SOFTWARE

Open Access



WebGIVI: a web-based gene enrichment analysis and visualization tool

Liang Sun^{1,6}, Yongnan Zhu^{2,3}, A. S. M. Ashique Mahmood⁴, Catalina O. Tudor⁴, Jia Ren⁵, K. Vijay-Shanker⁴, Jian Chen² and Carl J. Schmidt^{1*}

Abstract

Background: A major challenge of high throughput transcriptome studies is presenting the data to researchers in an interpretable format. In many cases, the outputs of such studies are gene lists which are then examined for enriched biological concepts. One approach to help the researcher interpret large gene datasets is to associate genes and informative terms (iTerm) that are obtained from the biomedical literature using the eGIFT text-mining system. However, examining large lists of iTerm and gene pairs is a daunting task.

Results: We have developed WebGIVI, an interactive web-based visualization tool (http://raven.anr.udel.edu/webgivi/) to explore gene:iTerm pairs. WebGIVI was built via Cytoscape and Data Driven Document JavaScript libraries and can be used to relate genes to iTerms and then visualize gene and iTerm pairs. WebGIVI can accept a gene list that is used to retrieve the gene symbols and corresponding iTerm list. This list can be submitted to visualize the gene iTerm pairs using two distinct methods: a Concept Map or a Cytoscape Network Map. In addition, WebGIVI also supports uploading and visualization of any two-column tab separated data.

Conclusions: WebGIVI provides an interactive and integrated network graph of gene and iTerms that allows filtering, sorting, and grouping, which can aid biologists in developing hypothesis based on the input gene lists. In addition, WebGIVI can visualize hundreds of nodes and generate a high-resolution image that is important for most of research publications. The source code can be freely downloaded at https://github.com/sunliang3361/WebGIVI. The WebGIVI tutorial is available at http://raven.anr.udel.edu/webgivi/tutorial.php.

Keywords: Visualization, eGIFT, Gene iTerm, Gene enrichment, Web development

Background

High-throughput technologies provide biologists with large lists of genes or proteins when they compare expression data between two biological states (e.g., normal tissue vs. cancer tissue). Grouping enriched genes to known biological processes and pathways is a common strategy for understanding the biology that underlies the differences between the two states. Approaches include GO enrichment analysis such as DAVID [1, 2], GOEAST [3] and Gorilla [4], and pathway analysis such as KEGG [5] and Reactome [6].

Full list of author information is available at the end of the article



eGIFT [7] uses a text-mining method to identify informative terms (iTerms) for individual genes. iTerms are not limited to gene ontology (GO) terms; they also capture more detailed biological knowledge. Consequently, eGIFT provides a finer grained interpretation of gene lists than GO analysis. The current gene analysis results of eGIFT provide users with a list of ranked iTerms and their associated genes in a tabular format. A graphic representation of these gene and iTerm relations would allow biologists to better interpret their input gene lists or gene-iTerm pair lists. This often captures the biological concept enriched in the input data.

Visualization tool

An effective visualization of large data sets can provide biologists with means to discover buried relationships in complex data sets. Currently, several different visualization



© The Author(s). 2017 **Open Access** This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

^{*} Correspondence: schmidtc@udel.edu

 $^{^{1}\}mbox{Department}$ of Animal and Food Sciences, University of Delaware, Newark, DE, USA



tools are used to capture the relationship between genes, protein and networks, such as Arena 3D [8], Medusa [9], Ondex [10], Osprey [11], Pajek [12], BioLayout Express^{3D} [13], Cytoscape [14] and ProViz [15]. However, most of these tools need to be installed on a local computer and require plugins or third party software such as Java Runtime Environment. Installation and maintenance of such tools can be difficult for those unfamiliar with computer system administration. In contrast, web-based tools offer the

ability to visualize relationships without the overhead of having access to system administrators. Cytoscape.js [16] and D3.js [17] are the most popular visualization JavaScript libraries that can be applied to visualize network and biomolecule interactions. There are several successful web based visualization tools that use the Cytoscape and D3 JavaScript libraries in a biological context. For example, BNVC [18], a web-based visualization tool of biomolecular networks, can be used to compare two similar networks. PINV [19], also a web-based protein-protein interaction network visualizer, provides complex interaction networks with the ability to query, filter and group data of interest. However, complex graphs rendered by such tools are difficult to interpret when analyzing data with hundreds of nodes and edges. WebGIVI addresses the issue of visualizing large data sets by adopting a concept map method for visualization. The concept map aligns nodes on different layers and automatically calculates the distance between layers. This maximizes the amount of information that can be displayed despite limited physical size of the screen. In addition, a zoom-in feature of Concept Map allows the user to scale the graph for a better view. Now, a user can analyze hundreds of nodes in a high-resolution image produced by WebGIVI that can be saved and is suitable for publication.

Interactive WebGIVI provides an integrated graph to help users generate biological hypotheses. A database of rate-limiting genes, identified by text mining and manually



To the right of the page is a short video demonstrating various WebGIVI functionalities

verified, was also integrated into WebGIVI. All genes encoding rate-limiting products are colored in the visualization. Expression data that is uploaded to WebGIVI can also be color-coded according to expression levels in two different biological states. These features are accomplished by using PHP, Cytoscape.js and D3.js to generate a powerful and interactive web based visualization tool that implements gene enrichment analysis and gene and iTerms visualization.

Implementation

The flow chart of the data processing is depicted in Fig. 1. Currently, WebGIVI accepts multiple input data formats including: NCBI Entrez gene ID, UniProt [20], or Ensemble [21] Gene IDs. Prior to accessing WebGIVI, the user will have identified genes of interest in their biological system. The WebGIVI input list is used to retrieve the gene symbols and iTerms through an eGIFT API. In the retrieved list of gene symbol-iTerm pairs, the first column contains the gene-associated iTerms. Interactive WebGIVI also supports uploading and visualization of any two-column tab separated data. For

instance, gene symbols and related pathways data can be visualized.

WebGIVI has three major functions:

- 1. *Visualization*. Interactive table visualization includes sorting and deleting functions that help users reduce the graph to only iTerms and genes of interest. GUI functions in the Concept Map visualization include switching, filtering, sorting, searching, grouping and saving the data in tab delimited text files or as high-resolution images.
- 2. Integration of Rate-limiting and Expression Data. Rate limiting gene products determine the flow rate of metabolites or signal through pathways. We have identified 150 rate limiting gene products via text-mining of primary literature and biochemistry textbooks. This list was manually verified (CJS), and integrated into WebGIVI.

When a user enters an expression data list, genes that encode a rate limiting gene product are colored when visualizing gene:iTerm pair in the concept map. If such genes are differentially regulated they are likely have a significant impact on the overall rate of metabolite or signal flow through their

WebG	\overline{IVI} tool				Visualization of Gene and iTerm
Home	eGIFT	Tutorial	Contact		
nalysis Result:	ered data is larger th	an 1000 lines, nlease us	e concent man to vis	suallize!	
ort: p-values	Select: frequency <=		election: Clear Se		
		le blacklisted iTerms:	Reset table View	Download Concept Man View Cutosca	View.
			Reset table		
membran (no categor	ry)	stre (no category)		chaperon active (no category)	oxid (no category)
chaperon (no category)		cycl (no category)		hippocamp (no category)	missense mutat (no category)
stre protein (no category)		cli (no category)		gene (no category)	hormon (no category)
sever (no category)		hsc70 (gene/protein	1)	hsf1 (gene/protein)	p=0.0000905355411887 heat-shock (no HSP25
nonsense mutat (no c	category)	oxide stre (no categ	jory)	phenotyp (no category)	HSPA2 plasma membra _{HSPB8} ategory)
presynapse (no categ	gory)	splice vary (no cate	gory)	stre response (no category)	SERPINH1 cochaperone (no category)
affin (no category)		chaperone complex	(gene/protein)	heat stress (no category)	heat-inducible (no category)
protein homeostasis	(gene/protein)	contracture (medica	al condition)	triple helix (no category)	cytoprotection (function/process)
small heat-shock (no category)		shsp (gene/protein)		aggregate (function/process)	traffic (no category)
huntingtin (no category)		auxiliary subunit (no	category)	shock transcript (function/process	s) thermus (no category)
retin (no category)	retin (no category) n-terminus (no category)		gory)	shock response (no category)	synapse (cellular component)
zone (no category)		constitute express (function/process)	cytoprotective (no category)	raf1 (function/process)
misfold protein (gene	/protein)	catalyze (function/p	rocess)	agonist active (function/process)	protein aggregate (gene/protein)
secret (function/proce	ess)	modulate (function/	process)	primary culture (no category)	pyridox (no category)
channel (gene/proteir	n)	tetratricopeptide rep	peat (function/process)) steroid (pharmaceutical)	mitochondrium enzyme (no category)

Fig. 3 Results pages containing iTerms enriched in the input gene list. In this case the list was sorted by *p*-values as determined by Fisher's exact test. One can use the Frequency text box to eliminate iTerms that have fewer genes associated with them based on the user's cutoff. The Selection buttons allow one to "Select All", Clear the selections or Toggle selected iTerms. Also, individual iTerms can be selected by clicking. The selection features are used to remove irrelevant iTerms based on the users knowledge. One can include blacklisted iTerms (see text) by selecting the check box. The Reset Table button can be used to recreate the starting table should errors be made while deleting iTerms thought to be irrelevant. The Concept Map View and Cytoscape View open new windows with the corresponding representations of the iTerm:gene pairs. Hovering over an iTerm box will display the genes associated with that iTerm from the input list along with the *p*-value of that iTerm

respective pathways. Furthermore, to emphasize certain features of their data, users can upload expression files that highlight nodes that they consider informative.

3. *Link-out Functions*. Nodes and edges provide link outs to gene specific, iTerms or gene:iTerm pairs via the NCBI, UniPROT, or eGIFT database. For example, gene-iTerm pairs link to eGIFT sentence web page, which contain sentences with the gene and iTerm highlighted in the text.

Results and discussion

а

Data filtering functions

Sorting Functions: eGIFT uses a precomputed textmining database that has extracted all gene associated informative terms (iTerm) from PubMed abstracts. After submitting a gene list to WebGIVI, a table is returned to the user containing the iTerms associated with the input genes. By hovering over an iTerm, the user can see the genes associated with that iTerm. The default list is

Contac

sorted based on the Fisher's exact test p-value, but the user can choose to sort based on alphabetical order, the gene ontology group (process, function, compartment or unclassified) to which the iTerm has been classified or the frequency of appearance of each iTerm.

Editing Functions: Not all iTerms are informative in all use cases, but could be important to others. For example, 'in situ hybridization' is an irrelevant iTerm to our use case scenario but will be interesting to researchers who might want to apply this experimental method to their own work. However, some iTerms are highly likely to be non-informative. To remove such iTerms a "blacklist" has been developed that includes terms such as "some cell" or "10 fold" that are typically non-informative to the general WebGIVI user. Since the developers of WebGIVI cannot be certain that a given iTerm is irrelevant to all users, the returned iTerm list includes the blacklisted terms; a checkbox is provided that allows the user to hide any terms that are included in the blacklist. It is also beneficial to the user to also be



able to filter out irrelevant iTerms in the context of their study, and only save iTerms of direct relevance. Once data is submitted on WebGIVI's homepage, the returned list will allow the user to delete iTerms from the results table using deleting functions. If the user prefers, they can choose not to prefilter but visualize data in Concept Map or Cytoscape directly.

A biological use case scenario

We used the Sun et al. white-leghorn hepatocellular (LMH) cell heat stress dataset [22], which is a RNA-Seq study of LMH cells under heat stress. This study identified a total of 235 up-regulated and 578 down-regulated genes. Figure 2 shows a completed WebGIVI submission

page with a portion of the regulated genes from the LMH study (Additional file 1). In this case we used Entrez gene identification numbers. Following submission, an iTerm list (Fig. 3) is returned that can be sorted alphabetically, by frequency, Gene Ontology categories, or by p-value as determined by the Fisher's exact test. In this case the list is sorted by p-value. Hovering over an iTerm will show the corresponding p-value, along with the genes from the list associated with that iTerm. One can choose to display iTerms that have been blacklisted by checking the "Include blacklisted items". You can also select irrelevant iTerms by right clicking and delete them using the remove options. Users can view the output in either Cytoscape (Fig. 4) or as a Concept Map



Fig. 5 Concept Map view of iTerm:gene pairs. Genes from the input list with iTerms are in the vertical column at the center and iTerms are around the outside. The panel to the left is the legend explaining the graph, while the panel to the *right* allows manipulation of the output. This allows the user to switch the positions of the genes and iTerms, Search for specific genes or iTerms and to toggle the legend window on or off. The Load Expression File allows the user to load a file that indicates if a gene was up or down regulated in the experiment. *Yellow* corresponds to up while *green* to down regulated genes. The Save button allows the user to save either the graph text as a tab delimited two-column files or save the image. The Filter button allows the user to choose the minimum number of edges that a node must have to be visualized. See Fig. 6 for explanation of Subset Data

(Fig. 5) by selecting the appropriate buttons. The default mode in Cytoscape generates a force graph (Fig. 4a) and clicking on an edge connecting a gene product to an iTerm pops up a window that allows the user to connect to either NCBI or eGIFT (Fig. 4b). Additional view modes include tree or circle that are accessible by the Layout button.

While Cytoscape can be useful for small graphs, the Concept map view is generally easier to use for larger input data sets. In the default view, gene symbols are displayed in a column at the center of the view, while iTerms are displayed as a wheel around the gene symbols (Fig. 5). If necessary, additional layers are automatically added to display more gene:iTerm relations. Several attributes are visible in the Concept Map view. In this case we have uploaded a file indicating how the gene was regulated by heat stress using the Load Expression File button. Genes highlighted in yellow are enriched under heat stress while those in green are enriched in the control (thermoneutral) samples. In addition, genes encoding rate limiting gene products are indicated by a blue rectangle added to the symbol.

In the concept map view users can right click to select either genes or iTerms. Selecting one will create a gene:iTerm edge (Fig. 6) then clicking on the Subset Data button will create a new concept map with just the selected gene and iTerms (Fig. 7). This is useful to allow an investigator to link genes with similar iTerms for subsequent investigation. At any point, right clicking on an active edge will open a window that can be used to connect to PubMed, UniProt or eGIFT (Fig. 8). Linking out to NCBI database will search that database with the gene and iTerm, and retrieve links to abstracts that contain those two search terms. For example when searching for a gene:iTerm pair such as HSP90AA1 and the iTerm "chaperone" the search will be in the syntax "HSP90AA1 AND chaperone" and the results will





include all abstracts that include both the HSP90AA1 and chaperone. Linking to UniProt will access a search page allowing the user to view the UniProt entry for the gene product. Linking to eGIFT web pages provides users with the sentences extracted from the literature that contain the gene and iTerm pair (Fig. 9). This feature of WebGIVI greatly facilitates the user's understanding of the gene product's function. In addition, the PMID under a sentence links to the PubMed abstract pages from which the sentence was extracted. This further aids in placing the gene product in biological context. The ability to provide users with these sentences is a unique facet of WebGIVI functionality.

Interactive WebGIVI also provides a way to save the current view: users can export all data on the Concept Map as a two-column tab separated file, which can be resubmitted in WebGIVI to obtain the same graph. This feature allows a user to readily share the data with collaborators. Users can also export the graph as a scalable vector graphics (SVG) format, which can be transformed to high resolution image types via readily accessible image conversion web sites.

Comparison of WebGIVI, DAVID and AmiGO2 analysis

Incorporating the text-mining tool, WebGIVI, into the analysis of high-throughput transcriptome experiments complements Functional Annotation clustering provided by DAVID and GO analysis provided by AmiGO2 [23]. DAVID is an online knowledgebase that can output a list of enriched biological concepts from an input list of gene identifiers. The site makes use of multiple resources including the Gene Ontology, the KEGG pathway database, Interpro [24], along with several others. Amigo2 is a product of the Gene Ontology consortium and allows users to submit gene lists and identify enriched GO terms. We chose to compare WebGIVI with DAVID and Amigo2 because these resources are easy to use and have been widely adopted by the scientific community.

Heat stress has been implicated in affecting cell cycle regulatory processes including DNA synthesis, DNA repair, cell cycle checkpoints, cell proliferation, and spindle formation. The objective was to compare the ability of DAVID, AmiGO2 and WebGIVI to identify genes affecting cell cycle regulation that are up-regulated by heat



stress in the liver of chickens (Jastrebski et al. manuscript submitted). Genes whose expression was increased by heat stress were identified and submitted to Amigo2, DAVID, and WebGIVI for comparative analysis. Genes recognized by each analytical method as associated with the concept of cell cycle regulation (including cell cycle, DNA replication, DNA repair, checkpoint) were considered in this comparison of methods (see Additional file 2 for complete list of identified genes).

In combination DAVID, AMIGO2 and WebGIVI identified a total of 214 genes affecting cell cycle regulation as enriched by heat stress in the liver. WebGIVI identified the largest percentage of total genes (80%) and uniquely identified the greatest percentage of genes (30%) not recognized by either DAVID or AMIGO2 (Table 1). However, WebGIVI missed 7 genes that were captured by either DAVID or Amigo2 (Fig. 10). Taken together, this analysis indicates that multiple approaches are best for categorizing the biology embodied in gene lists, and that WebGIVI provides an important contribution to these analyses.

Future work

WebGIVI can be used to visualize not only eGIFT data, but also any two-column relationship data. For example, microRNA and target genes, kinase and substrate pairs, and protein-protein interactions could all be visualized with WebGIVI. Our tool can also be used as an extension for other tools. Compared to other visualization tools, our web-based gene iTerm visualization tool is highly customizable. Users can easily upload their own data and edit their data in the graph. No pre-installed or third party software such as Java Runtime Environment is required to visualize users' data. While we are developing a WebGIVI blacklist of iTerms we believe are not informative, users still need to examine all iTerms and remove ones they find uninformative manually. This inability of WebGIVI to learn as user's preferences and

		Home	Gene Search	Gene Analysis	/Term Search	Add Gene	Page Guide	Fee
Sent	ences containing "chaperon active" in the	BAG3	abstracts		see iTerms	see docum	ents for this /T	erm
	to select a particular species, use the drop-down	IIST DÉIC	JW.					
	select species							
1	BAG3 protein binds to and regulates Hsp70 c	hapero	ne activity .					
	PMID 17974966 - mice, human							
	Hand the second set it is to set (0)	-conta	ining proteins	depends on its	association wi	ith Bag3 , a	a stimulator o	of
2	macroautophagy .							

automatically remove iTerm provides room for future improvement. Potentially, machine-learning techniques could be applied to remove such iTerms. Future work will also integrate more bioinformatics databases such as protein kinase database and transcriptional factor database to enable the user to discover more interesting biological relationships and improve the usability of WebGIVI.

Conclusions

Interactive WebGIVI tool provides an integrated visualization and gene enrichment analysis tool. It helps biologists to visualize genes and iTerms online, makes sense of their biological data, and is useful to generating biological hypotheses from high throughput data.

Availability and requirements

Project name: WebGIVI: Web-based Gene and Iterm Visualization Tool

Table 1	Comparative	analysis	of WebGIVI,	DAVID	and Ar	miGO2
---------	-------------	----------	-------------	-------	--------	-------

Analysis tool	Percent of total	Percent unique
WebGIVI	80%	30%
DAVID	42%	6%
AmiGO2	50%	11%

The three tools identified a total of 214 genes associated with the biological concept of cell cycle regulation" (see text). The numbers indicate the percentage of the 214 genes identified by the different tools along with the number of genes uniquely identified by the corresponding tools. In this analysis, WebGIVI identified 64 (30%) genes as associated with cell cycle regulation that were not associated with this concept by either DAVID or AmiGO2 (see Fig. 10)

Project home page: http://raven.anr.udel.edu/webgivi/ Source code: https://github.com/sunliang3361/WebGIVI Operation system(s): Web based, Platform independent Programming language: HTML, CSS, JavaScript, PHP Other requirements: Modern Browser License: BSD License

Any restrictions to use by non-academics: None.



Additional files

Additional file 1: NCBI Gene Entrez ID List for the Case Scenario. This file contains NCBI gene Entrez ID list. The first column is Entrez ID, and the second column is the gene symbol. (XLSX 9 kb)

Additional file 2: NCBI Entrez IDs for Comparison of WebGIVI, DAVID and AmiGO2. This file contains NCBI gene Entrez ID list which is identifed by WebGIVI, DAVID and AmiGO2 to be associated with the concept of cell cycle regulation. (TXT 1 kb)

Abbreviations

GO: Gene ontology; iTerm: Informative term; LMH: White-leghorn hepatocellular cell; SVG: Scalable vector graphics

Acknowledgments

LS thanks Allen Hubbard for his help in editing this manuscript and Andrew Bancroft for his help in creating the tutorial video.

Funding

This work was supported by NSF grants NSF DBI-1147029 to CJS and NSF DBI-1260795 to JC.

Authors' contributions

LS, CJS, JC and YZ designed the software; LS, YZ and AM wrote the codes; COT, AM and KVS provided eGIFT data. JR helped design the Cytoscape network map. All authors helped revise the paper and implement the software. LS and CJS wrote the paper. All authors have read and approve of the final version of the manuscript.

Competing interests

The authors declare that they have no competing interests.

Consent for publication

Not applicable.

Ethics approval and consent to participate

Not applicable.

Publisher's Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details

¹Department of Animal and Food Sciences, University of Delaware, Newark, DE, USA. ²Department of Computer Science and Electrical Engineering, University of Maryland Baltimore County, 1000 Hilltop Circle, Baltimore, MD, USA. ³Department of Computer Science, Hangzhou Dianzi University, Hangzhou 310018, Zhejiang Province, People's Republic of China. ⁴Department of Computer and Information Sciences, University of Delaware, Newark, DE 19716, USA. ⁵Center for Bioinformatics and Computational Biology, University of Delaware, Newark, DE 19711, USA. ⁶Current address: Computing Service, The Samuel Roberts Noble Foundation, Ardmore, OK 73401, USA.

Received: 6 March 2017 Accepted: 28 April 2017 Published online: 04 May 2017

References

- Huang DW, Lempicki RA, Sherman BT. Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nat Protoc. 2009;4:44–57.
- Huang DW, Sherman BT, Lempicki RA. Bioinformatics enrichment tools: Paths toward the comprehensive functional analysis of large gene lists. Nucleic Acids Res. 2009;37:1–13.
- Zheng Q, Wang XJ. GOEAST: a web-based software toolkit for Gene Ontology enrichment analysis. Nucleic Acids Res. 2008;36:358–63.
- Eden E, Navon R, Steinfeld I, Lipson D, Yakhini Z. GOrilla: a tool for discovery and visualization of enriched GO terms in ranked gene lists. BMC Bioinformatics. 2009;10:48.

- Ogata H, Goto S, Sato K, Fujibuchi W, Bono H, Kanehisa M. KEGG: Kyoto encyclopedia of genes and genomes. Nucleic Acids Res. 1999;27:29–34.
- Croft D, O'Kelly G, Wu G, Haw R, Gillespie M, Matthews L, et al. Reactome: a database of reactions, pathways and biological processes. Nucleic Acids Res. 2011;39:691–7.
- 7. Tudor CO, Schmidt CJ, Vijay-Shanker K. eGIFT: mining gene information from the literature. BMC Bioinformatics. 2010;11:418.
- Pavlopoulos GA, O'Donoghue SI, Satagopam VP, Soldatos TG, Pafilis E, Schneider R. Arena3D: visualization of biological networks in 3D. BMC Syst Biol. 2008;2:104.
- 9. Hooper SD, Bork P. Medusa: a simple tool for interaction graph analysis. Bioinformatics. 2005;21:4432–3.
- Köhler J, Baumbach J, Taubert J, Specht M, Skusa A, Rüegg A, et al. Graphbased analysis and visualization of experimental results with ONDEX. Bioinformatics. 2006;22:1383–90.
- 11. Breitkreutz B, Stark C, Tyers M. Osprey: a network visualization system. Genome Biol. 2003;4:1–4.
- Batagelj V, Mrvar A. Pajek Program for Large Network Analysis. 1999;1–11 Downloaded May 2, 2017. http://vlado.fmf.uni-lj.si/pub/networks/doc/pajek.pdf.
- Freeman TC, Goldovsky L, Brosch M, van Dongen S, Mazière P, Grocock RJ, et al. Construction, visualisation, and clustering of transcription networks from microarray expression data. PLoS Comput Biol. 2007;3:2032–42.
- Shannon P, Markiel A, Ozier O, Baliga NS, Wang JT, Ramage D, et al. Cytoscape: a software environment for integrated models of biomolecular interaction networks. Genome Res. 2003;13:2498–504.
- Iragne F, Nikolski M, Mathieu B, Auber D, Sherman D. ProViz: protein interaction visualization and exploration. Bioinformatics. 2005;21:272–4.
- 16. Cytoscape.js. Available from: http://cytoscape.github.io/cytoscape.js/. Accessed 2 May 2017.
- Bostock M, Ogievetsky V, Heer J. D 3: Data-Driven Documents. IEEE Trans Vis Comput Graph. 2011;17:2301–9.
- Jiang X, Ronggui Y, Huiran Z, Wu Z, Kawata S. Biomolecular network visualization based on CPSE-Bio. Comput. Converg. Technol. (ICCCT), 2012 7th Int. Conf. 2012; 1488–92
- Salazar GA, Meintjes A, Mazandu GK, Rapanoël HA, Akinola RO, Mulder NJ. A web-based protein interaction network visualizer. BMC Bioinformatics. 2014;15:129.
- Bateman A, Martin MJ, O'Donovan C, Magrane M, Apweiler R, Alpi E, et al. UniProt: A hub for protein information. Nucleic Acids Res. 2015;43:D204–12.
- Kersey PJ, Allen JE, Armean I, Boddu S, Bolt BJ, Carvalho-Silva D, et al. Ensembl Genomes 2016: more genomes, more complexity. Nucleic Acids Res. 2015;44:D574–80.
- Sun L, Lamont SJ, Cooksey AM, McCarthy F, Tudor CO, Vijay-Shanker K, et al. Transcriptome response to heat stress in a chicken hepatocellular carcinoma cell line. Cell Stress Chaperones. 2015;20:939–50.
- Blake JA, Christie KR, Dolan ME, Drabkin HJ, Hill DP, Ni L, et al. Gene ontology consortium: going forward. Nucleic Acids Res. 2015;43:D1049–56.
- 24. Mitchell A, Chang HY, Daugherty L, Fraser M, Hunter S, Lopez R, et al. The InterPro protein families database: the classification resource after 15 years. Nucleic Acids Res. 2015;43:D213–21.

Submit your next manuscript to BioMed Central and we will help you at every step:

- We accept pre-submission inquiries
- Our selector tool helps you to find the most relevant journal
- We provide round the clock customer support
- Convenient online submission
- Thorough peer review
- Inclusion in PubMed and all major indexing services
- Maximum visibility for your research

Submit your manuscript at www.biomedcentral.com/submit

