# User's guide of DecodeSTORM

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## Contents

1. Requirements	2
2. How to install	2
2.1 How to install ImageJ	2
2.2 How to install DecodeSTORM	2
3. How to use	2
3.1 Change results save path	2
3.2 Format conversion	3
3.3 Channel registration	4
3.4 Artifact correction	4
3.4.1 Drift correction	4
3.4.2 Localization filtering	5
3.5 Quantitative analysis	6
3.5.1 Spatial distribution statistics	6
3.5.2 Segmentation and cluster	7
3.5.3 Colocalization	8
3.6 Visualization	9
4. Note	10
5. Testing of DecodeSTORM with experimental data	10
5.1 Artifact correction	11
5.1.1 Drift correction	11
5.1.2 Localization filtering	11
5.2 Quantitative analysis	12
5.2.1 Raw data	12
5.2.2 Artifact-corrected data	14

1. Requirements

Operation system: DecodeSTORM has been tested on Windows 7 (64-bit) and

Windows 10 (64-bit).

Software: ImageJ.

Note: please set at least 4 GB memory buffer for ImageJ.

2. How to install

2.1 How to install ImageJ

Users can download ImageJ at <a href="https://imagej.en.softonic.com/">https://imagej.en.softonic.com/</a>. After downloading,

double-click "ImageJ-win64.exe" to install.

Of course, users can also choose to download Fiji at <a href="https://fiji.sc/">https://fiji.sc/</a>. After

downloading, unpack it and use it directly. Fiji is an image processing package - a

"batteries-included" distribution of ImageJ, bundling a lot of plugins that facilitate

scientific image analysis.

We recommend that users download the latest version of ImageJ or Fiji.

2.2 How to install DecodeSTORM

DecodeSTORM is built for ImageJ independently. To install, simply copy four dynamic

link libraries (cudart64 110.dll, libopenblas.dll, opencv world450.dll, DecodeSTOR-

M CPPDLL.dll) files into the ImageJ installation folder. And then copy the

corresponding .jar plug-in file to the plugins folder of ImageJ.

3. How to use

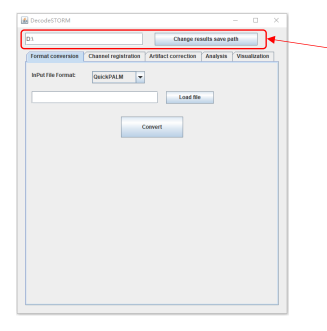
Open the DecodeSTROM: ImageJ → Plugins → DecodeSTORM

3.1 Change results save path

Click the "Change results save path" button to change the path to save the results of

format conversion, artifact correction, quantitative analysis, and visualization. The

default path is D:\.

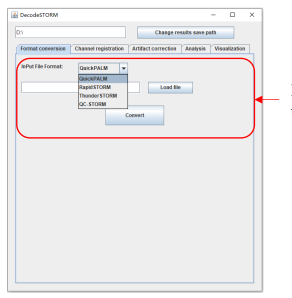


Change the path to save the results of format conversion, artifact correction, and quantitative analysis.

Figure 1: Change the path to save results.

## 3.2 Format conversion

Format conversion module is used to convert the localization table generated by QuickPALM, RapidSTORM, ThunderSTORM, or QC-STORM into the format suitable for DecodeSTORM. First, select the file format that users need to convert, then click the "Load file" button to upload the localization table, and finally click the "Convert" button. Results are saved at the selected path.



Format convert to get the localization table suitable for DecodeSTORM.

Figure 2: Format conversion module.

## 3.3 Channel registration

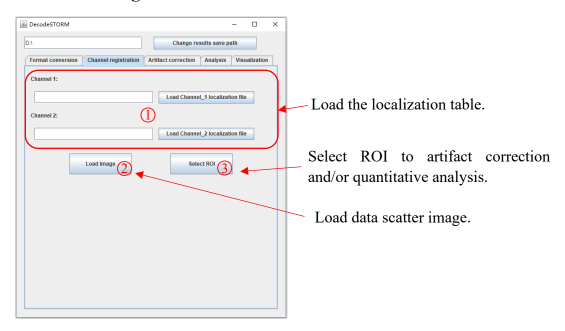


Figure 3: Channel registration module.

Load localization file: DecodeSTORM provides two channels for importing two-color SMLM data: channel 1 for green and channel 2 for red. For single-color SMLM data, users can click the "Load Channel\_1 localization file" or "Load Channel\_2 localization file" button to upload. For two-color SMLM data, users should click the "Load Channel\_1 localization file" and "Load Channel\_2 localization file" buttons to upload. Load image: Clicking the "Load Image" button will generate a scatter image of the data when the localization table is in the channel.

**Select ROI:** Select ROI for artifact correction and quantitative analysis. First, Click the ImageJ "Rectangle" button to select the ROI area. Then, use the keyboard shortcut Ctrl+T to load the ROI into the ROI manager and select. Lastly, click the "Select ROI" button of DecodeSTORM. DecodeSTORM allows the user to select multiple ROI areas to load into the ROI manager, and the user can select one of them for ROI artifact correction and quantitative analysis.

#### 3.4 Artifact correction

#### 3.4.1 Drift correction

DecodeSTORM provides redundant cross-correlation drift correction to correct artifacts caused by sample drift. A single frame contains too few localizations for

reliable correlation calculation, so users need to set the number of drift correction group frames (default: 500). In addition, users need to set the original image pixel size (default: 100), then select one or all channels and click on the "DrifCorr" button for drift correction, the results can be selected to be saved or not. Note that subsequent modules will continue to provide options for channel selection and whether to save the results. We also offer a GPU option for accelerated drift correction (the computer must have a Cuda-enabled GPU).

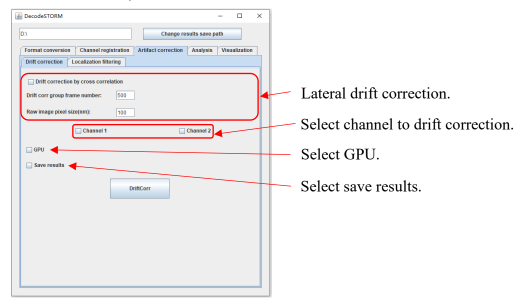


Figure 4: Drift correction in Artifact correction module.

#### 3.4.2 Localization filtering

Imprecise localizations, isolated localizations, duplicated localizations in the same frame, and localizations from molecules reappearing in subsequent frames result in imaging artifacts, which might impact quantitative analysis results, so the above localizations need to be removed or merged.

**Remove imprecise localizations:** This function requires localization tables to have localization precision or to be calculated using other information. To remove imprecise localizations, users need to set an imprecise localization threshold (default: 40).

**Remove isolated localizations:** Remove isolated localization based on local density. The parameters set by users are the radius to determine the size of the area to search for neighbor localization events (default: 30nm) and the minimum number of localization events within the radius (default: 5).

Remove duplicated localizations in the same frame: Duplicated localizations are determined according to the euclidean distance between localizations in the same frame. DecodeSTORM offers a distance threshold (default: 20nm) to remove duplicated localizations.

Merge localizations from molecules reappearing in subsequent frames: Localizations from molecules reappearing in subsequent frames are determined according to the euclidean distance between localizations in the subsequent frame. In DecodeSTORM, users need to set 2 parameters to merge localizations from molecules reappearing in subsequent frames: max frames of localizations reappearing (default: 1) and max distance between localizations (default: 10nm).

After the user selects the method, clicking the "Filtering" button will generate the filtered scatter plot. If users choose to save the results, the results users get using either method will be saved.

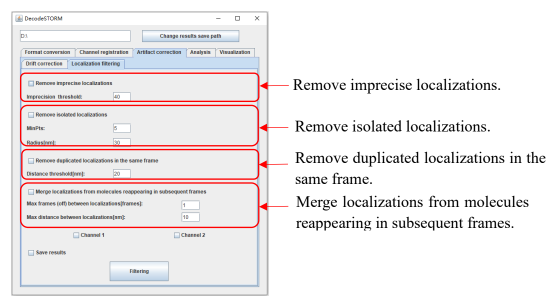


Figure 5: Localization filtering in Artifact correction module.

## 3.5 Quantitative analysis

#### 3.5.1 Spatial distribution statistics

In the spatial distribution statistics, the max analysis distance (default: 55nm) and ring width (default: 2nm) need to be set. Selecting the same channel means calculating the self-clustering of the channel data, and selecting different channels means the co-clustering of the two channels' data. Radial Distribution Function (G(r)) and Ripley's

H Function (H(r)) can be used separately or together. Clicking the "Statistics" button will generate their function graphs. If users choose to save the result, not only the function graph is saved, but also the function value and its corresponding radius r are saved in a CSV file. At the same time, the user can choose to link to the artifact correction module to use the corrected data or to use the raw data. Note, the subsequent modules in the "Analysis" also provide "Link artifact correction".

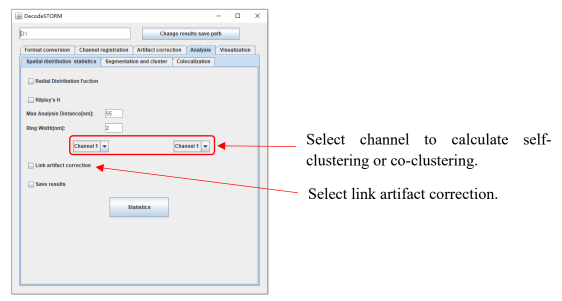


Figure 6: Spatial distribution statistics in the Quantitative analysis module.

#### 3.5.2 Segmentation and cluster

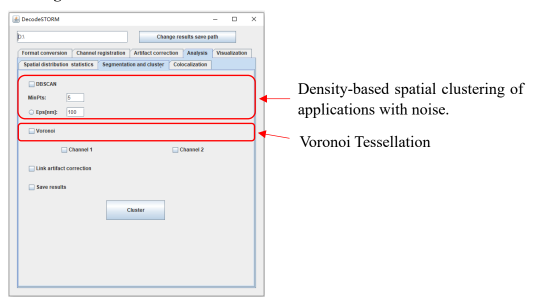


Figure 7: Segmentation and cluster in Quantitative analysis module.

DBSCAN has two parameters: the radius *eps* (default 100nm) that determines the size of the search nearest neighbor localization event region and the minimum number of

localization events *MinPts* (default 5) in the above region, which generates the molecular segmentation and cluster maps, meanwhile the number and density of clusters are obtained. In DecodeSTORM, *eps* can also be calculated automatically. Voronoi can generate a Voronoi diagram as well as obtain the number and density of clusters by cluster localization without setting any parameters.

DBSCAN and Voronoi can be used separately or together. Clicking the "Cluster" button will generate a DBSCAN clustering map and/or Voronoi diagram, a text box is appeared to show the number and density of clusters. If users choose to save the results, the DBSCAN clustering map and/or Voronoi diagram will be saved, in addition, the specific clustering information of DBSCAN will be saved in a CSV file.

#### 3.5.3 Colocalization

In DecodeSTORM, colocalization analysis is performed with three parameters to be set: maximum observation distance  $R_{\rm max}$ , search step Step and CBC threshold. The CBC threshold allows the user to autonomously segment colocalization and uncolocalization localizations. Clicking the "Colocalization" button will generate histograms of CH1  $\rightarrow$  CH2 and CH2  $\rightarrow$  CH1 colocalization values. If users choose to save the result, the above histogram and a CSV file are saved, the CSV file contains the percentage of the number of localizations whose colocalization value of CH1  $\rightarrow$  CH2 is greater than the threshold to the number of localization value of CH2  $\rightarrow$  CH1 is greater than the threshold to the number of localizations in CH2.

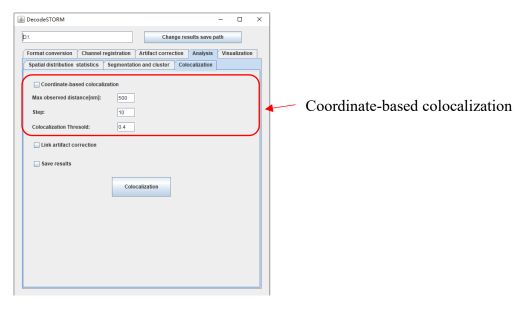


Figure 8: Colocalization in Quantitative analysis module.

## 3.6 Visualization

In addition to the generation of the scattered map, DecodeSTORM provides Gaussian rendering for SMLM data visualization, and users need to set the rendering pixel sizes, and raw image pixel size, and then click the "Rendering" button will generate a rendering image. We also provide an ROI option where users can render the selected ROI data.

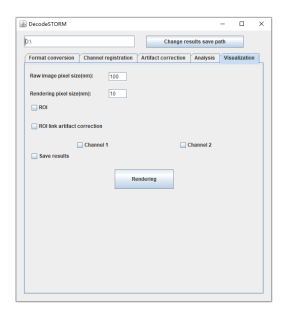


Figure 9: Visualization module.

## 4. Note

The default parameters in DecodeSTORM are set based on our test data. "Save results" is selected when using the "Filtering", "SpatialStatistics", "Segmentation and Cluster", "Colocalization" and "Rendering" module, the corresponding folders (Name: "FilteringResults", "SpatialStatisticsResults", "SegmentationAndClusterResults", "RenderingResults", "ColocalizationResults") will be generated for saving these results. These folders are located at the selected save path, drift correction and format convert results are located at the same path. In addition, the numbers in the CSV file generated by DecodeSTORM are presented in scientific notation.

## 5. Testing of DecodeSTORM with experimental data

We tested the DecodeSTORM using a part of the two-color data that Pageon et al. provided in their Github repository (<a href="https://github.com/PRNicovich/ClusDoC">https://github.com/PRNicovich/ClusDoC</a>) during the development of Clus-DOC. The two-color SMLM data were imaged from TCR (green) and pTCR (red), respectively, in activated Jurkat cells. The ROI we selected was also saved, and after importing the image, the saved ROI is loaded for testing.

Load localization file  $\rightarrow$  Load Image  $\rightarrow$  Click on the top right corner to enlarge the button  $\rightarrow$  "Ctrl + T"  $\rightarrow$  ROI Manager  $\rightarrow$  More  $\rightarrow$  Open  $\rightarrow$  Open 0430-0613.roi  $\rightarrow$  Click on the selected  $\rightarrow$  Select ROI. The specific operation results are shown in the following figures.

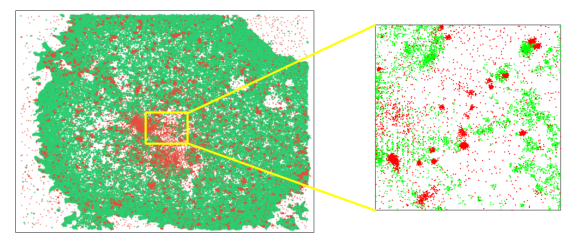


Figure 10: Select a noise-heavy part of the data provided by Pageon et al. for testing the software.

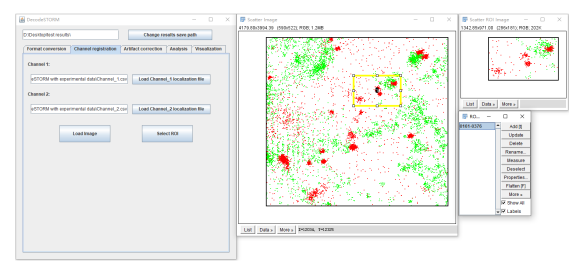


Figure 11: Upload data and select ROI.

## 5.1 Artifact correction

#### 5.1.1 Drift correction

This data lacks the information needed for the redundant inter-correlation drift correction method, so we did not perform a drift correction.

## 5.1.2 Localization filtering

We have filtered the data within channel 1 and channel 2 using all the methods in "Localization filtering".

Imprecision threshold: 40.

MinPts: 5.

Radius[nm]: 30.

Distance threshold[nm]: 20.

Max frames (off) between localizations[frames]: 1.

Max distance between localizations[nm]: 10.

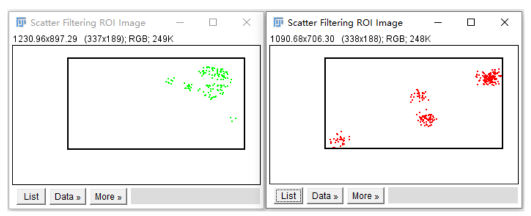


Figure 12: Scatter map of filtered data within channel 1 and channel 2.

## 5.2 Quantitative analysis

Since the data of channel 1 is irregularly distributed, we only performed spatial distribution statistics as well as segmentation and clustering on the data of channel 2. So the options in "spatial distribution statistics" and in "segmentation and clustering" should be selected "channel 2".

#### 5.2.1 Raw data

## (1) Spatial distribution statistics: G(r) and H(r)

Max analysis distance[nm]: 55.

Ring width[nm]: 2.

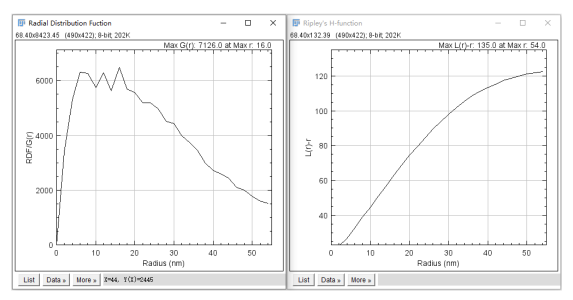


Figure 13: G(r) and H(r) of raw channel 2 data.

## (2) Segmentation and cluster: DBSCAN and Voronoi

MinPts: 5.

Eps: automatic calculation.

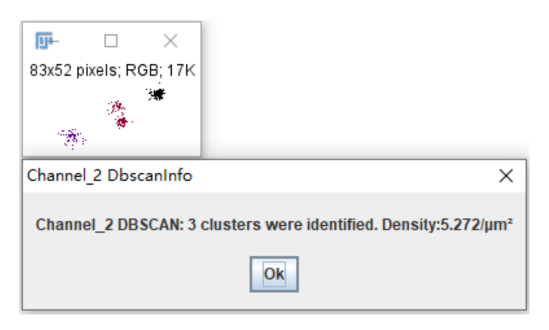


Figure 14: DBSCAN clustering maps of raw channel 1 and channel 2 data.

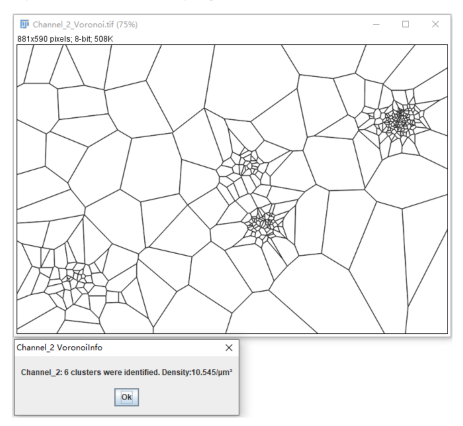


Figure 15: Voronoi diagram of raw channel 1 and channel 2 data.

## (3) Colocalization

Max observed distance[nm]: 500.

Step: 10.

Colocalization Thresold: 0.4.

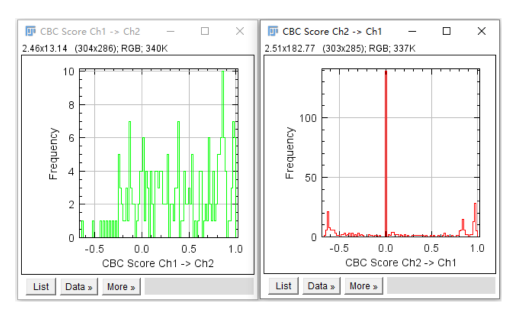


Figure 16: The histograms of CH1 → CH2 and CH2 → CH1 colocalization value without artifact correction.

#### 5.2.2 Artifact-corrected data

## (1) Spatial distribution statistics: G(r) and H(r)

Max analysis distance[nm]: 40.

Ring width[nm]: 2.

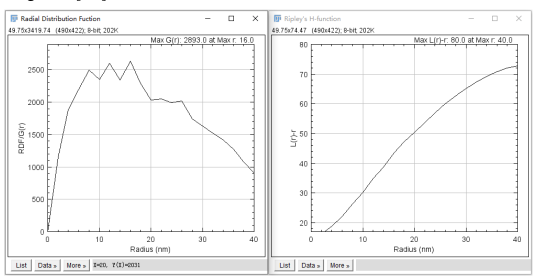


Figure 17: G(r) and H(r) of artifact-corrected channel 2 data.

## (2) Segmentation and cluster: DBSCAN and Voronoi

MinPts: 5.

Eps: automatic calculation.

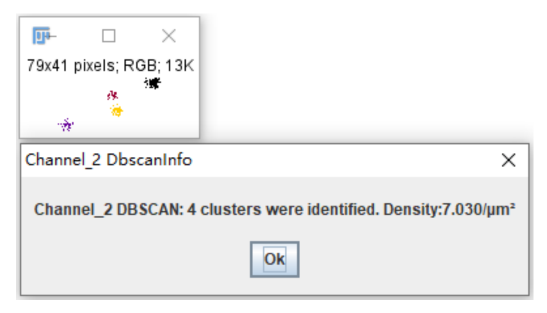


Figure 18: DBSCAN clustering maps of artifact-corrected channel 1 and channel 2 data.

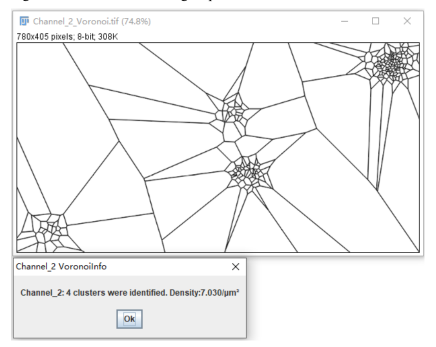


Figure 19: Voronoi diagram of artifact-corrected channel 1 and channel 2 data.

## (3) Colocalization

Max observed distance[nm]: 500.

Step: 10.

Colocalization Thresold: 0.4.

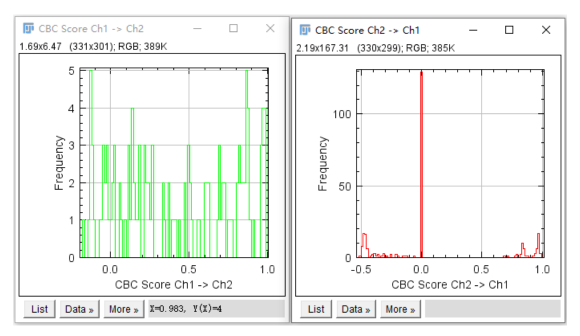


Figure 20: The histograms of CH1  $\rightarrow$  CH2 and CH2  $\rightarrow$  CH1 colocalization value with artifact correction.