## Beyond Small Graphs: Scale GNNs to Large Graphs

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CS598: Deep Learning with Graphs, 2024 Fall

https://ulab-uiuc.github.io/CS598/

- Recommender systems:
  - Amazon
  - YouTube
  - Pinterest
  - Etc.
- ML tasks:
  - Recommend items (link prediction)
  - Classify users/items (node classification)

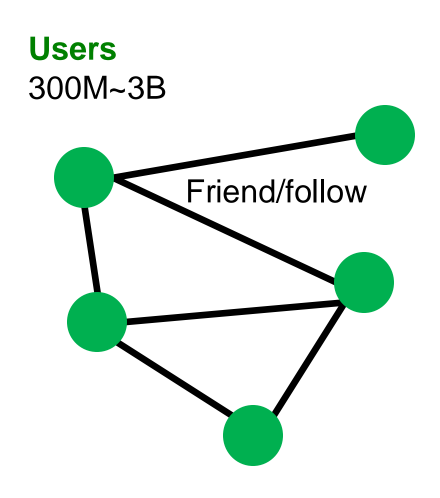
**Users Products/Videos** 100M~1B 10M ~ 1B Bought/saw

#### Social networks:

- Facebook
- Twitter
- Instagram
- Etc.

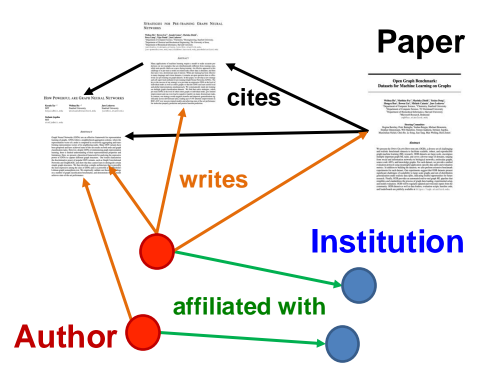
#### ML tasks:

- Friend recommendation (link-level)
- User property prediction (node-level)



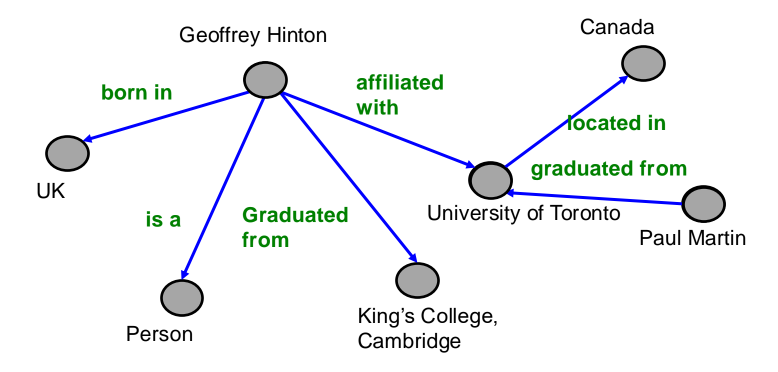
- Academic graph:
  - Microsoft Academic Graph
- ML tasks:
  - Paper categorization (node classification)
  - Author collaboration recommendation
  - Paper citation recommendation (link prediction)

Papers Authors 120M 120M



- Knowledge Graphs (KGs):
  - Wikidata
  - Freebase
- ML tasks:
  - KG completion
  - Reasoning

#### Entities 80M—90M



#### What is in Common?

#### Large-scale:

- #nodes ranges from 10M to 10B.
  - E.g., 1B nodes, 20 dim float32 feat -> 1B\*20\*4 = 80GB memory for node features
- #edges ranges from 100M to 100B.
  - E.g., 10B edges, int32 -> 10B\*2\*4 = 80GB memory for sparse adj matrix

#### Tasks

- Node-level: User/item/paper classification.
- **Link-level**: Recommendation, completion.

#### Todays' lecture

Scale up GNNs to large graphs!

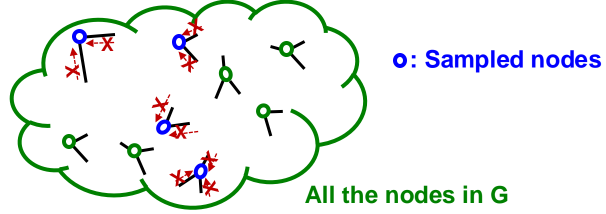
- Recall: How we usually train an ML model on large data (N=#data is large)
- Objective: Minimize the averaged loss

$$\ell(\boldsymbol{\theta}) = \frac{1}{N} \sum_{i=0}^{N-1} \ell_i(\boldsymbol{\theta})$$

- $\theta$ : model parameters,  $\ell_i(\theta)$ : loss for i-th data point.
- We perform Stochastic Gradient Descent (SGD).
  - Randomly sample  $M (\ll N)$  data (mini-batches).
  - Compute the  $\ell_{sub}(\boldsymbol{\theta})$  over the M data points.
  - Perform SGD:  $\theta \leftarrow \theta \nabla \ell_{sub}(\theta)$

#### What if we were to use the standard SGD for GNN?

• In mini-batch, we sample  $M (\ll N)$  nodes independently:



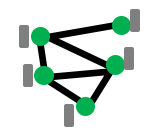
- Sampled nodes tend to be isolated from each other.
- Recall: GNN generates node embeddings by aggregating neighboring node features.
  - GNN does not access to neighboring nodes within the mini-batch!
- Standard SGD cannot effectively train GNNs.

 Naïve full-batch implementation: Generate embeddings of all the nodes at the same time:

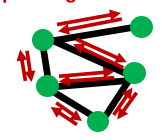
$$H^{(k+1)} = \sigma(\tilde{A}H^{(k)}W_k^{\mathrm{T}} + H^{(k)}B_k^{\mathrm{T}})$$

- Load the entire graph A and features X. Set  $H^{(0)} = X$ .
- At each GNN layer: Compute embeddings of all nodes using all the node embeddings from the previous layer.
- Compute the loss
- Perform gradient descent

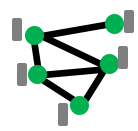
Given **all node embeddings** at layer K



Perform messagepassing



Obtain **all node embeddings** at layer K+1



- However, Full-batch implementation is not feasible for a large graphs.
  Why?
- Because we want to use GPU for fast training, but GPU memory is extremely limited (only 10GB--80GB).
  - The entire graph and the features cannot be loaded on GPU.

Slow computation, large memory

CPU 1TB—10TB Fast computation, limited memory

GPU 10GB—80GB

#### Today's Lecture

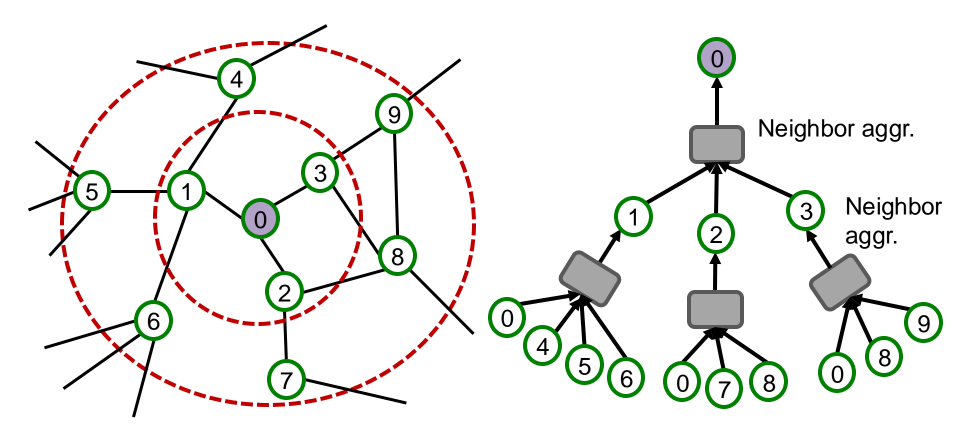
#### We introduce three methods for scaling up GNNs:

- Two methods perform message-passing over small subgraphs in each mini-batch; only the subgraphs need to be loaded on a GPU at a time.
  - Neighbor Sampling [Hamilton et al. NeurIPS 2017]
  - Cluster-GCN [Chiang et al. KDD 2019]
- One method simplifies a GNN into feature-preprocessing operation (can be efficiently performed even on a CPU)
  - Simplified GCN [Wu et al. ICML 2019]

# Beyond Small Graphs: Scale GNNs to Large Graphs GraphSAGE Neighbor Sampling: Scaling up GNNs

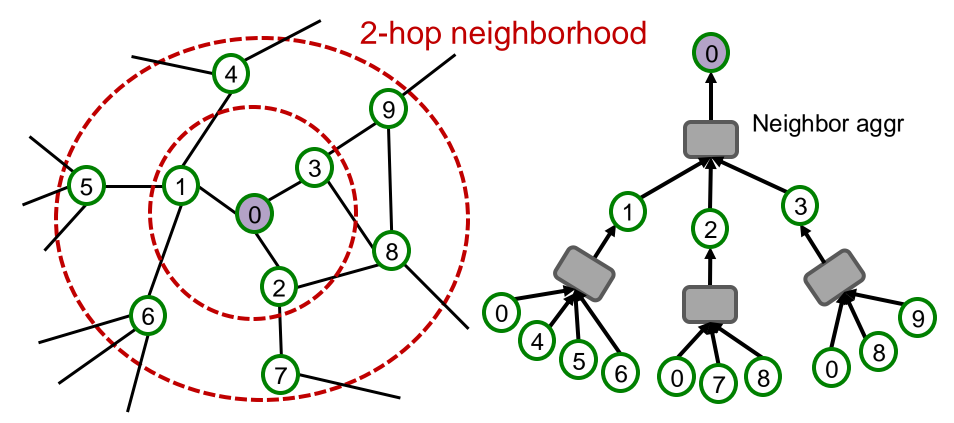
### Recall: Computational Graph

- Recall: GNNs generate node embeddings via neighbor aggregation.
  - Represented as a computational graph (right).



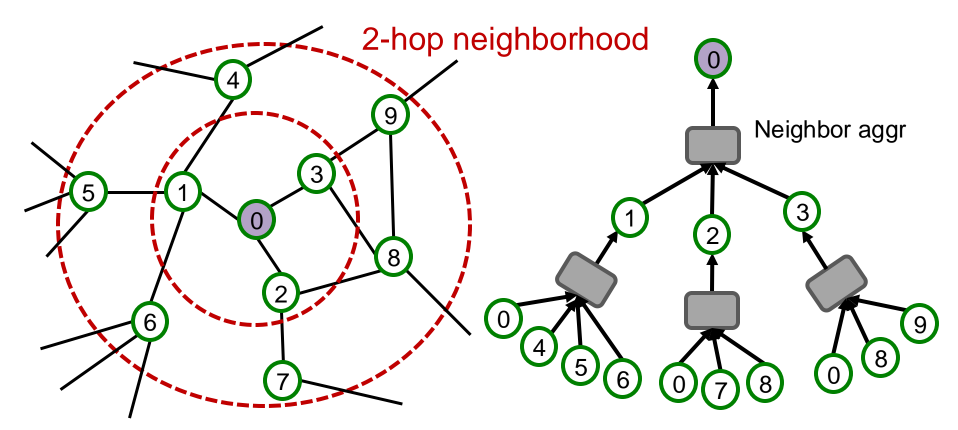
#### Recall: Computational Graph

Observation: A 2-layer GNN generates embedding of node "0" using 2-hop neighborhood structure and features.



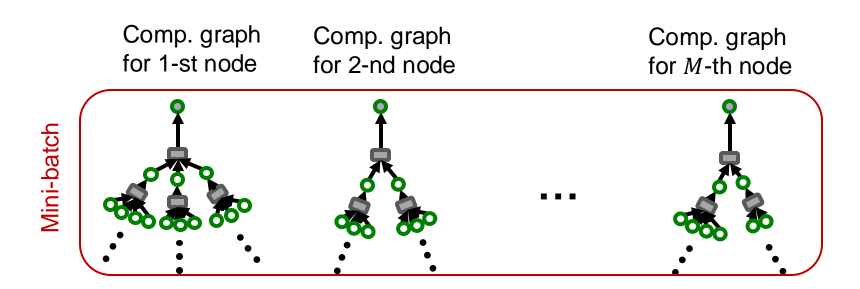
#### Recall: Computational Graph

• Observation: More generally, K-layer GNNs generate embedding of a node using K-hop neighborhood structure and features.



### Computing Node Embeddings

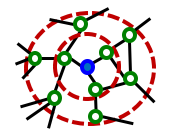
- Key insight: To compute embedding of a single node, all we need is the K-hop neighborhood (which defines the computation graph).
- Given a set of M different nodes in a mini-batch, we can generate their embeddings using M computational graphs. Can be computed on GPU!



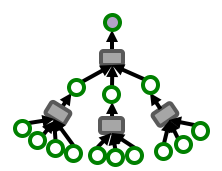
### Stochastic Training of GNNs

- We can now consider the following SGD strategy for training K-layer GNNs:
  - Randomly sample  $M (\ll N)$  nodes.
  - For each sampled node v:
    - Get K-hop neighborhood, and construct the computation graph.
    - Use the above to generate v's embedding.
  - Compute the loss  $\ell_{sub}(\boldsymbol{\theta})$  averaged over the M nodes.
  - Perform SGD:  $\theta \leftarrow \theta \nabla \ell_{sub}(\theta)$

*K*-hop neighborhood

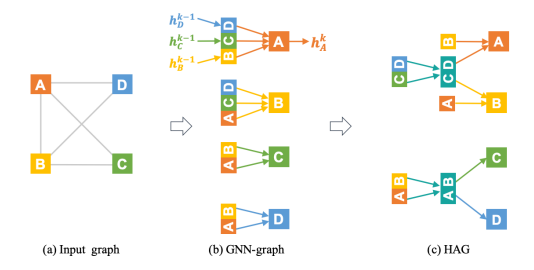


Computational graph



## Issue with Stochastic Training (1)

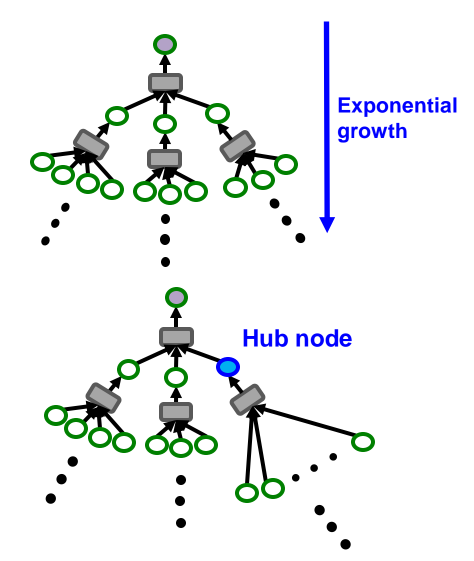
- For each node, we need to get the entire K-hop neighborhood and pass it through the computation graph.
- We need to aggregate lot of information just to compute one node embedding.
- Issue 1: some computational redundancy:



#### Issue with Stochastic Training (2)

#### 2<sup>nd</sup> issue:

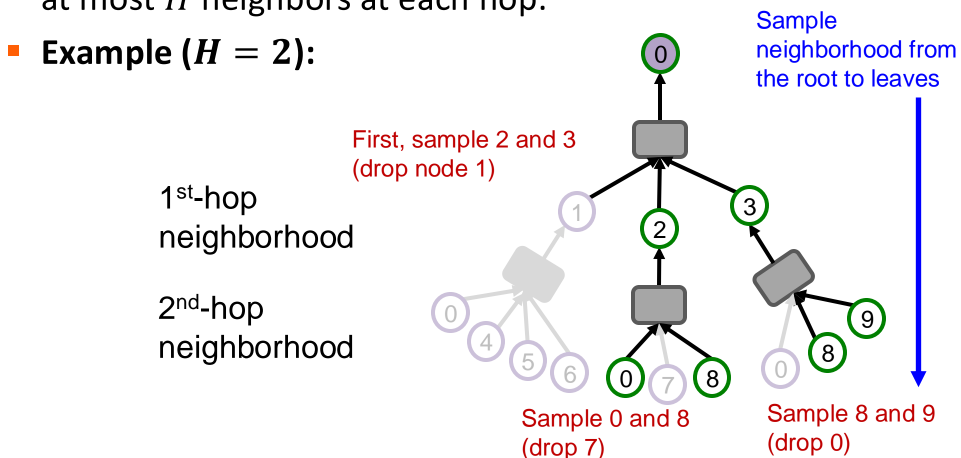
- Computation graph becomes exponentially large with respect to the layer size K.
- Computation graph explodes when it hits a hub node (highdegree node).
- Next: Make the comp. graph more compact!



### Neighborhood Sampling

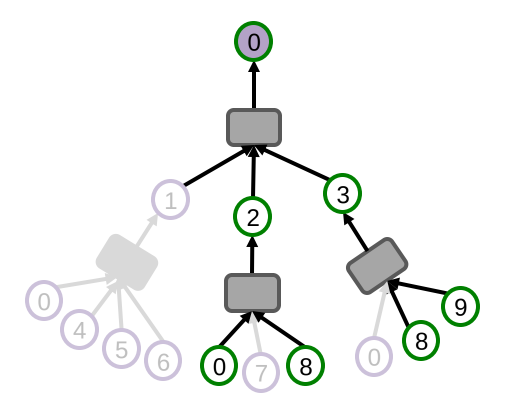
Key idea: Construct the computational graph by (randomly) sampling

at most *H* neighbors at each hop.



## Neighborhood Sampling

 We can use the pruned computational graph to more efficiently compute node embeddings.



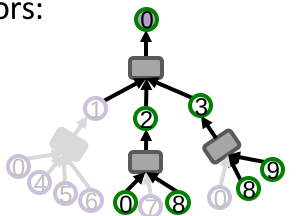
## Neighborhood Sampling Algorithm

#### Neighbor sampling for K-layer GNN

- For k = 1, 2, ..., K:
  - For each node in k-hop neighborhood:

• (Randomly) sample at most  $H_k$  neighbors:

1st-hop<br/>neighborhoodSample  $H_1 = 2$ <br/>neighbors2nd-hop<br/>neighborhoodSample  $H_2 = 2$ <br/>neighbors



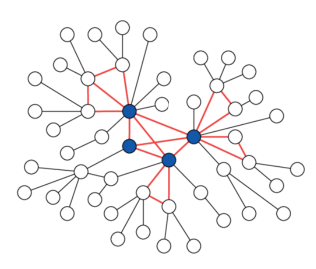
• K-layer GNN will at most involve  $\prod_{k=1}^K H_k$  leaf nodes in comp. graph.

### Remarks on Neighbor Sampling (1)

- Remark 1: Trade-off in sampling number H
  - Smaller H leads to more efficient neighbor aggregation, but results in more unstable training due to the larger variance in neighbor aggregation.
- Remark 2: Computational time
  - Even with neighbor sampling, the size of the computational graph is still exponential with respect to number of GNN layers K.
  - Adding one GNN layer would make computation H times more expensive.

## Remarks on Neighbor Sampling (2)

- Remark 3: How to sample the nodes
  - Random sampling: fast but many times not optimal (may sample many "unimportant" nodes)
  - Random Walk with Restarts:
    - Natural graphs are "scale free", sampling random neighbors, samples many low degree "leaf" nodes.
    - Strategy to sample important nodes:
      - Compute Random Walk with Restarts score  $R_i$  starting at the **green** node
      - At each level sample H neighbors i with the highest  $R_i$
    - This strategy works much better in practice.



### Summary: Neighbor Sampling

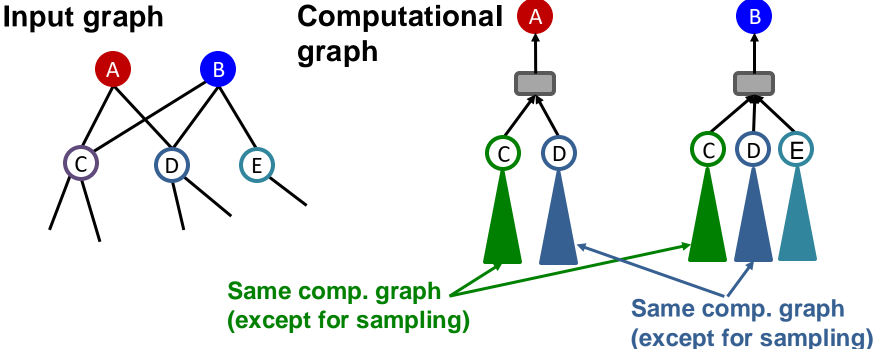
- A computational graph is constructed for each node in a mini-batch.
- In neighbor sampling, the comp. graph is pruned/sub-sampled to increase computational efficiency.
- The pruned comp. graph is used to generate a node embedding.
- However, computational graphs can still become large, especially for GNNs with many message-passing layers.

#### Beyond Small Graphs: Scale GNNs to Large Graphs

## Cluster-GCN: Scaling up GNNs

#### Issues with Neighbor Sampling

- The size of computational graph becomes exponentially large w.r.t. the #GNN layers.
- Computation is redundant, especially when nodes in a mini-batch share many neighbors.

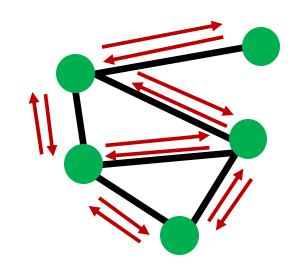


#### Recall: Full Batch GNN

- In full-batch GNN implementation, all the node embeddings are updated together using embeddings of the previous layer.
- Update for all  $v \in V$

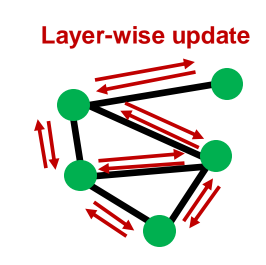
$$h_v^{(\ell)} = COMBINE\left(h_v^{(\ell-1)}, AGGR\left(\left\{\begin{matrix} \boldsymbol{h_u^{(\ell-1)}} \\ \boldsymbol{h_u^{(\ell)}} \end{matrix}\right\}_{u \in N(v)}\right)\right)$$

- In each layer, only 2\*#(edges) messages need to be computed.
- For K-layer GNN, only 2K\*#(edges) messages need to be computed.
- GNN's entire computation is only linear in #(edges) and #(GNN layers). Fast!



#### Insight from Full-batch GNN

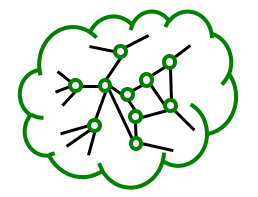
- The layer-wise node embedding update allows the re-use of embeddings from the previous layer.
- This significantly reduces the computational redundancy of neighbor sampling.
  - Of course, the layer-wise update is not feasible for a large graph due to limited GPU memory.
    - Requires putting the entire graph and features on GPU.



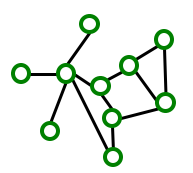
### Subgraph Sampling

Key idea: We can sample a small subgraph of the large graph and then perform the efficient layer-wise node embeddings update over the subgraph.

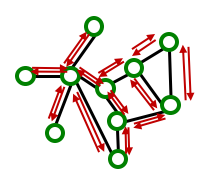
Large graph



Sampled subgraph (small enough to be put on a GPU)



Layer-wise node embeddings update on the GPU



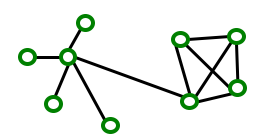
## Subgraph Sampling

- Key question: What subgraphs are good for training GNNs?
  - Recall: GNN performs node embedding by passing messages via the edges.
    - Subgraphs should retain edge connectivity structure of the original graph as much as possible.
    - This way, the GNN over the subgraph generates embeddings closer to the GNN over the original graph.

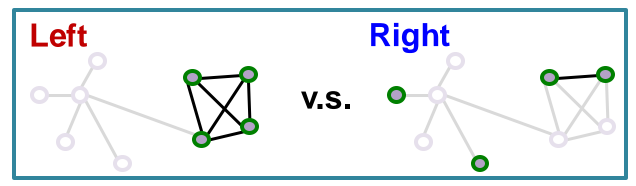
## Subgraph Sampling: Case Study

Which subgraph is good for training GNN?

#### **Original graph**



**Subgraphs** (both 4-node induced subgraph)



- Left subgraph retains the essential community structure among the 4 nodes → Good
- Right subgraph drops many connectivity patterns, even leading to isolated nodes → Bad

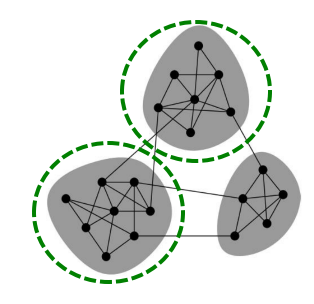
### **Exploiting Community Structure**

Real-world graph exhibits community structure

- A large graph can be decomposed into many small communities.
- Key insight [Chiang et al. KDD 2019]:

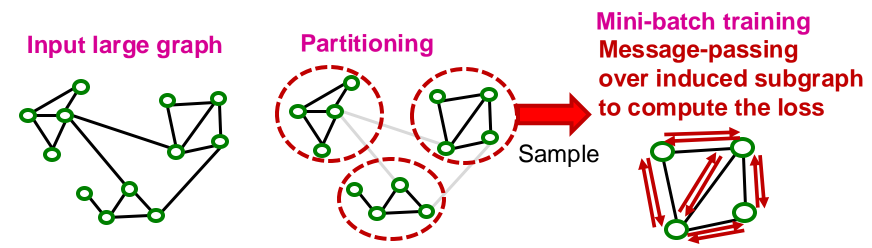
Sample a community as a subgraph.

Each subgraph retains essential local connectivity pattern of the original graph.



#### Cluster-GCN: Overview

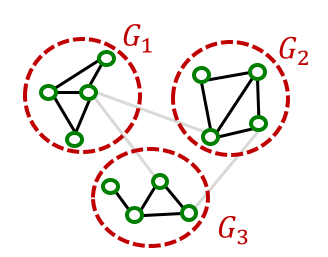
- We first introduce "vanilla" Cluster-GCN.
- Cluster-GCN consists of two steps:
  - Pre-processing: Given a large graph, partition it into groups of nodes (i.e., subgraphs).
  - Mini-batch training: Sample one node group at a time. Apply GNN's message passing over the induced subgraph.



## Cluster-GCN: Pre-processing

- Given a large graph G = (V, E), partition its nodes V into C groups:  $V_1, \dots, V_C$ .
  - We can use any scalable community detection methods, e.g., Louvain, METIS [Karypis et al. SIAM 1998].
- $V_1, \dots, V_C$  induces C subgraphs,  $G_1, \dots, G_C$ ,
  - Recall:  $G_c \equiv (V_c, E_c)$ ,
  - where  $E_c = \{(u, v) | u, v \in V_c\}$

Notice: Between-group edges are *not* included in  $G_1, ..., G_C$ .



### Cluster-GCN: Mini-batch Training

- For each mini-batch, randomly sample a node group  $V_c$ .
- Construct induced subgraph  $G_c = (V_c, E_c)$

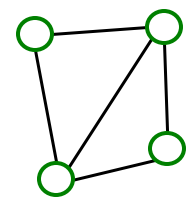
Sampled node group  $V_c$ 

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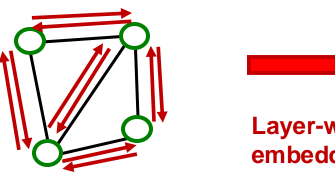
 $\circ$ 

Induced subgraph  $G_c$ 



#### Cluster-GCN: Mini-batch Training

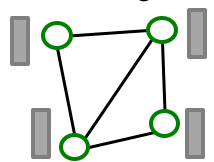
- Apply GNN's layer-wise node update over  $G_c$  to obtain embedding  $h_v$  for each node  $v \in V_c$ .
- Compute the loss for each node  $v \in V_c$  and take average:  $\ell_{sub}(\theta) = (1/|V_c|) \cdot \sum_{v \in V_c} \ell_v(\theta)$
- Update params:  $\theta \leftarrow \theta \nabla \ell_{sub}(\theta)$ Induced subgraph  $G_c$





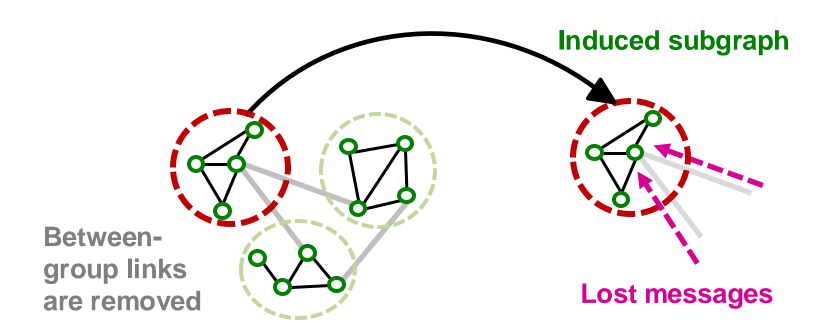
Layer-wise node embedding update

#### **Embedding**



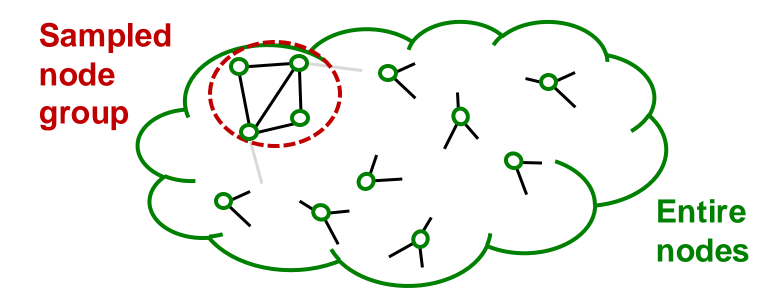
#### Issues with Cluster-GCN (1)

- The induced subgraph removes between-group links.
- As a result, messages from other groups will be lost during message passing, which could hurt the GNN's performance.



#### Issues with Cluster-GCN (2)

- Graph community detection algorithm puts similar nodes together in the same group.
- Sampled node group tends to only cover the small-concentrated portion of the entire data.



#### Issues with Cluster-GCN (3)

# Sampled nodes are not diverse enough to be represent the entire graph structure:

- As a result, the gradient averaged over the sampled nodes,  $\frac{1}{|V_c|}\sum_{v\in V_c}\ell_v(\boldsymbol{\theta})$ , becomes unreliable.
  - Fluctuates a lot from a node group to another.
  - In other words, the gradient has high variance.
- Leads to slow convergence of SGD

#### Advanced Cluster-GCN: Overview

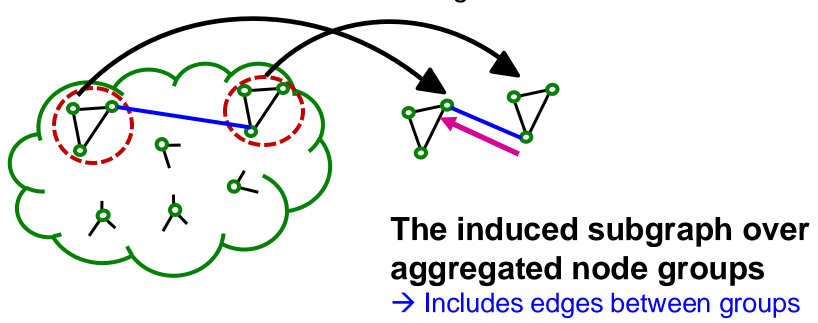
- Solution: Aggregate multiple node groups per mini-batch.
- Partition the graph into relatively-small groups of nodes.
- For each mini-batch:
  - Sample and aggregate multiple node groups.
  - Construct the induced subgraph of the union of node groups.
  - The rest is the same as vanilla Cluster-GCN (compute node embeddings and the loss, update parameters)

#### Advanced Cluster-GCN: Overview

Why does the solution work?

#### Sampling multiple node groups

→ Makes the sampled nodes more representative of the entire nodes. Leads to less variance in gradient estimation.



→ Message can flow across groups.

#### Advanced Cluster-GCN

Similar to vanilla Cluster-GCN, advanced Cluster-GCN also follows 2-step approaches.

#### **Pre-processing step:**

- Given a large graph G = (V, E), partition its nodes V into C relatively-small groups:  $V_1, \dots, V_C$ .
  - $V_1, ..., V_C$  needs to be small so that even if multiple of them are aggregated, the resulting group would not be too large.

#### Advanced Cluster-GCN

#### Mini-batch training:

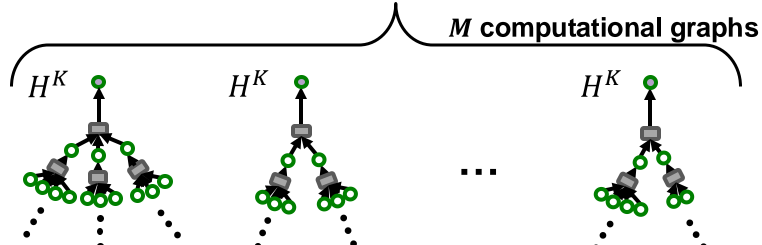
- For each mini-batch, randomly sample a set of q node groups:  $\{V_{t_1}, ..., V_{t_q}\} \subset \{V_1, ..., V_C\}.$
- Aggregate all nodes across the sampled node groups:  $V_{aggr} = V_{t_1} \cup \cdots \cup V_{t_a}$
- Extract the induced subgraph

$$G_{aggr} = (V_{aggr}, E_{aggr}),$$
 where  $E_{aggr} = \{(u, v) \mid u, v \in V_{aggr}\}$ 

•  $E_{aggr}$  also includes between-group edges!

### Comparison of Time Complexity

- Generate M ( $\ll N$ ) node embeddings using K-layer GNN (N: #all nodes).
- Neighbor-sampling (sampling H nodes per layer):
  - For each node, the size of K-layer computational graph is  $H^K$ .
  - For M nodes, the cost is  $M \cdot H^K$



## Comparison of Time Complexity

• Generate M ( $\ll N$ ) node embeddings using K-layer GNN (N: #all nodes).

#### Cluster-GCN:

- Perform message passing over a subgraph induced by the M nodes.
- The subgraph contains  $M \cdot D_{avg}$  edges, where  $D_{avg}$  is the average node degree.
- K-layer message passing over the subgraph costs at most  $K \cdot M \cdot D_{avq}$ .

### Comparison of Time Complexity

- In summary, the cost to generate embeddings for M nodes using K-layer GNN is:
  - Neighbor-sampling (sample H nodes per layer):  $M \cdot H^K$
  - Cluster-GCN:  $K \cdot M \cdot D_{avg}$
- Assume  $H=D_{avg}/2$ . In other words, 50% of neighbors are sampled.
  - Then, Cluster-GCN (cost: 2MHK) is much more efficient than neighbor sampling (cost:  $MH^K$ ).
  - Linear (instead of exponential) dependency w.r.t. K.

#### Cluster-GCN: Summary

- Cluster-GCN first partitions the entire nodes into a set of small node groups.
- At each mini-batch, multiple node groups are sampled, and their nodes are aggregated.
- GNN performs layer-wise node embeddings update over the induced subgraph.
- Cluster-GCN is more computationally efficient than neighbor sampling, especially when #(GNN layers) is large.
- But Cluster-GCN leads to systematically biased gradient estimates (due to missing cross-community edges)

Beyond Small Graphs: Scale GNNs to Large Graphs

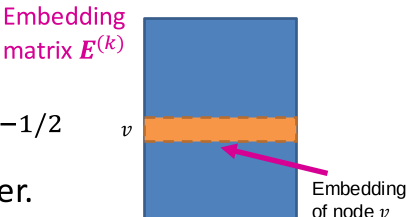
Scaling up by Simplifying GNN Architecture

## Roadmap of Simplifying GCN

- We start from Graph Convolutional Network (GCN) [Kipf & Welling ICLR 2017].
- We simplify GCN ("SimplGCN") by removing the non-linear activation from the GCN [Wu et al. ICML 2019].
  - SimplGCN demonstrated that the performance on benchmark is not much lower by the simplification.
  - Simplified GCN turns out to be extremely scalable by the model design.
  - The simplification strategy is very similar to the one used by LightGCN for recommender systems.

## Quick Overview of LightGCN (1)

- Adjacency matrix: A
- Degree matrix: D
- Normalized adjacency matrix:  $\widetilde{\boldsymbol{A}} \equiv \boldsymbol{D}^{-1/2} \boldsymbol{A} \boldsymbol{D}^{-1/2}$
- Let  $E^{(k)}$  be the embedding matrix at k-th layer.
- Let E be the input embedding matrix.
  - We backprop into E.
- GCN's aggregation in the matrix form
  - $\mathbf{E}^{(k+1)} = \text{ReLU}(\widetilde{\mathbf{A}}\mathbf{E}^{(k)}\mathbf{W}^{(k)})$



## Quick Overview of LightGCN (2)

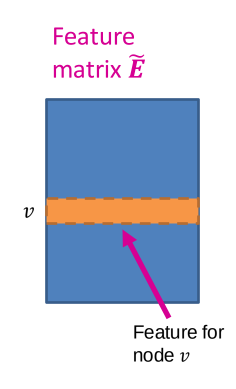
- Removing ReLU non-linearity gives us
  - $E^{(K)} = \widetilde{A}^K E$  W, where  $W \equiv W^{(0)} \cdots W^{(K-1)}$ Diffusing node embeddings along the graph
- Efficient algorithm to obtain  $\widetilde{\pmb{A}}^K \pmb{E}$ 
  - Start from input embedding matrix E.
  - Apply  $\mathbf{E} \leftarrow \widetilde{\mathbf{A}} \; \mathbf{E} \; \text{for } K \; \text{times.}$
- Weight matrix W can be ignored for now.
  - lacktriangledown W acts as a linear classifier over the diffused node embeddings  $m{\widetilde{A}}^K \, m{E}$  .

### Differences to LightGCN

- SimplGCN adds self-loops to adjacency matrix A:
  - $\blacksquare A \leftarrow A + I$
  - Follows the original GCN by Kipf & Welling.
- SimplGCN assumes input node embeddings E to be given as features:
  - Input embedding matrix E is fixed rather than learned.
  - Important consequence:  $\widetilde{A}^K E$  needs to be calculated only once.
    - Can be treated as a pre-processing step.

## Simplified GCN: "SimplGCN"

- Let  $\widetilde{\boldsymbol{E}} = \widetilde{\boldsymbol{A}}^K \boldsymbol{E}$  be pre-processed feature matrix.
  - Each row stores the pre-processed feature for each node.
  - $\widetilde{E}$  can be used as input to any scalable ML models (e.g., linear model, MLP).
- SimplGCN empirically shows learning a linear model over  $\widetilde{E}$  often gives performance comparable to GCN!



#### Comparison with Other Methods

- Compared to neighbor sampling and cluster-GCN, SimplGCN is much more efficient.
  - SimplGCN computes  $\widetilde{E}$  only once at the beginning.
    - The pre-processing (sparse matrix vector product,  $E \leftarrow \widetilde{A} E$ ) can be performed efficiently on CPU.
  - Once  $\widetilde{E}$  is obtained, getting an embedding for node v only takes constant time!
    - Just look up a row for node v in  $\widetilde{\boldsymbol{E}}$ .
    - No need to build a computational graph or sample a subgraph.
- But the model is less expressive (next).

#### Potential Issue of Simplified GCN

 Compared to the original GNN models, SimplGCN's expressive power is limited due to the lack of non-linearity in generating node embeddings.

## Performance of Simplified GCN

Surprisingly, in semi-supervised node classification benchmark,
 SimplGCN works comparably to the original GNNs despite being less expressive.

Why?

#### **Graph Homophily**

- Many node classification tasks exhibit homophily structure, i.e., nodes connected by edges tend to share the same target labels.
- Examples:
  - Paper category classification in paper-citation network
    - Two papers tend to share the same category if one cites another.
  - Movie recommendation for users in social networks
    - Two users tend to like the same movie if they are friends in a social network.

#### When does Simplified GCN Work?

- Recall the preprocessing step of the simplified GCN: Do  $E \leftarrow \widetilde{A} E$  for K times.
  - E is node feature matrix E = X
- Pre-processed features are obtained by iteratively averaging their neighboring node features.

  Average
- As a result, nodes connected by edges tend to have similar pre-processed features.

#### When does Simplified GCN Work?

- Premise: Model uses the pre-processed node features to make prediction.
- Nodes connected by edges tend to get similar pre-processed features.
  - → Nodes connected by edges tend to be predicted the same labels by the model
- Simplified SGC's prediction aligns well with the graph homophily in many node classification benchmark datasets.

#### Simplified GCN: Summary

- Simplified GCN removes non-linearity in GCN and reduces to the simple pre-processing of node features.
- Once the pre-processed features are obtained, scalable mini-batch SGD can be directly applied to optimize the parameters.
- Simplified GCN works surprisingly well in node classification benchmark.
  - The feature pre-processing aligns well with graph homophily in real-world prediction tasks.