## Introduction to Pathway and Network Analysis

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- High-throughput genetic/genomic technologies enable comprehensive monitoring of a biological system
- Analysis of high-throughput data typically yields a list of differentially expressed genes, proteins, metabolites...
  - Typically provides lists of single genes, etc.
  - Will use "genes" throughout, but using interchangeably mostly
- This list often fails to provide mechanistic insights into the underlying biology of the condition being studied
- How to extract meaning from a long list of differentially expressed genes → pathway/network analysis

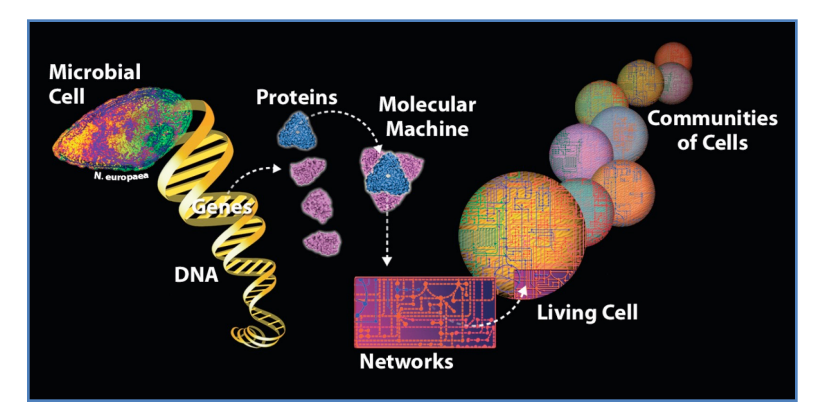
#### What makes an airplane fly?



Chas' Stainless Steel, Mark Thompson's Airplane Parts, About 1000 Pounds of Stainless Steel Wire, and Gagosian's Beverly Hills Space

#### From components to networks

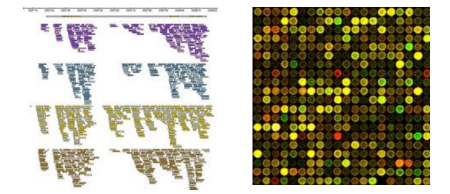
A biological function is a result of many interacting molecules and cannot be attributed to just a single molecule.

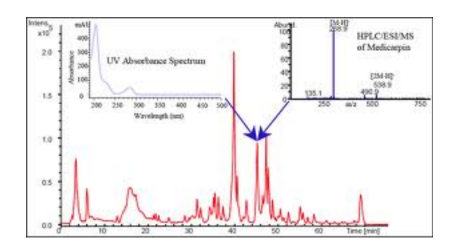


- One approach: simplify analysis by grouping long lists of individual genes into smaller sets of related genesreduces the complexity of analysis.
  - a large number of knowledge bases developed to help with this task
- Knowledge bases
  - describe biological processes, components, or structures in which individual genes \are known to be involved in
  - how and where gene products interact with each other

- Analysis at the functional level is appealing for two reasons:
  - First, grouping thousands of genes by the pathways they are involved in reduces the complexity to just several hundred pathways for the experiment
  - Second, identifying active pathways that differ between two conditions can have more explanatory power than a simple list of genes

- What kinds of data is used for such analysis?
  - Gene expression data
    - Microarrays
    - RNA-seq
  - Proteomic data
  - Metabolomics data
  - Single nucleotide
    polymorphisms (SNPs)





 What kinds of questions can we ask/answer with these approaches?



- The term "pathway analysis" gets used often, and often in different ways
  - applied to the analysis of Gene Ontology (GO) terms (also referred to as a "gene set")
  - physical interaction networks (e.g., protein–protein interactions)
  - kinetic simulation of pathways
  - steady-state pathway analysis (e.g., flux-balance analysis)
  - inference of pathways from expression and sequence data
- May or may not actually describe biological pathways

- For the first part of this module, we will focus on methods that exploit pathway knowledge in public repositories rather than on methods that infer pathways from molecular measurements
  - Use repositories such as GO or Kyoto Encyclopedia of Genes and Genomes (KEGG)

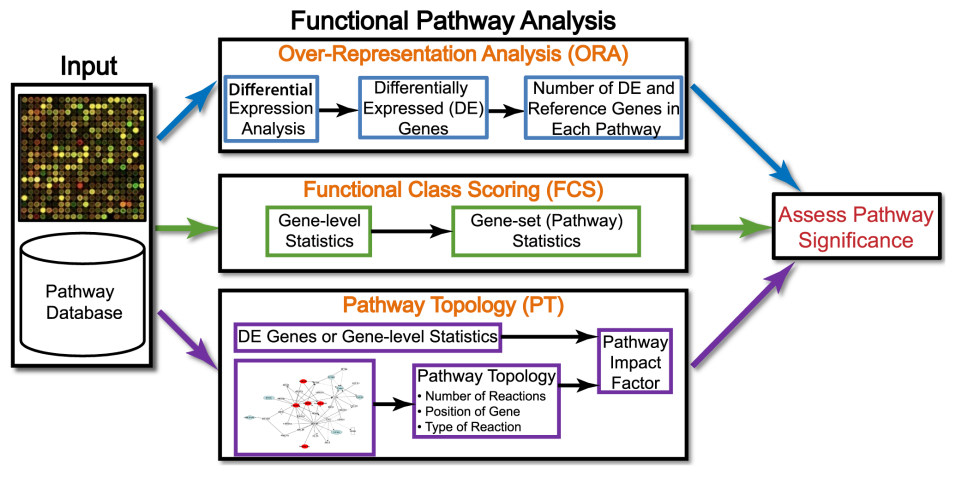
→ knowledge base-driven pathway analysis

#### A History of Pathway Analysis Approaches

 Over a decade of development of pathway analysis approaches

- Can be *roughly* divided into three generations:
  - 1<sup>st</sup>: Over-Representation Analysis (ORA)
    Approaches
  - 2<sup>nd</sup> : Functional Class Scoring (FCS) Approaches
  - 3<sup>rd</sup> : Pathway Topology (PT)-Based Approaches

Khatri P, Sirota M, Butte AJ. Ten years of pathway analysis: current approaches and outstanding challenges. PLoS Comput Biol. 2012;8(2):e1002375.



- The data generated by an experiment using a high-throughput technology (e.g., microarray, proteomics, metabolomics), along with functional annotations (pathway database) of the corresponding genome, are input to virtually all pathway analysis methods.
- ORA methods require that the input is a list of differentially expressed genes
- FCS methods use the entire data matrix as input
- PT-based methods additionally utilize the number and type of interactions between gene products, which may or may not be a part of a pathway database.
- The result of every pathway analysis method is a list of significant pathways in the condition under study.

#### Over-Representation Analysis (ORA) Approaches

- Earliest methods → over-representation analysis (ORA)
- Statistically evaluates the fraction of genes in a particular pathway found among the set of genes showing changes in expression
- It is also referred to as "2×2 table method" in the literature

#### **Over-Representation Analysis (ORA)**

- Uses one or more variations of the following strategy:
  - First, an input list is created using a certain threshold or criteria
    - For example, may choose genes that are differentially over- or under-expressed in a given condition at a false discovery rate (FDR) of 5%
  - Then, for each pathway, input genes that are part of the pathway are counted
  - This process is repeated for an appropriate background list of genes
    - (e.g., all genes measured on a microarray)
  - Next, every pathway is tested for over- or underrepresentation in the list of input genes
    - The most commonly used tests are based on the hypergeometric, chi-square, or binomial distribution

ORA tools
Onto-Express

GenMAPP

GoMiner

FatiGO

GOstat

**FuncAssociate** 

**GOToolBox** 

GeneMerge

GOEAST

ClueGO

FunSpec

GARBAN

GO:TermFinder

WebGestalt

agriGO

GOFFA

WEGO

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Web (http://vortex.cs.wayne.edu)

Web (http://gostat.wehi.edu.au)

Standalone (http://www.genmapp.org)

Web (http://babelomics.bioinfo.cipf.es)

Web (http://llama.mshri.on.ca/funcassociate/)

Web (http://omicslab.genetics.ac.cn/GOEAST/)

Web (http://bioinfo.vanderbilt.edu/webgestalt/)

Standalone (http://www.ici.upmc.fr/cluego/)

Web (http://funspec.med.utoronto.ca/)

Web (http://bioinfo.cau.edu.cn/agriGO/)

Web

Web (http://genome.crg.es/GOToolBox/)

Standalone, Web (http://discover.nci.nih.gov/gominer)

Standalone, Web (http://genemerge.cbcb.umd.edu/)

Standalone (http://search.cpan.org/dist/GO-TermFinder/)

Standalone, Web (http://edkb.fda.gov/webstart/arraytrack/)

Web (http://wego.genomics.org.cn/cgi-bin/wego/index.pl)

- First, the different statistics used by ORA are independent of the measured changes
  - (e.g., hypergeometric distribution, binomial distribution, chisquare distribution, etc.)
- Tests consider the number of genes alone but ignore any values associated with them
  - such as probe intensities
- By discarding this data, ORA treats each gene equally
  - Information about the extent of regulation (e.g., fold-changes, significance of a change, etc.) can be useful in assigning different weights to input genes/pathways
  - This can provide more information

- Second, ORA typically uses only the most significant genes and discards the others
  - input list of genes is usually obtained using an arbitrary threshold (e.g., genes with fold-change and/or p-values)
- Marginally less significant genes are missed, resulting in information loss
  - (e.g., fold-change = 1.999 or p-value = 0.051)
  - A few methods avoiding thresholds
    - They use an iterative approach that adds one gene at a time to find a set of genes for which a pathway is most significant

- Third, ORA assumes that each gene is independent of the other genes
- However, biology is a complex web of interactions between gene products that constitute different pathways
  - One goal might be to gain insights into how interactions between gene products are manifested as changes in expression
  - A strategy that assumes the genes are independent is significantly limited in its ability to provide insights
- Furthermore, assuming independence between genes amounts to "competitive null hypothesis" testing (more later), which ignores the correlation structure between genes
  - the estimated significance of a pathway may be biased or incorrect

- Fourth, ORA assumes that each pathway is independent of other pathways → NOT TRUE!
- Examples of dependence:
  - GO defines a biological process as a series of events accomplished by one or more ordered assemblies of molecular functions
  - The cell cycle pathway in KEGG where the presence of a growth factor activates the MAPK signaling pathway
    - This, in turn, activates the cell cycle pathway
- No ORA methods account for this dependence between molecular functions in GO and signaling pathways in KEGG

#### Functional Class Scoring (FCS) Approaches

- The hypothesis of functional class scoring (FCS) is that although large changes in individual genes can have significant effects on pathways, weaker but coordinated changes in sets of functionally related genes (i.e., pathways) can also have significant effects
- With few exceptions, all FCS methods use a variation of a general framework that consists of the following three steps.

- First, a gene-level statistic is computed using the molecular measurements from an experiment
  - Involves computing differential expression of individual genes or proteins
- Statistics currently used at gene-level include correlation of molecular measurements with phenotype
  - ANOVA
  - Q-statistic
  - signal-to-noise ratio
  - t-test
  - Z-score

- Choice of a gene-level statistic generally has a negligible effect on the identification of significantly enriched gene sets
  - However, when there are few biological replicates, a regularized statistic may be better
- Untransformed gene-level statistics can fail to identify pathways with up- and down-regulated genes
  - In this case, transformation of gene-level statistics (e.g., absolute values, squared values, ranks, etc.) is better

- Second, the gene-level statistics for all genes in a pathway are aggregated into a single pathwaylevel statistic
  - can be multivariate and account for interdependencies among genes
  - can be univariate and disregard interdependencies among genes
- The pathway-level statistics used include:
  - Kolmogorov-Smirnov statistic
  - sum, mean, or median of gene-level statistic
  - Wilcoxon rank sum
  - maxmean statistic

- Irrespective of its type, the power of a pathwaylevel statistic depends on
  - the proportion of differentially expressed genes in a pathway
  - the size of the pathway
  - the amount of correlation between genes in the pathway
- Univariate statistics show more power at stringent cutoffs when applied to real biological data, and equal power as multivariate statistics at less stringent cutoffs

- Assessing the statistical significance of the pathway-level statistic
- When computing statistical significance, the null hypothesis tested by current pathway analysis approaches can be broadly divided into two categories:
  - i) competitive null hypothesis
  - ii) self-contained null hypothesis
- A self-contained null hypothesis permutes class labels (i.e., phenotypes) for each sample and compares the set of genes in a given pathway with itself, while ignoring the genes that are not in the pathway
- A competitive null hypothesis permutes gene labels for each pathway, and compares the set of genes in the pathway with a set of genes that are not in the pathway

#### FCS tools

GSEA	Standalone (http://www.broadinstitute.org/gsea/)
sigPathway	Standalone (BioConductor)
Category	Standalone (BioConductor)
SAFE	Standalone (BioConductor)
GlobalTest	Standalone (BioConductor)
PCOT2	Standalone (BioConductor)
SAM-GS	Standalone (http://www.ualberta.ca/~yyasui/software.html)
Catmap	Standalone (http://bioinfo.thep.lu.se/catmap.html)
T-profiler	Web (http://www.t-profiler.org)
FunCluster	Standalone (http://corneliu.henegar.info/FunCluster.htm)
GeneTrail	Web (http://genetrail.bioinf.uni-sb.de)
GAzer	Web

Khatri P, Sirota M, Butte AJ. Ten years of pathway analysis: current approaches and outstanding challenges. PLoS Comput Biol. 2012;8(2):e1002375.

## Advantages of FCS Methods

FCS methods address three limitations of ORA

1. Don't require an arbitrary threshold for dividing expression data into significant and non-significant pools. Rather, FCS methods use all available molecular measurements for

Rather, FCS methods use all available molecular measurements for pathway analysis.

- 2. While ORA completely ignores molecular measurements when identifying significant pathways, FCS methods use this information in order to detect coordinated changes in the expression of genes in the same pathway
- 3. By considering the coordinated changes in gene expression, FCS methods account for dependence between genes in a pathway

## Limitations of FCS Methods

- First, similar to ORA, FCS analyzes each pathway independently
  - Because a gene can function in more than one pathway, meaning that pathways can cross and overlap
  - Consequently, in an experiment, while one pathway may be affected in an experiment, one may observe other pathways being significantly affected due to the set of overlapping genes
- Such a phenomenon is very common when using the GO terms to define pathways due to the hierarchical nature of the GO

## Limitations of FCS Methods

- Second, many FCS methods use changes in gene expression to rank genes in a given pathway, and discard the changes from further analysis
  - For instance, assume that two genes in a pathway, A and B, are changing by 2-fold and 20-fold, respectively
  - As long as they both have the same respective ranks in comparison with other genes in the pathway, most FCS methods will treat them equally, although the gene with the higher fold-change should probably get more weight
- Importantly, however, considering only the ranks of genes is also advantageous, as it is more robust to outliers.
  - A notable exception to this scenario is approaches that use gene-level statistics (e.g., t-statistic) to compute pathway-level scores.
  - For example, an FCS method that computes a pathway-level statistic as a sum or mean of the gene-level statistic accounts for a relative difference in measurements (e.g., Category, SAFE).

- A large number of publicly available pathway knowledge bases provide information beyond simple lists of genes for each pathway
  - KEGG
  - MetaCyc
  - Reactome
  - RegulonDB
  - STKE
  - BioCarta
  - PantherDB
  - ...
- Unlike GO and MSigDB, these knowledge bases also provide information about gene products that interact with each other in a given pathway, how they interact (e.g., activation, inhibition, etc.), and where they interact (e.g., cytoplasm, nucleus, etc.)

- ORA and FCS methods consider only the number of genes in a pathway or gene coexpression to identify significant pathways, and ignore the additional information available from these knowledge bases
  - Even if the pathways are completely redrawn with new links between the genes, as long as they contain the same set of genes, ORA and FCS will produce the same results
- Pathway topology (PT)-based methods have been developed to use the additional information
  - PT-based methods are essentially the same as FCS methods in that they perform the same three steps as FCS methods
  - The key difference between the two is the use of pathway topology to compute gene-level statistics

- Rahnenfuhrer et al. proposed ScorePAGE, which computes similarity between each pair of genes in a pathway (e.g., correlation, covariance, etc.)
  - similarity measurement between each pair of genes is analogous to gene-level statistics in FCS methods
  - averaged to compute a pathway-level score
- Instead of giving equal weight to all pairwise similarities, ScorePAGE divides the pairwise similarities by the number of reactions needed to connect two genes in a given pathway

- Impact factor (IF) analysis
  - IF considers the structure and dynamics of an entire pathway by incorporating a number of important biological factors, including changes in gene expression, types of interactions, and the positions of genes in a pathway

#### Ali will talk more about these approaches in detail!!!

## IF Analysis

- Briefly...
  - Models a signaling pathway as a graph, where nodes represent genes and edges represent interactions between them
  - Defines a gene-level statistic, called perturbation factor (PF) of a gene, as a sum of its measured change in expression and a linear function of the perturbation factors of all genes in a pathway
  - Because the PF of each gene is defined by a linear equation, the entire pathway is defined as a linear system
    - addresses loops in the pathways
  - The IF of a pathway (pathway-level statistic) is defined as a sum of PF of all genes in a pathway

 FCS methods that use correlations among genes implicitly assume that the underlying network, as defined by the correlation structure, does not change as the experimental conditions change

 This assumption may be inaccurate → PT approaches improve on this

- NetGSA accounts for the the change in correlation as well as the change in network structure as experimental conditions change
  - like IF analysis, models gene expression as a linear function of other genes in the network
- it differs from IF in two aspects
  - First, it accounts for a gene's baseline expression by representing it as a latent variable in the model
  - Second, it requires that the pathways be represented as directed acyclic graphs DAGs
    - If a pathway contains cycles, NetGSA requires additional latent variables affecting the nodes in the cycle.
    - In contrast, IF analysis does not impose any constraint on the structure of a pathway

### Limitations of PT-based Approaches

- True pathway topology is dependent on the type of cell due to cell-specific gene expression profiles and condition being studied
  - information is rarely available
  - fragmented in knowledge bases if available
  - As annotations improve, these approaches are expected to become more useful
- Inability to model dynamic states of a system
- Inability to consider interactions between pathways due to weak inter-pathway links to account for interdependence between pathways

PT-based tools	
ScorePAGE	No implementation available
Pathway-Express	Web (http://vortex.cs.wayne.edu)
SPIA	Standalone (BioConductor)
NetGSA	R package netgsa

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## **Outstanding Challenges**

- Broad Categories:
- 1. annotation challenges
- 2. methodological challenges

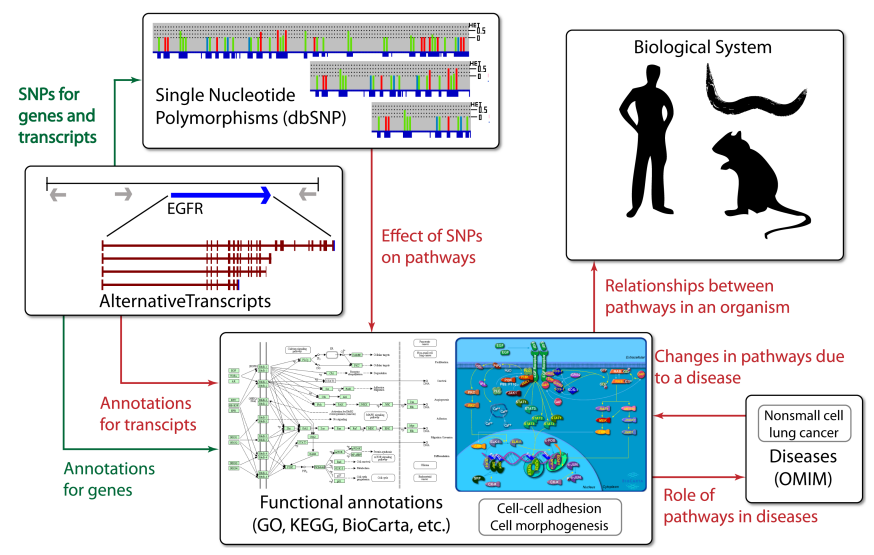
## **Outstanding Challenges**

- Next generation approaches will require improvement of the existing annotations
  - necessary to create accurate, high resolution knowledge bases with detailed condition-, tissue-, and cell-specific functions of each gene
    - PharmGKB ....
  - these knowledge bases will allow investigators to model an organism's biology as a dynamic system, and will help predict changes in the system due to factors such as mutations or environmental changes

### **Annotation Challenges**

- Low resolution knowledge bases
- Incomplete and inaccurate annotations
- Missing condition- and cell-specific information

Green arrows represent abundantly available information, and red arrows represent missing and/or incomplete information. The ultimate goal of pathway analysis is to analyze a biological system as a large, single network. However, the links between smaller individual pathways are not yet well known. Furthermore, the effects of a SNP on a given pathway are also missing from current knowledge bases. While some pathways are known to be related to a few diseases, it is not clear whether the changes in pathways are the cause for those diseases or the downstream effects of the diseases.



## Low Resolution Knowledge Bases

- Knowledge bases not as high resolution as technologies
  - using RNA-seq, more than 90% of the human genome is estimated to be alternatively spliced
  - multiple transcripts from the same gene may have related, distinct, or even opposing functions
  - GWAS have identified a large number of SNPs that may be involved in different conditions and diseases.
  - However, current knowledge bases only specify which genes are active in a given pathway
  - Essential that they also begin specifying other information, such as transcripts that are active in a given pathway or how a given SNP affects a pathway

### Low Resolution Knowledge Bases

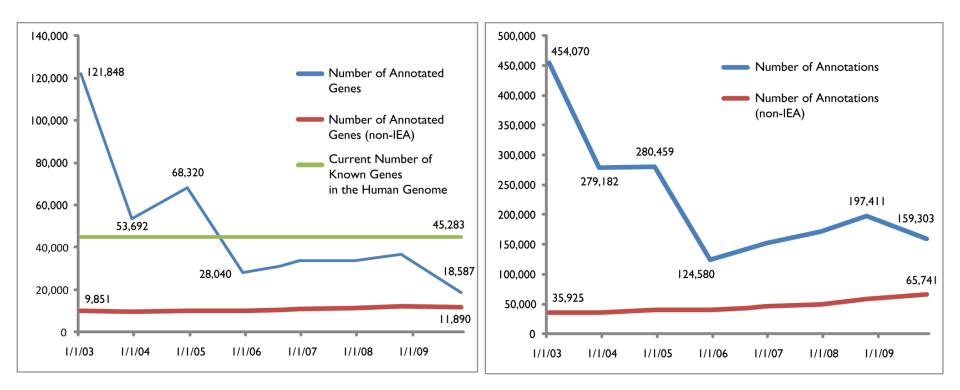
- Because of these low resolution knowledge bases, every available pathway analysis tool first maps the input to a non-redundant namespace, typically an Entrez Gene ID
  - this type of mapping is advantageous, although it can be nontrivial, as it allows the existing pathway analysis approaches to be independent of the technology used in the experiment
  - However, mapping in this way also results in the loss of important information that may have been provided because a specific technology was used
    - XRN2a, a variant of gene XRN2, is expressed in several human tissues, whereas another variant of the same gene, XRN2b, is mainly expressed in blood leukocytes
    - Although RNA-seq can quantify expression of both variants, mapping both transcripts to a single gene causes loss of tissue-specific information, and possibly even condition-specific information

### Low Resolution Knowledge Bases

- Therefore, before pathway analysis can exploit current and future technological advances in biotechnology, it is critically important to annotate exact transcripts and SNPs that participate in a given pathway
- While new approaches are being developed in this regard, they may not yet be adequate
  - Braun et al. proposed a method for analyzing SNP data from a GWAS
  - Still relies on mapping multiple SNPs to a single gene, followed by gene-to-pathway mapping

- A surprisingly large number of genes are still not annotated
- Many of the genes are hypothetical, predicted, or pseudogenes
  - Although the number of protein-coding genes in the human genome is estimated to be between 20,000 and 25,000, according Entrez Gene, there are 45,283 human genes, of which 14,162 are pseudogenes
  - One could argue that the pseudogenes should not be included when evaluating functional annotation coverage
  - pseudogene-derived small interfering RNAs have been shown to regulate gene expression in mouse oocytes
  - GO provides annotations for 271 pseudogenes
  - A widely used DNA microarray, Affymetrix HG U133 plus 2.0, contains 1,026 probe sets that correspond to 823 pseudogenes
  - Should pseudogenes be included in the count when estimating annotation coverage for the human genome?

Number of GO-annotated genes (left panel) and number of GO annotations (right panel) for human from January 2003 to November 2009.As the estimated number of known genes in the human genome is adjusted (between January 2003 and December 2003) and annotation practices are modified (between December 2004 and December 2005, and between October 2008 and November 2009), one can argue that, although the number of annotated genes and the annotations are decreasing (which is mainly due to the adjusted number of genes in the human genome and changes in the annotation process), the quality of annotations is improving, as demonstrated by the steady increase in non-IEA annotations and the number of genes with non-IEA annotations. However, the increase in the number of genes with non-IEA annotations is very slow. In almost 7 years, between January 2003 and November 2009, only 2,039 new genes received non-IEA annotations. At the same time, the number of non-IEA annotations increased from 35,925 to 65,741, indicating a strong research bias for a small number of genes.doi:10.1371/journal.pcbi.1002375.g003



- Additionally, many of the existing annotations are of low quality and may be inaccurate
  - >90% of the annotations in the October 2015 release of GO had the evidence code "inferred from electronic annotations (IEA)"
  - the only ones in GO that are not curated manually
  - Annotations inferred from indirect evidence are considered to be of lower quality than those derived from direct experimental evidence
  - If the annotations with IEA code are removed, the number of genes with good quality annotations in the November 2015 release of human GO annotations is reduced from ~18K to ~12K

- It is very likely that the reduced number of annotations and annotated genes since January 2003 is an indicator of improving quality
- This is due in part to the fact that the number of genes in a genome are continuously being adjusted and the functional annotation algorithms are being improved
  - the number of non-IEA annotations is continuously increasing
- However, the rate of increase for non-IEA annotations is very slow (approximately 2,000 genes annotated in 7 years)

- Manual curation of the entire genome is expected to take a very long time (~13–25 years)
- Entire research community could participate in the curation process
- One approach to facilitate participation of a large number of researchers is to adopt a standard annotation format similar to Minimum Information About a Microarray Experiment (MIAME)
  - should this be required like GEO?
- A format for functional annotation can be designed or adopted from the existing formats (e.g., BioPAX, SBML)
  - Such a format could allow researchers to specify an experimentally confirmed role of a specific transcript or a SNP in a pathway along with experimental and biological conditions

- Most pathway knowledge bases are built by curating experiments performed in different cell types at different time points under different conditions
- These details are typically not available in the knowledge bases!
- One effect of this omission is that multiple independent genes are annotated to participate in the same interaction in a pathway
- This effect is so widespread that many pathway knowledge bases represent a set of distinct genes as a single node in a pathway

- Example: *Wnt/beta-catenin pathway in STKE* 
  - the node labeled "Genes" represents 19 genes directly targeted by Wnt in different organisms (Xenopus and human) in different cells and tissues (colon carcinoma cells and epithelial cells
  - these non-specific genes introduce bias for these pathways in all existing analysis approaches
  - For instance, any ORA method will assign higher significance (typically an order of magnitude lower pvalue) to a pathway with more genes
  - Similarly, more genes in a pathway also increase the probability of a higher pathway-level statistic in FCS approaches, yielding higher significance for a given pathway.

- This contextual information is typically not available from most of the existing knowledge bases
- A standard functional annotation format discussed above would make this information available to curators and developers
  - For instance, the recently proposed Biological Connection Markup Language (BCML) allows pathway representation to specify the cell or organism in which each pathway interaction occurs.
  - BCML can generate cell-, condition-, or organism-specific pathways based on user-defined query criteria, which in turn can be used for targeted analysis

- Existing knowledge bases do not describe the effects of an abnormal condition on a pathway
  - For example, it is not clear how the Alzheimer's disease pathway in KEGG differs from a normal pathway
  - Nor it is clear which set of interactions leads to Alzheimer's disease
- We are now understanding that context plays an important role in pathway interactions
- Information about how cell and tissue type, age, and environmental exposures affect pathway interactions will add complexity that is currently lacking

## Methodological Challenges

Benchmark data sets for comparing different methods

Inability to model and analyze dynamic response

• Inability to model effects of an external stimuli

# **Comparing Different Methods**

- How do we compare different pathway analysis methods?
- Simulated data
  - Advantages:
    - Real signal is simulated, so "true" answer is known
  - Disadvantages
    - Cannot contain all the complexity of real data
    - The success of the methods can reflect the similarity of how well the simulation matches the knowledgebase structure used

# **Comparing Different Methods**

- Benchmark data
  - Advantages:
    - Can compare sensitivity and specificity
    - Several datasets have been consistently used in the literature
    - Includes all the complexity of real biological data
  - Disadvantages
    - Affected by confounding factors
      - absence of a pure division into classes
      - presence of outliers
      - ....
    - No true answer known for grounded comparisons actual biology isnt known

# **Comparing Different Methods**

- A general challenge: *Different definitions of the same pathway in different knowledge bases can affect performance assessment* 
  - GO defines different pathways for apoptosis in different cells
    - (e.g., cardiac muscle cell apoptosis, B cell apoptosis, T cell apoptosis)
    - Further distinguishes between induction and regulation of apoptosis
  - KEGG defines a single signaling pathway for apoptosis
    - does not distinguish between induction and regulation
  - An approach using KEGG would identify a single pathway as significant, whereas GO could identify multiple pathways, and/or specific aspects of a single apoptosis pathway

# Inability to model and analyze dynamic response

- No existing approach can collectively model and analyze high-throughput data as a single dynamic system
- Current approaches analyze a snapshot assuming that each pathway is independent of the others at a given time
  - measure expression changes at multiple time points, and analyze each time point individually
  - Implicitly assumes that pathways at different time points are independent
- Need models that accounts for dependence among pathways at different time points
  - Much of this limitation is due to technology/experimental design → not all bioinformatics limitations

# Inability to model effects of an external stimuli

- Gene set—based approaches often only consider genes and their products
- Completely ignore the effects of other molecules participating in a pathway
  - such as the rate limiting step of a multi-step pathway.
- Example:
  - The amount/strength of Ca<sup>2+</sup> causes different transcription factors to be activated
  - This information is usually not available.

## Summary

- In the last decade, pathway analysis has matured, and become the standard for trying to dissect the biology of high throughput experiments.
- Many similarities across the three main generations of pathway analysis tools.
- Will discuss more details of some of these choices, knowledge bases, and specific approaches next.
- Many open methods development challenges!

## **Overview of Module**

- First Half:
  - Overview of gene set and pathway analysis
    - Commonly used databases and annotation issues
    - 1<sup>st</sup> and 2<sup>nd</sup> generation tools
      - Basic differences in methods
      - Details on very popular methods
    - Issues with different "omics" datatypes
- Second Half
  - "3<sup>rd</sup> generation" methods
  - Network analysis modeling

#### Questions?

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