A quantum leap in the reproducibility, precision, and sensitivity of gene-expression-profile analysis even when sample size is extremely small

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(Based on the PhD dissertation of Kevin Lim)



Why small sample size?



- Biological constraint
 - Comparing cell lines
 - Comparing mutants vs wildtype
- Rare-sample constraint
- Population-size constraint
 - Singapore is small, we often wait a long time for enough patients presenting the desired phenotype
- Cost constraint

Outline



- Ideals of a perfect method for gene selection in gene expression profile analysis
- Failure of commonly-used methods
- Reproducible precise & sensitive selection of genes, even when sample size is extremely small
- Reliable accurate cross-batch classification, even when batch effect is severe and sample size is small



THE IDEAL

A perfect method for identifying causal factors of a disease



- A perfect method should ...
 - Report all causal factors present in a dataset
 - Not report any non-factor
- ⇒ When it is applied to two representative datasets of the disease, the two sets of identified factors should be the same
- ⇒ When it is applied to a subset of a dataset, the set of identified factors should be a subset of the set of identified factors when it is applied to the whole dataset



THE REALITY

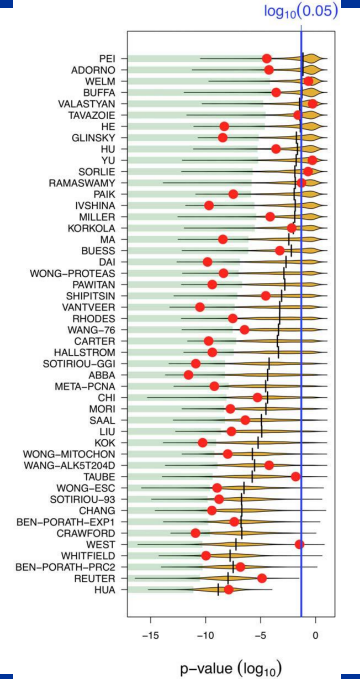
Percentage of overlapping genes



- Low % of overlapping genes from diff expt in general
 - Prostate cancer
 - Lapointe et al, 2004
 - Singh et al, 2002
 - Lung cancer
 - Garber et al, 2001
 - Bhattacharjee et al,
 2001
 - DMD
 - Haslett et al, 2002
 - Pescatori et al, 2007

Datasets	DEG	POG		
Prostate	Top 10	0.30		
Cancer	Top 50	0.14		
	Top100	0.15		
Lung	Top 10	0.00		
Cancer	Top 50	0.20		
	Top100	0.31		
DMD	Top 10	0.20		
DMD	Top 50	0.42		
	Top100	0.54		

Zhang et al, Bioinformatics, 2009



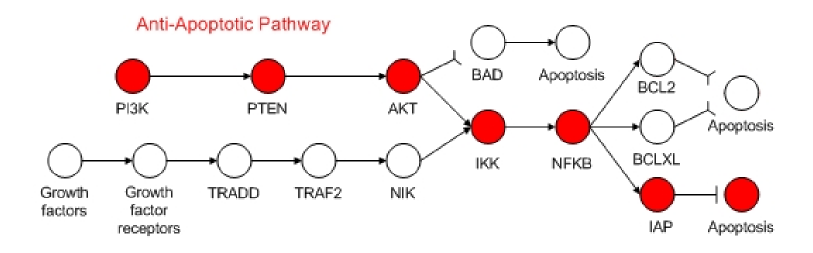


"Most random gene expression signatures are significantly associated with breast cancer outcome"

Venet et al., PLoS Comput Biol, 7(10):e1002240, 2011.

Gene regulatory circuits

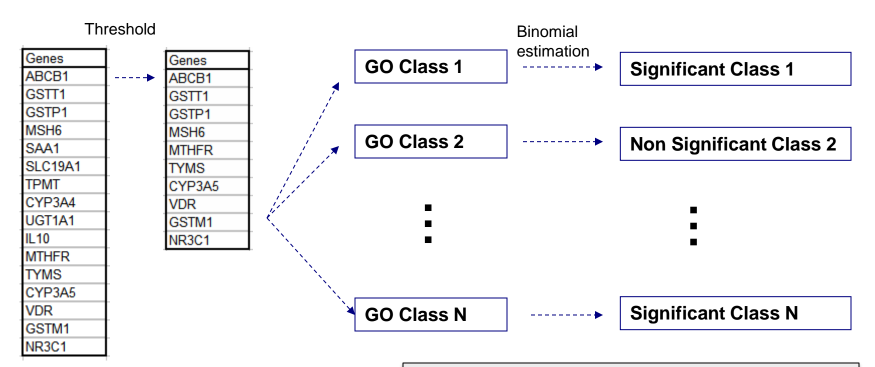




- Each disease has some underlying cause
- There is some unifying biological theme for genes that are truly associated with a disease

Overlap analysis: ORA





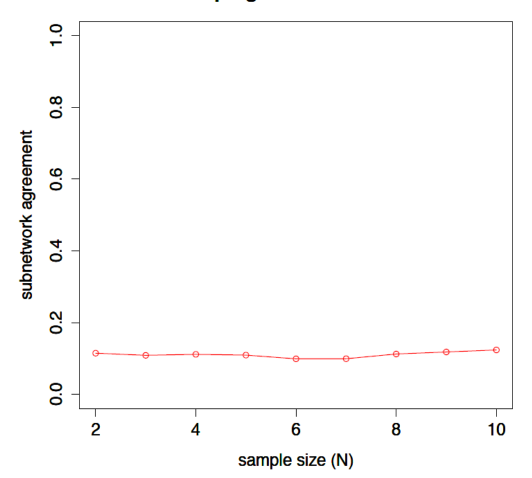
ORA tests whether a pathway is significant by intersecting the genes in the pathway with a pre-determined list of DE genes (we use all genes whose t-statistic meets the 5% significance threshold), and checking the significance of the size of the intersection using the hypergeometric test

S Draghici et al. "Global functional profiling of gene expression". *Genomics*, 81(2):98-104, 2003.

Disappointing performance



upregulated in DMD



DMD gene expression data

- Pescatori et al., 2007
- Haslett et al., 2002

Pathway data

• PathwayAPI, Soh et al., 2010

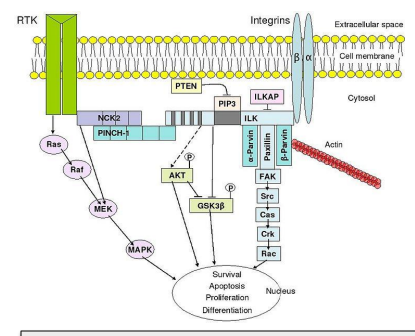


THE REASONS

Issue #1 with ORA



- Its null hypothesis basically says "Genes in the given pathway behaves no differently from randomly chosen gene sets of the same size"
- This null hypothesis is obviously false
- ⇒ Lots of false positives

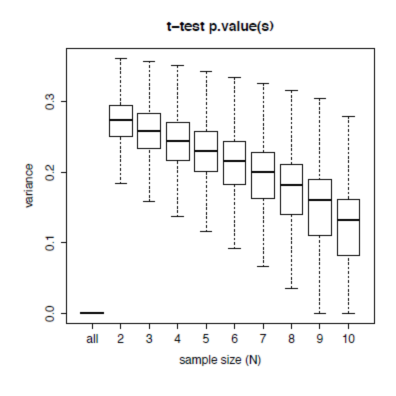


 A biological pathway is a series of actions among molecules in a cell that leads to a certain product or a change in a cell. Thus necessarily the behavour of genes in a pathway is more coordinated than random ones

Issue #2 with ORA



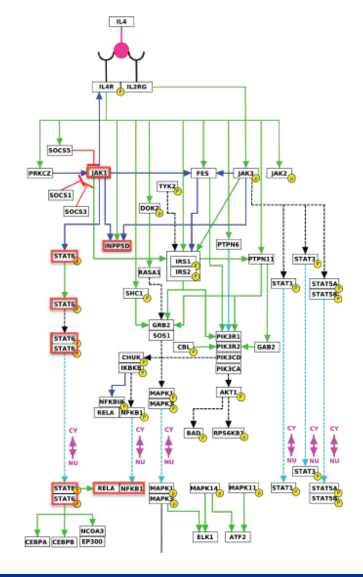
- It relies on a predetermined list of DE genes
- This list is sensitive to the test statistic used and to the significance threshold used
- This list is unstable regardless of the threshold used when sample size is small



Issue #3 with ORA

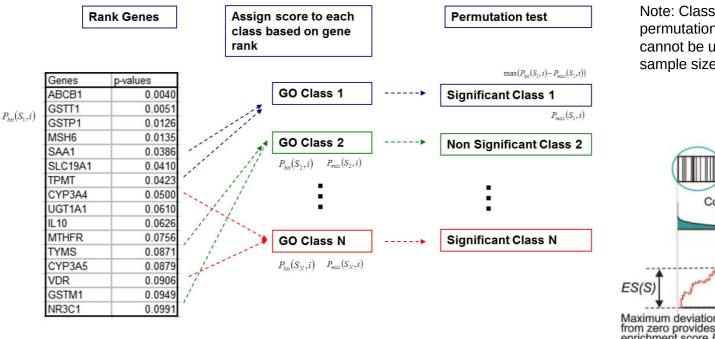


- It tests whether the entire pathway is significantly differentially expressed
- If only a branch of the pathway is relevant to the phenotypes, the noise from the large irrelevant part of the pathways can dilute the signal from that branch

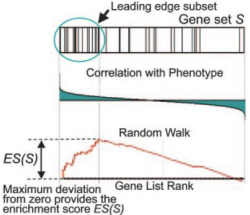


GSEA in gene-permutation mode





Note: Class label permutation mode cannot be used when sample size is small

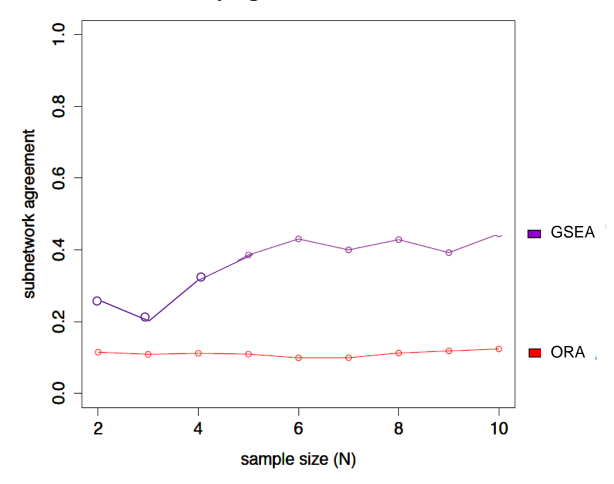


- Issue #2 & #3 solved to different degrees
 - Does not need pre-determined list of DE genes, but gene ranking (based on t-test p-value) is still unstable for small sample size
 - Irrelevant genes in pathway have only small effect on the ES(S) peak
- Issues #1 (when sample size is small) is unsolved

Better performance, but not great



upregulated in DMD



PFSNet: Exploiting subnetworks



 Induce subnetworks from pathways by considering only genes highly expressed in majority of patients in any class

$$\beta_1^*(g_i) = \sum_{\substack{p_j \in D}} \frac{fs(e_{g_i,p_j})}{|D|} \qquad \qquad \beta_2^*(g_i) = \sum_{\substack{p_j \in \neg D}} \frac{fs(e_{g_i,p_j})}{|\neg D|} \qquad \qquad \text{-ve class wt}$$
 Score of subnet S in patient k w/ +ve class wt +ve class wt}
$$Score_1^{p_k}(S) = \sum_{\substack{g_i \in S}} fs(e_{g_i,p_k}) * \beta_1^*(g_i) \qquad Score_2^{p_k}(S) = \sum_{\substack{g_i \in S}} fs(e_{g_i,p_k}) * \beta_2^*(g_i)$$

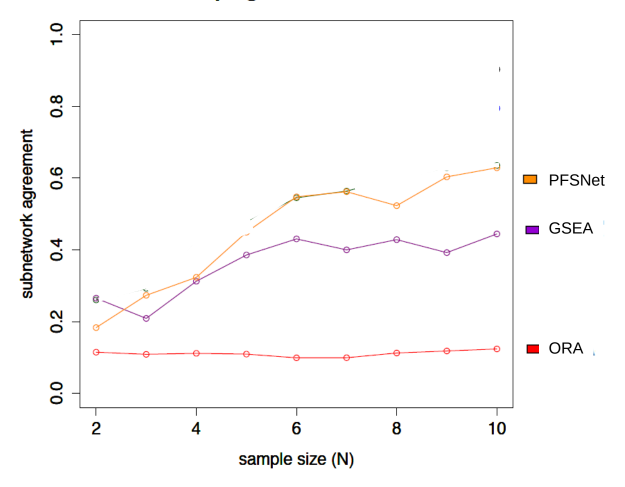
- For an irrelevant subnetwork S, the two scores above for each patient P_k should be roughly equal, regardless of his class
- Do a paired t-test to decide whether S is relevant
 - Get null distribution by permuting class labels

- β weights become unstable
- Cannot generate null distribution
- All 3 issues solved, but not when sample size is small

Much better performance, but still not great



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THE QUANTUM LEAP

EVEN WHEN SAMPLE SIZE IS EXTREMELY SMALL

ORA-Paired: Paired test and new null hypothesis



- Let g_i be genes in a given pathway P
- Let p_i be a patient
- Let q_k be a normal
- Let $\Delta_{i,j,k} = Expr(g_i,p_j) Expr(g_i,q_k)$
- Test whether $\Delta_{i,j,k}$ is a distribution with mean 0

Issue #1 is solved

 Null hypothesis is "Pathway P is irrelevant to the difference between patients and normals, and the genes in P behave similarly in patients and normals"

Issue #2 is solved

- No longer need a pre-determined list of DE genes
- Issue #3 is unsolved
- Is sample size now larger?
 - |patients| * |normals| * |genes in P|

Testing the null hypothesis



"Pathway P is irrelevant to the difference between patients and normals and so, the genes in P behave similarly in patients and normals"

Method #1

- T-test w/ a conservative degree of freedom
 - E.g., # normals + # patients

Method #2

- By the null hypothesis, a dataset and any of its class-label permutations are exchangeable
- ⇒ Get null distribution by class-label permutations
 - Only for large-size sample

Method #3

- Modified null hypothesis
 - "Pathway P induces genegene correlations, and genes in P behave according to these genegene correlations;
 - P is irrelevant to the diff betw patients and normals and so, genes in P behave similarly in patients and normals"
- ⇒ Get null distribution using datasets that conserve gene-gene correlations in the original dataset
 - E.g., array rotation

Array rotation



QR decomposition

$$X = X_{Q} \cdot X_{R}$$

Where

- X is gene expression array of n samples * m genes
- X_Q is n * r orientation matrix, r is rank of X
- X_R is sufficient statistics of covariance between the m genes

Rotation

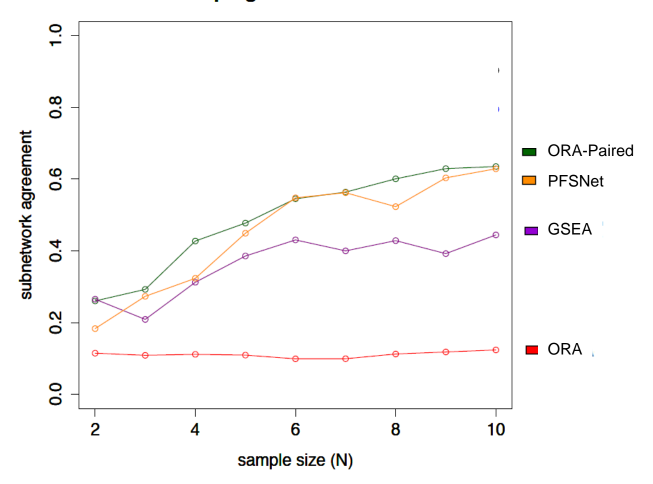
$$X' = R_Q \cdot X_Q \cdot X_R$$

Where

- R_Q is an n * n rotation operation
- ⇒ X' is rotation of X preserving gene-gene correlations
- ⇒ i.e., preserving constraints induced by pathways

Similar to PFSNet, good but not great National of Singar

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NEA-Paired: Paired test on subnetworks



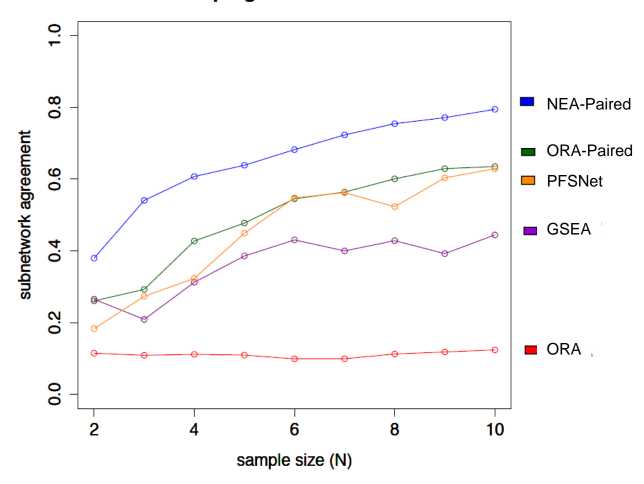
- Given a pathway P
- Let each node and its immediate neighbourhood in P be a subnetwork
- Apply ORA-Paired on each subnetwork individually

- Issues #1 & #2 are solved as per ORA-Paired
- Issue #3 is partly solved
 - Testing subnetworks instead of whole pathways
 - But subnetworks derived in a fragmented way

Much better performance



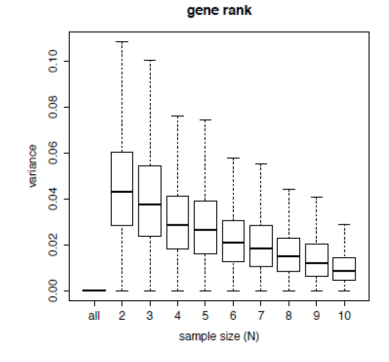
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ESSNet: Larger subnetworks



- Compute the average rank
 of a gene based on its
 expression level in
 patients in any class
- Use the top α% to extract large connected components in pathways
- Test each component using ORA-Paired

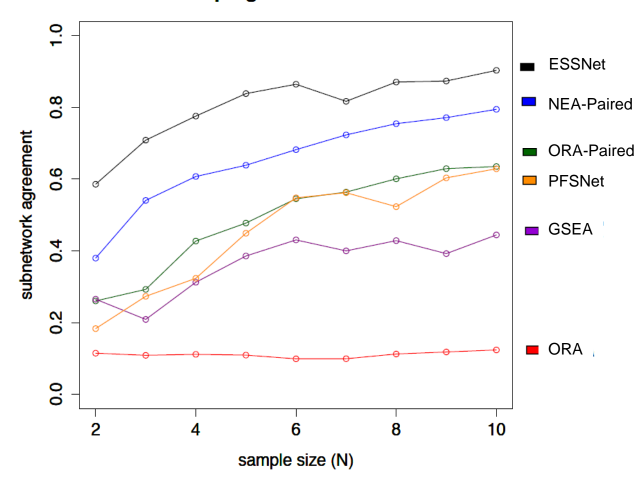


- Gene rank is very stable
- Issues #1 #3 solved

Fantastic performance

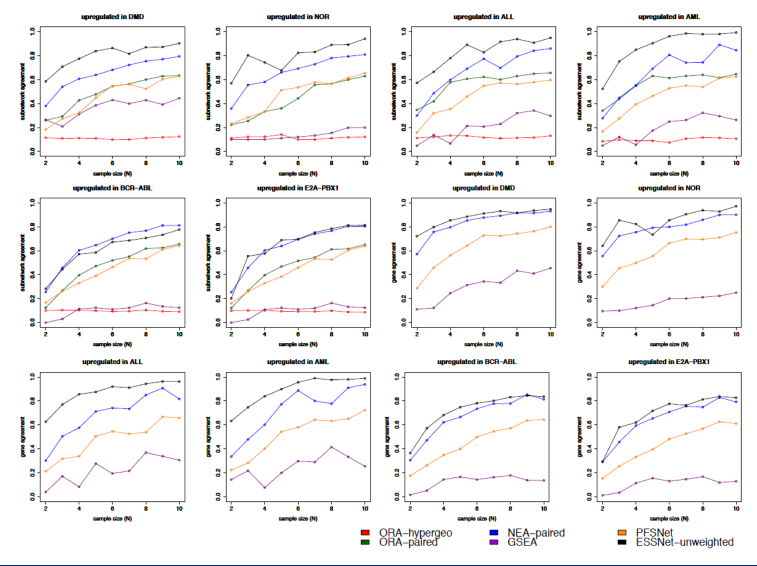


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More datasets tested





ESSNet is unlikely to report junk

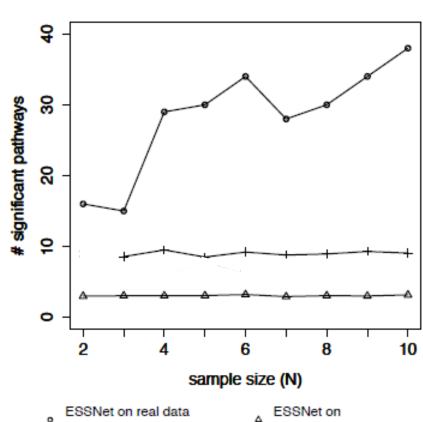


Table 4.2: Average number of subnetworks predicted by ESSNet over the sample sizes (N); the first number denotes the number of subnetworks in the numerator of the subnetwork-level agreement and the second number denotes the number of subnetworks in the denominator of the subnetwork-level agreement; cf. equation 4.5.

		DMD	ALL	BCR
	2	8.2/13.4	7.0/11.9	4.8/12.6
	3	11.1/15.9	11.3/17.9	5.0/11.7
Ξ	4	13.18/16.5	11.9/15.9	6.2/10.4
size	5	14.2/16.7	14.6/18.3	7.9/12.7
sample si	6	15.14/17.6	14.9/18.0	11.0/15.7
	7	15.2/17.4	16.1/19.2	12.9/17.5
	8	15.4/17.5	16.2/19.0	15.3/20.4
	9	16.6/18.8	17.0/19.8	15.8/20.8
	10	17.6/19.7	17.3/19.7	16.2/20.8



A negative-control experiment showing that ESSNet does not report junk



- (Pescatori et al. 2007)
- random negative control
- random negative control

ESSNet also dominates when sample size is large



Table 4.3: Number of subnetworks predicted by the various methods on a full dataset where the null distribution is computed using array rotation (rot), class-label swapping (cperm) and gene swapping (gswap); the first number denotes the number of subnetworks in the numerator of the subnetwork-level agreement and the second number denotes the number of subnetworks in the denominator of the subnetwork-level agreement; cf. equation 4.5.

	DI	MD	A	LL	BCR		
	rot	cperm	rot	cperm	rot	cperm	
ESSNet	20/23	13/15	22/24	25/27	24/29	30/32	
NEA-paired	77/98	91/115	140/163	109/119	176/192	37/43	
ORA-paired	30/62	30/62	34/74	34/74	53/99	53/99	
ORA-hypergeo	20/46	41/141	24/60	48/73	4/14	32/166	
	cperm	gswap	cperm	gswap	cperm	gswap	
GSEA	23/64	24/69	8/52	17/48	7/57	5/46	

Do ESSNet results agree on small datasets vs big datasets?



		Precision						Recall					
		DMD		ALL		BCR		DMD		ALL		BCR	
		D	$\neg D$	D	$\neg \mathbf{D}$	D	$\neg D$	D	$\neg \mathbf{D}$	D	$\neg \mathbf{D}$	D	$\neg \mathbf{D}$
	2	0.96	0.88	0.87	0.95	0.93	0.91	0.45	0.31	0.34	0.25	0.19	0.17
	3	0.93	0.86	0.99	0.89	0.90	0.87	0.56	0.45	0.56	0.41	0.21	0.16
(N)	4	0.88	0.88	0.97	0.92	0.91	0.87	0.67	0.50	0.51	0.53	0.35	0.48
	5	0.89	0.88	0.94	0.90	0.89	0.90	0.73	0.52	0.74	0.55	0.36	0.38
- 66	6	0.82	0.88	0.93	0.92	0.89	0.91	0.78	0.62	0.74	0.62	0.44	0.438
Ĕ	7	0.85	0.86	0.95	0.93	0.90	0.87	0.75	0.59	0.66	0.64	0.55	0.53
sample size	8	0.84	0.89	0.97	0.94	0.90	0.92	0.81	0.69	0.74	0.66	0.61	0.66
	9	0.88	0.90	0.94	0.92	0.89	0.89	0.90	0.67	0.76	0.74	0.65	0.67
	10	0.88	0.93	0.97	0.92	0.90	0.90	0.86	0.84	0.89	0.74	0.66	0.73

- Use ESSNet's results on entire datasets as the benchmark to evaluate ESSNet's results on small subsets of the datasets
- The precision (i.e., agreement) is superb, though some subnetworks are missed when smaller datasets are analysed

Leukemias: IL-4 signaling IL4R IL2RG LEGEND JAK1 JAK3 JAK2 FES Ligand mRNA TYK2 Enzyme complex Receptor DOK2 Protein-protein interaction Protein - protein dissociation eads to through unknown mechanis PTPN6 INPP5D ositive regulation of gene expressio STAT3 PTPN11 egative regulation of gene expressio IRS2 RASA1 Auto catalysis STATI, Acetylation STAT5A SHC1 Deacetylation STATS Phosphorylation Dephosphorylation Sumoviation GRB2 Desumoylation 5051 PIK3R1 Ubiquitination PIK3R2 GAB2 Deubiquitination CHUK, PIKECD Methylation IKBKB РІКЗСА Demethylation *almitoylation AKT1, roteolytic cleavag MAPK3 Inhibition NEKBIA

CY

NU WNU

STATI

STAT3

CY CY

Protein

MU

induced catalysis

Translocation

Cytoplasm Extracellular

Endosome

Nucleus

Plasma membrane

Endoplasmic reticulum Golgi apparatus Mitochondrion



For the Leukemia dataset (in which patients are either classified to have acute lymphoblastic leukemia or acute myeloid leukemia), one of the significant subnetworks that is biologically relevant is part of the Interleukin-4 signaling pathway; see figure 6b (supplementary material). The binding of Interleukin-4 to its receptor (Cardoso et al., 2008) causes a cascade of protein activation involving JAK1 and STAT6 phosphorylation. STAT6 dimerizes upon activation and is transported to the nucleus and interacts with the RELA/NFKB1 transcription factors, known to promote the proliferation of T-cells (Rayet and Gelinas, 1999). In contrast, acute myeloid leukemia does not have genes in this subnetwork up-regulated and are known to be unrelated to lymphocytes.

NCOA3

EP300

RELA NEKBI

RELA NFKB1

NU

MAPK1

BAD.

MAPK14

RPS6KB)

MAPK11

ATF2

SOC55

SOCS1

STATE.

STAT6

STAT6

STAT6

NU

STATE

STAT6

CEBPA CEBPB

SOCS3

PRKCZ

Remarks



- Consistent successful gene expression profile analysis needs deep integration of background knowledge
- Most gene expression profile analysis methods fail to give reproducible results when sample size is small (and some even fail when sample size is quite large)
- Logical analysis to identify key issues and simple logical solution to the issues can give fantastic results



DIFFICULTY OF CROSS-BATCH CLASSIFICATION

Batch effects



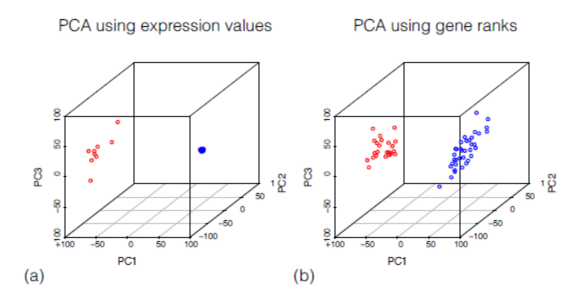
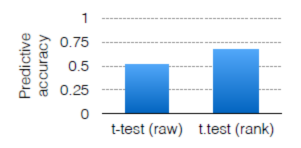


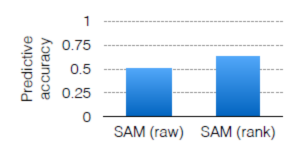
Figure 5.1: Batch effects in the DMD/NOR datasets, the blue and red color denote different data batches. (a) Scatterplot on the first 3 components using gene-expression values. (b) Scatterplot on the first 3 components using gene ranks.

- Batch effects are common
- Batch effects cannot always be removed using common normalization methods

Gene-feature-based classifiers do badly when there are batch effects, even after normalization







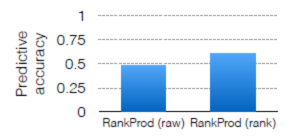
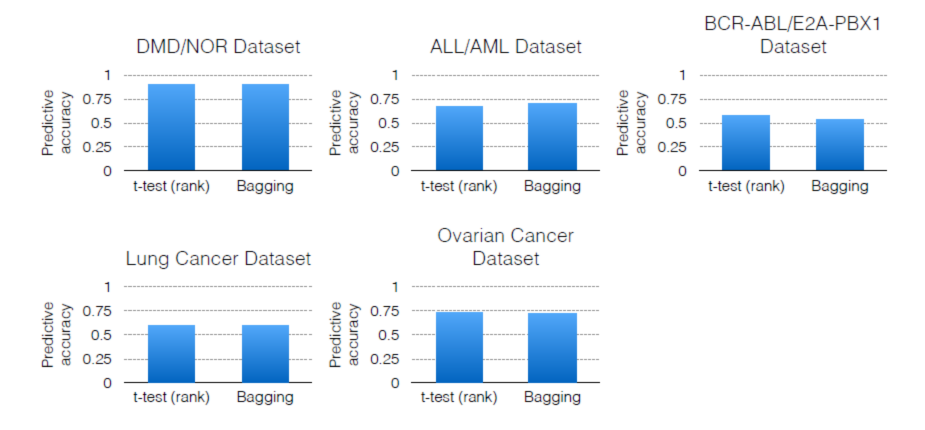


Figure 5.8: Predictive accuracy of gene-feature-based classifiers with and without rank normalization in the DMD/NOR dataset.

Gene selection by t-test, SAM, or rank product. Classifier by naïve Bayes

Ensemble classifiers can't always improve results of gene-feature-based classifiers with normalization





Genes from subnetworks produced by ESSNet can't help gene-featurebased classifiers



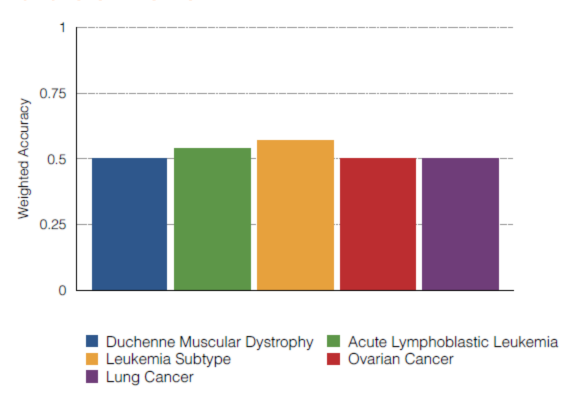


Figure 5.15: Predictive accuracy of gene-feature-based classifier using genes extracted from subnetworks in ESSNet; demonstrating that genes in the subnetworks by themselves are not a good discriminator for classification



SUCCESSFUL CROSS-BATCH CLASSIFICATION

WHEN SAMPLE SIZE IS LARGE

PFSNet-based features



PFSNet

- Induce subnetworks from pathways by considering only genes highly expressed in majority of patients in any class
- For each subnetwork S and each patient P_k, compute a pair of scores:

$$\beta_1^*(g_i) = \sum_{p_j \in D} \frac{fs(e_{g_i, p_j})}{|D|} \qquad \qquad \beta_2^*(g_i) = \sum_{p_j \in \neg D} \frac{fs(e_{g_i, p_j})}{|\neg D|}$$

$$Score_1^{p_k}(S) = \sum_{g_i \in S} fs(e_{g_i, p_k}) * \beta_1^*(g_i) \quad Score_2^{p_k}(S) = \sum_{g_i \in S} fs(e_{g_i, p_k}) * \beta_2^*(g_i)$$

Straightforward to use these scores as features

Successfully reducing batch effects



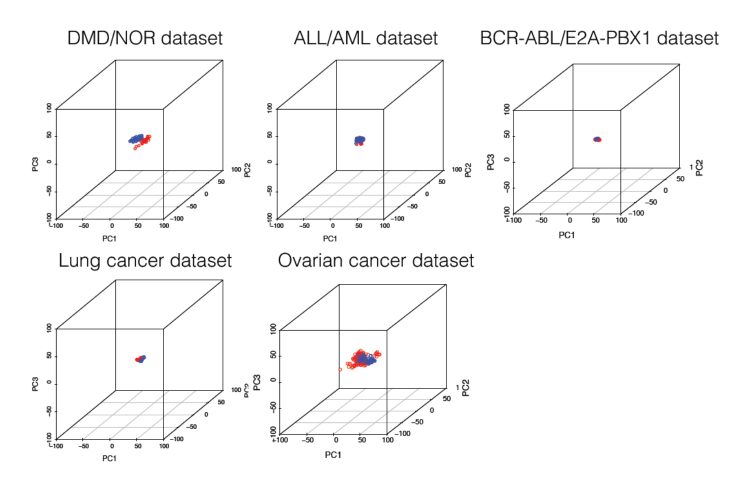


Figure 5.6: A figure showing that the batch effects are reduced by PFSNet subnetwork features. The colors red and blue represent different batches.

Successful cross-batch classification



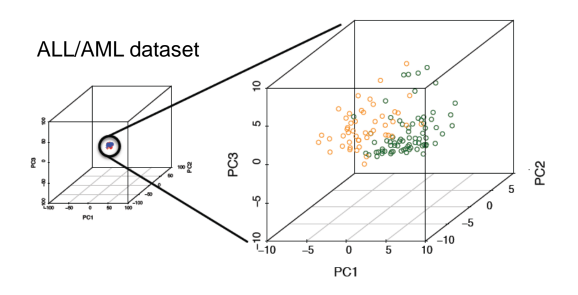


Figure 5.7: A figure showing that data points are separated by class labels instead of batch when PFSNet features are used. The colors green and orange represent different classes.



SUCCESSFUL CROSS-BATCH CLASSIFICATION

EVEN WHEN SAMPLE SIZE IS SMALL

ESSNet

- Induce subnetworks
 using genes highly
 expressed in majority of
 samples in any class
- Let g_i be genes in a given subnetwork S
- Let p_i be patients
- Let q_k be normals
- Let $\Delta_{i,j,k} = \text{Expr}(g_i,p_j) \text{Expr}(g_i,q_k)$
- Test whether $\Delta_{i,j,k}$ is a distribution with mean 0



ESSNet scores subnetworks but not patients.

How to produce feature vectors for patients?

ESSNet-based features



• The idea is to see whether the pairwise differences of genes with a subnetwork betw a given sample p_x and the two separate classes (D and $\neg D$) have a distribution around 0

$$\Delta_{(D)}(S, p_x) = \{e_{g_i, p_x} - e_{g_i, p'} \mid g_i \in S \text{ and } p' \in D\}$$

$$\Delta_{(\neg D)}(S, p_x) = \{e_{g_i, p_x} - e_{g_i, p'} \mid g_i \in S \text{ and } p' \in \neg D\}$$

- We expect $\Delta(D)(S,P_x)$ and $\Delta(\neg D)(S,P_x)$ to have +ve or –ve median for patients in one of the classes iff subnetwork S is useful for classification
 - The median and ± 2 std dev of $\Delta(D)(S,P_x)$ and $\Delta(\neg D)(S,P_x)$ give 6 features for P_x

ESSNet-based features



 We also obtain pairwise differences of genes within a subnetwork among all possible pairs of patients in D and ¬D

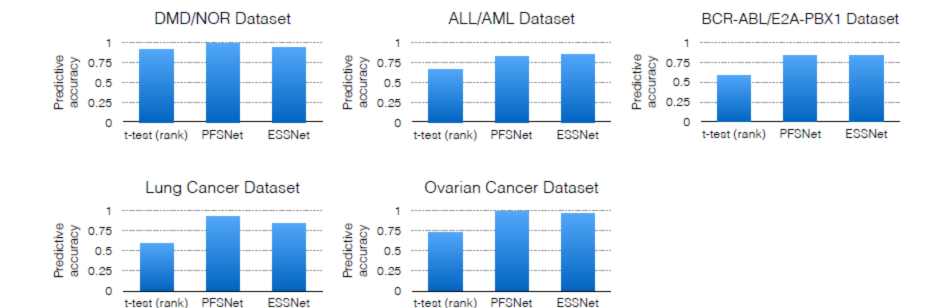
$$\Delta_{(D-\neg D)}(S) = \{e_{g_i,p'} - e_{g_i,p''} \mid g_i \in S \text{ and } p' \in D \text{ and } p'' \in \neg D\}$$
Similarly for $\Delta_{(\neg D-\neg D)}(S)$, $\Delta_{(\neg D-D)}(S)$, $\Delta_{(D-D)}(S)$

This gives 4 more features

$$\begin{split} ESSNet_feature_{7}^{p_{x},S} &= T_statistic(\Delta_{(\neg D)}(S,p_{x}),\Delta_{(D-\neg D)}(S))\\ ESSNet_feature_{8}^{p_{x},S} &= T_statistic(\Delta_{(\neg D)}(S,p_{x}),\Delta_{(\neg D-\neg D)}(S))\\ ESSNet_feature_{9}^{p_{x},S} &= T_statistic(\Delta_{(D)}(S,p_{x}),\Delta_{(D-D)}(S))\\ ESSNet_feature_{10}^{p_{x},S} &= T_statistic(\Delta_{(D)}(S,p_{x}),\Delta_{(\neg D-D)}(S)) \end{split}$$

ESSNet-based features lead to high cross-batch classification accuracy





ESSNet-based cross-batch hierachical clusterings



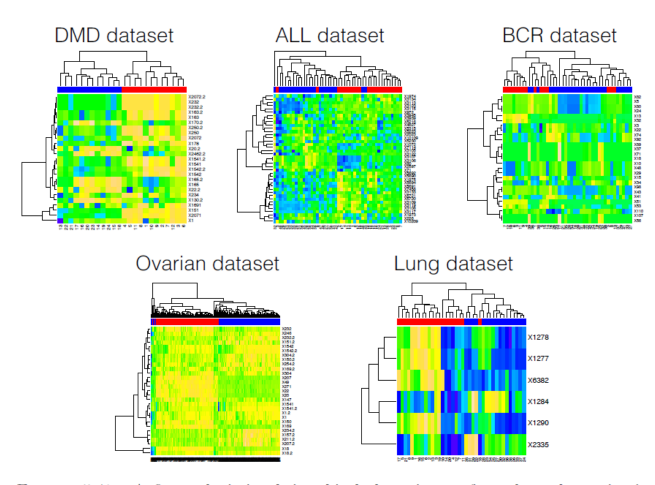
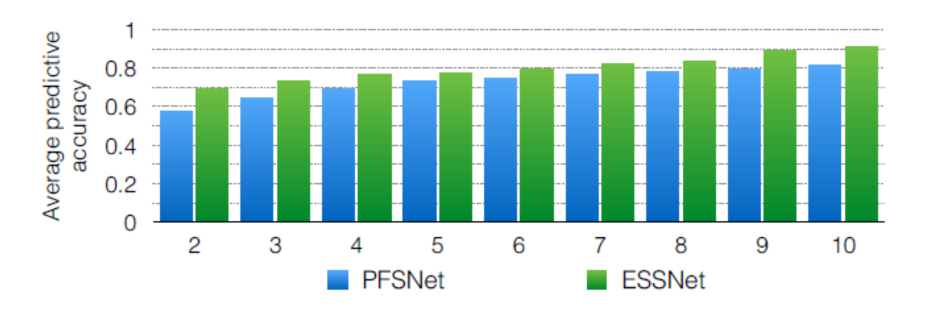


Figure 5.17: A figure depicting heirarchical clustering performed on the patient's subnetwork scores.

ESSNet-based features retain high cross-batch classification accuracy even when training-sample size is small





Remarks



- Traditional methods of classifying gene expression profiles often have difficulty predicting outcome of new batches of patients
 - Normalization does not always help
- ESSNet-based features are much more robust even when training-sample size is small
 - Subnetworks found by ESSNet are reproducible and gave high cross-batch classification accuracy
- ⇒ ESSNet is successful in isolating diseaserelevant subnetworks from pathways

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- Kevin Lim, Limsoon Wong. Finding consistent disease subnetworks using PFSNet. Bioinformatics, 30(2):189--196, January 2014
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