Novel formulations and methods for emerging problems in single cell data analysis

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- The wealth of data from various biological screens offers tremendous opportunities and challenges for Science of Information.
- Recent advances in single cell technologies enable us to probe dynamic states of individual cells.
- Single cell technologies are also redefining basic understanding of cell types, tissue organization, pathology, and response.
- Single cell technologies result in datasets, models, and information that are orders of magnitude more than conventional genomic/ transcriptomic/ interactomic technologies.
- Powerful new computational/ statistical/ modeling toolkits are necessary for single cell datasets.

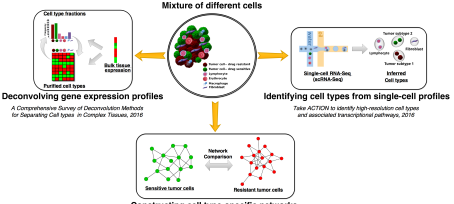


- DNA is the basic code that governs living systems.
- DNA is transcribed into RNA. This process of transcription is controlled by a number of transcriptional control mechanisms (Transcription Regulation, Post Transcriptional Regulation).
- RNA is translated into proteins the workhorses in living cells. The process of translation is controlled by a number of control mechanisms (Translational Controls).
- The activity of proteins is controlled by various post translational modifications (phosphorylation, methylation).



- Each cell in an organism (with some noted exceptions) inherits the same genetic code (its genome).
- Different cells exhibit different behavior (and function) as a result of different activity levels of genes.
- Cells exhibiting the same profile of genetic activity are generally believed to be of the same type.
- Within the set of genes, some genes are generally active across all cell types (housekeeping genes), other are selective to sets of cell types (tissue selective), others are specific to cell types (tissue specific).
- Genes whose activity is unique to cell types are called markers.
- The activity of genes in a cell is impacted by its state, stressors (external stimuli), disease, etc.
- The most common tool to interrogate the state of a cell is to study gene expression using microarrays or RNA Sequencing (RNASeq).
- Among the most common single cell technologies is single cell RNA Seq (scRNASeq).

Overview of Presentation



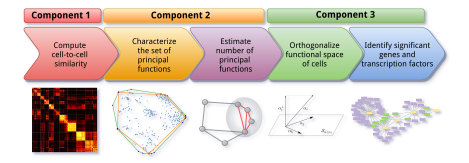
Constructing cell type-specific networks

A convex optimization approach for identification of human tissue-specific interactomes, 2016



Establishing functional identity of cells





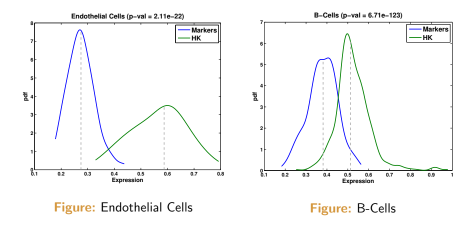


Underlying hypothesis

Transcriptional profile of cells is dominated by housekeeping genes, whereas their functional identity is determined by a combination of weak but preferentially expressed genes.

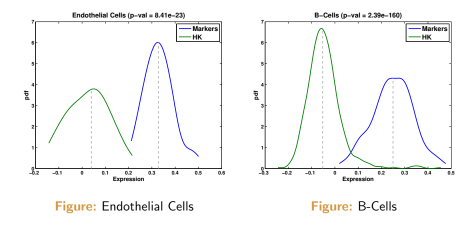


Component 1: New measures for cell-cell similarity Supporting evidence



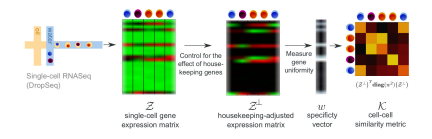


Component 1: New measures for cell-cell similarity Supporting evidence





Component 1: New measures for cell-cell similarity Cell similarity kernel in ACTION



The main steps involved in identifying similarity between cells



Component 1: New measures for cell-cell similarity Reducing the noise contributed by highly expressed but uninformative genes

Goal: Identify the shared subspace of genes

Low-rank decomposition

$$A = U_r \Sigma_r V_r = \sum_{i=1}^r \sigma_i u_i v_i^{T},$$

Example decomposition choices:

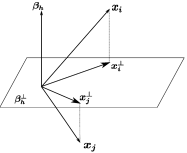
- Mean vector
 - Optimal in a least-square sense when the chance of observing a gene is uniform across all cells.
- Singular Value Decomposition (SVD)
- Nonnegative Matrix Underapproximation (NMU)
- Sparse NMU

Goal: Remove the effect of common subspace

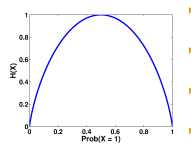
- x_i and x_j : tissues/cell types i and j
- z-score normalize x_i to compute z_i
- β_h : the common signature
- z-score normalize β_h to compute z_h
- Project to the orthogonal subspace:

$$\mathbf{z}_i^{\perp} = \left(\mathbf{I} - \frac{\mathbf{z}_h \mathbf{z}_h^T}{\|\mathbf{z}_h\|_2^2}\right) \mathbf{z}_i.$$

Similar in nature to the partial Pearson's correlation



Goal: Estimate expression-specificity of genes across different cells



- ► Entropy as a measure of expression uniformity: H(i) = -∑_i p_{ij}log(p_{ij})
- How informative is observing a gene with respect to the cell type that it came from
- Maximum entropy when probability of a gene coming from all cell types is equal
- For each gene *i*, compute a specificity factor *w_i*.

Similar formulations have been previously used for marker detection.

ACTION-adjusted cell signatures

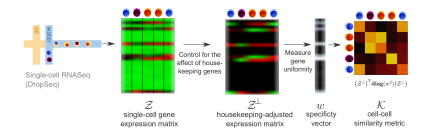
$$\mathbf{Y} = oldsymbol{diag}(oldsymbol{w}) \mathcal{Z}^ot$$

ACTION metric (kernel)

$$\mathbf{K}_{ACTION} = \mathbf{Y}^{T}\mathbf{Y} \\ = (\mathcal{Z}^{\perp})^{T} diag(\mathbf{w}^{2})(\mathcal{Z}^{\perp})$$



Component 1 Cell similarity kernel – revisited



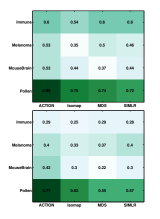
Now we have computed the ACTION kernel



- Immune: 1,522 immune cells from mouse hematopoietic system (30 different types of stem, progenitor, and fully differentiated cells)
- Melanoma: 4,645 malignant, immune, and stromal cells isolated from 19 freshly procured human melanoma tumors (7 major types, including T, B, NK, CAF, Endo, Macro, and Tumor)
- MouseBrain: 3005 cells from the mouse cortex and hippocampus (7 major types, including astrocytes-ependymal, endothelial-mural, interneurons, microglia, oligodendrocytes, pyramidal CA1, and pyramidal SS).
- Pollen: Small set of 301 cells spanning 11 different cell types in developing cerebral cortex



Performance of ACTION Kernel



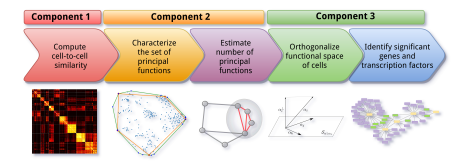
- Benchmarks:
 - SIMLR: Specifically designed for single-cell data
 - IsoMap,MDS: General purpose dimension reduction
- Tested a range of parameters (5:5:50). Reported best case for each method.

Ties:

- Immune (NMI: ACTION/MDS/SMLR, ARI: ACTON/MDS)
- Melanoma (ARI: ACTION/SIML)
- In all other cases, ACTION metric significantly outperforms all other methods.

Overall, ACTION metric performs better than other methods







General framework

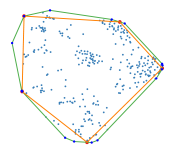
Various algorithms can be cast using this formulation

- K-means: $\mathbf{C} \in \mathbb{R}^+, \mathbf{H} \in \{0, 1\}$
- ▶ K-medoids: $C \in \{0, 1\}, H \in \{0, 1\}$

Convex NMF		1
argmin K,H	$\parallel \mathbf{Y} - \mathbf{Y}(:, \mathcal{S})\mathbf{H} \parallel$	
subject to:	$\parallel \mathbf{H}(:,i) \parallel_1 = 1, \mathbf{H} \in \mathbb{R}^+.$	

- ▶ It uses the same formulation as k-medoid, but relaxes the hard assignment of cells: $C \in \{0,1\}, H \in \mathbb{R}^n$
- Unlike k-medoid and k-means, it has an optimal global solution.
 - Under near-separability assumption: there exists, for each cell type, an ideal example in the population.
- A modification of the *Gram Schmidt* process.

Component 2: Characterizing principal functional profiles Convex NMF– Geometric interpretation



Geometry of functional space: each point is a cell and red points are the "pure cells"

- Picking k corner points/archetypes from the convex hull of the cells, such that they optimally "contain" the rest of cells.
- Each archetype is an ideal example of a cell type with a distinct set of principal functions.



- ▶ AA further relaxes matrix C: $C, H \in \mathbb{R}^+$.
- It can handle cases where pure pixel assumption is violated.
- \blacktriangleright But it no longer has global convergence guarantee \rightarrow it is also dependent on the initialization
 - To address this, we use the solution of convex NMF for initializing AA.
- ▶ In essence, this allows local adjustment of the Convex NMF solution.
- This can be thought of as a variant of block-coordinate descent for optimization.

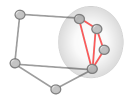


Goal: To identify when we should stop adding new archetypes.

- Underlying concept: add archetypes until we sense "oversampling."
- Oversampling happens when we start adding archetypes that are "too close" to each other.
- \blacktriangleright Each archetype is a cell \rightarrow we can compute their similarity of using the ACTION metric.



Component 2: Characterizing principal functional profiles Statistical significance of oversampling

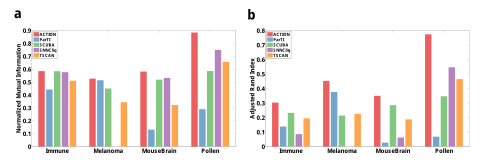


- We build and threshold an archetype-archetype similarity graph.
- For each connected component in this graph, we assess its statistical significance using an Erdos Renyi graph model.
- Probability that there exists in G a subgraph of density δ(Z) and size at least |Z|:

$$\Pr[\exists H \subseteq \boldsymbol{G}, |H| \ge |Z| : \delta(H) = \delta(Z)].$$



Component 2: Characterizing principal functional profiles Test 1: Identifying cell types using closest archetype



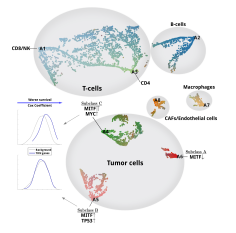
ACTION excels at identifying underlying cell types in all cases

PURDUE

- ► Use matrix **H** instead of **Y** in visualization:
 - We are interested in the relationship between cells and their surrounding archetypes.
- Initialize using Fiedler embedding
 - Position according to the dominant eigenvectors of the Laplacian matrix: L = diag(Δ_Y) – Y.
- Update using t-SNE



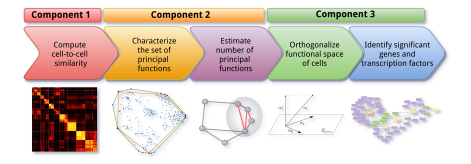
A continuous view of transcriptional profiles Case study in the Melanoma dataset



- T-cells reside in a continuum of states (Thogerson *et al.*).
- Tumor cells form compact groups.
- Two subclasses of MITF-associated tumors significantly differ in terms of their survival.

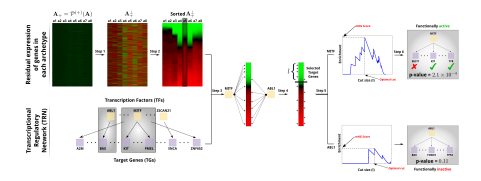
ACTION highlights the underlying topology of cell types







Component 3: Identifying the interactions underlying architypes Constructing TRN





Component 3: Identifying the interactions underlying architypes Constructing TRN

Goal: Identifying key regulatory elements that drive each cell type

1. Archetype Orthogonalization (\rightarrow Only over positive projection)

$$\boldsymbol{a}_{i}^{\perp} = \left(\boldsymbol{\mathsf{I}} - \boldsymbol{\mathsf{A}}_{-i} (\boldsymbol{\mathsf{A}}_{-i}^{\mathsf{T}} \boldsymbol{\mathsf{A}}_{-i})^{-1} \boldsymbol{\mathsf{A}}_{-i}^{\mathsf{T}} \right) \boldsymbol{a}_{i}$$

 $\mbox{2. Assessing significance of TFs/TGs}$

$$p\text{-value}(Z = b_l(\lambda)) = \operatorname{Prob}(b_l(\lambda) \leq Z)$$
$$= \sum_{x=b_l(\lambda)}^{\min(T,l)} \frac{\binom{T}{l}\binom{m-T}{l-x}}{\binom{m}{l}}$$

Use Dynamic Programming to compute exact *p*-value.



Grama et al.

CSol Annual Review, 2017

Key point!

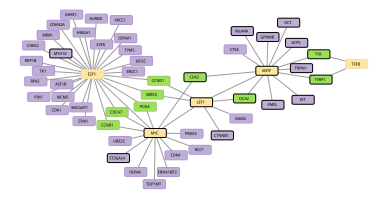
We identify "functional activity" of transcription factors (TFs) by aggregating transcriptional activity of their downstream targets, not the transcriptional level of TFs themselves. TFs can, and typically do, get regulated through post-translational mechanisms.



- ▶ Both *Subtype A* and *Subtype C* exhibit high activity of *MITF* and *Sox10* transcription factors, which are canonical markers for melanoma cells in the "proliferative" (as opposed to "invasive") state (Verfaiilie *et al.*).
- These two subtypes are significantly enriched for marker genes in the proliferative state:
 - ▶ *Subtype A:* 9.3 × 10⁻¹⁴
 - ► Subtype B: 7.9 × 10⁻¹¹
- Subtype A has higher MITF activity (according to its activated targets):
 - ► GPNMB, M1ANA, PMEL, and TYR are shared between two subtypes.
 - ► ACP5, CDK2, CTSK, DCT, KIT, and TRPM1/P1 are uniquely upregulated in subtype A.



Dissecting transcriptional controls of Melanoma subclasses Case study in MITF $\uparrow\uparrow/MYC\uparrow$ subtype



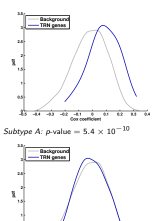
- ▶ 19 "functionally" active transcription factors in subtype A (*p*-value ≤ 0.05)
- We focus on the five most significant TFs and their targets (*p*-value $\leq 10^{-3}$)

TIRD

- MITF is among the best-known markers for classifying melanoma patients (Hartman *et al.*: MITF in melanoma: mechanisms behind its expression and activity).
- Overexpression of the E2F1 is common in high-grade tumors that are associated with poor survival in melanoma patients (Alla *et al.*: E2F1 in melanoma progression and metastasis).
- Melanoma cell phenotype switching, between proliferative an invasive states, is regulated by differential expression of LEF1/TCF4 (Eichhoff *et al.*:Differential LEF1 and TCF4 expression is involved in melanoma cell phenotype switching).
- Amplification and overexpression of the c-myc have been associated with poor outcome (Kraehn *et al.*: Extra c-myc oncogene copies in high risk cutaneous malignant melanoma and melanoma metastases).



Inferring transcriptional controls of Melanoma subtypes Survival analysis



OncoLnc (Jordan Anaya)

- Multivariate Cox regressions
- Gene expression, sex, age, and grade or histology as factors
- Genes associated with Subclass A have significantly worse outcome, compare to the background of all genes



Subtype C: p-value = 0.31

-0.4

-0.3 -0.2 -0.1

0.2 0.3

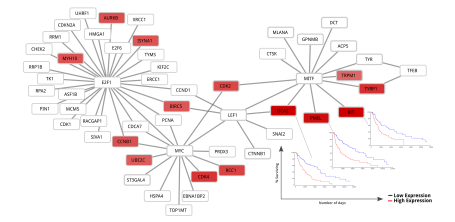
0.4

0.1

Cox coefficient

0.5

Case study in MITF // MYC hubbype Survival analysis revisited – Kaplan-Meier plots





- 1. A novel cell similarity metric that is robust to biological noise, while at the same time is sensitive enough to identify weak cell type-specific signals
- 2. New notion of functional identity of cells
 - Under the pure cell assumption, this metric induces a convex topology that embeds functional identity of cells
- **3.** Use functional identity of cells to identify both discrete cell types and continuous cell states
- 4. Identify driving transcriptional controls that mediate the functional identity of cells

Clinical significance: Characterization of two MITF-associated subclasses of Melanoma patients, one of which has substantially worse outcomes, along with their underlying regulatory elements.



- Use ACTION to infer cell types.
- Use inferred cell types to distinguish true zeros from missing values
 - There is a significant biological signal embedded merely within the sparsity pattern of the single-cell profiles.
- Use SVR to impute missing values.



- Identify stable attractor states within the continuous functional space of cells.
- Trace the most likely transition paths between the states.
- Identify regulatory factors that stimulate these transitions/fate decisions



- Use ACTION to identify cell types in human brain, construct cell type-specific region-region gene correlation networks, and compare them with the networks constructed from the resting state fMRI
- Impact of exposing RAW 264.7 macrophage cell line to exosomes from: (i) non-metastatic PEDF expressing A375 cells, and (ii) metastatic A375 melanoma cells (Joint project with Anindita Basu, University of Chicago).

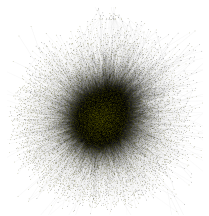


Constructing tissue-specific interactome



Motivation Global interactome is not context-specific

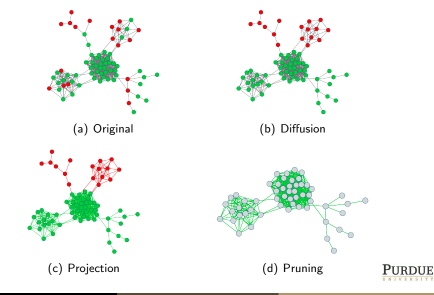
Global human interactome is a superset of all possible physical interactions that can take place in the cell. It does not provide any information as to which one of these interactions do take place in a given tissue/cell-type context.



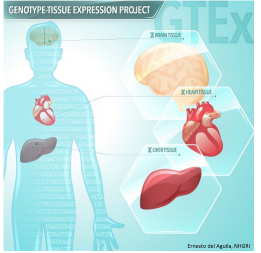
Can we predict which links/edge are active in a given context?



Roadmap Exemplar Networks



Genotype-Tissue Expression (GTEx) Project



- RNA-Seq dataset v4.0
- 2,916 samples
- 30 different tissues
- Processed each sample individually using UPC/SCAN



Adopted from: NIH CommonFund

Goal: Estimate functional activity of genes

Convex program

$$\begin{aligned} \mathbf{x}^* &= \underset{\mathbf{x}}{\operatorname{argmin}} \left\{ (1 - \alpha) \mathbf{x}^T \mathbf{L} \mathbf{x} + \alpha \parallel \mathbf{x} - \mathbf{z} \parallel_1 \right\} \\ \text{Subject to:} \begin{cases} \mathbf{1}^T \mathbf{x} = 1 \\ 0 \leq \mathbf{x} \end{cases} \end{aligned}$$

- Vector z encodes transcriptional activity of genes, estimated by UPC
- ► Matrix L is the Laplacian matrix, defined as A D, where d_{ii} is the weighted degree of ith vertex in the global interactome.
- \blacktriangleright Parameter α controls the relative importance of regularization

PURDUE

Convex program

$$\boldsymbol{x}^{*} = \underset{\boldsymbol{x}}{\operatorname{argmin}} \left\{ (1 - \alpha) \boldsymbol{x}^{\mathsf{T}} \boldsymbol{\mathsf{L}} \boldsymbol{x} + \alpha \parallel \boldsymbol{x} - \boldsymbol{z} \parallel_{1} \right\}$$

- The Laplacian operator L acts on a given function defined over vertices of a graph, such as x, and computes the smoothness of x over adjacent vertices.
- ▶ We can expand it as $\sum_{i,j} w_{i,j} (x_i x_j)^2$, which is the accumulated difference of values between adjacent nodes scaled by the weight of the edge connecting them.
- First term is a diffusion kernel. It propagates activity of genes through network links.

PURDUE UNIVERSITY

Convex program

$$\boldsymbol{x}^{*} = \underset{\boldsymbol{x}}{\operatorname{argmin}} \left\{ (1 - \alpha) \boldsymbol{x}^{\mathsf{T}} \boldsymbol{\mathsf{L}} \boldsymbol{x} + \alpha \parallel \boldsymbol{x} - \boldsymbol{z} \parallel_{1} \right\}$$

- The second term is a regularizer which penalizes changes or deviations
- We can expand it as ∑_i |x_i − z_i|, where x_i and z_i are the (inferred) functional and the transcriptional activity of gene i, respectively.
- It enforces sparsity over the vector of differences between transcriptional and functional activities.



What do we gain?

Tissue-specific networks have higher power/accuracy in predicting tissue-specific biology and pathobiology



Tissue-specific Pathology

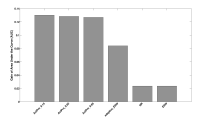
Predicting disease-related genes

	global	ActPro_0.15	ActPro_0.50	ActPro_0.85	ERW	NR
Alzheimer's disease	4.12E-3	6.96E-3	5.98E-3	5.44E-3	5.32E-3	9.60E-2
breast carcinoma	1.83E-3	1.11E-3	8.40E-4	8.30E-4	4.09E-3	8.15E-2
chronic lymphocytic leukemia	8.20E-4	7.40E-4	4.80E-4	5.10E-4	8.50E-4	2.94E-2
coronary artery disease	3.95E-1	1.58E-1	1.09E-1	1.03E-1	1.33E-1	1.93E-2
Crohn's disease	2.56E-2	1.93E-2	1.50E-2	1.44E-2	8.54E-2	4.14E-1
metabolic syndrome X	1.11E-2	1.09E-2	1.07E-2	1.12E-2	1.02E-1	7.39E-1
Parkinson's disease	1.59E-2	1.25E-2	9.89E-3	9.50E-3	1.34E-2	9.62E-2
primary biliary cirrhosis	7.20E-4	1.32E-3	3.16E-3	3.40E-3	2.80E-2	6.86E-1
psoriasis	2.10E-4	1.10E-3	1.16E-3	9.50E-4	4.67E-3	3.24E-1
rheumatoid arthritis	1.70E-2	9.28E-3	1.06E-2	1.10E-2	6.39E-2	3.61E-1
systemic lupus erythematosus	4.98E-2	1.19E-2	7.56E-3	7.22E-3	2.55E-3	1.60E-4
type 1 diabetes mellitus	2.64E-2	3.01E-2	2.38E-2	2.40E-2	2.64E-1	9.39E-1
type 2 diabetes mellitus	1.57E-3	2.90E-4	2.40E-4	1.80E-4	5.60E-4	7.90E-3
vitiligo	1.17E-3	2.13E-3	3.04E-3	3.54E-3	1.84E-2	5.69E-1
schizophrenia	3.47E-1	2.13E-1	1.93E-1	1.84E-1	1.40E-1	4.10E-2
combined	1.53E-13	1.24E-17	6.62E-19	3.70E-19	9.03E-14	2.43E-03

- 1. Symmetric random-walk as a measure of distance
- 2. Empirical *p*-value for each tissue
- 3. p-value combination using Edgington method
- ActPro excels in prioritizing disease-related genes



Grama et al.



- Edge Set Enrichment Analysis (ESEA).
- Differential correlation score:

$$\textit{EdgeScore} = \textit{MI}_{\textit{all}}(i, j) - \textit{MI}_{\textit{control}}(i, j)$$



Novel Insights Identifying disease-related pathways in brain



Prize Collecting Steiner Tree (PCST)

 $\underset{\langle v, e \rangle \in T}{\operatorname{argmin}} \left\{ \sum_{e} c_{e} - \lambda \sum_{v} b_{v} \right\}$

$$\bullet \ c_e = \frac{1}{w_e} \text{ and } b_v = \begin{cases} \infty; v \in markers \\ 1; O.W. \end{cases}$$

Red nodes are novel factors

ActPro identifies novel disease-related pathways

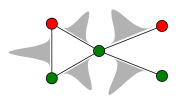


Grama et al.

Parkinson's Disease

CSol Annual Review, 2017

Goal: Identify driver network perturbations that mediate drug resistance.



- Use single-cell profiles to construct an ensemble of cell type-specific networks, one for before and one for after treatment.
- Combine individual networks within each ensemble to construct a meta-network with a distribution over each edge.
- Identify differential edges that are significantly rewired across conditions.

Key idea: A majority of perturbations do not disable proteins, but they affect individual interactions.

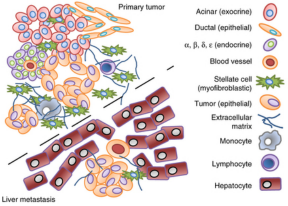


Constructing tissue-specific interactome



Motivation

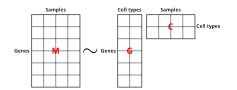
Tumor heterogeneity, including its internal diversity, as well as interaction with surrounding microenvironment, is one of the most fundamental determinants of treatment response, drug resistance, and patient relapse.



Adopted from Moffitt et al., 2015



Goal: To decompose a heterogeneous expression profile into its purified cell types



- ► M ∈ ℝ^{n×p}: Expression matrix of mixed samples
- G ∈ ℝ^{n×q}: Reference signature matrix of primary cell types.
- ► C ∈ ℝ^{q×p}: Relative proportions of each cell-type in mixture samples.

Given an observed mixture matrix **M**, find optimal **G** and **C** that approximate mixture matrix as closely as possible, according to a distance function δ , while satisfying a set of desired constraints:

Objective

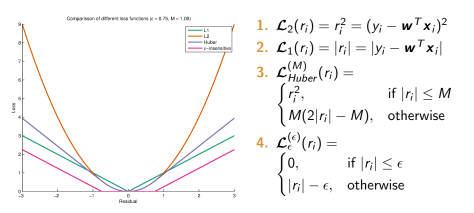
 $\min_{\mathbf{G},\mathbf{C}\in ext{feasible region}} \delta(\mathbf{GC},\mathbf{M})$



Goal: To systematically evaluate different configurations and their performance in gene expression deconvolution

- Different loss functions for evaluating estimation error
- Constraints on solutions
- Preprocessing and data filtering
- Feature selection
- Regularization







Shrinking/smoothing regression coefficients w:

$$oldsymbol{\mathcal{R}}_2(oldsymbol{w}) = \parallel oldsymbol{w} \parallel_2^2 = \sum_{i=1}^k w_i^2.$$

Sparsifying solutions :

$$\mathcal{R}_1(oldsymbol{w}) = \parallel oldsymbol{w} \parallel_1 = \sum_{i=1}^k ert w_i ert.$$



Ordinary Least Squares (OLS):

$$\min_{\boldsymbol{w}} \{ \sum_{i=1}^{m} \mathcal{L}_{2}(r_{i}) \} = \min_{\boldsymbol{w}} \{ \sum_{i=1}^{m} (y_{i} - \boldsymbol{w}^{T} \boldsymbol{x}_{i})^{2} \}$$
$$= \min_{\boldsymbol{w}} \| y - \boldsymbol{X} \boldsymbol{w} \|_{2}^{2}$$

Least Absolute Selection and Shrinkage Operator (LASSO) Regression:

$$\min_{\boldsymbol{w}} \{ \sum_{i=1}^{m} \mathcal{L}_{2}(r_{i}) + \lambda \mathcal{R}_{1}(\boldsymbol{w}) \}$$

=
$$\min_{\boldsymbol{w}} \parallel \boldsymbol{y} - \boldsymbol{X} \boldsymbol{w} \parallel_{2}^{2} + \lambda \parallel \boldsymbol{w} \parallel_{1}$$

Support Vector Regression (SVR):

$$\min_{\boldsymbol{w}} \{\sum_{i=1}^{m} \mathcal{L}_{\epsilon}(y_i - \boldsymbol{w}^T \boldsymbol{x}_i) + \lambda \mathcal{R}_2(\boldsymbol{w})\}$$

- Non-negativity (NN)
- Sum-to-one (STO)
- Similar cell quantity (SCQ)



Updating **C** is highly over-determined. We try to select genes to simultaneously minimize noise and enhance conditioning of the basis matrix \mathbf{G} :

- Range filtering
- Marker selection

New criteria: Sum-To-One (STO) violations

Violating reference gene:

$$\boldsymbol{m}(i) \leq \mathbf{G}_{min}(i); \forall 1 \leq i \leq n$$

Violating mixture gene:

$$\mathbf{G}_{max}(i) \leq \boldsymbol{m}(i); \forall 1 \leq i \leq n$$

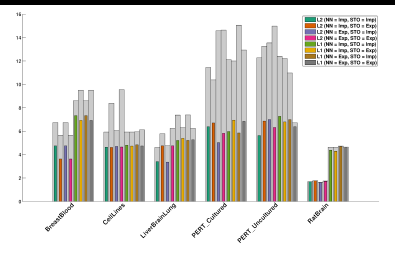


We performed comprehensive, unbiased evaluation for all combinations of these factors on the following datasets:

Dataset	# features	# samples	# references
BreastBlood	54675	9	2
CellLines	54675	12	4
LiverBrainLung	31099	33	3
PERT_Cultured	22215	2	11
PERT_Uncultured	22215	4	11
RatBrain	31099	10	4
Retina	22347	24	2



Summary of results Take home message

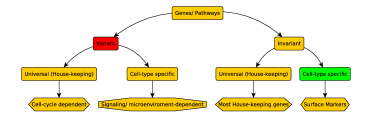


With the right choice of preprocessing and objective function, we can limit error levels in all test datasets
PURDUI

Grama et al.

CSol Annual Review, 2017

Selecting the "right" set of genes for deconvolution has one of the strongest effects on the overall deconvolution performance.



- Selecting genes that are not:
 - ► Time-dependent, such as cell cycle genes.
 - Microenvironment-dependent factors, such as genes involved in cell signaling pathways.



Motivation

- Bulk-tissue RNA-seq profiling is still more cost effective and the preferred choice for large population studies.
- Fresh specimens needed for single-cell profiling is not always available (for example in archived formalin fixed paraffin embedded (FFPE) tissue samples).
- There is a significant body of knowledge in existing databases using bulk-tissue profiling.



Thank you!

