Manual for TimePath

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Requirements and General instructions

- The software is available at www.sb.cs.cmu.edu/timepath
- The github is located at https://github.com/tmfs10/timepath
- You must have Java 7.0 or above and Python 2.7+ installed on your machine.
- You must have a GML file visualizer like Cytoscape to view the graph visualization

Data needed

1. A protein protein interaction network in the format

```
<protein> <interaction type> <protein> <probability>
:
```

The columns have to be separated by the **tab** character. The <interaction type> can be one of pp (protein-protein), ptm (post-translational modification), and pd (protein-dna). The latter two are directed whereas pp is undirected. A sample protein-protein interaction network with gene names under the HGNC symbol naming scheme is provided in the file ppi_ptm_pd_hgnc.txt.

- 2. A list of transcription factors (TFs). A sample list containing 348 TFs is provided.
- 3. A list of TF-gene interactions in the format

```
TF Gene Prior
<TF> <Gene> <Prior>
```

Again, the columns have to be tab separated. A sample TF-gene interaction file is provided in the file human_encode_100.txt.

4. A list of targets for a particular phase is in the format

```
<target 1 for phase 1> <target 1 for phase 2>...
<target 2 for phase 1> <target 2 for phase 2>...
```

and so on. The targets for each time point should be separated by a tab character. The targets should be ranked by how likely they are to be a target for that time point with the most likely target first. One can do this using p-values generated by a software like Deseq (for RNA-seq data). Another popular method is to rank the genes by most differentially expressed to least.

Note: The number of targets for each time point must be the *same*. If they are not, please add in the dummy target XXXX as needed to make them the same.

5. The time series file should be in the format <gene name> <log-fold change for timepoint 1> <log-fold change for timepoint 2> <gene name> <log-fold change for timepoint 1> <log-fold change for timepoint 2> It should also have a header row at the top. The log-fold change can be with respect to a 0 time point or with respect to a control condition depending how the exact needs.

Config file

Mandatory parameters

- 1. numTPGenes The number of targets to select for each time point
- 2. TPtargetsFile The path to the file containing targets for each time point as in point 4 in the "Data needed" section.
- 3. timeseriesFilepath The path to the file containing the time series gene expression data
- 4. ppiNetworkFilepath The path to the file containing the protein-protein interaction network
- 5. sourcesFilepath The path to the file containing the source proteins
- 6. tfGeneFile The path to the file containing the TF-gene interactions
- 7. pathFile Path to the file to which the pathways should be written to

Optional parametrs

- 1. maxNumPaths Maximum number of candidate pathways to extract per source and target gene pair. Default value is 100,000.
- 2. maxPathLength Maximum number of proteins per pathway. Default value is 10.
- 3. phases If you are merging time points together (we HIGHLY recommend merging down to 4-5 time points
- 4. rnaHitsFilepath Path to the file containing RNA hits if you have one
- 5. numProteinsPerPhaseToSelect Number of proteins to rank per phase
- 6. minRankingFoldChange Only proteins that show a big change in rank from one time point to another are selected (as we want to select the proteins that
- 7. numGenesToDisplay This is the number of intermediate/TF genes to display per time point in the graph visualization. Default is 15.
- 8. graphNodeWidth Width in pixels of the protein nodes in the graph in pixels. Default is 60.
- 9. graphNodeHeight Height in pixels of the protein nodes in the graph in pixels. Default is 40.
- 10. 11 This is the penalty for including too many genes in the network. Increase to reduce number of genes selected. Default is 1
- 11. 12 this is the penalty for including too few targets in the network. Increase to increase number of targets explained. Default is 0.3
- 12. L This is the size of the tabu list. Default is 200
- 13. N This is the total number of iterations. Default is 1000

Running the algorithm

The command to run the algorithm is as follows:-

```
java -jar -Xmx8g timepath.jar <config file>
```

Here <config file> is the is the configuration file which is explained in the previous section

Compute protein rankings

1. The command to compute the overall protein rankings is

```
python computeOverallProteinRanking.py <path file>
```

2. The command to compute time point specific protein ranking

```
python computeTopProteinsForTP.py <config file>
```

This will create a file <path file>.phasegenes.txt where <path file> is as entered in the config file

Visualization

You must compute the time point specific protein rankings (using computeTopProteinsForTP) before running the graph visualization. After doing that, run the following command

```
python createGraph.py <config file>
```

This will create a file named <path file>.graph.gml where <path file> is as entered in the config file. You can then load the gml file into a visualization software like Cytoscape to view the resulting network.