# find\_hostspots manual

### Background

In some application such as single-molecule FISH we obtain a spatial information about gene expression levels. The goal of this script is to identify a pattern of expression in 2D space, explore the reproducibility between biological replicates, and compare between few genes tested in the same experiment.

#### Data

Data was obtained from: Citri, Ami (2021), "smFISH data of IEG expression in the dorsal striatum after acute, repeated, and challenge cocaine exposures", Mendeley Data, V3, doi: 10.17632/p5tsv2wpmg.3

The data is smFISH results from striatum sections before and after 1h stimulation with cocaine. Three gene were tested: Egr2, Arc and Nr4a1. The dataset includes: number of replica, gene expression levels in each cells, x and y coordinates of the cell.

```
##
     rep_num Arc Egr2 Nr4a1 center_x
## 1
                          15 3019.079
           1
                8
                     4
                                        45.86756
## 2
           1
                1
                            0 2631.921
                                        62.09142
## 3
           1
              14
                     8
                          10 3773.377 101.17299
## 4
           1
               4
                     1
                            1 4018.893 93.77170
## 5
            1
              27
                     1
                            7 3709.124 104.29448
## 6
                            0 2538.018 89.10385
```

### Analysis guidelines

- 1. Find the high expressors cells based on the control samples.
- 2. Apply 2D kernel density estimation with contours on the highly expressors cells data
- 3. Select the highest contours that include at least 20% of the cells.
- 4. Apply the analysis separately for each replica. Overlap the results from all the replicas to see if we the selected contours are overlapping.
- 5. Calculate the average expression levels in the selected contours to gain information about the expression level of the genes at the hotspot.
- 6. Perform the same analysis with shuffled data. The expression levels of the original data are shuffled, while the x and y coordinates remained the same. This allows as to evaluate the specificity of your results.

## Example input

To use this script one must load files in the format described above. Two files are needed: one for the control (0h) and one from the treatment (1h). In addition, the list of genes to be analyzed should be included.

```
#setwd("/path/to/find_hotspot/dir")

#for control
path_data_for_bin = "input_example/_merged_0h_Arc_Nr4a1_.csv"

#for treatment
path_data_for_analysis = "input_example/_merged_1h_Arc_Nr4a1_.csv"

list_genes = c("Arc","Nr4a1","Egr2")

output_file_name = "output_example/hotspot_res.pdf"

shuffle_data = TRUE

shuffle_file_name = "output_example/hotspot_shuffled.pdf"
```