**Introduction**

**Multi-Geneplex** is a pipeline for the identification of communities on a gene multiplex composed of four layers: co-expression network, transcriptional and post-transcriptional co-targeting network and protein-protein interaction network.

The pipeline performs the following operartions:

1. Multiplex reconstruction

2. Layers filtering

3. Community detection on the multiplex

**The alpha thresholds for step 2. has to be chosen from the user. We suggest to folow the three criteria proposed in the paper.**

**The community detection algorithm for step 3. has to be chosen from the user, the suggested one is OSLOM.**

**Installation**

The following software components are required to run **Multi-Geneplex:**

* **Multi-Geneplex**
* R (<http://www.r-project.org/>)
* R packages: igraph and BioPhysConnectoR
* package Ganet [REF] (<https://sites.google.com/site/gokmenaltay/ganet>). To install the package download the file Ganet\_1.0.tar.gz at the above link. Set the working directory in the same folder that ganet is placed. Then, in the R console, write the following: install.packages(“GAnet\_1.0.tar.gz”, type=”source”).
* Python
* Python modules: networkx and scipy

In principle, **Multi-Geneplex** should run  under Linux and Mac.

**Usage**

To run **Multi-Geneplex**:

* Set the working directory in Multi-Geneplex
* ./ Multi-Geneplex\_run.sh

**Inputs**

While running, the program requires the following inputs:

* The path to the file containing the mRNA expression matrix in log2 with the first column containing genes names, the second column containing gene annotations (or gene names) and the first row containing sample names.
* **I**nteger corrisponding to the optimal alpha value, choosing among (0=0.005,1=0.01,2=0.02,3=0.03,4=0.05,5=0.1,6=0.2,7=0.3,8=0.4,9=0.5) for the espression network, transcription factor co-taregting network and microRNA co-taregting network. To choose the optimal alpha value you can use the three criteria suggested in the paper taking advantage of the files:
  + Multi-Geneplex/temp/selection\_links\_EXP.txt for the co-expression layer.
  + Multi-Geneplex/temp/selection\_links\_TF.txt for the transcription factor co-taregting layer.
  + Multi-Geneplex/temp/selection\_links\_MIRNA.txt for the microRNA co-taregting layer.
* The integer corrisponding to the algorithm you want to use for network clustering, choosing among:
  + 0: oslom undirected,
  + 2: infomap undirected
  + 4: louvain
  + 5: label propagation method,
  + 8: modularity optimization

**Outputs**

During **Multi-Geneplex** running some intermidiate outputs are created in Multi-Geneplex/temp/ the folder will be delated when thre execution is completed.

The final output comm\_final\_genesymbol.txt is in the folder Multi-Geneplex. The file reports in the first column an integer used as identifier for the communities. Then in each row the list of genes belonging to that community is reported. Communities are ordered in respect to thei dimension.

**Data**

The datasets to reproduce the results of our paper are available at:

* gastric xxx
* lung xxx,
* pancreas xxx
* colon xxx

**Contact**

Please feel free to contact us at

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Moreover, feel free to change the code according to your needs.

For every use of the original or modified pipeline cite us: