Motifs, Modules and Pathways

Proteomes Interactomes and Biological Networks

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http://biofold.org/

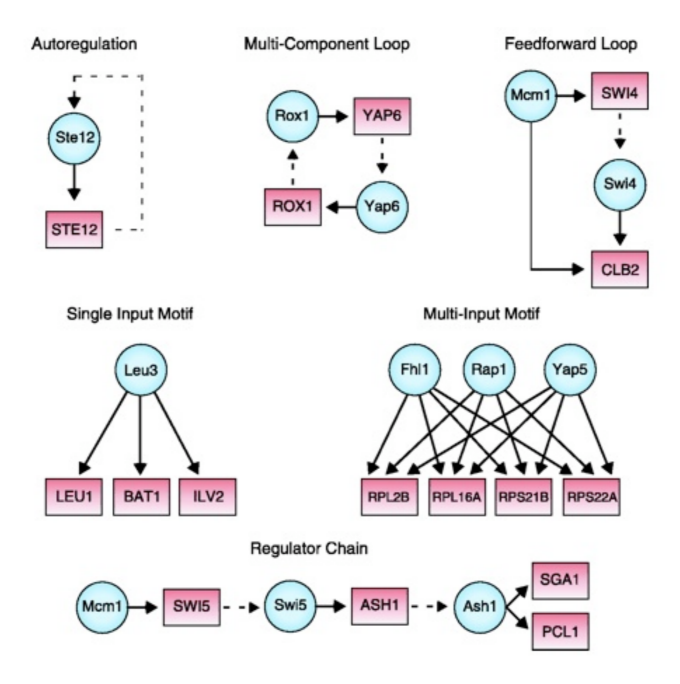


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Network Motifs

Network analysis is important for detecting network motifs, which are recurrent and statistically significant sub-graphs or patterns.



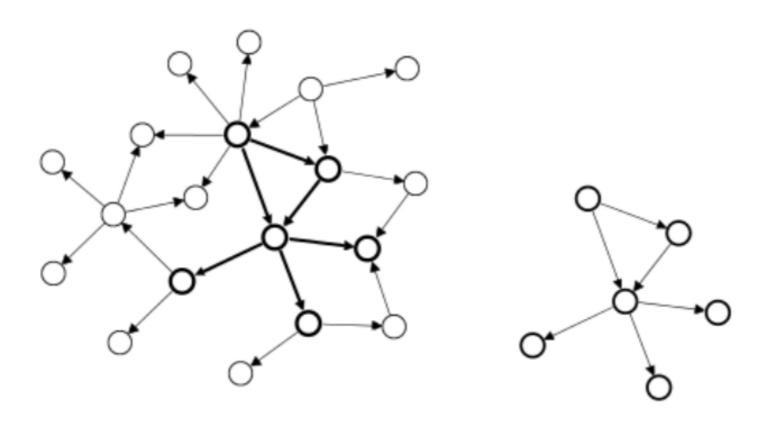
http://compbio.pbworks.com/

Motif Matching

A match of a motif G' in the target graph G = (V, E) is a subgraph G'' = (V'', E'') which is isomorphic to motif G'

Two graphs G' and G" are isomorphic if there is a bijective mapping between the edge and vertex identities

i.e. G' is transformed to G" by changing the vertex and edge identities



Problem Complexity

The complexity of graph isomorphism is in the 'grey area' of complexity:

- It belongs to NP class of problems (problems where solution is easy to verify once found)
- if the correspondence is known, the graph isomorphism belongs to P class of problems (problems that can be solved efficiently)
- f the correspondence is not known, the graph isomorphism is NP-complete (problems that are believed to be hard to solve but easy to verify)
- Subgraph isomorphism, checking if a subgraph G" that is isomorphic to given graph G' exists in a larger graph G, is known to be NPcomplete
- No hope for really fast algorithms for finding motifs.

Statistical Significance

A motif is a statistically overrepresented pattern of local interactions in the network

 Overrepresentation = occurring more frequently than expected by chance

 The motif has emerged several times therefore it has been conserved in the evolution of the network

 The rationale is that overrepresentation may denote possible conservation of the function

Significance tests

The statistical significance can be tested calculating the z-score of the presence of the motif on a set of randomly generated graphs obtained with

Generation of random networks with the Erdos-Renyi algorithm

Random shuffling of the edges

Detection of Motifs

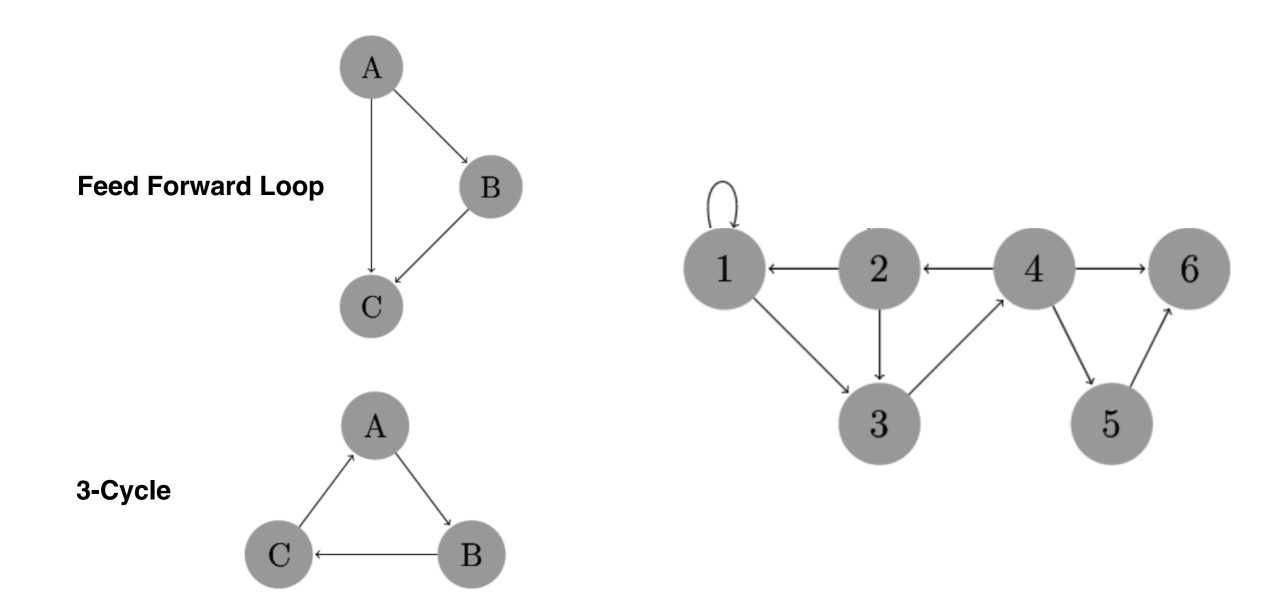
Networkx allow to select a subgraph of a the whole graph and verify if two graphs are isomorphic

```
>>> g = nx.Graph()
>>> g.add_edges_from([(1,2),(1,3)])
>>> mot = nx.Graph()
>>> mot.add_edges_from([("A","B")])
>>> g1 = g.subgraph([1,2])
>>> nx.is_isomorphic(g1,mot)
```

Exercise

Given the Feed Forward Loop (FFL) and 3-Cycle write the code to detect the motif in the graph with 6 nodes and 8 edges.

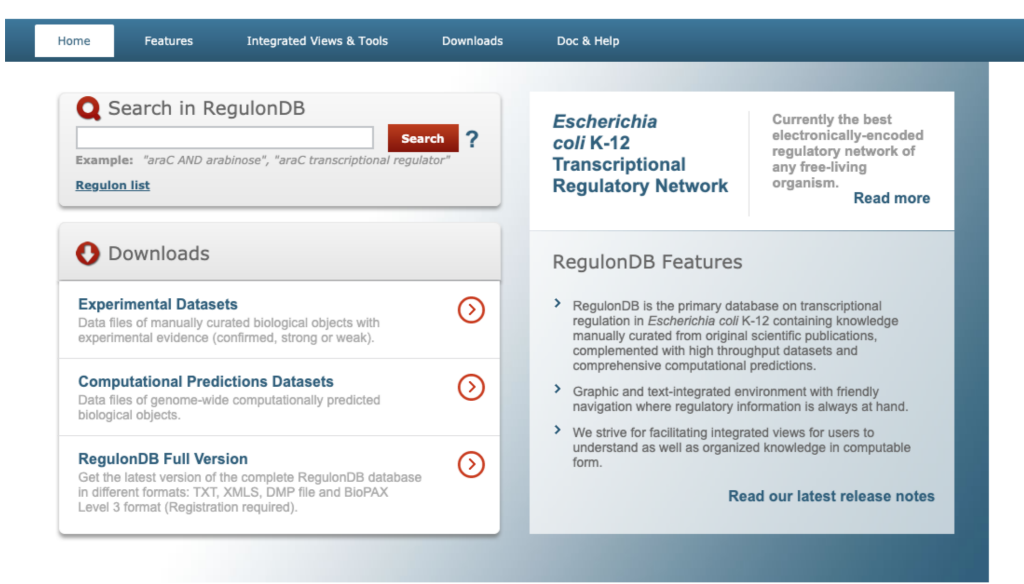
Calculate occurrence on random network and the z-score.



RegulonDB

Database of Escherichia coli K-12 Transcriptional Regulatory Network





Regulation Data

The regulation data includes information about the transcription factors (TF) that activate or repress the expression of the genes with associated supporting evidences.

```
# Release: 10.6.2 Date: 10-04-2019
# Columns:
# (1) Transcription Factor (TF) name
 (2) Gene regulated by the TF (regulated gene)
 (3) Regulatory effect of the TF on the regulated gene (+ activator, - repressor, +- dual, ? unknown)
  (4) Evidence that supports the existence of the regulatory interaction
                        [BCE, BPP, GEA, HIBSCS] Strong
AcrR
        acrA
                        [BCE, BPP, GEA, HIBSCS] Weak
AcrR
       acrB
       acrR
                        [AIBSCS, BCE, BPP, GEA, HIBSCS] Weak
AcrR
       flhC
                        [GEA, HIBSCS]
AcrR
                                       Weak
AcrR
       flhD
                        [GEA, HIBSCS]
                                       Weak
                        [BPP, GEA, HIBSCS]
AcrR
       marA
                                               Strong
                        [BPP, GEA, HIBSCS]
                                               Strong
AcrR
       marB
                       [BPP, GEA, HIBSCS]
                                               Strong
AcrR
        marR
       micF
                        [AIBSCS]
AcrR
                                       Weak
                        [BPP, GEA, HIBSCS]
AcrR
                                               Strong
        soxR
```

Nodes ad Edges

With networkx we can assign attributes to nodes and edges

```
>>> G=nx.DiGraph()
>>> G.add_node(1, color='blue')
>>> G.add_node(2, color='red')
>>> G.add_edge(1, 2, sign='+')
>>> G.node[1]
>>> G.edge[1][2]
```

Matches Node and Edges

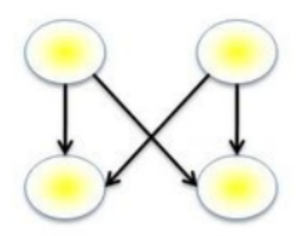
Matches can be performed based on node and edges attributes

```
>>> import networkx.algorithms.isomorphism as iso
>>> em=iso.categorical_edge_match('sign'='+')
>>> nm=iso.categorical_node_match('color'='red')
>>> nx.is_isomorphic(G1,G2,edge_match=em, node_match=nm)
```

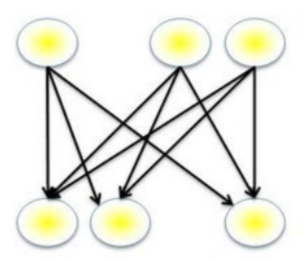
Exercise

Write a program to analyze the RegulonDB network considering only data with strong supporting information.

- Find the TF that regulates more genes (activation and suppression)
- Find the gene that is regulated by more TFs
- Find the Double-Positive Feedback loop and Multi-Input module



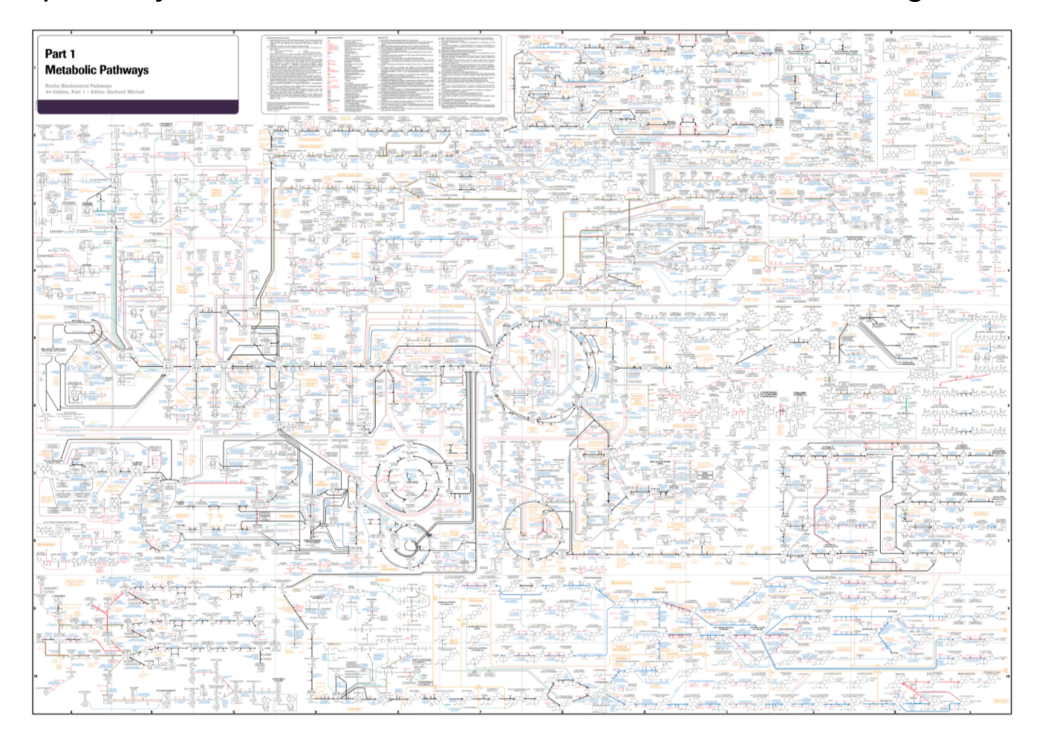
Double-Positive Feedback loop



Multi-Input module

Metabolic Pathway

Metabolic pathway is a linked series of chemical reactions occurring within a cell.



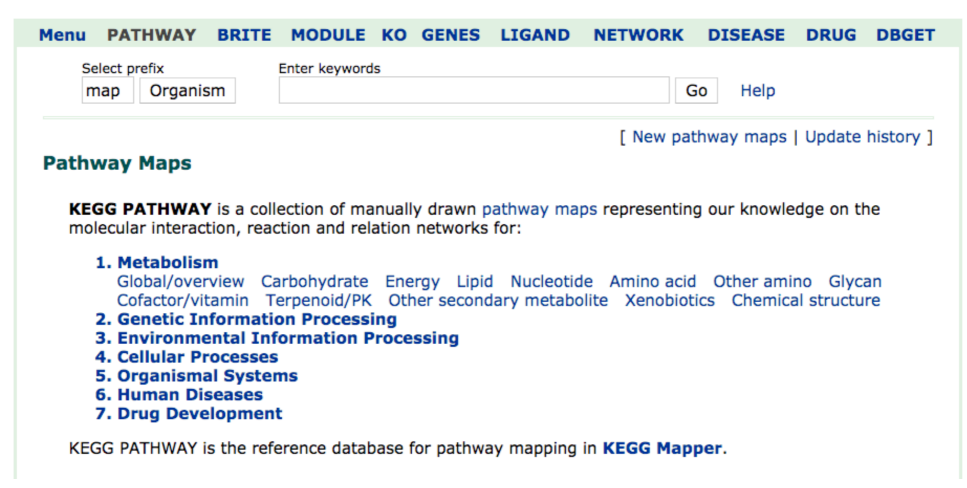
KEGG Database

It is the Kyoto Encyclopedia of Genes and Genomes. It collects many databases the most important one is KEGG Pathway which contains maps divided in 7 groups



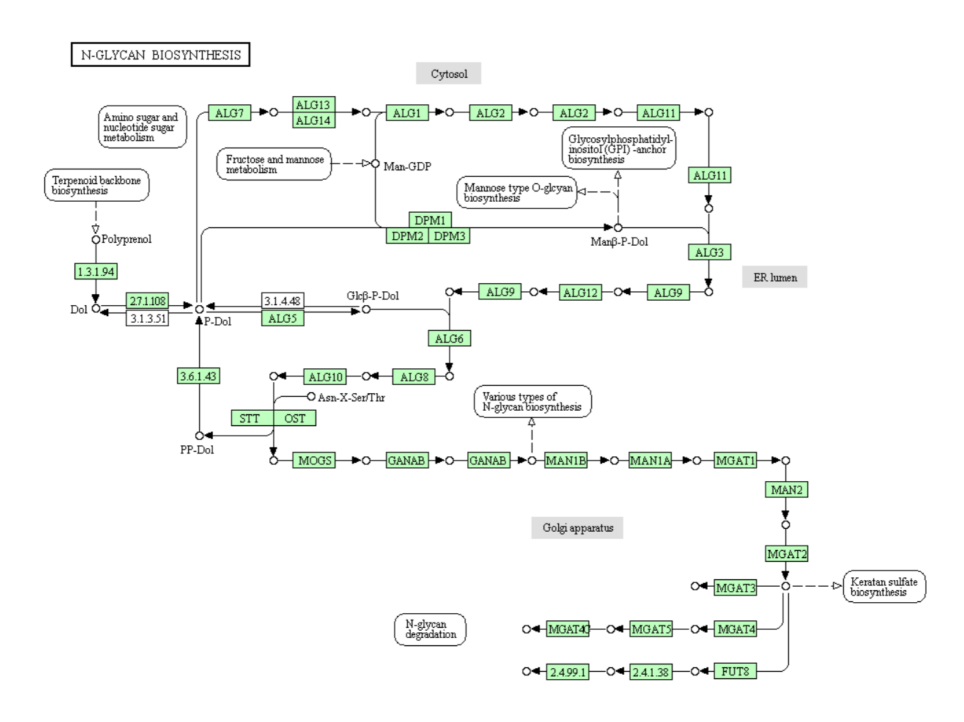
KEGG PATHWAY Database

Wiring diagrams of molecular interactions, reactions and relations



Pathway Map

It is a representation of a set of reactions in which the reactants, products, and intermediates of an enzymatic reaction, known as metabolites, are modified by a sequence of reactions catalyzed by enzymes.



KEGG Data

Given a pathway KEGG provide several information about the genes, the metabolites and the enzymes involved in the series of reactions

KFGG	PATH

PATHWAY: map00510

Help Entry map00510 Pathway N-Glycan biosynthesis Name Description N-glycans or asparagine-linked glycans are major constituents of glycoproteins in eukaryotes. N-glycans are covalently attached to asparagine with the consensus sequence of Asn-X-Ser/Thr by an Nglycosidic bond, GlcNAc b1- Asn. Biosynthesis of N-glycans begins on the cytoplasmic face of the ER membrane with the transferase reaction of UDP-GlcNAc and the lipid-like precursor P-Dol (dolichol phosphate) to generate GlcNAc a1- PP-Dol. After sequential addition of monosaccharides by ALG glycosyltransferases [MD:M00055], the N-glycan precursor is attached by the OST (oligosaccharyltransferase) complex to the polypeptide chain that is being synthesized and translocated through the ER membrane. The protein-bound N-glycan precursor is subsequently trimmed, extended, and modified in the ER and Golgi by a complex series of reactions catalyzed by membrane-bound glycosidases and glycosyltransferases. N-glycans thus synthesized are classified into three types: high-mannose type, complex type, and hybrid type. Defects in N-glycan biosynthesis lead to a variety of human diseases known as congenital disorders of glycosylation [DS:H00118 H00119]. Metabolism; Glycan biosynthesis and metabolism Class BRITE hierarchy

All links Pathway (5) KEGG MODULE (5) Disease (2) KEGG DISEASE (2) Chemical substance (38) KEGG COMPOUND (8) KEGG GLYCAN (30) Chemical reaction (72) KEGG ENZYME (35) KEGG REACTION (37) Gene (117824) KEGG ORTHOLOGY (45) RefGene (117779) Literature (6) PubMed (6) All databases (117947) Download RDF

KGML Format

The KEGG Markup Language (KGML) is an exchange format of the KEGG pathway maps.

The KGML files for metabolic pathway maps contain two types of graph object patterns:

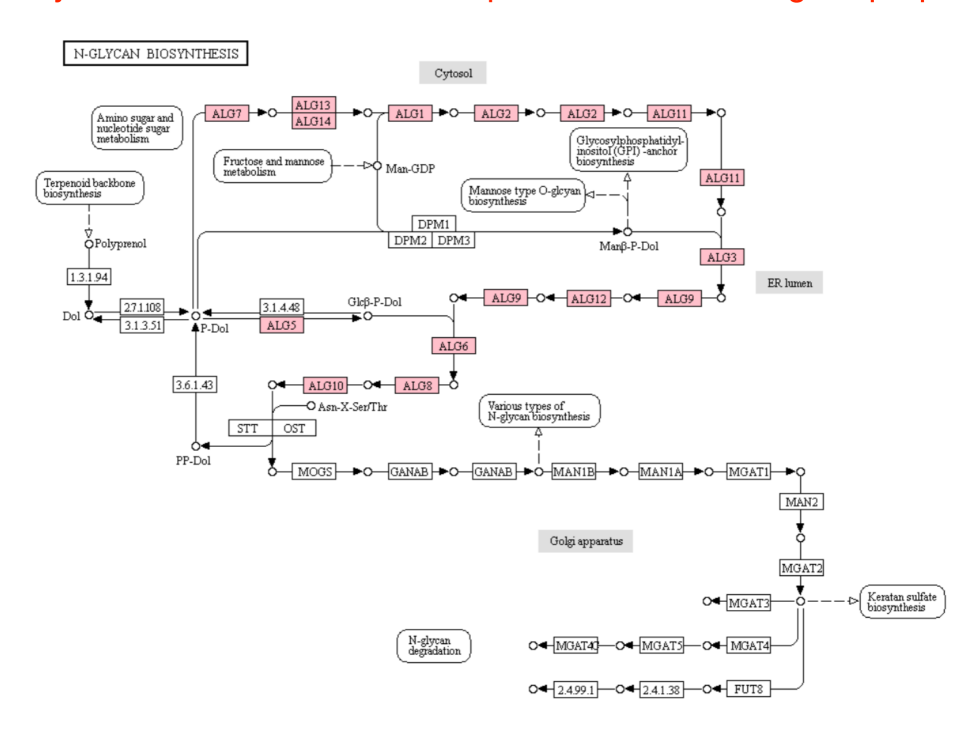
- boxes (enzymes) are linked by "relations"
- circles (chemical compounds) are linked by "reactions".

The information are provided in xml format. Enzymes are always indicated with Enzyme Commission number (EC number).

The EC number is composed by four numbers separated by periods. Those numbers represent a progressively finer classification of the enzyme.

Network Modules

A module is a set of genes/proteins performing a distinct biological function are characterized by coherent behavior with respect to certain biological property.



Exercise

Download kegg-kgml-parser-python from GitHub and install on your machine.

 Use the program to parse a KGML file and generate the network with networkx.

 Identify the module N-glycan precursor biosynthesis (M00055) and visualize it in the graph assigning a different color to its nodes.