SOFTWARE TOOL ARTICLE

Visualize omics data on networks with Omics Visualizer, a Cytoscape App [version 2; peer review: 2 approved]

Marc Legeay1, Nadezhda T. Doncheva1,2, John H. Morris3, Lars Juhl Jensen1

1Novo Nordisk Foundation Center for Protein Research, University of Copenhagen, Copenhagen, Denmark
2Center for Non-Coding RNA in Technology and Health, Department of Veterinary and Animal Sciences, University of Copenhagen, Copenhagen, Denmark
3Resource for Biocomputing, Visualization and Informatics, University of California, San Francisco, USA

First published: 28 Feb 2020, 9:157
https://doi.org/10.12688/f1000research.22280.1

Open Peer Review

Reviewer Status
Invited Reviewers
1
2

version 2
(revision)
30 Jun 2020

version 1
28 Feb 2020

1. Magnus Palmblad ID, Leiden University Medical Center, Leiden, The Netherlands
2. Ruth Isserlin ID, University of Toronto, Toronto, Canada

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Abstract
Cytoscape is an open-source software used to analyze and visualize biological networks. In addition to being able to import networks from a variety of sources, Cytoscape allows users to import tabular node data and visualize it onto networks. Unfortunately, such data tables can only contain one row of data per node, whereas omics data often have multiple rows for the same gene or protein, representing different post-translational modification sites, peptides, splice isoforms, or conditions. Here, we present a new app, Omics Visualizer, that allows users to import data tables with several rows referring to the same node, connect them to one or more networks, and visualize the connected data onto networks. Omics Visualizer uses the Cytoscape enhancedGraphics app to show the data either in the nodes (pie visualization) or around the nodes (donut visualization), where the colors of the slices represent the imported values. If the user does not provide a network, the app can retrieve one from the STRING database using the Cytoscape stringApp. The Omics Visualizer app is freely available at https://apps.cytoscape.org/apps/omicsvisualizer.

Keywords
Cytoscape, app, network visualization, omics data, network biology

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

**Corresponding authors:** Marc Legeay (marc.legeay@cpr.ku.dk), Lars Juhl Jensen (lars.juhl.jensen@cpr.ku.dk)

**Author roles:** Legeay M: Software, Writing – Original Draft Preparation, Writing – Review & Editing; Doncheva NT: Software, Writing – Review & Editing; Morris JH: Conceptualization, Writing – Review & Editing; Jensen LJ: Conceptualization, Writing – Review & Editing

**Competing interests:** No competing interests were disclosed.

**Grant information:** This work was funded by the Novo Nordisk Foundation (NNF14CC0001). NTD was also funded by Innovation Fund Denmark “Brainstem” (4108-00008B).

*The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.*

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**How to cite this article:** Legeay M, Doncheva NT, Morris JH and Jensen LJ. Visualize omics data on networks with Omics Visualizer, a Cytoscape App [version 2; peer review: 2 approved] F1000Research 2020, 9:157 https://doi.org/10.12688/f1000research.22280.2

**First published:** 28 Feb 2020, 9:157 https://doi.org/10.12688/f1000research.22280.1
Introduction

Cellular functions are mediated by complex networks of interactions between genes, proteins, and other molecular entities. Omics technologies are commonly used to measure the detailed regulation of these networks by quantifying changes of individual post-translational modification (PTM) sites, peptides, or splice isoforms across different experimental conditions. However, it is not easy to visualize such data sets, which have multiple values per gene or protein, onto the networks using existing network visualization tools such as Cytoscape\(^1\) or Gephi\(^2\).

To address this, we present the new Omics Visualizer app for Cytoscape\(^3\). The app allows users to import data tables with several rows referring to the same node and to visualize such data on networks; while designed with omics data in mind, the app is data agnostic. These values can be shown directly on the nodes of the networks as pie or donut visualizations, in which the color of each slice represents a different value for the same node. The Omics Visualizer app was implemented using the API that Cytoscape makes available to developers, and the data visualization builds upon the enhancedGraphics app\(^4\). The app furthermore integrates with the stringApp\(^5\) to facilitate easy visualization of data onto networks from the STRING database\(^6\) and supports Cytoscape Automation to facilitate integration with other tools\(^7\).

The workflow for visualizing data with the app is outlined in Figure 1. The first key step is to obtain an Omics Visualizer table that is connected with a network in Cytoscape. There are three ways to do this: 1) import a table from a file and explicitly connect it to an existing network in Cytoscape, 2) import a table from a file and retrieve a STRING network based on the table, and 3) import selected columns from the node table of an existing Cytoscape network into an Omics Visualizer table. Once the table and the network are connected, it is possible to create data visualizations on the network and then generate a legend.

Methods

Implementation

The typical Omics Visualizer\(^3\) workflow consists of four steps: importing data as a table, optionally filtering the data, connecting it to one or more networks, and finally visualizing the connected data onto the networks.

There are two ways to import an Omics Visualizer table, the first one is from a file, the second one is from a node table. The Omics Visualizer table import from file mimics the Cytoscape default import process. The app can handle text files (e.g. comma- or tab-delimited values) as well as spreadsheet files from e.g. Microsoft Excel. The app auto-detects the data type of each column, and the user can subsequently select which columns to import and modify the auto-detected data type for each column if needed. To also allow visualization of data from a network’s node table, selected columns can be imported as an Omics Visualizer table with three columns such that each row lists...
the identifier of a node (node column), the name of a selected column from the node table (source column), and the imported value (values column). Both import methods create two private unassigned tables in Cytoscape: one that contains the data imported from the file, and another that stores all associated Omics Visualizer properties. These tables are private, which means that the user cannot interact with them directly in the Cytoscape UI. Instead, the data can be viewed (but not edited) through the Omics Visualizer panel located in the table panels. The tables are also accessible via the command interface and the Cytoscape API.

The imported file should be formatted as indicated in Figure 2, which is a sample of the table used in Figure 4 and Figure 5. One column must contain the node identifier value (‘UniProt’ in Figure 2), which will be used to connect the Omics Visualizer table with the node table. The other columns contain the data to be visualized. In case of a pie visualization, all values are in one column and the slices for one pie are contained in different rows with the same node identifier (‘Cluster’ column in Figure 2 A, C). In a case of donut visualizations, the values can be contained in several columns (‘EOC vs FTE’ and ‘EOC vs OSE’ columns in Figure 2 B, C). It is possible to have additional columns in the file, which can be imported and used to filter the rows, as protein identifiers for retrieving a STRING network or as labels (‘AA position’ in Figure 2).

If the data was not already filtered before importing it into Cytoscape, Omics Visualizer enables users to do so afterwards. Omics Visualizer can filter the rows based on selected columns with string, numeric and Boolean values and offers several operators depending on the data type:\texttt{EQUALS}, \texttt{NOT\_EQUALS}, \texttt{NULL} and \texttt{NOT\_NULL} for all types; \texttt{CONTAINS}, \texttt{NOT\_CONTAINS} and \texttt{MATCHES} for string values; \texttt{LOWER}, \texttt{LOWER\_EQUALS}, \texttt{GREATER} and \texttt{GREATER\_EQUALS} for numeric values. Similar to the Cytoscape selection filters, Omics Visualizer interface allows the user to create Boolean formulas with nested criteria. Once the filter is applied, rows that do not satisfy the filter are hidden in the panel, and the active filter is stored in the properties table to allow it to be changed later.

To visualize the data table onto a network, the Omics Visualizer table must first be connected to the network by specifying matching key columns in the node and Omics Visualizer tables. This connection information is stored in the network table, which is used to recreate the link when a session file is loaded. An Omics Visualizer table can be connected to several networks, but a network can only be connected to one Omics Visualizer table.

<table>
<thead>
<tr>
<th>A</th>
<th>UniProt</th>
<th>AA position</th>
<th>Cluster</th>
</tr>
</thead>
<tbody>
<tr>
<td>P20700</td>
<td>S391</td>
<td>B</td>
<td></td>
</tr>
<tr>
<td>P20700</td>
<td>S404</td>
<td>C</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S983</td>
<td>A</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S1385</td>
<td>A</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S1545</td>
<td>C</td>
<td></td>
</tr>
<tr>
<td>O15164</td>
<td>S722</td>
<td>C</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>B</th>
<th>UniProt</th>
<th>AA position</th>
<th>EOC vs FTE</th>
<th>EOC vs OSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>P20700</td>
<td>S391</td>
<td>0.10</td>
<td>2.37</td>
<td></td>
</tr>
<tr>
<td>P20700</td>
<td>S404</td>
<td>2.77</td>
<td>3.82</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S983</td>
<td>-4.30</td>
<td>-3.98</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S1385</td>
<td>-3.89</td>
<td>-3.84</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S1545</td>
<td>4.66</td>
<td>4.16</td>
<td></td>
</tr>
<tr>
<td>O15164</td>
<td>S722</td>
<td>3.92</td>
<td>4.09</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>C</th>
<th>UniProt</th>
<th>AA position</th>
<th>EOC vs FTE</th>
<th>EOC vs OSE</th>
<th>Cluster</th>
</tr>
</thead>
<tbody>
<tr>
<td>P20700</td>
<td>S391</td>
<td>0.10</td>
<td>2.37</td>
<td>B</td>
<td></td>
</tr>
<tr>
<td>P20700</td>
<td>S404</td>
<td>2.77</td>
<td>3.82</td>
<td>C</td>
<td></td>
</tr>
<tr>
<td>Q9COC2</td>
<td>S983</td>
<td>-4.30</td>
<td>-3.98</td>
<td>A</td>
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<tr>
<td>Q9COC2</td>
<td>S1385</td>
<td>-3.89</td>
<td>-3.84</td>
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</tr>
<tr>
<td>Q9COC2</td>
<td>S1545</td>
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<td>4.16</td>
<td>C</td>
<td></td>
</tr>
<tr>
<td>O15164</td>
<td>S722</td>
<td>3.92</td>
<td>4.09</td>
<td>C</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2. Desired formats for table import. This shows three examples of how tables can be formatted before being imported with Omics Visualizer. One column contains the node identifier (‘UniProt’), and the other columns contain the values to be visualized (‘Cluster’ with pie visualization, ‘EOC vs FTE’ and ‘EOC vs OSE’ with donut visualization). It is possible to format your table for pie visualization (A), donut visualization (B), or both (C). You can have some extra columns that will not be used to visualize but for other purposes, for example, here the column ‘AA position’ identifies the specific site of the protein and are used as labels.
Visualizer table. If the user wants to use both types of visualization provided by Omics Visualizer (pie and donut), the data for both visualizations should be imported as a single table (Figure 2 C). A node table key can match the key in several Omics Visualizer table rows, in which case the node is connected to all of them. The number of connected rows for each node is stored in the node table; note that this number does not reflect any filtering. When the Omics Visualizer table is imported from the node table, the connection is automatically done using the identifier column (node column) as key.

Alternatively, Omics Visualizer can retrieve and automatically connect a network from the STRING database6 using the Cytoscape stringApp5. The user can select any column with gene/protein identifiers, which will be used to first query STRING to retrieve a network and subsequently as the key for connecting the network to the Omics Visualizer table.

Once connected, the user can show the data from the Omics Visualizer table on networks using either a pie or a donut visualization. The data values are mapped to colors using either a discrete mapping, where the user chooses the colors for each value, or a continuous mapping, where the user defines a color gradient. For the continuous mapping, the user specifies three values and corresponding colors, namely the minimum, middle, and maximum. Every value lower than the minimum or greater than the maximum value will be shown using the minimum and maximum color, respectively. Predefined color mappings can be used with the help of Cytoscape palettes such as ColorBrewer4 and Viridis (originally from Matplotlib4). The charts are drawn thanks to the enhancedGraphics app5. This app draws charts based on a description string, which is specified as a Cytoscape Custom Chart node style.

When the user creates a visualization, Omics Visualizer creates a column in the network table and several columns in the node table, and then it activates a Custom Chart node style: Omics Visualizer uses Custom Chart 7 to visualize pies and Custom Chart 8 to visualize donuts. The network table column is used to store the visualization properties. One node table column is created to store the enhancedGraphics string. Several node table columns are created to store the values of the visualizations: one column for pie visualization, and one column per ring for donut visualization. The values from the Omics Visualizer table rows are formatted into lists to fit the enhancedGraphics requirements. The enhancedGraphics continuous mapping always ranges between two defined values minimum and maximum, and a specific color can be given to zero. Omics Visualizer allows the user to define a minimum, middle and maximum value as it fits the data. Omics Visualizer centers the data around the middle value and adjusts the minimum and maximum accordingly. The values are only modified in the node table columns, not in the actual Omics Visualizer table.

Omics Visualizer can automatically generate legends from the visualizations in the form of Cytoscape annotations, which can be exported as part of the images if the user exports the network. The user may specify a title, font, font size and default location of the legend. To allow users to easily modify the legend, each element of the legend is a separate annotation. These are grouped to let the user move or delete the legend in one click, once the user has activated the Annotation Selection from Cytoscape (The “T” with a dotted border icon at the bottom of the network or the ‘Select → Mouse Drag Selects → Annotations Only’ menu). The name of the annotation group is used to differentiate the legend annotations from other annotations and should thus not be changed. When the legend is created, Omics Visualizer creates a network table column to store it.

All the columns created by Omics Visualizer in the node or network tables are in the namespace “Omics Visualizer”, enabling the user to easily identify and hide them if desired.

Omics Visualizer is built from Java 1.8 using Cytoscape API 3.7, meaning the minimum required version of Cytoscape is 3.7.0.

Operation
Cytoscape can be used with R or Python through to Cytoscape Automation7. Omics Visualizer implements commands in the specific ‘ov’ namespace, allowing Omics Visualizer to be used with the REST API. A full documentation of the commands is available at https://github.com/marcelegay/omics-visualizer/blob/master/automation_documentation.md. With the commands, the users can import a table, filter it, connect it with an existing network, retrieve a STRING network, create visualizations, and generate legends.

Use cases
We will illustrate how to use Omics Visualizer by visualizing site-specific proteomics data from a phosphoproteomics study of ovarian cancer by Francavilla et al.8. This study compares the phosphoproteome of primary cells derived from epithelial ovarian cancer (EOC) and two healthy tissues, namely ovarian surface epithelium (OSE) and distal fallopian tube epithelium (FTE). The goal of the study was to uncover cancer-specific changes in expression, phosphorylation state, and kinase signatures by comparing cancer and two healthy tissues (EOC vs. OSE, and EOC vs. FTE) for each site. The sites were then clustered into three clusters: A) sites abundant in healthy (OSE and FTE) but not cancerous tissues (EOC); B) sites abundant in FTE and EOC, but not OSE; and C) sites abundant only in cancerous tissue (EOC).

In this example, we want to visualize this site-specific data set on a network in which each node is a protein. We start from Table S3 from Francavilla et al., where each line represents a phosphorylation site in a protein, as opposed to a protein. We will first load the table file into Cytoscape thanks to Omics Visualizer and then create a STRING network from the data. We will finally apply some style to the network to visualize the data.

Importing the data from a file
Table S3 only contains the expression in the different samples, but not the individual comparisons between EOC and healthy
tissues, so we modified it to add the comparison columns ‘EOC vs OSE’ and ‘EOC vs FTE’. The modified file is available as Underlying data.

To load the data, we will use the specific import feature from Omics Visualizer, that can be accessed from the ‘File → Import → Omics Visualizer table from File...’ menu, or the ‘Apps → Omics Visualizer → Import table from file’ menu. This opens a custom dialog, which is very similar to the standard Cytoscape import table dialog. The user can name the table (or a default name will be given) and select the columns and their type to import. By default, all columns are imported and their type is inferred according to the first hundred lines of the file.

Here, we specifically want to import the ‘UniProt’ column to retrieve a STRING network afterwards, the ‘AA Position’ column to label the sites, the comparison columns ‘EOC vs OSE’ and ‘EOC vs FTE’ with log-ratios to be visualized, the ‘Adj p-value’ column to filter the table by significant sites, and the cluster assignment column ‘Cluster’ to visualize it. The file can be loaded with the following command:

```
ov load file= "/path/to/Francavilla2017CellRep.tsv"
```

Once the table is imported, the Omics Visualizer panel appears (Figure 3, left table). The top part of the panel has a row of icons giving access to the different features, displays the number of rows of the table, and enables access to the different Omics Visualizer imported tables. The second part of the panel is the table itself.

Filtering the table

It is possible to filter the table with the GUI or with the automation command. The filter GUI can be accessed via the filter icon from the Omics Visualizer panel (Figure 3, left), or with the ‘Apps → Omics Visualizer → Filter table’ menu. We filter the table so that we select only sites that have an adjusted p-value lower or equal to 0.01. This filter can be applied to the current table with the following automation command:

```
ov filter filter="(Adj p-value,LOWER_EQUALS,0.01)"
```

After the filter has been applied, the filter icon changes colors and the number of rows before and after filtering is displayed (Figure 3, left table).

Connecting a network

In this use case, we will use a STRING network corresponding to our data. With the help of the stringApp, we perform a ‘protein query’ with the list of the UniProt identifiers of proteins from our table that have at least one phosphorylation site with an adjusted p-value lower or equal to 0.01.

The current version of the stringApp retrieves a STRING v11 network with default confidence of 0.4 that consists of 237 nodes and 1020 edges. To reduce the size of the network, we cluster it using the Markov clustering (MCL) from the cluster-Maker2 app. We used the inflation value of 2.5 and the stringdb score as array sources. For illustrations purposes, we show only the second biggest cluster, which contains 40 nodes and 107 edges.

Once the STRING network is imported in Cytoscape by the stringApp, we connect the network with the table. The link icon from the Omics Visualizer panel (Figure 3, left) or the ‘Apps → Omics Visualizer → Manage table connections’ menu give access to the connect dialog. The dialog shows already connected networks and gives the possibility to modify or delete them. You can also create a new connection by selecting the network to connect to, then the two key columns from the network and from the table. In our case, the network was retrieved from the UniProt identifiers stored in the ‘UniProt’ column of our table. The stringApp stores the query into the column ‘query term’, so we use it as ‘key column from Network’ and we use ‘UniProt’ as the ‘key column from Table’. Figure 3 illustrates the mapping between the two tables. The “P20700” (blue box) from the node table is linked to two rows of the Omics Visualizer table which have the same value in the UniProt column.

The connection between the current network and the current table can be performed with the following automation command:

```
ov connect mappingColNet="query term" mappingColTable="UniProt"
```
Figure 4. STRING network of proteins with significantly regulated phosphorylation sites detected in an ovarian cancer study.\textsuperscript{10} Log-ratios of protein abundance between cancer and healthy tissues were mapped around the nodes using a blue–white–red gradient. Each slice of the rings represents a significantly regulated phosphorylation site. The inner ring is the comparison between epithelial ovarian cancer and distal fallopian tube epithelium (EOC vs FTE), the outer ring is the comparison between epithelial ovarian cancer and ovarian surface epithelium (EOC vs OSE). This subnetwork is the second biggest cluster obtained using Markov clustering on the STRING network of all proteins from the study that have at least one phosphorylation site with an adjusted p-value lower or equal to 0.01.

**Retrieving a STRING network**

It is possible to do the previous step more quickly by automatically retrieving a STRING network from the Omics Visualizer table. We can retrieve a STRING network with the STRING icon from the Omics Visualizer panel (Figure 3, left) or the ‘Apps → Omics Visualizer → Retrieve STRING network’ menu. We have to select the species, the column that contains the identifiers to query, the confidence cutoff and, if filtering was applied as in our case, whether to retrieve the identifiers only from filtered rows. By default the species is human, the cutoff is 0.40, and the query column is identified by a case-insensitive search for “uniprot” among the column names.
Once the STRING network is retrieved thanks to stringApp automation, the network is automatically connected to the Omics Visualizer table with the 'query term' column from the node table with the query column selected by the user. The network can also be retrieved from the current Omics Visualizer table with the following automation command:

```
ov retrieve taxonID=9606 filteredOnly=true queryColumn="UniProt"
```

**Donut visualization**

The donut visualization can be configured via the donut icon from the Omics Visualizer panel (Figure 3, left) or the 'Apps → Omics Visualizer → Create donut visualization' menu.

Here, we want to visualize the two disease vs. healthy comparisons for each site. We select as Values the two comparison columns (EOC vs FTE and EOC vs OSE) that contain the numerical values we want to visualize. Because the values are log-ratios, we apply a continuous mapping and use a color
scale. To be able to identify the individual phosphorylation sites in the network, we label the charts with the position of the amino acid (AA position column) of the site in the sequence of the protein. We customize the chart so that the first slice starts at a 3 o’clock position and the label font size is changed to 15. The next dialog enables the user to configure the color scale. By default, a diverging palette is selected, and the color scales from the minimum to the maximum value and is centered on zero. The minimum and maximum bound values are directly computed from the values of the table: the maximum bound is the maximum of the absolute value of the minimum and maximum data values; the minimum bound is the opposite of the maximum bound. The user can change both the values and the colors associated. In our case, we will change the range and set it from -8 to 8. If a table value is lower than the minimum value of the scale or larger than the maximum, the value will be associated with the color associated with the minimum or maximum value, respectively. The visualization can be modified or deleted by accessing the dialog again. The resulting network can be seen in Figure 4. We have two donuts around each node representing the two comparisons. One slice of a donut is one value from the Omics Visualizer table associated with the node: in our case, it is the log-ratio of a specific site. It is possible to flip the visualization, and have as many donuts as sites, with two slices for each comparison, by selecting “Ring is row” instead of the default “Ring is column” in the last dialog.

All the commands related to the donut visualization are associated with the keyword “outer” since the visualization is around the node. The visualization of the current table can be obtained with the following automation command:

```
ov viz apply outer continuous
attributes="EOC vs FTE,EOC vs OSE"
labels="AA position" filteredOnly=true
rangeMin=-8 rangeMax=8
chartSettings="arcstart:0,labelsize:15"
```

**Pie visualization**

If we are not interested in the individual log-ratios from the comparisons, we can instead summarize the results as a pie visualization of the expression clusters.

The pie visualization can be configured via the pie icon from the Omics Visualizer panel (Figure 3, left) or the ‘Apps → Omics Visualizer → Create pie visualization’ menu. The pie visualization dialog looks similar to one for the donut visualization dialog, and the user must first select the column that contains the values to draw. The difference here is that the user can only select one column, because the different slices of the pie come from the different values associated with the same node. In our case, we want to visualize the “Cluster” column and apply a discrete mapping. As for the donut visualization, we will label the chart slices with the amino acid position (AA position). We also customize the chart to have the first slice starting at a 3 o’clock position and change the label font size to 15. The next dialog enables us to configure the color mapping. By default, Omics Visualizer selects a qualitative palette. Here the user may change the color associated with each value detected in the table. The default colors are too similar to each other and we changed them by clicking on the color square. The pie visualization can also be modified or deleted by accessing the dialog again.

All the commands related to the pie visualization are associated with the keyword “inner” since the visualization is inside the node. The resulting network is shown in Figure 5 and can be obtained with the following command:

```
ov viz apply inner discrete
attributes="Cluster" labels="AA position"
colorMapping="A:#1F78B4,B:#FF7F00,C:#E31A1C" filteredOnly=true
chartSettings="arcstart:0,labelsize:15"
```

**Legend**

The user can generate a legend by clicking the map icon from the Omics Visualizer panel (Figure 3, left), or the ‘Apps → Omics Visualizer → Manage legend’ menu, and choosing which legends to generate (pie and/or donut visualizations), the title, the font, and the position of the legend. Omics Visualizer will then create Cytoscape annotations corresponding to the current visualizations, displaying names of the columns above the color legends. In the case of the donut visualization, a list of column names shows the order of the columns. Because legends are annotations, they can be moved, edited, and exported with the network as an image. The user can modify an annotation by selecting it from the Annotation panel and choosing “Modify Annotation ...” from the contextual menu. Note that the name of an annotation can be the same as its textual content, so the annotation needs to be modified for its content to change. To move an annotation, the user has to activate the Annotation selection (Select → Mouse Drag Selects → Annotations Only).

The legends in Figure 4 and Figure 5 were automatically generated with the following command:

```
ov legend draw position="EAST_TOP"
title=""
```

**Conclusions**

Omics Visualizer is a Cytoscape app that improves how omics data can be visualized on molecular networks. A key feature of Omics Visualizer is that, unlike the standard Cytoscape node table, it enables the user to import files with several rows of data related to the same node, with each row representing a different site, isoform, or experimental condition. This data can subsequently be visualized in or around the nodes as charts with a separate slice for each row that is connected to the node. These slices can be colored to show both continuous and discrete values, and Omics Visualizer can automatically produce a color legend to help in preparation of publication-ready figures. All the actions of Omics Visualizer can be done...
using the Cytoscape GUI or the commands, making it easy to use Omics Visualizer also in R or Python via the Cytoscape REST API.

Data availability

Underlying data
Zenodo: Modified Table S3 - Francavilla et al. 2017. https://doi.org/10.5281/zenodo.3629220

Data is available under the terms of the Creative Commons Attribution 4.0 International.

Software availability

The software, source code, and tutorial are available at the Cytoscape App Store: http://apps.cytoscape.org/apps/omicsvisualizer.

Archived source code at the time of publication: https://doi.org/10.5281/zenodo.3631584

License: BSD 2-Clause “Simplified” License.

Author contributions

JHM and LJJ designed the app. ML is the main developer of the app, helped by NTD. NTD, JHM and LJJ contributed to the manuscript. ML wrote the manuscript.

Acknowledgements

The authors thank Katerina Nastou and Giulia Corsi for beta-testing the app.

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   Publisher Full Text
   Published Abstract | Publisher Full Text | Free Full Text
   Published Abstract | Publisher Full Text | Free Full Text
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   PubMed Abstract | Publisher Full Text | Free Full Text
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Version 1

Reviewer Report 15 April 2020

https://doi.org/10.5256/f1000research.24577.r61035

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Ruth Isserlin

Donnelly Centre for Cellular and Biomolecular Research, Donnelly Centre for Cellular and Biomolecular Research, University of Toronto, Toronto, ON, Canada

Authors Legeay et al. introduce a new Cytoscape app to enable easy import and visualization of multi data type -omics data onto networks with added features to quickly import interaction data from String. A great new addition to the cytoscape app set.

Detailed comments:

In the method section -

“the data can be viewed (but not edited) through the Omics Visualizer panel located next to the node, edge, and network table panels." - not sure what you mean by this. Is it an additional tab in the table panel or is it its own panel next to the table pane? After running the app I see that it is an additional tab. it might sound better as “the data can viewed (but not edited) through the Omics Visualizer panel in the table panel.” (Just for aesthetics I really like it if the app icon can be included in the tab name as well. Makes it easier to spot and click on.)

“but a network can only be connected to one Omics Visualizer table.” - Does this mean that you can't annotate a network with isoform data and phosphorylations at the same time? You can use both pie and donut visualizations at the same time so as long as the different data types are represented in the same table you can use multiple types. Maybe specify that an network can only be associated with one omics set but if you integrate multiple omics data into one table prior to loading into cytoscape you can integrate different data types.

The paragraph starting with “To visualize the data table onto a network,” that outlines how the different tables are connected and the relationships of those connections would be much clearer with a figure outlining all the different connections described.

It would be helpful to have a description of the type of table the app is expecting and how the two different charts can be used. I.e. I loaded in a table with genes vs. Patients expecting to see all my
patient data on the node but I think that it only maps one column for the pie chart and multiple columns for the donut so the expectation is that there will be duplicate genes in a given column for the pie.

What is the maximum number of conditions/instances that can be mapped to an individual node and visualization type?

“The enhancedGraphics continuous mapping always ranges from minimum to zero, and zero to maximum.” - are minimum values calculated based on the filtered set of -omics data or the entire dataset loaded?

“Omics Visualizer can automatically generate legends from the visualizations in the form of Cytoscape annotations, which can be exported as part of the images if the user exports the network.” - it was very difficult to find the legend. I had to come back to the text a few times to figure out that it was somewhere on the network canvas and compared to the network it was tiny so it was hard to find.

**Use cases:**

“To reduce the size of the network, we cluster it using the Markov clustering from the clusterMaker2 app” - in the clustermaker2 app Markov clustering is called MCL. Might be good to add MCL in brackets for users who don't know they are equivalent.

“For illustrations purposes, we here show only the second biggest cluster, which contains 40 nodes and 107 edges.” - it is not clear how you limit to just that cluster. Do you filter the network by _mclCluster or do you create a network of the clusters directly from the cluster maker2 app?

“It is possible to flip the visualization, and have as many donuts as sites, with two slices for each comparison.” - how could this be done? Does it involve changing our choice of column to EOC and FTE and OSE separately?

Inner and outer visualization can be confusing. At first I thought you were referring to the inner and outer rings of the donut. Maybe changing or adding Pie and Donut to Outer visualization and Inner visualization headers will be helpful.

“The difference here is that the user can only select one column, because the different slices of the pie come from the different values associated with the same node. “ - This is an important distinction that needs to be specified earlier. Whether it is in the introduction or in the methods but I think that the different type of data Omics is expecting and the data types best used for each visualization types needs to be expanded on sooner in the paper.

In the inner visualization section - “In our case, we want to visualize the “Cluster” column and apply a discrete mapping.” - it is unclear what the meaning for this “Cluster” column that was in the initial file loaded.

In the legend section - “Because legends are annotations, they can be moved, edited, and exported with the network as an image.” - how can the legends be edited or moved? I tried right clicking on the legend and selected Edit -> modify annotation but nothing happened. I figured out
that you modify the annotation through the annotation panel but might be good to add how you do it here. (I couldn't figure out how to move it though)

**Minor grammatical suggestions:**

“For illustrations purposes, we here show only the second biggest cluster, which contains 40 nodes and 107 edges.” - remove the “here” “For illustrations purposes, we show only the second biggest cluster, which contains 40 nodes and 107 edges.”

“The visualization is configured thanks to a specific dialog” - not sure what this is saying.

“The inner visualization can be configured thanks to the pie icon or the ‘Apps → Omics Visualizer → Create pie visualization’ menu. “ - sounds better as “The inner visualization can be configured using the pie icon or the ‘Apps → Omics Visualizer → Create pie visualization’ menu.”

**Is the rationale for developing the new software tool clearly explained?**
Yes

**Is the description of the software tool technically sound?**
Yes

**Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?**
Yes

**Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?**
Yes

**Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** bioinformatics

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

Author Response 16 Jun 2020

**Marc Legeay,** Novo Nordisk Foundation Center for Protein Research, University of Copenhagen, Copenhagen, Denmark

We would like to thank you for your comments and suggestions. We have addressed them in the new version of the manuscript, which also describes a new feature added to the app
during review, namely to import columns from the node table of a network loaded in Cytoscape. We also added two new figures to describe the general Omics Visualizer workflow and to clarify the format needed for input tables.

We took into account all your comments regarding the phrasing. For other comments, we give our answers below (we labeled your comments with “Reviewer” and ours with “Response”).

Reviewer: “but a network can only be connected to one Omics Visualizer table.” - Does this mean that you can’t annotate a network with isoform data and phosphorylations at the same time? You can use both pie and donut visualizations at the same time so as long as the different data types are represented in the same table you can use multiple types. Maybe specify that an network can only be associated with one omics set but if you integrate multiple omics data into one table prior to loading into cytoscape you can integrate different data types.

Response: Right now, a network can only be connected to one Omics Visualizer table, so indeed if the user wants to have both visualizations, all the data has to be merged into one table (see new Figure 2C). We are currently working on a new version of the app that would enable a network to be connected to several tables.

Reviewer: It would be helpful to have a description of the type of table the app is expecting and how the two different charts can be used. I.e. I loaded in a table with genes vs. Patients expecting to see all my patient data on the node but I think that it only maps one column for the pie chart and multiple columns for the donut so the expectation is that there will be duplicate genes in a given column for the pie.

Response: We added a new Figure 2 giving three examples of tables that could be imported into Omics Visualizer. We hope it makes things easier to understand.

Reviewer: What is the maximum number of conditions/instances that can be mapped to an individual node and visualization type?

Response: There is no technical limit to the number, and it is thus up to the user to decide how many slices or rings to make. However, to create easily understandable visualizations, we would advise against having excessively many slices or rings.

Reviewer: “The enhancedGraphics continuous mapping always ranges from minimum to zero, and zero to maximum.” - are minimum values calculated based on the filtered set of omics data or the entire dataset loaded?

Response: The user can put a minimum and maximum value, all values below the minimum will be colored as the minimum, and all values above the maximum will be colored as the maximum. Those values are, by default, the min and max of the filtered values. The color gradient from enhancedGraphics ranges from a negative value to zero, then zero to a positive value. Thus, Omics Visualizer has to change the table values to center the data. We modified the manuscript accordingly.
**Reviewer:** “Omics Visualizer can automatically generate legends from the visualizations in the form of Cytoscape annotations, which can be exported as part of the images if the user exports the network. “ - it was very difficult to find the legend. I had to come back to the text a few times to figure out that it was somewhere on the network canvas and compared to the network it was tiny so it was hard to find.

**Response:** By default, the font size of the legend is the same as the font size of the node labels, meaning that if you have a huge network, the legend with default font size can be hard to find. When the legend is generated, we change the view of the network to fit everything (including the legend).

**Reviewer:** “For illustrations purposes, we here show only the second biggest cluster, which contains 40 nodes and 107 edges.” - it is not clear how you limit to just that cluster. Do you filter the network by _mclCluster or do you create a network of the clusters directly from the cluster maker2 app?

**Response:** We enabled the option “Create new clustered network” when we ran clusterMaker and extracted the second largest cluster by selecting all nodes and edges belonging to it and creating a new network from the selection.

**Reviewer:** “It is possible to flip the visualization, and have as many donuts as sites, with two slices for each comparison.” - how could this be done? Does it involve changing our choice of column to EOC and FTE and OSE separately?

**Response:** No, you can keep the same columns, it is just a matter of whether each ring represents a condition (“ring as column”) or a site (“ring as row”).

**Reviewer:** In the legend section - “Because legends are annotations, they can be moved, edited, and exported with the network as an image.” - how can the legends be edited or moved? I tried right clicking on the legend and selected Edit -> modify annotation but nothing happened. I figured out that you modify the annotation through the annotation panel but might be good to add how you do it here. (I couldn’t figure out how to move it though)

**Response:** We added an explanation on how to move and change the text of the legend using Cytoscape’s Annotations control panel.

**Competing Interests:** No competing interests were disclosed.
Magnus Palmblad
Center for Proteomics and Metabolomics, Leiden University Medical Center, Leiden, The Netherlands

To evaluate the user experience, I first downloaded and used the Omics Visualizer, following the on-line tutorial. The interface is intuitive (like how Cytoscape itself works), exposing just the right amount of information and options to the user. It is also clear to the user what the app does.

The manuscript partially overlaps with but complements the on-line tutorial with background and some technical detail how the app integrates in Cytoscape. The Omics Visualizer fills a clear niche in the Cytoscape appverse, allowing tables to be imported surjectively (mapping multiple rows in the input to the same network node) as opposed to bijectively (with a one-to-one correspondence between the rows in the input table and nodes in the network). I suppose this could be achieved by manipulating tables (connecting row information in the data to columns in a node table) in R or Python, pushing the necessary data to Cytoscape through RCy3 or py2cytoscape (Omics Visualizer commands are also exposed via the Cytoscape REST API). But the Omics Visualizer is completely interactive and requires no scripting or formulae to be entered in the Cytoscape tables. The menus are clear and self-explanatory. The visualizer provides pie and donut visualizations for displaying multiple node attributes, such as post-translational modification sites and occupancy onto the same node. As would be expected from this team, Omics Visualizers also works well with the Cytoscape STRING app. I find this manuscript a very valuable companion to the app and on-line tutorial.

The manuscript is very clearly written, and the length appropriate for the subject matter. I only have two minor comments:

1. "3 o'clock angle" should be "3 o'clock position" (in the "Outer" and "Inner visualization" paragraphs)?

2. The "Filter table rows" option referred to in the on-line tutorial is called "Filter table" in Cytoscape.

Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Yes

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Yes

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Yes

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Analytical chemistry, molecular biology, mass-spectrometry based proteomics, bioinformatics.

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

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**Author Response 16 Jun 2020**

Marc Legeay, Novo Nordisk Foundation Center for Protein Research, University of Copenhagen, Copenhagen, Denmark

We would like to thank you for your comments and we have modified the manuscript to take them into account. In the meantime, we also released a new version of the app with the additional feature to import columns from the node table of a network loaded in Cytoscape. We thus further modified the manuscript to describe the new feature. We also added two new figures to describe the general Omics Visualizer workflow and to clarify the format needed for input tables.

**Competing Interests:** No competing interests were disclosed.

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