# Graph Neural Networks Demystified

An overview of the essential concepts in Stanford CS224W (Lectures 1~9) with only oversimplified examples

### Ng Yen Kaow

### **Embeddings**

- Relatively small vectors associated with each object where similar objects have similar embeddings
- Using the embeddings of graph elements, various tasks can be performed
  - Cluster nodes in a graph
  - Predict properties of a node
  - Predict if two nodes may be connected
  - Classify entire graphs
- To perform each task, use the embedding with a suitable ML method
  - e.g. clustering can be performed with k-means

### Obtaining embeddings

- Embeddings can be formed with or learned from features
  - Node-level features
    - Degree
    - Centrality (eigenvector/ betweenness/ closeness)
    - Clustering coefficient
    - Graphlets
    - Structure-based features
  - Link-level features
    - Distance-based features
    - Local/global neighborhood overlap
  - Graph-level features
    - Graph kernels
- Task-independent embeddings can be learned from unsupervised learning

### Task-independent embeddings

Unsupervised extraction by random walks

### DeepWalk

- Estimate pairwise distance between nodes (hence their co-occurrence probability)
  - Usable for finding product relatedness in recommender
- Node embeddings
  - 1. Estimate node distances with random walks
  - 2. Train a neural network (with node input and embedding output) such that distances between embeddings agree with estimated distances

### Anonymous Walk

- Embeddings for entire graphs
- Simpler method: just add up neighbors

### Embeddings by adding neighbors

- Sum up the features of (self and) neighbor nodes
  - Features of nodes in close proximity will become similar

Example: Let  $h_i^j$  denote features of node i at iteration j and let  $h_1^0 = (1 \ 0 \ 0)$ ,  $h_2^0 = (0 \quad 1 \quad 0)$ , and  $h_3^0 = (0 \quad 0 \quad 1)$ 

#### **Initial state**



#### 1st iteration



$$h_{1}^{2} = h_{1}^{1} + h_{2}^{1} = 2 2 2 0$$

$$h_{2}^{2} = h_{1}^{1} + h_{2}^{1} = 2 2 2 0$$

$$v_{2}$$

$$v_{3}$$

 $h_1 \equiv h_2$  after only 1 iteration

CS224W Lecture 5

## Embeddings by adding neighbors

- To cluster nodes in a graph, will it work if we
  - 1. Start with a unique feature for each node, and
  - Repeatedly add up neighboring features, and
  - 3. Finally, cluster the resultant features with some method like *k*-means?



Let's try with karate club network

### Embeddings by adding neighbors





- Let matrix H be a matrix where each row is a node and each column is a feature
  - $\blacksquare$  *H* have dim  $|V| \times d$
- □ Let *A* be an adjacency matrix
  - Let  $\hat{A} = A + I$  where I is the identify matrix
- $\Box$  Then, **sum** is simply  $\hat{A}H$

$$\begin{pmatrix} a & b & c \\ & \dots & \\ & \dots & \end{pmatrix} \begin{pmatrix} h_1 \\ h_2 \\ h_3 \end{pmatrix} = \begin{pmatrix} ah_1 + bh_2 + ch_3 \\ & \dots \\ & \dots & \end{pmatrix}$$

Permutation invariant so that the outcome is the same regardless of node order within matrix

e.g. 
$$v_1$$
  $v_2$   $v_3$ 

$$\begin{pmatrix} 1 & 1 & 0 \\ 1 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix} \begin{pmatrix} h_1 \\ h_2 \\ h_3 \end{pmatrix} = \begin{pmatrix} h_1 + h_2 \\ h_1 + h_2 \\ h_3 \end{pmatrix}$$

- Let matrix H be a matrix where each row is a node and each column is a feature
  - $\blacksquare$  *H* have dim  $|V| \times d$
- □ Let *A* be an adjacency matrix
  - Let  $\hat{A} = A + I$  where I is the identify matrix
- $\square$  Further **normalize** each row of  $\hat{A}$  to sum to 1

$$\begin{pmatrix} 1/3 & 1/3 & 1/3 \\ & \dots & \\ & \dots & \end{pmatrix} \begin{pmatrix} h_1 \\ h_2 \\ h_3 \end{pmatrix} = \begin{pmatrix} (h_1 + h_2 + h_3)/3 \\ & \dots & \\ & \dots & \end{pmatrix} \begin{bmatrix} \\ \\ \\ \\ \\ \end{bmatrix}$$

Note that normalize does the same thing as mean

- Let matrix H be a matrix where each row is a node and each column is a feature
  - H have dim  $|V| \times d$
- □ Let A be an adjacency matrix
  - Let  $\hat{A} = A + I$  where I is the identify matrix
- $\ \square$  Further **normalize** each row of  $\hat{A}$  to sum to 1
  - To perform this normalization, it suffices that we let  $\hat{A} \leftarrow D^{-1}\hat{A}$  where D is the diagonal node degree matrix
    - □ In PyTorch, use torch.nn.functional.normalize(A, p=1, dim=1)
  - Or, use  $\hat{A} \leftarrow D^{-\frac{1}{2}} \hat{A} D^{-\frac{1}{2}}$ , the spectral variant
    - □ In PyTorch, use

```
D = torch.diag(torch.sum(A, 1)).inverse().sqrt()
```

```
D = torch.mm(torch.mm(D, A), D)
```

Normalized  $\hat{A}$  is in general **not symmetric** 

Redo karate club with normalized  $\hat{A} \leftarrow D^{-1}\hat{A}$ 









### Adding neighbors: evaluation

- Why do we need normalization
  - Without normalization, feature values for the nodes of high centrality would quickly add up, making them distinct from the nodes of low centrality
- □ How many iterations should be used?
  - Each iteration would "bunch up" neighboring features of 1 hop away (receptive field)
  - With that in mind, we should determine the number of iterations by the nature of the graph
  - The earliest (RNN-like) GNNs are iterated until convergence but these ideas were quickly replaced by (CNN-like) GNNs where the number of iterations is fixed as defined by the number of layers

### Relationship to PageRank

For simplicity consider eigenvector centrality problem, that is, the undirected version of PageRank

 $\ \square$  Problem Statement. Suppose each node v corresponds to a value  $x_v$  which is the sum of its neighbors' values

$$x_v = \frac{1}{\lambda} \sum_{u \in N(v)} x_u$$

where  $\lambda$  is some given constant and positive factor Given adjacency matrix A of a graph G, solve  $x_v$  for all  $v \in G$ 

- Problem is equivalent to that of finding vector x such that  $\lambda x = Ax$ 
  - Solutions are all the eigenvectors that maximize  $x^T A x$
- Problem is also equivalent to that of adding up all neighboring single-valued features, but excluding that of self, until convergence

### Adding neighbors: evaluation

- Benefits of strategy
  - Simplicity
  - Efficiently computed with adjacency matrix
- Disadvantage of strategy
  - Embeddings produced are of size of the number of nodes in the graph
    - ⇒ Learn a transformation matrix  $W: R^{|V|} \to R^d$  for some smaller d

### Transformation matrix W

- $\square$  W is typically a linear transformation layer of size  $|V| \times d$  where d is the target dimensionality of the embeddings
- $lue{}$  Combined with the adjacency matrix  $\hat{A}$ , we now have a complete matrix formulation for computing embedding  $h_v$  of a node v from (itself and) its neighbors, in the form of

$$h_v \leftarrow (\hat{A})_v HW$$

#### where

- $(\hat{A})_v$  is the row in  $\hat{A}$  for the node v, and
- H is a matrix containing the features/embeddings of all the nodes (of course, only the rows in H with non-zero entries in  $(\hat{A})_n$  are needed for computing  $h_v$ )
- □ Variations in this formula lead to various frameworks © 2021. Ng Yen Kaow

### **Variations**

- Message-aggregation (MSG-AGG)
  - First transform features/embeddings (MSG),
     then aggregate transformed embeddings (AGG)

$$h_v \leftarrow (\hat{A})_v (HW)$$
aggregate

Separate computation of self and neighbors

Exclude entry for v from  $(\hat{A})_{v}$ , and let

Aggregate only neighbors Self Learn a different transformation for self 
$$h_v \leftarrow \mathrm{AGG}\left((\hat{A})_v^{HW}, h_v^{W'}\right)$$
 Also denoted as  $B$ 

where AGG is, for instance, concatenation

### Frameworks

Graph Convolutional Network (GCN)

$$h_v \leftarrow (\hat{A})_v(HW)$$
 (basically just MSG-AGG)

### □ GraphSAGE

Exclude entry for v from  $(\hat{A})_v$ 

$$h_v \leftarrow \left( \frac{\mathsf{CONCAT}\left(\mathsf{AGG}\left((\hat{A})_v H\right), h_v\right)}{\mathsf{Concatenate self \& aggregated neighbors}} \right) W$$

AGG can be one of many options including MLP, LSTM, etc.

⇒ AGG is learnable

(Why use these? See Graph Isomorphism Network)

### Frameworks

- □ Graph Attention Networks (GAN)
  - Instead of learning AGG, learn  $\hat{A}$ 
    - Generalize the adjacency matrix  $\hat{A}$  to **attention** weights  $A = (\alpha_{vu})$

$$h_v \leftarrow (\hat{A})_v HW \Rightarrow h_v \leftarrow (A)_v HW$$
 where  $\alpha_{vu} = \frac{\exp(e_{vu})}{\sum_{x \in N(v)} \exp(e_{vx})}$ , and

 $e_{vu}$  is a measure of how related u and v are

- $e_{vu}$  is usually computed as LINEAR(CONCAT( $h_vW$ ,  $h_uW$ ))
- Do not confuse with Generative Adversarial Networks which is for generating anime pics
- Implemented in PyTorch Geometric (PyG) as GCNConv (GCN), SAGEConv (GraphSAGE), and GATConv (GAN)
  - See https://pytorch-geometric.readthedocs.io/en/latest/modules/nn.html

### Frameworks

### ■ Message Passing Neural Network (MPNN)

Involve N(v) in the transformation W for v

$$h_v \leftarrow (\hat{A})_v HW$$
  

$$\Rightarrow h_v \leftarrow H \bigoplus_{u \in N(v)} \phi(h_v, h_u)$$

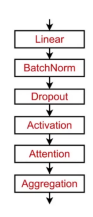
This change allows us to incorporate edge features in the embedding

$$\Rightarrow h_v \leftarrow H \bigoplus_{u \in N(v)} \phi(h_v, h_u, e_{vu})$$

- How to compute  $\phi(h_v, h_u, e_{vu})$  algebraically?
  - Let edge features be in a 3D matrix E
  - Then,  $(\hat{A})_v H$  and  $(\hat{A})_v (E)_v$  gives us two matrices with matching rows (each row corresponding to  $h_u$  and  $e_{vu}$  respectively)
  - Concatenate  $(\hat{A})_v H$  and  $(\hat{A})_v (E)_v$  and give as input to an NN
- A similar framework, Principal Neighborhood Aggregation (PNAConv), is implemented in PyG (these frameworks are not discussed in CS224W)

### In practical use

- At this point we have not mentioned activation function or other elements of DL
  - For activation function just let  $h_v \leftarrow \sigma(h_v)$
  - Mix and match as you like



- Embeddings can be used for many downstream tasks
  - We have earlier used k-means for clustering the final output
  - Better performed by constructing a neural network directly with the GNN layers

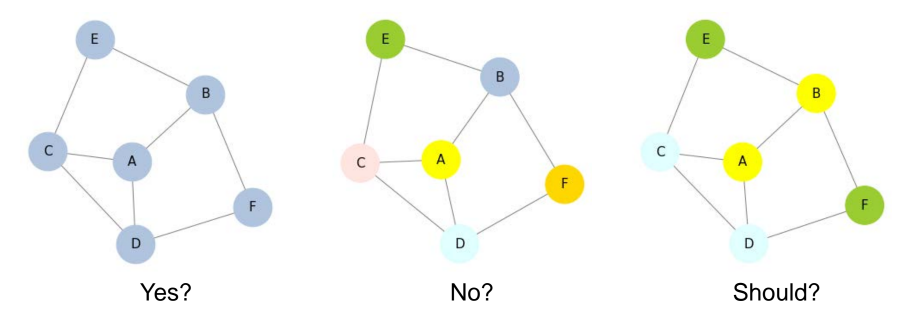
### In practical use

- Adding graph elements
  - Features
    - Similar to feature engineering
  - Virtual nodes
    - Connecting all the nodes in a sparse but apparent subgraph to a virtual node will allow those nodes to better communicate
  - Virtual edges
    - Create new graph by systematically adding edges
    - Example: Given a bipartite graph, breaking the graph into two of only nodes of the same type is good for some analyses
      - Let *A* be the adjacency matrix of the bipartite graph *G*
      - $A^2$  then gives the number of paths of distance 2 between nodes in G ⇒ an adjacency matrix between nodes of the same type
        - $\Rightarrow$  allows us to separate G into two graphs, each of same node type
      - $\blacksquare$   $A + A^2$  can form an adjacency matrix with heterogeneous edges

## Training GNNs

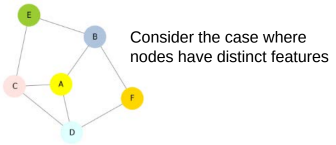
- Using node embeddings as input to a prediction function
  - Embedding of 1 node can be used directly
  - Embedding of 2 nodes can be
    - Concatenated to form an edge embedding
    - Projected on each other to get their similarity
  - Embeddings of nodes of the entire graph can be
    - □ Summed, averaged, searched for max/min, etc.
    - Clustered, then the clusters summed, average, etc., in a hierarchical fashion
- Edge embeddings from edge features are also possible, though not discussed in CS224W
  - The framework Node and Edge features in graph Neural Networks (NENN) (not yet in PyG)

 Should C and D have the same embedding in the following graphs? Given that features are given by the colors and mutually exclusive (orthogonal)

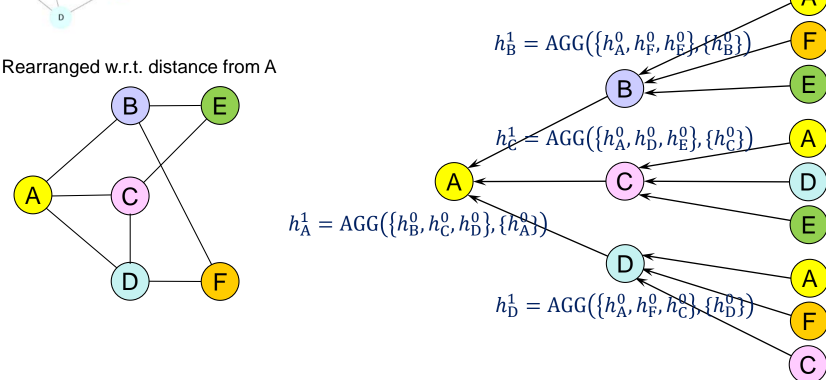


- □ How about A and B?
- Idea: Two nodes should have the same embedding if they have the same feature and neighborhood structure, and vice versa

 Given a GCN of 2 layers, the embedding of A is computed as follows



Computation graph of A's embedding



- Given a GCN of 2 layers, the embedding of A is computed as follows
  - Let  $h_A^0 = 100000$ ,  $h_B^0 = 010000$ ,  $h_C^0 = 001000$ ,  $h_D^0 = 000100$ ,  $h_D^0 = 000100$ ,  $h_D^0 = 000010$ , and let AGG be **addition**. Then
    - $h_{B}^{1} = AGG(\{h_{A}^{0}, h_{F}^{0}, h_{E}^{0}\}, \{h_{B}^{0}\}) = 1|1|0|0|1|1$   $h_{C}^{1} = AGG(\{h_{A}^{0}, h_{D}^{0}, h_{E}^{0}\}, \{h_{C}^{0}\}) = 1|0|1|1|1|0$
    - $h_{\rm D}^{1} = AGG(\{h_{\rm A}^{0}, h_{\rm F}^{0}, h_{\rm C}^{0}\}, \{h_{\rm D}^{0}\}) = 1|0|1|1|0|1$

Compute AGG(X) as  $\hat{A}H$ , where  $\hat{A}$  is the adjacency matrix (with self loop), and H is a matrix containing all the embeddings in X

Finally the embedding of A is

$$h_{A}^{2} = AGG(\{h_{B}^{1}, h_{C}^{1}, h_{D}^{1}\}, \{h_{A}^{1}\}) = 4|2|3|3|2|2$$

Similarly, 
$$h_{\rm B}^2 = 2|4|2|2|2|2$$
  
 $h_{\rm C}^2 = 3|2|4|3|2|1$   
 $h_{\rm D}^2 = 3|2|3|4|1|2$   
 $h_E^2 = 2|2|2|1|3|1$   
 $h_E^2 = 2|2|1|2|1|3$ 

- Given a GCN of 2 layers, the embedding of A is computed as follows
  - Let  $h_{A}^{0} = \boxed{100000}$ ,  $h_{B}^{0} = \boxed{010000}$ ,  $h_{C}^{0} = \boxed{001000}$ ,  $h_{D}^{0} = \boxed{000100}$ ,  $h_{D}^{0} = \boxed{000100}$ ,  $h_{D}^{0} = \boxed{000100}$ , and let AGG be **addition**. Then

$$h_A^2 = 4|2|3|3|2|2$$

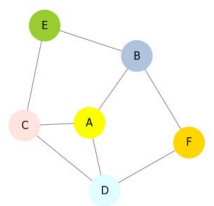
$$h_B^2 = 2|4|2|2|2|2$$

$$h_C^2 = 3|2|4|3|2|1$$

$$h_D^2 = 3|2|3|4|1|2$$

$$h_E^2 = 2|2|2|1|3|1$$

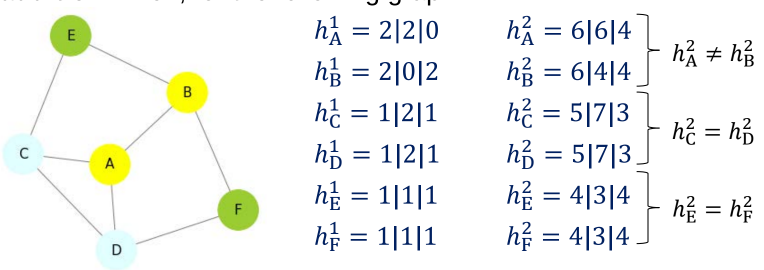
$$h_F^2 = 2|2|1|2|1|3$$



By induction they will be distinct for all subsequent iterations

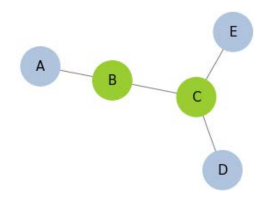
- For a graph with distinct node features, the embeddings will be distinct under addition regardless of neighborhood structure or iterations
  - With the exception of "twin nodes" that are connected only to each other (in which case they will become equal after the first iteration)

- Given a GCN of 2 layers, the embedding of A is computed as follows
  - Let  $h_A^0 = h_B^0 = \boxed{1 \odot 0}$ ,  $h_C^0 = h_D^0 = \boxed{0 \odot 1}$ ,  $h_E^0 = h_F^0 = \boxed{0 \odot 1}$ , and let AGG be addition. Then, for the following graph



- Two nodes with the same feature will always have the same embedding under addition if and only if they have the same neighborhood structure
  - What about other AGG functions, e.g. mean?

Let  $h_A^0 = h_D^0 = h_E^0 = \boxed{10}$ ,  $h_B^0 = h_C^0 = \boxed{01}$ , and let AGG be **mean**. Then, for the following graph



$$h_{\rm A}^1 = 0.5 \mid 0.5$$
  
 $h_{\rm B}^1 = 0.33 \mid 0.67$   
 $h_{\rm C}^1 = 0.5 \mid 0.5$   
 $h_{\rm D}^1 = 0.5 \mid 0.5$   
 $h_{\rm E}^1 = 0.5 \mid 0.5$ 

 $\hat{A}H$  with normalized  $\hat{A}$ 

- As expected,  $h_{\rm A}^1 = h_{\rm D}^1 = h_{\rm E}^1$  due to the same feature and neighborhood structure (within 1 hop)
- However,  $h_A^1 = h_C^1$  in spite of their differences in both features and neighborhood structure
  - ⇒ mean cannot get distinct embeddings for distinct nodes
  - Even though this is true only for the first iteration in this example, similar examples can be obtained for any number of layers

- While our earlier examples did not consider the transformation W or the activation function  $\sigma$ , the arguments are just as valid with them considered
- A function that can distinguish the nodes of distinct feature and neighborhood structure is one that is **injective**
  - mean and max are not injective
  - On the other hand, sum has problems as mentioned
- Theorem (Xu et al. 2019). Any injective AGG function can be expressed as  $\Phi(\sum_{x \in S} f(x))$  for some non-linear  $\Phi$  and linear f
- Since MLP is able to approximate any function, we can learn  $\Phi$  and f with non-linear  $MLP_{\Phi}$  and linear  $MLP_{f}$

$$AGG = MLP_{\Phi} \left( \sum_{x \in S} MLP_f(x) \right)$$

⇒ Graph Isomorphism Network (GIN)