

Gene Ontology解析

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What is Gene Ontology (GO)?

- ▶ GO project describes gene products from all organisms using a consistent and computable language.
- ▶ GO produces sets of explicitly defined, structured vocabularies in both a computer- and human-readable manner.
- ▶ 3 categories
 - ▶ Biological processes
 - ▶ Molecular functions
 - ▶ Cellular components
- ▶ 2 components
 - ▶ Ontology: term definition and the structured relationships between them
 - ▶ Associations between gene products and the GO terms.

<http://www.geneontology.org/>

Two components of GO

- ▶ Ontology
- ▶ Gene associations

Gene Ontology Consortium

Search GO data

Search

Ontology

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Gene Ontology: the framework for the model of biology. The GO defines concepts/classes used to describe gene function, and relationships between these concepts. It classifies functions along three aspects:

molecular function
molecular activities of gene products

cellular component
where gene products are active

biological process
pathways and larger processes made up of the activities of multiple gene products.

[more](#)

Annotations

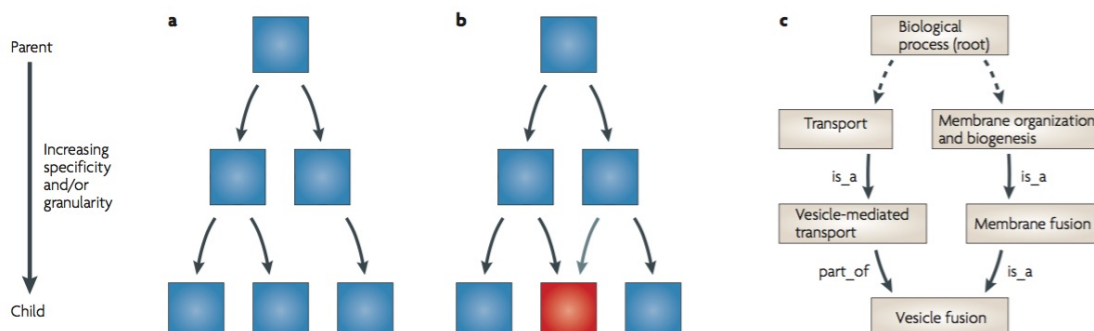
[Download annotations](#) (standard files)

[Filter and download](#) (customizable files <100k lines)

GO annotations: the model of biology. Annotations are statements describing the functions of specific genes, using concepts in the Gene Ontology. The simplest and most common annotation links one gene to one function, e.g. FZD4 + Wnt signaling pathway. Each statement is based on a specified piece of evidence. [more](#)

Ontology structure

- ▶ Ontologies are represented as a directed acyclic graph (DAG).
- ▶ Parent-child relationship
 - ▶ is_a
 - ▶ part_of
- ▶ Ontology can be changed / updated




Rhee et al., 2008

Gene association

- ▶ Gene <=> GO
- ▶ A gene may associate with multiple GO terms.
- ▶ Evidence codes.


Evidence code	Evidence code description	Source of evidence	Manually checked
IDA	Inferred from direct assay	Experimental	Yes
IEP	Inferred from expression pattern	Experimental	Yes
IGI	Inferred from genetic interaction	Experimental	Yes
IMP	Inferred from mutant phenotype	Experimental	Yes
IPI	Inferred from physical interaction	Experimental	Yes
ISS	Inferred from sequence or structural similarity	Computational	Yes
RCA	Inferred from reviewed computational analysis	Computational	Yes
IGC	Inferred from genomic context	Computational	Yes
IEA	Inferred from electronic annotation	Computational	No
IC	Inferred by curator	Indirectly derived from experimental or computational evidence made by a curator	Yes
TAS	Traceable author statement	Indirectly derived from experimental or computational evidence made by the author of the published article	Yes
NAS	Non-traceable author statement	No 'source of evidence' statement given	Yes
ND	No biological data available	No information available	Yes
NR	Not recorded	Unknown	Yes

nanos

Gene Product Information 

http://amigo.geneontology.org/amigo/gene_product/FB:FBgn0002962

Symbol nos
Name(s) nanos

Total annotations: 29; showing: 1-10
Results count 

[First](#) [Prev](#) [Next](#) [Last](#) [Download \(up to 100000\)](#)

<input type="checkbox"/> Gene/product	Gene/product name	Annotation qualifier	GO class (direct)	Annotation extension	Contributor	Organism	Evidence	Evidence with	PANTHER family	Isoform	Reference
<input type="checkbox"/> nos	nanos		germ cell migration		FlyBase	Drosophila melanogaster	TAS		nanos protein pthr12887		FB:FBn0107500 PMID:9988212
<input type="checkbox"/> nos	nanos		oogenesis		FlyBase	Drosophila melanogaster	IMP		nanos protein pthr12887		FB:FBn0107609 PMID:10101171
<input type="checkbox"/> nos	nanos		spermatogenesis		FlyBase	Drosophila melanogaster	IMP		nanos protein pthr12887		FB:FBn0107609 PMID:10101171
<input type="checkbox"/> nos	nanos		pole plasm		FlyBase	Drosophila melanogaster	TAS		nanos protein pthr12887		FB:FBn0110978 PMID:10449356
<input type="checkbox"/> nos	nanos		anterior/posterior axis specification, embryo		FlyBase	Drosophila melanogaster	TAS		nanos protein pthr12887		FB:FBn0111327 PMID:10494038
<input type="checkbox"/> nos	nanos		oocyte anterior/posterior axis specification		FlyBase	Drosophila melanogaster	NAS		nanos protein pthr12887		FB:FBn0128774 PMID:10878576
<input type="checkbox"/> nos	nanos		protein binding		FlyBase	Drosophila melanogaster	IPI	FB:FBgn0000392	nanos protein pthr12887		FB:FBn0131417 PMID:11060247
<input type="checkbox"/> nos	nanos		germ-line stem cell division		FlyBase	Drosophila melanogaster	NAS		nanos protein pthr12887		FB:FBn0132358 PMID:11131516
<input type="checkbox"/> nos	nanos		protein binding		UniProt	Drosophila melanogaster	IPI	FB:FBgn0010300	nanos protein pthr12887		FB:FBn0135777 PMID:11274060
<input type="checkbox"/> nos	nanos		female meiosis chromosome segregation		FlyBase	Drosophila melanogaster	IMP		nanos protein pthr12887		FB:FBn0135802 PMID:11290718

How to annotate GO for non-model organisms?

- ▶ Ortholog grouping with a model organism and then transfer the GO terms from the reference organism to your target organism.
- ▶ BLAST2GO
- ▶ InterProScan

Gene Ontology enrichment analysis

- ▶ What is GO enrichment analysis?
- ▶ Why GO enrichment analysis is required in DEG studies?
- ▶ Type of GO enrichment analysis.
 - ▶ gene set
 - ▶ gene score
- ▶ Software
 - ▶ gene set type: DAVID (web), metasplice (web), goseq (R), GOstat (R)
 - ▶ gene score: GSEA, roast, camera
 - ▶ both: ErmineJ

Basic over-representation test: 2 x 2 table and Fisher's exact test

- ▶ Suppose we perform a test of DE and find a list of 200 significant genes out of 10,000
- ▶ Consider a specific GO term, apoptosis. Among the 200 DE genes, 20 genes are annotated as apoptosis related, while 300 / 10,000 are associated with apoptosis in the whole gene set.
- ▶ Question: Is the gene set “apoptosis” over-represented among “significant” genes?

	apoptosis	non-apoptosis	total
DE	20	180	200
non-DE	280	9,520	9,800
total	300	9,700	10,000

```
> mat <- matrix(c(20,200-20,300-20, 10000-300-(200-20)),
nrow=2, byrow=T)
> fisher.test(mat, alternative="greater")

Fisher's Exact Test for Count Data

data:  mat
p-value = 2.269e-06
alternative hypothesis: true odds ratio is greater than 1
95 percent confidence interval:
 2.418508      Inf
sample estimates:
odds ratio
 3.777069
```

Try 演習問題 ex10

Gene score type enrichment analysis

- ▶ Drawback of basic 2x2 table method

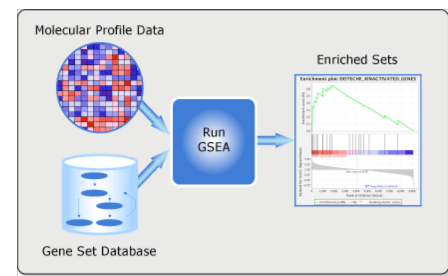
- ▶ Threshold value is arbitral
- ▶ Magnitude of significance is ignored

- ▶ GSEA

- ▶ <http://software.broadinstitute.org/gsea/index.jsp>

- ▶ ROAST, CAMERA

- ▶ implemented within edgeR



Tutorial: ErmineJ

演習問題 exII

- ▶ <http://erminej.chibi.ubc.ca/>



- ▶ Easy to use Java software with both GUI and CUI
- ▶ Three enrich methods supported
 - ▶ ORA: overrepresentation analysis
 - ▶ GSR: gene score resampling
 - ▶ ROC: rank-based gene score in receiver-operator curves