# Comp790-166: Computational Biology

Lecture 27

April 19, 2022

#### Announcements

- Homework 2 is due Friday.
- Project presentations start on Monday.
- Please send me your slides (preferably google slides!) prior to presentation.
- Project writeups due on the exam day, May 5
- Rubric for project writeups and presentations, https://docs.google.com/document/d/1FKh4\_ 9VK6CHwqLs2V3mTo457FwELEFYNrXwEqNbG1Bk/edit?usp=sharing

# Today

- Technical Writing in Computational Biology
- Summary of themes we have discussed wrt graph-based techniques.

# Technical Writing Motivation

- We are all busy. Make your paper clear and easy to understand.
- In Comp Bio we write for two different audiences.
- Notation, figure presentation, publicly available code goes a long way.
- Communication is Your Job!
  - Good writing through simple language and organization
  - Well-documented publicly available code

#### Question

What part of technical writing do you find the most challenging?

## Abstract: A Self Contained Story

- An elevator pitch of the main points
- Someone should read this and know exactly what your paper is about.
- Sentence breakdown
  - 1 sentence background
  - 1 sentence about what is still missing
  - 1 sentence about what you did
  - 1 sentence about what results suggest
  - 1 inspirational sentence about how this advances the field.

#### Introduction

General sections of an introduction.

- Problem motivation- what are we even talking about?
- Description of previous approaches to the problem.
  - Always highlight the work of others in a positive way
- A paragraph where you compare and contrast previous solutions. You
  can still discuss limitations by spinning them in relation to all of the
  positive things that the other authors have done.
- Paragraph giving an overview of your contributions. Someone might only read this section of your paper. You need to sell your contribution in a human-readable way.

#### Methods: First Defining Your Notation

- Notation needs to be clearly defined. There should never be a symbol in an equation that has not been properly defined.
- Keep bolding, italics, upper-case and lower-case consistent
- Dimensions of matrices need to be consistent represented with the same letter (usually p, d, or m)
- Indices should always map the same thing throughout the paper (for example *i* referring to cells and *j* referring to a feature of a cell)

## **Example Defining Notation**

We start with some notation. We assume that we have an undirected graph G=(V,E), where there are n=|V| nodes with features on each node represented by a matrix  $X\in\mathbb{R}^{n\times p}$ . Let A be the adjacency matrix of the graph, D be the diagonal degree matrix, and S be the normalized adjacency matrix  $D^{-1/2}AD^{-1/2}$ . For the prediction problem, the node set V is split into a disjoint set of unlabeled nodes U and labeled nodes L, which are subsets of the indices  $\{1,\ldots,n\}$ . We will further split the labeled nodes into a training set  $L_t$  and validation set  $L_v$ . We represent the labels by a one-hot-encoding matrix  $Y\in\mathbb{R}^{n\times c}$ , where c is the number of classes (i.e.,  $Y_{ij}=1$  if  $i\in L$  is known to be in class j, and 0 otherwise, where the ith row of Y is all zero if  $i\in U$ ), Our problem is transductive node classification: assign each node  $j\in U$  a label in  $\{1,\ldots,c\}$ , given G,X, and Y.

Figure: from Huang et al. ICLR 2021.

#### Methods: Problem Formulation

- A section where you mathematically define your problem with the notation you introduced.
- What are your inputs and outputs? What are the dimensions of the inputs and outputs and what do they represent?
- Even if you write out your problem in text format, reference the variables that you defined in the text.

For example: 'For each cell,  $\mathbf{x}_i \in \mathbb{R}^d$ , we wish to learn its label,  $y_i$  through the use of the graph,  $\mathcal{G}$ .

## Tip: Give Reminders

- It is good to keep reminding readers what notations and abstractions represent.
- For example, defining a graph? It doesn't hurt to remind them that nodes are cells and edges represent sufficient similarity between cells.
- Connect problem formulation to 'Figure 1'. In defining the overview of your problem, reference sub-panels of figure 1 of interest.

## Example of a Comprehensive Figure 1

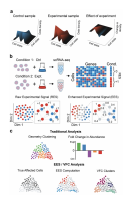


Figure: from Burkhardt et al. Nature Biotech. 2021.

#### Schematic Illustrations

If you draw cells, or patients, make sure these are carried through the entire figure.

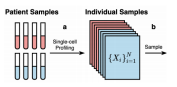


Figure: from Haidong Yi. https://www.biorxiv.org/content/biorxiv/early/2021/04/14/2021.04.13.439702.full.pdf

#### Pseudo-Code

Writing good pseudo code is extremely helpful. It can often by more helpful than the entire methods section.

```
Algorithm 2 xNetMF (G_1, G_2, p, K, \gamma_s, \gamma_a)
 1: ----- STEP 1. Node Identity Extraction -----

 for node u in V₁ ∪ V₂ do

                                        > counts of node degrees of k-hop neighbors of u
         for hop k up to K do
             \mathbf{d}_{u}^{k} = \text{CountDegreeDistributions}(\mathcal{R}_{u}^{k})
                                                                     ▶ 1 \le K \le \text{graph diameter}
         end for
                                                                    \triangleright discount factor δ ∈ (0, 1]
 8: ---- STEP 2. Efficient Similarity-based Representation -----
 9: ====== STEP 2a, Reduced n×p Similarity Computation =======

 £ = ChooseLandmarks(G<sub>1</sub>, G<sub>2</sub>,p)

    choose p nodes from G<sub>1</sub>, G<sub>2</sub>

 for node u in V do

         for node v in f do
             c_{uv} = e^{-\gamma_S \cdot ||\mathbf{d}_u - \mathbf{d}_v||_2^2 - \gamma_a \cdot \text{dist}(\mathbf{f}_u, \mathbf{f}_v)}
         end for
15: end for
                       ▶ Used in low-rank approx. of similarity graph (not constructed)
16: ----- STEP 2b. From Similarity to Representation -----
17: W = C[\mathcal{L}, \mathcal{L}]
                                             ▶ Rows of C corresponding to landmark nodes
18: [\mathbf{U}, \Sigma, \mathbf{V}] = \text{SVD}(\mathbf{W}^{\dagger})
19. \tilde{\mathbf{Y}} = \mathbf{C}\mathbf{H}\boldsymbol{\Sigma}^{-\frac{1}{2}}
                                  > Embedding: implicit factorization of similarity graph
20: \tilde{Y} = Normalize(\tilde{Y}) \Rightarrow Postprocessing: make embeddings have magnitude 1
21: \tilde{\mathbf{Y}}_1, \tilde{\mathbf{Y}}_2 = \operatorname{Split}(\tilde{\mathbf{Y}})
                                            ▶ Separate representations for nodes in G1, G2
22: return Y1, Y2
```

Figure: from https://arxiv.org/pdf/1802.06257.pdf

#### Results

- Figure/table legends should be self-contained. For example, if there is some kind of confidence interval around your curve, tell us what it represents
- Plotting: try to choose appropriate axis to capture all of the datapoints. Don't just plot for example between 0 and 1 on the y-axis by default.
- Make sure that each panel of your results figures are clearly referenced in the text.
- Avoid sloppiness. Don't let a table flow over the margin. Try to avoid different fonts and font sizes between figures.
- Colors: choose them well. Try changing default colors and removing grids from plots, etc.

#### Information to Include in Results

- Baselines: How were the baseline methods used? Did you use default parameters?
- (In real life...) you should be testing your method on several datasets (3 in biology is good).
- **Dataset description:** Describe these datasets, any pre-processing you did, and where the information can be accessed.
- **Description of Experiments:** Experiments need to be clearly described, including small details like the number of times you repeated such experiment. Always reference the figure or table where the results appear wrt a given experiment.

#### Discussion

- Recap what you have done with an overall summary
- Explain how your work complements or addresses some unmet need in the field
- Summarize your results again
- Discuss limitations and future work
- **Inspirational Parting Thought:** What is the main reason people should care and why does your work advance the field?

# Publishing in Comp Bio

- Conferences
  - ISMB
  - RECOMB
  - ACM BCB
- Journals
  - Bioinformatics
  - Cell Systems
  - Nature Journals (Nature Methods, Nature Biotech, Nature Communications)

#### Writing a Conference Paper

- Self-contained, well-structured, making it easy to read and write
- Much faster in terms of review, revision
- Appealing to CS audience.

#### Writing a Journal Article

- The main text is selling an algorithm to a broad audience.
- Heavily relies on supplemental text to get all of the relevant details.
- Very slow process. From initial submission to publication can take 1 year.
- Not as appealing to a CS audience.
- More appealing to biology audience.
- Very expensive to publish

# Providing Code

- It is good to provide code with your paper starting at the time of submission
- Repository should contain a pre-processed version of the data and instructions about how to run code on these data.

## From the Point of View of a Paper Consumer

- It is great to publish in fancy interdisciplinary journals
- It becomes less valuable to us on the CS side if the method is scattered over 100 pages of supplement
- Writing a version of your paper with all of the technical details for ArXiv is very good practice.

## A Word of Advice for Being a PhD Student in Comp Bio

Protect your expertise and your time. You are not a core facility.

- Prioritize collaborations that are mutually beneficial
- Make sure you publish your own papers without too many distractions of analyzing random datasets.
- Check where your potential collaborators put their comp bio people in the author list.

## Communicating Between Fields

- People will care about different things, between biology and computer science- tailor your details accordingly.
- You need to translate your complex model to a series of steps that don't involve mathematical phrases that we all take for granted. For example, don't say phrases like 'L1 penalty'

## Choosing What to Work On

Inspired by the talk of Quaid Morris
https://www.youtube.com/watch?v=xueh6WnpRDQ

- Don't be the state-of-the-art, be the benchmark (aka ask a new question)
- Choose hard problems rooted in biology that other people wouldn't have thought to ask because they don't read the biological literature.
- Watch the superstars who speak both languages. Watch how they publish and what they choose to work on.

## Transitioning and Summarizing What we Have Covered

We have focused on representing data as graphs and using the graphs to help us to answer questions.

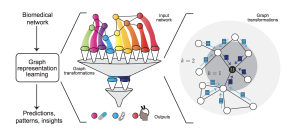
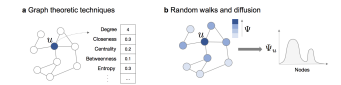


Figure: From https://arxiv.org/abs/2104.04883. For example. Assigning proteins to groups or people to outcomes.

## Class 1: Graph Summary Statistics and Diffusion

Summary statistics and diffusion can describe patterns in the graph, importance of nodes,

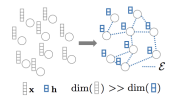


#### Graph Structure and Diffusion and Papers

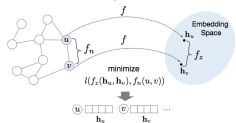
- PhenoGraph: Partition cells to cell clusters
- BigClam: For overlapping clustering
- MAGIC: for imputation in single cell data.
- **MELD:** for predicting the specificity of each cell to each condition.
- Conos: Combining multiple single cell datasets
- REGAL: graph alignment based on structural properties

#### Node Embedding Theme

#### e Manifold learning



#### f Shallow network embeddings



## Class 2: Node Embedding Theme

- Node2Vec for node embedding (embedding)
- SUGAR for data augmentation in single-cell analysis (manifold)
- SLICER for trajectory inference (manifold)
- Grassmann Embedding for combining multiple datasets (manifold)
- Mashup for embedding nodes according to multiple relational definitions (embedding)

## Class 3: Machine Learning on Graphs

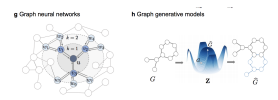


Figure: We haven't seen so much here.....

#### Seen in ML on Graphs

- Correct and Smooth for predicting labels of nodes based on simple base predictor for node features.
- More next year.....