WEB TOOL

**BioJS DAGViewer**: A reusable JavaScript component for displaying directed graphs [v1; ref status: indexed, http://f1000r.es/2ut]

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**Abstract**

**Summary**: The DAGViewer BioJS component is a reusable JavaScript component made available as part of the BioJS project and intended to be used to display graphs of structured data, with a particular emphasis on Directed Acyclic Graphs (DAGs). It enables users to embed representations of graphs of data, such as ontologies or phylogenetic trees, in hyper-text documents (HTML). This component is generic, since it is capable (given the appropriate configuration) of displaying any kind of data that is organised as a graph. The features of this component which are useful for examining and filtering large and complex graphs are described.

**Availability**: http://github.com/alexkalderimis/dag-viewer-biojs; http://github.com/biojs/biojs; http://dx.doi.org/10.5281/zenodo.8303.

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Introduction

The graph abstract data type is an important concept in mathematics and computer science, and is the most appropriate representation for several classes of real world phenomena and scientific constructions. Some examples of these include phylogenetic trees, protein-protein interaction networks and scientific ontologies such as the Gene Ontology\(^1\) and the Sequence Ontology\(^2\). One feature of this type of data structure is that they are much easier for humans to understand when presented as a graphical network which preserves the structured nature of the data, than when they are displayed flattened in tabular or list format. The component described here is capable of displaying graphs of data, in particular Directed Acyclic Graphs (DAGs), efficiently using JavaScript to calculate the layout, and features of modern web browsers for rendering, and is designed to integrate with other components in the BioJS\(^3\) project.

Current methods and implementations

Two commonly used approaches for representing graphs in two dimensions, allowing display in HTML documents, are the force-directed layout, and the Sugiyama layout. These differ in the way that they represent the hierarchical organisation of elements within a graph, and are each suitable for different kinds of data.

Force-directed layouts distribute nodes throughout the available co-ordinate space, placing related nodes closer to each other, and unrelated nodes further away from each other. A typical method of achieving this is to model the layout as a two-dimensional particle simulation, where nodes exert a repulsive force upon each other, and edges between nodes exert an attractive force. Stable layouts are those representing local energy minima of the simulation.

This method is straightforward to implement (see Cytoscape\(^4\) and D3 project\(^5\) for example JavaScript implementations), and is a suitable representation of graphs where we care more about the existence of edges than their directions, and more about identifying clusters of nodes than elucidating the internal structure of such clusters. For example in a protein-protein interaction network (see Figure 1) force-directed layouts are often used since they are good at indicating highly connected interactors and clusters of interactors, thus highlighting centrally significant parts of the graph.

The other commonly-used approach to rendering graphs visually is Sugiyama-style graph drawing\(^6\). This method also attempts to group related elements, but in addition it assigns significance to the structure of relationships by introducing the concept of root and rank. Rank is defined as the number of edges in the shortest path from a node to a root. A root is defined as a node of rank 0. When rendered, nodes of the same rank within a graph are aligned visually, either horizontally or vertically, producing a structured hierarchical layout of the graph.

This method requires edges to have a direction that indicates which side of the relationship is closer to the root. Such graphs are typically described as trees, and the nodes furthest from the root as leaves. This kind of representation is suitable for graphs in which the structure of relationships is important, which is a feature of several types of graphs, such as ontologies, and phylogenetic trees. Singly rooted, acyclic trees are the most straightforward structures to lay out and display, but this method can be applied to multiply rooted directed graphs with cycles (such as biochemical pathways).

Until now, using the Sugiyama method has required the generation of image files, either on demand or through batch preprocessing, and then sending them out over the network to a suitable display device. Several tools exist for this purpose, including GraphViz\(^7\), which is used by several projects for rendering Gene Ontology

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**Figure 1.** Using Cytoscape to display an interaction network with a force-directed layout.
graphs. This requires any group wishing to employ this method for graphical network analysis to have access to the resources and expertise to manage either a server capable of dynamically generating such images, or to produce the images required in advance. In either case, user interaction is very limited.

What is new about the DAGViewer component is the use of a JavaScript Sugiyama layout engine to eliminate the image file generation, which cannot be done within a browser. Modern web browsers have advanced to the point where it is now practicable to calculate layouts for graphs of moderate size (in the order of around 200 to 500 nodes, depending on the density of connections) and render them in a dynamic hyper-text page, using tools such as JavaScript and Scaled Vector Graphics (SVG). This accounts for the great majority of networks that one might want to visualise, particularly since networks of greater information densities are very difficult for humans to interpret when rendered. We have taken advantage of the opportunity afforded by modern browser tools to produce a generic network display tool that does not require any server-side resources, and that is suitable for a variety of scientific purposes. This approach provides a much greater degree of customisation, interaction and flexibility than approaches based on image generation.

Features
The graph viewer presented here uses a collection of publically available, open-source JavaScript tools, including the Backbone framework, the dagre-d3 layout engine, and D3 data-binding and presentation library. The combination of these tools make it possible to build a tool in JavaScript and running in modern browsers that provides rich interaction and graphical analysis possibilities, allowing users to focus on the data, e.g. in the Gene Ontology Annotation display in Figure 2.

The current implementation allows a JavaScript component to be placed on any page and be provided with any kind of linked network data; the data are rendered to the screen in the familiar box and line style of a Sugiyama graph drawing. Unlike static images, this graph can be zoomed, panned, reorientated and rescaled, allowing users to make sense of dense networks. Since the graph is rendered with SVG technology, rescaling does not lower graphical resolution, and text legibility is preserved over a wide range of zoom levels.

The user can interact more deeply with this representation than they could with a standard fixed image. Individual nodes and edges can each have their own styles and behaviour, allowing contextual tooltips and mouseover effects to provide information even when zoomed out. Since the information composing the graph is available to the page at runtime as a data-structure, it can be searched and filtered, and the graph can be zoomed and scaled to highlight particular nodes and edges that interest the user.

A control panel element (see Figure 2) provides access to this functionality, allowing users to search for nodes within the graph, and filter the graph to focus on relevant sub-sets of the available information. Figure 3 illustrates the display of one particular subgraph of the information presented in Figure 2, reorientated to make the best use of the available screen space. This particular subgraph is defined as those nodes reachable from one particular high-level ontology term, developmental process.

Figure 2. The DAGViewer, displaying annotations (left) from the Gene Ontology for the Drosophila melanogaster gene cdc2, and the control panel (right).
Installation
As a BioJS JavaScript component, the intended audience is web developers aiming to provide functionality for life scientists. It is expected to be deployed within HTML pages and rendered in modern browsers. As such, installation means indicating which resources a page needs to load. The DAGViewer tool is a modular javascript component, making use of other existing resources (Supplementary materials A); these dependencies need to be included on the page before the component itself can be used. Once these are loaded the BioJS DAGViewer component itself can be included (see code sample 1). This should be downloaded from the BioJS repository and hosted locally.

Listing 1. Loading the DAG-Viewer Library
<script src="Biojs.DAGViewer.js"></script>

Usage
With these elements available, a user is then able to instantiate a new DAGViewer component pointing at a defined element in the document object model (DOM), or page:

Listing 2. Instantiating a new DAGViewer Component
var viewer = new Biojs.DAGViewer({
   target: "element-id"
});

There are a large number of configurable parameters that can be provided at instantiation (or indeed, later). These mostly relate to configuring how to interpret the graph data provided. It is accepted that data may come in different formats, and rather than requiring users to convert their node and edge data to a predefined format, users can provide adapters that allow this component to read and display different kinds of graph data, while providing sensible defaults. More detail is provided on the BioJS registry documentation pages, but as an example consider a graph (representing a protein interaction network) which has nodes of the form:

Listing 3. Example Nodes
var nodes = [
   {primaryAccession: "P09089", name: "Protein zerknueLL1"},
   {primaryAccession: "A0ANL0", name: null}
];

Here we will want to identify each node by its accession number (here from Uniprot) and label it by its name, if it has one, or by its accession if it does not. This behaviour can be defined by passing a couple of parameters:

Listing 4. Node Adaptor Example
var viewer = new Biojs.DAGViewer({
   target: "element-id",
   nodeLabels: ["name", "primaryAccession"],
   nodeKey: function (n) { return n.primaryAccession; }
});

Here the nodeLabels parameter indicates which fields should be read to obtain a label for this node, and the nodeKey parameter is a function that takes a node and returns an identifier (possibly computed). Similar configuration options exist for interpreting edges, determining the list of graph roots, providing style classes to nodes and edges and other functions.

Figure 3. The graph viewer, displaying a subgraph of annotations from the Gene Ontology for the Drosophila melanogaster gene cdc2, selected using the control panel.
Once configured, the component must be given the definition of the graph it is meant to visualise. A graph here is defined as two collections, one of nodes, and the other of edges between nodes. These can be unconnected data structures, such as loaded from JSON files, without interior references, or they may be circular self-referential data-structures, with edges pointing to their nodes. A small graph that represents a (grossly simplified) portion of the *H. sapiens* family tree, and the viewer to display it, could be configured as follows:

Listing 5. *H. sapiens* phylogenetic tree sample graph

```javascript
var species = [
    {name: "H. sapiens", status: "extant"},
    {name: "H. neanderthalensis", status: "extinct"},
    {name: "H. heidelbergensis", status: "extinct"},
    {name: "H. erectus", status: "extinct"},
    {name: "H. ergaster", status: "extinct"},
    {name: "H. habilis", status: "extinct"}
];

var relationships = [
    {subject: species[0], ancestor: species[2]},
    {subject: species[1], ancestor: species[2]},
    {subject: species[2], ancestor: species[4]},
    {subject: species[3], ancestor: species[4]},
    {subject: species[4], ancestor: species[5]}
];

var viewer = new Biojs.DAGViewer({
    target: "element-id",
    nodeLabels: ["name"],
    edgeProps: {"subject", "ancestor"}
});
viewer.setGraph(
    {nodes: species,
     edges: relationships});
```

As well as defining the data model, this component allows applications to respond to user input. An example of this is responding when a user clicks on a node in the graph. In the case of our human ancestry graph, that might look like this:

Listing 6. Listening for Events

```javascript
viewer.addListener(
    "click:node",
    function (name, species) {
        alert(name + " is " +
            species.get("status"));
    });
```

Beyond simplifying this task for developers wishing to get started in graphical network visualisation and analysis, by being built from open web-standard technologies this tool can be used to interoperate with existing and future applications in ways impossible for static image rendering tools. The graph definition can be fetched from a remote networked web service, for example, thus integrating with a large number of existing browser accessible tools.

Because of its flexible data definition, this component is able to consume data from a wide variety of different sources with minimal parsing effort. Since the standard node and edge representation is generally in the form of subject-predicate-object, this component would integrate very well into semantic web tools serving triples as their data representation.

### Conclusion

This component addresses an important need in the bioinformatics community for an effective, attractive and usable visualisation tool for a broad variety of directed acyclic graphs. It is therefore anticipated that this tool will be of use to those developing tools for researchers in the life sciences. A great deal of effort has gone into creating, curating and integrating high quality data sets, and there already exist many services which expose these data-sets to the world through networked web services. This tool is designed to plug in seamlessly with existing technologies, helping to maximise the value of existing and future curated data sets by bringing enhanced visualisation and exploration functionality. By publishing this component freely within the BioJS project we expect that a great deal of duplicated effort can be avoided, saving significant amounts of time and money for researchers and their funding bodies.

### Software availability

Zenodo: BioJS DAG-Viewer Component, doi: 10.5281/zenodo.830313.

GitHub: BioJS, http://github.com/biojs/biojs

### Author contributions

Alex Kalderimis wrote the manuscript and implemented the component, under the supervision of Gos Micklem, to a set of user
specifications supplied by Julie Sullivan. Rachel Lyne, Radek Štěpán and Mike Lyne contributed to the component design and revised the manuscript. All authors have approved the manuscript.

Competing interests
No competing interests were disclosed.

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Supplementary materials

A Dependencies

Note that the foundation scripts at least must be loaded in the body section of the HTML page, since they require the Document Object Model (DOM) to be ready.

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Referee Responses for Version 1

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Approved: 20 February 2014

Referee Report: 20 February 2014
This article describes a JavaScript component that allows web developers to render node/edge-type graphs in a web page, rather than in a static image, and interact with them dynamically via a tool bar and control panel. This component is a really nice addition to the world of graph visualization because it separates the definition of the graph from the displaying of it; it allows a user to programatically define the node/edge relationships and is extremely lightweight relative to most similar software tools.

The Sugiyama method used by the authors to determine node and edge placement seems to work well. I only tested the demo graph, but I couldn't think of any way to improve the layout - unlike most other tools I've tried. The implementation also seems quite straightforward and clear, very human-readable for source code.

There are, however, a couple issues that I would like the authors to address:
1. Even though I didn't think moving nodes around was necessary in the demo it was still the first thing I wanted to do (e.g., clicking on node and dragging it into a new position) and I can imagine other graphs where this might feel a bit more urgent. I suspect this sort of functionality is beyond the scope of the JS component, but could the authors either convince me that I'm being old-fashioned and this is something I don't really want to do or mention that this is not an intended use of tool?
2. I can imagine situations in which a user would like to generate a slide or publication-ready image from the graph that they've rendered and lovingly customized. Is there a way to export an image? Or is that in contradiction to the imageless nature of the HTML-based rendering?
3. Will there be a tutorial or walk through of the control panel features? I think this would be welcomed by future users.

Minor issues:

- In paragraph 3 of the 'Features' section the phrase “the graph can been zoomed” should read the graph can be zoomed.
- Demo code - in demo.html one of the script src links is localhost. I had to change this to match the other src link before the script would work so this needs to be corrected.

Overall, this seems like a sound and useful contribution to the field of computational biology.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.
Referee Report: 20 February 2014
The work is an interesting and well written article and reports a resource that should be of interest to numerous developers.

The comments below are intended to help make the work more accessible to others:

- It would be useful to define what some of the graph terms mean such as acyclic/cycles for those who are not familiar with the terminology. Similarly, some background information about the InterMine web services would be useful.
- How does one reproduce the examples in Figures 2 and 3?
- Where do the graph size numbers 200-500 nodes for modern browsers come from? Maybe the authors could qualify which with browser, system RAM and which JavaScript API the numbers are for (presumably D3). I would also disagree with statement that ‘networks of greater information densities are very difficult for humans to interpret when rendered’. See our tool BioLayout Express3D.
- How do I run the examples at http://zenodo.org/record/8303? It might be useful to host the examples somewhere that we can just click and run the demo? (jsFiddle is a great place to host JavaScript examples).
- There are a few typos - paragraph 7 of the ‘Current methods and implementations’ section "Scaled Vector Graphics (SVG)" should read Scalable Vector Graphics (SVG). In paragraph 3 of the ‘Features’ section should "mouseover" read mouseover? In the same paragraph the phrase “the graph can been zoomed” should read the graph can be zoomed.

We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Competing Interests: No competing interests were disclosed.