Estimating Drivers Cell State Transitions using Gene Regulatory Network Models

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Abstract

In the language of systems biology, the state of a cell can be represented by a gene regulatory network that characterizes the gene transcriptional processes that are active in that cell type. And transitions that occur in a wide range of biological processes, ranging from development to disease, can be thought of as transformation of the gene regulatory network from its initial state to its final state. Here we propose a regression-based generalization of the PANDA method for gene regulatory network inference for individual states, and a linear algebra approach to modeling cell state transitions, identifying transcription factors that alter the network structure as cell states change.

The problem of gene regulatory network inference

One of the fundamental problems is biology is modeling the transition between biological states such as that which occurs during development or as a healthy tissue transforms into a disease state. While it is appealing to conceive of this process as deterministic, reflecting a change from one well-defined phenotype to another, in truth the situation is much more complex. Neither the initial nor the final phenotype is discrete, but each falls into a continuum of states, which, on average, captures features of that phenotype. Indeed, within each tissue—with unique patterns of gene expression and individual regulatory

processes. The same is true when considering individuals as each healthy and disease state is unique to each member of a study population. One way to conceptualize the state transition problem is to imagine that each phenotype has a characteristic gene regulatory network and that there are a set of processes that are either activated or inactivated to transform the network in the initial state into that characterizing the final state. Identifying those changes could, in principle, help us to understand not only the processes that drive the state change, but also how one might intervene to either promote or inhibit such a transition.

As our ability to generate large-scale, integrative multi-omic datasets has grown, there has been an increased interest in using those data to infer gene regulatory networks to model fundamental biological processes. While there have been many network inference (NI) methods published, each of which uses a different approach to estimating the "strength" of interactions between genes (or between transcription factors and their targets), they all suffer from the same fundamental limitation. Every method relies on estimating weights that represent the likelihood of an interaction between two genes and then setting a threshold to identify "real" (high confidence) edges. While setting edge confidence thresholds allows us to graphically represent networks and allows us to compare networks based on the presence or absence of edges, it ultimately requires that we discard information regarding those "weak" edges that fail to reach significance. Now, one could argue that discarding low significance edges is sensible as one common goal in network inference is to deduce a single, high confidence network model that represents a particular phenotype under study or, in some cases, a transition between phenotypes. An alternative view is that these edges contain estimates for gene coexpression that may be muted due to the heterogeneity of the biological sample and that networks are best described by edgeweights on a continuous scale. Furthermore, we argue that the inclusion of all network edgeweights in downstream analyses allows for the collection of systematic, though subtle, differences in coexpression which may otherwise have been washed out by extraneous noise.

BERE: A regression-based approach to modeling gene regulatory networks

In 2013, we described PANDA [3], a method [11] for estimating gene regulatory networks that uses "message passing" a [2] to integrate multiple types of genomic data. PANDA begins with a prior regulatory network based on mapping transcription factor motifs to a reference genome and integrates other sources of data, such as protein-protein interaction and gene expression profiles, to estimate individual sample networks. While PANDA has proven to be very useful in a number of applications [4,5,8], its iterative approach to edge-weight optimization limits its utility in situations requiring a large number of network bootstrap estimations. To address this limitation, we developed BERE, Bi-

partite Edge Reconstruction from Expression. BERE approaches the network inference problem by considering the available evidence of an edge for each possible TF-gene pair. This evidence can be divided into two components, referred to here as direct and indirect. Consider the edge between a TF and a gene, referred to here as TF_i and g_j , respectively. The direct evidence, $d_{i,j}$, consists of the squared conditional correlation of the g_i and g_j given all other regulators of g_i . Where g_i is the gene which encodes TF_i

$$d_{i,j} = cor(g_i, g_j | \{g_{k,-j} : k \neq j, k \in \mathbf{TF}\})^2$$

Naturally, the use of direct evidence inadequately captures regulatory relationships due to the impacts of technical noise and numerous biological external factors such as stable or transient protein-protein interactions, post-translational modifications, etc. which may confound or modify a regulatory effect. These sources of confounding and variability in the expression pattern of a gene coding a TF may obscure the effects it has on all of its target genes. Therefore it is of value if we can complement our estimate of the likelihood of a regulatory mechanism by aggregating the information from the gene expression patterns of all suspected targets of transcription factors. PANDA achieves its superior performance in part by convergence towards "agreement", whereby large collections of gene expression patterns must agree with the proposed regulatory structure in order to claim an interaction. Similarly, BERE looks for agreement between the gene expression patterns of large sets of co-targeted genes. We refer to this feature as indirect evidence and can achieve this by again utilizing our set of regulatory priors. In this portion of the analysis we suspend the recognition of a TF as a member of the gene list and instead consider each of the m TFs to be binary classifications across the entire gene list. Class labels are determined by the presence or absence of a sequence binding motif for that TF in the vicinity of the gene.

The indirect evidence between the two nodes, $e_{i,j}$, represents the fitted probability that g_i belongs to the class of genes targeted by TF_j . g_i is considered to be a new observation placed into the n-dimensional space separated by transcription factor targets and non-targets. To divide up the space, BERE uses a regularized logistic regression on the gene expression data with the training set taken to be all genes and the training labels taken to be the existence or non-existence of a known sequence motif for TF_j upstream of g_i . The penalized model matrix comes from the recognition that correlations between co-regulated genes will be most strong when the TF_j is most prevalent. We therefore use the abundance of TF_j to weight the penalized model matrix, providing increased sensitivity for detecting coexpression for those samples in which we most expect it to occur. To build each of our classifiers we use the L2 regularization with the penalized model matrix, \mathbf{Q} , a diagonal matrix with weights equal to the the inverse expression value of the transcription factor. Effectively, we maximize the penalized logistic likelihood function

$$\sum_{i=1}^{n} log \left[exp \left(\beta' \mathbf{x_i} \right)^{Y_i} \left\{ 1 - exp \left(\beta' \mathbf{x_i} \right) \right\}^{1-Y_i} \right] - \lambda \beta' \mathbf{Q} \beta$$

This computation is run using the R package "penalized", with the penalty term lambda estimated via default 5 fold cross validation.

By scoring each gene according to the strength of indirect evidence for a regulatory response to each of the TFs, we can combine this with the direct evidence of regulation (squared conditional correlation of expression for genei and TF_i). The appropriate manner in which to combine direct and indirect evidence remains an open question. Though both measures are bounded by [0,1] their interpretation is quite different. The direct evidence can be considered in terms of it's conditional gene expression R^2 between nodes, while the indirect evidence is interpreted as a probability. We use a non-parametric approach to combine evidence. The targets of each TF are then ranked and combined as a weighted sum, $w_i = (1 - \alpha) [rank(d_i)] + \alpha [rank(e_i)], i \in \{1, ..., n\}.$ Our choice of the weight, α , here is based on empirical evaluation, and perhaps not surprisingly, is loosely correlated with organism complexity. In validation sets from Yeast, the optimal alpha was observed near $\alpha = .9$ while simpler E. coli datasets saw an optimal value of $\alpha = .6$ and an in silico dataset, optimality was achieved at $\alpha < .5$ This naturally reflects the fact that the increased complexity of the network necessitates the use of larger scale agreement between genes, rather than a reliance on pairwise correlations between potentially noisier and more complex expression patterns.

Modeling cell state transitions as a problem in gene regulatory network transition

Cell state transitions—such as those that occur during development, or as healthy tissue transforms into a disease phenotype—are fundamental properties of biological systems. Understanding what drives these transitions, and modeling the processes, is one of the great open challenges in modern biology. One way to conceptualize the state transition problem is to imagine that each phenotype has its own characteristic gene regulatory network, and that there are a set of processes that are either activated or inactivated to transform the network in the initial state into that which characterizes the final state. Identifying those changes could, in principle, help us to understand not only the processes that drive the state change, but also how one might intervene to either promote or inhibit such a transition. The starting point for modeling cell state transitions is to model the initial and final cell states. One might imagine that the initial and final cell states consist of characteristic processes, some of which are shared (sometimes referred to as "housekeeping" functions) and others which are unique to the particular state. The way we understand these processes is that they are controlled by gene regulatory networks in which transcription factors (and other regulators) moderate the transcription of individual genes whose expression characterizes the state. One way to represent such processes is to draw a directed network graph, in which transcription factors and genes are nodes network in the network, and edges represent the regulatory interactions between transcription factors and their target genes that are active in, and characteristic of, a particular cellular state. One way of representing such a network, with interactions between m transcription factors and n target genes, is as a binary $m \times n$ "adjacency matrix," with 1's representing active transcription factor-target interactions, and 0's representing the lack of a transcription factor-target gene regulatory interaction. One can then think of a cell fate transition as the process that transforms the network in its initial state to its final state form, adding and deleting edges to remake the network that characterizes one phenotype into that which characterizes the other. Using the adjacency matrix formalism, one can think of this as a problem in linear algebra in which we attempt to find an $m \times m$ "transition matrix" \mathbf{T} , subject to a set of constraints that approximates the conversion from the initial network's adjacency matrix \mathbf{A} into the final network's adjacency matrix \mathbf{B} , or



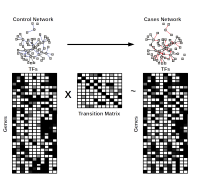


Figure 1: Overview of the Transition Matrix problem. Our approach seeks to find the $TF \times TF$ matrix which best characterizes the transition in TF targetting between cases and controls. We can think about this problem as the estimation of the set of TF-TF interactions which minimizes edgeweight error in the cases network when applied to the controls network.

While it is appealing to conceive of this process as deterministic, reflecting a change from one well-defined phenotype to another, in truth the situation is much more complex. Neither the initial nor the final phenotype is discrete, but each falls into a continuum of states, which, on average, captures the features of that phenotype. Indeed, within each tissue there are many, many cells, each of which is its own particular instance of that tissue—with unique patterns of gene expression and individual regulatory processes. In the language of adjacency matrices, what this means is that rather than representing each state by a matrix with binary entries, what one should do is use a representation in which entries are continuous, representing the strength of the transcription factor-target gene interaction averaged over the collection of samples (or cells) representing each state. And consequently, the problem of estimating the transition matrix is

generalized to solving, where E is an $m \times n$ error matrix representing the uncertainty in the estimation of the individual edges. In this formalism, modeling the cell state transition is equivalent to estimating the appropriate transition matrix T that maps how the transcription factor-target gene interactions are "rewired" between states. And one could hypothesize that the drivers of the cell state transition are those transcription factors that are have the greatest change in the targets that they regulate. In evaluating the state transitions, we recognize the limitations of current network inference methods to predict individual edgeweights. It's therefore of interest to combine measurements across sets of edgeweights in order to extract meaningful signal from a network perturbation. Effectively, we approached the problem as a dimension reduction problem with the goal of identifying high-influence systematic regulatory network alterations rather than isolated independent events. There are many existing methods for reducing a high-dimensional matrix such as a gene regulatory adjacency matrix. Commonly, Principal Components Analysis (PCA) identifies eigenvectors which can reconstruct the greatest degree of variance from the original data. One drawback of this approach is the lack of interpret-ability of these vectors. Our transition matrix approach can be considered as a data reduction method which (1) preserves the intuitive interpretation of its vectors and (2) utilizes our expectation that meaningful network transitions will occur via biologically systematic alterations and not via random, independent edge alterations.

Transition Analysis

A primary purpose of reconstructing GRNs is the understanding of the biological mechanisms which characterize disease. Identifying differential TF targeting may suggest a therapeutic target, uncover a disease substructure or identify biomarkers for early detection. However, significant limitations exist with respect to generating reliable context-specific GRNs. Substantial advances have been made in this area with algorithms like SEREND and PANDA, but these methods rely more heavily on the static sequence motif data rather than gene expression data collected across groups, such as in a case-control study. Both methods, along with BERE, use gene expression data as a refinement of the more accurate motif priors. It is therefore a side effect that any GRN inference method that relies on motifs directly for edgeweight calculation will be susceptible to having the bulk of its predictive power stripped when making comparisons between networks. Consequently, it is of critical importance to choose a network inference method that uses the context-specific data to most accurately refine above what information is gleaned from our static data.

The task of identifying meaningful network transitions then becomes an evaluation of the relative refinement of edgeweights. Since the majority of the predictive power for each edge is contained in the motif contribution, we are left with relative edgeweight refinement that have a low signal, high noise. In other words, we have a very large number of individually unreliable edgeweights. In effort to extract the maximum effect, we seek to combine the information

contained in each edge via a novel dimension reduction approach.

Consider two adjacency matrices, **A**, **B** representing the two GRNs estimated from a case-control study. Each matrix has dimensions $(p \times m)$ representing the set of p genes targeted by m TFs. We seek a matrix, **T**, such that

$$B = AT + E$$

Where **E** is our error matrix, which we want to minimize. Intuitively, we may frame this as a set of m independent regression problems, where m is the number of transcription factors and also the column rank of $\mathbf{A}, \mathbf{B}, \mathbf{T}$ and \mathbf{E} . For a column in \mathbf{B} , \mathbf{b}_i , we note that a corresponding column in \mathbf{T} , τ_i , represents the OLS solution to

$$E\left[\mathbf{b}_{i}\right] = \tau_{i1}\mathbf{a}_{1i} + \tau_{i2}\mathbf{a}_{2i} + \dots + \tau_{im}\mathbf{a}_{mi}$$

or alternatively expressed

$$\begin{bmatrix} \mathbf{b}_{i1} \\ \mathbf{b}_{i2} \\ \vdots \\ \mathbf{b}_{ip} \end{bmatrix} = \tau_{1,i} \begin{bmatrix} \mathbf{a}_{11} \\ \mathbf{a}_{21} \\ \vdots \\ \mathbf{a}_{p1} \end{bmatrix} + \tau_{2,i} \begin{bmatrix} \mathbf{a}_{12} \\ \mathbf{a}_{22} \\ \vdots \\ \mathbf{a}_{p2} \end{bmatrix} + \cdots + \tau_{p,i} \begin{bmatrix} \mathbf{a}_{1p} \\ \mathbf{a}_{2p} \\ \vdots \\ \mathbf{a}_{pp} \end{bmatrix} + \begin{bmatrix} e_{i1} \\ e_{i2} \\ \vdots \\ e_{ip} \end{bmatrix}$$

where $E[e_{ij}] = 0$

This can be solved with normal equations,

$$\tau_i = (\mathbf{A}^T \mathbf{A})^{-1} \mathbf{A}^T \mathbf{b}_i$$

$$\mathbf{T} = [\tau_1, \tau_2, \dots, \tau_n]$$

Which produces the least square estimate. I.e. loss function $L(\mathbf{T}) = \sum_{gene=1}^{N} ||\mathbf{B}_{gene} - \mathbf{A}_{gene}\mathbf{T}||^2$ is minimized. We further extend this method to include a penalty term [14]. An L_1 regularization is used by creating an identity penalty model matrix for each column regression such that only the k^{th} diagonal element is 0 and all other diagonals are 1. This gives priority for the k^{th} regression coefficient in the k^{th} regression model.

$$\mathbf{Q}_{i,j} = \begin{cases} 1 & for \ i = j \neq k \\ 0 & elsewhere \end{cases}$$

This solution is obtained using the "penalized" library in R as the minimization of the penalized squared loss function

$$\sum_{i=1}^{p} \left(\mathbf{B}_{i,k} - \sum_{j=1}^{m} A_{i,j} \mathbf{T}_{j,k} \right)^{2} + \lambda \sqrt{\beta' \mathbf{Q} \beta}$$

This example illustrates a key feature of this method. Specifically, that the transition matrix is a data reduction method that reduces the case-control network transformation from a set of $2 \times p \times m$ estimates to a set of $m \times m$

estimates that are easily interpreted. We can think of a column, τ_i , on the matrix \mathbf{T} as containing the linear combination of regulatory targets of TF_i in \mathbf{A} that best approximates the regulatory targets of TF_i in \mathbf{B} . As one would expect, a large proportion of the matrix "mass" would be on the diagonal for those TFs which do not change regulatory behavior between case and control. It is therefore of interest to evaluate values off of the diagonal as indications of a network transition.

Transition Matrix Analysis

Many mechanisms which may be differentially present, such as RNA degredation, post-translational modification, protein-level interactions and epigenetic alterations have the ability to impact downstream targeting without impacting the expression level of the TF itself. It may be of particular scientific or therapeutic interest to identify those TFs which have undergone significant overall changes in behavior between controls and cases. With that objective in mind, we express the statistic- differential Transcription Factor Involvement (DTFI), as a measure for quantifying this property.

$$s_{j} = \frac{\sum_{i=1}^{m} I(i \neq j) \tau_{i,j}^{2}}{\sum_{i=1}^{m} \tau_{i,j}^{2}}$$

DTFI can be loosely interpreted as the proportion of TF targeting patterns which is explained by the targeting patterns of other available TFs. This measure, a statistic on the interval [0,1] seeks to elucidate transitions which are systematic, informative, and non-arbitrary in nature by capturing only the edgeweight signal for which there is an attributable regulatory pattern. The distribution of this statistic under the null has a mean and standard deviation which depend on the motif structure. In particular, both mean and standard deviation are increased for TFs which have fewer prior regulatory targets. From a statistical perspective, TFs with relatively more targets are able to generate more stable targeted expression patterns, which leads to more consistent estimates in "agreement" algorithms such as PANDA and BERE. From a biological perspective, increased motif presence may indicate that the TFs are more likely to be ubiquitous housekeeping proteins that do not meaningfully alter their involvement between cases and controls. The dependence of the null distribution on the motif structure is addressed via the following resampling procedure.

- 1. Gene expression samples are randomly assigned to case and control forming the null-case and null-control with group sizes preserved.
- 2. GRNs are reconstructed for the null-case and null-control with the same prior regulatory structure.
- 3. The transition matrix algorithm is applied for the two null networks.
- 4. The differential TFI is calculated for each TF.

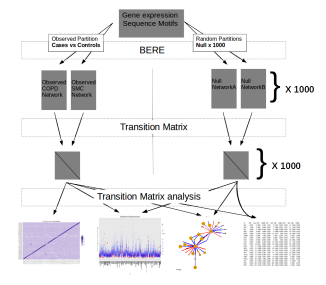


Figure 2: Overview of Transition Matrix analysis workflow. (1) BERE is applied separately to subsets of the gene expression data including the cases group, the controls group and 1000 permutations of the cases and controls labels. (2) Transition matrix is estimated between the cases and controls and each of the pair or permuted "cases" and "controls".

5. Repeat 1-4 1000 times.

TM significantly improves TF-TF edge estimation from simulated gene expression data

To evaluate the ability of our method to recover edges between transcription factors, we generated simulated gene expression data. We began by generating a true controls adjacency matrix, $M_{0(p\times q)}$, describing the weighted edges between q transcription factors and p genes. A state transition was generated by sampling 100 TF-TF pairs and adjusting the edgeweight at the corresponding point on the true cases adjacency matrix, $M_{1(p\times q)}$. These TF-TF pairs ultimately represent the edges that we seek to recover and the size of the adjustments are the parameters of interest. We sampled from a multivariate Gaussian distribution with the off-diagonal of the variance-covariance matrix, Σ , defined as the M_0M_0' . Furthermore, we scaled the magnitude of the diagonal of Σ to achieve the desired proportion of noise. We aimed for an area under the curve of the receiver-operator characteristic of approximately 0.70 as this has reasonably been achieved in existing biological studies [3]. We note that the two simulated regulatory priors have AUC-ROC of .570 and .547, which feeds the BERE

and PANDA algorithms with priors which are substantially less predictive than sequence motif priors commonly used for network inference methods.

This sampling represented our simulated control samples. The adjusted adjacency matrix, M_1 , was similarly used to generate simulated expression data for the cases group. Next, we reconstructed the networks from our expression data using a set of commonly used network inference methods - Weighted Gene Correlation Network Analysis (WGCNA) [6] [7], Topological Overlap Measure (TOM) [13], Algorithm for the Reconstruction of Gene Regulatory Networks (ARACNE) [10], Context Likelihood of Relatedness (CLR) [1], Passing Attributes between Networks for Data Assimilation (PANDA) [3] and simple Pearson correlation (PC).

We applied the transition matrix with default parameters on each case-control pair of networks. For comparison, we estimated the difference from case to control in edgeweights derived from the direct edge prediction using each network inference method. The predictions for the TM approach and the direct approach were evaluated by the area-under-the-curve of the receiver-operator-characteristic (AUCROC) with the true transition adjustments taken as the gold standard. For each of the network inference methods tested, we found substantial improvement in the predicted transitions over the direct network inference method. In many cases, the edgeweight difference (column 2) was not statistically significant for predicting transitions, but when the TM was applied (column 3) a strong predictive signal appeared. In other cases, an existing signal was observed using the direct approach, but was dramatically improved with the application of the TM.

The intuition behind the improvement is simple. While the estimation of a TF-TF edge is typically evaluated via some pairwise gene expression pattern which may be rife with technical and biological noise, the TM approach borrows information from all downstream targets in estimating the relative change in relationship between the TFs.

Transition Matrix finds concordance in independent datasets for COPD

We applied our method to four case-control datasets for Chronic Obstructive Pulmonary Disease (COPD)- Evaluation of COPD Longitudinally to Identify Predictive Surrogate Endpoints (ECLIPSE), the COPDGene study, Lung Genomics Research Consortium (LGRC) and Lung Tissue Chronic Obstructive Pulmonary Disease (LTCOPD). Each of these studies consisted of gene expression assays obtained from patients with COPD and a set of smoker controls. The tissue used in the ECLIPSE and COPDGene study was peripheral blood mononuclear cell (PBMC), while lung tissue was sampled for LGRC and LT-COPD. It is therefore reassuring that agreement is more strongly achieved between studies of the same tissue origin than acros tissues. Each of the four studies was most closely correlated with studies of the same tissue. However it

AUC-ROC for Edgeweight differences vs Transition Matrix using various NI methods

	0 0	8					
NI Method	Network AUC	Edgeweight	Transition				
		differences	Matrix				
Pearson	.704	.510 (p=.72)	.802 (p<.0001)				
WGCNA(6)	.704	.512 (p=.61)	.688 (p<.0001)				
WGCNA(12)	.704	.52 (p=.10)	.589 (p=.02)				
ARACNE	.515	.523 (p=.58)	.566 (p=.09)				
CLR	.694	.57 (p=.19)	.814 (p<.0001)				
TOM	.703	.51 (p=.62)	.689 (p<.0001)				
PANDA*	.747	.520 (p=.13)	.793 (p<.0001)				
PANDA**	.652	.509 (p=.43)	.66 (p<.0001)				
BERE*	.813						
BERE**	.697						

Table 1: Comparison of edgeweight difference to Transition Matrix in simulated case-control gene expression. Several network inference methods were run on our *in silico* case-control data. The overall network area under the curve of the receiver-operator characteristic (AUC-ROC) was performed for each method averaged across cases and controls. For PANDA* and PANDA**, which additionally utilizes motif prior information, motif priors with AUC-ROC of .570 and .547 were used. The naive TF-TF transitions were calculated as the difference in TF-TF edgeweight between cases and controls. The transition matrix TF-TF transitions used the absolute transition matrix values.

is quite notable that the we do see much of the same dTFI signal across studies involving different tissue types.

We separately applied our BERE network inference approach on cases and controls and computed the transition matrix. Top significance hits for dTFI showed strong concordance between each of the datasets. Out of 166 TF used in this study, seven were among top 10 most differentially involved in both the ECLIPSE and COPDGene studies. Furthermore, three of these seven TFs (GABPA, ELK4, ELK1) also appeared as significant in the LGRC results with FDR<.01.

Overall, there was a strong correlation between results in the two PBMC studies ($r_s = 0.87$) and the two lung tissue studies ($r_s = 0.71$), and a weaker correlation between cross-tissue results ($\bar{r}_s = 0.63$).

Of further interest is the particular TFs which are differentially observed across studies. For example, ERS1, the gene encoding Estrogen receptor alpha (ES- α) is likely to behave differently in males compared with females. We found that this TF is significantly differentially involved in the COPDGene study - identified as the 4th most differentially involved TF, but not the ECLIPSE study. Accordingly, the COPDGene study had a greater gender disparity (Odds Ratio = 1.126) than the ECLIPSE study (Odds Ratio = 1.067).

Notably in the LGRC dataset, we discovered a differential targetting pattern involving the TFs RFX1 and RFX2. Both of these transcription factors were

	ECLIPSE			COPDGene			LGRC					
TF	dTFI	rank	FDR	LIMMA	dTFI	rank	FDR	LIMMA	dTFI	rank	FDR	LIMMA
SP2	.0371	1	<.0001	.1756	.0120	9	.9717	.6517	.0164	11	.7642	.0075
E2F4	.0360	2	<.0001	.3913	.0132	7	.0003	.9367	.0110	41	.039	.0878
SP1	.0286	3	.0004	.3634	.0136	5	.6088	.0838	.0113	38	.9236	.4242
NRF1	.0282	4	<.0001	.0966	.0133	6	.0068	.0045	.0104	43	.0379	.2974
EGR1	.0265	5	<.0001	.4379	.0094	21	.9717	.8540	.0112	39	.9236	.7979
GABPA	.0253	6	<.0001	.4650	.0189	1	<.0001	.5138	.0210	7	<.0001	.3868
ELK4	.0245	7	<.0001	.0001	.0159	3	<.0001	.8057	.0178	9	<.0001	.0183
ELK1	.0233	8	<.0001	.0913	.0185	2	<.0001	.9010	.0122	32	.0011	.7968
KLF4	.0227	9	<.0001	.1915	.0119	10	.4573	.0025	.0129	22	.437	.0526
E2F1	.0207	10	<.0001	.6929	.0089	23	.4024	.6465	.0079	66	.3789	.0022
ESR1	.0140	17	.0002	.9598	.0142	4	.0049	.5853	.0125	26	.0033	.7246
NFIL3	.0115	29	.8698	.0404	.0061	49	.4404	.1191	.0239	4	.0418	.7605
FOXQ1	.0114	31	.9643	.4543	.0106	14	.9752	.5314	.0196	8	.1031	.0503
IRF2	.0108	37	.4149	.1914	.0090	22	.531	.8237	.0168	10	.0083	.2469
PLAG1	.0103	44	.4706	.0384	.0127	8	.4024	.0008	.0154	13	.0541	.0371
JUNB	.0065	75	.8736	.0197	.0047	79	.5281	.9526	.0262	3	<.0001	.3996
FOSL1	.0062	80	.8595	.5850	.0055	56	.6088	.6995	.0238	5	<.0001	.8708
FOS	.0054	90	.7639	.5156	.0043	86	.4576	.6668	.0211	6	.0024	.9500
RFX1	.0014	162	.5736	.0361	.0011	164	.4404	.0885	.0401	1	<.0001	.0175
RFX2	.0014	161	.5736	.0109	.0015	157	.4404	.0059	.0333	2	<.0001	.0004

Table 2: Combined list of TFs which were among the top 10 hits (out of 166 available TFs) in any of the 3 studies, ordered by the dTFI in the ECLIPSE study. For each study, columns indicate the TF's (1) differential TF Involvement, (2) dTFI Rank within list of TFs, (3) Significance of dTFI by false discovery rate, and (4) p-value for LIMMA differential gene expression analysis.

highly statistically significant (FDR<.0001) and ranked as the top two results in the LGRC study. However, their signal was muted in the ECLIPSE and COPDGene studies, neither of which identified these transcription factors as drivers of the Smoker Control to COPD transition. There are many possible explanations for this result, but it is reasonable to speculate that the dramatically different result is due to the fact that the tissue of origin for the LGRC differs from the ECLIPSE and COPDGene tissue.

The top hits which are most consistent across studies have been implicated in independent studies for the development of COPD. Two of the top 3 hits, NRF1 and GABPA have been implicated in a mitochondrial mechanism for disease progression [Cloonan, Suzanne M et al]. Interestingly, the majority of the TFs identified as differentially involved do not exhibit significant differential gene expression. This suggests that for these proteins, their role in the disease may not occur until the post-transcription stage. It also suggests that conventional gene expression analysis is insufficient for identifying many of the TF drivers of disease.

Supplemental Methods

The computations in this paper were run on the Odyssey cluster supported by the FAS Division of Science, Research Computing Group at Harvard University.

Transition matrix finds significant protein-protein interaction

As noted above, there are numerous biological regulatory mechanisms which may yield detectable transitions. Of particular interest are those which are less readily detectable via conventional methods, such as differential gene expression analysis. One mechanism studied here involves one TF binding to another TF to promote, suppress or alter one or both of their regulatory patterns. These multiprotein interactions create combinatorial complexity that can explain much of the variation in organism complexity which is unexplained by number of genes alone [9].

To test the ability to detect protein level interactions, we compared our estimated transitions to a set of known protein-protein interactions [12]. This set contained 223 pairwise interactions between a total of 189 transcription factors. Of interest was the effectiveness of identifying these interactions via the transition from one phenotypic state to another. This is a challenging task for several reasons, (1) protein-protein interaction is merely one of a myraid of detectable transition mechanisms, (2) it is reasonable to assume that only a small subset of the known PPI are actually differentially present between case and control and (3) technological limitations in the active field of proteomics cannot be expected to identify all interactions with a reasonable degree of certainty.

For the transition from Smoker control to COPD, we compared for the transition estimates to those generated by the transition from randomly resampling the phenotypic labels. The AUCROC for the prediction of the 223 known protein-protein interactions was $0.5695\,(p=0.018)$, suggesting that our approach is successful at detecting this highly obscured signal.

Impact of homogeneity of cases and controls on statistical inference

Many network inference methods use a measurement of pairwise co-expression as a sufficient statistic for building gene networks based on gene expression data. The ability to detect coexpression in a sample is a function of both the level of extraneous noise and the biological variability within the sample. In our analysis, we compared the transition from the observed controls to observed cases and compared it to the null distribution estimated from a randomly sampled partition of mixed phenotypes. It is therefore of interest to consider the impact of generating gene regulatory networks from samples with heterogeneous versus homogeneous phenotypes. We consider whether transitions between any two homogeneous networks yields increased variance of test statistics and inflation of type I error when using heterogeneous populations for estimating the null distribution of these test statistics.

To explore, we sampled 100 cases and 100 controls. Within both cases and controls, we split each into two groups, denoted Cases A, Cases B, Con-

trols_A, Controls_B, each of size 50. We performed BERE network inference on each group and ran the transition matrix for each pairwise network transition. The null networks were generated via a heterogeneous sampling without replacement of 50 samples for the null cases and null controls.

Of interest is the relative distribution of test statistics in transitions across-phenotypes compared to within-phenotype, relative to the null (heterogeneous) transitions.

We find that the distribution of dTFI for homogeneous within-phenotype transitions (Cases_A \rightarrow Cases_B, Controls_A \rightarrow Controls_B) closely matched the distribution of dTFI under the heterogeneous null. Comparatively, across-phenotype transitions showed strongly significant results compared to the heterogeneous null. We view these results as strong evidence that our findings in this manuscript are not inflated due to the comparison of homogeneous observed networks relative to heterogeneous null networks.

Efficiency of estimation

Let \mathbf{x}_p be a Gaussian p-vector representing a sample of gene expression data containing q transcription factors and p-q non-transcription factor genes.

$$\mathbf{x}_p \sim N\left(\mu, \Sigma\right)$$

where μ is the *p*-vector of mean gene expression values and Σ is the $p \times p$ variance-covariance matrix. In this scenario, Σ may be regarded as a combination of two independent variance-covariance sources- (1) biological signal, (2) biological noise and technical noise.

In investigating gene regulation, many network inference methods are constructed for the estimation of the $p \times q$ subset of Σ pertaining to the effect of the q TFs on the p genes. In identifying drivers of state transitions, we seek to focus on the $q \times q$ matrix of TF-TF effects. We show that our method vastly ourperforms commonly used network inference methods in estimating these specific effects.

Consider a state change between two experimental conditions, A and B, characterized by an alteration of size δ to the biological signal component of the TF-TF variance-covariance matrix at point $\Sigma_{i,j}$ where i and j are indices for two TFs in Σ .

Using a univariate coexpression calculation (the basis for Pearson networks and WGCNA estimates), the estimated variance of our estimate of δ can be calculated:

$$-\rho_A < \delta < \rho_A, \delta + \rho_A \le 1$$

$$Var(\hat{\rho}_{i,j,A} - \hat{\rho}_{i,j,B}) = Var(\hat{\delta}_{cor})$$

$$Var(\hat{\rho}_{i,j,A}) + Var(\hat{\rho}_{i,j,B})$$

$$= \frac{1 - \rho_{i,j,A}^2}{n_A - 2} + \frac{1 - \rho_{i,j,B}^2}{n_B - 2}$$

$$= \frac{1}{n_A - 2} + \frac{1}{n_B - 2} - \frac{\rho_{i,j,A}^2}{n_A - 2} - \frac{\rho_{i,j,B}^2}{n_B - 2}$$

Meanwhile, in condition B the new correlation of TF_i with some gene, $gene_k$ $k \in 1, 2 \dots p$, denoted cor^* , becomes

$$cor^*(TF_i, gene_k) = cor(TF_i, gene_k) + \delta cor(TF_j, gene_k)$$

where the term $\delta cor(TF_j, gene_k)$ is the change due to the interaction of TF_i and TF_j .

The variance of our estimate using the transition matrix can be expressed as follows:

$$Var\left(TM_{i,j}\right) = Var\left(\hat{\delta}_{TM}\right)$$

$$= \frac{\left(\frac{1}{p}\right)\sum_{k=1}^{p} vVar\left(\hat{\rho}_{i,k,A} - \hat{\rho}_{i,k,B}\right)}{\sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{j}}\right)^{2}}$$

$$= \frac{\left(\frac{1}{p}\right)\sum_{k=1}^{p} \left[Var\left(\hat{\rho}_{i,k,A}\right) + Var\left(\hat{\rho}_{i,k,B}\right)\right]}{\sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{i}}\right)^{2}}$$

$$\leq \frac{\left(\frac{1}{p}\right)\sum_{k=1}^{p} \left[\frac{1}{n_{A}-2} + \frac{1}{n_{B}-2}\right]}{\sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{i}}\right)^{2}}$$

$$\leq \frac{\frac{1}{n_{A}-2} + \frac{1}{n_{B}-2}}{\sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{i}}\right)^{2}}$$

$$\leq \frac{Var\left(\hat{\delta}_{cor}\right) + \frac{\rho_{i,j,A}^{2}}{n_{A}-2} + \frac{\rho_{i,j,B}^{2}}{n_{B}-2}}{\sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{i}}\right)^{2}}$$

$$\leq Var\left(\hat{\delta}_{cor}\right) + \frac{Var\left(\hat{\delta}_{cor}\right)\left(1 - \sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{i}}\right)^{2} + \frac{\rho_{i,j,A}^{2}}{n_{A}-2} + \frac{\rho_{i,j,B}^{2}}{n_{B}-2}}{\sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho_{i}}\right)^{2}}$$

So we have that $Var\left(TM_{i,j}\right) < Var\left(\hat{\delta}_{cor}\right)$ when

$$Var\left(\hat{\delta}_{cor}\right)\left(1 - \sum_{k=1}^{p} \left(\rho_{j,k} - \bar{\rho}_{i}\right)^{2}\right) < \frac{\rho_{i,j,A}^{2}}{n_{A} - 2} + \frac{\rho_{i,j,B}^{2}}{n_{B} - 2}$$

Since each term except $\left(1 - \sum_{k=1}^{p} (\rho_{j,k} - \bar{\rho}_i)^2\right)$ is strictly non-negative, we see that this inequality holds when

$$\sum_{k=1}^{p} (\rho_{j,k} - \bar{\rho_i})^2 < 1$$

Thus, we have a more efficient estimator of δ when

$$p > \frac{1}{Var\left(\rho_{j,k}\right)}$$

In practice, we typically have a large number of genes, p, so that our transition matrix estimator will be expected to be dramatically more efficient than the commonly used Pearson or WGCNA estimators.

References

- [1] Jeremiah J Faith, Boris Hayete, Joshua T Thaden, Ilaria Mogno, Jamey Wierzbowski, Guillaume Cottarel, Simon Kasif, James J Collins, and Timothy S Gardner. Large-scale mapping and validation of escherichia coli transcriptional regulation from a compendium of expression profiles. *PLoS Biol*, 5(1):e8, 2007.
- [2] Brendan J Frey and Delbert Dueck. Clustering by passing messages between data points. *science*, 315(5814):972–976, 2007.
- [3] Kimberly Glass, Curtis Huttenhower, John Quackenbush, and Guo-Cheng Yuan. Passing messages between biological networks to refine predicted interactions. *PloS one*, 8(5):e64832, 2013.
- [4] Kimberly Glass, John Quackenbush, Edwin K Silverman, Bartolome Celli, Stephen I Rennard, Guo-Cheng Yuan, and Dawn L DeMeo. Sexuallydimorphic targeting of functionally-related genes in copd. BMC systems biology, 8(1):118, 2014.
- [5] Kimberly Glass, John Quackenbush, Dimitrios Spentzos, Benjamin Haibe-Kains, and Guo-Cheng Yuan. A network model for angiogenesis in ovarian cancer. *BMC bioinformatics*, 16(1):115, 2015.
- [6] Peter Langfelder and Steve Horvath. Wgcna: an r package for weighted correlation network analysis. *BMC Bioinformatics*, (1):559, 2008.
- [7] Peter Langfelder and Steve Horvath. Fast R functions for robust correlations and hierarchical clustering. Journal of Statistical Software, 46(11):1–17, 2012.
- [8] Taotao Lao, Kimberly Glass, Weiliang Qiu, Francesca Polverino, Kusha-gra Gupta, Jarrett Morrow, John Dominic Mancini, Linh Vuong, Mark A Perrella, Craig P Hersh, et al. Genome medicine. 2015.

- [9] Michael Levine and Robert Tjian. Transcription regulation and animal diversity. *Nature*, 424(6945):147–151, 2003.
- [10] Adam A Margolin, Ilya Nemenman, Katia Basso, Chris Wiggins, Gustavo Stolovitzky, Riccardo D Favera, and Andrea Califano. Aracne: an algorithm for the reconstruction of gene regulatory networks in a mammalian cellular context. *BMC bioinformatics*, 7(Suppl 1):S7, 2006.
- [11] Catharina Olsen, Kathleen Fleming, Niall Prendergast, Renee Rubio, Frank Emmert-Streib, Gianluca Bontempi, Benjamin Haibe-Kains, and John Quackenbush. Inference and validation of predictive gene networks from biomedical literature and gene expression data. *Genomics*, 103(5):329–336, 2014.
- [12] Timothy Ravasi, Harukazu Suzuki, Carlo Vittorio Cannistraci, Shintaro Katayama, Vladimir B Bajic, Kai Tan, Altuna Akalin, Sebastian Schmeier, Mutsumi Kanamori-Katayama, Nicolas Bertin, et al. An atlas of combinatorial transcriptional regulation in mouse and man. Cell, 140(5):744-752, 2010.
- [13] Erzsébet Ravasz, Anna Lisa Somera, Dale A Mongru, Zoltán N Oltvai, and A-L Barabási. Hierarchical organization of modularity in metabolic networks. *science*, 297(5586):1551–1555, 2002.
- [14] Robert Tibshirani. Regression shrinkage and selection via the lasso. *Journal of the Royal Statistical Society. Series B (Methodological)*, pages 267–288, 1996.

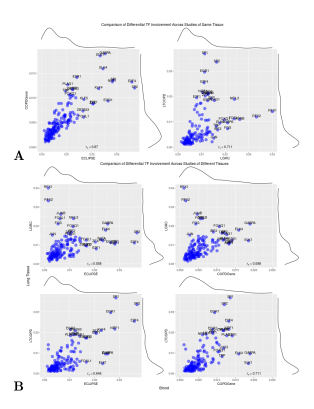


Figure 3: Comparison of dTFI for 166 TFs across four independent studies in two tissues. ECLIPSE and COPGene data were obtained via PBMC and LGRC and LTCOPD were obtained via lung tissue. Results for two studies with gene expression data obtained from the same source (A), PBMC (left) and lung tissue (right) show strong concordance with a high degree of agreement, particularly for top hits. Spearman correlation for each of these tissue types was $r_s = .87$ and $r_s = .71$, respectively. Correlations for across tissue study comparisons demonstrated a weaker, but still measureable agreement. Each of the four studies was most consistent with the study of the same tissue type as its own.

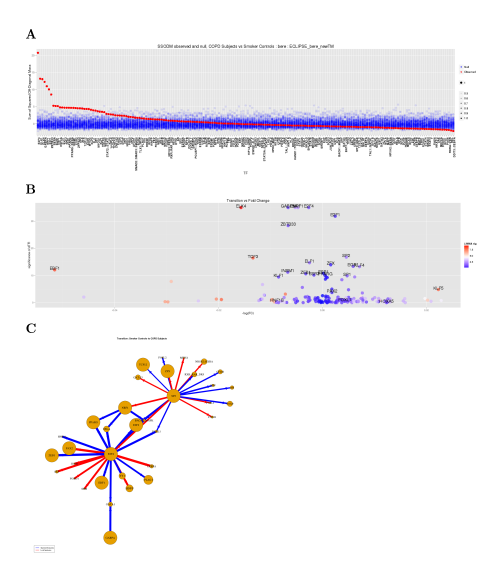


Figure 4: (A) Differential transcription factor involvement for the observed case-control (red) scaled for each TF by the distribution under the randomized case-control. (B) Comparison of TF differential involvement vs differential expression. TFs which are differentially involved are not necessarily differentially expressed and vice versa. Many TFs can be observed which have significantly different targetting patterns, but which are not statistically significantly differentially expressed. This suggests that our method finds transcription factors which are differentially affected at a post-transcription stage. (C) Driver network transitions. Network transitions are depicted here with arrows indicating the flow of targetting patterns from one transcription factor to another. Edges are sized according to the magnitude of the transition and nodes (TFs) are sized by the overall dTFI for each TF. The gain of targetting features is indicated by the color blue while the loss of features is indicated by red.

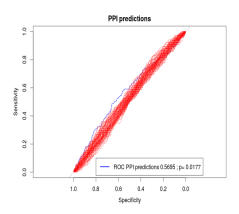


Figure 5: ROC curve for prediction of PPI based on a transition from Smoker control to COPD (blue) compared to a random case-control partition of the ECLIPSE data.