# Unsupervised and selfsupervised Learning for Graphs

CPSC483: Deep Learning on Graph-Structured Data

Rex Ying

# Project Proposal

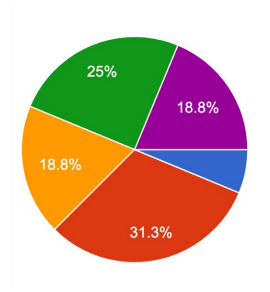
- No late days (give yourself a fall break)
- Make sure that you do preliminary analysis on dataset!
  - Not enough: Size, number of graphs
  - Reasonable: Degree distribution, centrality measures, PageRank ...
  - Great: Spectral embeddings, degree of homophily (Homework 2, Qn 2), motifs, clustering ... Innovative methods to analyze the dataset from different perspectives
- Graduate and undergrad course versions have different requirements
- Suggestion: mention how you plan to split the work among group members
  - This is necessary if the group consists of students taking undergraduate as well as graduate versions of the course

### Comments on Colabs

- Colab filenames for submission: colab2\_firstname\_lastname.ipynb
- Coding convention
  - \_\_getitem\_\_\_, \_\_setitem\_\_\_ (use slicing [])
    - See many different <u>slicing</u> functionalities
    - We usually inherit \_\_item\_\_ method to empower these slicing functionalities
  - \_\_call\_\_ : directly call the class
  - Call forward function of nn.Module (use model())
  - Similar functionalities: \_\_len\_\_(), \_\_str\_\_(), \_\_contains\_\_()
- Try to respect the prompt (e.g. write within the designated blanks)
  - If possible, don't alter the return values, arguments etc. You could simply name your variables with the specified names of the return values / arguments

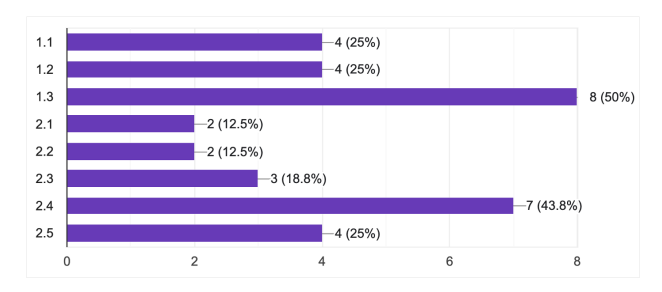
# Feedback of HW2 (1)

# Q1: How difficult do you think HW2 overall is?





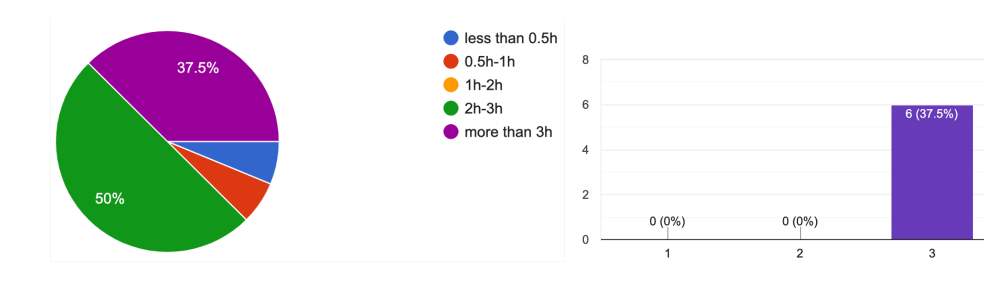
# Q2:Which questions are too difficult for you?



# Feedback of HW2 (2)

#### Q3: How much time do you spend on it?

Q4: How helpful do you think Homework 2 is?



3 (18.8%)

7 (43.8%)

## Comments on Spectral GNN Lecture

- Graph Laplacian (Homework 2)
  - Spectral embeddings: formed by first few eigenvectors of graph laplacian
- DFT: matrix multiplication (that takes  $O(n \log n)$  time)
- **Diffusion** wavelets (graph diffusion multi-han attention) Recall:  $A_{ppr} = \alpha \sum_{i=0}^{\infty} (1-\alpha)^i A^i = \sum_{i=0}^{\infty} \alpha (1-\alpha)^i A^i$

**Spectral** analysis ( $\sum_{\hat{L}_{sym}}^{\text{Proposition 2.}} \sum_{i=1}^{L} \hat{\lambda}_{i}^{g}$  and  $\lambda_{i}^{g}$  be the i-th eigeinvalues of  $\hat{L}_{sym}$  and  $\hat{L}_{sym}$ .

$$\frac{\hat{\lambda}_i^g}{\lambda_i^g} = \frac{1 - \frac{\alpha}{1 - (1 - \alpha)(1 - \lambda_i^g)}}{\lambda_i^g} = \frac{1}{\frac{\alpha}{1 - \alpha} + \lambda_i^g}.$$
 (9)

## Unsupervised and Self-supervised Learning for Graphs

- Network Community Detection
  - Network Communities
  - Louvain Algorithm
- Strategies for Pre-training Graph Neural Networks

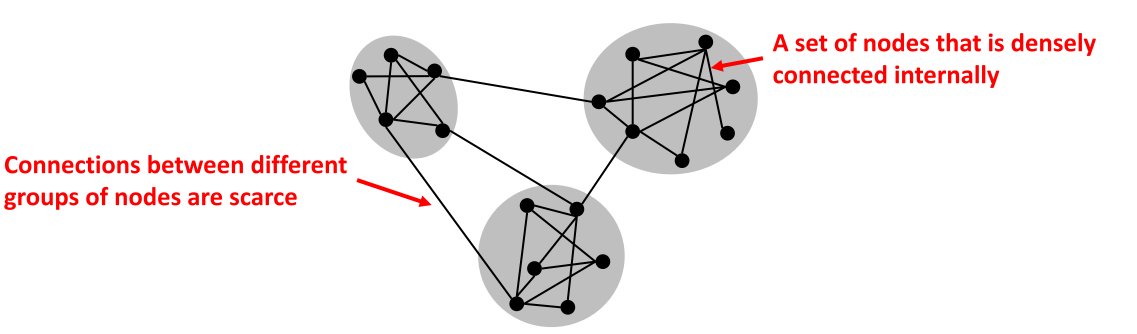
## Unsupervised and Self-supervised Learning for Graphs

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  - Louvain Algorithm

Strategies for Pre-training Graph Neural Networks

### Networks & Communities

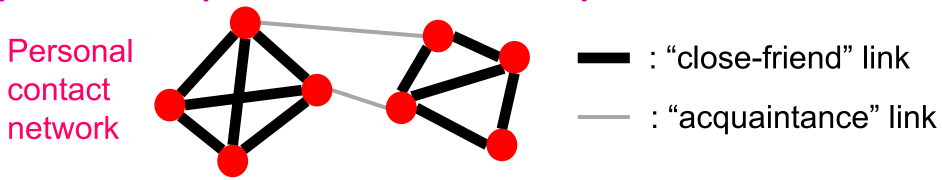
We often think of networks "looking" like this:



What led to such a conceptual picture?

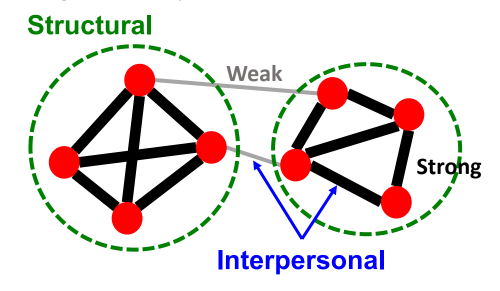
## Flow of Job Information

- How do people find out about new jobs?
  - This question is studied by Mark Granovetter, part of his PhD in 1960s
  - People find the information through personal contacts
- But: Contacts are often acquaintances rather than close friends
  - This is surprising: One would expect your friends to help you out more than casual acquaintances
- Why is it that acquaintances are most helpful?



# Granovetter's Answer (1)

- Two perspectives on friendships
  - Structural: Friendships span different parts of the network
  - Interpersonal: Friendship between two people is either strong or weak
    - Structurally embedded (tightly-connected) edges are also socially **strong**
    - Long-range edges spanning different parts of the network are socially weak

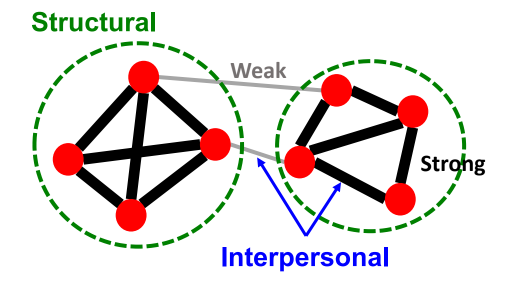


# Granovetter's Answer (2)

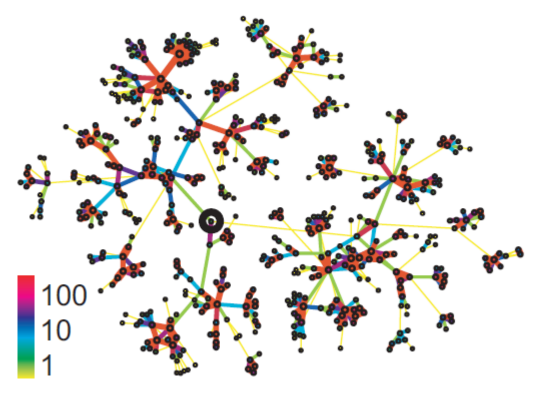
- Why is it that acquaintances are most helpful?
  - Long-range edges allow you to gather information from different parts of the network and get a job

Structurally embedded edges are heavily redundant in terms of information

access



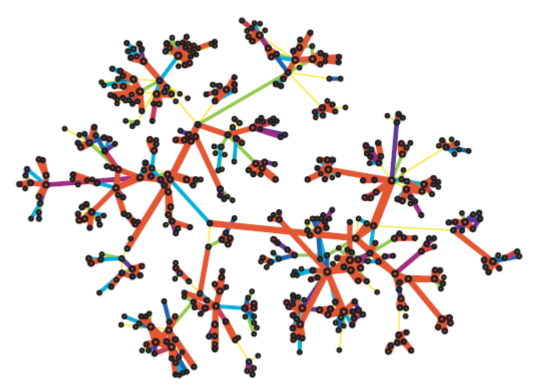
# Real Network, Real Edge Strengths



"Structure and tie strengths in mobile communication networks" Onnela et al. (2007)

- Real edge strengths in mobile call graph
  - Strong ties are more embedded (have higher overlap)

# Real Network, Permuted Tie Strengths

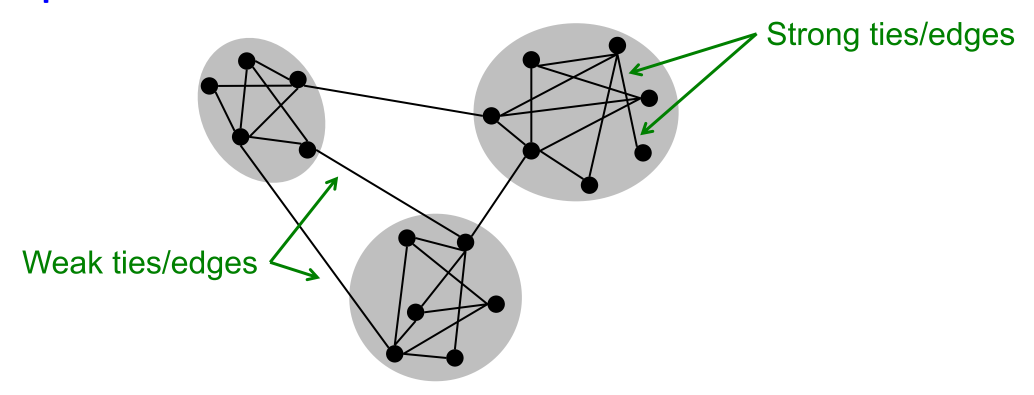


"Structure and tie strengths in mobile communication networks" Onnela et al. (2007)

• Same network, same set of edge strengths but now strengths are randomly shuffled

# Conceptual Picture of Networks

Granovetter's theory leads to the following conceptual picture of networks



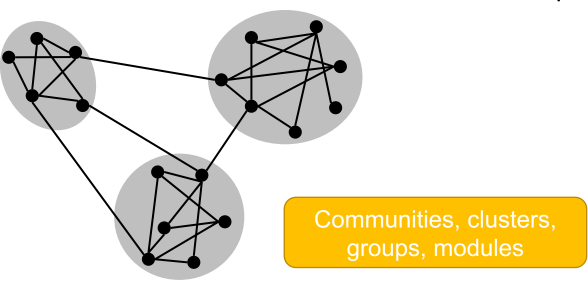
## Network Communities

 Granovetter's theory suggests that networks are composed of tightly connected sets of nodes

Network communities:

• Sets of nodes with lots of internal connections and few external ones (to the rest of

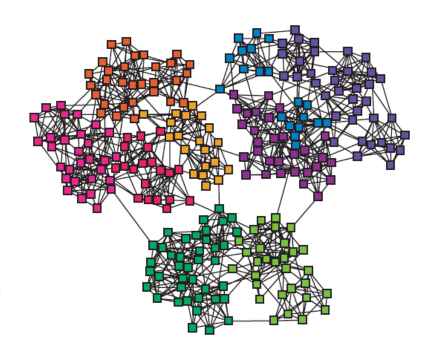
the network).



## How to Find Network Communities

 How do we automatically find such densely connected groups of nodes?

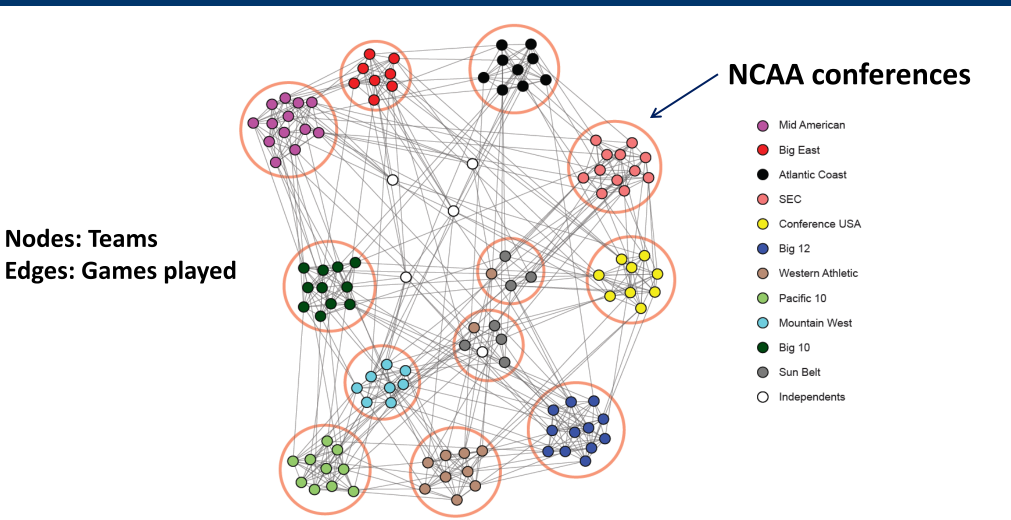
 Ideally such automatically detected clusters would then correspond to real groups



For example:

Communities, clusters, groups, modules

# Example: NCAA Football Network



**Nodes: Teams** 

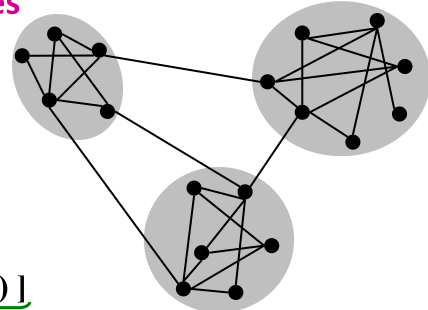
### Network Communities Measure

Communities: sets of tightly connected nodes

- Define: Modularity Q
  - A measure of how well a network is partitioned into communities
  - Given a partitioning of the network into groups disjoint  $s \in S$ :

$$Q \propto \sum_{s \in S} [ (\# \text{ edges within group } s) - (\text{expected } \# \text{ edges within group } s) ]$$

Need a null model



# Null Model: Configuration Model (1)

- Given real G on n nodes and m edges, how to know the expected number of edges between nodes i and j?
  - We can construct a network G' with uniformly random connections that has the same degree distribution as G
    - Consider G' as a multigraph (multiple edges can exist between nodes)
- The expected number of edges between nodes i and j of degrees  $k_i$  and  $k_j$

equals: 
$$k_i \cdot \frac{k_j}{2m} = \frac{k_i k_j}{2m}$$

- There are 2m directed edges (counting i->j and j->i) in total.
- For each of  $k_i$  out-going edges from node i, the chance of it landing to node j is  $k_i/2m$ , hence  $k_ik_j/2m$ .

# Null Model: Configuration Model (2)

• The expected number of edges between nodes i and j of degrees  $k_i$  and  $k_j$ 

equals: 
$$k_i \cdot \frac{k_j}{2m} = \frac{k_i k_j}{2m}$$

• The expected number of edges in (multigraph) G':

• = 
$$\frac{1}{2}\sum_{i\in N}\sum_{j\in N}\frac{k_ik_j}{2m} = \frac{1}{2}\cdot\frac{1}{2m}\sum_{i\in N}k_i(\sum_{j\in N}k_j) =$$
  
• =  $\frac{1}{4m}2m\cdot2m = m$  Note:  

$$\sum_{u\in N}k_u = 2m$$

 Under null model, both the degree distribution and the total number of edges are preserved.

Notice: This model applies to both weighted and unweighted networks. For weighted networks we use the weighted degree (sum of the edge weights).

# Modularity (1)

Modularity of partitioning S of graph G:

 $\mathbf{Q} \propto \sum_{s \in S} [$  (# edges within group s) – (expected # edges within group s) ]

$$Q(G,S) = \frac{1}{2m} \sum_{s \in S} \sum_{i \in s} \sum_{j \in s} \left( A_{ij} - \frac{k_i k_j}{2m} \right)$$

$$A_{ij} = \begin{cases} 1, \text{ if } i \to j \\ 0, \text{ otherwise} \end{cases}$$

$$\text{(if G is weighted then } A_{ij} \text{ is the edge weight)}$$

- Modularity values take range [-1,1]
  - It is positive if the number of edges within groups exceeds the expected number
  - Q greater than 0.3-0.7 means significant community structure
  - Notice Modularity applies to weighted and unweighted networks.

# Modularity (2)

$$Q(G,S) = rac{1}{2m} \sum_{s \in S} \sum_{i \in s} \sum_{j \in s} \left( A_{ij} - rac{k_i k_j}{2m} \right)$$
 For each group s

#### Equivalently modularity can be written as:

$$Q = rac{1}{2m} \sum_{ij} igg[ A_{ij} - rac{k_i k_j}{2m} igg] \delta(c_i, c_j)$$

- $A_{ij}$  represents the edge weight between nodes i and j;
- $k_i$  and  $k_j$  are the sum of the weights of the edges attached to nodes i and j, respectively;
- 2m is the sum of all of the edge weights in the graph;
- ullet  $c_i$  and  $c_j$  are the communities of the nodes; and
- $\delta$  is a simple delta function.  $\delta(c_i, c_j) = 1$  if  $c_i = c_j$  else 0

# Idea: We can identify communities by maximizing modularity

## Unsupervised and Self-supervised Learning for Graphs

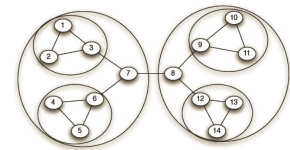
- Network Community Detection
  - Network Communities
  - Louvain Algorithm

Strategies for Pre-training Graph Neural Networks

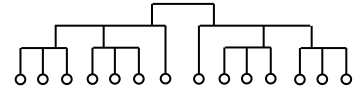
# Louvain Algorithm (1)

- Greedy algorithm for community detection
  - $O(n \log n)$  run time
- Supports weighted graphs
- Provides hierarchical communities
- Widely utilized to study large networks because:
  - Fast
  - Rapid convergence
  - High modularity output (i.e., "better communities")

#### Network and communities:



#### Dendrogram:



"Fast unfolding of communities in large networks" Blondel et al. (2008)

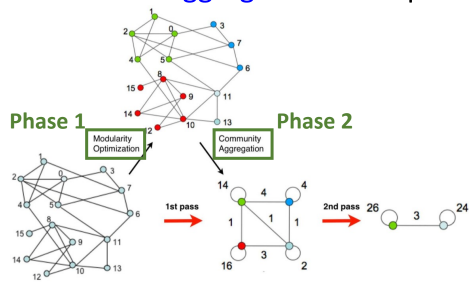
# Louvain Algorithm (2)

- Louvain algorithm greedily maximizes modularity
- Each pass is made of 2 phases:
  - Phase 1 (Partitioning): Modularity is optimized by allowing only local changes to node-communities memberships

• Phase 2 (Restructuring): The identified communities are aggregated into super-nodes

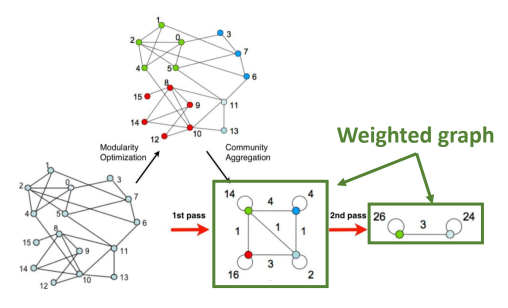
to build a new network

- Go to Phase 1
- The passes are repeated iteratively until no increase of modularity is possible.



# Louvain Algorithm (3)

- Louvain algorithm considers graphs as weighted
  - The original graph can be unweighted (i.e., edge weights are all 1)
  - As the communities get identified and aggregated into super-nodes, weighted graphs are created (weights count the number of edges in the original graph)
  - Weighted version of Modularity is applied



# Louvain: 1st phase (Partitioning)

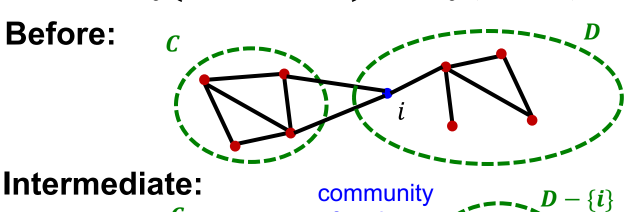
- Put each node in a graph into a distinct community (one node per community)
- For each node i, the algorithm performs two calculations
  - Compute the modularity delta ( $\Delta Q$ ) when putting node i into the community of some neighbor j
  - Move i to a community of node j that yields the largest gain in  $\Delta Q$
- Phase 1 runs until no movement yields a gain

Note that the output of the algorithm depends on the order in which the nodes are considered. Research indicates that the ordering of the nodes does not have a significant influence on the overall modularity that is obtained.

# Louvain 1st phase: Modularity Gain

#### What is $\Delta Q$ if we move node i from community D to C?

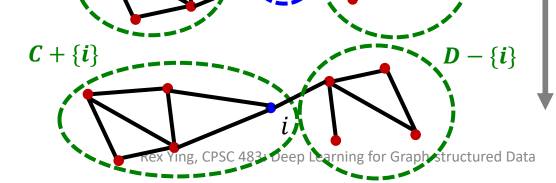
$$\Delta Q(D \to i \to C) = \Delta Q(D \to i) + \Delta Q(i \to C)$$



Removing *i* from *D* 

$$\Delta Q(D \rightarrow i)$$

After:



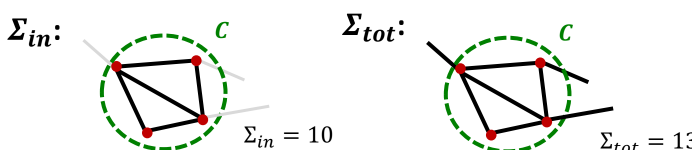
of node i

Merging *i* into *C* 

$$\Delta Q(i \rightarrow C)$$

# Louvain 1<sup>st</sup> phase: Deriving $\Delta Q(i \rightarrow C)$ (1)

- Let's derive  $\Delta Q(i \rightarrow C)$
- First, we derive modularity within C, i.e., Q(C).
- Define:
  - $\Sigma_{in} \equiv \sum_{i,j \in C} A_{ij}$ 
    - sum of link weights <u>between</u> nodes in *C*
  - $\Sigma_{tot} \equiv \sum_{i \in C} k_i$ 
    - sum of <u>all</u> link weights of nodes in *C*



# Louvain 1<sup>st</sup> phase: Deriving $\Delta Q(i \rightarrow C)$ (2)

#### • Define:

- $\Sigma_{in} \equiv \sum_{i,j \in C} A_{ij}$ ... sum of link weights <u>between</u> nodes in C
- $\Sigma_{tot} \equiv \sum_{i \in C} k_i$ ... sum of <u>all</u> link weights of nodes in C
- Then, we have

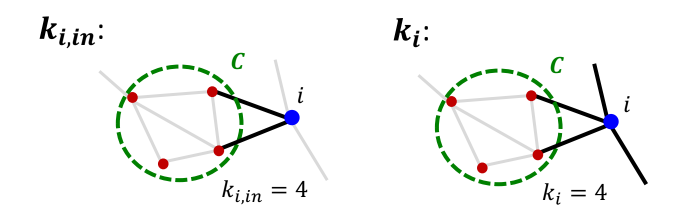
$$Q(C) \equiv \frac{1}{2m} \sum_{i,j \in C} \left[ A_{ij} - \frac{k_i k_j}{2m} \right] = \frac{\sum_{i,j \in C} A_{ij}}{2m} - \frac{(\sum_{i \in C} k_i)(\sum_{j \in C} k_j)}{(2m)^2}$$
Links within the community  $\frac{\sum_{in}}{2m} - \frac{\sum_{tot}}{2m}$  Total links

Q(C) is large when most of the total links are within-community links

# Louvain 1<sup>st</sup> phase: Deriving $\Delta Q(i \rightarrow C)$ (3)

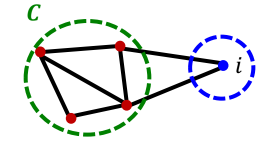
#### Further define:

- $k_{i,in} \equiv \sum_{j \in C} A_{ij} + \sum_{j \in C} A_{ji}$  ... sum of link weights connecting node i and C
  - Note that each edge gets counted twice, see formula
- $k_i$ ... sum of <u>all</u> link weights (i.e., degree) of node i



# Louvain 1<sup>st</sup> phase: Deriving $\Delta Q(i \rightarrow C)$ (4)

#### Before merging



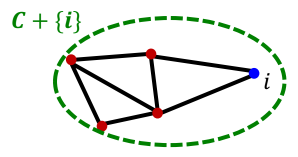
Isolated community of node *i* 

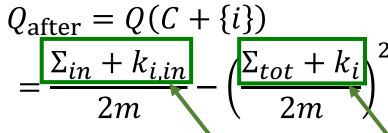
$$Q_{\text{before}} = Q(C) + Q(\{i\})$$

$$= \left[\frac{\Sigma_{in}}{2m} - \left(\frac{\Sigma_{tot}}{2m}\right)^{2}\right] + \left[0 - \left(\frac{k_{i}}{2m}\right)^{2}\right]$$

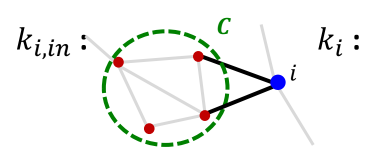


#### After merging

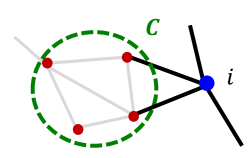








 $k_{i,in} = 4$ 



$$k_i = 4$$

"
$$\Sigma_{in}$$
" of  $C + \{i\}$  " $\Sigma_{tot}$ " of  $C + \{i\}$ 

# Louvain 1st phase: Modularity Gain

• 
$$\Delta Q(i \to C) = Q_{\text{after}} - Q_{\text{before}}$$

$$= \left[ \frac{\sum_{in} + k_{i,in}}{2m} - \left( \frac{\sum_{tot} + k_i}{2m} \right)^2 \right]$$

$$- \left[ \frac{\sum_{in} - \left( \frac{\sum_{tot}}{2m} \right)^2 - \left( \frac{k_i}{2m} \right)^2 \right]$$

- $\Delta Q(D \rightarrow i)$  can be derived similarly.
- In summary, we can compute:

$$\Delta Q(D \to i \to C) = \Delta Q(D \to i) + \Delta Q(i \to C)$$

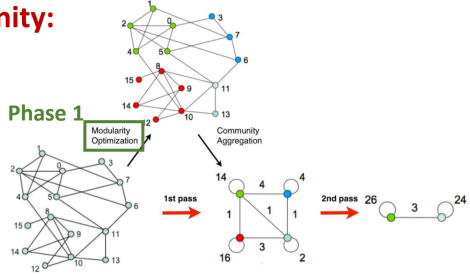
# Louvain 1st Phase: Summary

- Iterate until no node moves to a new community
  - For each node  $i \in V$  currently in community C, compute the best community C':

$$C' = \operatorname{argmax}_{C'} \Delta Q(C \rightarrow i \rightarrow C')$$

• If  $\Delta Q(C \rightarrow i \rightarrow C') > 0$ , then update the community:

- $C \leftarrow C \{i\}$
- $C' \leftarrow C' + \{i\}$

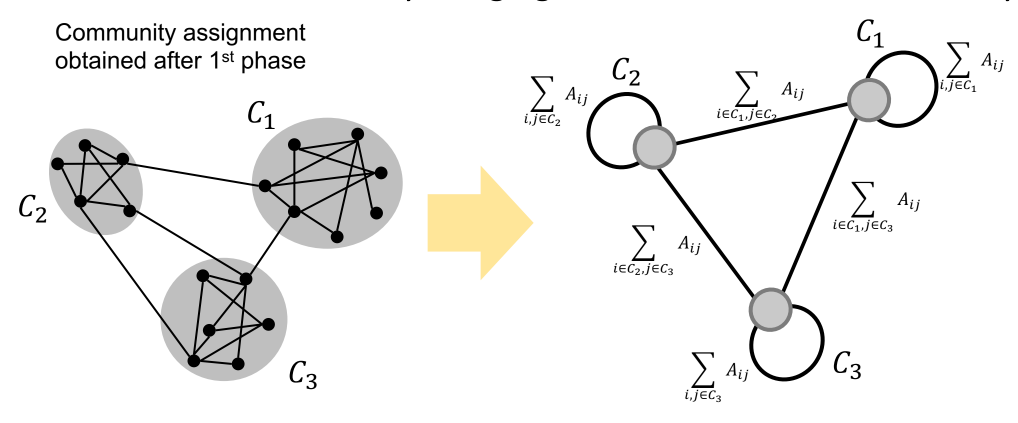


# Louvain: 2<sup>nd</sup> phase (Restructuring)

- The communities obtained in the first phase are contracted into super-nodes, and the network is created accordingly:
  - Super-nodes are connected if there is at least one edge between the nodes of the corresponding communities
  - The weight of the edge between the two super-nodes is the sum of the weights from all edges between their corresponding communities
- Phase 1 is then run on the super-node network

#### Louvain 2st Phase: Summary

Super nodes are constructed by merging nodes in the same community.



#### Interesting Questions

- Can we use GNNs to perform clustering / community detection?
  - Direct classification of nodes into clusters
  - Link prediction
- Can we use community information to improve GNN models
  - Additional node features
  - ClusterGCN
  - DiffPool
- Theoretical questions
  - Investigate the relation between clustering and over-smoothing phenomenon

#### Unsupervised and Self-supervised Learning for Graphs

- Network Community Detection
  - Network Communities
  - Louvain Algorithm

Strategies for Pre-training Graph Neural Networks

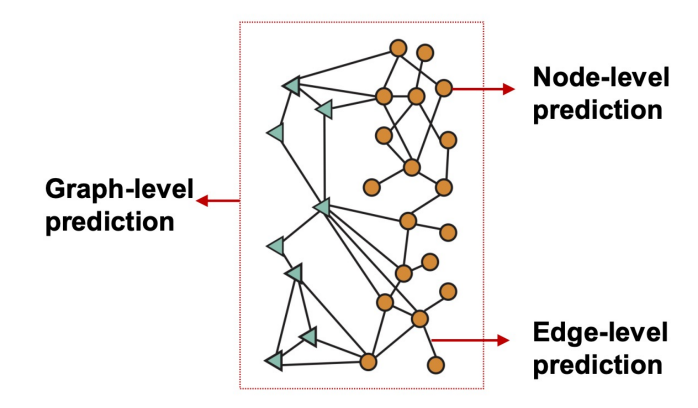
#### Supervised Learning on Graphs (1)

- Usually we train a GNN with some supervised labels:
  - Node labels  $y_v$ : in a citation network, which subject area does a node belong to
  - Edge labels  $y_{uv}$ : in a transaction network, whether an edge is fraudulent
  - Graph labels  $y_G$ : among molecular graphs, the drug likeness of graphs
- And apply a loss function  $l(\cdot)$  to optimize the parameters  $\min l(y, \widehat{y})$

**Predicted labels** 

# Supervised Learning on Graphs (2)

Different supervised labels on graph



#### Challenge of Supervised Learning on Graphs

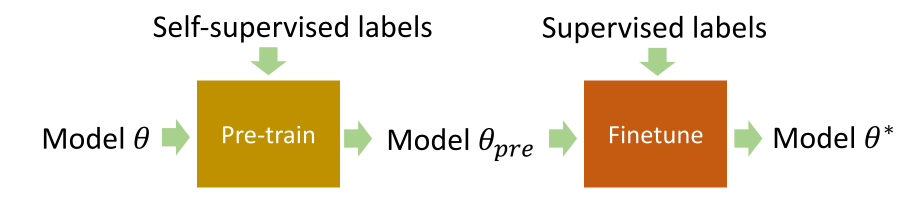
- Task-specific labels can be extremely scarce
  - Obtaining labels requires expensive lab experiments and human annotation
    - Machine learning models easily overfit to small training data
  - E.g., testing the chemical property of a molecule in a wet laboratory
- Graph data often contains out-of-distribution samples
  - Graphs in the training set are structurally very different from graphs in the test set
  - E.g., predict chemical properties of a new molecule which is different from all molecules in training set

#### Self-supervised Learning on Graphs (1)

- How to improve model's out-of-distribution prediction performance even with limited data?
- The solution: self-supervised learning!
  - We can find abundant supervision signals within the graph
    - Node-level  $y_v$ . Node statistics: such as clustering coefficient, PageRank
    - Edge-level  $y_{uv}$ . Link prediction: hide the edge between two nodes, predict if there should be a link
    - Graph-level  $y_G$ . Graph statistics: for example, predict if two graphs are isomorphic
  - Self-supervised learning can inject domain knowledge into a model
    - Model can generalize well without many task-specific labeled data

# Self-supervised Learning on Graphs (2)

- Key idea: Use self-supervised signals to pre-train a model, then finetune the model with (scarce) training data
  - Pre-train: train the model on relevant tasks with self-supervised signals
  - Finetune: adapt the model to downstream task by using task-specific labels to tune the pretrained model



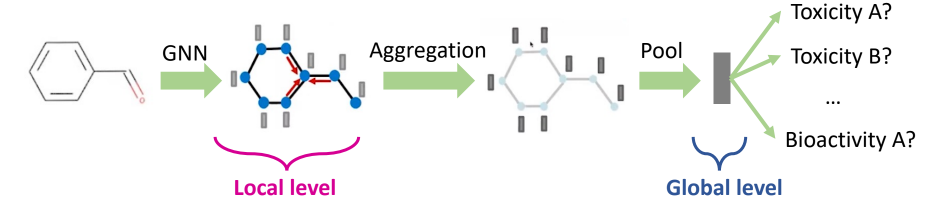
# Pre-training GNNs (1)

- How to design pre-training strategies for GNNs?
- Let's think about molecular property prediction task
  - Node: atom
  - Edge: interaction between two atoms
  - Task: Given a molecular graph, predict its corresponding chemical property

$$f(\bigcirc) = \begin{cases} Toxicity A? \\ Toxicity B? \\ \vdots \\ Bioactivity A? \end{cases}$$

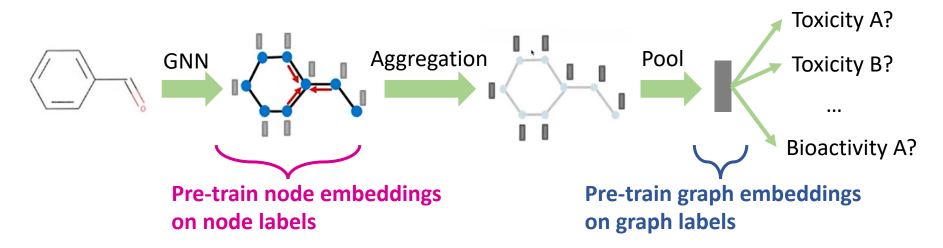
# Pre-training GNNs (2)

- We can apply GNN to learn the molecular graph representation
  - Local level: Iteratively perform neighbor aggregation to obtain node representation
  - Global level: Use pooling operation to obtain graph representation



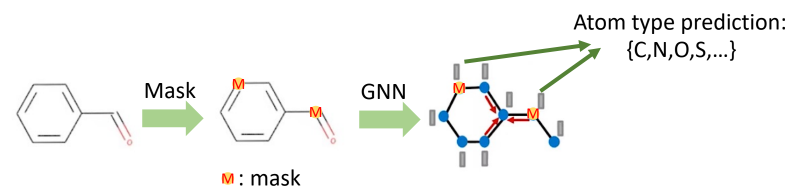
# Pre-training GNNs (3)

- How to pre-train GNN for molecular graph?
  - We can pre-train both node and graph embeddings
  - GNN can capture domain-specific knowledge of both local and global structure



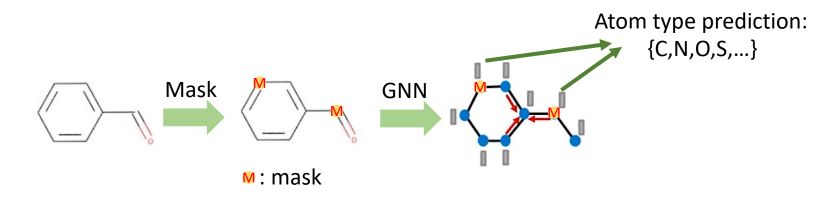
# Node-level Pre-training: Attribute Masking (1)

- Node-level self-supervised task: Attribute Masking
  - Self-supervised signal: original node feature (atom type)
  - Mask some node features
  - Use GNN to generate node embeddings
  - Apply a linear model on the embeddings to predict the masked node feature



# Node-level Pre-training: Attribute Masking (2)

- Intuition: atoms on the molecular graph follow chemistry rules
- Attribute masking can enforce GNN to learn such domain knowledge
  - Valency
  - Electronic or steric properties of functional groups

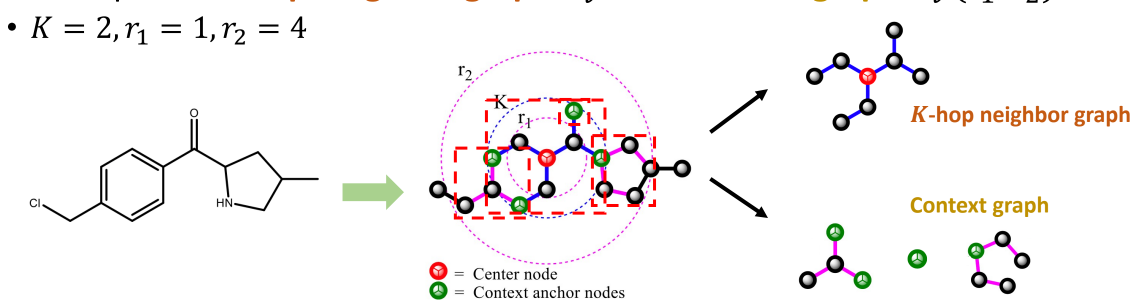


#### Node-level Pre-training: Context Prediction (1)

- Node-level self-supervised task: Context prediction
  - Self-supervised signal: context graph
- K-hop neighbor graph  $G_v^K$ 
  - The subgraph containing all nodes and edges that are at most K-hop away from v
- Context graph  $G_v^c(r_1, r_2)$ 
  - The subgraph containing all nodes and edges that are between  $r_1$ -hop and  $r_2$ -hop away from v
  - It is a ring of width  $r_2 r_1$
- Given a center node v, we require  $r_1 < K$  and the nodes which are shared between neighbor and context graph are context anchor nodes

# Node-level Pre-training: Context Prediction (2)

• An example of K-hop neighbor graph  $G_v^K$  and Context graph  $G_v^C(r_1, r_2)$ 



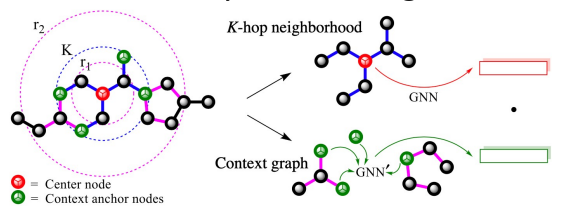
# Node-level Pre-training: Context Prediction (3)

- Node-level self-supervised task: Context prediction
  - Key idea: use subgraphs to predict their surrounding graph structures
  - Encode the  $G^K_v$  using the main GNN to obtain center node embedding  $\mathbf{h}^K_v$
  - Encode the  $G_v^c(r_1,r_2)$  using context GNN to obtain the context anchor node embeddings
    - Context GNN is an auxiliary GNN
  - Simply average all the context anchor nodes' embedding to obtain context embedding  $\mathbf{h}_{v}^{c}$
  - Minimize the distance between  $\mathbf{h}_v^K$  and  $\mathbf{h}_v^c$ :

$$\min \sum_{v} 1 - \sigma(\mathbf{h}_{v}^{K} \cdot \mathbf{h}_{v}^{c})$$
Inner product
Inner product

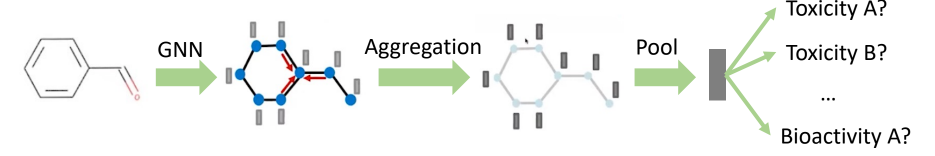
# Node-level Pre-training: Context Prediction (4)

- Intuition: subgraphs that are surrounded by similar contexts are semantically similar
- Pre-trained with Context prediction task, GNN can map nodes appearing in similar structural contexts to nearby embeddings



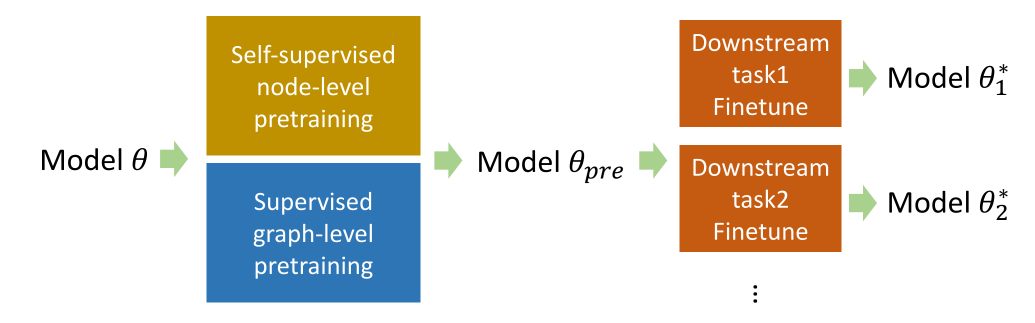
#### Graph-level Pre-training: Attribute Prediction

- Graph-level supervised task: Attribute prediction
  - Multi-task supervised pretraining: Predict a diverse set of supervised labels
    - Toxicity A? Toxicity B? Bioactivity A? Bioactivity B?
  - Each property corresponds to a binary classification task
  - Training GNN on many relevant tasks simultaneously



#### Pre-training on Graph: Overview

- Pretraining a GNN on molecular graph
  - Node-level pretraining: attribute masking and context prediction
  - Graph-level pretraining: attribute prediction



#### Summary

- Unsupervised learning for graph: community detection
  - What is Network Communities?
    - Network community measure Modularity
  - Detection method: Louvain Algorithm
    - Phase 1 (Partitioning) and Phase 2 (Reconstructing)
- Strategies for pretraining Graph Neural Networks
  - Node-level pretraining
  - Graph-level pretraining