### Network analysis with Cytoscape

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in lecture on systems bioinformatics

02/08/2012

#### Outline

- Introduction
- 2 Code
- Results
- 4 Literature

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#### Introduction I

 $\ldots$  why should we care about glycolysis and gluconeogeneses

#### Introduction

- using the tool CellDesigner to build a glycolysis/gluconeogenese network,
- import it into the tool Cytoscape,
- analyse the imported network and
- implement missing centralities and indices by coding some plugins

### Glycolysis/Gluconeogenese

- catabolic linear pathway of glycolysis deals with the breakdown and extraction of energy from glucose
- the reverse anabolic process gluconeogenesis is equally important
- gluconeogenesis helps to keep blood glucose levels within critical limits
- these processes provide many points for regulation

### Glycolysis

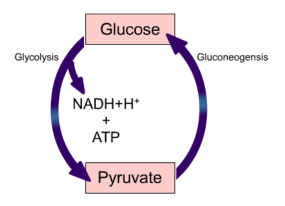


Fig. 1: Glycolysis vs. Gluconeogenese

### Glycolysis

The overall process of gluconeogenesis is energy intensive:

Compare this with glycolysis:

Fig. 2: Glycolysis vs. Gluconeogenese

### Glycolysis and Disease

- Genetic disease mutations are generally rare due to importance of the metabolic pathway
- Cancer typically tumor cells have glycolytic rates that are up to 200 times higher
- Other disease disfunctioning glycolysis or glucose metabolism has been associated with some other diseases

### Glycolysis and Cancer

- Ubiquitous gene overexpression appears to be restricted to glycolysis, in conclusion, Glycolysis is indeed special.
- this may also be of some interest for therapy, increased Glucose consumption can be observed with clinical tumour imaging
- Gene expression patterns in general can be modified by external factors such as drugs or components of nutrition
- one may envision substances that modify expression of glycolysis genes as complementary to conventional cancer therapies

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#### Cell Designer

- tried to model a glycolysis network
- tool very user unfriendly
- export into a format for cytoscape resulted in "strange" networks
- this task was skipped

### Cytoscape

- provides many useful plugins
- plugin KGMLReader used to import KEGG Glycolysis network
- modified the network to make it user readable

0

#### Introduction II

... basic setup of our two networks ... how we modelled them and what did not work (CellDesigner) ...

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## Plugin for calculation of Katz's Index

#### Definition (Katz's Index)

Let A denote the adjacency matrix of a **directed** or **undirected** graph with  $dim(A) = n \times n$  and let  $\alpha$  denote a non-negative damping factor.

$$I_{Katz}(i) = \sum_{k=1,\dots,\infty} \sum_{j=1,\dots n} \alpha(A^k)_{ji}$$
 (1)

 $\Leftrightarrow$ 

$$I_{Katz} = ((\mathbb{I}_m - \alpha A^T)^{-1} - \mathbb{I}_m)\mathbb{I}_v$$
 (2)

For calculations we used Eq. 2 and considered two distinct cases:

- **1**  $\alpha$  small  $\Rightarrow$  emulation of degree centrality
- $\alpha \in ]0, \frac{1}{|\lambda_{\max}|}] \Rightarrow \text{ influence of pathes weighted}$



#### Listing 1: Necessary import statements

```
1 package katz.plugin;
2 import java.util.List;
4 import java.awt.event.ActionEvent;
5 import javax.swing.JOptionPane;
6 import javax.text.DecimalFormat;
7
8 import cytoscape.plugin.CytoscapePlugin;
9 import cytoscape.util.CytoscapeAction;
10 import cytoscape.CyNetwork;
11 import cytoscape.CyNetwork;
12 import cytoscape.CyNetworkview;
14
15 import Jama.Matrix;
```

#### Listing 2: Basic class definition

```
public class KatzPlugin extends CytoscapePlugin {
16
17
18
       public KatzPlugin() {
19
            KatzPlugin. MolbiPluginAction action = new KatzPlugin. MolbiPluginAction();
20
            action.setPreferredMenu("Plugins.Molbi");
21
            Cytoscape . getDesktop() . getCyMenus() . addAction(action):
22
23
24
       public String describe() {
25
            return "Plugin to calculate the topological Katz Index for networks":
26
27
28
       public class MolbiPluginAction extends CytoscapeAction {
29
30
            public MolbiPluginAction() {
31
                super("Katz's Index (alpha = 0.1)"):
32
33
34
            OOverride.
35
            public void actionPerformed(ActionEvent ae) {
36
                run();
37
```

#### Listing 3: Run method (I)

```
38
           private void run() {
39
40
                CyNetwork network = Cytoscape.getCurrentNetwork();
                CyNetworkView view = Cytoscape.getCurrentNetworkView();
41
42
                int N = network.getNodeCount() + 1:
43
44
                if (N = 1) {
45
                    JOptionPane.showMessageDialog(view.getComponent(), "No network/view
                         loaded."):
46
                    return:
47
48
49
                double[][] A = new double[N][N];
50
                for (double [] row : A) Arrays, fill (row, 0.0):
51
52
                for (CyEdge edge : (List < CyEdge >) network.edgesList()) {
53
                    int i = Math.abs(edge.getSource().getRootGraphIndex());
                    int i = Math.abs(edge.getTarget().getRootGraphIndex());
54
55
                    A[i][i] = 1:
56
```

#### Listing 4: Run method (II)

```
57
               Matrix M = new Matrix(A);
58
                Matrix I = Matrix.identity(N. N):
59
                Matrix IVec = new Matrix(N, 1, 1.0):
60
61
               double alpha = 0.1; // emulates degree centrality
62
               double [][] values = ((I.minus(M.transpose().times(alpha))).inverse()).
                     minus(I).times(IVec).getArrayCopy();
63
64
                StringBuilder sb = new StringBuilder();
65
               DecimalFormat df = new DecimalFormat("#0.000"):
66
               for (int i = 1: i < N: i++) {
67
                    if (i % 10 == 0) sb.append("\n");
68
                    sb.append("C(").append(i).append("): ").append(df.format(values[i
69
                         [[0])).append(" ");
70
71
72
               JOptionPane.showMessageDialog(view.getComponent(), "All done.\n" + sb);
73
               view.redrawGraph(false. true):
74
75
76 }
```

# Plugin for calculation of Katz's Index - $\alpha \equiv \frac{1}{|\lambda_{max}|}$

#### Listing 5: Run method (II)

```
57
               Matrix M = new Matrix(A);
58
               Matrix I = Matrix.identity(N, N);
               Matrix IVec = new Matrix(N. 1. 1.0):
59
60
               double[] eigs = M. eig().getRealEigenvalues();
61
               double alpha;
62
63
               Arrays.sort(eigs):
64
65
               if (eigs[eigs.length - 1] == 0) alpha = 0.1; // emulates degree
                     centrality
66
                else alpha = 1.0 / Math.abs(eigs[eigs.length - 1]);
67
68
               double [][] values = ((I.minus(M.transpose().times(alpha))).inverse()).
                     minus(I).times(IVec).getArrayCopy();
69
                StringBuilder sb = new StringBuilder();
               DecimalFormat df = new DecimalFormat("#0.000"):
70
71
72
               for (int i = 1; i < N; i++) {
73
                    if (i % 10 == 0) sb.append("\n");
                    sb.append("C(").append(i).append("): ").append(df.format(values[i
74
                         [[0])).append(" ");
75
76
77
               JOptionPane.showMessageDialog(view.getComponent(), "All done.\n" + sb);
78
               view.redrawGraph(false, true);
79
80
81 }
                                                            4 D > 4 B > 4 B > 4 B >
```

#### Definition (Randics's Index)

Let v(i) denote the #neighbors of vertex v with index i and N #nodes.

$$I_{Randic} = \sum_{i=0}^{N-1} \frac{1}{\sqrt{v(i)}} \tag{3}$$

#### Cave

Randic's Index works only for undirected and connected graphs.

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#### Cave

Randic's Index works only for undirected and connected graphs.

#### **Application**

Currently there is no known application of Randic's Index for the analysis of biological networks.

#### Listing 6: Necessary import statements

```
1 package randic.plugin;
2
3 import java.util.List;
4 import java.awt.event.ActionEvent;
5 import javax.swing.JOptionPane;
6 import java text.DecimalFormat;
7
8 import cytoscape.plugin.CytoscapePlugin;
9 import cytoscape.util.CytoscapeAction;
10 import cytoscape.CyNetwork;
11 import cytoscape.CyNetwork;
12 import cytoscape.CyNetwork;
13 import cytoscape.View.CyNetworkView;
14
15 import Jama.Matrix;
```

#### Listing 7: Basic class definition

```
public class RandicPlugin extends CytoscapePlugin {
16
17
18
       public RandicPlugin() {
19
            RandicPlugin . MolbiPluginAction action = new RandicPlugin . MolbiPluginAction();
20
            action.setPreferredMenu("Plugins.Molbi");
21
            Cytoscape . getDesktop() . getCyMenus() . addAction(action):
22
23
24
       public String describe() {
25
            return "Plugin to calculate the Index of Randic for networks":
26
27
28
       public class MolbiPluginAction extends CytoscapeAction {
29
30
            public MolbiPluginAction() {
31
                super("Randic's Index"):
32
33
34
            OOverride.
35
            public void actionPerformed(ActionEvent ae) {
36
                run();
37
```

#### Listing 8: Run method

```
38
           private void run() {
39
                CyNetwork network = Cytoscape.getCurrentNetwork();
                CyNetworkView view = Cytoscape.getCurrentNetworkView();
40
41
                int N = network.getNodeCount();
42
43
                if (N = 0) {
                    JOptionPane.showMessageDialog(view.getComponent(), "No network/view
44
                         loaded."):
45
                    return:
46
47
48
                Double IRandic = 0.0:
49
                for (CyNode node : (List < CyNode >) network.nodesList())
                   // otherwise Randic's Index will be +infinity!
50
                    if (network.getDegree(node) != 0)
51
52
                      IRandic += Math.pow(network.getDegree(node), -0.5);
53
54
                JOptionPane.showMessageDialog(view.getComponent(), "All done.\nRandic
                     Index " + IRandic):
55
                view.redrawGraph(false, true);
56
57
58
```

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#### Results - Centralities

- determine the relative importance of a vertex within a graph
- centraltity concepts were first developed in social network analysis
  - -> e.g. how influential a person is within a social network

#### Definition (Four measures of centrality)

- degree centrality
- eccentricity centrality
- closeness centrality
- betweeness centrality

### Results - Degree Centrality

#### Definition (Degree Centrality)

$$C_{deg}(v) = \{e | e \in E \text{ and } v \in e\}$$

- counts the number of edges attached to a vertex
- a local centrality measure
  - -> only direct neighborhood considered
- directed graphs:
  - -> indegree (interpreted as "popularity")
  - -> outdegree (interpreted as "gregariousness")

## Results - Degree Centrality

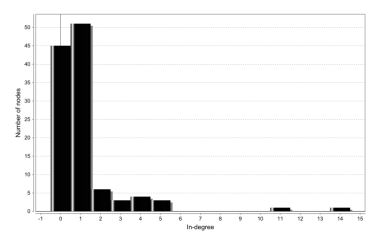


Fig. 3: Indegree Centrality

## Results - Degree Centrality

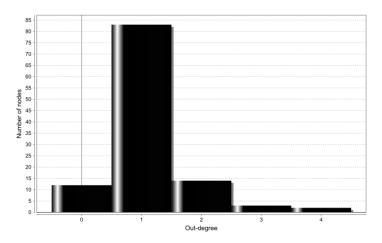


Fig. 4: Outdegree Centrality

### Results - Eccentricity Centrality

#### Definition (Eccentricity Centrality)

$$C_{ecc}(s) = \frac{1}{\max\{d_{st}|t \in V\}}$$

- determine the maximum distance between every two vertices
- central vertices get low values
- centralities require high centrality values
  - -> reciprocal is used as centrality value
- only for CONNECTED networks

## Results - Eccentricity Centrality

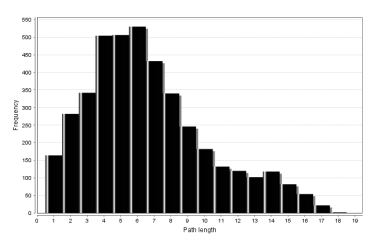


Fig. 5: Eccentricity Centrality

### Results - Closeness Centrality

#### Definition (Closeness Centrality)

$$C_{clo}(s) = \frac{1}{\sum\limits_{t \in V} d_{st}}$$

- $C_{clo}$  of a vertex s is the sum of shortest path form s to all other vertices
- degree of interaction in the network
- the closer a point is to all the rest, the more effective and independent he can reach them

## Results - Closeness Centrality

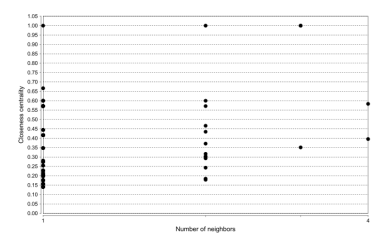


Fig. 6: Closeness Centrality

### Results - Betweenness Centrality

#### Definition (Betweenness Centrality)

$$C_{spb}(v) = \sum\limits_{s \in V \; and \; s 
eq v \; t \in V \; and \; s 
eq v} \delta_{st}(v)$$

- center of attention: indirect relationships
  - -> Control of interaction
- v is the more powerful the more shortest paths between other vertices it can interrupt

## Results - Betweenness Centrality

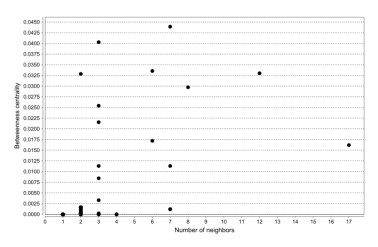


Fig. 7: Closeness Centrality

#### Results - Conclusion

- the analysis tools are not very helpful
  - -> cannot see which vertex has which centrality
  - -> only give a statistical overview of the whole network
- but you can select a vertex in the network window and get an overview of all centralities from the selected vertex

## Results - Betweenness Centrality

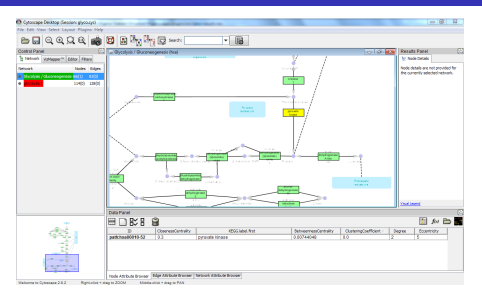


Fig. 8: Centrality Overview

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#### Literature



#### [Gatenby and Gillies]

Why do cancers have high aerobic glycolysis? Nature Review Cance, Vol. 4, p. 891-899, November 2004.



#### [Altneberga and Greulich]

B. Altenberga and K.O. Greulich, Genes of glycolysis are ubiquitously overexpressed in 24 cancer classes.

Genomics Vol. 84, p. 1014-1020, September 2004.