MATLAB package: Cell-Cycle-Chromatin-Dynamic-Organization, CCCDO, user guide

General:

Live 3D measurement of telomere diffusion offers an effective method to investigate Chromatin dynamics and reveal cell-cycle-dependent levels of motion-constraint, caused by various cellular processes.

CCCDO is a MATLAB package using ensemble MSD analysis in order to quantify cell-cycle dependant chromatin dynamics during the interphase.

Part 0: installation of Code package folder

unzip "CCCDO_startup .m" and the folder CCCDO_mater to your MATLAB folder path, usually: "C:\ ... \ MATLAB "

Part 1: Modify the start-up file

- 1. Open the file "CCCDO_startup .m"
- 2. For the first installation only change: "InstallRun = 1"

this action will create the folder tree required for analysis, as described in the part 2. Return: "InstallRun = 0;" immediately after installation.

3. Define directory were the data should be stored. In first installation the parent folder will be created in this folder.

4. Define the MATLAB folder path,

5. Define parent folder name. {"ParentFolderName = 'ExampleFolder';"} this folder will be created in the target folder user defined earlier.

```
%% %%%%%%% make Data storage folder if InstallRun is true
%%%%%%%%%%%%%%
if InstallRun==1
    % cell cycle phase names
    PhaseNames = {'GO','S','G2','LateG2','UT'};
    % cell lines names
    CellTypeName = {'MEF3T3','MEFLmnaKO'};
    % parent folder name
```

ParentFolderName = 'ExampleFolder';
Part 2: User need to arrange the parent folder content, before starting running any analysis, as follow:

Name	Туре	Name	Туре
Level 1			
MEF3T3_G0	Experiment folder	MEFLmnaKO_G0	Experiment folder
MEF3T3_S	Experiment folder	MEFLmnaKO_S	Experiment folder
MEF3T3_G2	Experiment folder	MEFLmnaKO_G2	Experiment folder
MEF3T3_LateG2	Experiment folder	MEFLmnaKO_LateG2	Experiment folder
MEF3T3_UT	Experiment folder	MEFLmnaKO_UT	Experiment folder
EnsembleAnalysis	Analysis folder		
Level 2: content of an Experiment folder			
Analysis	Level 3 folder		
Level 4 folders : "Cell_*", single cell data, includes csv files resulted from trajectory reconstruction, example file name for cell 6: "MEF3T3_S_Cell_6_*.csv"			
localizations.csv	File in Cell	Localization results: output of the DeepStorm3D net	
homographies.csv	File in Cell	Affinity matrixes, for each frame, which compensate for the lateral nucleus drift	
_localizations_warped.csv	File in Cell_	Compensated localization -after Affinity matrixes been applied on the raw localization	
trajectories.csv	File in Cell	Reconstructed trajectories of the compensated localization	
msds.csv	File in Cell	Mean square displacement vector (3D)	
Level 2: content of the folder "EnsembleAnalysis"			
Axial	Ensemble analysis based on the axial (Z) information		
Lateral	Ensemble analysis based on the lateral (XY) information		
Full	Ensemble analysis based on 3D information		
SummaryAnalysis	Constraining volume analysis		

Part 3: User parameters definition file

Open the file "UserParameter.m"

Part A: define general parameters for all experiment

- 1. Define the length of imaging (in frames)
- 2. Define the threshold for minimal length of a trajectory
- 3. Define the parent folder where all data and analysis results are saved. This folder should be arranged precisely as "Example folder" attached to this package

Part B: define the privet parameters for each experiment

Task indexes: 1,2

This part has 10 section: 2 cell lines ('MEF 3T3','MEF LmnaKO') and 5 phases ('G0','S','G2','LateG2','UT').

For each section user need to define:

- 1. Excluded cells ("ExcludedCells"): cells to be excluded from analysis. If none define =[];
- 2. Number of cells ("numCells"): total number of cells, including those need to be excluded.
- 3. Trajectories to be excluded ("ExcludeTrj"): after running QC analysis user may exclude manually specific trajectories due to not accurate linear fit. If none, as during the first run, leave the struct empty("ExcludeTrj=struct();")

Part C: define folders for ensemble MSD analysis. **User do not need to edit this section!**

Task index: 3

Part D: define time for constraining volume analysis

Task index: 4

User need to specify time in sec for calculate the constraining volume ("TimeAve = 25;")

Part 4: analysis pipeline

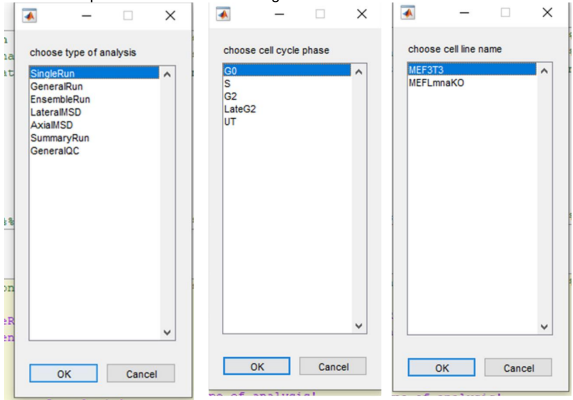
Step0: save all CSV files with trajectories information as described in part 2.

Step 1: Run mode: 'SingleRun'

Rearrange and save each cell data, for specific "experiment folder" (see part 2).

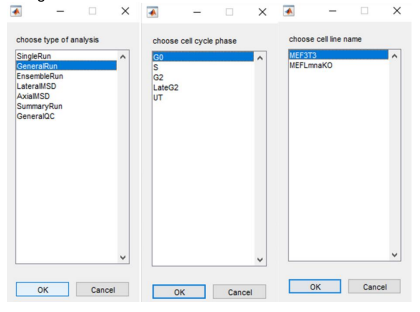
Run file: "CCCDO_startup.m" then run function: "Main.m"

Answer the questions of the user dialog boxes:



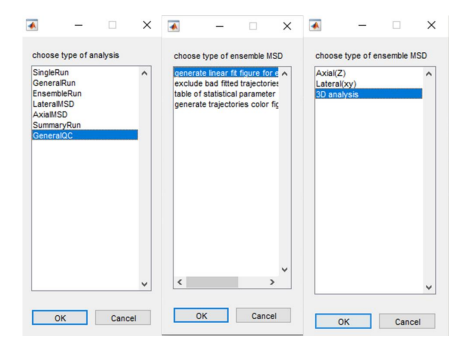
Step 2: RunMode: 'GeneralRun'

This function summarizes all trajectories data from single experiment. Run file: "CCCDO_startup.m" then run function: "Main.m". Answer the questions of the user dialog boxes:



Step 2QC1: RunMode: 'GeneralQC'

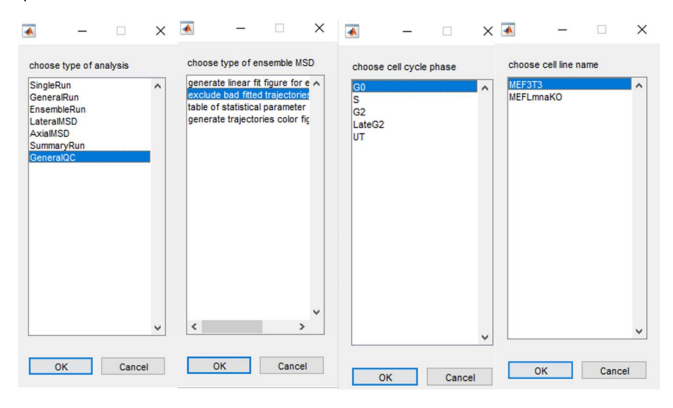
At this point user may run QC analysis which generate linear fit figures for all the trajectories in single experiment, in order to manually detect bad fitted trajectories and exclude them from analysis. Run file: "CCCDO_startup.m" then run function: "Main.m". Answer the questions of the user dialog boxes:



Step 2QC2: RunMode: 'GeneralQC'

After running Step 2QC1, user should update the file: "UserParameter.m" and specify the trajectories to be excluded as follow:

Then run Main.m to remove these trajectories from analysis; answer the dialog boxes questions:

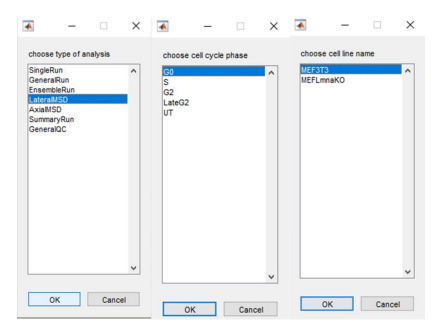


Step 2QC3: RunMode: 'GeneralRun'

When QC procedures are done. Run again general analysis.

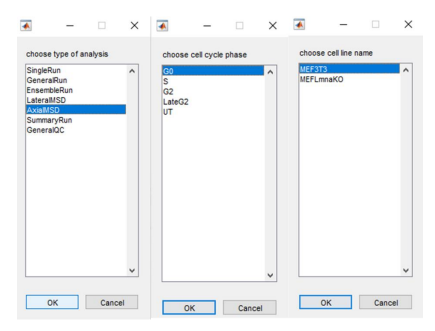
Step 3a: RunMode: 'LateralMSD'

This function summarizes all trajectories data from single experiment, using only XY coordinates. Run file: "CCCDO_startup.m" then run function: "Main.m". Answer the questions of the user dialog boxes:



Step 3b: RunMode: 'AxialMSD'

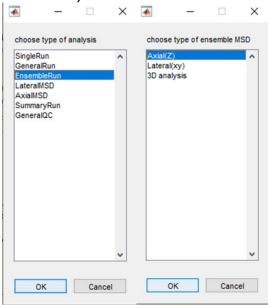
This function summarizes all trajectories data from single experiment, using only Z coordinates. Run file: "CCCDO_startup.m" then run function: "Main.m". Answer the questions of the user dialog boxes:



Step 4: RunMode: 'EnsembleRun'

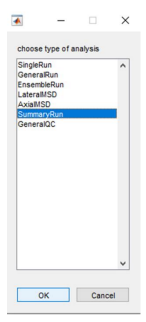
calculates anomalous exponent and diffusion coefficient.

compare between cell cycle phase and between cell lines (MEF 3T3 and MEFLmnaKO) user shall run this section 3 times for each type of ensemble MSD analysis (3D, axial and lateral).



Step 5: RunMode: 'SummaryRun'

This function calculates the "constraining volume"- the mean volume "visited" by a random telomere in each phase of the cell cycle. This value represents the constraint level of the specific cell cycle phase.



Step 6: RunMode: 'GeneralQC'

Final step is statistical tests for all analyses. This run will calculate the results significance using two-sample t-test, under the assumption that different variance is allowed. In addition, a table similar to supplementary table S2 in our article will be printed in MATLAB command line.

