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The **GeneralGeneletModel.py** package (imported as GGM in accompanying scripts) includes a GeneletNetwork class that for initializing, simulating, and compiling sequences for genelet networks. The general genelet model functions at the beginning of **GeneralGeneletModel.py** are helper functions for the GeneletNetwork class. Below are the descriptions for the initialization of GeneletNetwork and how to use its associated functions.

1. Class initialization definition:

CLASS: GeneletNetwork(act_vec,prod_vec,blk_vec,indc_vec=0)

Initializes the genelet network instance to be simulated

Input definitions:

For act vec and blk vec:

List the length of ind nodes (total nodes)

Numbers represent which orthogonal activators/blockers correspond to which nodes

O indicates no activator/blocker for a given node

For prod vec:

List the length of ind nodes (total nodes)

Numbers represent which RNA repressors and/or RNA coactivators are produced by which nodes

-ve values are repressors
+ve values are coactivators
0 = no production of an inducer RNA

For indc_vec:

List the length of ind nodes (total nodes)

Numbers represent which inducer RNAs are produced by which nodes

-ve values are inducers that bind repressors (thereby activating)

+ve values are inducers that bind coactivators (thereby BLKing)

0 = no production of an inducer RNA

indc vec is optional is default to zeros if not provided

Output definitions:

After initializing the GeneletNetwork class the user has access to the following Attributes [where: model = GeneletNetwork()]

model.ortho_nodes
model.ind_nodes
model.ind_nodes
model.act_vec
model.prod_vec

model.prod_vec

model.blk_vec
model.indc_vec
model.indc_vec

model.indc_vec

model.topology_mat

model.I topology mat

- number of orthogonal genelet nodes
- number of total (individual) genelet nodes
- activator vector mapping ind_nodes to orthogonal activators
- RNA production vector mapping ind_nodes to which RNAs they produce (repressors/coactivators)
- blocker vector mapping ind_nodes to orthogonal blockers
- RNA inducer production vector mapping ind_nodes to which inducers they produce
- topology matrix (an alternative representation of act_vec/prod_vec notation)
- inducer topology matrix (an alternative representation of

	<pre>act_vec/indc_vec notation)</pre>
model.act_mat	 activator connection matrix used in ODEs to map activators to ind_nodes
model.rep_mat	 repressor connection matrix used in ODEs to map repressors to ind_nodes
model.blk_mat	 blocker connection matrix used in ODEs to map blockers to ind_nodes
model.ca_mat	 coactivator connection matrix used in ODEs to map coactivators to ind_nodes
model.Rprod_mat	 repressor production matrix used in ODEs to map which ind_nodes produce which repressors
model.Cprod_mat	 coactivator production matrix used in ODEs to map which ind nodes produce which coactivators
model.Rindc_mat	- repressor inducer production matrix used in ODEs to map which ind nodes produce which repressor inducers
model.Cindc_mat	 coactivator inducer production matrix used in ODEs to map which ind_nodes produce which coactivator inducers

2. Function: initial conditions() definition:

Input definitions:

For dA_tot and G_tot:

numpy arrays the length of ortho_nodes and ind_nodes, respectively

Represent total concentrations (nM) of activators and genelets, respectively

For G int vec:

List the length of ind_nodes (total nodes)

Represents the initial state of a node

- 1 for ON
- -1 for BLK
- 0 for OFF

For dB_added:

All BLK genelets are assumed to be annealed with 50% excess dB Thus for 25 nM of a BLK there will be 32.5 nM total dB and 12.5 nM free dB to as the initial concentration

To specify additional free dB modify the dB_added input (in nM)

EX: model.initial_conditions(dA_tot,G_tot,G_int_vec,dB_added=[0,150,150])

For all other species the default values are 0:

To modify these species input a list of concentrations for any other species by name (in nM):

EX: model.initial_conditions(dA_tot,G_tot,G_int_vec,rCin =[0,150,150])
EX: model.initial_conditions(dA_tot,G_tot,G_int_vec,rRin=[1000,0,0])

Output definitions:

After calling initial_conditions the user will have access to the following attributes:

3. Function: simulate() definition:

Simulates the network with the above initial conditions

Simulate can be called numerous times and specific conditions can be updated by name \mbox{Use} 'NA' for values that should not be updated for a given species

For updating a species conditions, the list has to be the length of the total number of that species and input in $\ensuremath{\text{nM}}$

 \mbox{dR} is a special option where DNA repressors can be added to the reaction to permanently remove DNA activators

dR species are currently not included in the symbolic equations that can be output as they are not really part of the networks

 $t_{\rm vec2}$ should start at the last timepoint from $t_{\rm vec1}$ and go to a later timepoint

Input definitions:

For rnase:

```
Selects the rnase to use in the simulations
'RnH' = RNase H (degrade RNA in RNA:DNA duplex)
```

'RnA' = RNase A (degrade ssRNA and RNA in RNA:DNA duplex (likely with slower rate than RNase H))

'Both' = RNase H and A

This input is optional and 'RnH' is the default

For leak:

Represents the leak transcription rate of BLK genelets as a fraction of the ON transcription rate between 0 and 1 $\,$

Can be entered as a single value applied to all BLK genelets or as a list for different leak rates for each RNA production:

leak = [0.05, 0.1, 0.075, 0.06]

There should be ind_node # of leak inputs as in this model each individual genelet has its own production rate even if two genelets produce the same RNA

For rate constants:

Use different rate constants than the default values

Import as a list of single values (1/M-s for $2^{\rm nd}$ order reactions and 1/s for $1^{\rm st}$ order reactions):

[kpr,kpc,kpi,kd_H,kd_A,kga,kgar,kar,kgb,kgab,kgbc,kbc,kir]

```
Each individual rate can be a list of length ind_nodes for production rates
             Or a list of length ortho nodes for other rates
             Or a single value which will be assumed to be the same for all nodes
EX: model.simulate(t_vec1,1)
EX: model.simulate(t_vec2,2,rIr=['NA','NA',10000])
EX: model.simulate(t_vec3,3,rR=[1000,'NA','NA'])
EX: model.simulate(t_vec1,1,rnase='RnA') # use RNase A
EX: model.simulate(t vec1,1,rnase='both',leak=0.1) # use both RNase A and H
                                                   # 10% leak transcription
EX: model.simulate(t_vec1,1,rnase='both',leak=0.1,rate_constants=rate_list)
                                                 # use both RNase A and H
                                                 # 10% leak transcription for all RNAs
                                                 # user input rate constants
EX: model.simulate(t vec1,1,leak=[0.05,0.1,0.075]) # different leak rates for 3 RNAs
Output definitions:
After running simulate the user has access to the following useful attributes:
             -sol.y contains all of the concentration date and sol.t contains the time
model.sol
             in seconds
model.(rate constants) -all of the individual rate constants can be accessed by name
                              -the dictionary containing all of the concentrations
model.output concentration
                               with specific name keys
EX: G1 = model.output_concentration['GdA1']
EX: A1 = model.output concentration['A1']
EX: R4 = model.output concentration['rR4']
4. Function: export sym eqs() definition:
export_sym_eqs(file_name='sym_eqs',path_name='')
Exports the symbolic ODE equations of the model to a text file
Both of the inputs are optional and have default values
Input definitions:
For file name:
      User defined name of the .txt file (do not include .txt in the name)
For path_name:
      User defined path to where the file should be saved (include \ at the end)
      If user does not supply a path the file is saved in current folder
EX: model.export sym eqs(file name='TSN eqs',path name='C:\\Desktop\\')
```

Output definitions:

The output from this is the text file with the symbolic equations in it with the specified or default filename in the specified or current (default) folder

5. Function: plot topology() definition:

This considers both genelet nodes and RNA coactivators/repressors as topological nodes which allows inducer RNA connections to be represented

If this is called right after initializing the network the initial states will be inferred

If this is called after the initial_conditions function has been called the user defined initial states will be displayed

Input definitions:

For pos:

This is an optional input of each nodes position in the plot

It has to be a dictionary with keys that are the names of the topological nodes and the desired [x,y] position values

For layout:

This is an optional input that selects the algorithm to organize the nodes in the plot

```
Default is 'spring'
Other options:
    'spectral'
    'circular'
    'shell'
```

See the networkX package layout options for further details

For plot title:

This is an optional title for the topology plot

For show rnas:

This is necessary if you want to plot topologies where there are nodes that produce inducers

If this is set to 0 then the RNAs will not be plotted and any inducer nodes will appear free standing $\,$

EX: model.plot_topology(layout='shell',plot_title='IFFL network')

Output definitions:

The output from this is the topology plot

model.net edges gives the user access to all the network edges used in the plot

6. Function: compile network sequences() definition:

compile_network_sequences(input_file_loc,desired_nodes=[],min_design=0,bth=1,\
save_file_name='genelet_seqs',save_path_name='')

Exports genelet sequences for a specified topology to a text file

Algorithm for domain selection for unspecified nodes:

First, nodes that must both be coactivated AND repressed are populated with domains that meet this criteria selected in order of their OCRW score in Supp. Section 12

Second, nodes that do not have ANY coactivator or repressor inputs are Populated

Since these nodes are the likely place a network will be expanded they are:

First populated with domains that can be coactivated AND repressed selected in order of their OCRW score in Supp. Section 12

Second populated with domains that can be coactivated selected in order of their OCRW score in Supp. Section $12\,$

Third populated with domains that can be coactivated selected in order of their OCRW score in Supp. Section 12

Third, nodes that have ONLY coactivator inputs are populated

If min_design = 0 the first available domain that can be coactivated will be selected in order of their OCW score in Supp. Section 12

Fourth, nodes that have ONLY coactivator inputs are populated

If min_design = 0 the first available domain that can be repressed will

be selected in order of their OCW score in Supp. Section 12

Input definitions:

For input_file_loc:

The file path to the Excel file holding all of the viable genelet node Sequences (all_genelet_sequences.xlsx)

For desired_nodes:

An optional input that specifies which input domains should be used for each ${\tt TN}$ in the network

By default, this is empty, algorithm will select its own nodes if not provided

Provide as a list of length ortho_nodes with the names of nodes to be used to compile the sequences (['G1','G5','G8'])

TN1 TN2 TN3

The order of the list of nodes is in the order of the topological nodes as

shown above

An error will occur if you select a domain that cannot conduct the regulation required at that position

i.e. if the TN needs to be coactivated and repressed and you select a domain that only represses $\,$

If only some nodes are specified and others are to be selected, leave the unspecified positions blank: (['G1',' ','G8']) $${\tt TN1}$$ TN2 TN3

The unspecified positions will be filled with other domains using the same algorithm as when no nodes are specified

If there is an unspecified position and min_design = 1 then the unspecified node will attempt to use a min design domain if possible

For min design:

An optional input to specified if sequences selected should adopt a minimum design

By default this is 0 so the first nodes that meet the needs of the network will be used as described for the algorithm

If $\min_{\text{design}} = 1$ the algorithm will select nodes with the minimal possible function in the network

For example, if a node is only coactivated in the network then the algorithm will use the first node that can only be coactivated at this position

If there are not enough minimal nodes to fill the network other nodes will be selected as described for the algorithm

For bth:

Optional input that specifies whether or not BTH domains should be included

This is 1 by default which keeps the 8 base 5' blocker toehold (BTH) on all the genelets in the network

This can be set to 0 and then the 8 base 5' blocker toehold will be removed from all genelets that are only repressed in a given network as these nodes do not need a blocker

This makes it possible to order shorter -nt sequences without having to truncate their 3' end (as for the TSN)

All such -nt sequences will be denoted with a * in the output file

For save file name:

User defined name of the .txt file to be saved (do not include .txt in the name) $\ \ \,$

If user does not supply a name it will be saved as 'genelet seqs.txt'

For save path name:

User defined path to where the file should be saved (include \ at the end)

If user does not supply a path the file is saved in current folder

EX.

model.compile_network_sequences(input_file_loc, save_file_name='TSN_sequences', save_pat
h_name='C:\\Desktop\\')

Output definitions:

The output from this is the text file with the genelet sequences in it

Each node is comprised of G-nt / G-t / activator / blocker sequences as needed for the simulated network

All -nt sequences that have had the BTH removed will be denoted with a * in the output file

Currently, any dummy reporter nodes (redundant nodes that do not produce any RNAs) will not be included as exported sequences

7. TROUBLESHOOTING

When doing iterative model.simulate calls make sure the t_vec units are correct, if $t_vec1 = linspace(0,5,1000)*3600$ then $t_vec2 = linspace(t_vec1[-1]/3600,10,1000)*3600$

The package currently does not populate sequences for any redundant dummy nodes in the system.