SOFTWARE TOOL ARTICLE

enhancedGraphics: a Cytoscape app for enhanced node graphics [version 1; referees: 2 approved]

John H. Morris¹, Allan Kuchinsky², Thomas E. Ferrin¹, Alexander R. Pico³

¹Resource for Biocomputing, Visualization and Informatics, University of California, San Francisco, CA 94143, USA
²Agilent Technologies, Santa Clara, CA 95051, USA
³Gladstone Institutes, San Francisco, CA 94158, USA

Abstract

enhancedGraphics (http://apps.cytoscape.org/apps/enhancedGraphics) is a Cytoscape app that implements a series of enhanced charts and graphics that may be added to Cytoscape nodes. It enables users and other app developers to create pie, line, bar, and circle plots that are driven by columns in the Cytoscape Node Table. Charts are drawn using vector graphics to allow full-resolution scaling.

This article is included in the International Society for Computational Biology Community Journal gateway.

This article is included in the Cytoscape Apps gateway.
Introduction
Cytoscape provides support for coloring and sizing nodes and node borders based on data values stored in the Node Table. This provides an extremely useful mapping between data values and a single visual property, but does not solve the need for more complex visualizations. Over the years, there have been attempts to support more complex mappings of multiple data values onto node visuals in Cytoscape. These include GOlorize, which maps GO terms to pie charts on nodes; GenePro, which visualizes groups of nodes as pie charts; VistaClará, which adds bar graphs to represent expression data; and more recently MultiColoredNodes. Each of these plugins and apps implemented their own graph and chart capabilities that are not accessible to other apps and in some cases not applicable outside of specific types of analyses. We felt that a better approach would be to implement an app that provided general support for graphs and charts to users as well as to the developers of other apps. The initial version of this approach was developed for Cytoscape 2.8 as nodeCharts, which was used by clusterMaker, for example, and numerous users (see Figure 3 in the Jäger, et al. paper). For Cytoscape 3, we reimplemented this approach as an app to take advantage of the new architecture and custom graphics API. The mechanism supports saving and restoring charts, as well as high-quality image file output suitable for publication. The intent is to provide a single, consistent, mechanism to draw charts and graphs on nodes as a general solution for diverse users and other app developers, mitigating the need to reinvent this capability in future apps.

Implementation
As part of the visual property mechanism, enhancedGraphics utilizes the Cytoscape 3 custom graphics API (org.cytoscape.view.presentation.customgraphics). To use the gradients and charts provided by enhancedGraphics, an app or user would create two things: a column that contains the instructions for creating the chart, and a passthrough visual mapping that maps that column to one of the custom graphics visual properties. The format of the instruction column is type: arglist, where type is the type of gradient or chart, and arglist is a list of name=value pairs that specify the arguments to create the gradient or chart (see details and examples in the tables below). The drawing and display of the chart or graph is handled by enhancedGraphics methods that are called by the Cytoscape rendering engine.

Internally, each enhancedGraphics chart type implements a CyCustomGraphicsFactory that is registered with OSGi. Each CyCustomGraphicsFactory informs the visual mapping mechanism of the chart type (e.g. ligrad) and method to create the CyCustomGraphics object given a String, which is the instruction column value. The CyCustomGraphics object parses the String as appropriate. Each CyCustomGraphics object implements a getLayers method that generates the appropriate list of CustomGraphicsLayers. The API defines three types of CustomGraphicsLayers: (1) the base interface, CustomGraphicsLayers, that provides a getPaint method to return a simple java.awt.Paint for the node; (2) ImageCustomGraphicLayer, that adds a getPaint method that returns a java.awt.TexturePaint suitable for painting an image on a node; and (3) PaintedShape which adds methods to return Shapes, Strokes, and Paints to draw arbitrary shapes. enhancedGraphics utilizes the base CustomGraphicsLayer for the two gradient types and PaintedShape for all of the charts.

Results
Figure 1 shows examples of all of the gradients and charts that are provided by enhancedGraphics. Up to nine different graphs can be combined on a single node by mapping different columns to different Custom Graphics properties and then offsetting the charts using the corresponding Custom Graphics Position properties. enhancedGraphics currently provides two different types of graphics options: gradients and charts.

Gradients
Gradients are simple paints on nodes. In order to provide the user with control over the exact paint to use, both linear and radial gradients allow the user to specify the gradient start and end (or center point and radius) and a stop list of color and opacity values. Table 1 provides the prefixes and arguments for the two gradient types.

Charts
enhancedGraphics currently provides six chart types: bar, circos, heat strip, line, pie, and stripe. Each chart type has its own set of

Figure 1. enhancedGraphics example Gradients and Charts.
arguments as shown in Table 3. In addition, there are a number of common options that are used by many of the charts. Table 2 provides the syntax and explanation for each of these common arguments.

**Examples**
The example charts shown in Figure 1 and provided in the Cytoscape session file Supplementary File 1 are generated from data columns. The instructions in the chart columns assume that the following columns exist: a, b, c, and d are integer columns in the default node table; Values is a list of Doubles also in the default node table, and Circle1 and Circle2 are also lists of Doubles. At this point, gradients are not dependent on any internal data. See Supplementary File 1 to see the instructions that generated Figure 1.

A more relevant biological example is shown in Figure 2. This image shows a portion of the galFiltered.cys network delivered as part of the sampleData with every Cytoscape download. The bar charts show the values of the expression data included as columns.

### Table 1. Gradient prefixes and arguments.

<table>
<thead>
<tr>
<th>Type</th>
<th>Prefix</th>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear Gradient</td>
<td>lingrad</td>
<td>start=&quot;x,y&quot; end=&quot;x,y&quot; stoplist=&quot;r,g,b,a,stop1</td>
<td>r,g,b,a,stop2</td>
</tr>
<tr>
<td>Radial Gradient</td>
<td>radgrad</td>
<td>center=&quot;x,y&quot; radius=&quot;r&quot; stoplist=&quot;r,g,b,a,stop1</td>
<td>r,g,b,a,stop2</td>
</tr>
</tbody>
</table>

### Table 2. Common arguments used by many charts.

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>attributelist=&quot;attr1.attr2...attrn&quot;</td>
<td>List of columns to use to get the values for the chart.</td>
</tr>
<tr>
<td>colorlist= [contrasting/modulated/rainbow/random updown colors]</td>
<td>The colorlist argument provides a number of options, including a series of keywords for automatically generating colors. updown colors is a specification of colors for positive, negative, and zero values: up:color, down:color[.zero:color] where the zero is optional. colors may be specified as a color name (red, green, blue) or an rgb or rgba color in hex notation, e.g. #FF0000 for red.</td>
</tr>
<tr>
<td>labels=&quot;label1,label2,..., labeln&quot;</td>
<td>List of labels for the chart. If not provided and attributelist is provided, the names of the attributes are used as labels.</td>
</tr>
<tr>
<td>labelcolor=color</td>
<td>Color of the labels</td>
</tr>
<tr>
<td>labelfont=font name</td>
<td>Font to use for the labels</td>
</tr>
<tr>
<td>labelsize=value</td>
<td>Size of the label font</td>
</tr>
<tr>
<td>labelstyle=[italics/bold/bolditalic/plain]</td>
<td>label style</td>
</tr>
<tr>
<td>range=&quot;min,max&quot;</td>
<td>The min and max range as floating point values. This is used to have consistent scaling across all nodes</td>
</tr>
<tr>
<td>scale=&quot;scale&quot;</td>
<td>A floating point value used to scale the chart.</td>
</tr>
<tr>
<td>showlabels=[true/false]</td>
<td>If false, labels aren’t drawn</td>
</tr>
<tr>
<td>values=&quot;v1,v2,...,vn&quot;</td>
<td>A list of values to use for the chart. One of values or attributelist will often be required, but not both</td>
</tr>
<tr>
<td>ybase=[top/middle/bottom]</td>
<td>The specified the base of the chart. Usually used to set the location of the 0 value for line and bar graphs.</td>
</tr>
</tbody>
</table>
Table 3. Charts and arguments.

<table>
<thead>
<tr>
<th>Chart Type</th>
<th>Prefix</th>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bar Chart</td>
<td>bchart</td>
<td>separation=value</td>
<td>The separation between bars</td>
</tr>
<tr>
<td>Circos chart</td>
<td>circoschart</td>
<td>arcstart=value</td>
<td>The start of each circle in degrees</td>
</tr>
<tr>
<td></td>
<td></td>
<td>arcwidth=value</td>
<td>The thickness of each of the rings</td>
</tr>
<tr>
<td></td>
<td></td>
<td>firstarc=value</td>
<td>The start of the first arc as a proportion of the entire node</td>
</tr>
<tr>
<td></td>
<td></td>
<td>firstarcwidth=width</td>
<td>The width of the first arc</td>
</tr>
<tr>
<td></td>
<td></td>
<td>labelcircles=</td>
<td>If true label each circle</td>
</tr>
<tr>
<td></td>
<td></td>
<td>[true</td>
<td>false]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sortslices=</td>
<td>If true sort the slices from largest to smallest</td>
</tr>
<tr>
<td></td>
<td></td>
<td>[true</td>
<td>false]</td>
</tr>
<tr>
<td>Heat strip chart</td>
<td>heatstripchart</td>
<td>colorlist=</td>
<td>Current gradient keywords include:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gradient keyword</td>
<td>yellowcyan, yellowblue, orangepurple, bluegreenyellow, purpleyellow,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>updown colors</td>
<td>greenpurple, redyellow, and if you absolutely must: redgreen. See Table 2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>range=</td>
<td>for a description of updown colors.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>separation=value</td>
<td>The separation between bars</td>
</tr>
<tr>
<td>Line chart</td>
<td>linechart</td>
<td>linewidth=value</td>
<td>The width of the lines on the plot</td>
</tr>
<tr>
<td>Pie chart</td>
<td>piechart</td>
<td>arcstart=value</td>
<td>The start of each circle in degrees</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sortslices=</td>
<td>If true sort the slices from largest to smallest</td>
</tr>
<tr>
<td></td>
<td></td>
<td>[true</td>
<td>false]</td>
</tr>
<tr>
<td>Stripe chart</td>
<td>stripechart</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

gal1RGexp, gal4RGexp, and gal80Rexp. A string column was created and all rows were filled with the enhancedGraphic arguments:

heatstripchart:

  `attributelist="gal1RGexp,gal4RGexp,gal80Rexp"
colorlist="yellowblue" range="-3.0,3.0"`

Conclusions

enhancedGraphics fills an important need for Cytoscape visualizations: the ability to display more complicated data relationships as graphical representations onto nodes. enhancedGraphics has been integrated into clusterMaker to show heatstrips on nodes corresponding to clusters, into upcoming apps such as cddApp, which connects to the NCBI CDD Database and uses enhancedGraphics to show pie charts of the domain coverage. We have also used enhancedGraphics to show sequence coverage histograms on nodes that represent sequence contigs. In the future, we want to improve the font handling and add support for small heatmaps painted on nodes. We also plan to add a graphical interface to help users construct these visualizations without having to write out instruction arguments. The concise syntax, however, will still be valuable to advanced users and other app developers seeking to create enhanced graphics. With enhancedGraphics, Cytoscape users and app developers can visualize multiple columns of data as graphs and charts on their network nodes.
Figure 2. Example of using enhancedGraphics to show expression data in the context of a protein-protein interaction network.

Software availability
Software available from: http://apps.cytoscape.org/apps/enhanced-Graphics
Latest source code: https://github.com/RBVI/enhancedGraphics
Source code as at the time of publication: https://github.com/F1000Research/enhancedGraphics/releases/tag/v1
Archived source code as at the time of publication: http://www.dx.doi.org/10.5281/zenodo.1042111
Software license:
Lesser GNU Public License 3.0: https://www.gnu.org/licenses/lgpl.htmlM

Author contributions
JHM, AK, and ARP wrote the original nodeCharts plugin. TEF and ARP supervised the project and contributed to the manuscript. JHM ported the app to Cytoscape 3 and wrote the manuscript.

Competing interests
No competing interests were disclosed.

Grant information
ARP and JHM were funded by NIGMS grant P41-GM103504. JHM and TEF were funded by NIGMS grant P41-GM103311.

Supplementary material
Example of the Instructions used to create Figure 1.
References


F1000Research 2014, 3:147 Last updated: 20 AUG 2018
Open Peer Review

Current Referee Status: ✔ ✔

Version 1

Referee Report 04 September 2014

doi:10.5256/f1000research.4771.r6001

Piet Molenaar
Academic Medical Center, Amsterdam, Netherlands

The app described in the article addresses a specific issue with the Cytoscape network visualization tool: drawing of multidimensional data on nodes is currently not possible by default. As such this is a very valuable addition to the Cytoscape app ecosystem. The article is well written but I do have some minor issues that can be considered for further improvement:

Minor issue
The implementation details are a too technical when an intended audience includes biologists (and might even scare them away). I would suggest to transfer these OSGI specific details to a supplement or add a protocol section before this section with a sample protocol (including installation details).

Minor issue
The current sample session file included in the supplemental data only contains example nodes. A session file or additional network showing actual data mapped on the gal dataset (as in the figure) would further clarify the actual usage of the app.

Minor issue
In the conclusions section references to cddApp and NCBI CDD Database are lacking.

Competing Interests: No competing interests were disclosed.

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.

Referee Report 17 July 2014

doi:10.5256/f1000research.4771.r5308

Lars Juhl Jensen
NNF Center for Protein Research, University of Copenhagen, Copenhagen, Denmark

General comments:

The authors describe a Cytoscape App, which provides a generic interface for displaying additional data
onto the nodes of a graph. I believe that this App will be highly useful to researchers working with transcriptomics and proteomics data, as these often have a need to display, for example, expression time-courses onto protein interaction networks. This is especially true if the App is used as the foundation for other more specialized Apps that make it easier for users to import and visualize specific data types in an appropriate manner.

Specific comments:

The *Integration* section is difficult to fully follow for people who are not Cytoscape developers, who cannot be assumed to know what, for example, a *CyCustomGraphicsFactory* is. It would in my opinion be good to revise this section to make it understandable to a broader audience.

In Figure 1, I find it strange that the pie charts are not circular and that the circos charts, unlike all the other chart types, are not embedded within a frame with rounded corners. Also, the figure would benefit from a figure caption explaining the difference between Pie Chart 1 and Pie Chart 2, as well as between Circos 1 and Circos 2. The differences between these chart subtypes are not clear to me.

**Competing Interests:** I have been and will be involved in teaching advanced courses together with the first author.

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.