Gene Set Enrichment – Introduction

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Objective

Is expression of genes in a gene set associated with experimental condition?

► E.g., Are there unusually many up-regulated genes in the gene set?

Many methods, a review is Kharti et al., 2012.

- Over-representation analysis (ORA) are differentially expressed (DE) genes in the set more common than expected?
- Functional class scoring (FCS) summarize statistic of DE of genes in a set, and compare to null
- Issues with sequence data?
- Issues with single-cell data?

What is a gene set?

Any *a priori* classification of 'genes' into biologically relevant groups

- Members of same biochemical pathway
- Proteins expressed in identical cellular compartments
- Co-expressed under certain conditions
- Targets of the same regulatory elements
- On the same cytogenic band

Sets do not need to be...

- exhaustive
- disjoint

Collections of gene sets

Gene Ontology (GO) Annotation (GOA)

- ► CC Cellular Components
- ► BP Biological Processes
- ► MF Molecular Function

Pathways

- ► MSigDb
- ▶ KEGG
- reactome
- ► PantherDB

Collections of gene sets

E.g., MSigDb

- c1 Positional gene sets chromosome & cytogenic band
- c2 Curated Gene Sets from online pathway databases, publications in PubMed, and knowledge of domain experts.
- c3 motif gene sets based on conserved cis-regulatory motifs from a comparative analysis of the human, mouse, rat, and dog genomes.
- c4 computational gene sets defined by mining large collections of cancer-oriented microarray data.
- c5 GO gene sets consist of genes annotated by the same GO terms.
- c6 oncogenic signatures defined directly from microarray gene expression data from cancer gene perturbations.
- c7 immunologic signatures defined directly from microarray gene expression data from immunologic studies.



Work flow

- 1. Experimental design
- 2. Sequencing, quality assessment, alignment
- 3. Differential expression

and then...

- 4. Perform gene set enrichment analysis
- 5. Adjust for multiple comparisons

Approach 1: hypergeometric tests

- 1. Classify each gene as 'differentially expressed' DE or not, e.g., based on p < 0.05
- 2. Are DE genes in the set more common than DE genes not in the set?
- Fisher hypergeometric test.
 GOstats; limma::goana()
- Conditional hypergeometric to accommodate GO DAG, GOstats
- ► But: artificial division into two groups (DE vs. not DE)

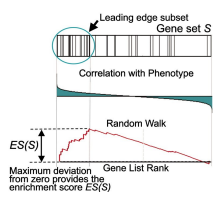
	In gene set?	
	Yes	No
DE	k	K
Not DE	n-k	N - K
NOT DE	11 – K	/v —

fisher.test()

Approach 2: enrichment score

Mootha et al., 2003; modified Subramanian et al., 2005.

- 1. Sort genes by log fold change
- Calculate running sum: incremented when gene in set, decremented when not.
- Maximum of the running sum is enrichment score ES; large ES means that genes in set are toward top of list.
- 4. Permuting subject labels for signficance

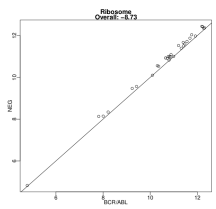


Subramanian et al., 2005, fig 1.

Approach 3: category *t*-test

E.g., Jiang & Gentleman, 2007; *Category*

- Summarize t (or other) statistic across genes in each set
- 2. Test for significance by permuting the subject labels
- 3. Much more straight-forward to implement



Expression in NEG vs BCR/ABL samples for genes in the 'ribosome' KEGG pathway; *Category* vignette.

Competitive versus self-contained null hypothesis

Goemann & Bühlmann, 2007

- ➤ Competitive null: The genes in the gene set do not have stronger association with the subject condition than other genes. Distinguishing more from less important sets. (Approach 1, 2)
- Self-contained null: The genes in the gene set do not have any association with the subject condition. Assessing individual sets. (Approach 3)
- Probably, self-contained null is closer to actual question of interest
- Permuting subjects (rather than genes) is appropriate

Approach 4: linear models

E.g., Hummel et al., 2008, GlobalAncova

- Colorectal tumors have good ('stage II') or bad ('stage III') prognosis. Do genes in the p53 pathway (just one gene set!) show different activity at the two stages?
- Linear model incorporates covariates sex of patient, location of tumor

limma

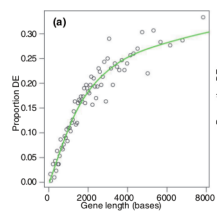
- Majewski et al., 2010 romer() and Wu & Smythe 2012 camera() for enrichment (competitive null) linear models
- Wu et al., 2010: roast(), mroast(), (and fry() − efficient) for self-contained null linear models

Approach 5: issues with sequence data?

- ► All else being equal, long genes receive more reads than short genes
- Per-gene P values proportional to gene size

E.g., Young et al., 2010, goseq

- Hypergeometric, weighted by gene size
- Substantial differences
- Better: read depth??



DE genes vs. transcript length. Points: bins of 300 genes. Line: fitted probability weighting function.

Approach 6: de novo discovery

- ➤ So far: analogous to supervised machine learning, where pathways are known in advance
- What about unsupervised discovery?

Example: Langfelder & Hovarth, WGCNA

- ► Weighted correlation network analysis
- Described in Langfelder & Horvath, 2008

Issues with single-cell data?

- Often, projections into reduced dimensions.
- Not genes *per se*, but weightings.
- ► An open issue, with opportunities for new methods!

Representing gene sets in R

- Named list(), where names of the list are sets, and each element of the list is a vector of genes in the set.
- data.frame() of set name / gene name pairs
- ► *GSEABase* input from standard file formats, representation as formal classes.

Benchmarks

A recent tweet from Levi Waldron provides a nice summary.

- ► *GSEABenchmarkR* for running benchmarks
- Self-contained tests often call random gene sets significant
- Hypergeometric test performs relatively well!

Conclusions

Gene set enrichment classifications

- ► Kharti et al: Over-representation analysis; functional class scoring; pathway topology
- ► Goemann & Bühlmann: Competitive vs. self-contained null

Selected Bioconductor packages (see biocViews)

Approach	Packages
Hypergeometric	GOstats, topGO, limma::goana()
Enrichment	<pre>limma::romer()</pre>
Category t-test	Category
Linear model	GlobalAncova, GSEAlm, limma::fry()
Pathway topology	SPIA
Sequence-specific	goseq
Visualization	pathview

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