# Machine Learning on Graphs



COMP9312\_23T2

# Outline

- Machine Learning on Graphs
- Node Feature Engineering



# **Data Structure & Algorithms**

#### Case studies on Community Detection:

Connected Component, K-Core, K-Truss, Clique, ... Clustering/partition algorithms, ...





# **Learning-based Algorithms**

- It is hard to define a good community.
- It is not hard to judge a community.



Efficiency VS Effectiveness



# **Application**



https://www.datacamp.com/tutorial/comprehensive-introduction-graph-neural-networks-gnns-tutorial



# **Application**

Node classification: Predict a property of a node Example: Categorize online users / items Link prediction: recommendation Example: Knowledge graph completion Graph classification: Categorize different graphs Example: Molecule property prediction Clustering: Detect if nodes form a community Example: Social circle detection Other tasks:

Graph generation: Drug discovery Graph evolution: Physical simulation



![](_page_5_Picture_4.jpeg)

#### **Application: Molecule Generation**

![](_page_6_Figure_1.jpeg)

Use case 1: Generate novel molecules with high drug likeness

Use case 2: Optimize existing molecules to have desirable properties

![](_page_6_Figure_4.jpeg)

# **Application: Drug Discovery**

Antibiotics are small molecular graphs

- Nodes: Atoms
- Edges: Chemical bonds

![](_page_7_Figure_4.jpeg)

Konaklieva, Monika I. "Molecular targets of  $\beta$ -lactam-based antimicrobials: beyond the usual suspects." Antibiotics 3.2 (2014): 128-142.

![](_page_7_Picture_6.jpeg)

# **Application: Drug Side Effects**

Many patients take multiple drugs to treat complex or co-existing diseases:

- 46% of people ages 70-79 take more than 5 drugs
- Many patients take more than 20 drugs to treat heart disease, depression, insomnia, etc.

Task: Given a pair of drugs predict adverse side effects

![](_page_8_Figure_5.jpeg)

# **Application: Google Map**

#### **Predict via Graph Neural Networks**

![](_page_9_Figure_2.jpeg)

![](_page_9_Picture_3.jpeg)

### ML/DL on traditional data

![](_page_10_Figure_1.jpeg)

# Challenges

#### Graphs are complex

- Arbitrary size and complex topological structure (*i.e.*, no spatial locality like grids)
- No fixed node ordering or reference point
- Often dynamic and have multimodal features

![](_page_11_Figure_5.jpeg)

![](_page_11_Picture_6.jpeg)

#### **Graph Neural Networks**

![](_page_12_Figure_1.jpeg)

![](_page_12_Picture_2.jpeg)

### How to get features

#### 1. Feature Engineering

Covered in this topic

2. Graph Representation Learning Optional topic of node embedding

![](_page_13_Figure_4.jpeg)

![](_page_13_Picture_5.jpeg)

# **Different types of graph features**

![](_page_14_Figure_1.jpeg)

![](_page_14_Picture_2.jpeg)

# **Traditional ML on Graphs**

Good features effectively represent the graph structure and achieve good performance.

- 1. Design features for **nodes**/edges/graphs.
- 2. Get features additional features from training data.
- 3. Use features to train parameters.

Testing: predict using the feature of query node/link/graph

![](_page_15_Picture_6.jpeg)

### **Node-Level Features**

#### Goal:

Characterize the structure and position of a node in the network:

![](_page_16_Picture_3.jpeg)

A typical application: node classification

![](_page_16_Picture_5.jpeg)

### **Adjacency** Matrix?

![](_page_17_Figure_1.jpeg)

Not working for big graphs!

![](_page_17_Picture_3.jpeg)

### Adjacency List?

![](_page_18_Figure_1.jpeg)

Feature dimension need to be consistent

![](_page_18_Picture_3.jpeg)

### Adjacency List?

#### How about this?

![](_page_19_Figure_2.jpeg)

v1	4	5	0	0
ν2	4	0	0	0
v3	4	5	0	0
ν4	1	2	3	5
ν5	1	3	4	0

Feature dimension need to be consistent

![](_page_19_Picture_5.jpeg)

### **Node-Level Features: Overview**

- Node degree
- Clustering coefficient
- Graphlets
- Node centrality
- •

![](_page_20_Picture_6.jpeg)

# **Node** Degree

Degree of a node: the number of neighbors. Treat all neighbors equally.

![](_page_21_Picture_2.jpeg)

![](_page_21_Picture_3.jpeg)

# **Node** Centrality: Clustering Coefficient

Measures how connected v's neighboring nodes are:

 $e_{v} = \frac{\#(\text{edges among neighboring nodes})}{\binom{k_{v}}{2}} \in [0,1]$ 

Can be also understand as #triangles/#possible triangles

Ego-network: the induced subgraph of the node and all its neighbors

![](_page_22_Figure_5.jpeg)

![](_page_22_Picture_6.jpeg)

# **Computing Clustering Coefficient**

Can you design an algorithm to compute the clustering coefficient of all nodes in a graph with *n* nodes and *m* edges?

![](_page_23_Picture_2.jpeg)

**Observation:** Clustering coefficient counts the #(triangles) in the ego-network.

![](_page_24_Picture_2.jpeg)

Three triangles in 6 possible triplets

We can generalize the above by counting #(pre-specified subgraphs, i.e., graphlets).

![](_page_24_Picture_5.jpeg)

**Graphlets** are small subgraphs.

We aim to describe network structure around the node based on graphlets.

Analogy:

Degree

counts #(edges) that a node touches.

**Clustering coefficient** 

counts #(triangles) that a node is involved.

**Graphlet Degree Vector (GDV):** 

Graphlet-base features for nodes

**GDV** counts **#(graphlets)** that a node is involved.

![](_page_25_Picture_11.jpeg)

How to represent a node by graphlets?

Let's start by considering (connected) graphlets with three nodes:

![](_page_26_Figure_3.jpeg)

![](_page_26_Picture_4.jpeg)

Choose a specific pattern (wedge)

How many subgraphs containing u that are isomorphic to the pattern?

![](_page_26_Picture_7.jpeg)

How many subgraphs containing *u* that are isomorphic to the pattern?

![](_page_27_Picture_2.jpeg)

#### **11** after removing symmetric cases

![](_page_27_Picture_4.jpeg)

We use **11** as the feature of **u** 

![](_page_27_Picture_6.jpeg)

Move forward by utilizing different types:

![](_page_28_Picture_2.jpeg)

![](_page_28_Figure_3.jpeg)

We use [6, 5] as the feature of u

![](_page_28_Picture_5.jpeg)

Move forward by only considering induced matching instances:

![](_page_29_Figure_2.jpeg)

![](_page_29_Picture_3.jpeg)

Move forward by only considering induced matching instances:

![](_page_30_Figure_2.jpeg)

![](_page_30_Picture_3.jpeg)

Move forward by only considering induced matching instances:

![](_page_31_Picture_2.jpeg)

![](_page_31_Picture_3.jpeg)

Choose a specific pattern (wedge)

#### We use [5, 3] as the feature of u

![](_page_31_Picture_6.jpeg)

Move forward by utilizing all 3-graphlets:

![](_page_32_Figure_2.jpeg)

6 for type 0 5 for type 1 1 for type 2

![](_page_32_Figure_4.jpeg)

There are three types in all 3-graphlets.

We use [5, 3, 1] as the feature of u

Type 0:

![](_page_32_Picture_7.jpeg)

Consider all graphlets with <= 5 nodes

How many node roles in all connected non-isomorphic subgraphs?

![](_page_33_Figure_3.jpeg)

There are **73** different graphlets of up to 5 nodes.

To get the node feature, compute the number of induced matching instances for each role id.

![](_page_33_Picture_6.jpeg)

**Graphlet Degree Vector (GDV)**: A count vector of graphlets rooted at a given node.

Considering graphlets of size 2-5 nodes we get:

 Vector of 73 coordinates is a signature of a node that describes the topology of node's neighborhood

Graphlet degree vector provides a measure of a node's local network topology:

 Comparing vectors of two nodes provides a more detailed measure of local topological similarity than node degrees or clustering coefficient.

Usually, we only compute up to 4 or 5 nodes . . .

![](_page_34_Picture_7.jpeg)

More examples: *GFV(u)* = [2,0,2,1]

![](_page_35_Picture_2.jpeg)

![](_page_35_Picture_3.jpeg)

![](_page_35_Picture_4.jpeg)

### **Node** Centrality

Node degree counts neighbors without capturing their importance. Node **centrality** takes the **node** importance in a graph into account

#### Different ways to model importance:

- PageRank
- Eigenvector centrality
- Betweenness centrality
- Closeness centrality
- many others...

![](_page_36_Figure_8.jpeg)

![](_page_36_Picture_9.jpeg)

# **Node** Centrality: Eigenvector

Motivation

A node is important if surrounded by important neighbors.

We model the centrality of node v as the sum of the centrality of neighbors:

$$c_v = rac{1}{\lambda} \sum_{u \in N(v)} c_u \stackrel{\lambda \text{ is normalization constant}}{( ext{it will turn out to be the largest}}$$

The above equation models centrality in a recursive manner. How do we solve it?

![](_page_37_Picture_6.jpeg)

# **Node Centrality: Eigenvector**

Rewrite the recursive equation in the matrix form.

$$c_{v} = \frac{1}{\lambda} \sum_{u \in N(v)} c_{u} \iff \lambda$$

 $\lambda$  is normalization const (largest eigenvalue of A)

#### $\lambda \boldsymbol{c} = \boldsymbol{A}\boldsymbol{c}$

- A: Adjacency matrix  $A_{uv} = 1$  if  $u \in N(v)$
- *c*: Centrality vector
- λ: Eigenvalue
- We see that centrality c is the eigenvector of A!
- The largest eigenvalue  $\lambda_{max}$  is always positive and unique (by Perron-Frobenius Theorem).
- The eigenvector  $c_{max}$  corresponding to  $\lambda_{max}$  is used for centrality.

Optional

Math Warning

# **Node** Centrality: Betweenness

#### **Betweenness centrality:**

A node is important if it lies on many shortest paths between other nodes.

 $c_{v} = \sum_{s \neq v \neq t} \frac{\#(\text{shortest paths betwen } s \text{ and } t \text{ that contain } v)}{\#(\text{shortest paths between } s \text{ and } t)}$ 

How to identity the bridge node

![](_page_39_Picture_5.jpeg)

#### Node Centrality: Betweenness (cont)

Example:

$$c_{v} = \sum_{s \neq v \neq t} \frac{\#(\text{shortest paths betwen } s \text{ and } t \text{ that contain } v)}{\#(\text{shortest paths between } s \text{ and } t)}$$

 $c_a = c_b = c_e = 0$ 

![](_page_40_Figure_4.jpeg)

$$c_{c} = 3$$

$$(a-\underline{c}-b, \ a-\underline{c}-d, \ a-\underline{c}-d-e)$$

$$c_{d} = 3$$

$$(a-\underline{c}-\underline{d}-e, \ b-\underline{d}-e, \ c-\underline{d}-e)$$

![](_page_40_Picture_6.jpeg)

# **Computing Betweenness Centrality**

Exact solution:

O(nm) for unweighted graphs
O(nm+n²logn) for weighted graphs

https://kops.uni-konstanz.de/server/api/core/bitstreams/420590d1-3010-4eab-a585-6fa3eff46f9e/content

Approximate solution:

Sampling a set of shortest paths...

![](_page_41_Picture_6.jpeg)

### **Node** Centrality: Closeness

#### **Closeness centrality:**

A node is important if it closes to all other nodes.

 $c_{v} = \frac{1}{\sum_{u \neq v} \text{shortest path length between } u \text{ and } v}$ 

![](_page_42_Figure_4.jpeg)

 $c_a = 1/(2 + 1 + 2 + 3) = 1/8$ (a-c-b, a-c, a-c-d, a-c-d-e)

$$c_d = 1/(2 + 1 + 1 + 1) = 1/5$$
  
(d-c-a, d-b, d-c, d-e)

![](_page_42_Picture_7.jpeg)

# **Computing Closeness Centrality**

Can you design an algorithm to compute the closeness centrality of all nodes in a graph with *n* nodes and *m* edges?

![](_page_43_Picture_2.jpeg)

### **Node-Level Feature: Summary**

- Importance-based features:
  - Node degree
  - Different node centrality measures
- Structure-based features:
  - Node degree
  - Clustering coefficient
  - Graphlet count vector

![](_page_44_Picture_8.jpeg)

# **Node-level Feature: Summary**

Importance-based features: capture the importance of a node in a graph

- Node degree:
  - Simply counts the number of neighboring nodes
- Node centrality:
  - Model importance of neighbors in a graph
  - Different modeling choices: eigenvector centrality, betweenness centrality, closeness centrality

Useful for predicting influential nodes in a graph

• **Example:** predicting celebrity users in a social network

![](_page_45_Picture_9.jpeg)

# **Node-level Feature: Summary**

Structure-based features: Capture topological properties of local neighborhood around a node.

- Node degree:
  - Counts the number of neighboring nodes
- Clustering coefficient:
  - Measures how connected neighboring nodes are
- Graphlet degree vector:
  - Counts the occurrences of different graphlets

Useful for predicting a particular role a node plays in a graph:

Example: Predicting protein functionality in a protein-protein interaction network.

![](_page_46_Picture_10.jpeg)

#### Learning Outcome

- Traditional ML Pipeline
  - Hand-crafted feature + ML model
- Hand-crafted node features for graph data
  - Node degree, centrality, clustering coefficient, graphlets

![](_page_47_Picture_5.jpeg)