BIMM 143 Pathway Analysis and the Interpretation of Gene Lists

Lecture 15

Barry Grant UC San Diego

http://thegrantlab.org/bimm143



| x ‡ | baseMean 🗘 | log2FoldChange 🗘 | lfcSE 🗘 | stat 🗘 | pvalue 🗘 | padj 🗘 | symbol 🗘 |
|-----------------|-------------|------------------|------------|------------|--------------|--------------|----------|
| ENSG00000152583 | 954.77093 | 4.3683590 | 0.23713648 | 18.421286 | 8.867079e-76 | 1.342919e-71 | SPARCL1 |
| ENSG00000179094 | 743.25269 | 2.8638885 | 0.17555825 | 16.313039 | 7.972621e-60 | 6.037267e-56 | PER1 |
| ENSG00000116584 | 2277.91345 | -1.0347000 | 0.06505273 | -15.905557 | 5.798513e-57 | 2.927283e-53 | ARHGEF2 |
| ENSG00000189221 | 2383.75371 | 3.3415441 | 0.21241508 | 15.731200 | 9.244206e-56 | 3.500088e-52 | MAOA |
| ENSG00000120129 | 3440.70375 | 2.9652108 | 0.20370277 | 14.556557 | 5.306416e-48 | 1.607313e-44 | DUSP1 |
| ENSG00000148175 | 13493.92037 | 1.4271683 | 0.10036663 | 14.219550 | 6.929711e-46 | 1.749175e-42 | STOM |
| ENSG00000178695 | 2685.40974 | -2.4890689 | 0.17806407 | -13.978501 | 2.108817e-44 | 4.562576e-41 | KCTD12 |
| ENSG00000109906 | 439.54152 | 5.9275950 | 0.42819442 | 13.843233 | 1.397758e-43 | 2.646131e-40 | ZBTB16 |
| ENSG00000134686 | 2933.64246 | 1.4394898 | 0.10582729 | 13.602255 | 3.882769e-42 | 6.533838e-39 | PHC2 |
| ENSG00000101347 | 14134.99177 | 3.8504143 | 0.28490701 | 13.514635 | 1.281894e-41 | 1.941428e-38 | SAMHD1 |
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| ENSG00000166741 | 7542.25287 | 2.2195906 | 0.16673544 | 13.312050 | 1.970000e-40 | 2.486304e-37 | NNMT |
| ENSG00000125148 | 3695.87946 | 2.1985636 | 0.16700546 | 13.164621 | 1.402400e-39 | 1.633797e-36 | MT2A |
| ENSG00000162614 | 5646.18314 | 1.9711402 | 0.15020631 | 13.122885 | 2.434854e-39 | 2.633990e-36 | NEXN |
| ENSG00000106976 | 989.04683 | -1.8501713 | 0.14778657 | -12.519211 | 5.861471e-36 | 5.918132e-33 | DNM1 |
| ENSG00000187193 | 199.07694 | 3.2551424 | 0.26090711 | 12.476250 | 1.006146e-35 | 9.523804e-33 | MT1X |
| ENSG00000256235 | 1123.47954 | 1.2801193 | 0.10547438 | 12.136779 | 6.742862e-34 | 6.007096e-31 | SMIM3 |
| ENSG00000177666 | 2639.57020 | 1.1399947 | 0.09606884 | 11.866436 | 1.768422e-32 | 1.487930e-29 | PNPLA2 |
| ENSG00000164125 | 7257.00808 | 1.0248523 | 0.08657600 | 11.837603 | 2.494830e-32 | 1.988642e-29 | FAM198B |
| ENSG00000198624 | 2020.04495 | 2.8141014 | 0.24063429 | 11.694515 | 1.359615e-31 | 1.029569e-28 | CCDC69 |
| ENSG00000123562 | 5008.55294 | 1.0045453 | 0.08901501 | 11.285123 | 1.554241e-29 | 1.120904e-26 | MORF4L2 |
| ENSG00000144369 | 1283.77980 | -1.3090041 | 0.11714863 | -11.173875 | 5.473974e-29 | 3.768333e-26 | FAM171B |
| ENSG00000196517 | 241.91536 | -2.3456877 | 0.21047366 | -11.144804 | 7.591120e-29 | 4.998588e-26 | SLC6A9 |
| ENSG00000135821 | 19973.40000 | 3.0413943 | 0.27601796 | 11.018828 | 3.100706e-28 | 1.956675e-25 | GLUL |



Log2(FoldChange)

My high-throughput experiment generated a long list of genes/proteins...



What do I do now?

Pathway analysis! (a.k.a. geneset enrichment)

Use bioinformatics methods to help extract biological meaning from such lists...



Basic idea

Differentially Expressed Genes (**DEGs**)

| x ÷ | baseMean 🗘 | log2FoldChange 🗘 | lfcSE [‡] | stat 🗘 | ¢ pvalue | padj [‡] | symbol |
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Gene-sets (Pathways, annotations, etc...)



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Gene-sets (Pathways, annotations, etc...)





Pathway analysis (a.k.a. geneset enrichment) Principle



- DEGs come from your experiment
- Pathway genes ("geneset") come from annotations > Important, but typically not a competitive advantage
- > Critical, needs to be as clean as possible
- Variations of the math: overlap, ranking, networks... > Not critical, different algorithms show similar performances

Pathway analysis (a.k.a. geneset enrichment) Limitations

Geneset annotation bias: can only discover what is already known

Side note.

- Non-model organisms: no high-quality genesets available
- Post-transcriptional regulation is neglected
- Tissue-specific variations of pathways are not annotated
 - e.g. NF-κB regulates metabolism, not inflammation, in adipocytes
- Size bias: stats are influenced by the size of the pathway
 - Many pathways/receptors converge to few regulators
 e.g. Tens of innate immune receptors activate four TFs: NF-kB, AP-1, IRF3/7, NFAT

Starting point for pathway analysis: Your gene list

- You have a list of genes/proteins of interest
- You have quantitative data for each gene/protein
 - Fold change
 - p-value
 - Spectral counts
 - Presence/absence



Translating between identifiers

- Many different identifiers exist for genes and proteins, e.g. UniProt, Entrez, etc.
- Often you will have to translate one set of ids into another
 - A program might only accept certain types of ids
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Translating between identifiers: UniProt < <u>www.uniprot.org</u> >

| UniProt | | Downloads · Contact · Documentation/Help |
|---------------------------------------------------|------------------------------------------------------------------------------------|------------------------------------------|
| Search in Query Protein Knowledgebase (UniProtKB) | Search Clear Fields » Search Blast Align | Retrieve ID Mapping |
| WELCOME | NEWS | |
| Identifiers | From EMBL/GenBank/DDBJ To UniProtKB AC or Choose File no file selected | Map Swap Clear |

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VLOOKUP in Excel - good if you are an excel whizz - I am not!

 Download flat file from Entrez, Uniprot, etc; Open in Excel; Find columns that correspond to the 2 IDs you want to convert between; Sort by ID; Use vlookup to translate your list

Translating between identifiers: Excel VLOOKUP

VLOOKUP(lookup_value, table_array, col_index_num)

| | Edit | | | Font | | | Aligr | ment | | Nu | umber |
|----|-----------------------------------------------------------|------------|------------|------------|------------|---|--------------|-----------|-------------|-----------|--------|
| f | 🖣 🚽 💽 Fi | II 🔻 Cali | bri (Body) | v 12 | • A• A• | | ≡ abo | c 🔻 📆 Wra | ap Text 🔻 🛛 | General | |
| Pa | Paste V Clear • B I U · · · · · · · · · · · · · · · · · · | | | | | | | | | | |
| | B3 | | | | | | | | | | |
| | A | B | C | D | E | F | G | Н | | J | K |
| 1 | Data Table | | | | | | Annotation T | able | | | |
| 2 | RefSeq | Symbol | Exp1 | Exp2 | Exp3 | | RefSeq | Symbol | Entrez ID | Unigene | RefSeq |
| 3 | NM_153103 | Kif1c , | 2.31975457 | 1.24558927 | 2.78816871 | | NM_001001 | Zfp85-rs1 | 22746 | Mm.288396 | NM_001 |
| 4 | NM_146017 | Gabrp | 4.15029735 | 3.08055836 | 1.18919962 | | NM_001001 | Scap | 235623 | Mm.288741 | NM_001 |
| 5 | NM_018883 | Camkk1 | 3.83282512 | 0.0522951 | 0.64684259 | | NM_001001 | Scap | 235623 | Mm.288741 | NM_001 |
| 6 | NM_145936 | Tspyl2 | 0.45449369 | 1.62761318 | 7.59770627 | | NM_001001 | Fbxo41 | 330369 | Mm.38777 | NM_001 |
| 7 | NM_026599 | Cgnl1 | 4.84541871 | 2.84751796 | 1.61595768 | | NM_001001 | Taf9b | 407786 | Mm.19440 | NM_001 |
| 8 | NM_013926 | Cbx8 | 1.22903318 | 0.2863077 | 0.02952665 | | NM_001001 | Taf9b | 407786 | Mm.19440 | NM_001 |
| 9 | NR_015566 | A330023F24 | 1.44695053 | 0.98809479 | 1.59330144 | | NM_001001 | BC051142 | 407788 | Mm.73205 | NM_001 |
| 10 | NM_008623 | Mpz | 0.50749263 | 0.94350028 | 6.10581569 | | NM_001001 | BC051142 | 407788 | Mm.73205 | NM_001 |
| 11 | NM_183127 | Fate1 | 2.45672795 | 4.87960794 | 3.60759511 | | NM_001001 | BC048546 | 232400 | Mm.259234 | NM_001 |
| 12 | NM_008943 | | 4.78701069 | 4.15302647 | 0.85432314 | | NM_001001 | Zfp941 | 407812 | Mm.359154 | NM_001 |
| 13 | NM_025382 | | 0.66397344 | 1.40664187 | 3.09539802 | | NM_001001 | BC031181 | 407819 | Mm.29866 | NM_001 |
| 14 | NM_182841 | | 1.25528938 | 0.20505996 | 2.76879488 | | NM_001001 | Baz2b | 407823 | Mm.486364 | NM_001 |
| 15 | NM_030061 | | 0.17670108 | 2.75415469 | 2.98900691 | | NM_001001 | Tmem204 | 407831 | Mm.34379 | NM_001 |
| 16 | NM_133216 | | 6.572343 | 0.59671282 | 3.84650536 | | NM_001001 | Ccdc111 | 408022 | Mm.217385 | NM_001 |
| 17 | NM 030063 | | 7.05132762 | 0.65043627 | 1.68111836 | | NM 001001 | BC048507 | 408058 | Mm.177840 | NM 001 |

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 - Download flat file from Entrez, Uniprot, etc; Open in Excel; Find columns that correspond to the two ids you want to convert between; Use vlookup to translate your list
- Use the merge() or mapIDs() functions in R <u>fast</u>, *versatile* & reproducible!
 - Also clusterProfiler::bitr() function and many others... [Link to clusterProfiler vignette]



> merge(mygenes, anno, by.x="row.names", by.y= "ensgene")

This is our differential expressed genes

Pennincier.

- # Using the merge() function
- > anno <- read.csv("data/annotables_grch38.csv")</pre>
- > merge(mygenes, anno, by.x="row.names", by.y= "ensgene")



bitr: Biological Id TranslatoR

clusterProfiler provides bitr and bitr_kegg for converting ID types. Both bitr and bitr_kegg support many species including model and many non-model organisms.

| x <− c(| "GPX3", | "GLRX", | "LBP", | "CRYAB", | "DEFB1", | "HCLS1", | "SOD2", | "HSPA2", |
|----------------|------------|-------------|-----------|-----------|-----------|-------------|-----------|----------|
| | "ORM1", | "IGFBP1", | "PTHLH", | "GPC3", | "IGFBP3" | ,"TOB1", | "MITF", | "NDRG1", |
| | "NR1H4", | "FGFR3", | "PVR", | "IL6", | "PTPRM", | "ERBB2", | "NID2", | "LAMB1", |
| | "COMP", | "PLS3", | "MCAM", | "SPP1", | "LAMC1", | "COL4A2", | "COL4A1", | "MYOC", |
| | "ANXA4", | "TFPI2", | "CST6", | "SLPI", | "TIMP2", | "СРМ", | "GGT1", | "NNMT", |
| | "MAL", | "EEF1A2", | "HGD", | "TCN2", | "CDA", | "PCCA", | "CRYM", | "PDXK", |
| | "STC1", | "WARS", | "HMOX1", | "FXYD2", | "RBP4", | "SLC6A12", | "KDELR3", | "ITM2B") |
| eg = bi | .tr(x, fro | omType="SYI | MBOL", to | Type="ENT | REZID", O | rgDb="org.H | s.eg.db") | |
| head(eg |) | | | | | | | |

| ## | | SYMBOL | ENTREZID |
|----|---|--------|----------|
| ## | 1 | GPX3 | 2878 |
| ## | 2 | GLRX | 2745 |
| ## | 3 | LBP | 3929 |
| ## | 4 | CRYAB | 1410 |
| ## | 5 | DEFB1 | 1672 |
| ## | 6 | HCLS1 | 3059 |

See package vignette:

https://bioconductor.org/packages/release/bioc/html/clusterProfiler.html

What functional set databases do you want?

- Most commonly used:
 - Gene Ontology (GO)
 - KEGG Pathways (mostly metabolic)
 - GeneGO MetaBase
 - Ingenuity Pathway Analysis (IPA) INGENUITY
- Many others...
 - Enzyme Classification, PFAM, Reactome,
 - Disease Ontology, MSigDB, Chemical Entities of Biological Interest, Network of Cancer Genes etc...
 - See: Open Biomedical Ontologies (<u>www.obofoundry.org</u>)



GO < <u>www.geneontology.org</u> >

- What function does HSF1 perform?
 - response to heat; sequence-specific DNA binding; transcription; etc

 Ontology => a structured and controlled vocabulary that allows us to annotate gene products consistently, interpret the relationships among annotations, and can easily be *handled by a computer*

 GO database consists of 3 ontologies that describe gene products in terms of their associated biological processes, cellular components and molecular functions

GO Annotations

- GO is <u>not</u> a stand-alone database of genes/proteins or sequences
- Rather gene products get annotated with **GO terms** by UniProt and other organism specific databases, such as Flybase, Wormbase, MGI, ZFIN, etc.
- Annotations are available through AmiGO < <u>amigo.geneontology.org</u> >

| DA analadas DA apakaging DA apakadas gan da apakaging DA apakadas gan da apakaging DA apakadas gan da apakadas | Pringer the second | e Gen | e Ont | tology | Α | miGO | |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------|--------|----------|---------------------|------------------------------------------------------------------------|----------------------------------------------------------------------------|--|
| | Search | Browse | BLAST | Homolog Annotations | Tools & Resources | Help | |
| | Search the Gene Ontology database | | | | | | |
| | | 0 | GO terms | • genes or proteins | exact match | Beta Beta | |
| AmiGO versio Try AmiGO La | on: <u>1.8</u> abs | | | | GO datab <u>Cite this data</u> • <u>Terms</u> Copyright © 1999-2 | ase release 2013-10-05 s of use • GO helpdesk 2010 the Gene Ontology | |

GO is structured as a "directed graph"



GO evidence codes

| Evidence code | Evidence code description | Source of evidence | Manually checked | Current number of annotations* |
|---------------|-------------------------------------------------|------------------------------------------------------------------------------------------------------------|---------------------|--------------------------------|
| IDA | Inferred from direct assay | Experimental | Yes | 71,050 |
| IEP | Inferred from expression pattern | Experimental | Yes | 4,598 |
| IGI | Inferred from genetic interaction | Experimental | Yes | 8,311 |
| IMP | Inferred from mutant phenotype | Experimental | Yes | 61,549 |
| IPI | Inferred from physical interaction | Experimental | Yes | 17,043 |
| ISS | Inferred from sequence or structural similarity | Computational | Yes | 196,643 |
| RCA | Inferred from reviewed computational analysis | Computational | Yes | 103,792 |
| IGC | Inferred from genomic context | Computational | Yes | 4 |
| IEA | Inferred from electronic annotation | Computational | No | 15,687,382 |
| IC | Inferred by curator | Indirectly derived from experimental or computational evidence made by a curator | Yes | 5,167 |
| TAS | Traceable author statement | Indirectly derived from experimental or computational evidence made by the author of the published article | Yes | 44,564 |
| NAS | Non-traceable author statement | No 'source of evidence' statement given | Yes | 25,656 |
| ND | No biological data available | No information available | Yes | 132,192 |
| NR | Not recorded | Unknown | Yes | 1,185 |

*October 2007 release

Use and misuse of the gene ontology annotations Seung Yon Rhee, Valerie Wood, Kara Dolinski & Sorin Draghici *Nature Reviews Genetics* **9**, 509-515 (2008) Experimental annotations by species



• See AmiGO for details: <u>http://amigo.geneontology.org/amigo/base_statistics</u>

Can now do gene list analysis with GeneGO online!

| | | pantherdb.org/webservices/go/overrep.jsp | • • • + |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------|
| GENEONTOLOGY Unifying Biology | Class | Ification System | . CONTACT US |
| Home About PANTHER Data New! PANTHER13.1 release | PANTHER Tools Workspa | ce Downloads Help/Tutorial | |
| Search | Gene List Analysis | Browse Sequence Search cSNP Scoring Keywo | ord Search |
| Go | Please refer to our article Error parsing request, no | in <u>Nature Protocols</u> for detailed instructions on how to use this page. <i>input specified</i> | |
| Quick linksWhole genome function viewsGenome statisticsData VersionHow to cite PANTHERNEW! Recent publication describing PANTHERNewsPANTHER13.1 ReleasedClick for additional info.Newsletter subscription | Help Tips Steps: 1. Select list and list type to analyze 2. Select Organism 3. Select operation | 1. Enter ids and or select file for batch upload. Else enter ids or select file or list from workspace for comparing to a reference list. Enter IDs: separate IDs by a space or comma Upload IDs: Choose File no file selected File format Please login to be able to select lists from your workspace. Select List Type: ID List Previously exported text search results Workspace list PANTHER Generic Mapping File VCF File Flanking region 20 Kb | |
| Enter your Email: Subscribe | | 2. Select organism. Homo sapiens Mus musculus Rattus norvegicus Gallus gallus Danio rerio 3. Select Analysis. | |
| | | 3. Select Analysis. Functional classification viewed in gene list | |

Another popular online tool: DAVID at NIAID < <u>david.abcc.ncifcrf.gov</u> >



DAVID

• Functional Annotation Chart

| Functional Annotation Chart | | | | | | | | Help and Manual |
|-------------------------------|---------------------------------------------------------------|----------------------------------------------------------------------------------------------------|-----------|-----------------|-------|-------------|---------|-----------------------|
| Current Current 2316 D/ | Gene List: Uploaded List Background: Homo sapi AVID IDs | ens | | | | | | <u>Hep and Manual</u> |
| Option | 15 | | | | | | | |
| Rerun Using | Options Create Sublist | | | | | | | 🔓 Download File |
| Sublist | Category | ≑ Term | RT | Genes | Count | \$ <u>%</u> | P-Value | ♦ <u>Benjamini</u> ♦ |
| | GOTERM_BP_5 | regulation of progression through cell cycle | RT | - | 98 | 4.2 | 3.3E-7 | 8.6E-4 |
| | GOTERM_BP_5 | apoptosis | RT | = | 131 | 5.7 | 1.6E-6 | 2.1E-3 |
| | GOTERM_BP_5 | <u>cell death</u> | <u>RT</u> | | 136 | 5.9 | 3.8E-6 | 3.3E-3 |
| | GOTERM_BP_5 | regulation of transcription from RNA polymerase II promoter | RT | = | 83 | 3.6 | 3.7E-5 | 2.4E-2 |
| | GOTERM_BP_5 | protein kinase cascade | RT | | 71 | 3.1 | 4.7E-5 | 2.4E-2 |
| | GOTERM_BP_5 | regulation of kinase activity | RT | E | 48 | 2.1 | 5.4E-5 | 2.3E-2 |
| | GOTERM_BP_5 | negative regulation of cell proliferation | RT | | 48 | 2.1 | 1.0E-4 | 3.7E-2 |
| | GOTERM_BP_5 | regulation of cell size | RT | ÷ | 41 | 1.8 | 1.2E-4 | 3.9E-2 |
| | GOTERM_BP_5 | monocarboxylic acid metabolic process | RT | 1. Alton (1997) | 48 | 2.1 | 1.3E-4 | 3.6E-2 |
| | GOTERM_BP_5 | positive regulation of nucleobase, nucleoside, nucleotide and nucleic acid metabolic process | RT | = | 61 | 2.6 | 1.5E-4 | 3.8E-2 |
| | GOTERM_BP_5 | positive regulation of cellular metabolic process | RT | = | 72 | 3.1 | 1.7E-4 | 3.8E-2 |

Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources Da Wei Huang, Brad T Sherman & Richard A Lempicki *Nature Protocols* **4**, *44 - 57 (2009)*

Overlapping functional sets

Many functional sets overlap

• In particular those from databases that are hierarchical in nature (e.g. GO)

Hierarchy enables:

- Annotation flexibility (e.g. allow different degrees of annotation completeness based on what is known)
- Computational methods to "understand" function relationships (e.g. ATPase function is a subset of enzyme function)

Unfortunately, this also makes functional profiling trickier

• Clustering of functional sets can be helpful in these cases

DAVID

• DAVID now offers functional annotation clustering:

| | Help and Tool Manual |
|------------------|----------------------------------|
| 2320 DAVID IDs | |
| Check Defaults 🗹 | Clear All |
| | |
| | |
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| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | |
| | 2320 DAVID IDs Check Defaults |

DAVID Functional Annotation Clustering

Based on shared genes between functional sets

| Functional Annotation Clustering | | | | | | | | | | | |
|----------------------------------|--------------------------------------------------|----------------------------------------------------------------------------------------------------|----|----------|----------|-------|---------|-------------|--|--|--|
| Curre | Current Gene List: Uploaded List 3 | | | | | | | | | | |
| 2320 | 2320 DAVID IDs | | | | | | | | | | |
| 🗄 Opt | Options Classification Stringency Medium | | | | | | | | | | |
| Rerur | Rerun using options Create Sublist | | | | | | | | | | |
| | Annotation Cluster 1 | Enrichment Score: 3.72 | G | | 1 | Count | P_Value | e Benjamini | | | |
| | GOTERM_BP_5 | regulation of transcription from RNA polymerase II promoter | RT | = | | 83 | 3.7E-5 | 2.4E-2 | | | |
| | GOTERM_BP_5 | positive regulation of nucleobase, nucleoside, nucleotide and nucleic acid metabolic process | RT | • | | 61 | 1.5E-4 | 3.8E-2 | | | |
| | GOTERM_BP_5 | positive regulation of cellular metabolic process | RT | Ξ. | | 72 | 1.7E-4 | 3.8E-2 | | | |
| | GOTERM_BP_5 | positive regulation of transcription | RT | - | | 58 | 3.8E-4 | 5.0E-2 | | | |
| | GOTERM_BP_5 | positive regulation of transcription, DNA- dependent | RT | Ξ. | | 48 | 7.4E-4 | 7.6E-2 | | | |
| | Annotation Cluster 2 | Enrichment Score: 3.54 | G | | | Count | P_Value | e Benjamini | | | |
| | GOTERM_BP_5 | regulation of cell size | RT | E | | 41 | 1.2E-4 | 3.9E-2 | | | |
| | GOTERM_BP_5 | regulation of cell growth | RT | ÷ | | 33 | 3.7E-4 | 5.1E-2 | | | |
| | GOTERM_BP_5 | cell morphogenesis | RT | = | | 81 | 5.2E-4 | 5.7E-2 | | | |
| | Annotation Cluster 3 | Enrichment Score: 3.37 | G | | | Count | P_Value | e Benjamini | | | |
| | GOTERM_BP_5 | apoptosis | RT | - | | 131 | 1.6E-6 | 2.1E-3 | | | |
| | GOTERM_BP_5 | <u>cell death</u> | RT | = | | 136 | 3.8E-6 | 3.3E-3 | | | |
| | GOTERM_BP_5 | regulation of programmed cell death | RT | = | | 88 | 3.2E-4 | 5.8E-2 | | | |
| | GOTERM_BP_5 | positive regulation of apoptosis | RT | 1 | | 48 | 3.3E-4 | 5.6E-2 | | | |
| | GOTERM_BP_5 | regulation of apoptosis | RT | = | | 87 | 3.5E-4 | 5.2E-2 | | | |
| | GOTERM_BP_5 | positive regulation of programmed cell death | RT | = | | 48 | 4.0E-4 | 5.0E-2 | | | |

Want more?



- GeneGO < portal.genego.com >
 - MD/PhD curated annotations, great for certain domains (eg, Cystic Fibrosis)
 - Nice network analysis tools
 - Email us for access
- Oncomine < www.oncomine.org >
 - Extensive cancer related expression datasets
 - Nice concept analysis tools
 - Research edition is free for academics, Premium edition \$\$\$
- Lots and lots other R/Bioconductor packages in this area!!!

Hands-on time!

Do it Lourser

https://bioboot.github.io/bimm143_S19/lectures/#15



counts + metadata

<u>countData</u>

| gene | ctrl_1 | ctrl_2 | exp_1 | exp_2 | |
|-------|--------|--------|-------|-------|--|
| geneA | 10 | 11 | 56 | 45 | |
| geneB | 0 | 0 | 128 | 54 | |
| geneC | 42 | 41 | 59 | 41 | |
| geneD | 103 | 122 | 1 | 23 | |
| geneE | 10 | 23 | 14 | 56 | |
| geneF | 0 | 1 | 2 | 0 | |
| | | | | | |

<u>countData</u> is the count matrix (Number of reads coming from each gene for each sample) <u>colData</u>

| id | treatment | sex | |
|--------|-----------|--------|------|
| ctrl_1 | control | male | |
| ctrl_2 | control | female | |
| exp_1 | treated | male | |
| exp_2 | treated | female | •••• |

<u>colData</u> describes metadata about the *columns* of countData

N.B. First column of colData must match column names (i.e. sample names) of countData (-1st)





- Detailed, high-confidence consensus
- Biochemical reactions
- Small-scale, fewer genes
- Concentrated from decades of literature
- Simplified cellular logic, noisy
- Abstractions: directed, undirected
- Large-scale, genome-wide
- Constructed from omics data integration

Goal

1

Enrichment of fixed gene sets

Identification of pre-built pathways or networks that are enriched in a set of mutated or differentially expressed genes

2 De novo sub-network construction and clustering

Norte Class

Construction of specific sub-networks from the set of mutated or differentially expressed genes to identify an extended list of putative cancer genes



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1

Enrichment of fixed gene sets

Identification of pre-built pathways or networks that are enriched in a set of mutated or differentially expressed genes

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Pathway analysis (a.k.a. geneset enrichment) Limitations

Geneset annotation bias: can only discover what is already known

Side note.

- Non-model organisms: no high-quality genesets available
- Post-transcriptional regulation is neglected
- Tissue-specific variations of pathways are not annotated
 - e.g. NF-κB regulates metabolism, not inflammation, in adipocytes
- Size bias: stats are influenced by the size of the pathway
 - Many pathways/receptors converge to few regulators
 e.g. Tens of innate immune receptors activate four TFs: NF-kB, AP-1, IRF3/7, NFAT

Pathway & Network Analysis Overview



R Knowledge Check For BIMM-143

Do it Louiseir

This will be marked but not graded (*i.e.* will not factor into your course grade)

Time Limit: 40 mins