







Knowledge-guided Algorithms in Systems Biology

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Computational Genomics Course

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http://www.ece.mcgill.ca/~aemad2/











Plan for this Lecture

Topic: Methods for analyzing omics datasets while integrating prior knowledge

- Systems Biology and Knowledge Networks
- Sample Clustering
- Gene Prioritization
- Gene Set Characterization

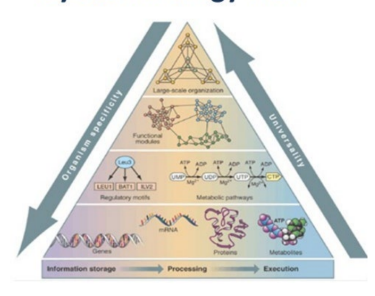
Emphasis: tools that take advantage of prior knowledge networks (KnowEnG)

Goal: understand basic concepts and aware of approaches and resources

Systems Biology

 Systems biology is the computational and mathematical modeling of complex biological systems.

System biology view



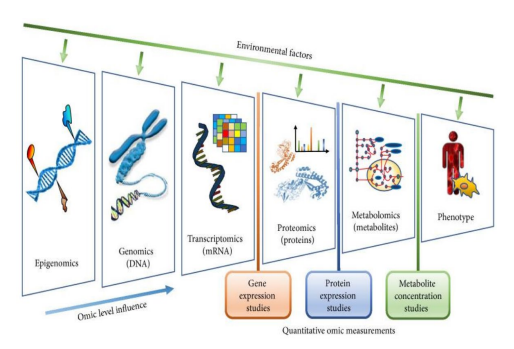


Figure from Oltvai, Z.N. and Barabasi Life's complexity pyramid.

Figure from Angione, C. Human Systems Biology and Metabolic Modelling: A Review-From Disease Metabolism to Precision Medicine. Biomed Res Int 2019.

 Studies the interactions between the components of biological systems such as genes, proteins, metabolites, etc. (i.e. biological networks), and how these interactions give rise to the function and behavior of that system (phenotype)

Statistical and Machine Learning Methods

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Applied to heterogeneous 'omics and phenotype data and prior knowledge

Unsupervised Learning

Supervised Learning

- No training example exists and the goal is to learn structure in the data
- Training examples are provided with desired inputs and outputs to help learning the desired rule

Clustering (subtyping)

Classification (resistance group)

Regression (survival time)

Dimensionality Reduction
(data visualization)

Supervised Feature Selection
(biomarkers)

Some Example Applications

Clustering (subtyping)

Identifying the subtypes of a disease

Supervised Feature Selection (biomarkers)

 Identifying genes associated with a disease

Classification (resistance group)

 Predicting whether a patient is sensitive or resistant to a drug

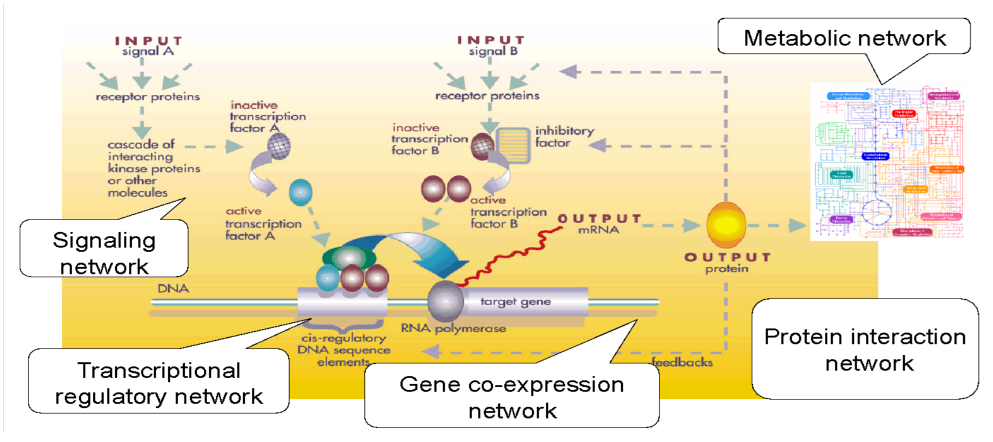
Regression (survival time)

- Predicting the survival probability of a cancer patient
- etc.

Prior Knowledge as Biological Networks

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- Existing prior knowledge in literature captures known interactions within and across different levels of the biological systems
- Knowledge Network a graphical representation of the interactions of the components of a biological systems

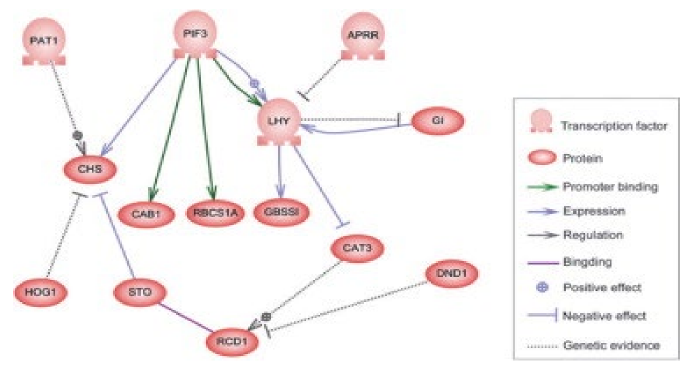


Directed Biological Networks

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Gene regulatory networks

- Nodes represent genes, proteins, etc.
- Edges show regulatory relationships between the nodes
- The network shows which entities (e.g. transcription factors) regulate the expression of each gene
- Edges can have meaningful weights



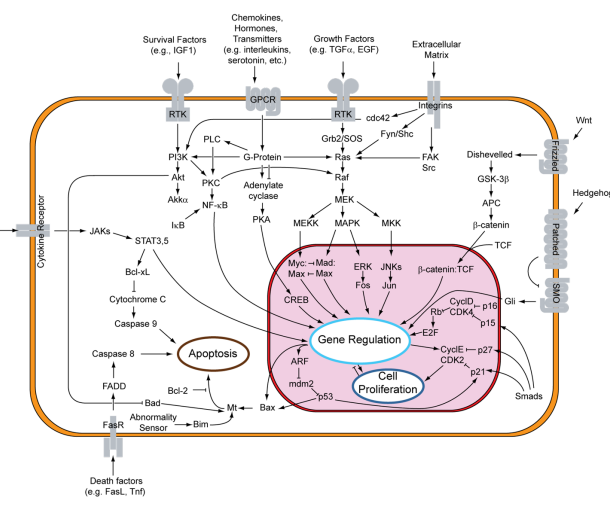
Directed Biological Networks

Signaling Networks

 Represents communications within and between cells

 Responsible for receiving, transmitting and processing information

 The network is a graphical representation of the interactions of the components of a biological systems



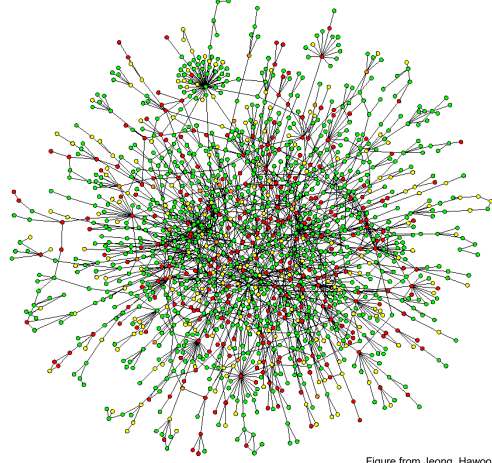
Signal Transduction Pathway

Experimental Networks

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Protein-protein interaction networks

- Nodes represent proteins
- Edges show interactions between proteins
- Interactions usually refer to different levels of physical contact and proximity of protein molecules



Experimental Networks

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Gene co-expression networks

- Nodes represent genes
- An edge exists between two genes that are highly co-expressed across different samples

BMC Bioinformatics. 2008; 9: 559. Published online 2008 Dec 29. doi: 10.1186/1471-2105-9-559

PMCID: PMC2631488

WGCNA: an R package for weighted correlation network analysis

Reviewed by Peter Langfelder¹ and Steve Horvath ^{™2}

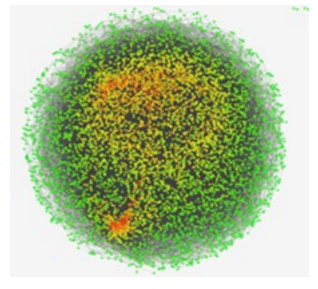


Figure from https://commons.wikimedia.org/wiki/File:Gene coexpression network with 7221 genes for 18 gastric cancer patients.png

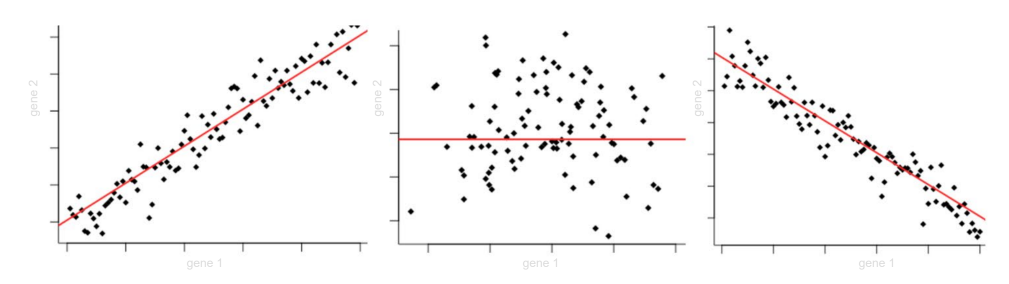


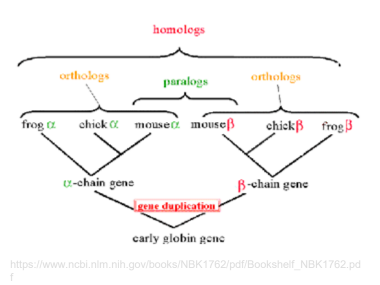
Figure from https://www.freecodecamp.org/news/how-machines-make-predictions-finding-correlations-in-complex-data-dfd9f0d87889/

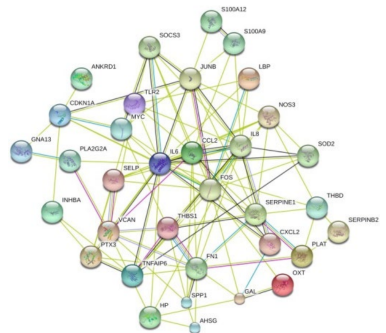
Computational Networks

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Evolutionary Conservation networks

- Nodes represent gene DNA or protein amino acid sequences
- Edges represent the similarity between the pair of sequences, the more similarly the more recently the nodes share an evolutionary history





Text Mining networks

- Nodes represent gene entities
- Edges represent the frequency names, aliases, and synonyms for a pair of genes co-occur in literature abstracts

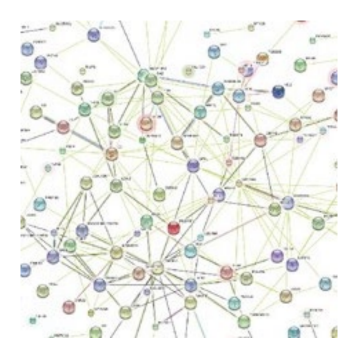
Figure from Yahaya, et al. "Gene expression changes associated with the airway wall response to injury." *PloS one* 8.4 (2013).

Computational Networks

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Integrated networks

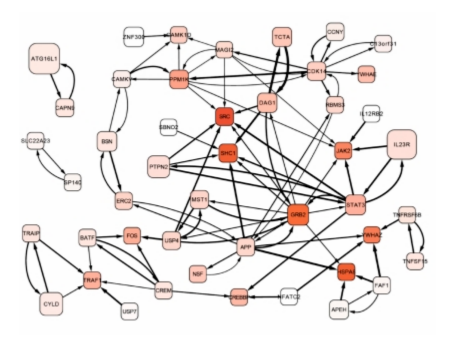
- Nodes represent gene or proteins
- Edges represent the weighted combination of normalized edge weights from many different types of network edges based on some predetermined criteria



Nucleic Acids Res. 2015 Jan;43(Database issue):D447-52. doi: 10.1093/nar/gku1003. Epub 2014 Oct 28.

STRING v10: protein-protein interaction networks, integrated over the tree of life.

Szklarczyk D1, Franceschini A1, Wyder S1, Forslund K2, Heller D1, Huerta-Cepas J2, Simonovic M1, Roth A1, Santos A3, Tsafou KP3, Kuhn M4, Bork P5, Jensen LJ6, von Mering C7.



Genome Res. 2011 Jul;21(7):1109-21. doi: 10.1101/gr.118992.110. Epub 2011 May 2.

Prioritizing candidate disease genes by network-based boosting of genome-wide association data.

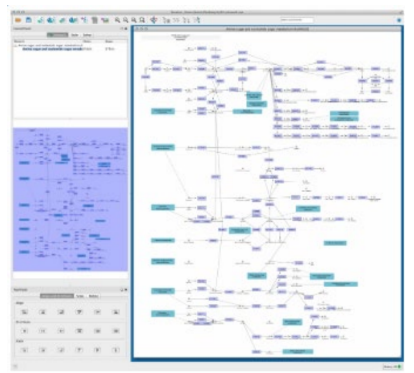
Lee I1, Blom UM, Wang PI, Shim JE, Marcotte EM.

Visualizing / Sharing Biological Networks

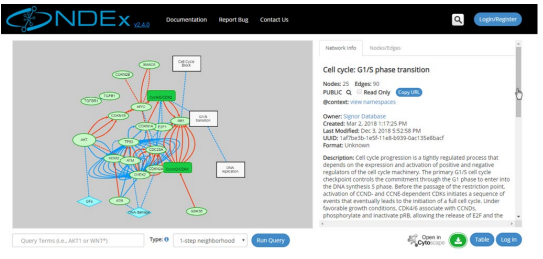








https://cytoscape.org/release_notes_3_2_1.html



https://home.ndexbio.org/quick-start/

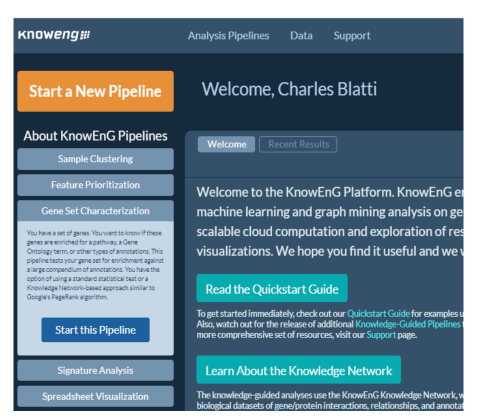


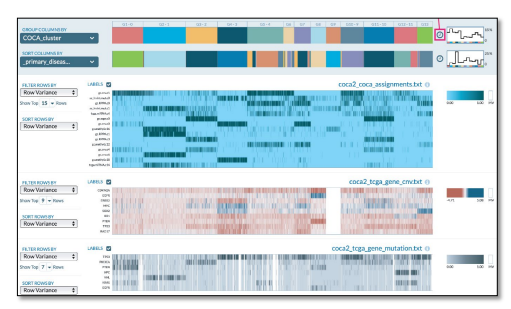






KnowEnG: Platform for Networkguided Analysis

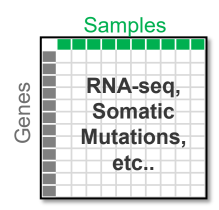




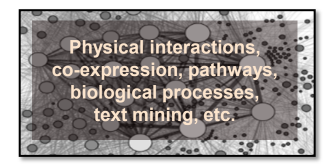
KnowEnG: Knowledge Engine for Genomics

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'omics Data Analysis Pipelines



Using Prior Knowledge



In a Scalable Cloud Platform



KnowEnG Pipelines and User Interface



Sample Clustering

 What are the separate transcriptomic subtypes of patients and how do they relate to outcome?

Feature(Gene) Prioritization

What genes are differentially expressed with respect to viral shedding

Gene Set Characterization

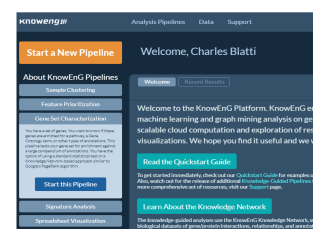
What pathways do these differentially expressed genes relate to?

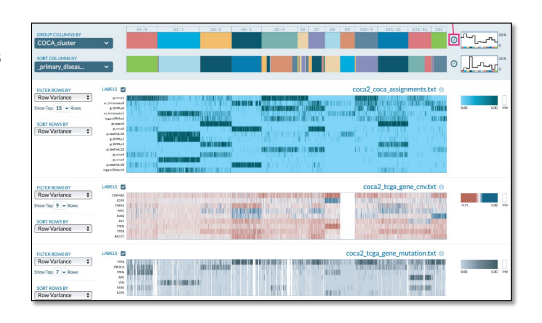
Signature Analysis

 Given a new patient, what subtype does their profile most resemble?

Spreadsheet Visualization

 Given multiple omics and clinical datasets on patient samples, what features relate to selected phenotypes?

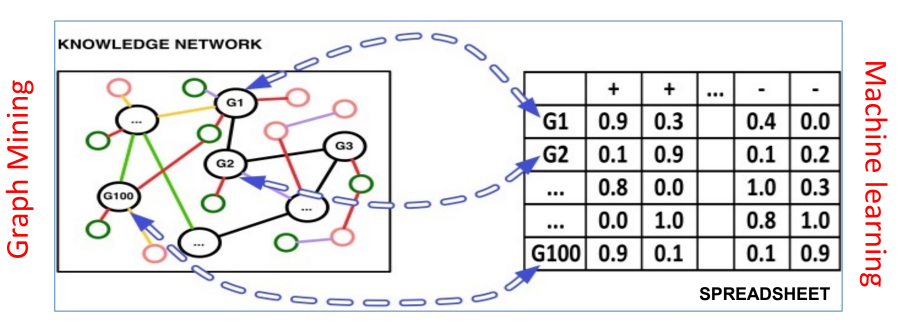




Analysis Pipelines Using Prior Knowledge

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- Knowledge Network (KN): heterogeneous graph whose nodes and edges encodes major public data sets as a network represented by genes/proteins, their properties, and relationships
- Omics data: a spreadsheet (rows = genes or proteins) to be analyzed

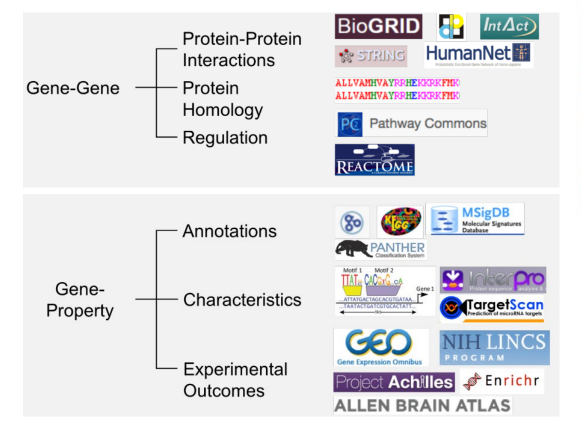


Knowledge network + user spreadsheet

KnowEnG Prior Knowledge Networks

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| KNOWLEDGE NETWORK CONTENTS: | | | | | |
|----------------------------------|---------------|--|--|--|--|
| Version: | KN-20rep-1702 | | | | |
| Number of Species: | 20 | | | | |
| Number of Resources: | 13 | | | | |
| Number of Datasets: | 159 | | | | |
| Number of Edge Types: | 43 | | | | |
| Number of Edges: | 233,459,368 | | | | |
| Number of Nodes: | 594,474 | | | | |
| Number of Gene Nodes: | 404,868 | | | | |
| Number of Property Nodes: | 189,605 | | | | |



| Edge Type Collection 💠 | Human Network Edges (millions) | Human Datasets \$ | All Network Edges (millions) | All Datasets |
|--------------------------|---|----------------------|---------------------------------------|--------------|
| Text_Mining/Integrated | 9.0 | 2 | 130.6 | 19 |
| Coexpression | 7.3 | 2 | 119.8 | 19 |
| Experimental_Interaction | 5.4 | 4 | 108.7 | 21 |
| Conservation/Proximity | 1.6 | 2 | 26.1 | 36 |
| Pathway_Database | 1.1 | 3 | 63.4 | 20 |
| Total | 24.3 | 8 | 448.7 | 42 |

| Edge Type Collection | Human Network Edges (millions) | Human Property Nodes (thousands) | ♦ Human Datasets | All Network Edges (millions) | All Property Nodes (thousands) |
|-------------------------|---|---|---------------------|---------------------------------------|---|
| Tissue_Expression | 13.7 | 25.9 | 32 | 13.7 | 25.9 |
| Disease/Drug | 6.0 | 82.3 | 13 | 6.3 | 83.4 |
| Regulation | 4.4 | 3.3 | 10 | 4.4 | 3.3 |
| Pathways | 0.6 | 16.9 | 5 | 1.4 | 34.6 |
| Ontologies | 0.3 | 17.2 | 5 | 1.8 | 23.5 |
| Protein_Domains | 0.0 | 6.2 | 2 | 0.5 | 7.8 |
| Total | 25.0 | 151.7 | 67 | 28.1 | 178.5 |

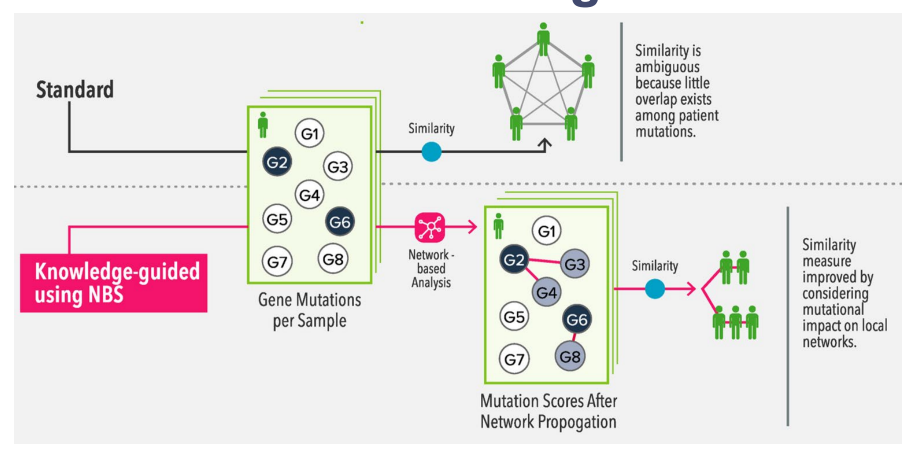








Network-guided Sample Clustering



Network-Guided Sample Clustering

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Goal:

Stratification (clustering) of tumor samples based on somatic mutation profiles

Main Issue:

- The mutation data is very sparse and most conventional clustering techniques fail to identify reasonable patterns
- Although two tumors may not share the same somatic mutations, they
 may affect the same pathways and interaction networks

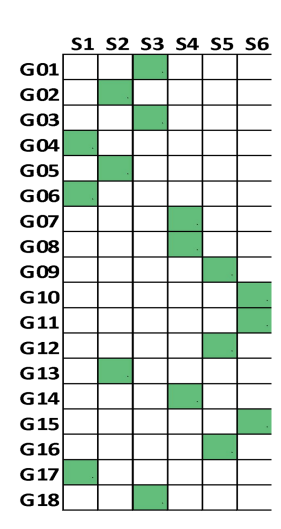
Knowledge-Guided Sample Clustering

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Problem: Data sparsity in gene-level somatic mutation data

Toy Example

 Due to the sparsity of the data, all samples are at equal distance of each other



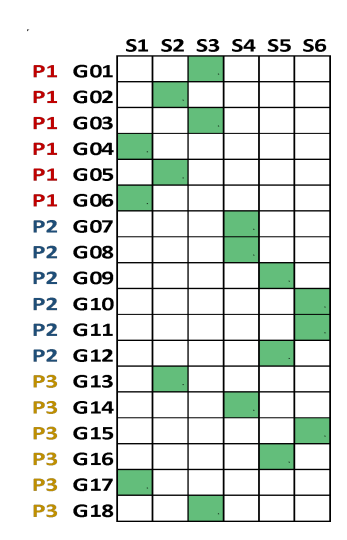
Knowledge-Guided Sample Clustering

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Problem: Data sparsity in gene-level somatic mutation data

Toy Example

- Due to the sparsity of the data, all samples are at equal distance of each other
- Pathway information clarifies the similarity among some samples



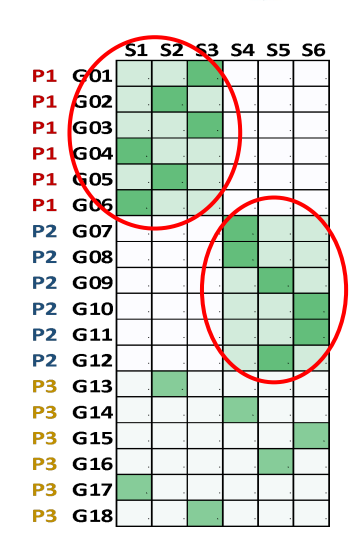
Knowledge-Guided Sample Clustering

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Problem: Data sparsity in gene-level somatic mutation data

Toy Example

- Due to the sparsity of the data, all samples are at equal distance of each other
- Pathway information clarifies the similarity among some samples
- Conventional clustering methods can then identify clusters based on networksmoothed features



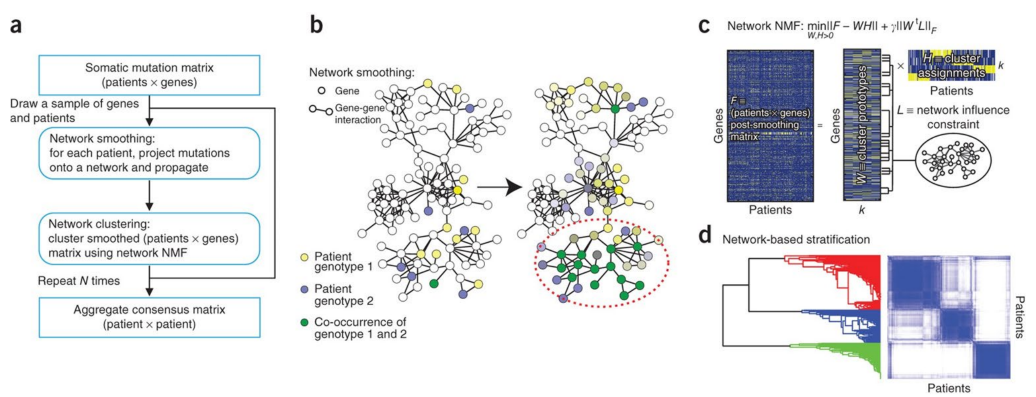
Network-based Stratification (NBS)

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Nat Methods. 2013 Nov;10(11):1108-15. doi: 10.1038/nmeth.2651. Epub 2013 Sep 15.

Network-based stratification of tumor mutations.

Hofree M¹, Shen JP, Carter H, Gross A, Ideker T.

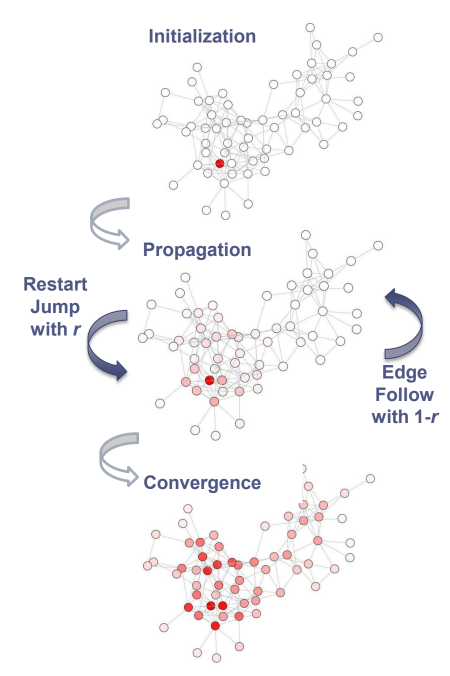


- Network Smoothing Random Walk with Restart
- Patient Sampling for Robust Clustering

Random Walk With Restart Algorithm

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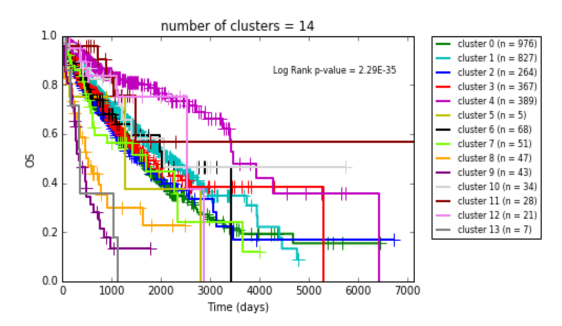
- Fast, scalable guilt-byassociation method
 - Same ideas as personalized PageRank
- Intuition
 - Walker at a node either
 - With probability 1-r, follows an outgoing edge
 - With restart probability r, returns to node in restart set
 - Converges to long run "stationary" distribution of the walker over the nodes
- Final node ranking based on distribution incorporates
 - Connectedness of node in network
 - Proximity of node to restart set



NBS Sample Clustering with KnowEnG

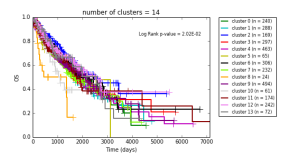
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- 3276 tumor samples from TCGA from 12 cancer projects with sparse non-synonymous somatic mutation
- Perform standard and network-guided Sample
 Clustering in platform
- Knowledge-guided clusters significantly relate to survival outcome

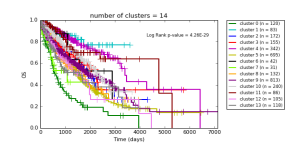


Knowledge-guided analysis of "omics" data using the KnowEnG cloud platform

 Much better than standard methods that do not incorporate prior knowledge

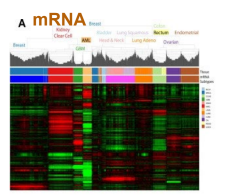


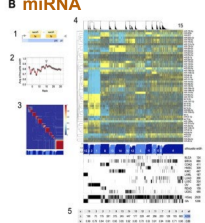
 In line with specialized method developed in TCGA paper that would be very difficult to reproduce

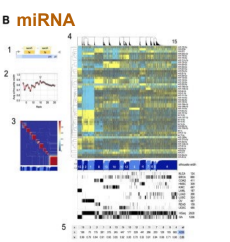


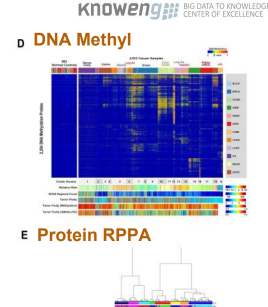
Integrating Experimental Assays for Stratification

- Data from each experimental assay is subjected to sample clustering to find cancer subtypes per assay
- Mutation data required specialized knowledge guided methods (panel F)

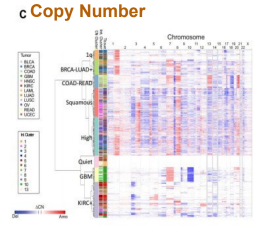


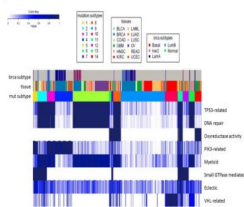










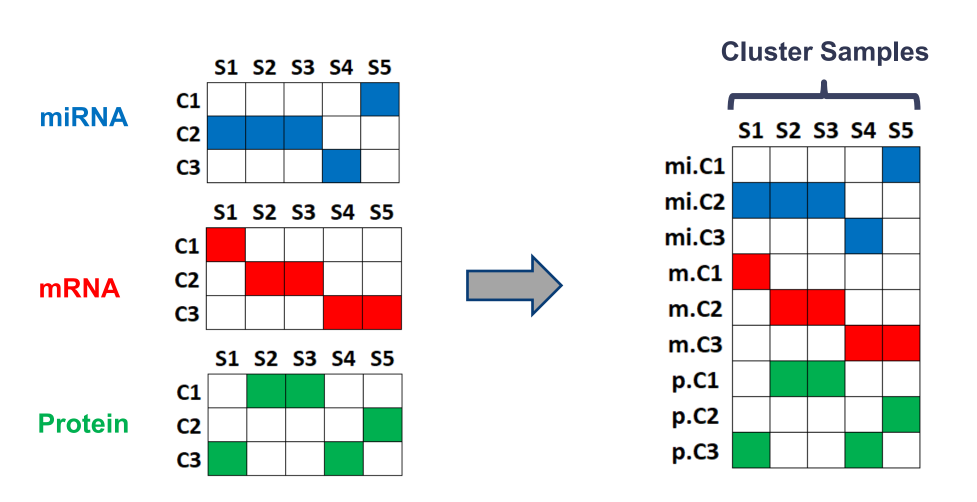




Cluster-Of-Cluster-Assignments (COCA)

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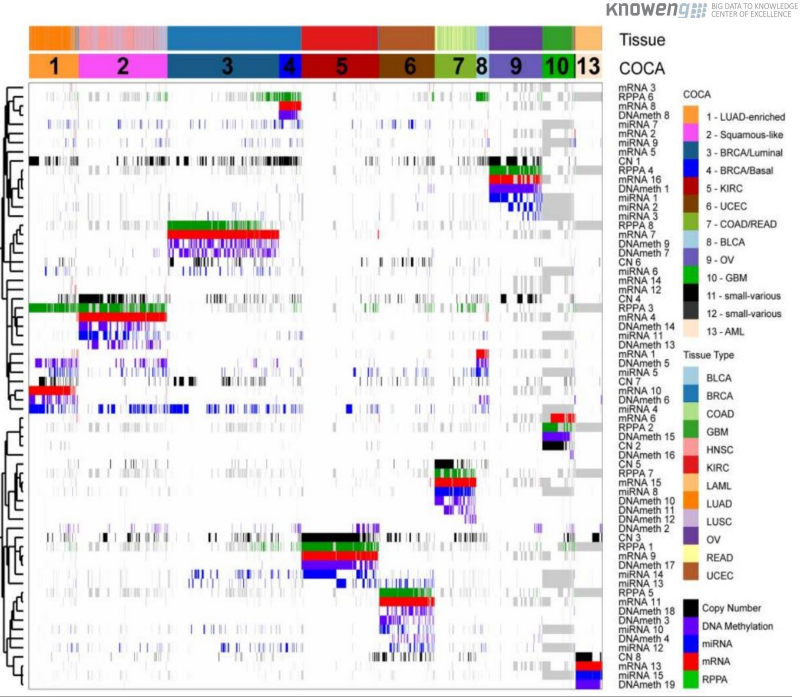
- Merge cluster assignments x samples matrices
- Cluster the samples in the multi-omics matrix



13 Cancer Subtypes from 6 Assays

Strong
 relationship
 between
 subtypes &
 disease

 Interesting relations between clusters of different data types



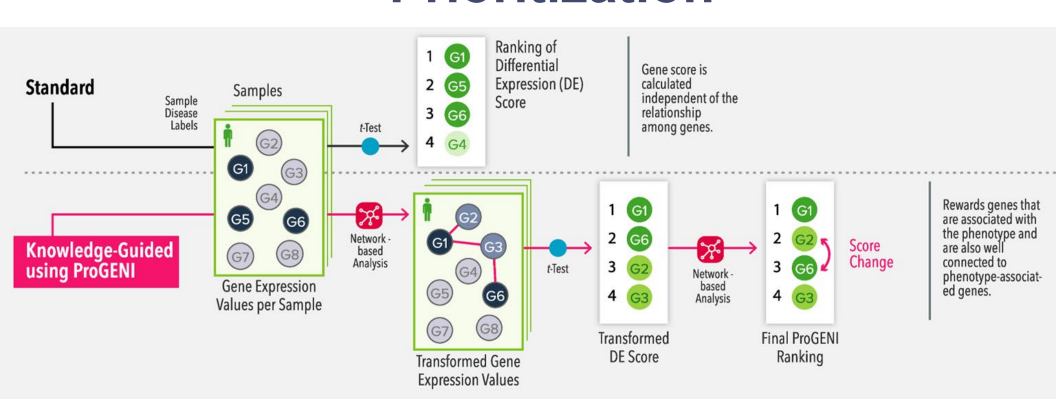






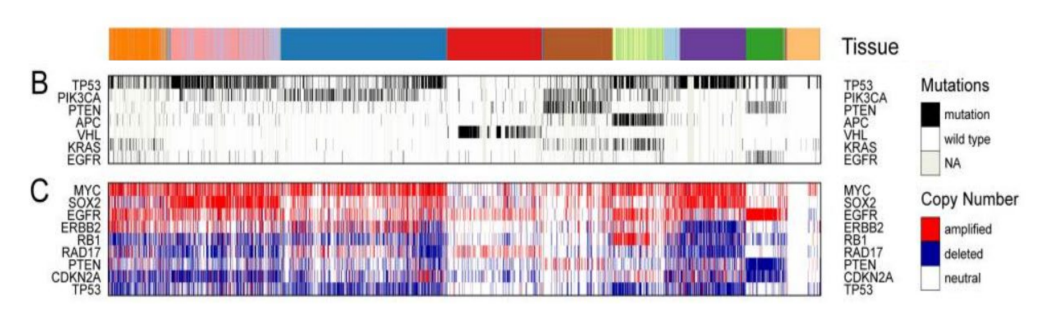


Network-Guided Gene Prioritization



Characterizing Cancer Subtypes

- Find top related mutations and copy number alterations
- Compare each subtype vs `all others`
- KnowEnG calls this `Gene Prioritization`



Towards Network-Guided Gene Prioritization

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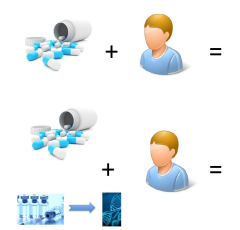
Drug Sensitivity Example

Goal:

 Identifying genes whose basal mRNA expression determines the drug sensitivity in different samples (supervised feature selection)

Motivations:

- Overcoming drug resistance
- Revealing drug mechanism of action
- Identifying novel drug targets
- Predicting drug sensitivity of individuals



Standard Gene Prioritization

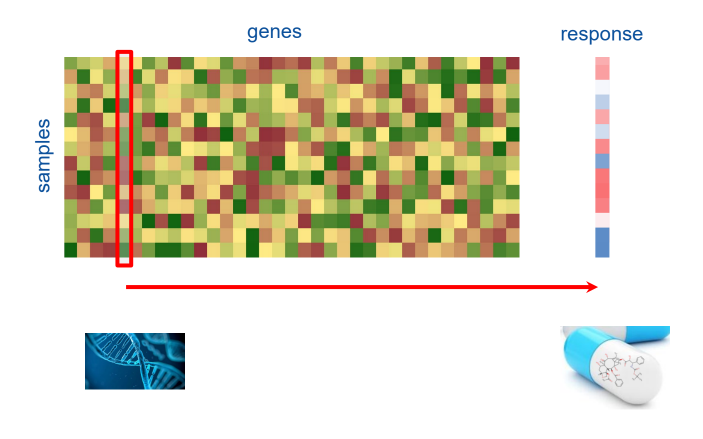


Examples of current methods:

 Score each gene based on the correlation of its expression with drug response Nat Chem Biol. 2016 Feb;12(2):109-16. doi: 10.1038/nchembio.1986. Epub 2015 Dec 14.

Correlating chemical sensitivity and basal gene expression reveals mechanism of action.

 $\begin{array}{l} \underline{\text{Rees MG}^1, \underline{\text{Seashore-Ludlow B}^{1,2}, \text{Cheah JH}^{1,2}, \underline{\text{Adams DJ}^{1,2}, \text{Price EV}^{1,2}, \underline{\text{Gill S}^1, \underline{\text{Javaid S}^3, Coletti}} \\ \underline{\text{ME}^1, Jones VL}^1, \underline{\text{Bodycombe NE}^{1,2}, \underline{\text{Soule CK}}^{1,2}, \underline{\text{Alexander B}^1, \underline{\text{Li A}^1, Montgomery P}^1, \underline{\text{Kotz JD}^1, \underline{\text{Hon CS}^1, Munoz B}^1, \underline{\text{Liefeld T}^{1,2}, \underline{\text{Dančik V}^1, Haber DA}^3, \underline{\text{Clish CB}^1, \underline{\text{Bittker JA}^1, \underline{\text{Palmer M}^{1,2}, \underline{\text{Wagner BK}^1, Clemons PA}^1, \underline{\text{Shamji AF}^1, \underline{\text{Schreiber SL}^1}}.} \end{array}$



Standard Gene Prioritization



Examples of current methods:

- Score each gene based on the correlation of its expression with drug response
- Use multivariable regression algorithms such as Elastic Net to relate multiple genes' expression values to drug response

Nat Chem Biol. 2016 Feb;12(2):109-16. doi: 10.1038/nchembio.1986. Epub 2015 Dec 14.

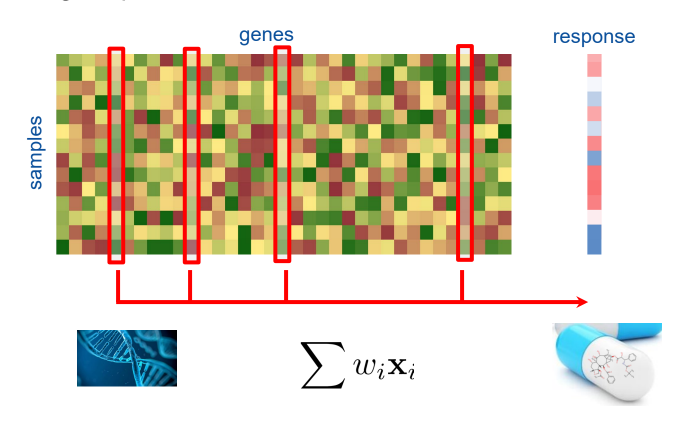
Correlating chemical sensitivity and basal gene expression reveals mechanism of action.

Rees MG¹, Seashore-Ludlow B^{1,2}, Cheah JH^{1,2}, Adams DJ^{1,2}, Price EV^{1,2}, Gill S¹, Javaid S³, Coletti ME¹, Jones VL¹, Bodycombe NE^{1,2}, Soule CK^{1,2}, Alexander B¹, Li A¹, Montgomery P¹, Kotz JD¹, Hon CS¹, Munoz B¹, Liefeld T^{1,2}, Dančík V¹, Haber DA³, Clish CB¹, Bittker JA¹, Palmer M^{1,2}, Wagner BK¹, Clemons PA¹, Shamji AF¹, Schreiber SL¹.

Nature, 2012 Mar 28;483(7391):603-7. doi: 10.1038/nature11003.

The Cancer Cell Line Encyclopedia enables predictive modelling of anticancer drug sensitivity.

Barretina J¹, Caponigro G, Stransky N, Venkatesan K, Margolin AA, Kim S, Wilson CJ, Lehár J, Kryukov GY, Sonkin D, Reddy A, Liu M, Murray L, Berger MF, Monahan JE, Morais P, Meltzer J, Korejwa A, Jané-Valbuena J, Mapa FA, Thibault J, Bric-Furlong E, Raman P, Shipway A, Engels IH, Cheng J, Yu GK, Yu J, Aspesi P Jr, de Silva M, Jagtap K, Jones MD, Wang L, Hatton C, Palescandolo E, Gupta S, Mahan S, Sougnez C, Onofrio RC, Liefeld T, MacConaill L, Winckler W, Reich M, Li N, Mesirov JP, Gabriel SB, Getz G, Ardlie K, Chan V, Myer VE, Weber BL, Porter J, Warmuth M, Finan P, Harris JL, Meyerson M, Golub TR, Morrissey MP, Sellers WR, Schledel R, Garraway LA.



Augmenting Gene Prioritization



Examples of current methods:

- Score each gene based on the correlation of its expression with drug response
- Use multivariable regression algorithms such as Elastic Net to relate multiple genes' expression values to drug response

Nat Chem Biol. 2016 Feb;12(2):109-16. doi: 10.1038/nchembio.1986. Epub 2015 Dec 14.

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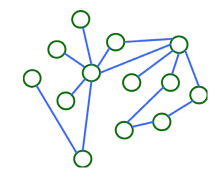
Nature, 2012 Mar 28;483(7391):603-7. doi: 10.1038/nature11003

The Cancer Cell Line Encyclopedia enables predictive modelling of anticancer drug sensitivity.

Barretina J¹, Caponigro G, Stransky N, Venkatesan K, Margolin AA, Kim S, Wilson CJ, Lehár J, Kryukov GV, Sonkin D, Reddy A, Liu M, Murray L, Berger MF, Monahan JE, Morais P, Meltzer J, Korejwa A, Jané-Valbuena J, Mapa FA, Thibault J, Bric-Furlong E, Raman P, Shipway A, Engels IH, Cheng J, Yu GK, Yu J Aspesi P Jr, de Silva M, Jagtap K, Jones MD, Wang L, Hatton C, Palescandolo E, Gupta S, Mahan S, Sougnez C, Onofrio RC, Liefeld T, MacConaill L, Winckler W, Reich M, Li N, Mesirov JP, Gabriel SB, Get G, Ardlie K, Chan V, Myer VE, Weber BL, Porter J, Warmuth M, Finan P, Harris JL, Meyerson M, Golub TR, Morrissey MP, Sellers WR, Schlegel R, Garraway LA.

Shortcoming:

 These methods do not incorporate prior information about the interaction of the genes

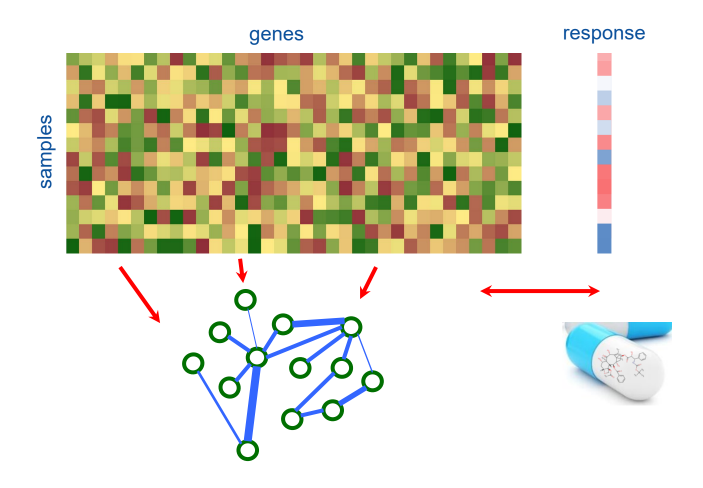


Network-Guided Gene Prioritization

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Hypothesis:

 Since genes and proteins involved in drug MoA are functionally related, prior knowledge in the form of gene interaction network (e.g. PPI) can improve accuracy of the prioritization task



ProGENI



ProGENI: Network-guided gene prioritization

An algorithm that incorporates gene network information to improve prioritization accuracy



Featured article: new insights into mechanisms of chemoresistance



Emad et al. Genome Biology (2017) 18:153 DOI 10.1186/s13059-017-1282-3

Genome Biology

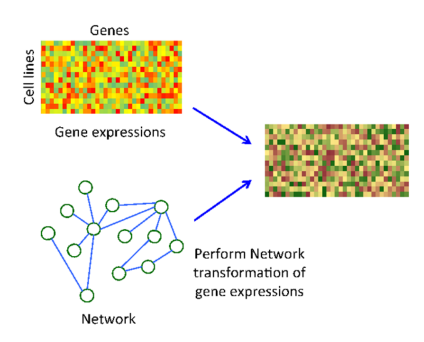
RESEARCH

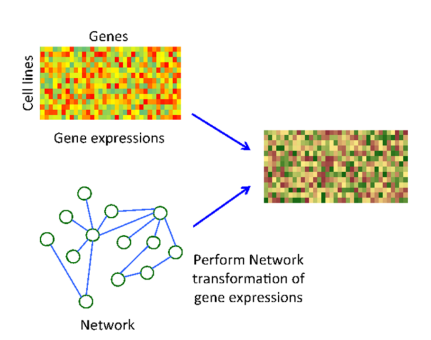
Open Access

CrossMark

Knowledge-guided gene prioritization reveals new insights into the mechanisms of chemoresistance

Amin Emad¹, Junmei Cairns², Krishna R. Kalari³, Liewei Wang^{2*} and Saurabh Sinha^{4*}





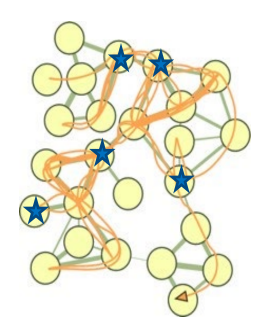
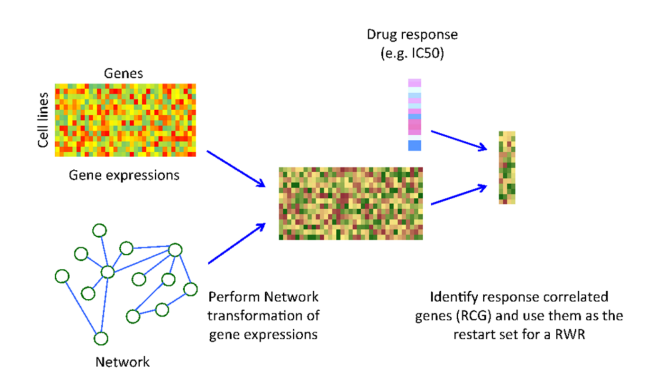


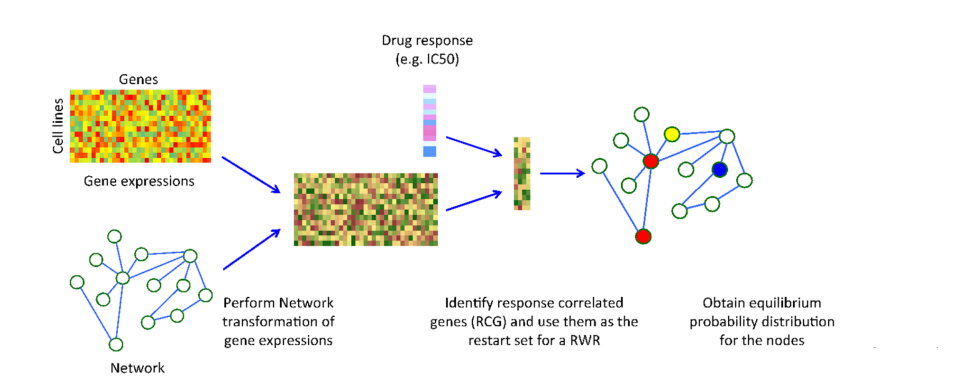
Figure from Rosvall and Bergstrom. "Maps of random walks on complex networks reveal community structure." *Proceedings of the national academy of sciences* 105.4 (2008).

Step 2: Find genes most correlated with drug response (RCG set)



Step 2: Find genes most correlated with drug response (RCG set)

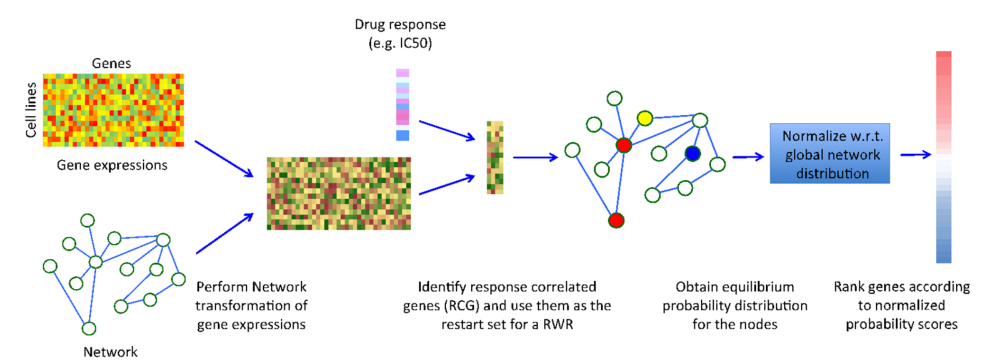
Step 3: Score genes based on their relevance to the RCG set



Step 2: Find genes most correlated with drug response (RCG set)

Step 3: Score genes based on their relevance to the RCG set

Step 4: Remove network bias by normalizing scores w.r.t. scores corresponding to global network topology



- Human lymphoblastoid cell lines (LCL)
 - Gene expression (~17K genes of ~300 cell lines)
 - Drug response of 24 cytotoxic treatments



- Publicly available dataset from GDSC
 - Gene expression (~13K genes of ~600 cell lines from 13 tissues)
 - Drug response of 139 cytotoxic treatments



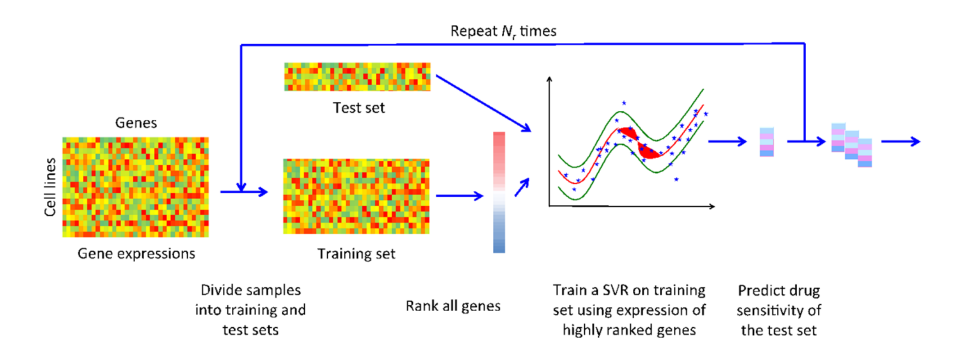
- Publicly available prior knowledge
 - Network of gene interactions (PPI and genetic interactions) from STRING (~1.5M edges, ~15.5K nodes)



Validation Using Drug Response Prediction

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 Genes ranked highly using a good prioritization method are good predictors of drug sensitivity

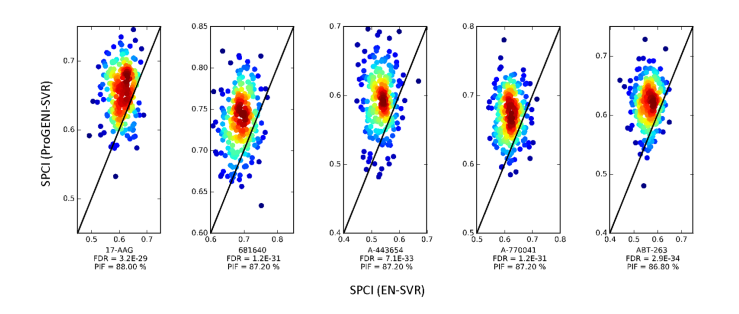


Validation Using Drug Response Prediction

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| LCL Dataset | Pearson | Elastic Net |
|---|---------|-------------|
| Num. Drugs (out of 24) ProGENI > Baseline | 14 | 20 |
| FDR (Wilcoxon signed-rank test) | 6.5 E-3 | 9.6 E-5 |

| GDSC Dataset | Pearson | Elastic Net |
|--|---------|-------------|
| Num. Drugs (out of 139) ProGENI > Baseline | 66 | 110 |
| FDR (Wilcoxon signed-rank test) | 9.1 E-4 | 4.0 E-21 |



Functional Validation

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We validated role of 33 (out of 45) genes (73%) for three drugs.

| Gene Symbol | Rank (ProGENI) | Rank (Pearson) | Absolute value of Pearson correlation coefficient | Evidence | |
|------------------------------------|-------------------------------------|------------------------------------|--|---------------------|--|
| ATF1 | 1 | 1 | 0.2000 | Direct (this study) | |
| MIS12 | 2 | 4 | 0.1887 | Direct (this study) | |
| OSBPL2 | 5 | 6 | 0.1865 | Direct (this study) | |
| CSNK2A1 | 7 | 1587 | 0.0752 | Direct (literature) | |
| PSIP1 (LEDGF) | 8 | 46 | 0.1537 | Direct (literature) | |
| CAMK2A | 9 | 6991 | 0.0157 | Direct (literature) | |
| CSNK2A2 | 10 | 4870 | 0.0347 | Direct (literature) | |
| GOSR1 | 11 | 6867 | 0.0167 | Direct (this study) | |
| MAPK8 | 13 | 7574 | 0.0112 | Direct (literature) | |
| SPI1 | 14 | 6287 | 0.0217 | Direct (literature) | RRS1 |
| CREB1 | 15 | 665 | 0.1000 | Direct (literature) | |
| NOC3L | 3 | 3 | 0.1893 | Not found | HEATR |
| IL27RA | 4 | 2 | 0.1911 | Not found | NOC3L POUR (NOL3) |
| MGEA5 | 6 | 7 | 0.1814 | Not found | MIS12 NIFK MGEA5 OGT |
| WAPAL | 12 | 8 | 0.1805 | Not found | NOL6 GTPBP4 LEO1 SUI/O2 CASP2 |
| | BT549 | | | | SUGTI SIN3A ZNF45 |
| - | | p-value < 0.0001 | | | PXN PRICO. |
| 0.8- 0.6- 0.4- 0.2- Negsi | | Negsi siOSBPL2 | DONOMONIO HOLINO | L27R/ | CSNK2AZ USC SPII VWHAO EDFI MAPK8 SPC PDPK1 ATF1 PAK1 CREM CRESS CRESS CRESS CAN GOSR1 CREM CRESS CRESS CRESS CAN CRESS CRESS CRESS CAN CRESS CRESS CRESS CRESS CRESS CRESS CRESS CRESS CAN CRESS C |
| 0.0 -2 - | 0.0 1 0 1 -2 ncentration (μΜ) | siOSBPL2 -1 0 concentration (µM) | | EBB | FNBP1 RPS6KA1 PRKACA RPS6KA1 |

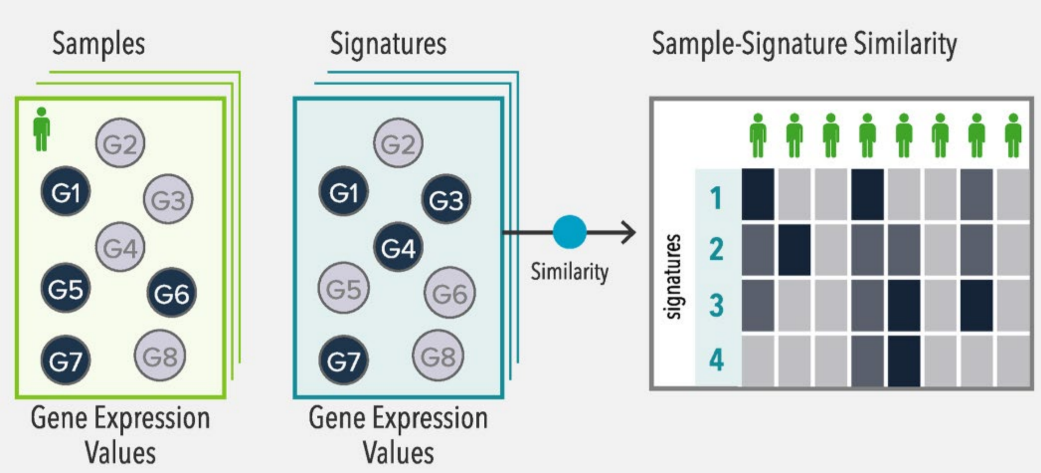




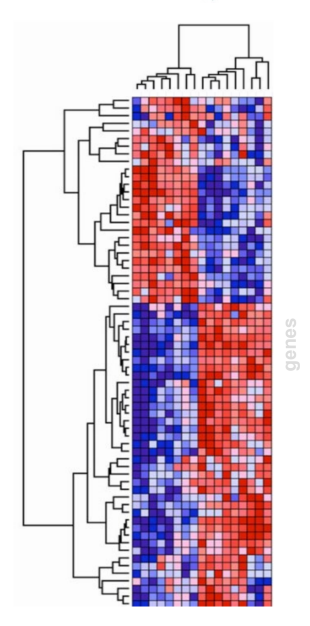




Gene Expression Signatures



- Massive Transcriptomic Profiling Projects
 - TCGA and ICGC
 - GTEX and CCLE
 - LINCS
- Definitions
 - Projects produce expression vectors for samples (e.g. gene expression levels)
 - Scoring the difference in expression between samples of two (or more) conditions produces differential expression vectors
- Signature (of a biological state):
 - Gene Set differentially, characteristically expressed genes in that state relative to some reference (control or population)
 - Differential Expression Vector the differential expression scores for the subset of genes in the same comparison

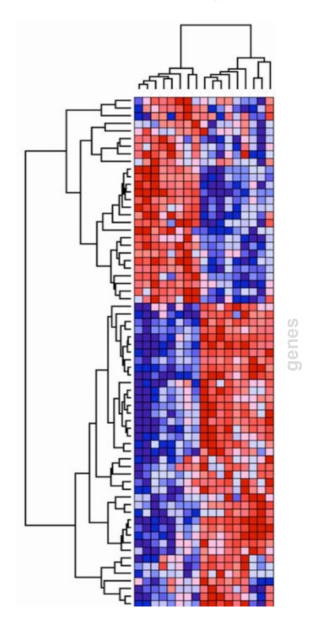


samples

Gene Expression Signatures

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- Example Comparisons
 - Mutated vs Wild-Type
 - Metastatic vs Primary
 - Tumor vs Normal
 - Perturbagens
 - Drug Treatment vs Placebo
 - Environmental Stimuli vs Control
- Gene Signatures provide a uniquely characteristic pattern of gene expression that is tied to its studied biological or medical phenomenon
 - Enable researchers to relate samples and other phenomenon by finding the similarity to the gene signatures
 - Focus understanding on underlying mechanism for phenomenon to a subset of gene behaviors



samples

Public Resources for Gene Signatures

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- There are many public resources for acquiring gene expression signatures
 - Extracting signatures yourself





Libraries of Curated Signatures





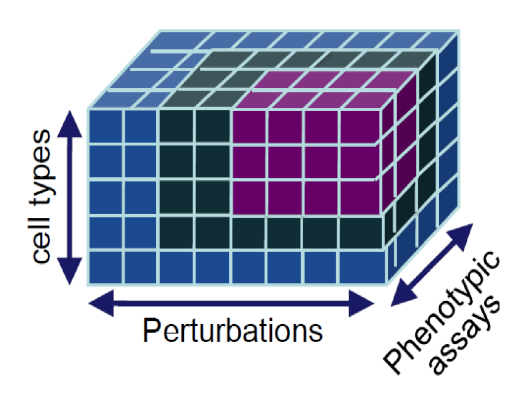
 Lab will use signatures from the Library of Integrated Network-Based Cellular Signatures (LINCS)



The LINCS DataCube of Signatures

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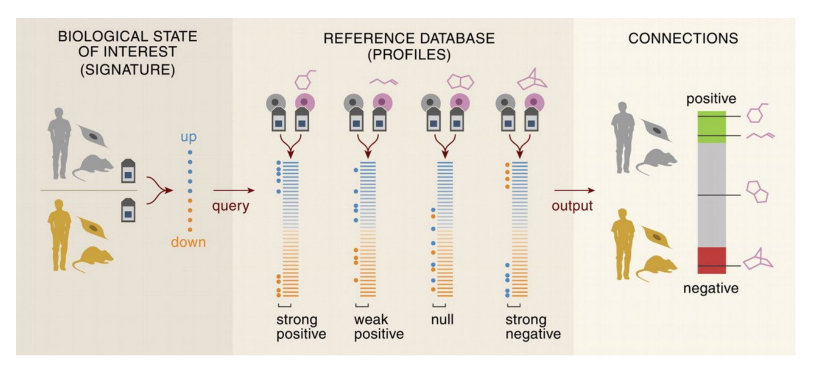
- Gathering a data cube of gene signatures
- Using many different:
 - Cell Types
 - Dozens of cell lines
 - Induced pluripotent stem cells
 - Primary Cells
 - Perturbagens
 - Small molecules / Drugs
 - CRISPR overexpression and
 - shRNA knockdown
 - Microenvironments
 - Ligands
 - Experimental Assays
 - Gene expression: microarray, RNA-seq, L1000
 - Protein expression: RPPA, P100 mass spectrometry
 - Morphological and Proliferation: biochemical and imaging assays



Given a query signature and a library of reference signatures, how do you find the similar signatures?

A gene signature-based approach identifies mTOR as a regulator

Rosenbluth JM1, Mays DJ, Pino MF, Tang LJ, Pietenpol JA.



Types of Similarity Comparisons

Gene Set & Differential Expression Vector Differential Expression Vector & Differential Expression Vector Gene Set & Gene Set

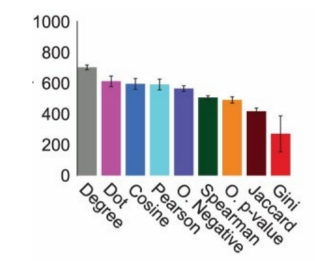
Standard Similarity Measures



 When both signatures are represented as differential expression vectors:

| | Correlation | Formula (x, y) | Description | Study |
|---|-------------|--|---|--------------------|
| 1 | Pearson | $\frac{\sum_{i} (x_{i} - \overline{x})(y_{i} - \overline{y})}{\sqrt{\sum_{i} (x_{i} - \overline{x})^{2}} \sqrt{\sum_{i} (y_{i} - \overline{y})^{2}}}$ | Linear similarity measure that uses mean-centering and normalization of the profiles. | Pearson 1920 [29] |
| 2 | Cosine | $\frac{\sum_{t} x_{t} y_{t}}{\sqrt{\sum_{t} x_{t}^{2}} \sqrt{\sum_{t} p_{t}^{2}}}$ | Linear similarity measure that uses normalization of the profiles. | |
| 3 | Spearman | $\frac{\sum_{t} (r_t - r)(s_t - s)}{\sqrt{\sum_{t} (r_t - r)^2} \sqrt{\sum_{t} (s_t - s)^2}} $ where r_t is rank of x_t in \mathbf{x}_t s_t is rank of y_t in \mathbf{y}_t | Spearman correlation is Pearson correlation on the ranks of elements in the profile. | Spearman 1904 [34] |

 In one analysis, they did not observe a large performance difference between the possible measures



PLoS One. 2013 Jul 10;8(7):e68664. doi: 10.1371/journal.pone.0068664. Print 2013.

Comparison of profile similarity measures for genetic interaction networks.

Deshpande R1, Vandersluis B, Myers CL

Gene Set Enrichment Analysis

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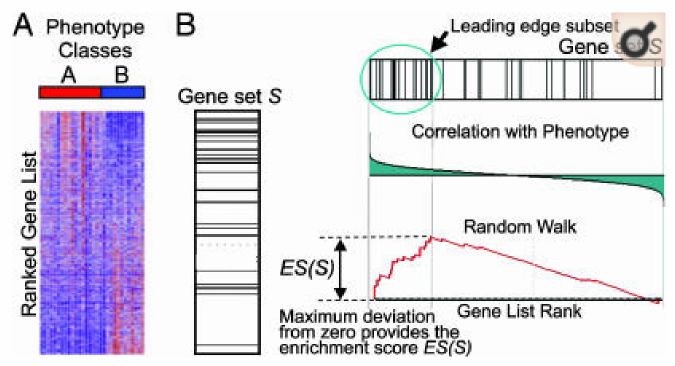
- When sample signature is vector and library signature is gene set
 - GSEA http://software.broadinstitute.org/gsea/index.jsp

Proc Natl Acad Sci U S A. 2005 Oct 25:102(43):15545-50. Epub 2005 Sep 3

Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles.

Subramanian A¹, Tamayo P, Mootha VK, Mukherjee S, Ebert BL, Gillette MA, Paulovich A, Pomeroy SL Golub TR, Lander ES, Mesirov JP.





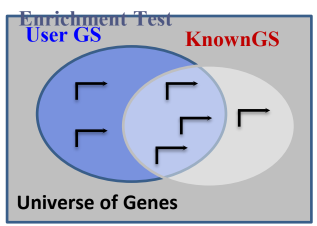
- Modification of the Kolmogorov-Smirnov Statistic
 - Calculate the enrichment score (ES) that represents the amount the genes in the gene set are over-represented in the top or the bottom of the signature vector
 - Estimate statistical significance of the ES by permuting the mappings between the data
 - Adjust for multiple hypothesis testing when analyzing a large number of gene sets

- For use when both signatures are gene sets
 - Also known as Gene Set Characterization
- One-sided exact Fisher / Hypergeometric distribution tests
 - Covered by Saurabh this morning
- Available through tools like:
 - DAVID https://david.ncifcrf.gov/
 - Enrichr http://amp.pharm.mssm.edu/Enrichr/
 - Metascape http://metascape.org/gp/index.html





Standard



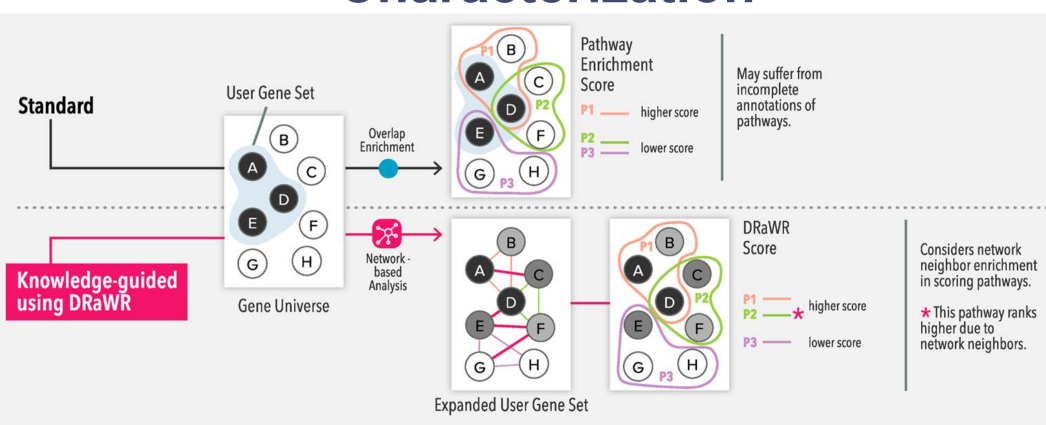








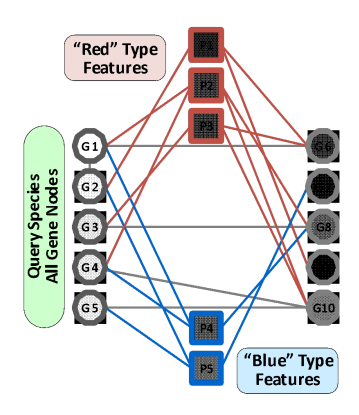
Network-Guided Gene Set Characterization



Idea for a Network-based Method

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- Use guilt-by-association principles to find out which annotations are well connected to the query genes in a heterogeneous network.
- These well connected annotations should be specific to the query genes, and not simply hub nodes in the network.
- Developed Discriminative Random Walks with Restart (DRaWR)



Bioinformatics. 2016 Jul 15;32(14):2167-75. doi: 10.1093/bioinformatics/btw151. Epub 2016 Mar 19.

Characterizing gene sets using discriminative random walks with restart on heterogeneous biological networks.

Blatti C1, Sinha S2.

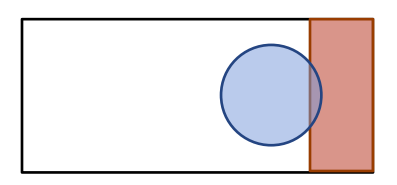
Value of Network-Guided Analysis

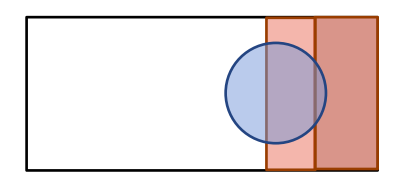
Take advantage of gene neighbors

User Set

Apoptosis Genes

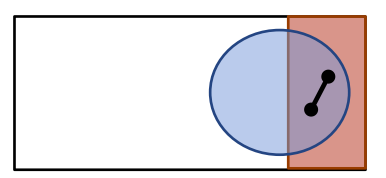
Genes That Bind To Apoptosis Genes



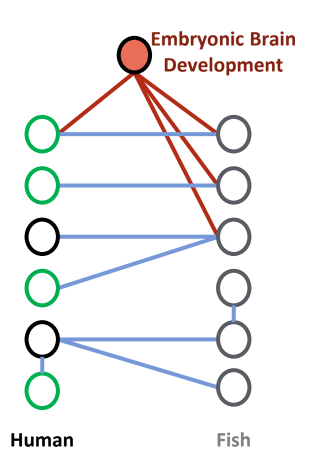


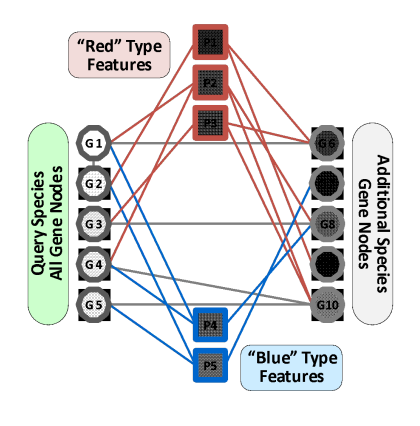
Incorporate dependencies from separate knowledge in

analysis



Extension to poorly annotated domains





Integrating multiple data types

Network-based DRaWR Method

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- DRaWR using random walks on a network
 - Construct a heterogeneous network of interest

"Red" Type **Features** All Gene Nodes **Query Species** Gene Nodes "Blue" Type **Features**

Heterogeneous Edge Types

> type_A type_B type_C

Bioinformatics. 2016 Jul 15;32(14):2167-75. doi: 10.1093/bioinformatics/btw151. Epub 2016 Mar 19.

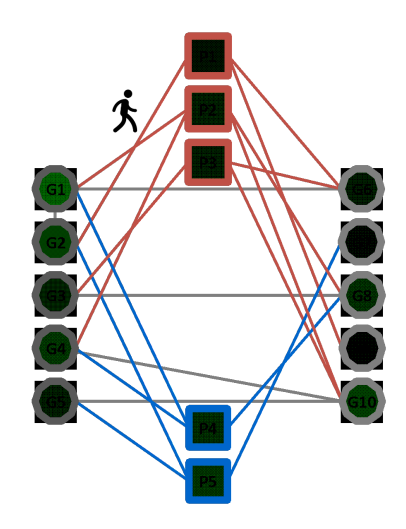
Characterizing gene sets using discriminative random walks with restart on heterogeneous biological networks.

Blatti C¹, Sinha S².

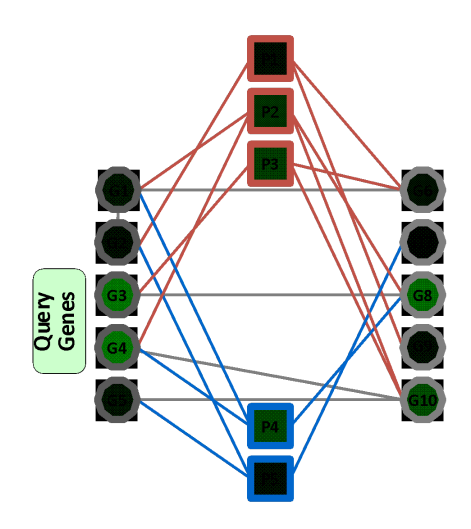
Network Methods for GSC

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- DRaWR using random walks on a network
 - Construct a network of interest
 - Find stationary distribution on network



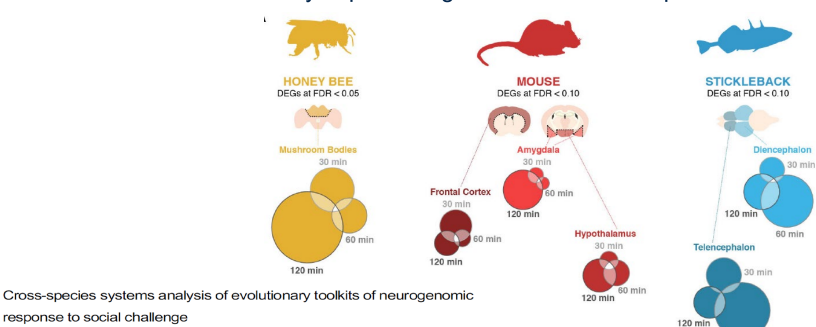
- DRaWR using random walks on a network
 - Construct a network of interest
 - Find stationary distribution on network
 - Find gene set specific distribution
 - Return annotation nodes that are especially related to the query



Social Aggression Study Application

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- Idea: Evolutionary "toolkits" genes and modules with lineage-specific variations but deep conservation of function
- Questions: Are there toolkits that underlie social behaviors
 - Such as aggressive response to territorial intrusions?
- Study: gather brain transcriptomic responses to social challenge from three social species – honey bees, mice, and stickleback fish
 - With and without exposure to intraspecies intruder
 - From different brain regions and/or durations after event
- Results: sets of differentially expressed genes across three species

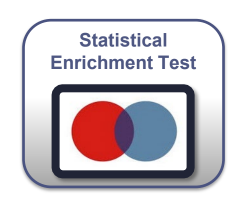


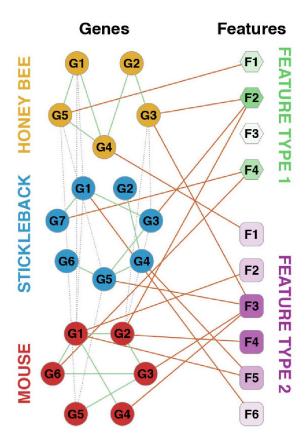


Michael C. Saul¹, Charles Blatti^{1,2}, Wei Yang^{1,2}, Syed Abbas Bukhari^{1,3}, Hagai Y. Shpigler^{1,4}, Joseph M. Troy^{1,3}, Christopher H. Seward^{1,5}, Laura Sloofman^{1,6}, Sriram Chandrasekaran⁷, Alison M. Bell^{1,3,8,9}, Lisa Stubbs^{1,3,5,9}, Gene E. Robinson^{1,9,10}, Sihai Dave Zhao^{1,11,*}, and Saurabh Sinha^{1,2,10,*}

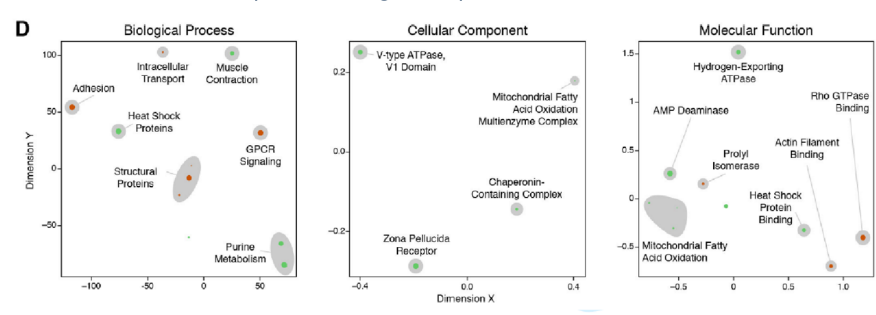
response to social challenge

- Would like to find Gene Ontology annotations that:
 - Relate to DE gene sets of all three species
 - However, Gene Ontology annotation quality varies greatly in three species
 - Or relate to DE genes sets of the Mouse
 - However, the corresponding sets from the other species might have greatly different function
- Solution:
 - Integrate Orthology and Gene Ontology information in a three species network
 - Find Gene Ontology terms that are strongly connected to the DE gene sets of all three species simultaneously





Annotations of two (red and green) conserved Gene Modules



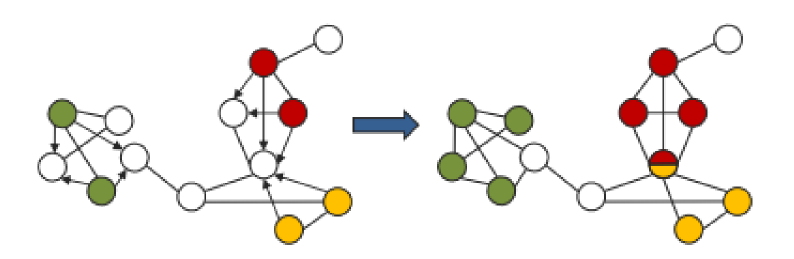
Specific results for red module

| : 14 | | | #Annotated | | | DRaWR GO Term Rank | | | | | Fisher Pvalue | | | |
|--------|------------|---|------------|-----|----|--------------------|-------|------|------|-------|---------------|-------|-------|-------|
| Branch | GO ID | GO Description | НВ | MM | SB | Combo | НВ | MM | SB | Max | НВ | MM | SB | Min |
| BP | GO:0032366 | intracellular sterol transport | | 2 | | 0.3% | 1.6% | 0.1% | 0.4% | 1.6% | | 0.040 | | 0.040 |
| BP | GO:0071704 | organic substance metabolic process | 3 | 5 | 4 | 2.3% | 2.2% | 0.3% | 0.4% | 2.3% | 0.134 | 0.040 | | 0.040 |
| BP | GO:0016043 | cellular component organization | 4 | 9 | 12 | 2.3% | 2.2% | 2.9% | 0.8% | 2.9% | 0.175 | 0.151 | 0.002 | 0.002 |
| BP | GO:0007160 | cell-matrix adhesion | 5 | 74 | 16 | 2.5% | 0.4% | 3.5% | 1.8% | 3.5% | 0.002 | 0.001 | | 0.001 |
| MF | GO:0017048 | Rho GTPase binding | 6 | 30 | 13 | 3.1% | 2.0% | 3.9% | 0.8% | 3.9% | 0.020 | 0.024 | 0.002 | 0.002 |
| BP | GO:0038032 | termination of G-protein coupled receptor | 11 | 1 | 44 | 1.6% | 6.8% | 1.4% | 0.3% | 6.8% | | | 0.000 | 0.000 |
| MF | GO:0051015 | actin filament binding | 17 | 114 | 9 | 7.6% | 4.0% | 8.0% | 8.3% | 8.3% | 0.013 | 0.125 | | 0.013 |
| MF | GO:0003755 | peptidyl-prolyl cis-trans isomerase activit | 22 | 42 | 17 | 4.7% | 2.1% | 9.1% | 1.3% | 9.1% | 0.031 | | 0.108 | 0.031 |
| BP | GO:0031032 | actomyosin structure organization | 2 | 18 | | 1.8% | 0.4% | 2.7% | 9.6% | 9.6% | 0.047 | | | 0.047 |
| MF | GO:0003779 | actin binding | 48 | 284 | 78 | 8.7% | 10.0% | 6.9% | 8.3% | 10.0% | 0.086 | 0.021 | 0.001 | 0.001 |

Gene Ranking / Function Prediction

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- Given:
 - Novel gene set(s) generated by a genomic researcher
- Task:
 - Rank genes for the strength of their relationship to the user's gene set(s)...
 - ... in order to assess the coherence of the genes in the experimental gene set or identify putative related genes



GeneMANIA Approach

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- GeneMANIA stands for
 - Multiple Association Network Integration Algorithm
- Main Idea
 - Given a gene set with a known functions
 - And several gene-gene interaction affinity networks
 - Find genes that relate to the functional set through the edges of the given networks
- Approach
 - Find out how well each network predicts the membership of the given set
 - A linear regression-based algorithm that calculates a single composite functional association network from multiple data sources
 - Do label propagation guilt-by-association algorithm on the composite functional association network

Genome Biol. 2008;9 Suppl 1:S4. doi: 10.1186/gb-2008-9-s1-s4. Epub 2008 Jun 27.

GeneMANIA: a real-time multiple association network integration algorithm for predicting gene function.

Mostafavi S1, Ray D, Warde-Farley D, Grouios C, Morris Q.

GeneMANIA Performance



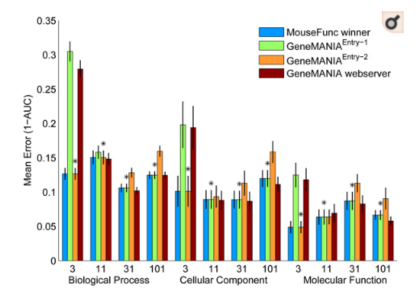
 Participated in grand challenge for this function prediction task on Mouse genes

A critical assessment of Mus musculus gene function prediction using integrated genomic evidence.

Peña-Castillo L1, Tasan M, Myers CL, Lee H, Joshi T, Zhang C, Guan Y, Leone M, Pagnani Krumpelman C, Tian W, Obozinski G, Qi Y, Mostafavi S, Lin GN, Berriz GF, Gibbons FD, J, Grant C, Barutcuoglu Z, Hill DP, Warde-Farley D, Grouios C, Ray D, Blake JA, Deng M, Morris Q, Klein-Seetharaman J, Bar-Joseph Z, Chen T, Sun F. Troyanskaya OG, Marcotte EM

Did extraordinary well in the competition and has

improve method since then



 Has easy to use webserver for running functional prediction with small genesets









In this Lecture and the Lab

- Biological Knowledge Networks
 - KnowEnG Platform
- Network-Guided Sample Clustering
 - Network Based Stratification, COCA
- Network-Guided Gene Prioritization
 - ProGENI
- Gene Signatures and Similarity Methods
 - LINCS, GSEA, Enrichr, DAVID
- Network-based Gene Set Characterization
 - DRaWR
- Network-based Function Prediction
 - GeneMANIA



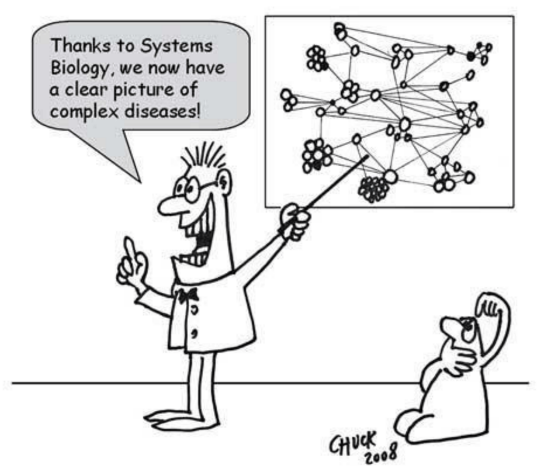








Thank you, Any Questions?





KnowEnG Resources

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Also Check Out:

- Network Preparation for uploading your custom network to the platform for analysis
- Signature Analysis for mapping samples to signatures by correlation of omics profiles

Tutorials:

- Quickstarts: https://knoweng.org/quick-start/
- YouTube: https://www.youtube.com/channel/UCjyllolCaZIGtZC20XLBOyg

Resources:

- Data Preparation Guide: https://github.com/KnowEnG/quickstart-demos/blob/master/pipeline readmes/README-DataPrep.md
- Knowledge Network Contents:
 - Summary: https://knoweng.org/kn-data-references/
 - Download: https://github.com/KnowEnG/KN Fetcher/blob/master/Contents.md

Research

- Knowledge-guided analysis of omics Data (KnowEng cloud platform paper): https://journals.plos.org/plosbiology/article?id=10.1371/journal.pbio.3000583
- TCGA Analysis Walkthrough: https://github.com/KnowEnG/quickstart-demos/tree/master/publication_data/blatti_et_al_2019

Source Code:

- Docker Images: https://hub.docker.com/u/knowengdev/
- Github Repos: https://knoweng.github.io/

Other Cloud Platforms

- https://cgc.sbgenomics.com/public/apps#q?search=knoweng
- Contact Us with Questions and Feedback: knoweng-support@illinois.edu

Using A Permanent KnowEnG Account

KNOWENS BIG DATA TO KNOWLEDGE CENTER OF EXCELLENCE

- For permanent account:
 - Go to https://knoweng.org/analyze/
 Click on "Create an account"
 - Follow the instructions

PLATFORM IS NOW AVAILABLE!

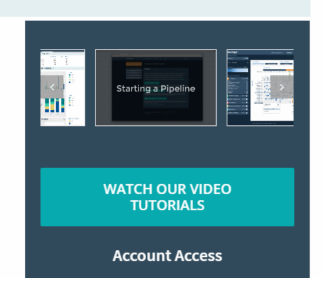
LOGIN OR REGISTER

Welcome to the KnowEnG Platform!

KnowEnG enables knowledge-guided machine learning and graph mining analysis on genomic datasets using scalable cloud computation and exploration of results with interactive visualizations.

KNOWLEDGE-GUIDED PIPELINES

Researchers can upload their data in form of a spreadsheet and choose from several analysis



 Lasso: learns a linear model from the training data using only a few features (sparse linear model)

$$\hat{\boldsymbol{\beta}} = \arg\min_{\boldsymbol{\beta}} (||\mathbf{y} - \mathbf{X}\boldsymbol{\beta}||^2 + \lambda_1 ||\boldsymbol{\beta}||_1)$$

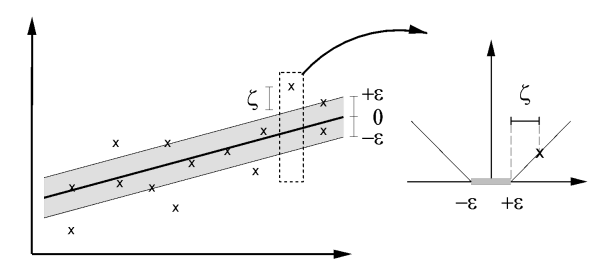
 Elastic Net: learns a linear model from the training data by linearly combining ridge and Lasso regression regularization terms (a generalization of both Lasso and ridge regression)

$$\hat{\boldsymbol{\beta}} = \arg\min_{\boldsymbol{\beta}} \left(||\mathbf{y} - \mathbf{X}\boldsymbol{\beta}||^2 + \lambda_2 ||\boldsymbol{\beta}||_2 + \lambda_1 ||\boldsymbol{\beta}||_1 \right)$$

Regression algorithms

Kernel-SVR:

 Linear SVR learns a linear model such that it has at most ε-deviation from the response values and is as flat as possible



(Smola and Schölkopf, 1998)

 Kernel-SVR generalizes the idea to nonlinear models by mapping the features to a high-dimensional kernel space

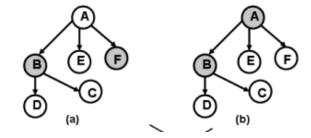
Other Network Based Characterization Methods

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Bioinformatics, 2009 Jan 1;25(1):75-82. doi: 10.1093/bioinformatics/btn577. Epub 2008 Nov 5.

A novel signaling pathway impact analysis.

Tarca AL1, Draghici S, Khatri P, Hassan SS, Mittal P, Kim JS, Kim CJ, Kusanovic JP, Romero R.



SPIA Idea:

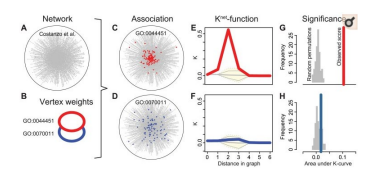
- Combine with standard enrichment p-value that asks about the significance of the number of perturbed genes in the pathway
- Perturbagen p-value, which asks if the amount of total accumulated perturbation after one network propagation step is significant when considering the value it takes with random controls

PLoS Comput Biol. 2014 Sep 11;10(9):e1003808. doi: 10.1371/journal.pcbi.1003808. eCollection 2014 Sep.

SANTA: quantifying the functional content of molecular networks.

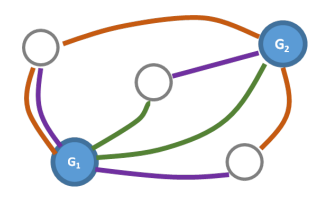
Cornish AJ1, Markowetz F2.

, Shortest Path Length criteria



Incorporating Meta-Paths

 DRaWR random walks on heterogeneous networks make no consideration / memory of the edge *types* they have followed



Paths from G1 -> G2: type_A type_A - type_B type_C- type_C type_B - type_C (x2)

meta-path:

a path defined by sequence of edges types between two nodes

 Explore if similarity in a gene set can best be described by particular types of meta-paths amongst its genes.

Initial Study:

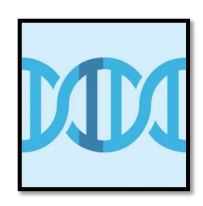
53 MSigDB DE gene sets from separate cancer studies



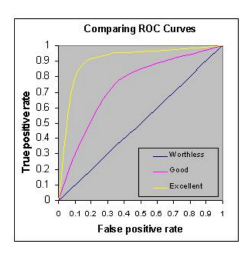
 If we hide a subset of genes disrupted by the development of cancer, what types of networks are best suited to recover them?

Evaluation:

- Partition 75% of DE genes for training, 25% for testing
- Use DRaWR on KnowNet subnetworks and training data to rank genes
- Report average AUCs of ranking using test genes as truth







Networks Under Consideration

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- Gene-Gene Edge Types
 - H: Homology
 - CoEx: Co-Expression
 - TM: Text Mining
 - Exp: Experimental Interaction
- Gene-Property Edge Types
 - PD: Protein Domains
 - GO: Gene Ontology

- Number of Species
 - Human: only
 - 2sp: Human and Mouse
- Specificity of the edges
 - Specific: high confidence edges
 - Loose: all edges of that types
- Combinations of Edge Types
 - 1ty: One primary type
 - 2ty: Primary type + homology
 - Many: 3+ edge types

- Gene Ontology annotations and Text Mining relations are the best edge types for recovering cancer set DE genes
- Networks with all edges (Loose) are better at recovering gene than networks with only high confidence edges
- Protein Domain annotations are poor predictors for cancer DE genes, but great for embryonic development

| Species | ▼ NEdgeT ■ | EdgeType | 1 | idgeThresh | Ţ | avg 🔻 | min 🔻 | max 🔻 |
|---------|------------|-----------------|---|------------|---|-------|-------|-------|
| Human | many | GO.TM.H | i | .oose | П | 0,723 | 0,610 | 0.847 |
| Human | many | All | | .oose | г | 0.722 | 0.614 | 0.863 |
| 2sp | many | GO.TM.H | | .oose | | 0.721 | 0.610 | 0.843 |
| 2sp | many | All | | .oose | г | 0.714 | 0.606 | 0.852 |
| 2sp | 2ty | GO.H | | .oose | | 0.706 | 0.578 | 0.862 |
| 2sp | 2ty | TM.H | | .oose | г | 0.701 | 0.567 | 0.813 |
| Human | many | All | | Specific | | 0.701 | 0.590 | 0.838 |
| Human | many | GO.TM.H | | Specific | г | 0.701 | 0.584 | 0.855 |
| Human | many | GO.TM | | .oose | | 0.701 | 0.545 | 0.870 |
| 2sp | many | GO.TM.H | | Specific | Г | 0.699 | 0.579 | 0.848 |
| 2sp | many | All | | Specific | | 0.698 | 0.594 | 0.824 |
| 2sp | many | GO.TM | | oose | Г | 0.695 | 0.537 | 0.863 |
| 2sp | 2ty | GO.H | | Specific | | 0.694 | 0.555 | 0.853 |
| Human | 1ty | Text Mining | | .oose | Г | 0.693 | 0.544 | 0.838 |
| Human | 1ty | Gene Ontolog | , | .oose | | 0.690 | 0.541 | 0.851 |
| 2sp | 1ty | Gene Ontolog | | .oose | Г | 0.689 | 0.538 | 0.848 |
| Human | many | GO.TM | | Specific | | 0.675 | 0.539 | 0.831 |
| 2sp | 2ty | TM.H | | Specific | Г | 0.673 | 0.563 | 0.797 |
| 2sp | many | GO.TM | | Specific | | 0.671 | 0.541 | 0.823 |
| 2sp | 2ty | PPI.H | | .oose | Г | 0.668 | 0.557 | 0.800 |
| 2sp | 1ty | Gene Ontolog | , | Specific | | 0.666 | 0.515 | 0.844 |
| Human | 1ty | Gene Ontolog | ı | Specific | Г | 0.664 | 0.534 | 0.842 |
| 2sp | 2ty | CoE.H | | .oose | | 0.663 | 0.508 | 0.827 |
| 2sp | 2ty | Exp.H | | Specific | | 0.656 | 0.549 | 0.769 |
| Human | 1ty | Text Mining | | Specific | | 0.656 | 0.555 | 0.812 |
| 2sp | 2ty | Exp.H | | .oose | | 0.647 | 0.533 | 0.763 |
| 2sp | 2ty | PPI.H | | Specific | | 0.644 | 0.515 | 0.746 |
| Human | 1ty | Co-expression | | oose | | 0.629 | 0.498 | 0.840 |
| Human | 1ty | Experimental | | Specific | | 0.604 | 0.455 | 0.756 |
| Human | 1ty | Co-expression | | Specific | | 0.601 | 0.353 | 0.875 |
| Human | 1ty | Prot-Prot Inter | | oose | | 0.598 | 0.475 | 0.730 |
| zsp | zty | COE.H | | ресітіс | | U.598 | U.4// | U./25 |
| 2sp | 2ty | PD.H | | .oose | | 0.592 | 0.481 | 0.701 |
| Human | 1ty | Experimental | i | .oose | | 0.589 | 0.424 | 0.778 |