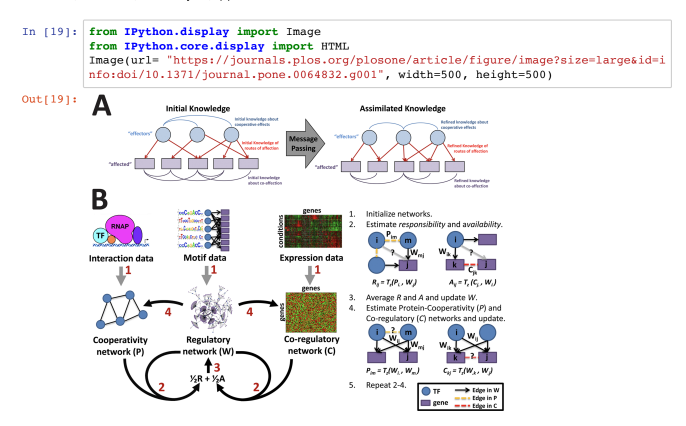
Up and running with PANDA and netZooPy

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Regulatory network reconstruction is a fundamental problem in computational biology. There are significant limitations to such reconstruction using individual datasets, and increasingly people attempt to construct networks using multiple, independent datasets obtained from complementary sources, but methods for this integration are lacking. We developed PANDA (Passing Attributes between Networks for Data Assimilation), a message-passing model using multiple sources of information to predict regulatory relationships, and used it to integrate protein-protein interaction, gene expression, and sequence motif data to reconstruct genome-wide, condition-specific regulatory networks in yeast as a model. The resulting networks were not only more accurate than those produced using individual data sets and other existing methods, but they also captured information regarding specific biological mechanisms and pathways that were missed using other methodologies. PANDA is scalable to higher eukaryotes, applicable to specific tissue or cell type data and conceptually generalizable to include a variety of regulatory, interaction, expression, and other genome-scale data.

Glass K, Huttenhower C, Quackenbush J, Yuan GC. Passing Messages Between Biological Networks to Refine Predicted Interactions, PLoS One, 2013 May 31;8(5):e64832



Installation and Setup

```
In [20]: %%bash
         cd ~
         # git clone https://github.com/netZoo/netZooPy.git
         cd netZooPy
         pip3 install -e .
         Obtaining file:///Users/redmo/netZooPy
         Requirement already satisfied: pandas in /usr/local/lib/python3.7/site-packages
         (from netZooPy==0.1.1) (0.25.2)
         Requirement already satisfied: numpy in /usr/local/lib/python3.7/site-packages
         (from netZooPy==0.1.1) (1.17.0)
         Requirement already satisfied: networkx in /usr/local/lib/python3.7/site-package
         s (from netZooPy==0.1.1) (2.4)
         Requirement already satisfied: matplotlib in /usr/local/Cellar/graph-tool/2.29 1
         /libexec/lib/python3.7/site-packages (from netZooPy==0.1.1) (3.1.1)
         Requirement already satisfied: scipy in /usr/local/lib/python3.7/site-packages
         (from netZooPy==0.1.1) (1.3.0)
         Requirement already satisfied: python-dateutil>=2.6.1 in /usr/local/lib/python3.
         7/site-packages (from pandas->netZooPy==0.1.1) (2.8.0)
         Requirement already satisfied: pytz>=2017.2 in /usr/local/Cellar/graph-tool/2.29
         1/libexec/lib/python3.7/site-packages (from pandas->netZooPy==0.1.1) (2019.1)
         Requirement already satisfied: decorator>=4.3.0 in /usr/local/lib/python3.7/site
         -packages (from networkx->netZooPy==0.1.1) (4.4.0)
         Requirement already satisfied: cycler>=0.10 in /usr/local/Cellar/graph-tool/2.29
         1/libexec/lib/python3.7/site-packages (from matplotlib->netZooPy==0.1.1) (0.10.
         0)
         Requirement already satisfied: kiwisolver>=1.0.1 in /usr/local/Cellar/graph-tool
         /2.29 1/libexec/lib/python3.7/site-packages (from matplotlib->netZooPy==0.1.1)
         (1.1.0)
         Requirement already satisfied: pyparsing!=2.0.4,!=2.1.2,!=2.1.6,>=2.0.1 in /usr/
         local/Cellar/graph-tool/2.29_1/libexec/lib/python3.7/site-packages (from matplot
         lib->netZooPy==0.1.1) (2.4.0)
         Requirement already satisfied: six>=1.5 in /usr/local/lib/python3.7/site-package
         s (from python-dateutil>=2.6.1->pandas->netZooPy==0.1.1) (1.12.0)
         Requirement already satisfied: setuptools in /usr/local/lib/python3.7/site-packa
         qes (from kiwisolver>=1.0.1->matplotlib->netZooPy==0.1.1) (41.2.0)
         Installing collected packages: netZooPy
           Found existing installation: netZooPy 0.1.1
             Uninstalling netZooPy-0.1.1:
               Successfully uninstalled netZooPy-0.1.1
           Running setup.py develop for netZooPy
         Successfully installed netZooPy
```

or with conda

```
In [21]: # source activate myenv
# conda install git pip
# pip install git+git://github.com/netZoo/netZooPy.git

In [22]: import os
# os.getcwd()
# os.chdir("~/")
os.getcwd()

Out[22]: '/Users/redmo'

In [23]: from netZooPy.panda.panda import Panda
import pandas as pd
import matplotlib.pyplot as plt
```

Parameter Setting & Exploring the Data

First, we start by setting the path to the 1) motif prior network, 2) the gene expression data, and 3) the ppi network data. The motif prior network is typically a TF-by-gene binary matrix where 1 indicates the presence of sequence (motif) of a TF in the gene regulatory region and 0 otherwise. Gene expression data is typically a gene-by-sample matrix containing expression data. PPI network is a TF-by-TF binary matrix, where 1 indicates a physical interaction between two TFs and 0 otherwise. If two TFs are likely to binding, they are likely to form regulatory complexes for the same genes.

```
In [24]: expression_data='netZooPy/tests/ToyData/ToyExpressionData.txt'
    motif_data='netZooPy/tests/ToyData/ToyMotifData.txt'
    ppi_data='netZooPy/tests/ToyData/ToyPPIData.txt'
    panda_output='netZooPy/tests/panda/output_panda.txt'
```

There are 1000 genes and 51 samples in our toy data. This is your novel input. The remaining files are known interaction lists.

```
In [25]: motif_data=pd.read_csv(motif_data,sep="\t",header=None)
    motif_data[0].unique().size

Out[25]: 87

In [26]: motif_data[1].unique().size
Out[26]: 913
```

Since the first column is TF, you thus have 87 TF and 913 genes are returned from the second column, with their interaction weights in the third column (motif_data[2]). Now lets check out the ppi data, another interaction list with three columns, with 238 interactions between the TF.

Calling PANDA

One can chose to run in terminal simply by pointing to the input files

```
In [28]: # %%bash
# cd netZooPy
# pip3 install -e .
# python netZooPy/panda/run_panda.py -e netZooPy/tests/ToyData/ToyExpressionData.t
xt -m netZooPy/tests/ToyData/ToyMotifData.txt -p netZooPy/tests/ToyData/ToyPPIDat
a.txt -f True -o test_panda.txt
```

Alternatively one can continue running in Jupyter, using all data sources:

```
In [40]: expression data='netZooPy/tests/ToyData/ToyExpressionData.txt'
         motif_data='netZooPy/tests/ToyData/ToyMotifData.txt'
         ppi_data='netZooPy/tests/ToyData/ToyPPIData.txt'
         panda_obj = Panda(expression_data, motif_data, ppi_data, save_tmp=True, save_memory
         = False, remove_missing=False, keep_expression_matrix = False)
         panda_obj.save_panda_results(panda_output)
         Loading motif data ...
         Unique TFs: 87
           Elapsed time: 0.01 sec.
         Loading expression data ...
         Expression matrix: (1000, 50)
           Elapsed time: 0.02 sec.
         Loading PPI data ...
         Number of PPIs: 238
           Elapsed time: 0.00 sec.
         Calculating coexpression network ...
           Elapsed time: 0.01 sec.
         Creating motif network ...
           Elapsed time: 0.01 sec.
         Creating PPI network ...
           Elapsed time: 0.00 sec.
         Normalizing networks ...
           Elapsed time: 0.03 sec.
         Saving expression matrix and normalized networks ...
           Elapsed time: 0.00 sec.
         Running PANDA algorithm ...
         /Users/redmo/opt/anaconda3/lib/python3.7/site-packages/scipy/stats/stats.py:231
         5: RuntimeWarning: divide by zero encountered in true divide
           return (a - mns) / sstd
         Saving PANDA network to netZooPy/tests/panda/output_panda.txt ...
           Elapsed time: 0.22 sec.
In [30]: import sys
         sys.getsizeof(panda_obj)
```

Out[30]: 64

In [31]: panda_obj.top_network_plot(top=10)

/Users/redmo/opt/anaconda3/lib/python3.7/site-packages/networkx/drawing/nx_pylab.py:579: MatplotlibDeprecationWarning:

The iterable function was deprecated in Matplotlib 3.1 and will be removed in 3.

3. Use np.iterable instead.

if not cb.iterable(width):

/Users/redmo/opt/anaconda3/lib/python3.7/site-packages/networkx/drawing/nx_pylab.py:585: MatplotlibDeprecationWarning:

The iterable function was deprecated in Matplotlib 3.1 and will be removed in 3.

3. Use np.iterable instead.

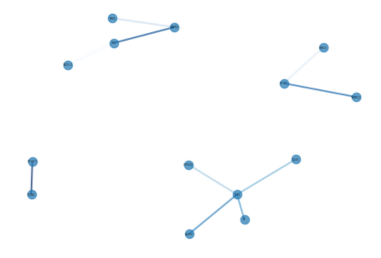
and cb.iterable(edge_color) \

/Users/redmo/opt/anaconda3/lib/python3.7/site-packages/networkx/drawing/nx_pylab.py:595: MatplotlibDeprecationWarning:

The iterable function was deprecated in Matplotlib 3.1 and will be removed in 3.

3. Use np.iterable instead.

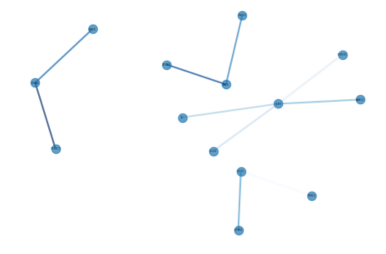
for c in edge_color]):



using only the motif prior

```
In [32]: expression_data=None
         motif_data='netZooPy/tests/ToyData/ToyMotifData.txt'
         ppi_data=None
         panda_obj = Panda(expression_data, motif_data, ppi_data,remove_missing=True)
         panda_obj.save_panda_results(panda_output)
         Loading motif data ...
         Unique TFs: 87
           Elapsed time: 0.02 sec.
         No Expression data given: correlation matrix will be an identity matrix of size
         No PPI data given: ppi matrix will be an identity matrix of size 87
         Remove motif not in expression data:
            0 rows removed from the initial 14597
         Calculating coexpression network ...
           Elapsed time: 0.00 sec.
         Creating motif network ...
           Elapsed time: 0.01 sec.
         Normalizing networks ...
           Elapsed time: 0.03 sec.
         Saving expression matrix and normalized networks \dots
           Elapsed time: 0.00 sec.
         Running PANDA algorithm ...
         Saving PANDA network to netZooPy/tests/panda/output panda.txt ...
           Elapsed time: 0.22 sec.
```

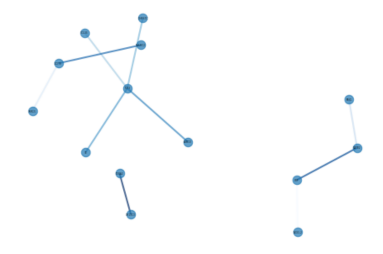
In [33]: panda obj.top network plot(top=10)



without the expression matrix

```
In [34]: expression data=None
         motif_data='netZooPy/tests/ToyData/ToyMotifData.txt'
         ppi_data='netZooPy/tests/ToyData/ToyPPIData.txt'
         panda_obj = Panda(expression_data, motif_data, ppi_data,remove_missing=True)
         panda_obj.save_panda_results(panda_output)
         Loading motif data ...
         Unique TFs: 87
           Elapsed time: 0.02 sec.
         No Expression data given: correlation matrix will be an identity matrix of size
         913
         Loading PPI data ...
         Number of PPIs: 238
           Elapsed time: 0.00 sec.
         Remove motif not in expression data:
            0 rows removed from the initial 14597
         Remove ppi not in motif:
            0 rows removed from the initial 238
         Calculating coexpression network ...
           Elapsed time: 0.00 sec.
         Creating motif network ...
           Elapsed time: 0.01 sec.
         Creating PPI network ...
           Elapsed time: 0.00 sec.
         Normalizing networks ...
           Elapsed time: 0.03 sec.
         Saving expression matrix and normalized networks ...
           Elapsed time: 0.00 sec.
         Running PANDA algorithm ...
         Saving PANDA network to netZooPy/tests/panda/output_panda.txt ...
           Elapsed time: 0.21 sec.
```

In [35]: panda_obj.top_network_plot(top=10)



and without using a motif prior

```
In [36]: expression data='netZooPy/tests/ToyData/ToyExpressionData.txt'
         motif_data=None
         ppi_data='netZooPy/tests/ToyData/ToyPPIData.txt'
         panda_obj = Panda(expression_data, motif_data, ppi_data)
         panda_obj.save_panda_results(panda_output)
         Loading expression data ...
         Expression matrix: (1000, 50)
           Elapsed time: 0.02 sec.
         Loading PPI data ...
         Number of PPIs: 238
           Elapsed time: 0.00 sec.
         Calculating coexpression network ...
           Elapsed time: 0.01 sec.
         Returning the correlation matrix of expression data in <Panda_obj>.correlation m
         atrix
         Saving PANDA network to netZooPy/tests/panda/output panda.txt ...
           Elapsed time: 1.99 sec.
```

and saving memory by deleting intermediary files

```
In [38]: | expression_data='netZooPy/tests/ToyData/ToyExpressionData.txt'
         motif data='netZooPy/tests/ToyData/ToyMotifData.txt'
         ppi data='netZooPy/tests/ToyData/ToyPPIData.txt'
         panda obj = Panda(expression_data, motif_data, ppi_data, save_memory=True)
         panda obj.save panda results(panda output)
         Loading motif data ...
         Unique TFs: 87
           Elapsed time: 0.02 sec.
         Loading expression data ...
         Expression matrix: (1000, 50)
           Elapsed time: 0.01 sec.
         Loading PPI data ...
         Number of PPIs: 238
           Elapsed time: 0.00 sec.
         Calculating coexpression network ...
           Elapsed time: 0.01 sec.
         Creating motif network ...
           Elapsed time: 0.01 sec.
         Creating PPI network ...
           Elapsed time: 0.00 sec.
         Normalizing networks ...
           Elapsed time: 0.03 sec.
         Clearing motif and ppi data, unique tfs, and gene names for speed
         Saving expression matrix and normalized networks ...
           Elapsed time: 0.01 sec.
         Running PANDA algorithm ...
         Saving PANDA network to netZooPy/tests/panda/output panda.txt ...
           Elapsed time: 0.08 sec.
```

Basic follow up analysis is also possible, such as degree calculation per gene

```
In [46]: panda_obj.return_panda_indegree()
Out[46]:
                        force
              gene
              41157
                     1.139713
             AACSL
                     -5.090725
              AAK1 -10.923515
           ABCA17P
                     -8.919393
             ABCB8
                     9.091765
                 ...
             ZNF826
                     -7.009150
             ZNF845
                     -7.942670
             ZNF878 -11.919082
            ZSWIM3
                    38.185909
            ZWILCH
                    -1.784825
          1000 rows × 1 columns
In [51]:
          panda_obj.save_panda_results()
          Saving PANDA network to panda.npy ...
             Elapsed time: 0.14 sec.
```