Quantification of smFISH images (How to use)

Before start

- Download the repository or the content of this folder.
- unzip compressed tif image files.

Descriptions

This quantification pipeline is divided into 5 parts:

- 1st part: oocytes outlines delimitation and extraction.
- 2nd part: image preprocessing.
- 3rd part: condensates segmentation.
- 4th part: generation of outlines and filtered images for smFISH spots detection in the cytosol.
- 5th part: smFISH spots detection and filtering.

Requirements:

- MATLAB (tested v R2018b or higher)
 - Required toolboxes:

Image Processing Toolbox

Statistics and Machine Learning Toolbox

Parallel Computing Toolbox

Curve Fitting Toolbox

Optimization Toolbox

• Additional requirements:

FISH-quant (1,2,#)

https://bitbucket.org/muellerflorian/fish_quant/src/master/

- Fiji (ImageJ)
- R
- Optional: RStudio

Set up:

- Copy the scripts provided (*/scripts_smFISH_quantification) to the MATLAB folder (usually located in Documents directory).
 - Note: overlapping code scripts from other quantifications do not need to be replaced.
- Install FISH-quant:
 - Go to https://bitbucket.org/muellerflorian/fish_quant/src/master/ (1,2).
 - Click Downloads and then download the repository (1,2).
 - Unzip the file in the MATLAB folder (Documents directory) (1,2).
 - Go to ./Documents/MATLAB/FISH_quant/Documentation (1,2).
 - Open FISH-QUANT__Tutorials.pdf (1,2).
 - Follow the section 1.1 Install FISH-quant for Matlab (1,2).

Set up:

How does this pipeline work?

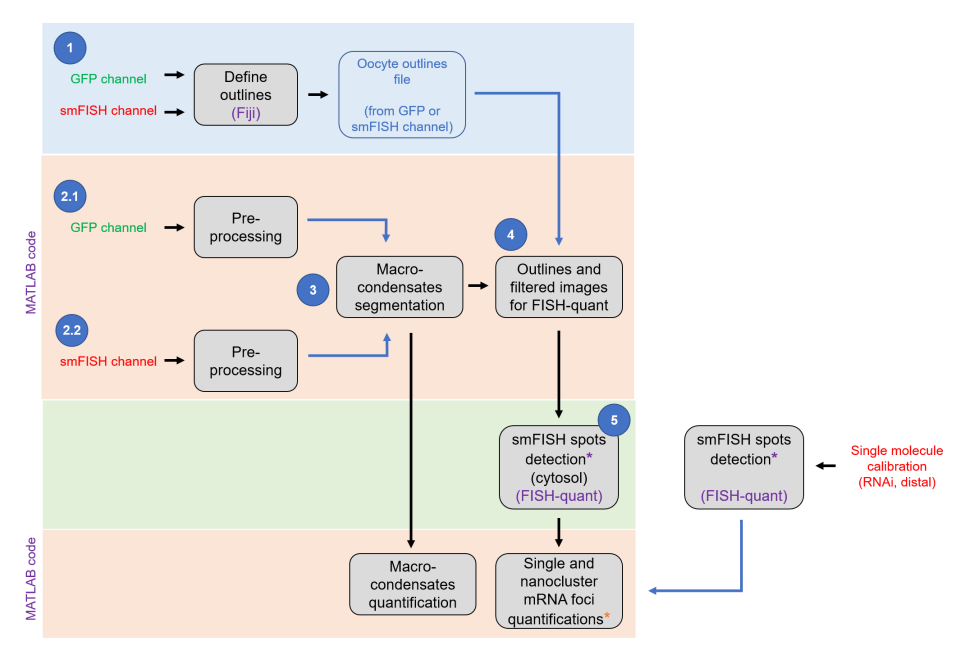


Figure 1. smFISH image quantification pipeline. It uses Fiji, FISH-quant, and custom MATLAB code.

(*) These steps can be done in batch. For the rest of steps, several MATLAB sections can be opened simultaneously.

Provided image examples

The image data is provided at ./example image

It consists of two channels (already separated as independent images):

- GFP channel: "w4_GFP.tif"
- smFISH channel: "w4_Spn4.tif"

An example oocyte outlines file (see Fig. 1) is also provided:

• Oocyte outlines: "w4_GFP.zip"

User guide

1st part: oocyte outlines delimitation and extraction

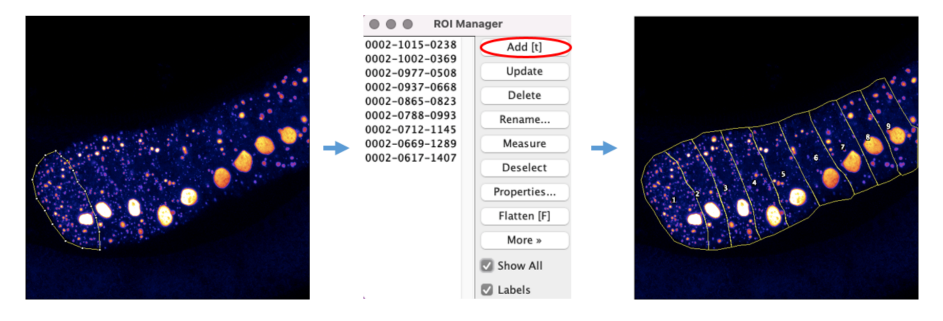
Requirements:

- Fiji (Image J)
- R
- Optional: RStudio

Outlines delimitation

Procedure:

- 1. Open Fiji (ImageJ).
- 2. Open RIO Manager
 - $\circ \ \ \mathsf{Analyze} > \mathsf{Tools} > \mathsf{RIO} \ \mathsf{Manger...}$
- 3. Using Fiji, open the image you want to analyze.
- 4. Use the "polygon selections" tool to manually draw outlines that delimit the oocyte borders.
 - Each **oocyte outline** can be treated as an individual **R**egion **O**f **I**nterest (**ROI**).
 - To delimit oocytes, a max projection could be useful as gonad dimensions are variable along the z-plane.
 - Each oocyte outline (ROI) can be saved by clicking on the "Add [t]" bottom located in the RIO Manager window, as shown below:



- 5. Once all outlines are added, select them and save them as a ".zip" file. **Important:** name the file with the same name as the image where the outlines are coming from.
 - o In the case of only one outline, it will be saved as a "roi" file.

Extraction of xy coordinates from outlines (required for 4rd part)

Example data and description:

Example images and outline file for this guide can be found in the folder ./example_image
Images were obtained imaging a smFISH experiment. It consists of two channels (already separated as independent images):

- GFP channel: "w4_GFP.tif"
- smFISH channel: "w4_Spn4"

An example oocyte outlines file (see Fig. 1) is also provided. You can obtain it as explained in the previous section (Outlines determination)

• Oocyte outlines: "w4_GFP.zip"



Procedure:

6. Data should be organized in a folder as just shown above:

For each "name".zip outline file, a matching "name".tif image must be in the folder.

- For this example, only one outline file will be processed, alternatively several outline files with their corresponding images can be processed simultaneously.
- 7. Using Fiji, open the script BATCH_improved_outlines_oocyte_extraction.ijm (location: ./image_outlines)
- 8. Run it and select the folder containing the images and oocyte outlines (ROIs) to analyze.

This script extracts the xy coordinates from the ROIs delimited and saved using Fiji.

• Output: "name"_Cells.csv files with xy coordinates of ROIs.



- 9. Using R or RStudio, open the script BATCH_automatic_outlines_reorganization_by_columns.R (location: ./image_outlines)
- 10. Run the script and select any file within the folder containing the images and ROIs to analyze.

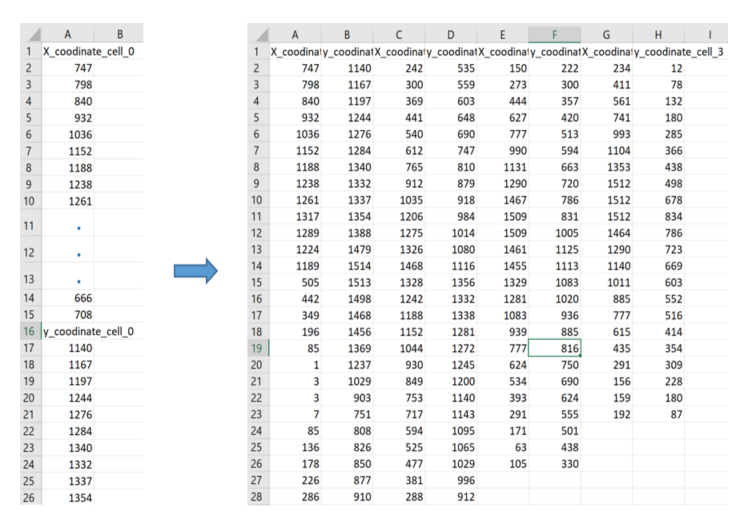
This script reorganizes the outlines files, so they are readable in the image processing step.

• Output: MOD_"name"_Cells.csv files with reorganized xy coordinates of ROIs.



Optionally

You can do this reorganization manually as shown below:



How to modify BATCH_improved_outlines_oocyte_extraction.ijm

```
//... Modify if needed (File Identifiers)...
file_ext = "zip"; // ROI file extension
//...
// Only line 2 should be modified. If needed replace "zip" by "roi".
```

2nd part: image preprocessing

Procedure. For each channel, GFP and smFISH, do the following:

- 11. Open MATLAB
- 12. In the command window type:
 - edit smFISH_preprocessing_Script

This file will open:

```
----- smFISH preprocessing
    % Description of the stack
    % There are two possibilities:
    % = 1 if stack has beginning and ending frames out of focus (whole gonad)
    % = 0 if stack is in focus (usually a z-section within the gonad)
    is_stack_out_of_focus = 1; %
    smFISH_channel = 0; % type of image: if FISH = 1, if GFP = 0
    % Parameters
    quantile_BGD_subtraction = 0.9; % quantile to subtract BGD value (initial filtering), 0 if it's unknown
    % if quantile_BGD_subtraction = 0, two BGD approximations are computed:
    different_BGD_approximations = 1; % 1 for different functions, 2 for same function with different mask
sizes
    % image ranking
    block_size_for_image_ranking = 2; % faster results with larger values like 4, 8, 16; but less detail
    % run
    smFISH_image_Preprocessing
```

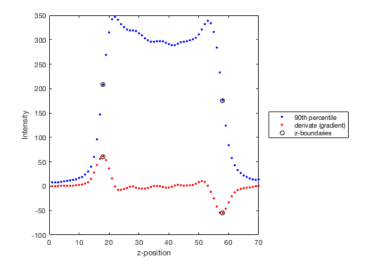
- 13. If needed, modify the parameters for preprocessing, then save changes:
 - Line # 6: is_stack_out_of_focus
 This scrip is optimized for two options either

This scrip is optimized for two options, either a complete gonad in the z-plane with out-of-focus planes (is_stack_out_of_focus = 1), or a z-section with NO out-of-focus planes (is_stack_out_of_focus = 0).

```
o Line # 7: smFISH_channel
Make smFISH_channel = 1, for smFISH channel
Make smFISH_channel = 0, for GFP channel
```

Line # 10: quantile_BGD_subtraction
 quantile_BGD_subtraction = 0.9 is recommended for most cases.
 To select a value during the preprocessing make quantile_BGD_subtraction = 0

- Line # 15: block_size_for_image_ranking = 2
 Faster results with larger values like 4, 8, 16; but less detail.
- 14. In the command window type: smFISH_preprocessing_Script
- 15. The script will ask you to select either a GFP or smFISH image.
 - The preprocessing steps are the same, the distinction is only used to name the outputs properly.
- 16. If <u>is_stack_out_of_focus</u> = 1, The script determines the oocyte z-boundaries of the stack (oocytes z-edges), as follows:

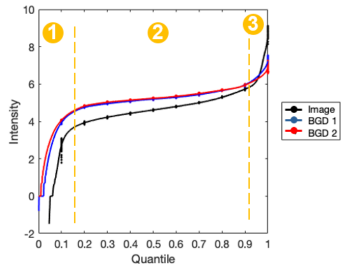


*** This method won't work if the z-stack acquisition exhibits bleaching in the z-plane or if it doesn't have out-of-focus frames both at the beginning and end of the z-stack.

17. Then:

if quantile_BGD_subtraction = 0.9 if quantile_BGD_subtraction = 0

- The script will perform two background approximations (BGD1 and BGD2):



- The script will subtract the background.
- Image ranking will be performed.
- And output will be saved.

The curve displays three regions:

- 1. slide background,
- 2. oocyte background.
- 3. brighter objects like smFISH clusters.

quantile_BGD_subtraction should be the quantile that separates region 2 from region 3.

By default, quantile_BGD_subtraction = 0.9

- Type in the command window the quantile you want.
- The script will subtract the background.
- Image ranking will be performed.
- And output will be saved.

By channel the script will save a MATLAB workspace. For instance:

- WS_name-smFISH-cahnnel_FISH.mat
- WS_name-GFP-cahnnel_GFP.mat

3rd part: condensates segmentation

Example data and description

For this step, the following files are required:

```
    location: ./example_image

            GFP channel image
            smFISH channel mage

    location: ./example_image/output_image_preprocessing (2nd part)

            WS_name-smFISH-cahnnel_FISH.mat
            WS_name-GFP-cahnnel_GFP.mat

    location: ./example_image/output_outlines_reorganization (1st part)

            MOD_"name-channel"_Cells.csv
```

For this example, the, the following files are provided:

```
w4_Spn4.tif
w4_GFP.tif
WS_w4_Spn4_FISH.mat
WS_w4_GFP_GFP.mat
MOD_w4_GFP_Cells.csv
```

Procedure

- 18. Make a folder with the 5 elements described in "example data and description".
- 19. Type in the command window:

```
edit smFISH_Segmentation_Outlines_Script
```

20. The script below will appear (If needed, modify accordingly to instructions). Then save changes.

```
% ----- smFISH image analysis (segmentation)
%% File ID indicators
files = struct; % do not modify
files.FISH_img = '*Spn4*.tif'; % file identifier for smFISH image
files.FISH_ws = '*FISH*.mat'; % file identifier for smFISH workspace
files.GFP_img = '*GFP*.tif'; % file identifier for GFP image
files.GFP_ws = '*GFP*.mat'; % file identifier for GFP workspace
files.outlines = 'MOD*Cells.csv'; % file identifier for cell outlines
%% Microscope parameters
% Define pixel size in nm
pixel_size = struct; % do not modify
pixel_size.xy = 49; % pixel size in x and y
pixel_size.z = 250; % pixel size in z
% Acquisition parameters
s = struct; % do not modify
s.Em = 568; % cy3 Emission
s.Ex = 554; % cy3 Excitation
s.NA = 1.4; % Numerical aperture
s.RI = 1.518; % Diffractive index
s.type = 'confocal'; % Microscope type
s.Pixel = pixel_size;
%% Segmentation parameters
options = struct; % do not modify
options.Int_Threshold_GFP = 0.6; % Normalized [0 - 1] Intensity Threshold: GFP
options.Int_Threshold_FISH = 0.8; % Normalized [0 - 1] Intensity Threshold: smFISH
options.Volume_Threshold = 32; % Volume threshold for segmentation (in pixels)
                              = 1; % if 1, condensate are dilated to include edges
options.Dilate_Granule
% Outlines settings
options.FQ_outline
                           = 1; % if 1, outlines for FQ are an output
options.Exclude_Granules = 1; % if 1, creates images and outlines without granules
options.Save_Coordinates = 1; % if 1, segmented objects coordinates for are saved
%% Alternative masking
Filt_options = struct; % do not modify
Filt options.gauss2x = 1;
Filt_options.bgd_xy = 5;
Filt_options.bgd_z
Filt options.spots xy = 0.1;
Filt_options.spots_z = 0.1;
Filt_options.output = 0;
options.Filt
                        = Filt_options;
```

```
%% RUN
Segmentation_Outlines_run
```

• The following values should be adapted to different experimental conditions, for example; a different mRNA target, type of microscopy, among others. However, should be maintained if you are analyzing the same experiment.

```
o options.Int_Threshold_GFP = 0.6
```

Normalized intensity threshold for segmentation of the GFP image.

Values accepted from 0 to 1. Values close to 0 would be background while values close to 1 will be condensates.

o options.Int Threshold FISH = 0.8

Normalized intensity threshold for segmentation of the smFISH image.

Values accepted from 0 to 1. Values close to 0 would be background while values close to 1 will be condensates.

o options.Volume_Threshold = 32

Volume threshold for segmentation (in pixels).

o options.Dilate_Granule = 1

if = 1, condensates are dilated to make sure edges are included. This is a very important step as edges not segmented might generate false positives while detecting cytosolic spots. **IMPORTANT:** Dilation needs to be done by condensate, so this step might take time depending on computer specifications.

• If you need to make test without running the whole script (specially the dilation that takes most of the time), in step 18 type in the command window:

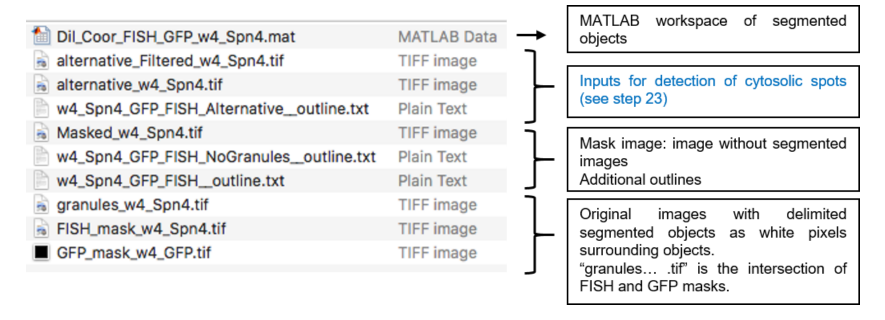
```
edit smFISH_Segmentation_Script_TEST
```

21. Accordingly, type in the command window:

```
smFISH_Segmentation_Outlines_Script
or
smFISH_Segmentation_Script_TEST
```

22. Select the folder created in step 18.

Outputs



4th and 5th part:

Generation of outlines and filtered images for smFISH spots detection in the cytosol, smFISH spots detection and filtering.

The segmentation script also generates the outlines, raw image, and filter image required to run FISH-quant (1-3) to detect smFISH spots. For more information, refer to the already published manuals (1-3).

- 23. In order to detect cytosolic smFISH spots after running this segmentation (Fig. 1), please use the following files as FISH-quant inputs (**outputs from 3rd part: condensates segmentation**):
- alternative... .tifimage
- alternative_Filtered... .tifimage
- ...Alternative__outline.txt outline file
- 24. Once in FISH-quant(1-3), single or batch analysis modes, do not perform the filtering step. If you want to modify the filtering parameters go to step **20** and make changes as follows:

```
%% Alternative masking
Filt_options = struct; % do not modify
Filt_options.gauss2x = 1;
Filt_options.bgd_xy = 5; % Kernel BGD [pixel]: XY (see FISH-quant)
```

```
Filt_options.bgd_z = 5; % Kernel BGD [pixel]: Z (see FISH-quant)
Filt_options.spots_xy = 0.1; % Kernel SNR [pixel]: XY (see FISH-quant)
Filt_options.spots_z = 0.1; % Kernel SNR [pixel]: Z (see FISH-quant)
Filt_options.output = 0;
options.Filt = Filt_options
```

25. Use FISH-quant as previously described (1-3)

- Recommendations
 - Set analysis parameters with the original images
 - Create your file with settings (see FISH-quant documentation1,2,3)
 - Use batch mode to run the images (step 23)
 - Select Use filtered images

Integration with FISH-quant

Condensates quantification (Fig.1, see Supplementary Information (4)) is performed after the detection of cytosolic smFISH spots.

To avoid any detection of spots within condensates by FISH-quant, the output images (step 22 and 23) are labeled, so:

- The only possible detection of false positive smFISH sports occurs at condensate edges.
- After detection in FISH-quant (1-3), false positive spots at the condensate edges can be filtered out because they would have a unique and very high "Pixel intensity (Raw)" value that: (1st) it is not present in the image and (2nd) it is biased to the right side as shown in the following example:

Before filtering out false positives from condensates edges

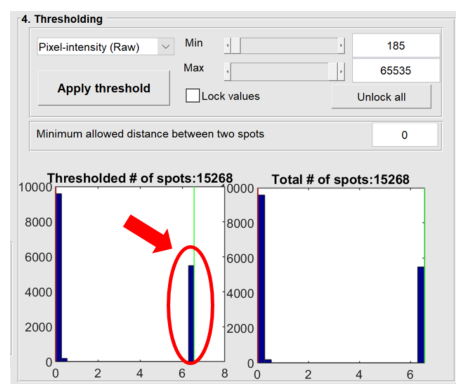


Image from FISH-quant v3 main interface (1-3).

After filtering out false positives from condensates edges

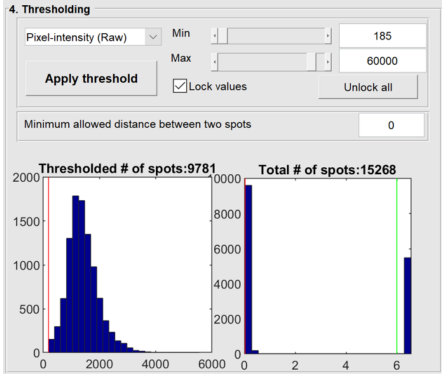


Image from FISH-quant v3 main interface (1-3).

References

- 1. Mueller, F., Senecal, A., Tantale, K. et al. FISH-quant: automatic counting of transcripts in 3D FISH images. Nat Methods 10, 277–278 (2013).
- 2. Tsanov, N., Samacoits, A., Chouaib, R. et al. smiFISH and FISH-quant a flexible single RNA detection approach with super-resolution capability, Nucleic Acids Research 44 (22), e165 (2016).
- 3. https://bitbucket.org/muellerflorian/fish_quant/src/master/
- 4. Cardona et al., Self-demixing of mRNA copies buffers mRNA:mRNA and mRNA:regulator stoichiometries, Cell (2023), https://doi.org/10.1016/j.cell.2023.08.018