

Comp683: Computational Biology

Lecture 19

April 8, 2024

Today

- Departure from single-cell
- Begin multi-modal integration for biomedical datasets
- Specifying a *joint subspace* for multiple samples across several modalities
- Linear algebra tricks - Rayleigh Ritz Theorem

Classical Omics Integration Problem

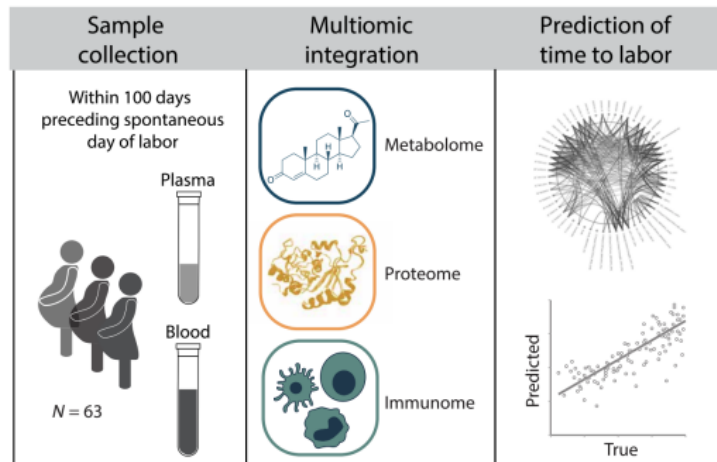


Figure: Figure from Stelzer *et al.* Science Translational Medicine. 2021. How do we leverage disparate modalities to predict something about patients, given inherent properties and quirks of each dataset?

The Cancer Genome Atlas (TCGA)

The cancer Genome Atlas was one of the first major profiling efforts, collecting diverse types of data across many patients, cancers, and biological modalities.

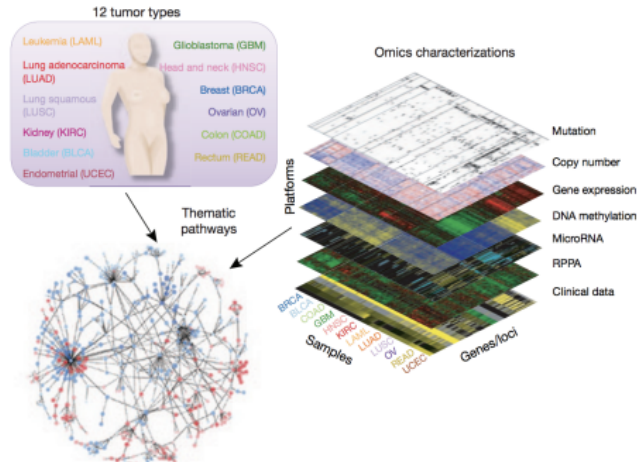


Figure: from TCGA, Nature Genetics. 2013.

FYI: LinkedOmics for Ready-to-use data with minimal pre-processing

- Download TCGA data here across many different cancers
- <http://www.linkedomics.org/login.php>

LinkedOmics "OMICS" Datatype

• Clinical Data : It includes attributes like age, overall survival, pathological stage (I, II, III, IV), TNM staging, Clinical subtype, Molecular Subtype, number of lymph nodes, radiation therapy.

• Copy Number (Level: Focal, Gene) : Normalized copy number (SNPs) and Copy number alterations for aggregated/segmented regions, per sample

• miRNA (Level: Gene, Isoform) : Normalized signals per probe or probe set for each participant's tumor sample

• Mutation (Level: Site, Gene) : Mutation calls for each participant

• Methylation (Level: Site, Gene) : Average methylation level for each participant genome, per sample

• RNAseq (Level: Gene) : The normalized expression signal of individual Gene (transcripts), per sample

• RPMA (Level: Analyte, Gene) : Normalized protein expression for each gene, per sample

• Proteomics (Level: Gene) : Average log-ratio of sample reporter-ion to common reference of peptide ions associated with the gene in acquisitions from a specific biological Sample (Unshared Log Ratio-Average log-ratio of sample reporter-ion to common reference of peptide ions of unshared peptides only associated with the gene in acquisitions from a specific biological sample).

• Phospho-Proteomics (Level: Site) : Average log-ratio of sample reporter-ion to common reference of peptide ions associated with phosphorylated site combinations in acquisitions from a specific biological sample (CDAP Protein Report).

• Glyco-Proteomics (Level: Site) : Average log-ratio of sample reporter-ion to common reference of peptide ions associated with deglycosylated N-glycosylation site combinations in acquisitions from a specific biological sample (CDAP Protein Report).

For more information ([Click here](#)) ↗

LinkedOmics Data Source

Cancer Type	Cohort Source	Cancer ID	Samples	Death Events	Median OS (yrs)	Permissions	Link	Data Download
Adrenocortical carcinoma	TCGA	ACC	92	33	NA	Y	TCGA ↗, GDAC ↗	Download ↘
Bladder urothelial carcinoma	TCGA	BLCA	412	178	2.84	Y	TCGA ↗, GDAC ↗	Download ↘
Breast invasive carcinoma	TCGA	BRCA	1097	151	10.81	Y	TCGA ↗, GDAC ↗, CPTAC ↗	Download ↘
Cervical and endocervical cancers	TCGA	CESC	307	71	8.48	Y	TCGA ↗, GDAC ↗	Download ↘

The Problem Also Comes up for Single-Cell

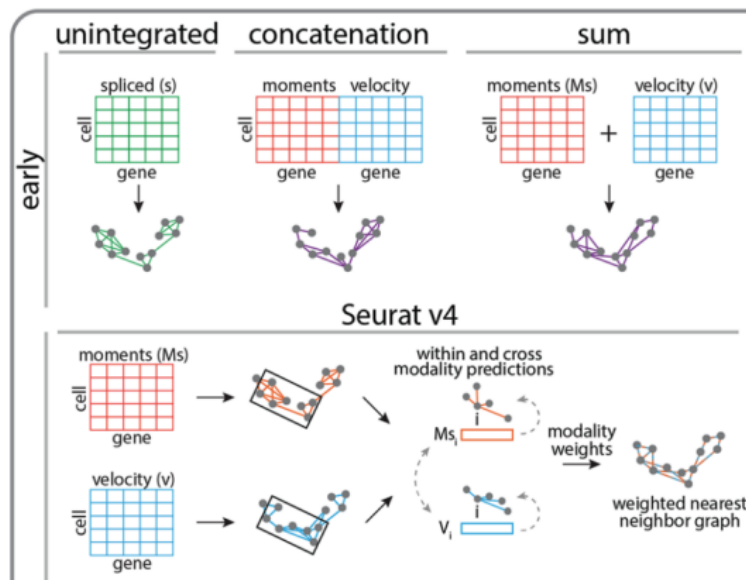


Figure: from Ranek *et al.* Genome Biology. 2022. How do we best combine various single-cell measurements to (for example) predict the label of the sample?

Notation and Problem Formulation

- Consider M types of omics data measurements $\{\mathbf{X}^m\}_{m=1}^M$ from the same set of N patients.
- For a modality, m , there are p_m measured features and the dimensions of the data matrix are therefore $p_m \times N$
- We will let G^m be the graph for modality m
- **Goal:** We seek a joint subspace embedding, $\mathbf{U} \in \mathbb{R}^{N \times k}$ that is representative of all modalities.

Comment

Before we had node2vec, we just used nice theorems from linear algebra!
:D (the OG graph embedding)

Overview of Subspace Merging

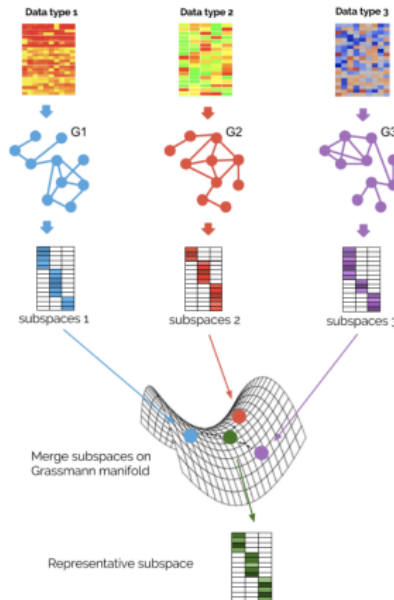


Figure: from Ding *et al.* Bioinformatics. 2019.

What is a Grassmann Manifold?

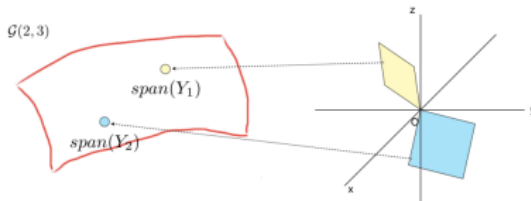


Figure: Example of $\mathcal{G}(2,3)$

- Notation, $\mathcal{G}(k, n)$ is the set of k -dimensional linear subspaces in \mathbb{R}^n , such that each subspace is a point.
- So, each point on \mathcal{G} can be represented by an orthonormal matrix $\mathbf{Y} \in \mathbb{R}^{n \times k}$
- **Selling Point:** We know how to talk about how geometrically close the subspaces are, based on principle angles

Why is this useful to our problem?

- **Each Modality Graph As A Subspace:** From each modality, we create a graph. We can ultimately compute the joint subspace or *embedding* given individual subspaces.
- **Well-Defined Distances Measures:** We know how to compute distances between subspaces on the Grassmannian. The representative subspace, \mathbf{U} should be equidistant from the per-modality subspaces (\mathbf{U}^m s).

Build a Similarity Graph Between Patients in Each Modality

Use our ‘favorite’ rule for calculating edge weights as,

$$S_{ij}^m = \exp\left(-\frac{\|\mathbf{x}_i^m - \mathbf{x}_j^m\|^2}{2t^2}\right), i = 1, \dots, N, j = 1, \dots, N$$

From here, retain the top k edges for each node based on S_{ij} and use W_{ij} for the notation of the edge weights retained, such that, $W_{ij}^m = S_{ij}^m$

Quadratic form helps with optimization problem

We already talked about the total variation of a signal in terms of the Graph Laplacian, or the variation of a signal around neighbors as,

$$\mathbf{x}^T \mathbf{L} \mathbf{x} = \frac{1}{2} \sum_{i=1}^N \sum_{j=1}^N A_{ij} (x_i - x_j)^2 \quad (1)$$

This is useful for us, because you can think of \mathbf{x} as a dimension of the embedding coordinates, which should not vary too much across the graph.

Pause for Rayleigh Ritz Theorem

Let \mathbf{A} be a square, symmetric matrix, $N \times N$ matrix with eigenvalues, $\lambda_1 \leq \lambda_2 \cdots \leq \lambda_n$ and corresponding eigenvectors $\{\mathbf{v}_1, \mathbf{v}_2, \dots, \mathbf{v}_n\}$. Then define

$$R_{\mathbf{A}}(\mathbf{x}) = \frac{\mathbf{x}^T \mathbf{A} \mathbf{x}}{\mathbf{x}^T \mathbf{x}}. \quad (2)$$

Then the minimum value of $R_{\mathbf{A}}(\mathbf{x})$ is λ_1 and it's taken for $\mathbf{x} = \mathbf{v}_1$

Matrix Extension

We will be seeing a lot on the form of $\mathbf{X}^T \mathbf{L} \mathbf{X}$. We can talk about the trace of that matrix product as the distance in vectors of adjacent nodes.

$$\text{trace}(\mathbf{X}^T \mathbf{L} \mathbf{X}) = \frac{1}{2} \sum_{i=1}^N \sum_{j=1}^N A_{ij} \|\mathbf{x}_i - \mathbf{x}_j\| \quad (3)$$

An extension of Rayleigh Ritz says that the minimum k -dimensional matrix \mathbf{X} of $\text{trace}(\mathbf{X}^T \mathbf{L} \mathbf{X})$ is $\lambda_1 + \lambda_2 + \dots + \lambda_k$ and is obtained using the first k eigenvectors of \mathbf{L} , as $\{\mathbf{v}_1, \mathbf{v}_2, \dots, \mathbf{v}_k\}$.

Specify Optimization Problem in terms of Normalized Graph Laplacian

$$\mathbf{L}^m = \mathbf{D}^{m^{-\frac{1}{2}}} (\mathbf{D}^m - \mathbf{W}^m) \mathbf{D}^{m^{-\frac{1}{2}}}$$

Written out this gives us,

$$L_{i,j}^{\text{sym}} := \begin{cases} 1 & \text{if } i = j \text{ and } \deg(v_i) \neq 0 \\ -\frac{1}{\sqrt{\deg(v_i)\deg(v_j)}} & \text{if } i \neq j \text{ and } v_i \text{ is adjacent to } v_j \\ 0 & \text{otherwise.} \end{cases}$$

Writing Down the Objective Function

The goal is to specify a \mathbf{U}^m for each modality. The optimal graph embedding in k dimensions can be written as,

$$\min_{\mathbf{U}^m \in \mathbb{R}^{N \times k}} \text{tr} \left(\mathbf{U}^{m'} \mathbf{L}^m \mathbf{U}^m \right), \quad \text{s.t. } \mathbf{U}^{m'} \mathbf{U}^m = I$$

- It turns out the solution is the first k eigenvectors of the Graph Laplacian \mathbf{L}^m by the Rayleigh–Ritz theorem

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¹Note that the $\mathbf{U}^{m'}$ refers to the transpose of \mathbf{U}^m

Defining a Projection Distance Between The Integrative Subspace and Individual Modality Subspaces

$$\begin{aligned} d_{\text{proj}}^2 \left(\mathbf{U}, \{\mathbf{U}^m\}_{m=1}^M \right) &= \sum_{m=1}^M d_{\text{proj}}^2 (\mathbf{U}, \mathbf{U}^m) \\ &= \sum_{m=1}^M [k - \text{tr} (\mathbf{U} \mathbf{U}' \mathbf{U}^m \mathbf{U}^{m'})] \\ &= kM - \sum_{i=1}^M \text{tr} (\mathbf{U} \mathbf{U}' \mathbf{U}^m \mathbf{U}^{m'}) \end{aligned}$$

The subspace, \mathbf{U} that minimizes this is close to all individual subspaces, $\{\mathbf{U}^m\}_{i=1}^M$

Optimization Problem for Multiple Subspaces

The optimization problem for merging multiple subspaces finally can be written as,

$$\min_{\mathbf{U} \in \mathbb{R}^{n \times k}} \sum_{m=1}^M \text{tr}(\mathbf{U}' \mathbf{L}^m \mathbf{U}) + \alpha \left[kM - \sum_{m=1}^M \text{tr}(\mathbf{U} \mathbf{U}' \mathbf{U}^m \mathbf{U}^{m'}) \right], \quad \text{s.t. } \mathbf{U}' \mathbf{U} = I$$

The authors showed that this simplifies to,

$$\min_{\mathbf{U} \in \mathbb{R}^{n \times k}} \text{tr} \left[\mathbf{U}' \left(\sum_{i=1}^M \mathbf{L}^m - \alpha \sum_{m=1}^M \mathbf{U}^m \mathbf{U}^{m'} \right) \mathbf{U} \right], \quad \text{s.t. } \mathbf{U}' \mathbf{U} = I$$

Rayleigh Ritz Again....

$$\min_{\mathbf{U} \in \mathbb{R}^{n \times k}} \text{tr} \left[\mathbf{U}' \left(\sum_{i=1}^M \mathbf{L}^m - \alpha \sum_{m=1}^M \mathbf{U}^m \mathbf{U}^{m'} \right) \mathbf{U} \right], \quad \text{s.t. } \mathbf{U}' \mathbf{U} = I$$

Hopefully you recognize the form of the objective. We can define a new matrix, \mathbf{L}_{mod} and again the first k eigenvectors are the optimal solution. Or,

$$\mathbf{L}_{mod} = \sum_{m=1}^M \mathbf{L}^m - \alpha \sum_{m=1}^M \mathbf{U}^m \mathbf{U}^{m'}$$

Recap

- ① **Per-Modality Graph:** Create a graph for each modality and compute corresponding Laplacians. These form the points on the Grassmannian.
- ② **Quadratic Form for Per-Modality Subspaces :** In general we want to ensure each subspace dimension respects each modality's graph structure and hence yields a small value for the quadratic form.
- ③ **Projection Distance:** The ultimate joint subspace, \mathbf{U} should be as close as possible to per-modality subspaces, \mathbf{U}^m 's
- ④ **Apply Rayleigh Ritz:** Objective is formulated in a way that we know the optimal solution is the first k eigenvectors.

Clustering on Merged Subspace

When you cluster on the merged subspace, you get groups with different prognostic interpretations.

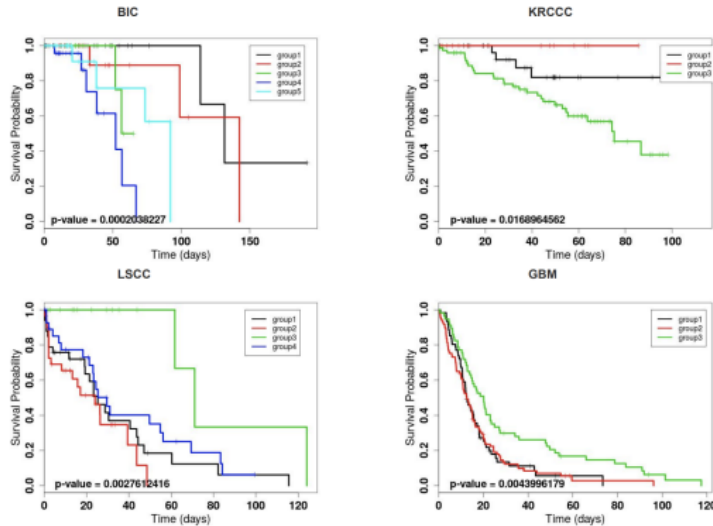


Figure: from Ding et al. Bioinformatics. 2018.

Another View : Between Patient Similarity

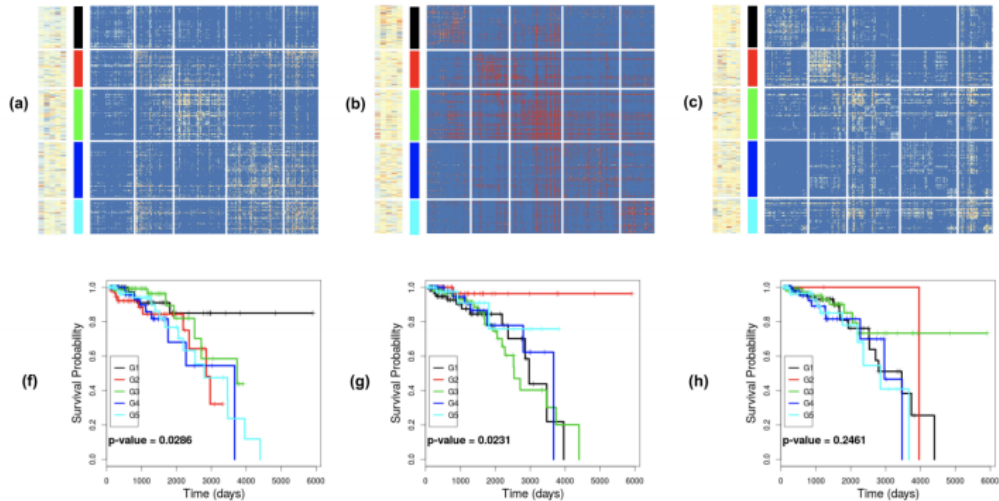


Figure: from Ding *et al.* Bioinformatics. 2018. Here we are viewing adjacency matrices between patients, based on all features jointly.

For Uncovering Trajectories Based on scRNA-seq Data

The joint subspace was used to infer the trajectory or ordering of cells.

A.

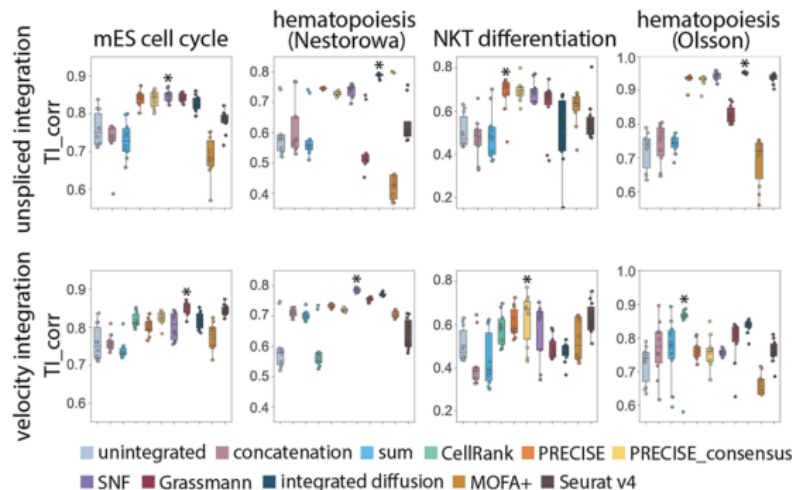


Figure: from Ranek *et al.* Genome Biology. 2022. Grassmann does pretty well, especially for integrating RNA velocity information.

Integrating Heterogeneous Information Sources

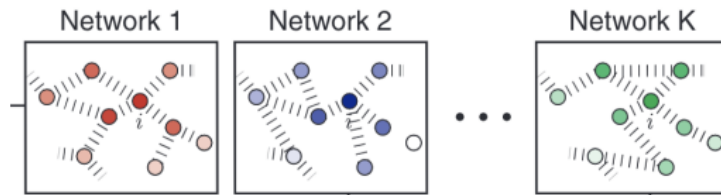


Figure: from Cho *et al.* Cell Systems. Each graph is representing a different relational definition between features.

Considering proteins, there are multiple methods for predicting whether these proteins interact .

- Physical binding
- gene expression
- co-localization
- experimentally determined
- text mined, etc.

We Seek a Unified Representations of these Nodes

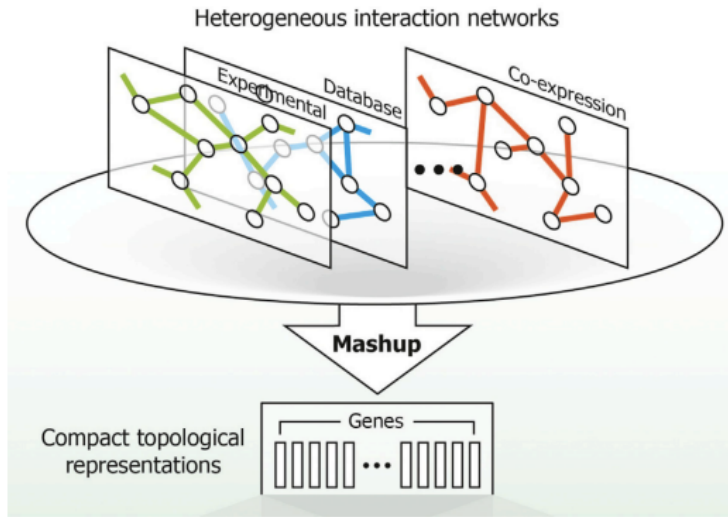


Figure: from Cho *et al.* Cell Systems. 2016.

Example from STRING

Using the STRING database, you can extract PPIs according to multiple relational definitions.

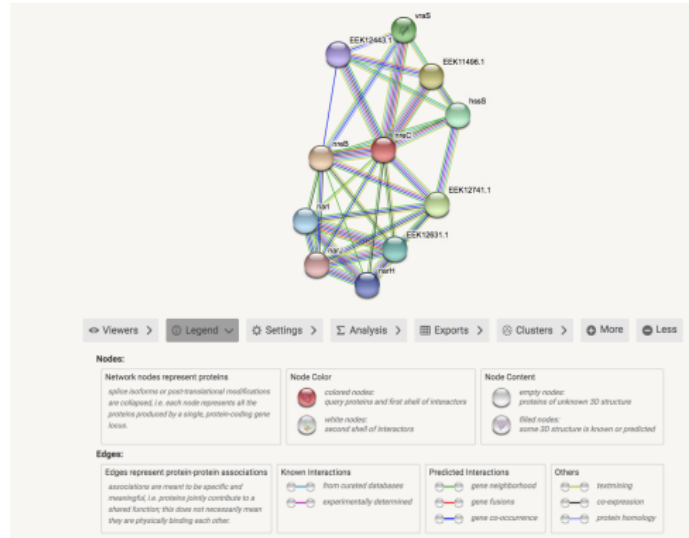


Figure: <https://string-db.org/>

Welcome Mashup

Given multiple relational definitions (e.g. multiple graphs) between a common set of nodes (features), define a consensus d -dimensional embedding vector for each node that aligns well with each individual graph (e.g. distinct relational definitions).

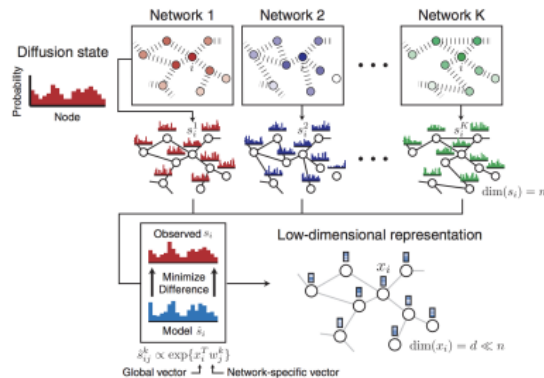


Figure: from Cho et al. Cell Systems. Each graph is representing a different relational definition between node (features).

Random Walk with Restart

- RWR is a way to account for both local and global ‘walk’ information in the graph by giving your walker the chance to restart

But first, let’s re-define the transition probability that a walker goes from node j to node i as,

$$B_{ij} = \frac{A_{ij}}{\sum_{i'} A_{i'j}}$$

RWR Formally Written

Given the transition matrix, B , the RWR from a node i is defined as,

$$s_i^{t+1} = (1 - p_r)Bs_i^t + p_r e_i$$

- p_r is the probability of restart
- e_i is an n -dimensional vector with $e_i(i) = 1$ and $e_i(j) = 0$ for $j \neq i$
- s_i^t is the vector of probabilities of each node being visited after t steps in the random walk, starting from node i

Clarifying What is Happening Here

$$s_i^{t+1} = (1 - p_r)Bs_i^t + p_r e_i$$

- The first term corresponds to following a random edge connected to the current node
- The second term corresponds to restarting from node i .
- At some point, this reaches a stationary distribution, s_i^∞ , or fixed point
- When the diffusion states between two nodes are close, this implies they have similar positions in the graph with respect to other nodes.

Quantifying Topological Overlap Between a Node Pair

Each node is given two vector representations, $\mathbf{w}_i, \mathbf{x}_i \in \mathbb{R}^d$

- Let \mathbf{w}_i refer to the context feature of a node (e.g. per relational definition)
- Let \mathbf{x}_i refer to the node feature of node i (e.g. overall)

Define a new similarity measure between nodes i and j as,

$$\hat{s}_{ij} = \frac{\exp\{\mathbf{x}_i^T \mathbf{w}_j\}}{\sum_{j'} \exp\{\mathbf{x}_i^T \mathbf{w}_{j'}\}}$$

Unpacking

$$\hat{s}_{ij} = \frac{\exp\{x_i^T w_j\}}{\sum_{j'} \exp\{x_i^T w_{j'}\}}$$

- If x_i and w_j are close in direction and hence have a large inner product, then node j should be frequently visited in the random walk starting from node i .

Recap of what is happening

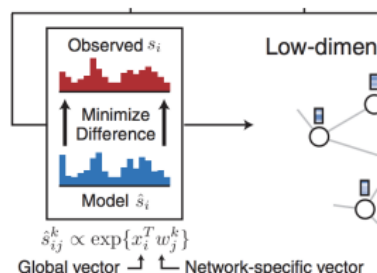


Figure: from Fig. 1. Given observed diffusion states from RWR, we should be able to find a global vector (x) and view-specific vector (w), such that a function of x and w gives a good diffusion state approximation.