Gene Set Enrichment

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What is Gene set enrichment?

Enrichment analysis is a technique commonly used to add biological context to a list of genes, such as list of significant genes filtered from differential analysis report. The procedure is based on assigning genes to groups and then finding overrepresented groups in filtered gene lists using a Fisher's exact test.

Running Gene set Enrichment

Gene set enrichment task can be invoked on a differential analysis output (or filtered differential analysis output) data node or filtered count matrix data node. Since the data node including all the features will serve as background, to get a meaningful result, always use a data node containing subset of features to invoke this task. Only gene names will be used in the computation.

- Click a Feature list data node
- · Click the Biological interpretation section of the toolbox
- Click Gene set enrichment
- ٠ There are two options for Database. KEGG database requires a special license (Figure 1).

Select gene set
Database
KEGG database Gene set database
KEGG database
Homo sapiens hsa_v1_22_08_24
Feature identifier
Select feature identifier
Choose the column that matches the values in the database. The match is case-insensitive. Otherwise, an annotation column will automatically be picked that matches the database.
Background
□ Specify background gene list
Enrichment analysis compares a list of observed genes. If specified, the background list of possible genes to determine whether groups of genes, such as pathways or GO terms, are overrepresented in the list of observed genes. If specified, the background list of genes is used as the list of possible genes.
Background

Figure 13. Selecting KEGG database

 Gene set database is user defined database, see more details in the Adding a Gene Set chapter. The gene sets available for the current Assembly are listed under the Gene set database drop-down list (Figure 2). The assembly is automatically selected, if possible. If the assembly cannot be detected, you can specify it using the drop-down.

Partek distributes Gene Ontology (GO) for human and mouse genomes, a bioinformatics initiative to unify the representation of gene and gene product attributes across various species [1, 2].

Select gene set
Database
KEGG database 🕘 Gene set database
Assembly
Homo sapiens (human) - hg38
Gene set database
2021 05 05 (Administrator) 🗸
Feature identifier
Select feature identifier
Choose the column that matches the values in the database. The match is case-insensitive. Otherwise, an annotation column will automatically be picked that matches the database.
Background
Specify background gene list
Enrichment analysis compares a list of observed genes to a list of possible genes to determine whether groups of genes, such as pathways or GO terms, are overrepresented in the list of observed genes. If specified, the background list of genes is used as the list of possible genes.

Figure 14. Selecting user defined gene set database

- Select feature identifier (optional) can be used to specify the feature format (e.g. Gene name, Gene ID, Feature ID).
- Specify the background gene list (optional) can be used for a feature list. Select the list using the drop-down. Click here for more information on List management.

The background gene list is used as the list of possible genes. By default, this is the genes included in the selected gene set database. If your assay limits the genes that could be detected, you may want to specify a background list.

• Click Finish to run

The result is stored under an *Enrichment task* node. To open it, **double click** on the node or select the respective **Task report** from the context sensitive menu.

Task report

Figure 3 shows an example *Gene set enrichment* task report using GO database. The table contains one gene set per row (*Gene set* column; the column entries are hyperlinks when using the distributed GO gene sets), with the category name in the *Description* column. The categories are ranked by the *Enric hment score*, which is the negative natural logarithm of the enrichment p-value (*P-value* column) derived from Fisher's exact test on the underlying contingency table. The higher the enrichment score, the more overrepresented the GO category is within the input list of significant genes. The columns

can be searched by typing in the search term in the respective box (and hitting Enter), or sorted by selecting the double arrow icon (

Data Viewer plots availa	ta Viewer plots available when table contains 100 or fewer rows.														
Gene set ↑↓	Description 1↓	Type ↑↓	Enrichment score ↑↓	P-value 17	FDR step up ↑↓	Rich factor ↑↓	Genes in set ↑↓	Genes in list ↑↓	Genes not in list ↑↓	Genes in list, not in set ↑↓	Genes not in list, not in set ↑↓	0			
GO:0043229	intracellular organelle	cellular component	16.23	8.9E-8	5.5E-4	0.9	210	186	24	71	42				
GO:0043231	intracellular membrane-bounded organelle	cellular component	14.44	5.4E-7	1.7E-3	0.9	181	162	19	95	47				
GO:0005634	nucleus	cellular component	10.58	2.5E-5	0.1	0.9	129	117	12	140	54				
GO:0005654	nucleoplasm	cellular component	10.06	4.3E-5	0.1	0.9	109	100	9	157	57				

Figure 15. Go enrichment report (truncated). Gene set column contains Gene Ontology identifiers (hyperlinks). Category labels are in the Description column. Enrichment score: negative natural logarithm of the enrichment P-value derived from the Fisher's exact test. Genes in list: number of genes that are present both in the list of significant genes and the gene set (GO category). Genes not in list: number of genes that are present in the list of significant genes. The column on the right contains links to gene breakdown chart and extra details

The contingency table (Figure 4) can be displayed by selecting the **View gene breakdown chart** icon on the right (**•**). The term "list" refers to the list of significant genes, while the term "set" refers to the respective GO category. The first row of the contingency table is also seen in the report, namely the *Gen es in list* and *Genes not in list* columns.



Figure 16. Contingency table used to calculate the enrichment p-value. List refers to the list of significant genes, set refers to the gene ontology category

The View extra details () button provides additional information on the GO category (Figure 5). In addition to the details already given in the report, a full list of *Genes in list* and *Genes not in list* can be inspected and downloaded (**Download data**) to the local computer as a text file. Use the arrow to expand these sections.

Gene set	GO:00432	29	Enrichment score	16.2343
Description	intracellul	ar organelle	P-value	8.90297E-8
iene breakd	lown			
	In list	Not in list		
In set	186	24		
Not in set	71	42		
> Genes I	n list			
✓ Genes n	ot in list			
🖪 Downlo	oad list			
ZNF479				
ZNF695				
FOXA1				
AR KRT15				
PTH1R				
TEX11				
WASF3				

Figure 17. Gene ontology enrichment extra details

As previously mentioned, if you are using the GO gene sets distributed by Partek, the GO identifiers in the first column are hyperlinks to the Gene Ontology web-site entries (an example shown in Figure 6).

				~																				
			(80 Ami	iGO 2	Home	Search	Too	ols & Res	sources	Help	Feedb	ck	About	AmiG	SO 1.8		Quick sear	ch	S	arch	0		
								alp	ha-	ami	no	acid	me	etak	olio	c pro	ces	S						
Term Informa	ation o																							
Accession	G0:19016	15 In acid metri	balic stoce																					
Ontology	biological	process	bowe probe.																					
Synonyms	alpha-amin	o acid meta	bolism																					
Definition	The chemi	all reactions	s and pathw	rays involvin	ig an alphi	a-amino a	icid. Source	GOCITE	ermGeni	e														
History	See term t	istory for G	0.1901605	at QuickGC																				
Subset	None																							
Community	Unavailable	3																						
Related	Link to all	genes and	gene prod	Jucts annot	tated to alp	pha-amini	o acid metal	bolic proc	cess.															
	Unk to all	direct and i	Adirect ann	otations to	arpha-am	Jino acid i	netabolic pr	ocess.			debe to a													
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Figure 18. Selecting a GO category in the table report opens up a browser and displays additional information on that category via GO web-page

Interactive KEGG pathway maps

When KEGG database is used, on the enrichment task report, when click on a pathway ID in the Gene set column, a KEGG pathway gene network picture is displayed (Figure 7).



Figure 19. KEGG pathway map

Each rectangle on the map represents a gene product in the pathway. Gene products are mostly proteins coded by a gene or group of genes, but they could be RNA too. Related pathways are shown as large rounded rectangles. Chemical compounds, DNA or other molecules are shown as circles.

Coloring the map

The pathway map is colored by the first fold-change column in the input *Feature list* data node. The control panel on the left can be used to configure the colors of the pathway map. In all options, rectangles colored white do not have gene information. Options for coloring include:

- Fixed color: all genes are colored black.
- Genes in list: all genes in the list are colored, default color is yellow, but this can be configured. Genes not in the list are black.
- Statistics in the gene list: .e.g FDR, p-value, Fold change etc. Colors can be customized by clicking on the color square to change.

Feature details

Mousing over a rectangle shows the genes indicated by the rectangle in the tooltip (Figure 8). Genes are listed on rows with all aliases in the KEGG database included on the row. Genes that are in the list are shown in bold. The gene being used to color the rectangle is shown in red.



Figure 20. Checking genes represented by a rectangle

On KEGG pathway maps that include chemical compounds, the chemical structure is shown in the tooltip on mouse-over (Figure 9).



Figure 21. Viewing chemical compound

Clicking a rectangle opens the page for that gene or group of genes on the KEGG website in a new tab in your web browser.

Click the Save image 💾 icon to download a PNG file showing the configured KEGG pathway map to your local computer.

Visualizing gene set enrichment results

If the gene set enrichment table has fewer than 100 results (rows), the GO categories can be visualized in the *Data Viewer*. Otherwise, a notification is displayed in the top left corner (Figure 10).

Data Viewer plots availab	le when table contains 100 or fewer rows.										
Gene set 1↓	Description 1	Enrichment score 1	P-value 17	FDR step up ↑↓	Rich factor ↑↓	Genes in set ↑↓	Genes in list ↑↓	Genes not in list ↑↓	Genes in list, not in set ↑↓	Genes not in list, not in set ↑↓	0
path:hsa04640	Hematopoietic cell lineage	15.01	3E-7	9.6E-5	0.3	31	9	22	31	1,103	
path:hsa05330	Allograft rejection	6.19	2.1E-3	0.2	0.2	17	4	13	36	1,112	
path:hsa05320	Autoimmune thyroid disease	6.19	2.1E-3	0.2	0.2	17	4	13	36	1,112	
path:hsa04940	Type I diabetes mellitus	5.96	2.6E-3	0.2	0.2	18	4	14	36	1.111	
path:hsa05412	Arrhythmogenic right ventricular cardiomyopathy	5.90	2.7E-3	0.2	0.3	9	3	6	37	1,119	

Figure 22. If the gene ontology table has more than 100 rows, visualization of results is not possible

If needed, filter down the number results, for instance by using a cut-off based on the enrichment score. Type in the cut-off value in the text box beneath the *Enrichment score* and hit **enter** (an example is shown in Figure 11). Once the number or results falls below 100, a link to the *Data Viewer* will be displayed (Figure 8). Click on the **View plots in Data Viewer** link to open a new *Data Viewer* session.

View plots in Data View	View plots in Data Viewer													
Gene set ↑↓	Description ↑↓	Enrichment score 1↓ P-value 1₽ >6		FDR step up ↑↓	Rich factor ↑↓	Genes in set ↑↓	Genes in list ↑↓	Genes not in list ↑↓	Genes in list, not in set ↑↓	Genes not in list, not in set ↑↓	0			
path:hsa04640	Hematopoietic cell lineage	15.01	3E-7	9.6E-5	0.3	31	9	22	31	1,103	==			
path:hsa05330	Allograft rejection	6.19	2.1E-3	0.2	0.2	17	4	13	36	1,112				
path:hsa05320	Autoimmune thyroid disease	6.19	2.1E-3	0.2	0.2	17	4	13	36	1,112				
	Rows per page 25v << (1 of 1) > >>													
🖪 Download table	🗄 Download gene sets													
Task details														

Figure 23. Use the View plots in Data Viewer link to visualize the gene ontology enrichment results. The link is not visible if the table contains more than 100 rows

Two plots are loaded into Data Viewer (Figure 12). Both plots show enrichment score on the horizontal axis and gene ontology categories (i.e. the ones present in the gene enrichment table) on the vertical axis. The plots show enrichments scores (*Enrichment score* column of the gene ontology table) and - in addition - the plot on the left uses color range to depict enrichment *P-value* (green = low, red = high P-value).

The same functionality is available for pathway enrichment results.



Figure 24. Visualizing gene ontology results. Vertical axis shows the gene ontology categories present in the underlying gene ontology table

References

- 1. Ashburner M, Ball CA, Blake JA et al. Gene Ontology: tool for the unification of biology. Nat Genetics. 2000; 25:25-29.
- 2. The Gene Ontology Consortium. Gene Ontology Consortium: going forward. *Nucleic Acids Res.* 2015; 43:D1049-1056.Recommended citations from the Geneontology.org website

Additional Assistance

If you need additional assistance, please visit our support page to submit a help ticket or find phone numbers for regional support.

