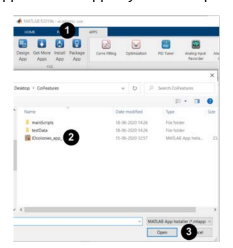
### **ColFeatures**

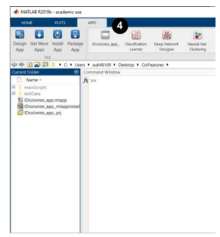
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### **GUI** interface

#### Installation

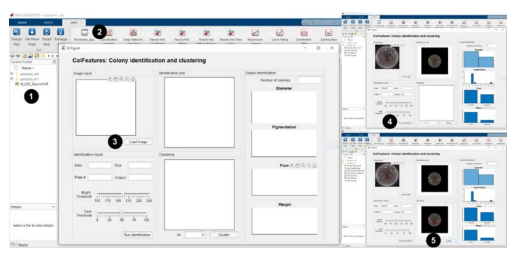
You need to have MATLAB installed in your computer. We were using 2019 version. For installation, after downloading the different files go to Apps and click on 'Install App' (1), browse the ColFeatures folder and select the MATLAB App Installation file (2), then open the file (3). After this the GUI interface will appear as an app in your APPS panel (4).





# Usage

Once installed, it is ready to be used. The first step is to go to the folder (do not forget to add the folder to the path) where you have your data, remember the GUI only works with single .tif files (1). Once in your folder, click on the App: *IDcolonies\_app\_* (2), the App will open right away. On the App, click on **Load image** and select the image to analyze (3), provide also de inputs for the fields: *Data, Plate # and File Output*. After clicking on *Run identification* (4) the App will show your image output and some of the metrics. You can either stop here or also try the *Cluster* option, after providing the number of clusters to partition your data and clicking in *Cluster* (5) the App will show how the colonies were clustered. In each step output files will be generated and stored in your main folder.

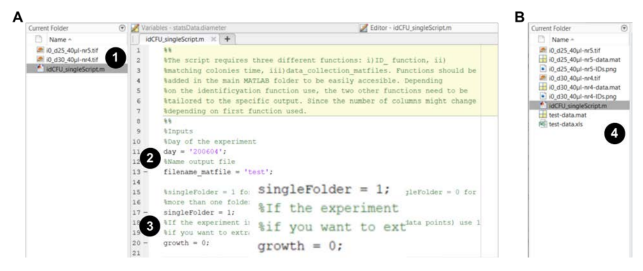


# **Batch analysis**

Download the MATLAB scripts IDcfu\_Final, dataCollection\_Final, dataCollectionGrowth\_Final and matchColonies\_Final, save them in your MATLAB folder. Also download the idCFU\_singleScript and depending on the analysis you will need to save in in specific locations.

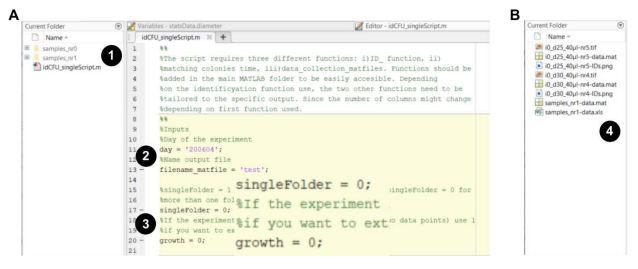
## One folder analysis

In this case, you are only interested in analyzing a set of images in a single folder. You need to save the idCFU\_singleScript in the same folder as your images (1). Provide the required inputs (2) and set the decision flags as follows: **single\_Folder = 1 and growth = 0 (3)**. After this you only need to run the script and three different types of files will be generated (4).



# Multiple folders analysis

There is also an option to analyze multiple folders at once. For this case, you need to save the  $idCFU\_singleScript$  in the core folder, which subfolders are the ones to be processed (1). It is important that the subfolders are numbered, such as nr0, nr1 and so for. The script requires the pattern "nr" to go into each subfolder. Provide the required inputs (2) and set the decision flags as follows:  $single\_Folder = 0$  and growth = 0 (3). After this you only need to run the script and three different types of files will be generated in each folder (4).



## Multiple folders analysis and growth rate calculation

When analyzing multiple folders, we can also obtain apparent growth rate based on two pictures taken at different time points. One has to be careful to have a way to make sure that the colony centroids will not change much, in our case, we found it useful to draw a line in one side of the plate and also mark a line on the tray to take pictures. For this case, you need to save the idCFU\_singleScript in the core folder, which subfolders are the ones to be processed (1). It is important that the subfolders are numbered, such as **nr0**, **nr1** and so for. The script requires the pattern "**nr**" to go into each subfolder. Provide the required inputs (2) and set the decision flags as follows: **single\_Folder = 0 and growth = 0 (3)**. After this you only need to run the script and three different types of files will be generated in each folder. This option will also generate a final data frame where all the data points from each folder will be saved (4).

