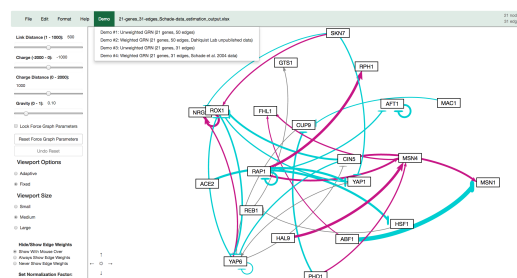
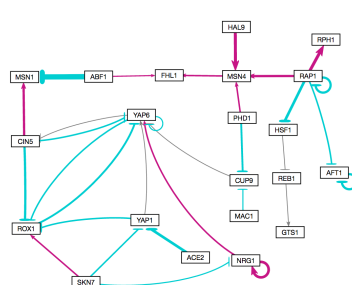


# GRNsight: a web application and service for visualizing models of small- to medium-scale gene regulatory networks

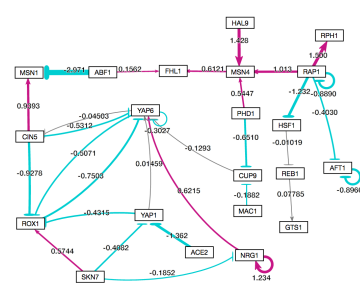
Anonymous Author(s)



(a) Screenshot of GRNsight



(b) Weighted graph after manual manipulation.



(c) Weighted graph with edge weights displayed.

**Figure 1:** GRNsight’s automatic graph layout of Demo #4: Weighted GRN (21 genes, 31 edges, Schade et al. 2004 data) in (a) is manipulated through manual node dragging and adjusting force graph parameters to produce the graphs in (b) and (c).

## ABSTRACT

We present new features in v2 of GRNsight, a web application and service for interactive visualization of small- to medium-scale gene regulatory networks (GRNs). A GRN consists of genes, transcription factors, and the regulatory connections between them which govern the level of expression of mRNA and protein from genes. GRNsight produces weighted or unweighted network graphs from an Excel spreadsheet containing an adjacency matrix where regulators are named in the columns and target genes in the rows, a Simple Interaction Format (SIF) text file, or a GraphML XML file. GRNsight represents genes as nodes and regulatory connections as edges with colors, end markers, and thicknesses corresponding to the sign and magnitude of activation or repression. GRNsight visualizations can be modified through manually dragging nodes or adjusting sliders that change the force graph parameters. GRNsight is best-suited for visualizing networks of fewer than 35 nodes and 70 edges, and has general applicability for displaying any small, unweighted or weighted network with directed edges for systems biology or other application domains. The GRNsight application (<http://dondi.github.io/GRNsight/>) and code (<https://github.com/dondi/GRNsight>) are available under the open source BSD license.

## CCS CONCEPTS

• Human-centered computing → Scientific visualization; Visualization toolkits;

## KEYWORDS

Scientific Visualization, Software Engineering, Bioinformatics

## ACM Reference format:

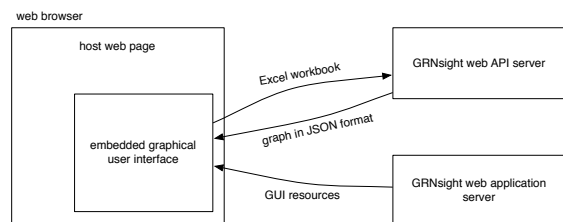
Anonymous Author(s). 2017. GRNsight: a web application and service for visualizing models of small- to medium-scale gene regulatory networks. In *Proceedings of SIGGRAPH 2017 Posters, Los Angeles, CA, USA, August 2017*, 2 pages. [https://doi.org/10.475/123\\_4](https://doi.org/10.475/123_4)

## 1 INTRODUCTION AND MOTIVATION

GRNsight is a web application and service for the interactive visualization of small- to medium-scale gene regulatory networks (GRNs), optimized for use by novice and experienced biologists alike to quickly and easily view unweighted and weighted network graphs. Visual inspection has long been recognized as distinct from other forms of purely numeric, computational, or algorithmic data analysis [Card et al. 1999; Tufte 1986], and GRNsight enables the potential for insight derived specifically by visual inspection. The following are the requirements for GRNsight:

- (1) Exist as a web application without the need to download and install specialized software;
- (2) Be simple and intuitive to use;
- (3) Automatically lay out and display small- to medium-scale, unweighted and weighted, directed network graphs in a way that is familiar to biologists and adds value to the interpretation of the modeling results.

**Software Architecture:** GRNsight has a service-oriented architecture, consisting of separate server and web client components. The server provides a web API that accepts files in Microsoft Excel workbook (.xlsx), SIF (.sif), and GraphML (.graphml) formats and converts them into a unified JSON representation. A converse import function accepts this JSON representation and converts it into either SIF (.sif) or GraphML (.graphml) formats for export. The web application server provides code and resources for the graphical user interface that displays this JSON representation of the graph.



**Figure 2: GRNsight architecture and component interactions.**

**Graph Customizations and User Interface:** Graph visualization is facilitated by the Data-Driven Documents JavaScript library [Bostock et al. 2011]. D3.js provides data mapping and layout routines which GRNsight heavily customizes in order to achieve the desired graph visualization. The resulting graph is a Scalable Vector Graphics (SVG) drawing in which D3.js maps gene objects from the JSON representation provided by the web API server onto labeled rectangles. Edge weights are mapped into Bezier curves. The resulting graph is interactive, initially using D3.js’s force graph layout algorithm to automatically determine the positions of the gene rectangles. The GRNsight user interface includes a menu/status bar and sliders that adjust D3.js’s force graph layout parameters, to refine the automated visualization. Design decisions for the user interface were driven by applicable interaction design guidelines and principles [Nielsen 1994; Norman 2013; Shneiderman 2010] in alignment with the mental model and expectations of the target user base, consisting primarily of biologists.

## 2 EXTENSIONS TO GRNSIGHT IN V2

Since the release of GRNsight v1, further research, as well as feature requests from peer review have motivated the following improvements to GRNsight v2. We focus on features which have enhanced the visualization and display of the network graph GRNsight produces.

### 2.1 Separation of Viewport from Graph Bounding Box

The bounding box and viewport for the graph have been separated allowing for the following new features

- (1) The bounding box can now be fixed to the size of the viewport or adapted to the size of the graph
- (2) The viewport size can be selected from among small, medium, and large options
- (3) The best viewport size is automatically detected from the browser
- (4) The viewport can be fit to the size of the browser window
- (5) Zooming and scrolling have been enabled

### 2.2 Edge Weight Display Options

A new visualization feature for weighted graphs introduced in GRNsight v2 is an options menu for displaying the edge weights of the graph. By default, edge weights are shown only upon mouse-over of the edge. Users can now choose to always show or always

hide weights. Edge Options exist in the sidebar menu, and under the Format tab in the File menu.

## 2.3 Graph Normalization

To allow for the comparison of weighted network graphs, a feature request was made for the user to have the option to customize the normalization factor applied to the edge thickness of the graph. By default, GRNsight detects the maximum and minimum relationship values of a network, and normalizes the data to fit within 12 distinct edge thicknesses corresponding to the strength of its regulatory relationship. By adding the option to user-specified minimum and maximum values, the edge thicknesses can be consistently normalized in a weighted graph, so the user can compare different graphs on the same scale.

## 3 CONCLUSION AND FUTURE WORK

We have successfully implemented GRNsight, a web application and service for visualizing small- to medium-scale GRNs that is simple and intuitive to use. GRNsight reads a weighted or unweighted representation of a GRN, and automatically lays out and displays unweighted and weighted network graphs, enabling interpretation of the weight parameters more easily than one could from an adjacency matrix alone. Extensions to GRNsight in v2 have expanded GRNsight’s visualization and layout capabilities, giving the user more control over the visual display of the network graph.

GRNsight is in active development, we would like to further enhance the tool to compute and display graph statistics, as betweenness centrality measure. Further research will be conducted to evaluate different graph layout options, such as a hierarchical or block layout and the effectiveness that alternate visualization paradigms for the biologist...

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