Log window "end of simulation:" together with time and displays the result images (three dots in the name means the name of the original image):
 - 1. Two Images showing minimally segregated cases (best mixed): segmin ch1 bin ..., segmin ch2 bin ...
 - 2. Two Images showing maximally segregated cases (worst mixed): segmax_ch1_bin_..., segmax_ch2_bin_...
 - 3. One Image showing the mask (i.e. the image that shows the area, where particles were allowed to be placed. The pixel value of 1 (black) indicates this area. If no restriction (ROI) of simulation space was selected (see step 7) then this image will be black:
 Mask_
 - 4. Two Images showing the original input images through the mask (if the mask was uniform black (no restrictions) than these images are the same as original input images):
 M orig ch1 bin ..., M orig ch2 bin ...
 - 5. One image showing overlap of *segmin_ch1_bin_...* and *segmin_ch2_bin_...* images (useful to see how severe overlap is, if it was permitted- black colour (pixel value 512) indicates overlapping of *ch1* and *ch2*, grey (pixel value 255) means particle is there, and white (pixel value 0) means empty space):

overlap of ch1 min seg and ch2 min seg

6. In addition, Log window is displayed, where most important data of simulation is displayed.

All of these images are stored as .tif files beginning with the above names in the save directory selected in step 5, together with Log window, stored as: $Log_Sim_seg_extremes_ver1.3.txt$. The image files from 1. and 2., together with $ch1_bin_$, $ch2_bin_$ image files serve as input files for macro $MSS_calc_ver1.3$, where Multiscale spatial segregation analysis is performed.

Macro 3: MSS_calc_ver1.34.ijm

Description

This script runs as a macro in Fiji (ImageJ) environment. Its purpose is to calculate the measures of segregation in images of two components system (mixture of two particles, particle and background....). Two components are called ch1 and ch2 (particles). The main quantity to be calculated is spatial size scale dependent segregation level (Sd^{\wedge}), i.e. Sd^{\wedge} as a function of size, d, of field of view (FOV). This is obtained by scanning the image by randomly placing the square of size $d \times d$ (= FOV) and calculating the dominance of one particle over the other in each placed FOV. This is done n times and then averaged to calculate Sd^ (for the details, please check the reference Dogsa et al., 2021, once it is published). To do this the two input images (8-bit binary format, black and white) are required, each representing one type of the particle (ch1 bin , ch2 bin) in the same space (like an image of oats-ch1 and raisins-ch2 in the same plate). The two images can be obtained by the help of macro Convert to bin ver1.3.ijm, but the user can obtain them also in other ways (e.g. manually doing the splitting and converting via ImageJ tools). To compare Sd^ of your sample to the positive and negative controls, i.e. to extremely segregated and extremely non-segregated cases in the same space by the same particles the images displaying these situations are required for ch1 and ch2 particles. These images (segmin ch1 bin, segmax ch1 bin and segmin ch2 bin , segmax ch2 bin) can be obtained by the macro Sim seg extremes ver1.3.ijm. The stacks of images are supported by the macro. Besides calculating the segregation level (Sd^) as a function of dimension of field of view (FOV), the macro can also calculate:

- multiscale spatial segregation level (MSSL), which is Sdⁿ averaged over all d (from dmin to dmax) of FOV to give distance averaged segregation level for users (sample) image and for min and max segregation images.
- 2. multiscale spatial segregation distance (*MSSD*), which is *Sd^* integrated over all *d* of FOV It can be interpreted as the distance (dimension) over which segregation level is 1 (i.e. maximal segregation level), if we theoretically compact all segregation levels, *Sd^*, from *dmin* to *dmax* to the distances from *dmin* to *MSSD*. This distance roughly corresponds to dimension of particles in the hypothetical image, where these particles form homogenous aggregates of *MSSD* and contain either one or the other type of the particle.
- Relative multiscale spatial segregation level (rMSSL). This segregation measure takes into
 account MSSL of users (sample) image and MSSL of min and max segregation images to
 calculate relative FOV independent segregation measure.

Step by step guide

1. If you do not already have Fiji, download and install it from: https://imagej.net/software/fiji/downloads

2. Run Fiji (ImageJ-win64.exe file, if you are running 64-bit version).

If asked to make an update, you can choose »Remind me later«, the update is usually not needed to successfully run MSS calc ver1.3, unless you are running really old version of Fiji ImageJ.

A. Installation

3. In Windows explorer (or equivalent File browser) go to the folder where you have installed Fiji and find \plugins folder within Fiji folder. In our case:

C:\Users\iztokd\Downloads\fiji-win64\Fiji.app\plugins

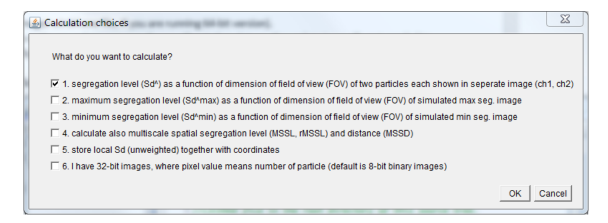
Create folder (if it is not there already there) \MSSegregation within plugins folder

In running Fiji app press Ctrl+shift+M (=Install Macro) and choose a file MSS_calc_ver1.34.ijm

Save it to directory \MSSegregation, created above.

Close and restart Fiji app. If asked to save LUTMenuTool.ijm, just say "No".

- 4. Now you can run the macro from the Menu Bar: Plugins> MSSegregation> MSS_calc_ver1.34
 - B. Direct run of the Macro
- 5. Open Windows explorer (or equivalent File browser) and "drag and drop" the MSS_calc_ver1.34.ijm to the running Fiji application.
- 6. New window with displaying the code of *MSS_calc_ver1.34.ijm* will open (script editor) and you can run it by pressing Ctrl+R or by choosing Run>Run from menu of the script editor.
- 7. Now you will be asked to choose a directory to save the output calculations, which are the result of the simulation. Select directory and press "Select".
- 8. Then you will be asked what to calculate.



Option 1:

This option is by default checked, as this is the most fundamental purpose of this macro. To calculate segregation level (Sd^{Λ}) , you will need two binary images, each representing one type of the particle or component in the common mixture, which can be obtained by splitting and converting the original sample image by $Convert_bin_ver1.3.ijm$ macro.

Option 2:

This option is used to compare calculated segregation level of sample image to the positive control, i.e. to the segregation level of hypothetically maximally segregated sample image. This image can be obtained by Sim_seg_extremes_ver1.3.ijm macro.

Option 3:

This option is used to compare calculated segregation level of sample image to the negative control, i.e. to the segregation level of hypothetically minimally segregated sample image. This image can be obtained by Sim_seg_extremes_ver1.3.ijm macro.

Option 4:

If the Option 1 to 3 is checked the macro will calculate the segregation measures derived from segregation level, Sd^{Λ} .

Option 5:

If you select this option the local Sd (unweighted) will be stored for each field of view (FOV) together with the center coordinates of FOV and corresponding weights. For each size, d, of FOV, a separate output file will be written to the disk. Note that Sd^{Λ} is weighted average of local Sd.

Option 6:

By default the input images need to be 8-bit with pixel value of 255 indicating the presence of the particle and 0 of its absence. If this option is checked, the user can also input 32-bit images, where pixel value represents the number of particles in the location of the pixel (currently, this option is under development and is not fully tested).

Pressing "OK" will get you to the next step, while pressing "Cancel" will exit the macro.

- 9. Now you will be asked to select a directory to save the calculation results. Click on the chosen directory and press "Select".
- 10. Next you will be asked to select an image file that represent ch1 particles. The image needs to be in 8-bit format (unless you have checked Option 5 in step 6). If you had used the macro Convert_to_bin.ijm than this is the file that ends with _ch1_bin.tif. Click "OK" and select the file. Repeat the same for the image file that represents ch2 particles.

- 11. If you have checked Option 3 and 4 in step 6, you will be asked to choose corresponding 8-bit (32-bit if you have checked option 5 in step 6) images, representing maximal and minimal segregation cases. If these images were obtained by the macro <code>Sim_seg_extremes_ver1.3.ijm.</code>, then the corresponding files are named as: <code>segmax_ch1_bin_....tif</code>, <code>segmax_ch2_bin_....tif</code> and <code>segmin_ch1_bin_....tif</code>, <code>segmin_ch2_bin_....tif</code>). For each image click "OK" and select the file.
- 12. In this step you can adjust calculation parameters.

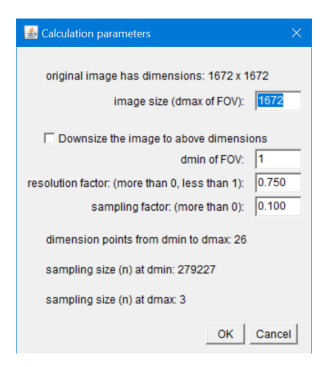


Image size (dmax of FOV) [pixels]:

By default, image size in pixels is displayed corresponding to the maximum dimension (dmax) of FOV that will be applied to an image to calculate FOV dimension dependant segregation level, Sd^{Λ} . The user can decrease the dmax of FOV for segregation level calculation, but in this case (s)he will not obtain the segregation level, Sd^{Λ} of entire area of image.

Downsize the image to above dimensions:

If the time taken for the calculation is very long and the image has unnecessary high pixel resolution, the user can downsize the input images to new *dmax* [pixels]. In this case the *Sd*^ of entire area of image can be obtained.

dmin of FOV [pixels]:

The user can set the smallest dimension (dmin) of FOV. This can improve the calculation speed significantly, if Sd^{Λ} at dmin of 1 pix is not necessary to be calculated, than this number should be increased. For example, in the mixture of oats & raisins that are in size about 50 pix, there the Sd^{Λ} at dmin of 1 pix will for sure be 1 as the two constituents cannot interpenetrate each other.

resolution factor:

This factor determines the number of different dimensions, d of FOV at which Sd^{Λ} will be calculated and by which factor the values of d will be spaced, in the direction from dmax to dmin. For example factor of 0.99, means that starting from dmax, every next d will be smaller by factor of 0.99 in the direction to dmin. Unless the range from dmax to dmin is very low, this means a lot of d points.

sampling factor:

This factor determines the number, n, of local segregation levels will be calculated from randomly placed FOVs of dimension d in the image , which get then averaged to obtain Sd^n at d. The more the better the statistics, but of course more time is required for the calculation. Maximal sampling size per d is 1 000 000, but often a few thousands will be enough. This factor impacts the duration of the calculation significantly, so try first with n about few thousands and check the estimated experimental error of the calculated quantities in the end of simulation, maybe they are small enough and you will not need to increase sampling factor.

Note that the total time of calculation depends on the number of points to be calculated (distance points from *dmax* to *dmin*), the number of sampling at each point (sampling size, *n*, and dmin and dmax) and the resolution of image.

If the changes were made to whatever parameters default values in this menu, the parameters (distance points from dmax to dmin), (sampling size at dmin and dmax) will get refreshed after clicking "OK". Once no more changes to the parameters values (and parameter values are within their ranges) had been made, clicking "OK" will start the calculations and calculation start time will appear in the Log window.

During the calculations, the user can follow the progress in Log window. For each calculation point the data about the dimension, d, and number, n, of FOV (sampling size) will be displayed in Log window. Once the Sd[^] at certain d is calculated, the macro proceeds to calculate next Sd[^] at next d, until all Sd^ from dmax to dmin are calculated. When this happens the calculation end time will appear in the Log window and the calculation results will be displayed in Results window, including all the segregation measures selected in step 6 together with their standard errors. The Result table and Log window are saved to the saving directory specified in step 6, having the names Sd_MSSDL_ver1.3.txt Log_Sd_ver1.3.txt. addition, the file and In statistics of segregation Sd ver1.3.txt containing additional segregation indices that can help very advanced users to further analyse their segregation cases is also stored to the saving directory.

Step 11: As a last step, the user can choose that the segregation level as a function of *d* of FOV gets plotted, which will be than displayed.

Examples

This section comprises three examples of MSSegregation package application. For each macro applied the input images entered at certain step along with input parameters are given. Also, the output images and most important results at certain steps of the macros are shown. The order of appearance of macros within the particular examples is the same as the run order of the macros. To understand how to run and use certain macros, refer to the previous sections of this manual. The input/output files for all three examples are given on github: https://github.com/lztokD/MSSegregation-package. You can also watch online movies, where we have shown how to remake this available on FigShare (the links are given in the text below). In Example 3 we have used free online external application to plot choropleth maps (https://www.datawrapper.de), but in general the user can obtain the choropleth maps also in other ways (e.g. by other software, drawing by hands, copying from digital sources...).

Example 1: Oats & Raisins

Convert_to_bin_ver1.3.ijm

Parameters used: step 5 (*Option 1 and 2*), step 7 (*Option 1*), step 9: (closed image = blue), step 11 for image (red) = ch1: *Option 1, 2*, lower-threshold [upper slider] = 1, upper-threshold [lower slider] = 61; for image (green) = ch2: *Option 1, 3*, lower-threshold = 67, upper-threshold = 255

Movie showing the procedure:

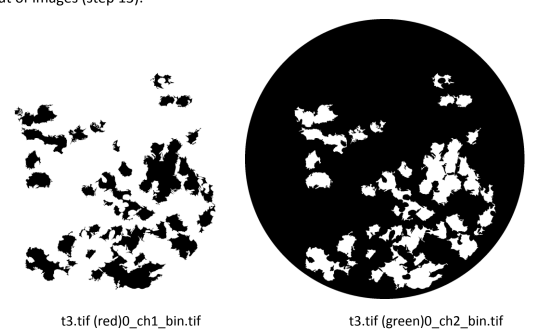
Convert_bin_oats_rasins.wmv at https://figshare.com/s/5a08f7b84a76586e1601

Input images (step 8):



t3.tif (RGB image)

Output of images (step 13):



Sim_seg_extremes_ver1.3.ijm

Parameters used: step 7 (Yes), step 8 (No), step 9: ch1 = 72 pix, ch2 = 1 pix, step 10: Randomly placed particles in the image (stack): ch1 and ch2 particles can not overlap, ch1 and ch2 particles will be present in the image (stack), step 11, (No, after >99% ch2 particles placed).

Movie showing the procedure: segregation_extremes_oats_raisins.wmv at https://figshare.com/s/5a08f7b84a76586e1601

Input images (step 6):

Output of Convert_to_bin_ver1.3.ijm:

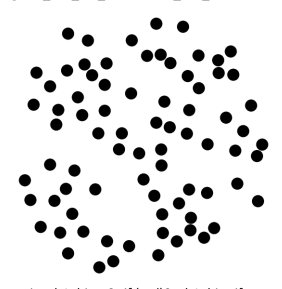
t3.tif (red)0_ch1_bin.tif

t3.tif (green)0_ch2_bin.tif

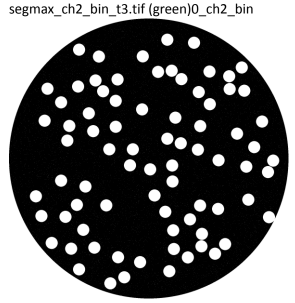
Output images (step 15):



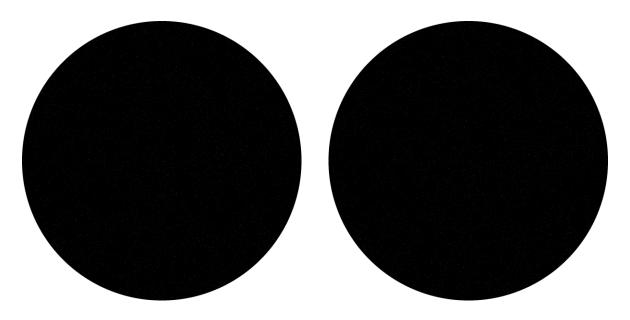
segmax_ch1_bin_t3.tif (red)0_ch1_bin



 $segmin_ch1_bin_t3.tif (red)0_ch1_bin.tif$

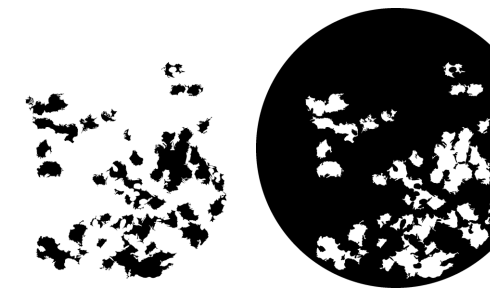


 $segmin_ch2_bin_t3.tif (green)0_ch2_bin.tif$



overlap of ch1 min seg and ch2 min seg.tif

Mask_t3.tif (red)0_ch1_bin t3.tif (green)0_ch2_bin.tif



 $M_orig_ch1_bin_t3.tif (red)0_ch1_bin.tif$

M_orig_ch2_bin_t3.tif (green)0_ch2_bin.tif

MSS_calc_ver1.34.ijm

Parameters used: step 6 (Option 1 to 4), step 10 (dmax = 1663, dmin = 5, res_factor = 0.75, sampling_factor = 0.01), step 11 (Yes)

Movie showing the procedure:

MSS analysis oats raisins.wmv at https://figshare.com/s/5a08f7b84a76586e1601

Input images (step 8 and 9):

1. Output of Convert_to_bin_ver1.3.ijm:

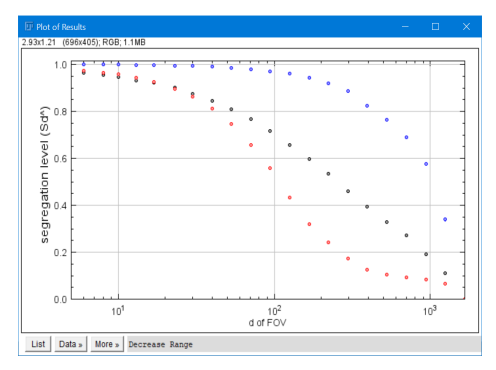
t3.tif (red)0_ch1_bin.tif t3.tif (green)0_ch2_bin.tif

2. Output of Sim_seg_extremes_ver1.3.ijm:

```
segmax_ch1_bin_t3.tif (red)0_ch1_bin.tif segmin_ch1_bin_t3.tif (red)0_ch1_bin.tif segmin_ch2_bin_t3.tif (green)0_ch2_bin.tif
```

Output result (most important):

-the graph of segregation level vs d of FOV for user's sample image (= Your image, black), max seg image (blue) and min seg image (red), (step 11, Yes):



• Sd^min, • Sd^ (user's sample image) and • Sd^max Part of the Results table showing average segregation measures:

	1.412E-1	min image:	5.753E-1	max image:	2.744E-1	your image:	multiscale spatial segergation level(MS	File Edit Font Results	□ Results
	2.141E-2	min image:	8.628E-2	max image:	4.127E-2	your image:	SL) standard ero		
	2.339E2	min image:	9.532E2	max image:	4.546E2	your image:	r multiscale spatial segergation distance (MSSD)[
	3.548E1	min image:	1.430E2	max image:	6.839E1	your image: 3.068E-1	multiscale spatial segergation level(MSSL) standard eror multiscale spatial segergation distance (MSSD)[pix] standard eror relative multiscale spatial segergation level (rMSSL-your im standard eror		
v [1]						1.105E-1	ur im standard eror 🔺		- 0 ×

Example 2: bacterial biofilm composed of two strains (microscopy)

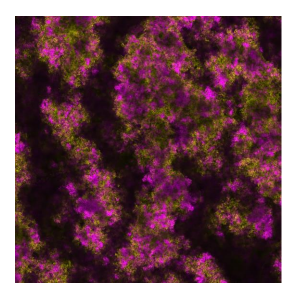
Convert_to_bin_ver1.3.ijm

Parameters used: step 5 (*Option 1 and 3*), step 7 (*Option 2*), step 11 (images ch1 and ch2: default threshold of Li Method applied on middle slice) (No other *Options*).

Movie showing the procedure:

Convert_bin_biofilms .wmv at https://figshare.com/s/5a08f7b84a76586e1601

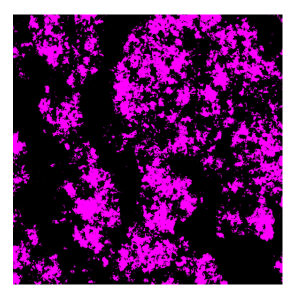
Input images (step 8):

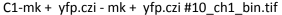


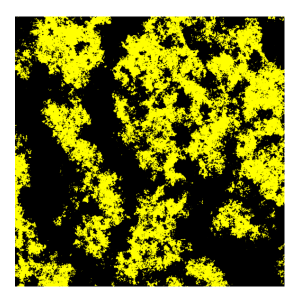
mk + yfp.czi (Zeiss. czi image stack of 17 slices, only middle slice (=9) is shown)

Note that images in this example where pseudolocored in Images by ch1 = pink and ch2 = yellow. This is not necessary in order to correctly perform MSS segregation analysis.

Output of images (step 13):







C2-mk + yfp.czi - mk + yfp.czi #10_ch2_bin.tif

Sim_seg_extremes_ver1.3.ijm

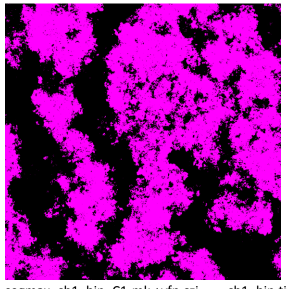
Parameters used: step 7 (*Yes*), step 8 (*No*), step 9: ch1 = 3 pix, ch2 = 3 pix, step 10: *Randomly placed particles in the image (stack): ch1 and ch2 particles can overlap, ch1 particles will replace ch2 particles in the image (stack)*, step 11, (*Yes*, after cca. 75% ch1 particles placed, the rest can be placed by partial overlap; the same for ch2).

Movie showing the procedure:

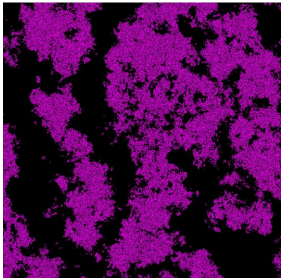
segregation_extremes_biofilms.wmv at https://figshare.com/s/5a08f7b84a76586e1601

Input images (step 6):

Output of Convert_to_bin_ver1.3.ijm:



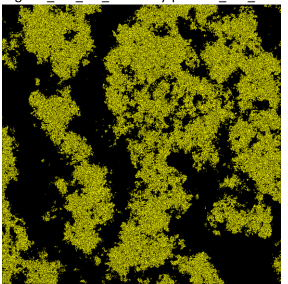
segmax_ch1_bin_C1-mk +yfp.czi -... _ch1_bin.tif



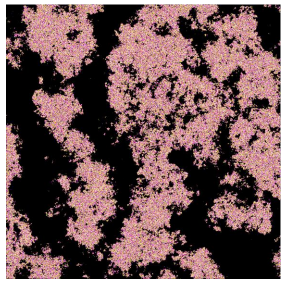
segmin_ch1_bin_C1-mk +yfp.czi -... _ch1_bin.tif

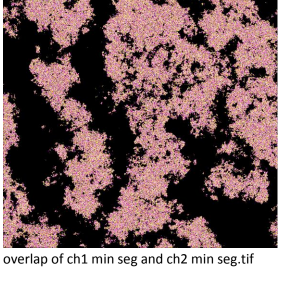


segmax_ch2_bin_C2-mk + yfp.czi - ..._ch2_bin.tif

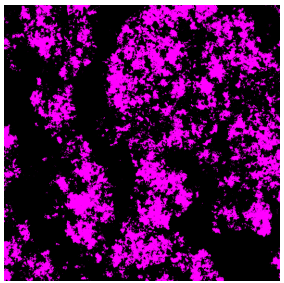


segmax_ch2_bin_C2-mk + yfp.czi - ..._ch2_bin.tif

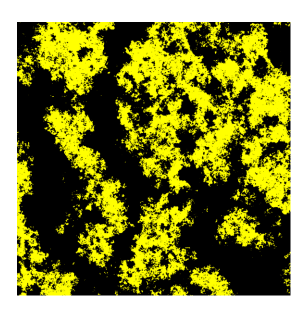




Mask_C1-mk + yfp.czi -mk + yfp... _ch2_bin.tif



 $M_orig_ch1_bin_C1-\ mk+yfp.czi-..._ch1_bin.tif$



 $M_orig_ch2_bin_C2\text{-}\,mk + \,yfp.czi\,\text{-}..._ch2_bin.tif$

MSS_calc_ver1.34.ijm

Parameters used: step 6 (Option 1 to 4), step 10 (dmax = 1964, dmin = 7, res_factor = 0.75, sampling_factor = 0.001), step 11 (Yes)

Movie showing the procedure:

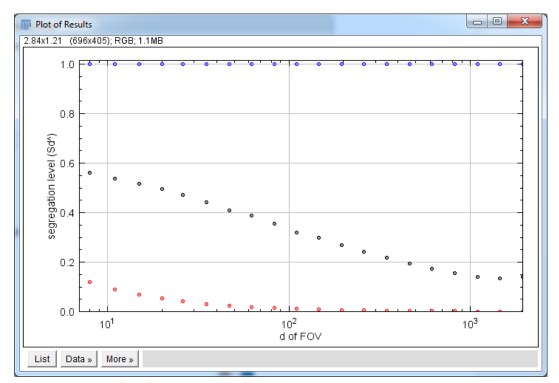
MSS_analysis_biofilms.wmv at https://figshare.com/s/5a08f7b84a76586e1601

Input images (step 8 and 9):

- 1. Output of *Convert_to_bin_ver1.3.ijm*: C1-mk + yfp.czi mk + yfp.czi #10_ch1_bin.tif C2-mk + yfp.czi mk + yfp.czi #10_ch2_bin.tif
- 2. Output of Sim_seg_extremes_ver1.3.ijm: segmax_ch1_bin_C1-mk +yfp.czi -..._ch1_bin.tif segmax_ch2_bin_C2-mk + yfp.czi ..._ch2_bin.tif segmin_ch1_bin_C1-mk +yfp.czi -..._ch1_bin.tif segmin_ch2_bin_C2-mk + yfp.czi ..._ch2_bin.tif

Output result (most important):

-the graph of segregation level vs d of FOV for user's sample image (= Your image, black), max seg image (blue) and min seg image (red), (step 11 , Yes) :



Sd[^]min,
 Sd[^] (user's sample image) and
 Sd[^]max

Part of the Results table showing average segregation measures:

File Edit Font Results multiscale spatial segergation level(MSSL) Standard error of MSSL your image: 1.788E-1
your image: 9.191E-3 max image: 0.000E0 min image: 5.296E-4
your image: 3.469E2 max image: 1.940E3 min image: 0.000E0 min image: 1.187E0 wour image: 1.783E1 your image: 1.783E0 1.783E0 1.027E0
relative multiscale spatial segergation level (rMSSL-y 1.757E-1
our Im standard eror of rMS 9241E-3