

Visualizing and Refining Connectivity Map Query Results

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Abstract

The Connectivity Map (CMap) is a database of gene expression signatures obtained from experiments in which cultured human cells are treated with pharmacologic and genomic perturbagens. A typical use case of this database is for a researcher to query with a signature of a cell state of interest and use the matching perturbagens to develop a functional hypothesis for follow-up. Current pattern matching algorithms that perform CMap queries suffer from a universal weakness – the enormous size and richness of signatures in CMap means that a query typically generates hundreds of strong connections. These connections are hard to distinguish, thereby making prioritization difficult. I hypothesize that one mode of prioritization is to highlight query results that are highly interconnected amongst themselves over singletons. The goal of this work is to provide a web-based tool for implementing an interconnectivity-based method of query result refinement and for visualizing CMap query results in a graph layout.

1 The Research Problem

The CMap database is a compendium of gene expression signatures resulting from the treatment of cultured human cells with small molecule compounds (CP), short hairpin RNAs (shRNA), or over-expression constructs (OE). The utility of the CMap database is that of a gene expression search engine. Users are able to pose questions about relationships between cellular states and formulate hypotheses based on similarities or differences in the states’ gene expression signatures.

Hypotheses are generated by posing search queries into the database and examining the query results. A CMap query is a focused question in which a user inputs a gene expression signature, called the query, and computes the similarity, or connectivity, between his/her query and other signatures in

the database. Positive connectivity indicates that two signatures' expression changes are similar and vice versa. Researchers can use CMap to find connections between signatures within or external to the database. Hypotheses may be in the form of "the shRNA knockdown of gene X connects to shRNA knockdown signatures of pathway Y members, so X is probably a member of Y." Or perhaps "the signature of compound Z connects to the knockdown signature of gene X, so perhaps X is the target of Z".

Lamb et al demonstrated a more directly therapeutically relevant use of the original incarnation of CMap when they discovered that the signature of sirolimus connected strongly to a signature of dexamethasone sensitivity. Dexamethasone is a treatment for acute lymphoblastic leukemia (ALL), but many patients eventually become resistant (Tissing, Meijerink, den Boer, & Pieters, 2003). The CMap connection between sirolimus and dexamethasone sensitivity suggested that sirolimus might be effective in reversing resistance in ALL patients who had become resistant to dexamethasone. A follow-up experiment confirmed that sirolimus conferred dexamethasone sensitivity to CEM-c1 cells, a previously dexamethasone-resistant cell line (Lamb et al., 2006).

The CMap database contains over 400,000 signatures spanning over 70 cell types. Because of the large size of the CMap database, interpreting and prioritizing query results has become a difficult task. For example, accepting only the top one percent of connections yields nearly 4,000 signatures. Follow-up on such a large number of primary hits is nearly impossible in

most cases. I propose that within a set of initial query results, there will frequently exist a set or sets of signatures that are more tightly interconnected with themselves than with other signatures. These interconnected sets are more likely to be indicative of robust biological signal and should therefore be prioritized over other singleton connections. The goal of this work is to build a web-based tool to implement an algorithm to identify subsets of high interconnectivity with lists of initial query results and to visualize the relationships between these subsets in a graph layout. I propose that this tool will be useful in refining initial CMap query results into smaller, more actionable lists of connections that can be further investigated in secondary assays.

2 Key Terms

1. gene expression profiling:
2. gene set enrichment analysis (GSEA):
3. Kolmogorov-Smirnov (KS) statistic:
4. graphical model:
5. CMap connection:

3 Background

3.1 Gene Expression Profiling

Gene expression profiling is the simultaneous measure of the RNA transcript levels of many genes within a cell or group of cells. These measurements can help to provide insight into the cellular state or states of the cells in question. For example, if many cell-cycle genes are observed to be active, it could suggest that the cells are actively dividing. Conversely, if many apoptotic genes are active, the cells might be dying. Frequently, the goal of gene expression profiling is to identify genes that are differentially regulated between one or more sets of conditions. For example, one might measure expression in cells that have and have not been treated with a drug of interest, and then compare the resulting expression profiles to identify genes that are substantially up- or down-regulated in the treated cells relative to the untreated. Current technology, such as the microarray, allow for many such gene expression experiments to be run in parallel, enabling the comparative analysis of hundreds or thousands of expression profiles corresponding to an equal number of experimental conditions. Similarly, expression profiling can be used to identify genes differentially regulated between disease and normal states. van't Veer et al used gene expression profiling to identify a set of genes, that were predictive of breast cancer metastasis (van 't Veer et al., 2002). Because of its ability to identify such signatures, gene expression profiling is a powerful and often-used tool in contemporary biology.

3.2 Gene Set Enrichment Analysis (GSEA)

Gene Set Enrichment Analysis (GSEA) is an analytical approach designed to extract biological insight from gene expression data (Subramanian et al., 2005). It leverages groups of genes, called gene sets, that share some biological commonality (i.e. members of a cellular signaling pathway) and computes their enrichment, or trend towards the top or bottom, of a ranked list of genes generated by comparing expression profiles across two experimental classes (i.e. tumor vs. normal). For example, one might define many sets of genes, each corresponding to a cellular pathway. One could then rank-order all genes by their differential expression when comparing profiles of tumor vs. normal samples. Lastly, one could compute the enrichment of each pathway in the rank-ordered list to attempt to identify pathways that might be active in the particular tumor in question.

Mechanically, GSEA computes a Kolmogorov-Smirnov (KS) statistic when comparing a given gene set to a given ranked list (Hollander & Wolfe, 1975). Effectively, this amounts to walking down the ranked list, increasing a running-sum statistic when we encounter a gene in the gene set and decreasing it when we encounter genes not in the gene set. The magnitude of the increment depends on the correlation of the gene with the phenotype. The enrichment score is the maximum deviation from zero encountered in the random walk (Subramanian et al., 2005). GSEA has been used extensively for identifying coherent sets of genes that are collectively modulated under certain disease and/or experimental conditions. In fact, a

GSEA software suite and an accompanying online database exist to facilitate comparisons between novel and curated gene sets. Cite MsigDB and GSEA.

3.3 The Connectivity Map

The Connectivity Map (CMap) is a database containing the gene expression signatures resulting from treating cultured cells with various chemical and genomic perturbations (Lamb et al., 2006). The purpose of CMap is to serve as a lookup table of functional annotation. These annotations might be derived by comparing signatures within the CMap database or by querying the database with externally generated signatures. The database itself can be thought of as a large matrix where each row is a gene and each column is an experiment in which a particular perturbagen was profiled under a given set of conditions (i.e. cell context, dose, treatment time, etc). The values in the matrix are differential expression measures generated by comparing the expression levels of the genes across perturbed and control states. Thus, each column of the matrix can be thought of as a given perturbagen’s expression signature.

3.3.1 Computing Connections in the Connectivity Map

A primary use of the CMap database is to compare the signatures of different perturbations and assess their similarity. Perturbagens that, when used to treat cultured cells, result in similar gene expression consequences will yield similar CMap signatures. Such signatures are said to be positively connected

in the CMap sense. Conversely, perturbagens that elicit inversely-related expression consequences are said to be negatively connected. For a given query signature Q and a reference signature R , the weighted connectivity score (WTCS) is computed by computing and integrating two KS statistics, one each for the n most up- and down-regulated genes in Q . The algorithm proceeds as follows:

1. Order Q
2. compute ES_{up} as the enrichment of the n most up-regulated genes in R
3. compute ES_{dn} as the enrichment of the n most down-regulated genes in R
4. compute WTCS as
 - (a) 0 if ES_{up} and ES_{dn} share the same sign
 - (b) $(|ES_{up}| + |ES_{dn}|/2)$, where the resulting WTCS is given the sign of ES_{up}

WTCS will be positive for signatures that are positively related and negative for those that are inversely related.

A common CMap use case is to select a given query signature Q from the database and compute its similarity to all other signatures. The remaining signatures can be ranked according to their connection strength with Q . The connections can be used to gain insight and form hypotheses about Q . For example, if Q is a signature of a novel compound and it connects strongly to signatures of compounds of a known pharmacological class, one

might hypothesize that the novel compound is also a member of this class. Similarly, if Q connects strongly to the knockdown signature of gene G, one might hypothesize that G is the novel compound's target. Q need to be a signature from the CMap database. For instance, it might be the signature of some disease and one might seek connections to genes whose modulation could be causing the disease or to compounds that could have therapeutic relevance.

3.4 Graphical Depictions of Biological Phenomena

In the worlds of computer science and mathematics, a graph is a means by which to represent a set of objects and relationships between them. It is frequently depicted as a set of nodes, where each node represents an object. Connections, where they exist, are represented by edges (1).

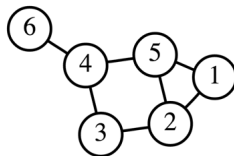


Figure 1: An example of a graph. The six nodes are connected by seven edges. All nodes other than node 6 have at least two edges.

Although they originated in the field of computer science, graphs have frequently been used as tools to model biological phenomena. Graphical models of protein interaction networks, gene expression networks, and other similar phenomena are commonplace. Friedman used graphical models to infer and visualize gene regulatory networks (Friedman, 2004). Lage et al used

graphical models to characterize existing and elucidate novel protein-protein interaction networks (Lage et al., 2007). Because of its widespread use and adoption, the graphical model is an appropriate, familiar, and effective means to depict connectivity between CMap signatures.

3.5 Approach for This Work

In this work, a graph will be used to depict the existence and strength of connections between gene expression signatures in the CMap database. The graph lends itself very well to this use case, as signatures will be represented as nodes and their pairwise connections by the edges. Edges will only be depicted between nodes with a non-zero connectivity score. Figure 2 depicts a mockup of what the final application might look like.

4 Methods

4.1 Computing Connections

Connections between CMap signatures will be computed using WTCS. In order to facilitate application performance, these connections will all be pre-computed and stored in a database. This way, the application can simply look up connectivity scores instead of computing them on-the-fly.

Users will interact with the application by inputting a list of signatures L that have resulted from running a CMap query. The application will sort

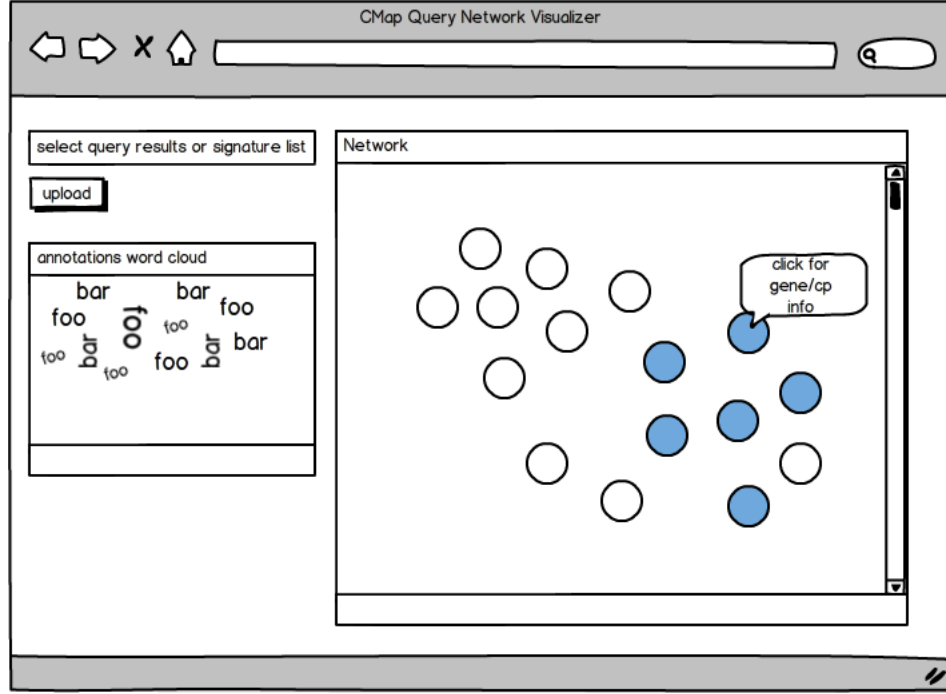


Figure 2: The App

L by the strength of connection to the user's query and then look up the connectivity scores between all pairwise combinations between the first N signatures within L, where N is a user-provided parameter. The application will then identify highly interconnected subsets of signatures by clustering the N signatures in pairwise connectivity space using k-means (Lloyd, 1982). Additionally, it will compute the optimal number of clusters using the GAP statistic and will report the within cluster sum of squares (WCSS) for each cluster.

The N signatures will be displayed as a graph, where each signature is a node. Node pairs with non-zero connectivity scores will have edges drawn between them and nodes will be colored according to their cluster membership. This will allow users to easily visualize cluster concordance and relationships between clusters. The application will allow users to mouse-over nodes and see additional information about the signatures, such as the experimental parameters under which they were generated. The user will also be able to select a cluster or other user-defined group of nodes and see a word cloud generated from their annotations. The word cloud might give insight to pathway or pharmacological class membership for the nodes in question.

Finally, the application will support export of the clusters or user-defined groups of nodes into a text file for download.

4.2 Software Components

4.2.1 Front End: HTML & D3.js

Hypertext markup language (HTML) is and has been the standard language for displaying information over the internet within web browsers. HTML5, the most recent revision of the HTML standard will be used as the framework of this application. HTML5 offers many useful features for application development and is supported by most modern web browsers (W3C, 2011).

To support user interaction, the graph and word cloud visualizations will be built using D3.js, a JavaScript library for data visualization. Created by

Mozilla in 1995, JavaScript is a programming language that is interpreted by most modern web browsers and allows developers to create interactive elements within web pages (Mozilla, 2013). D3, short for Data Driven Documents, is a JavaScript library written by Mike Bostock and designed specifically to enable rich and interactive data visualizations (Bostock, 2013). D3 is particularly well-suited for visualizing CMap connections because of its ability to easily integrate and bind data to on-screen elements. It has been used in many similar projects and is capable of generating the types of visualizations my application will require. Figure 3 and Figure 4 illustrate examples of D3 being used to generate interactive graph and word cloud visualizations.

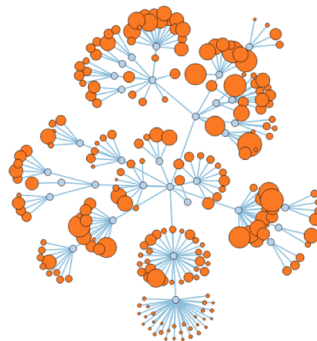
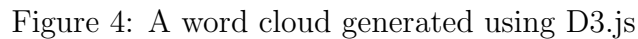


Figure 3: A graph generated using D3.js

4.2.2 Back End: Node.js & MongoDB

MongoDB is a database system developed by MongoDB Inc. Unlike traditional Structured Query Language (SQL) database systems that require rigid data storage schema, MongoDB's schema is very loose and fluid. Rather than



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be necessary when the application retrieves data from the database.

MongoDB Schema

MongoDB documents are simply JSON objects. These objects contain key-value pairs and the values can be accessed by providing the appropriate keys, or attributes. A CMap connection can be modeled as a very simple JSON object with the following attributes:

1. signature 1
2. signature 2
3. WTCS

An example CMap connection stored as a JSON document might look like this:

```
{  
    "signature_1": "signature_1_ID",  
    "signature_2": "signature_2_ID",  
    "wtcs": 0.65  
}
```

Providing the identifiers of signatures 1 and 2 are enough to uniquely identify this and any CMap connection. MongoDB allows for searching over the values of all documents that contain a given key or set of keys. I propose to store each CMap connection as a document in a single MongoDB collection. Based on the user's input set of query results (signature IDs), MongoDB will

easily be able to retrieve all connections between the query results by looking up all documents where the $signature_1$ and $signature_2$ fields are members of the input query result set.

5 Potential Challenges

6 Preliminary Timeline

2013-12-01 Receive approval of proposal

2013-12-15 Investigate and validate then notion of refining hit lists base on their interconnectivity

2013-12-22 Implement graph visualizer in D3

2013-12-29 Implement node selection and highlighting

2014-01-14 Implement Node.js and MongoDB backend

2014-02-07 Implement user-based input, uploading text files and CMap query results

2014-03-01 Implement word-cloud generation based on selected graph nodes

2014-03-28 Implement filtering, showing/hiding graph nodes, brushing to select nodes based on connectivity score

2014-04-15 Implement text or spreadsheet result export

2014-05-01 Implement linking to external sites (GeneBank, ChemBank, for clicking on graph nodes)

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