PathRings

Version 1.0 Manual

PathRings

PathRings is a web-based application for visualization and analysis of biological pathways. Users can visualize and search pathways from Reactome, as well as perform ortholog and gene expression analysis by uploading orthologous and gene expression data. This project is based on the PathBubbles project: http://sites.google.com/a/umbc.edu/pathbbubles/. The recommended web browser to use is Chrome.

PathRings is freely available at http://raven.anr.udel.edu/~sunliang/PathBubbles/. All code is available at Github here: https://github.com/ivcl/PathBubbles.

PathRings was submitted to Bioinformatics:

Yongnan Zhu, Liang Sun, Carl Schmidt, Keqin Wu, Zhigeng Pan, Jinglong Fang, J. C. PathRings: an web-based tool for Exploration of Ortholog and Expression Data in Biological Pathways[J] (submitted to BMC Bioinfomatics)

1. Overview

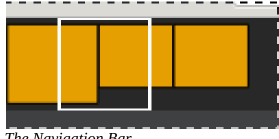


Figure 1: An overview of the PathRings interface. 1) The navigation bar. 2) The current view box. 3) The main workspace. 4) A sunburst (or treering) pathway visualization bubble. 5) A table bubble.

PathRings supports data exploration of four main types of pathway relationships. These are hierarchical, cross-talking, orthologous, and gene expression relationships. Each functional view is presented in a "bubble". These bubbles may be grouped together. Figure 1 provides an overview of the PathRings interface. At the top is a navigation bar (1) which provides an overview of the current layout, and along with (2) may be used to quickly scroll to different sections. The main workspace (3) is where the various bubbles are laid out. (4) and (5) are examples of two different bubble types.

2. Navigation Bar

The navigation bar spans the top of the PathRings interface. Each currently open bubble is displayed on the bar in a simplified form. The current view is represented with a white box, which may be dragged to change the current view.



The Navigation Bar

3. Main Workspace

Below the navigation bar is the main workspace, which is the main display area. Right-clicking on any open spot in the main workspace will bring up a context menu with the following options:

- Open Entire Pathway: Creates a new, default pathway bubble.
- Delete All: Will delete all bubbles.
- Open Help: Opens this document.

4. General Bubble Mechanics

All bubbles share a few common traits, and are manipulated in a similar manner. While the mouse is over an outside edge of a bubble, the cursor will change. Clicking and dragging will allow the user to resize the bubble. Clicking and dragging anywhere else on the bubble's border, including the title, will allow the user to move the bubble. Each bubble also has several buttons in the upper-right corner. All will have the 'X' button, for deleting the bubble, and the 'U' button, for ungrouping it. The 'M' button may be present as wello which will open another menu for further operations.

Bubbles may be grouped and moved as one unit. This is done by dragging and dropping one bubble ontop of another, or done implicitly when a bubble is created from a parent bubble. Bubbles may then be ungrouped by clicking and gragging the bubble, starting from the 'U' button in the top rigth corner.

4. Pathway Bubble

4.1 Main Bubble

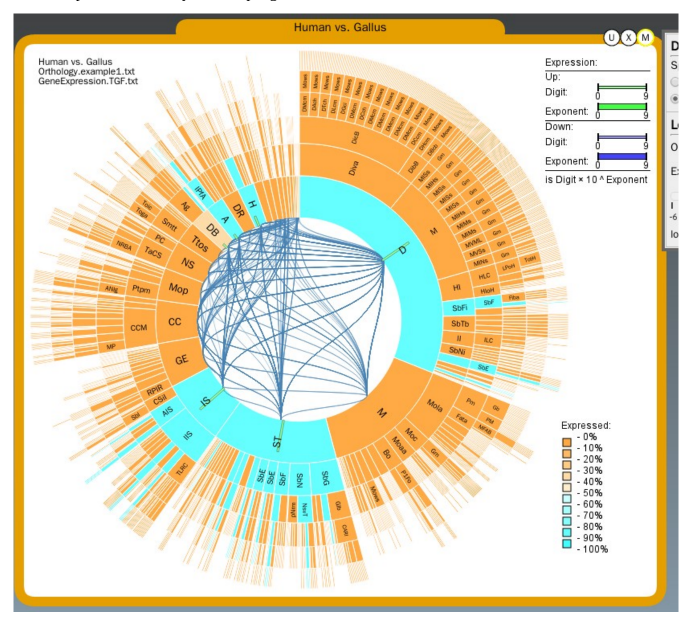


The Pathway Bubble visualizes up to the entire set of Reactome pathways. In the center of the bubble is a list of the top-level pathways being shown. There is a line drawn from each of these pathways to its actual representation in the graph. Hovering over the pathway name with the mouse will highlight this linking line, so that the actual pathway may easily be distinguished. Using the menu, you may opt to hide this information and instead display lines showing crosstalk between the different pathways.

The main visualization has a different meaning depending on whether or not expression data has been uploaded. Before it has been uploaded, each pathways is colored according to how closely the selected species or uploaded ortholog data matches that of humans. The key can be seen in the lower-

right, distinguishing between Complete, Partial, and Empty pathways. A Complete pathway has all of its genes also found in humans, and an Empty one has none.

The actual bars in the chosen pathway level show how many crosstalk genes are present in the given pathway. This is presented using a split vector in order to improve readability. The length of the thicker bar represents the order of magnitude of the value, and the thinner bar is the value of its leading digit. This is explained in the key in the top-right.



Once an expression file has been uploaded, the visualization changes. Each pathway is now color coded by what percentage of genes in that pathway is up or down expressed, as seen in the bottom-right. Each pathway in the selected level will show up to two different bars — one showing the number of up expressed genes and one showing the number of down expressed. These are displayed in the same split format and are explained in the upper-right.

The upper-left corner displays the currently selected species, as well as any loaded ortholog or

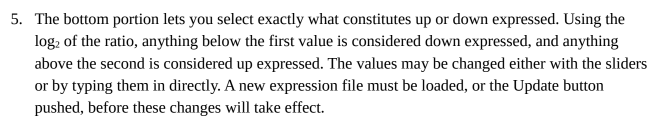
expression files.

The actual diagram itself can be clicked on to bring up a new bubble. Left clicking on a pathway section will bring that pathway section up in a new pathway bubble. Right clicking on a pathway section will bring up a new table bubble listing all of the components of that pathway. The crosstalk or expression bars may also be right clicked – in this instance, the resulting table bubble will only have those components which are part of the displayed bar. If crosstalk links are currently visible, moving the mouse over an expression or crosstalk bar will highlight those links. Any of these newly created bubbles will be visibly linked to the original bubble.

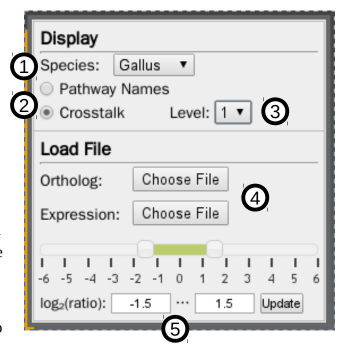
4.2 Bubble Menu

The menu opened with the circular 'M' button in the top right corner is used to load data files and otherwise manipulate the display:

- 1. This selects a species to compare vs. humans.
- 2. This controls what is displayed in the center of the diagram. Either the list of pathway names, or links showing crosstalk between pathways.
- 3. This selects at which level of the graph the bars showing crosstalk or expression are displayed, as well as what level the crosstalk links connect to if those are selected. Level 1 is the innermost and highest ring displayed. Higher levels move towards the edge.
- 4. This lets you choose an orthology or expression file to load.



In addition, any changes to species, ortholog, or expression data or display will be propogated to any linked bubbles as well.



5. Table Bubble

pathwayID	proteinID	uniprotID	symbol	displaySymbol	reactomeID	cellularLocation
3296482	Protein1	P27352	<u>GIF</u>	GIF	2980918	extracellular region
3296482	Protein10	P00167	CYB5A	CYB5A	198781	mitochondrial outer membrane
3296482	Protein100	Q05932-2	FPGS	FPGS-2	197968	cytosol
3296482	Protein101	Q9H2D1	SLC25A32	SLC25A32	200645	mitochondrial inner membrane
3296482	Protein102	Q05932-1	<u>FPGS</u>	FPGS-1	197981	mitochondrial matrix
3296482	Protein103	P11586	MTHFD1	MTHFD1	200641	cytosol
3296482	Protein104	P34896	SHMT1	SHMT1	71242	cytosol
3296482	Protein105	P42898	MTHFR	MTHFR	200663	cytosol
3296482	Protein106	Q9NZB8-1	MOCS1	MOCS1B	947547	cytosol
3296482	Protein107	Q9NZB8-5	MOCS1	MOCS1A	947532	cytosol
3296482	Protein108	095396	MOCS3	MOCS3	947570	cytosol
3296482	Protein109	095396	MOCS3	CysS-MOCS3	947553	cytosol
3296482	Protein11	P00387	CYB5R3	B5R	198771	mitochondrial outer membrane

The table bubble is comparatively simple. The columns may be left clicked to sort the table by that column. Clicking again will cycle through ascending and descending orders. Elements in the symbol column may be left clicked to bring up an NCBI information pane on them. Right clicking on a symbol element will bring up a new table bubble composed of just that symbol. If the crosstalk column is present, its entries may be clicked to highlight those crosstalking events in an attached Pathway Bubble, if possible.

6. Data Format

This section describes the file formats used in the uploading option for the Pathway Bubble.

6.1. Ortholog Gene Format

Your own ortholog gene data may be uploaded. It should be a tab-delimited, two column text file. The first column is the gene symbol, and the second is that gene's orthologous relationship between this species and human. If this gene cannot be found in any human pathway, \N should be used. Otherwise, it should be the Reactome human database identification number of that gene.

Symbol	DbId
ADA	\N
CDH2	414745
AK13	421497
MED6	426282
NRE3	39528

Table 1: Ortholog Data Format Example

6.2. Expression Data Format

Your own gene expression data may be uploaded as well. It should be a tab-delimited, three column text file. The first column is the Entrez gene identification number, the second is the gene symbol, and the third is the expression ratio.

Gene Id	Symbol	Ratio
374096	SMAD6	0.6892992938620315
395132	SMAD3	7.128911138923654
395247	SMAD2	3.370212765957447
395543	SMAD9	Infinity
395679	SMAD5	3.013579576317219

Table 2: Expression Data Format Example