Comp790-166: Computational Biology

Lecture 8

February 6, 2023

Announcement

Homework 1 is online and available, here, https://github.com/natalies-teaching/CompBio2023/tree/main/Homework1_2023. You can use the LaTeX template I provided or just submit as a PDF by 11:59pm **February 24.**

Today

- Data Augmentation for Single-Cell Data: Filling in the gaps across the cellular landscape.
- Start graph signal processing (GSP) an important tool for the upcoming attraction that is differential abundance analysis.

Do you Remember Questions

- What is the intuition for how MAGIC fixes noise and / or dropout? If given a cell, how would I adjust its features, given the rest of the data?
- What do the potential distances in PHATE do?

Rare + Sparse Cell Populations

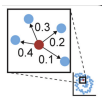


Figure: from MAGIC paper

- 'Dense' cell-populations vs 'Sparse' cell-populations.
- Low-frequency cell-populations and data artifacts can fail to be under-represented and not accurately reflect the underlying biology.
- Are members of a particular cell-type not there or are they just very infrequent?

General Problems with Downstream Tasks from Sparse Data

- Imbalanced classes can affect classification accuracy
- In clustering, imbalanced 'ground-truth' clusters can cause distortion or mis-representation of the clusters in the data
- Too few samples/observations can cause mis-representation of the dependencies or correlations between features.

Welcome SUGAR

Generate points uniformly from intrinsic data geometry / manifold:

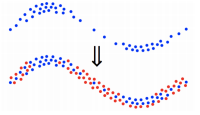


Figure: from Lindenbaum et al. NeurIPS. 2018

- Most generative modeling approaches seek to learn and replicate the data density
- SUGAR is a data-generation approach that uses the underlying geometry of the data (e.g. a random walk based approach)

Common Data Generation Approaches

- Older Techniques
 - Parametric Models: Specify a model and optimize the parameters through maximum likelihood optimization.
 - Use the learned model to generate new data
 - Use a histogram or kernel to estimate generating distributions
- Newer techniques to generate additional points from complicated data distributions
 - GAN (generative adversarial network) → https://papers.nips.cc/paper/2014/file/ 5ca3e9b122f61f8f06494c97b1afccf3-Paper.pdf
 - VAE (variational autoencoders) → https://arxiv.org/abs/1606.05908

Remembering our Good Friend Gaussian Kernel

For a pair of nodes, i and j, the strength of their connection can be computed as $\mathcal{K}(\mathbf{x}_i, \mathbf{x}_i)$, or

$$K_{ij} = \exp\left(-\frac{||\mathbf{x}_i - \mathbf{x}_j||^2}{2\sigma^2}\right) \tag{1}$$

We also remember our row-stochastic markov matrix, P computed from K.

Measure Based Gaussian Correlation (MGC)

- Given your data, X, define a set of reference points, r with $r \in X$.
- Define $\mu(\mathbf{r})$ as some measure over the reference points

Then define a new kernel, $\hat{\mathcal{K}}(\mathbf{x}_i, \mathbf{x}_j)$ as,

$$\hat{\mathcal{K}}(\mathbf{x}_i, \mathbf{x}_j) = \sum_{\mathbf{r} \in \mathbf{X}} \mathcal{K}(\mathbf{x}_i, \mathbf{r}) \mathcal{K}(\mathbf{x}_j, \mathbf{r}) \mu(\mathbf{r})$$
 (2)

For our purposes, $\mu(\mathbf{r})$ will be some value that relates to sparsity and is the inverse of node r's degree.

Problem Formulation for Data Generation Problem

- ullet Let ${\mathcal M}$ be a d-dimensional manifold such that ${\mathcal M} \in {\mathbb R}^d$
- Let $X \subset \mathcal{M}$ be a subset of points samples from \mathcal{M} with $\{x_1, x_2, \dots x_N\} \in X$
- Assuming that instances, \mathbf{X} are unevenly sampled from \mathcal{M} , we seek a set of new points, $\mathbf{Y} = \{\mathbf{y}_1, \mathbf{y}_2, \dots \mathbf{y}_M\}$ that also lie of \mathcal{M} and that $\mathbf{X} \cup \mathbf{Y}$ is uniform.

Synthesis Using Geometrically Aligned Random Walks

Algorithm 1 SUGAR: Synthesis Using Geometrically Aligned Random-walks

- 1: Compute the diffusion geometry operators K, P, and degrees $\hat{d}(i)$, i=1,...,N (see Sec. 3)
- 2: Define a sparsity measure $\hat{s}(i), i = 1, ..., N$ (Eq. 2).
- 3: Estimate a local covariance Σ_i , i = 1, ..., N, using k nearest neighbors around each x_i .
- 4: For each point i=1,...,N draw $\hat{\ell}(i)$ vectors (see Sec. 4.3) from a Gaussian distribution $\mathcal{N}(\boldsymbol{x}_i,\boldsymbol{\Sigma}_i)$. Let $\hat{\boldsymbol{Y}}_0$ be a matrix with these $M=\sum_{i=1}^N\hat{\ell}(i)$ generated vectors as its rows.

Figure: from Lindenbaum et al. NeurIPS. 2018. Let's walk through steps 1-4 for now.

Synthesis Using Geometrically Aligned Random Walks

- Specify Kernel: Initialized by forming a Gaussian Kernel over the input data, X, G_x.
 - Use G_x to estimate the degree of each node i, d(i).
 - The sparsity of each point, s(i) is defined as the inverse degree of node i or s(i) = 1/d(i).
- Sample According to Each Point Next, sample $\ell(i)$ points, $\mathbf{h}_j \in \mathbf{H}_i$ for $j = 1 \dots \ell_i$ around each $\mathbf{x}_i \in \mathbf{X}$ from a localized gaussian distribution (e.g. k-nearest points around i).
 - $G_i = \mathcal{N}(\mathbf{x}_i, \Sigma_i)$

Practical: Sampling from MV Gaussians

numpy.random.multivariate_normal

```
random.multivariate_normal(mean, cov, size=None,
check_valid='warn', tol=1e-8)
```

Draw random samples from a multivariate normal distribution.

The multivariate normal, multinormal or Gaussian distribution is a generalization of the one-dimensional normal distribution to higher dimensions. Such a distribution is specified by its mean and covariance matrix. These parameters are analogous to the mean (average or "center") and variance (standard deviation, or "width," squared) of the one-dimensional normal distribution.

Back to SUGAR

- Let \mathbf{Y}_0 be the set of all new $M = \sum_i \ell(i)$ points generated around each i, with $\mathbf{Y}_0 = \{\mathbf{y}_1, \mathbf{y}_2, \dots, \mathbf{y}_M\}$
- MGC Kernel: Now use affinities between points in **X** and **Y**₀. Here, points in **X** are used as reference.

$$\hat{\mathcal{K}}(\mathbf{y}_i, \mathbf{y}_j) = \sum_r \mathcal{K}(\mathbf{y}_i, \mathbf{x}_r) \mathcal{K}(\mathbf{x}_r, \mathbf{y}_j) s(r)$$
(3)

Pulling \mathbf{Y}_0 towards sparser regions of \mathcal{M}

Pasted psuedo code for the second part of SUGAR

- 5: Compute the sparsity based diffusion operator $\hat{\boldsymbol{P}}$ (see Sec 4.2).
- 6: Apply the operator \hat{P} at time instant t to the new generated points in \hat{Y}_0 to get diffused points as rows of $Y_t = \hat{P}^t \cdot Y_0$.
- 7: Rescale \boldsymbol{Y}_t to get the output $\boldsymbol{Y}[\cdot,j] = \boldsymbol{Y}_t[\cdot,j] \cdot \frac{\text{percentile}(\boldsymbol{X}[\cdot,j],99)}{\max \boldsymbol{Y}_t[\cdot,j]}, \ j=1,\ldots,D,$ in order to fit the original range of feature values in the data.

Figure: from Lindenbaum et al. NeurIPS. 2018.

Diffusion Operator Again

- Take affinities from $\hat{\mathcal{K}}$ and convert them to \mathbf{P} , the row-normalized Markov matrix.
- This will allow us to correct points in \mathbf{Y}_0 according to neighborhood regions

As you will all recognize, powering \mathbf{P} as \mathbf{P}^t estimates the probability of successfully traveling between nodes with t steps. The transformed matrix, \mathbf{Y}_t is computed as

$$\mathbf{Y}_t = \mathbf{P}^t \times \mathbf{Y}_o \tag{4}$$

Same Story Regarding t

Remember in PHATE, t was chosen according to the knee point of the Von-Neumann entropy of the normalized eigenvalues of \mathbf{P}^t .

 ${\sf Experiments}: \ {\sf Synthetic} \ {\sf and} \ {\sf Biological}$

Stanford Bunny

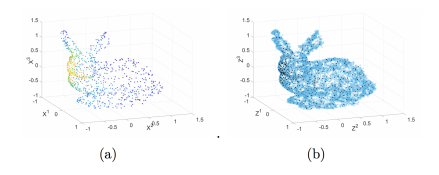


Figure: from Lindenbaum *et al.* NeurIPS. 2018. (a). The original set, \mathbf{X} of points, colored by node degree. (b) \mathbf{Y}_0 are generated points (blue) and original points, \mathbf{X} (black).

Stanford Bunny Part II

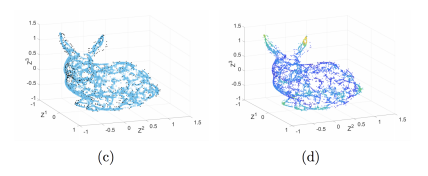


Figure: from Lindenbaum *et al.* NeurIPS. 2018. (c). Original and generated points, before MGC diffusion. (d) Generated points (Y) after MGC diffusion. Points are colored by degree.

Application : Augmented Clustering

SUGAR was used to generate additional data points to improve the quality of clusters identified with k-means.

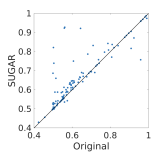


Figure: from Lindenbaum *et al.* NeurIPS. 2018. In 119 datasets, the data were augmented with additional datapoints using sugar. Adjusted Rand Index was computed for the original data vs data + SUGAR.

SUGAR on Single-Cell

SUGAR was using to augment cells in a single-cell RNAseq dataset.

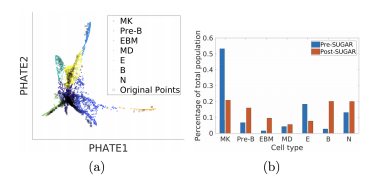


Figure: from Lindenbaum et al. NeurIPS. 2018

Maintaining Intra-Module Marker Co-Expression

Among cells assigned to the same module or cluster, after SUGAR led to higher intra-module between-marker correlation.

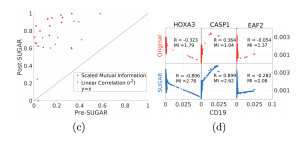


Figure: from Lindenbaum et al. NeurIPS. 2018

Graph Signal Processing

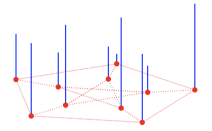


Figure: from Shuman *et al.* ArXiv. The purpose is to study the interplay between some signal and graph connectivity.

Piecewise Smooth Assumption

Translation: Nodes that are close (in terms of geodesic distance) on the graph should have similar signals. You can approximate the signal of a node, based on its neighbors.

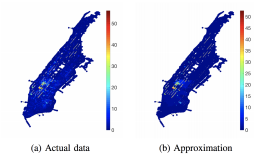


Figure: from https://arxiv.org/abs/1712.00468. Here the graph is street intersections in Manhattan and the signal is taxi pickups.

How Localized is the Signal?

Remember, our friend Graph Laplacian ($\mathbf{L} = \mathbf{D} - \mathbf{A}$),

- Some very nice theory falls out about based on the spectra of the Laplacian matrix, relating to how 'localized' a graph signal, f, is. For example f could be an expression of some protein.
 - First re-write **f** in terms of eigenvectors of the Laplacian
 - The eigenvectors corresponding to the first smallest eigenvalues of L
 are considered low frequency, and hence entries of the eigenvector
 entries corresponding to nodes that are connected should be similar
 - For higher **high frequencies** corresponding to larger eigenvalues, the values of the eigenvectors of adjacent nodes will be more different.

Signal Specificity

Here we visualize eigenvector entries at nodes $(\mathbf{u}_0, \mathbf{u}_1, \mathbf{u}_{50})$

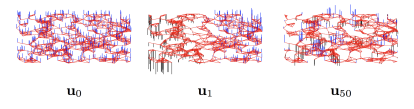


Figure: from GSP Review https://arxiv.org/abs/1211.0053

Similarly

Zero crossings mean that eigenvector entries are neighboring nodes will be different.

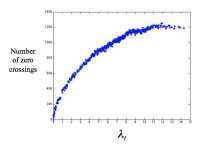


Figure: from GSP Review https://arxiv.org/abs/1211.0053

What is Graph Fourier Transform (on a high level?)

- Explain frequency content of the graph signal (e.g. experimental measurements/labels/etc) as a weighted sum of the eigenvectors of the Graph Laplacian
- The eigenvectors of the Graph Laplacian comprise the Graph Fourier Basis and can help to decouple high and low frequency signals

Local Variation of a Signal

The local variation of a signal or the sum of differences around a node can be written as,

$$(\mathcal{L}\mathbf{f})(i) = ([\mathbf{D} - \mathbf{A}]\mathbf{f})(i) \tag{5}$$

$$= d(i)\mathbf{f}(i) - \sum_{j} A_{ij}\mathbf{f}(j)$$
 (6)

$$=\sum_{i}A_{ij}(\mathbf{f}(i)-\mathbf{f}(j))\tag{7}$$

Local Variation Leads to Total Variation

The total variation of a signal on a graph is defined as follows and is also known as the Laplacian Quadratic Form

$$TV(\mathbf{f}) = \sum_{i,j} A_{ij} (\mathbf{f}(i) - \mathbf{f}(j))^2$$
 (8)

$$= \mathbf{f}^{T} \mathcal{L} \mathbf{f} \tag{9}$$

• Note here I have been assuming that we have an unweighted graph, but you could certainly substitute A_{ij} with a weighted version, W_{ij}

Getting to Graph Fourier Basis

- Start with the eigendecomposition of of **L** as $\mathbf{L} = \mathbf{\Psi} \mathbf{\Lambda} \mathbf{\Psi}^T$
- We can look at eigenvectors, $\mathbf{\Psi} = [\psi_1, \psi_2, \dots, \psi_N]$ of \mathcal{L}
- and eigenvalues, $\Lambda = [0 = \lambda_1 \leq \cdots \leq \lambda_N]$ of \mathcal{L}

The Graph Fourier Transform of a Signal

The *i*th frequency component of a signal, \mathbf{f} is the inner product between ψ_i and \mathbf{f} and can be written as,

$$\hat{\mathbf{f}}_i = \psi_i^T \mathbf{f} \tag{10}$$

The Graph Fourier Transform (GFT) is written as,

$$\hat{\mathbf{f}} = \mathbf{\Psi}^{\mathsf{T}} \mathbf{f} \tag{11}$$

GFT Will Be Used to Filter

- A filter on the graph will take in a signal and attenuate it according to a frequency response function.
- Low-Pass Filter: We filter or preserve only frequencies corresponding to eigenvalues below some threshold, λ_k . So, consider frequencies λ_b , with $\lambda_b < \lambda_k$
- **High-Pass Filters**: Preserve only frequencies corresponding to eigenvalues above some threshold, λ_k . So, consider frequencies λ_b , with $\lambda_b \geq \lambda_{k+1}$

A Simple Low-Pass Filter

Define some filter h as,

$$h: [0, \max(\mathbf{\Lambda})] \to [0, 1] \tag{12}$$

Assuming the cutoff is λ_k ,

$$h(x) > 0$$
, for $x < \lambda_k$ and $h(x) = 0$, otherwise

Defining Notation and Applying Filter to GFT

Define $h(\Lambda)$ as a diagonal matrix of eigenvalues with the filter applied. Based on what we computed with GFT, the filtered signal, $\hat{\mathbf{f}}_{filt}$ can be computed as,

$$\hat{\mathbf{f}}_{filt} = h(\mathbf{\Lambda})\hat{\mathbf{f}} \tag{13}$$

Applying a Filter in General

In general, you can filter an original signal, ${\bf f}$ in general as,

$$\underbrace{\Psi(\mathbf{I} + \alpha \mathbf{\Lambda})^{-1} \Psi^{T}}_{\text{Filtered Graph Laplacian}} \mathbf{f}.$$
(14)

Example Filters

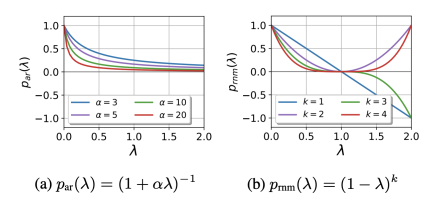


Figure: from https://openaccess.thecvf.com/content_CVPR_2019/papers/Li_Label_Efficient_Semi-Supervised_Learning_via_Graph_Filtering_CVPR_2019_paper.pdf