# CiliaGUI - Installation and User Guide

#### Installation

This video describes the installation of the CiliaDetect App.

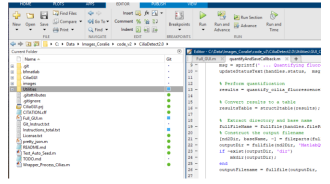
To run the toolbox, you need to:

- 1. Install MATLAB (2021 or later).
  - MATLAB is not free; check with your institution for access.
- If you cannot get a license, contact me we may try a standalone version of the program.
- Installation instructions: https://www.mathworks.com/help/install/ug/install-products-with-internet-connection.html
- During installation, make sure to install the following toolboxes:
- Statistics and Machine Learning Toolbox
- Image Processing Toolbox
- 2. Download the code from GitHub:
  - Repository: https://github.com/pmgarderes/CiliaDetect2.0
  - Click 'Download as ZIP file'.
- 3. Unzip and place the folder on your hard drive.
- Example: on Windows, I use WinRAR and put the files into a folder named ciliaGUI\_demo at the root of C:/.

### **Launching a Session**

- 4. Navigate to the folder and find Full\_GUI.m.
- Double-click it. If MATLAB is not open, this will automatically launch MATLAB and set the current folder correctly.

 $\Box$  Make sure the current folder in MATLAB (shown in the top bar) contains Full\_GUI.m. If it doesn't, adjust it manually.  $\rightarrow$  it should like like this (roughly)



- 5. Start a session:
  - Go to the Editor panel.
  - With Full\_GUI.m active (double click Full\_GUI in matlab to make sure it's active
  - click Run.

#### **Preprocessing – Z-Planes**

- 6. Prepare your microscopy images before using the GUI.
- The program only reads \*\_reduced.mat , i.e. files formatted for matlab. Thus we need to pre-process microscopy image and format them.
- Strategy: reduce the number of frames in a to a few averaged Z-planes ( down to a single Z-plane), so multiple Z-planes can be separated from the same Z-stack.

#### To preprocess:

- 1. Click the 'Pre-process' button.
- 2. Enter the downsampling factor (number of frames to average together).
  - Example: enter '1000' to collapse the full stack into a single Z-plane.
- 3. A file browser window will open. Navigate to the folder containing your microscopy files.
- $\hfill \square$  You will not see the individual files just select the directory where the files are located.
  - 4. The GUI text box will display messages ('WAIT') and begin downsampling.

Notes:

- Preprocessing large stacks may take a long time. Tip: you can batch-process many files overnight.
- Output: a new folder 'Matlab\_quantif' is created. Each original file generates a reduced version with the suffix '\_reduced.mat'.

#### **Loading a File**

- 7. Load files into the GUI:
- Only files ending in '\_reduced.mat' can be loaded.
- Click 'Select File', navigate to the directory, and choose the file.

### **GUI Display**

- 8. The central panel displays one Z-plane in one channel.
  - Adjust fluorescence intensity with numeric keypad: /, \*, -, +.
  - These settings apply across all Z-planes in the same channel but not across channels.
  - Navigate between channels: left/right arrow keys.
  - Navigate between Z-planes: up/down arrow keys.

#### **Detecting ROIs**

- 9. To add a new ROI:
  - Left-click directly on the cilium (not beside it).
  - Press Space.
  - The program segments the cilium around your click.
  - If detection fails (e.g., low contrast), it will appear as a single point.
  - Delete failed detections by pressing U once (undo).
- $\square$  Do not leave empty ROIs.

### **Parameters to Adjust Detection**

- 10. Sensitivity is the key parameter.
  - Recommended range: 0.3–0.7 (use 0.4–0.5 in most cases).

- Higher sensitivity: easier detection, less accurate ROIs.
- Lower sensitivity: stricter detection, fewer but more accurate ROIs.

#### **Merging ROIs**

- 11. If a cilium is split across Z-planes or incompletely detected:
- Click near the junction point.
- Press M to merge ROIs (works if they are close enough).

#### **Deleting ROIs**

- 12. To delete ROIs:
  - Single ROI: click near it and press S.
  - All ROIs: click 'Clear Detections'.

#### **Other Useful Buttons**

13. Refresh ROI display: press R.

## **Saving Detections**

- 14. Save your work:
  - Click 'Save Detections' at any time.
  - This creates a new file in 'Matlab\_quantif' inside 'reduced\_stack/'.
  - File name format: originalName\_reduced\_cilia\_detections.mat
  - Next time you open the reduced file, saved detections load automatically.
- □ Only load '\_reduced.mat' files. Do not load detection files directly.

#### **Visualizing Masks**

- 15. To view ROI and background masks:
- Click 'Visualize Masks'.
- Two images will appear:
- Left: cilia mask overlay.

- Right: background mask.
- Images are saved automatically.
- Adjustable parameters:
- Padding: gap between ROI and background (pixels).
- Background mask width (pixels).

#### **Quantify and Save Fluorescence**

16. To export quantified data:

- Click 'Quantify and Save Fluorescence'.
- Creates one Excel file (.xls) per image in 'Matlab\_quantif/'.
- Each row = one cilium.

Fluorescence outputs (per channel):

- F\_StackSum\_ch# sum across the Z-stack.
- F\_plan\_ch# mean in the Z-plane.

Morphological measures (if metadata available):

- Length (μm)
- Width (µm)
- Curviness (length/chord length)
- Length/Width ratio
- Area (µm²)

#### Notes:

- Both methods correct for background.
- Both are valid for comparisons if exposure and Z-range are similar.
- Recommendation:
- Use stack sum if Z-range > cilium span.
- Use mean method otherwise.

# **Editing Parameters**

- 17. Parameters are user-editable:
- Any parameter can be modified.
- Settings are automatically saved and reloaded in the next session.