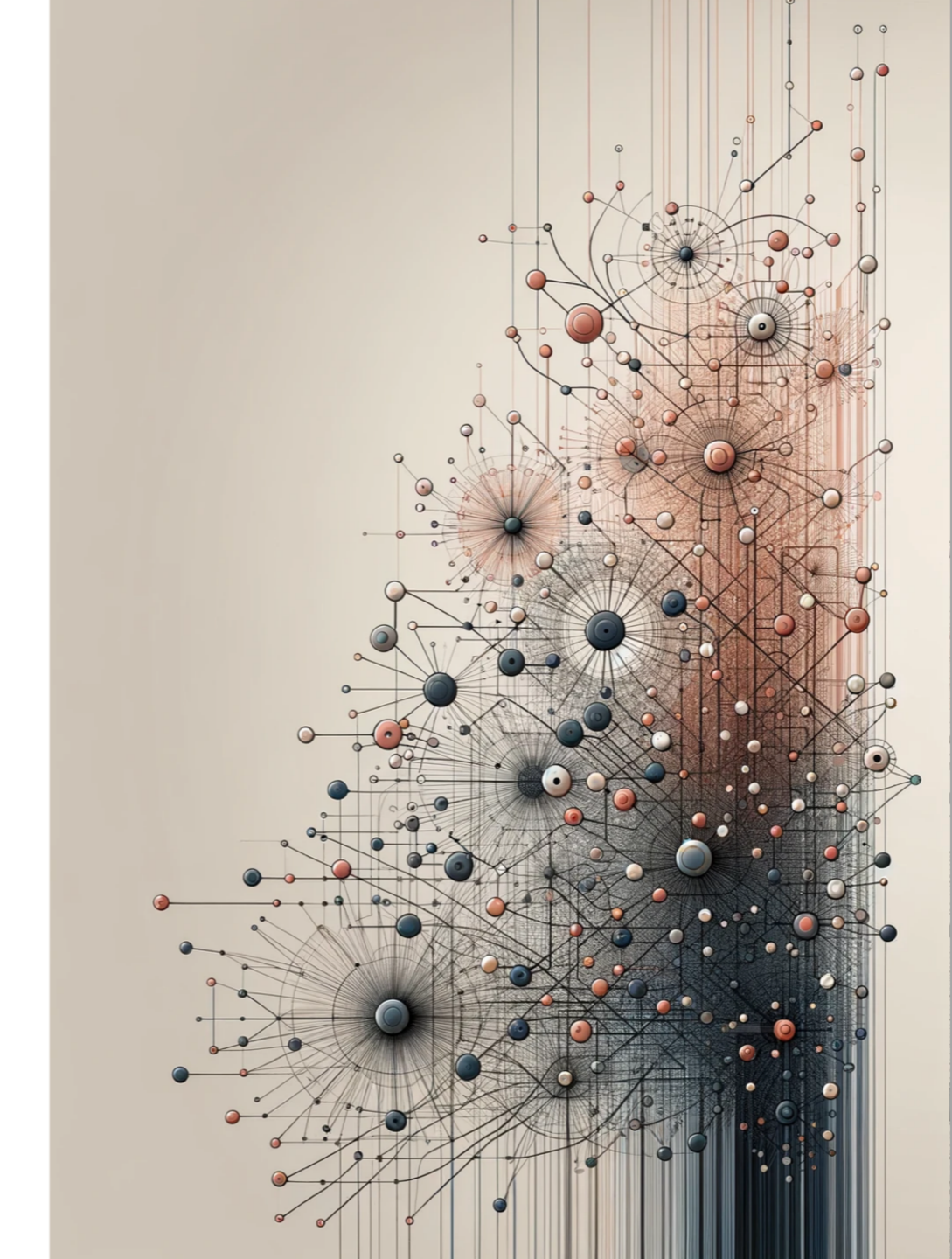


### Analisi e Visualizzazione delle Reti Complesse

NS22 - Traditional Machine Learning on Graphs

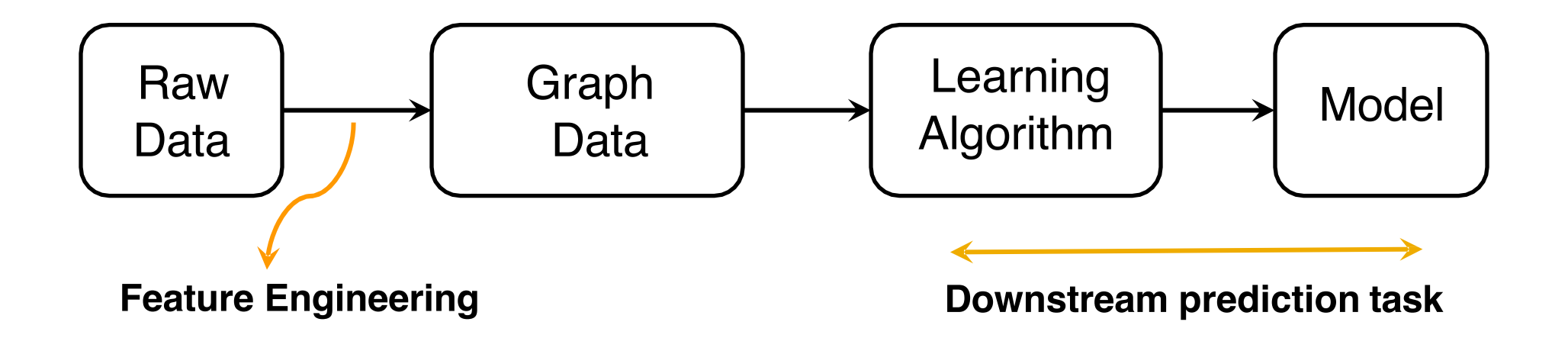
Prof. Rossano Schifanella



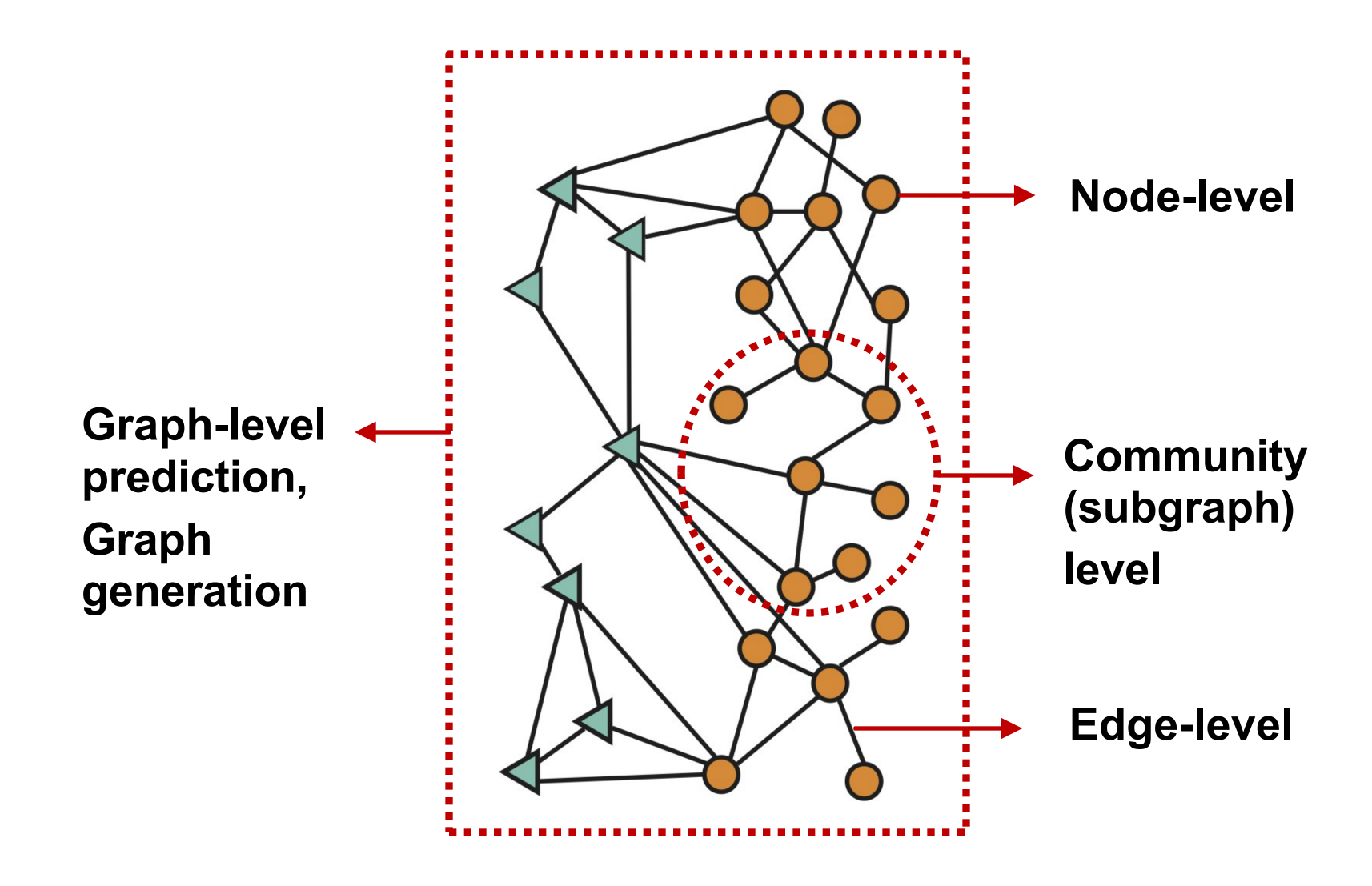
### General Pipeline

### (Supervised) Machine Learning Lifecycle

This feature, that feature. Every single time!



### Different types of tasks

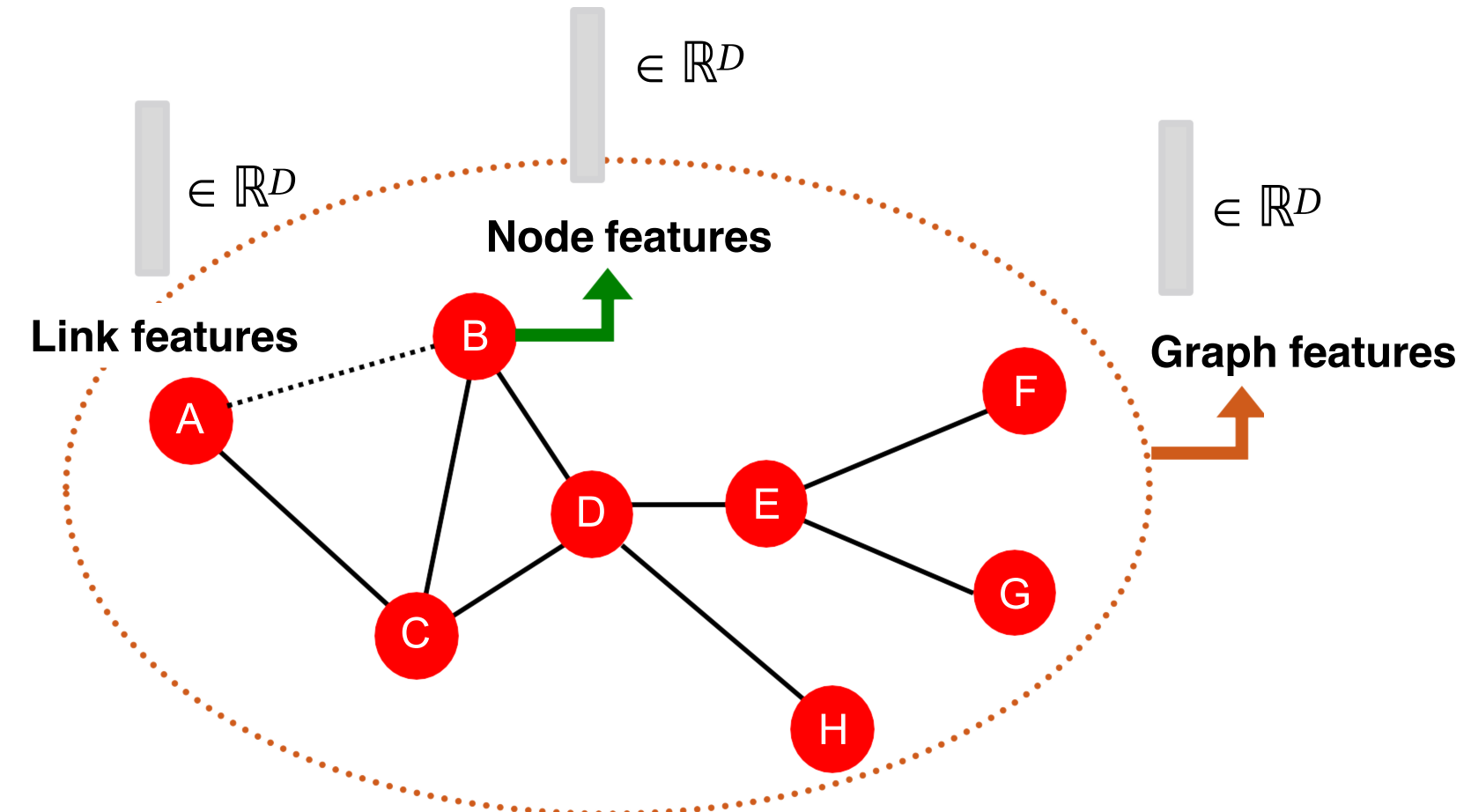


## Classic Graph ML Tasks

- Node classification: predict a property of a node
  - Example: Categorize online users/items
- Link prediction: predict whether there are missing links between two nodes
  - 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

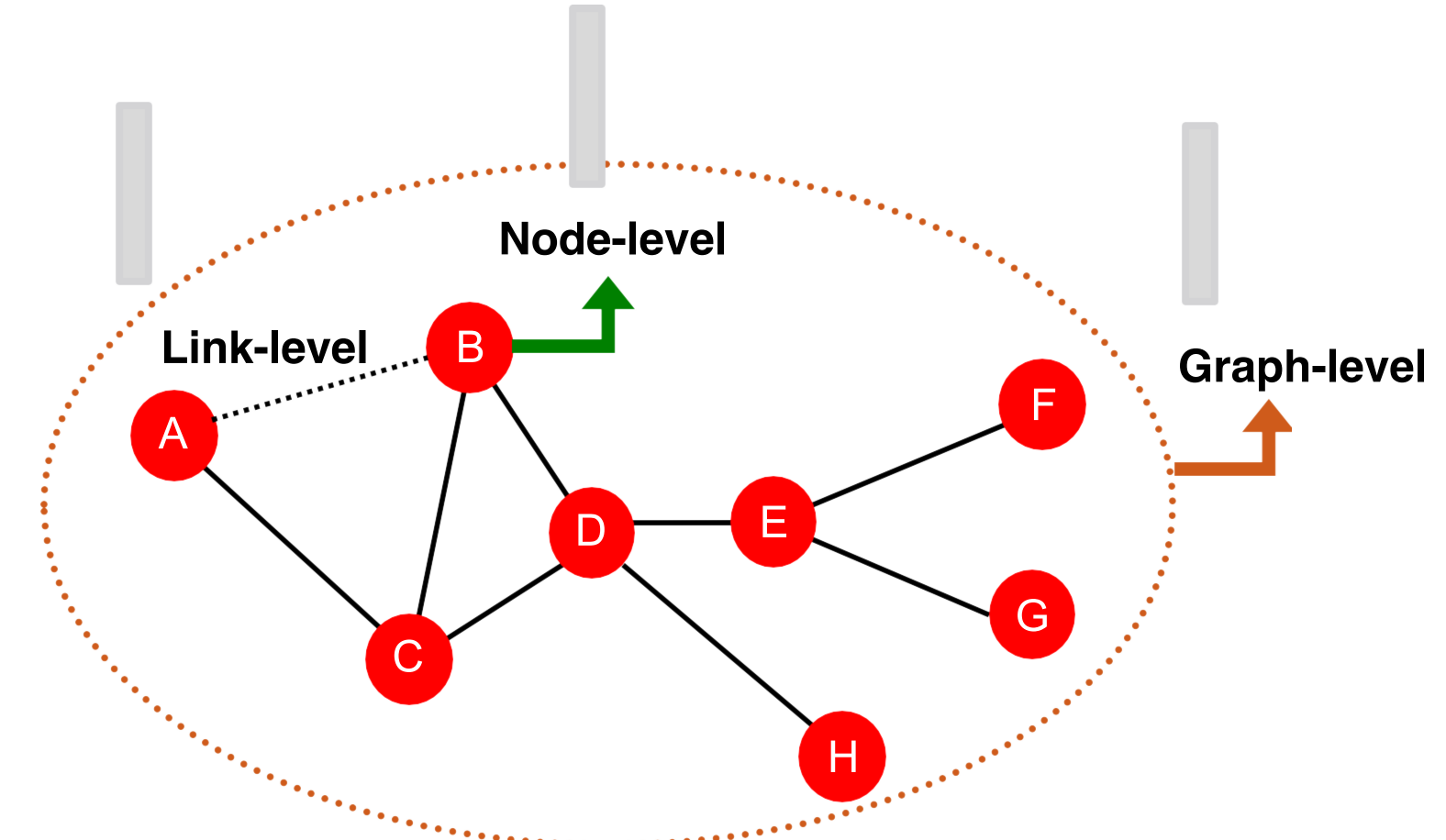
# Traditional LM Pipeline

- Design features for nodes/links/graphs
- Obtain features for all training data



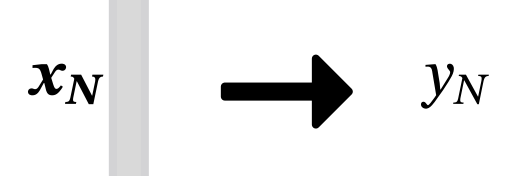
## Machine Learning Tasks: Review

- Node-level prediction
- Link-level prediction
- Graph-level prediction



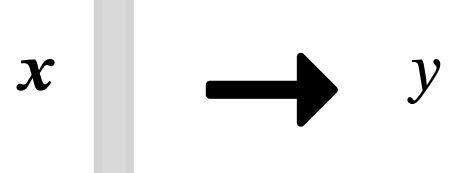
# Traditional LM Pipeline

- Train an ML model:
  - Logistic Regression
  - Random forest
  - Neural network, etc.



### Apply the model:

- Given a new node/link/graph
- Obtain its features
- Make a prediction



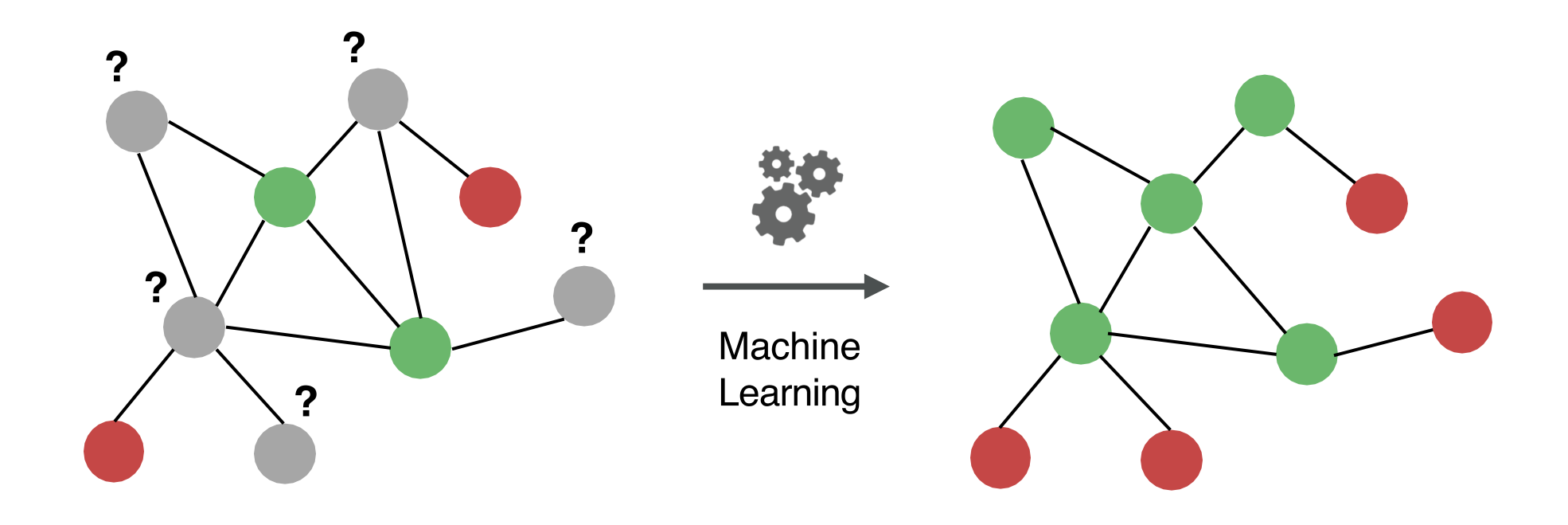
# Features Design

- Using **effective features** *x* over graphs is the key to achieving good model performance.
- Traditional ML pipeline uses hand-designed features.
- In this lecture, we overview the traditional features for:
  - Node-level prediction
  - Link-level prediction
  - Graph-level prediction
- For simplicity, we focus on undirected graphs.

# Machine Learning in Graphs

- Goal: Make predictions for a set of objects
- Design choices:
  - Features: d-dimensional vectors x
  - Objects: Nodes, edges, sets of nodes, entire graphs
  - Objective function:
    - What task are we aiming to solve?

### Node-Level Tasks



Node classification

ML needs features.

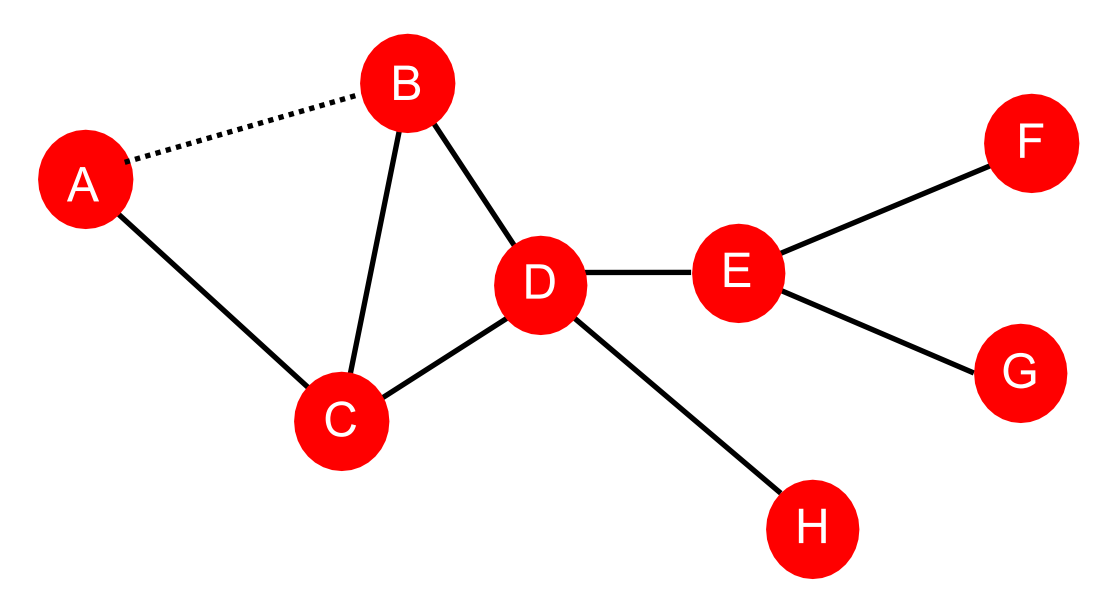
### Node-Level Overview

### Goal

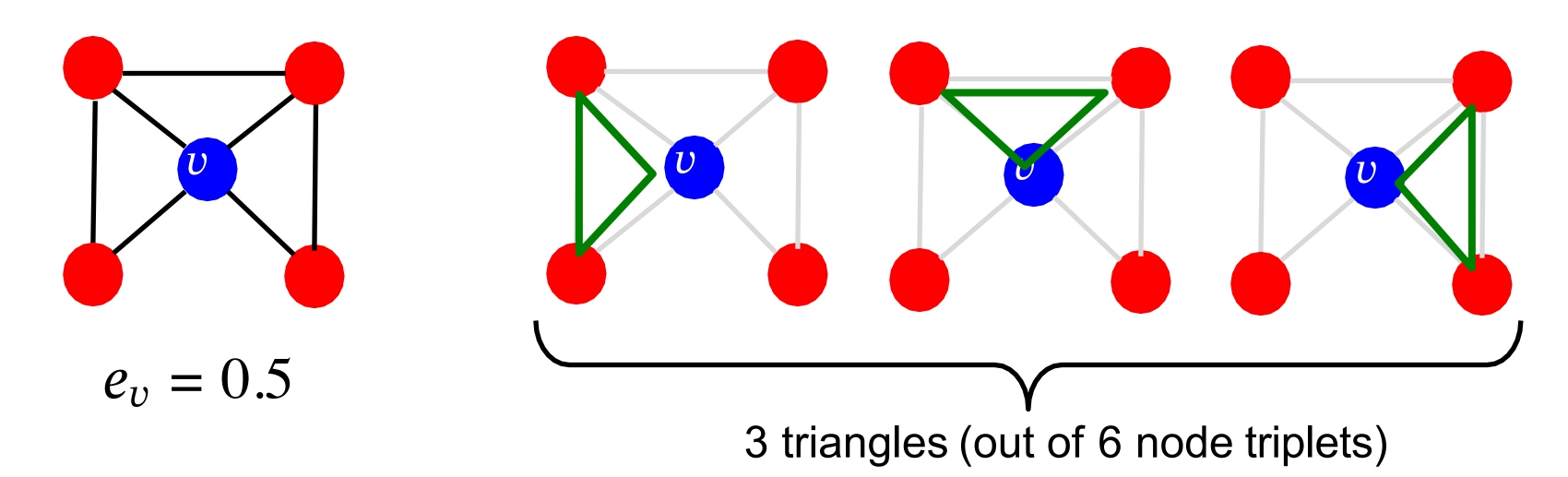
Characterize the structure and position of a node in the network:

- Node degree
- Node centrality
- Clustering coefficient
- Graphlets

#### **Node features**



• Observation: Clustering coefficient counts the #triangles in the ego network.

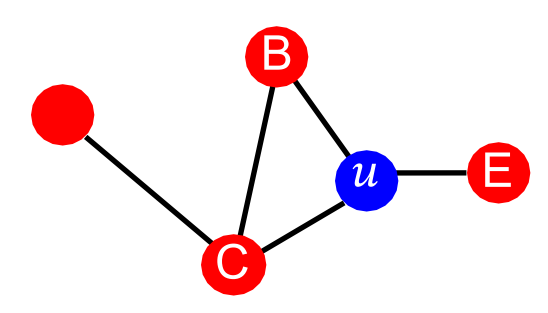


 We can generalize the above by counting #(pre-specified subgraphs, i.e., graphlets)

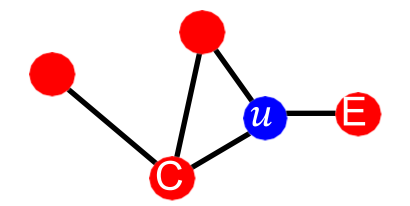
- Goal: Describe the network structure around node u
  - Graphlets are small subgraphs that describe the structure of node *u*'s network neighborhood.

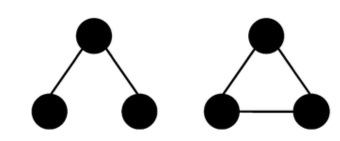
#### **Analogy:**

- Degree counts #(edges) that a node touches
- Clustering coefficient counts #(of triangles)
  that a node touches.
- Graphlet Degree Vector (GDV): Graphletbase features for nodes
  - GDV counts #(graphlets) that a node touches

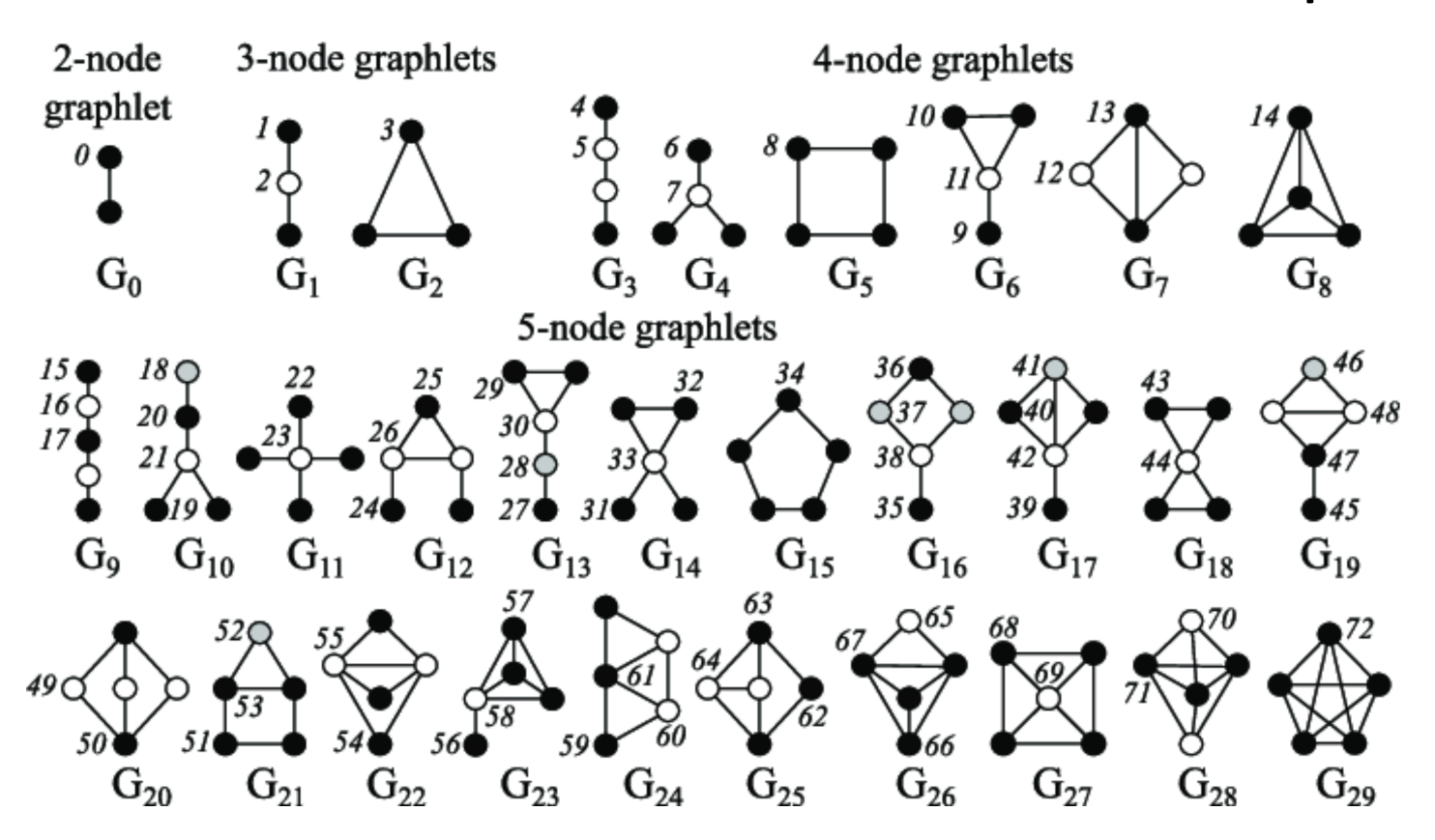


- Considering graphlets of size 2-5 nodes we get:
  - A vector of 73 coordinates is a signature of a node that describes the topology of the node's neighborhood
- Graphlet degree vector provides a measure of a node's local network topology:
  - Comparing vectors of two nodes provides a more detailed measure of local topological similarity than node degrees or clustering coefficient.





Rooted connected induced non-isomorphic subgraphs

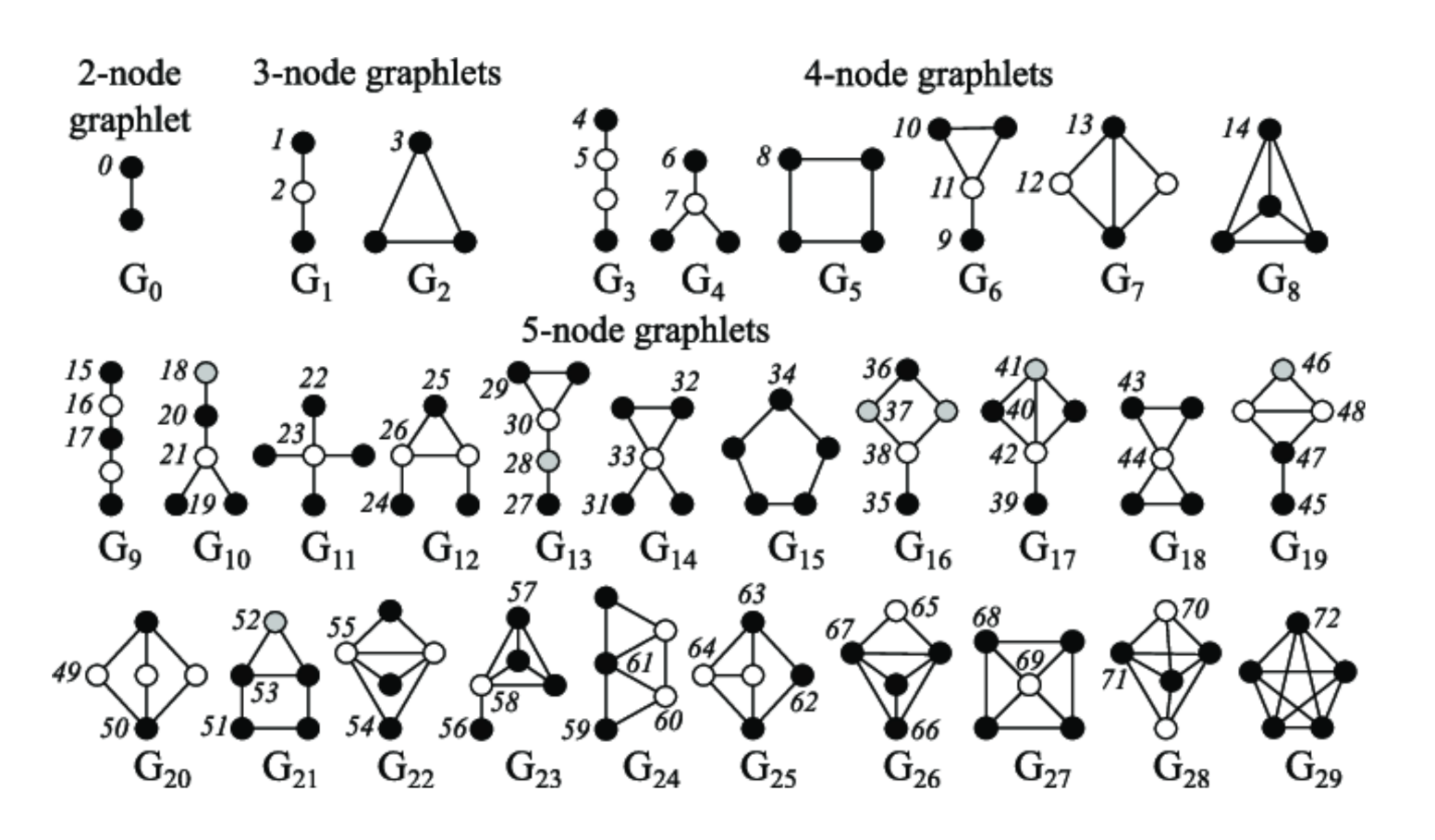


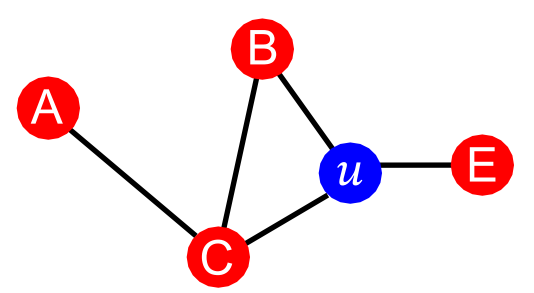
There are 30 different graphlets on up to 5 nodes

There are 73 distinct automorphism orbits (or simply orbits) for graphlets with up to 5 nodes.

Orbits are the distinct roles nodes play within those graphlets based on symmetry.

Rooted connected induced non-isomorphic subgraphs



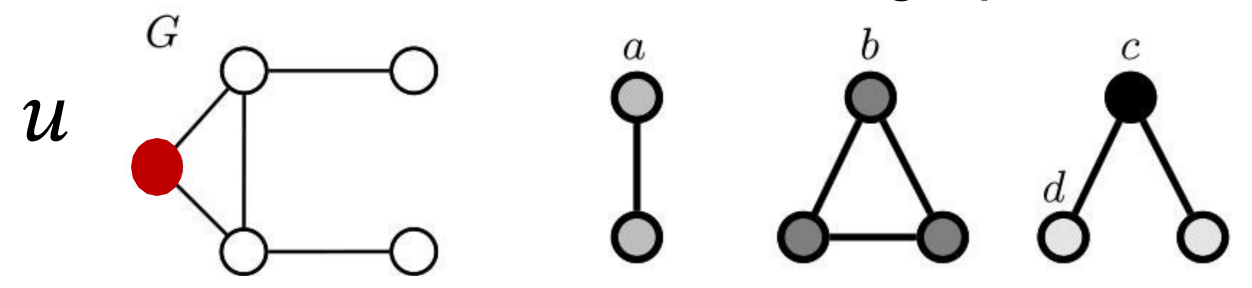


*u* has graphlets:0, 1, 2, 3, 5, 10, 11, ...

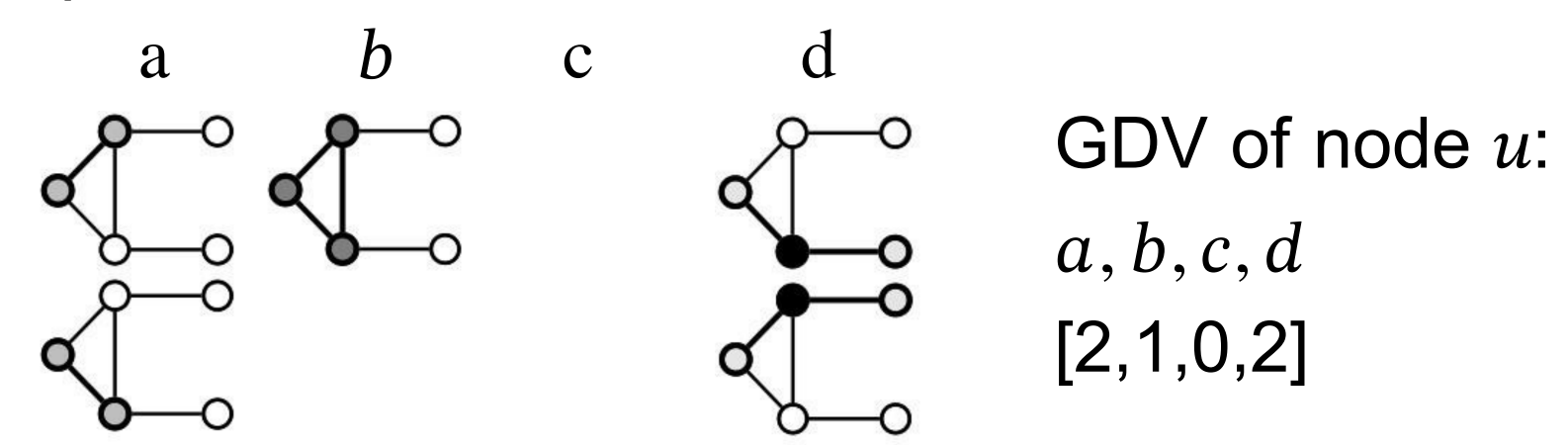
■ Graphlet Degree Vector (GDV): A count vector of graphlets rooted at a given node.

**■ Example:** 

Possible graphlets on up to 3 nodes



Graphlet instances of node u:



## Node-Level Features: Summary

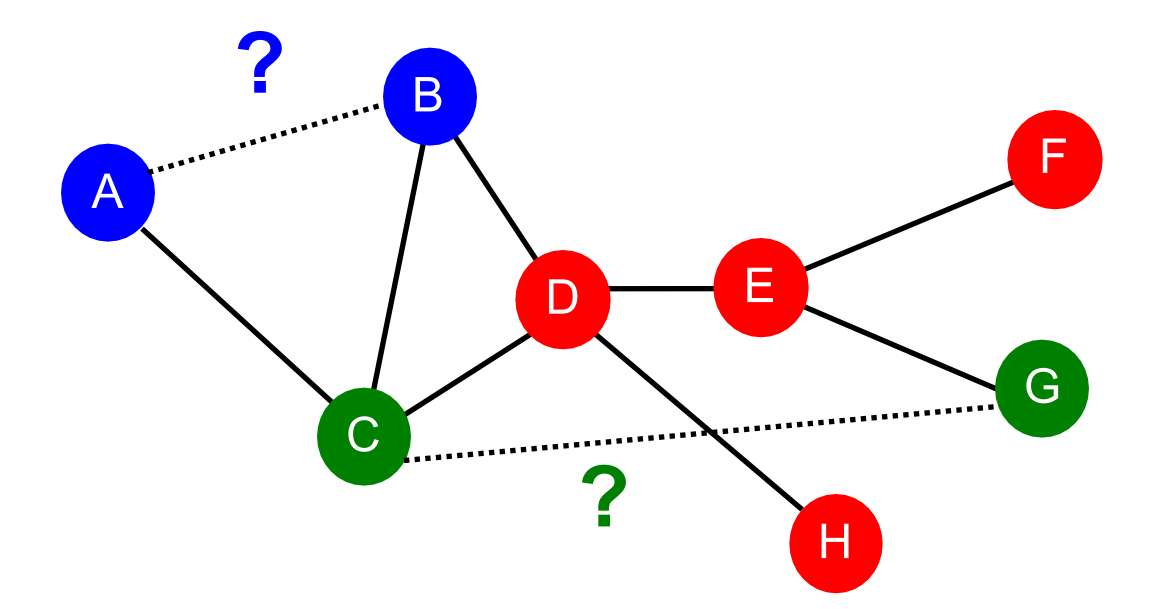
- Importance-based features: capture the importance of a node in a graph
  - Node degree:
    - Counts the number of neighboring nodes
  - Node centrality:
    - Models the importance of neighboring nodes in a graph
    - Different modeling choices: eigenvector centrality, betweenness centrality, closeness centrality
- Useful for predicting influential nodes in a graph
  - Example: predicting celebrity users in a social network

## Node-Level Features: Summary

- Structure-based features: capture topological properties of the 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.

### Link-Level Prediction Task: Recap

- The task is to predict new links based on the existing links.
- At test time, node pairs (with no existing links) are ranked, and top K node pairs are predicted.
- The key is to design features for a pair of nodes.



### Link Prediction as a Task

#### Two formulations of the link prediction task:

### • Links missing at random:

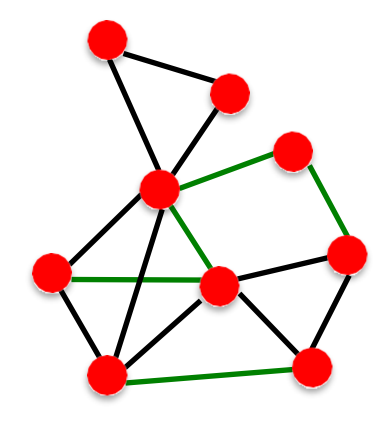
Remove a random set of links and then aim to predict them

#### Links over time:

• Given  $G[t_0,t_0']$  a graph defined by edges up to time  $t_0'$ , output a ranked list L of edges (not in  $G[t_0,t_0']$ ) that are predicted to appear in time  $G[t_1,t_1']$ 

#### Evaluation:

- $n=|E_{new}|$  : # new edges that appear during the test period  $G[t_1,t_1']$
- Take the top n elements of L and count the correct edges

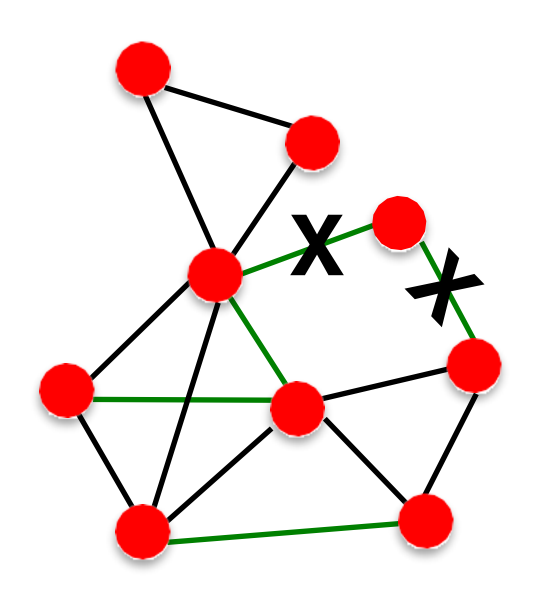


 $G[t_0, t'_0]$   $G[t_1, t'_1]$ 

# Link Prediction via Proximity

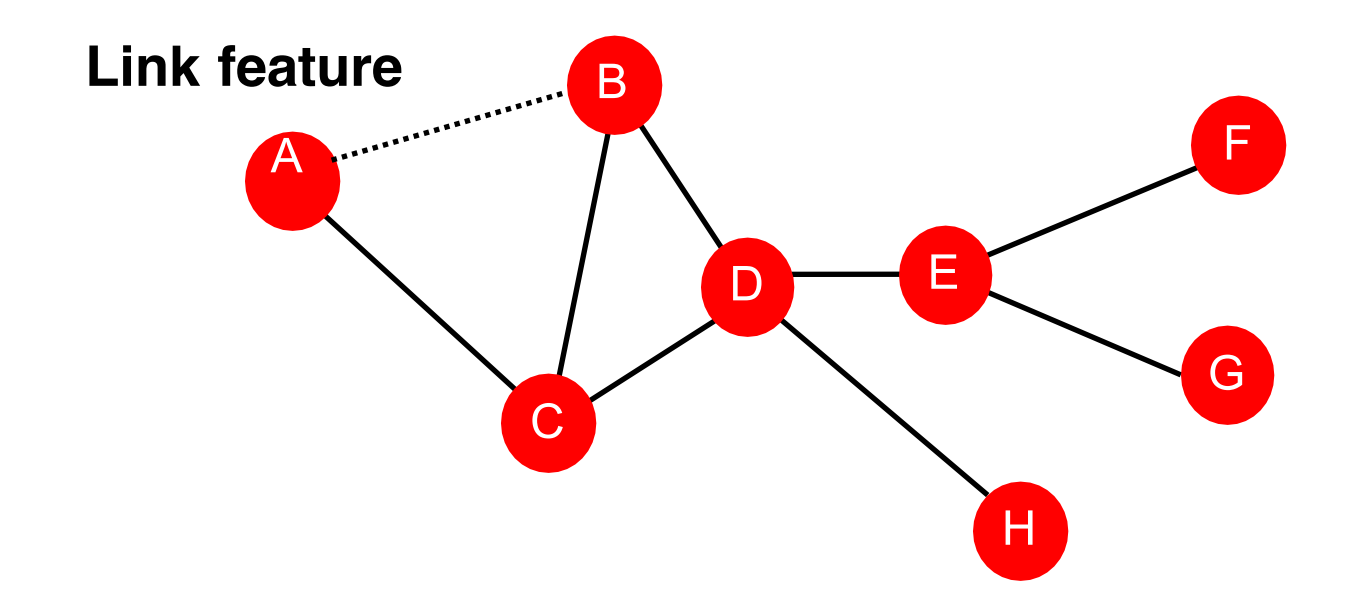
### Methodology:

- For each pair of nodes (x,y) compute score c(x,y)
  - For example, c(x,y) could be the # of common neighbors of x and y
- Sort pairs (x,y) by the decreasing score c(x,y)
- Predict top n pairs as new links
- See which of these links appear in  $G[t_1,t_1^\prime]$



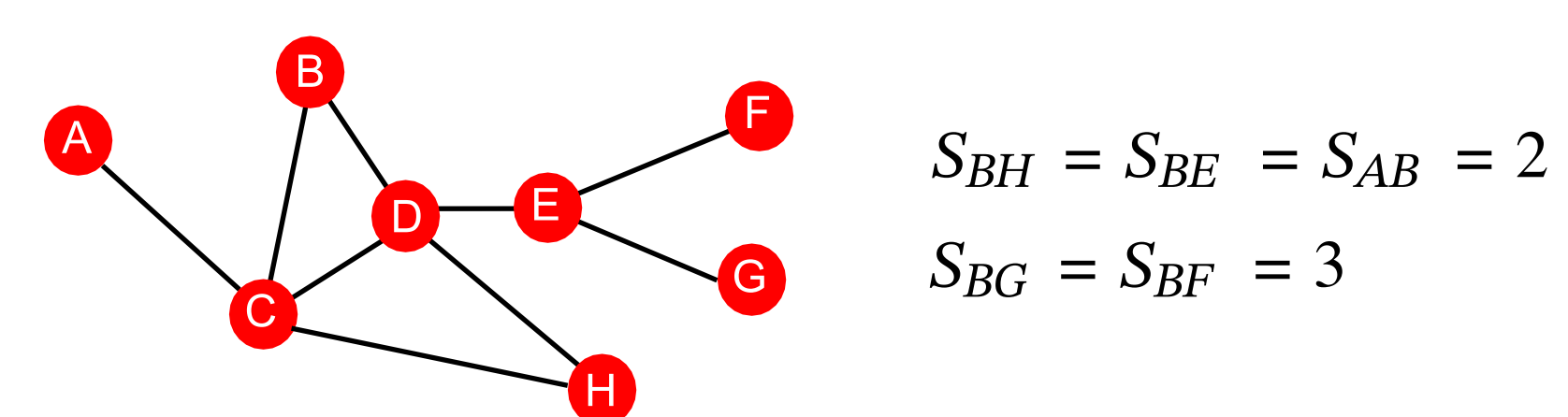
### Link-Level Features: Overview

- Distance-based feature
- Local neighborhood overlap
- Global neighborhood overlap



### Distance-based Features

- Shortest-path distance between two nodes
- However, this does not capture the degree of neighborhood overlap:
  - Node pair (B, H) has 2 shared neighboring nodes, while pairs (B, E) and (A, B) only have 1 such node.
- Example:



### Local Neighborhood Overlap

### Captures # neighboring nodes shared between two nodes v<sub>1</sub> and v<sub>2</sub>

Common neighbors  $|N(v_1) \cap N(v_2)|$ 

$$|N(v_1) \cap N(v_2)|$$

**Example:** 

$$|N(A) \cap N(B)| = |\{C\}| = 1$$

- Jaccard's coefficient
  - **Example:**

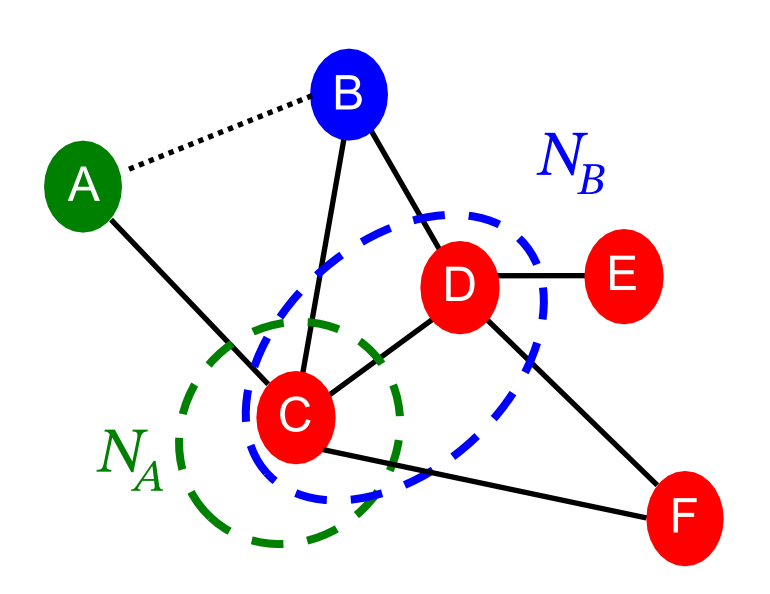
$$\frac{|N(v_1) \cap N(v_2)|}{|N(v_1) \cup N(v_2)|}$$

$$\frac{|N(A) \cap N(B)|}{|N(A) \cup N(B)|} = \frac{|\{C\}|}{|\{C,D\}|} = \frac{1}{2}$$

- Adamic-Adar index
  - **Example:**

$$\frac{1}{\log(k_C)} = \frac{1}{\log 4}$$

$$\sum_{u \in N(v_1) \cap N(v_2)} \frac{1}{\log(k_u)}$$



# Key differences

### Common Neighbors:

- Simple absolute count of shared neighbors.
- Higher count = higher similarity, but doesn't consider node size/graph density.

#### Jaccard Index:

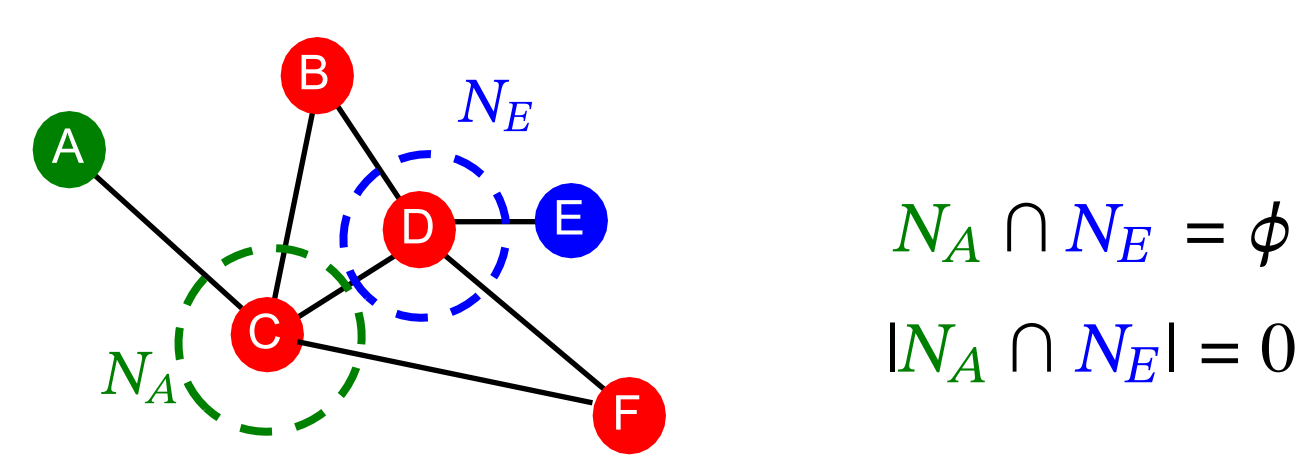
- Normalizes shared count by the total unique neighbors.
- Gives a relative proportion (0 to 1), better for comparing nodes of different sizes.

#### Adamic-Adar Index:

- Weights shared neighbors by the inverse of their degree.
- More weight given to common neighbors who are less connected themselves (rare).

### Global Neighborhood Overlap

- Limitations of local neighborhood features:
  - The metric is always zero if the two nodes do not have any neighbors in common.
  - However, the two nodes may still potentially be connected in the future.
- Global neighborhood overlap metrics resolve the limitation by considering the entire graph.



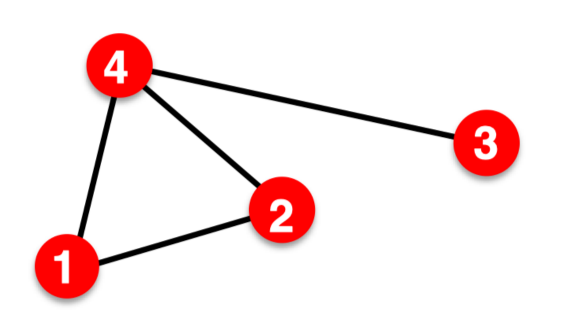
## Global Neighborhood Overlap

- Katz index: count the number of walks of all lengths between a given pair of nodes.
- Question: How to compute #walks between two nodes?

Use powers of the graph adjacency matrix!

### Intuition: Powers of Adj Matrices

- Computing #walks between two nodes
  - Recall:  $A_{uv} = 1$  if  $u \in N(v)$
  - Let  $P_{uv}^{(K)} = \text{#walks of length K between u and v}$
  - We will show  $P^{(K)} = A^k$
  - $P_{uv}^{(1)}$  = #walks of length 1 (direct neighborhood) between u and v =  $A_{uv}$



$$P_{12}^{(1)} = A_{12}$$

$$A = \begin{pmatrix} 0 & 1 & 0 & 1 \\ 1 & 0 & 0 & 1 \\ 0 & 0 & 0 & 1 \\ 1 & 1 & 1 & 0 \end{pmatrix}$$

## Intuition: Powers of Adj Matrices

- How to compute  $P_{uv}^{(2)}$ ?
- Step 1: compute #walks of length 1 between each of u's neighbors and v
- Step 2: Sum up these #walks across u's neighbors

$$P_{uv}^{(2)} = \sum_{i} A_{ui} * P_{iv}^{1} = \sum_{i} A_{ui} * A_{iv} = A_{uv}^{2}$$

Node 1's neighbors

#walks of length 1 between Node 1's neighbors and Node 2

$$A^2 = \begin{pmatrix} 0 & 1 & 0 & 1 \\ 1 & 0 & 0 & 1 \\ 0 & 0 & 0 & 1 \\ 1 & 1 & 1 & 0 \end{pmatrix} \times \begin{pmatrix} 0 & 1 & 0 & 1 \\ 1 & 0 & 0 & 1 \\ 0 & 0 & 0 & 1 \\ 1 & 1 & 1 & 0 \end{pmatrix} = \begin{pmatrix} 2 & 1 & 1 & 1 \\ 1 & 2 & 1 & 1 \\ 1 & 1 & 1 & 0 \\ 1 & 1 & 0 & 3 \end{pmatrix}$$
 adjacency

## Global Neighborhood Overlap

- Katz index: count the number of walks of all lengths between a pair of nodes.
- How to compute #walks between two nodes?
- Use adjacency matrix powers!
  - $A_{uv}$  specifies #walks of length 1 (direct neighborhood) between u and v
  - $A_{uv}^2$  specifies #walks of length 2 between u and v
  - •
  - $A_{uv}^{\,l}$  specifies #walks of length l between u and v

# Global Neighborhood Overlap

• Katz index between  $v_1$  and  $v_2$  is calculated as the sum over all walks lengths:

$$S_{v_1v_2} = \sum_{l=1}^{\infty} \beta^l A_{v_1v_2}^l$$

- $0 < \beta < 1$ : attenuation factor (longer paths counts less)
- Katz index matrix is computed in closed form:

$$S = \sum_{i=1}^{\infty} \beta^i A^i = (I - \beta A)^{-1} - I$$
 by geometric series of matrices

# Link-Level Features: Summary

#### Distance-based features:

• It uses the shortest path length between two nodes but does not capture how the neighborhood overlaps.

### Local neighborhood overlap:

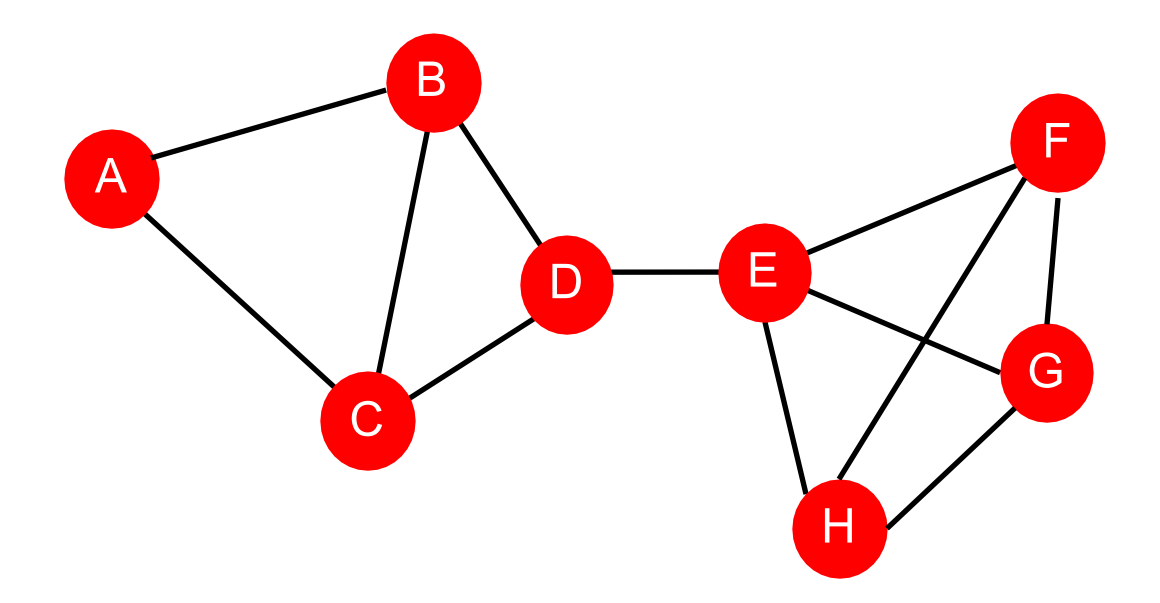
- Captures how many neighboring nodes are shared by two nodes.
- Becomes zero when no neighbor nodes are shared.

### Global neighborhood overlap:

- It uses a global graph structure to score two nodes.
- Katz index counts #walks of all lengths between two nodes.

### Graph-Level Features

- Goal: We want features that characterize the structure of an entire graph.
- For example:



### Background: Kernel Methods

- Goal: We want to use machine learning (ML) to make predictions about entire graphs (e.g., classify molecules, detect communities).
  - Standard ML Approach: ML models usually work with feature vectors lists of numbers describing each data point.
  - The Problem with Graphs: How do you represent a complex graph as a simple list of numbers (a feature vector)?
  - Designing good, informative features by hand that capture the graph's structure can be very difficult and time-consuming.
- The Kernel Idea: What if, instead of defining features first, we directly define a way to measure similarity between two graphs? This similarity measure is called a **kernel**.

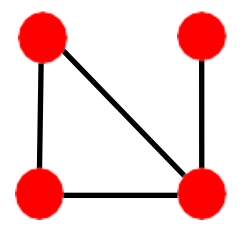
### Graph-Level Features: Overview

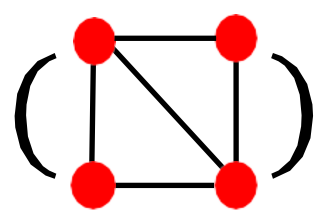
### **Graph Kernels**:

- Measure the similarity between two graphs:
  - Graphlet Kernel [1]
  - Weisfeiler-Lehman Kernel [2]
  - Other kernels are also proposed in the literature
  - (beyond the scope of this lecture)
    - Random-walk kernel
    - Shortest-path graph kernel
    - And many more...
- 1. Shervashidze, Nino, et al. "Efficient graphlet kernels for large graph comparison." Artificial Intelligence and Statistics. 2009.
- 2. Shervashidze, Nino, et al. "Weisfeiler-lehman graph kernels." Journal of Machine Learning Research 12.9 (2011).

# Graph Kernel: Key Idea

- Goal: Design graph feature vector  $\Phi(G)$
- Key idea: <u>Bag-of-Words</u> (BoW) for a graph
  - Recall: BoW uses the word counts as document features (no ordering considered).
  - Naïve extension to a graph: regard nodes as words.
  - Since both graphs have 4 red nodes, we get the same feature vector for two different graphs...





## Graph Kernel: Key Idea

What if we use **Bag of node degrees**?

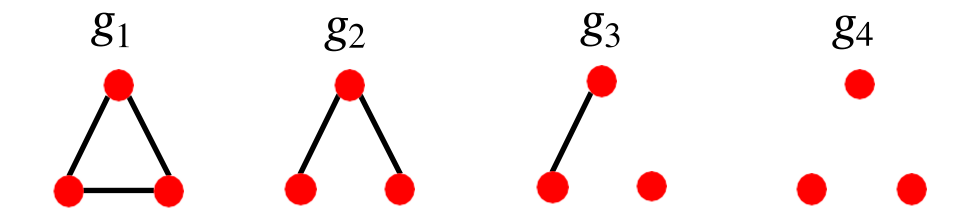
Deg1: Deg2: Deg3: 
$$\phi(1) = \text{count}(1) = [1, 2, 1]$$
Obtains different features for different graphs! 
$$\phi(1) = \text{count}(1) = [0, 2, 2]$$

Both Graphlet Kernel and Weisfeiler-Lehman (WL) Kernel use Bagof-\* representation of graph, where \* is more sophisticated than node degrees!

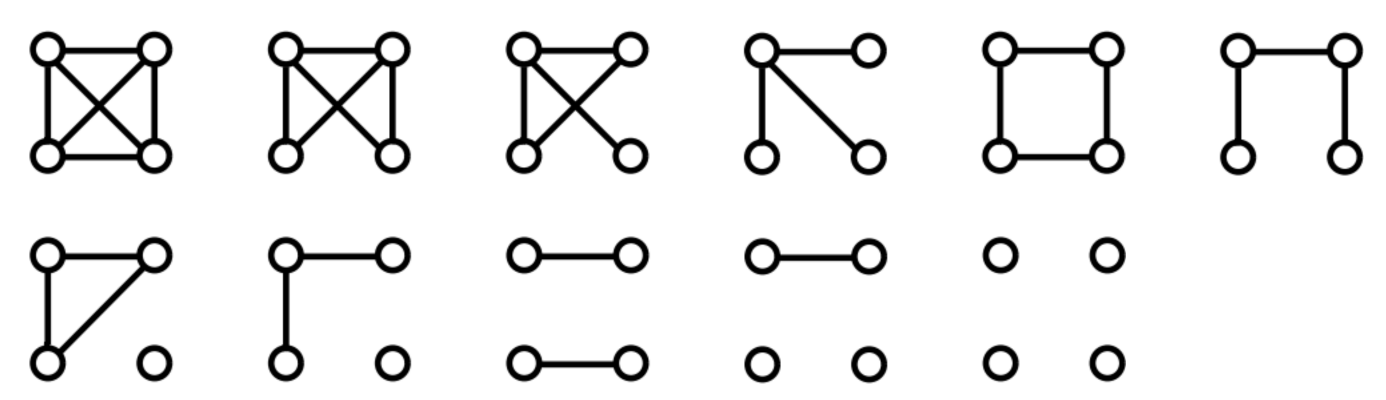
- Key idea: Count the number of different graphlets in a graph.
  - Note: The definition of graphlets here differs slightly from the node-level features.
  - The two differences are:
    - •Nodes in graphlets here do not need to be connected (allows for isolated nodes)
    - The graphlets here are not rooted.
    - Examples in the next slide illustrate this.

Let  $gk = (g1, g2, ..., gn_k)$  be a list of graphlets of size k.

• For k = 3, there are 4 graphlets.



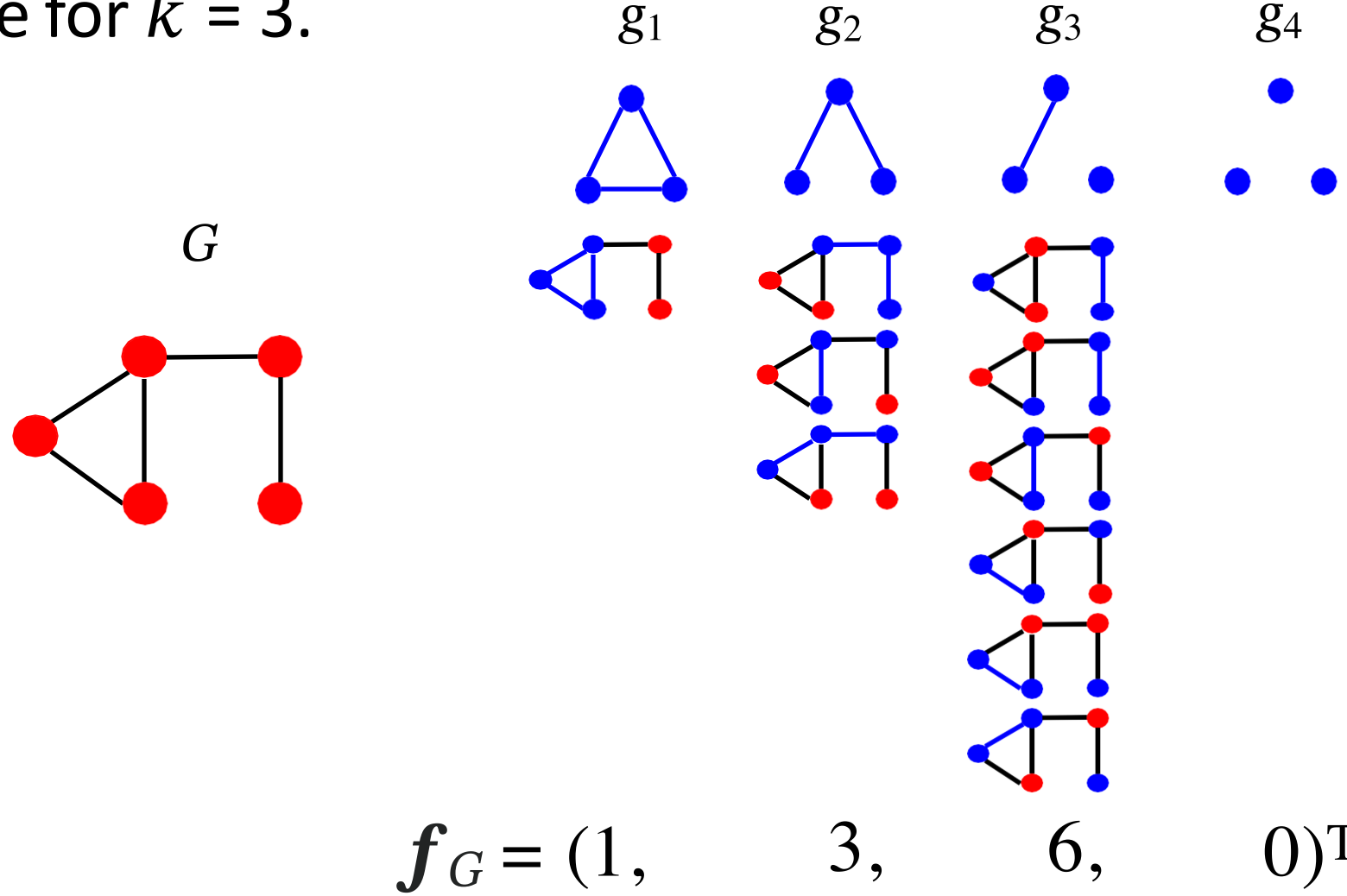
• For k = 4, there are 11 graphlets.



Given graph G and a graphlet list  $G_k = (g_1, g_2, ..., g_{n_k})$  define the graphlet count vector  $f_G \in \mathbb{R}^{n_k}$  as

$$f(G)_i = \#(g_i \subseteq G) \text{ for } i = 1, 2, ..., n_k$$

Example for k = 3.



## Graph-Level Graphlet Kernel

Given two graphs, G and G', graphlet kernel is computed as the dot product

$$K(G, G') = f(G)^T f(G')$$

**Problem:** if G and G' have different sizes, that will greatly skew the value.

Solution: normalize each feature vector

$$h(G) = \frac{f_G}{sum(f_G)}$$

# The Graphlet Kernel

- Limitations: Counting graphlets is expensive!
- Counting size-k graphlets for a graph with size n by enumeration takes  $n^k$ .
- This is unavoidable in the worst case since the subgraph isomorphism test (judging whether a graph is a subgraph of another graph) is **NP-hard**.
- If a graph's node degree is bounded by d, an  $O(nd^{k-1})$  algorithm exists to count all the graphlets of size k.
- Can we design a more efficient graph kernel?

### Weisfeiler-Lehman Kernel

- Goal: Design an efficient graph feature descriptor  $\Phi(G)$
- <u>Idea</u>: Use neighborhood structure to enrich node vocabulary iteratively.
  - Generalized version of Bag of node degrees since node degrees are one-hop neighborhood information.
- Algorithm to achieve this:
  - Color refinement

#### Color Refinement

**Given:** A graph *G* with a set of nodes *V*.

- Assign an initial color  $c^{(0)}(v)$  to each node v.
- Iteratively refine node colors by

$$c^{k+1}(v) = HASH(c^{(k)}(v), \{c^{(k)}(u)\}_{u \in N(v)})$$

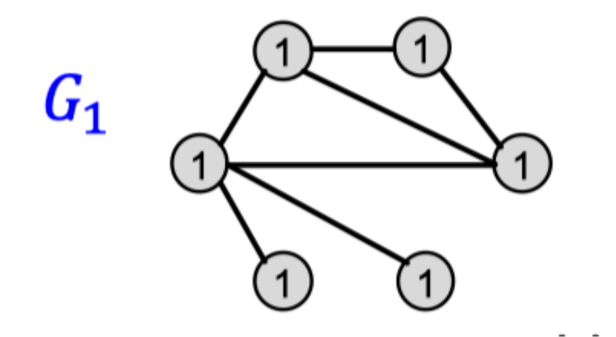
where HASH maps different inputs to different colors

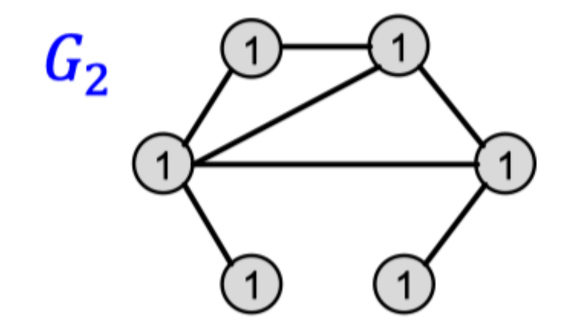
- After k steps of color refinement,  $c^{(K)}(v)$  summarizes the structure of the K-hop neighborhood

### Color Refinement (1)

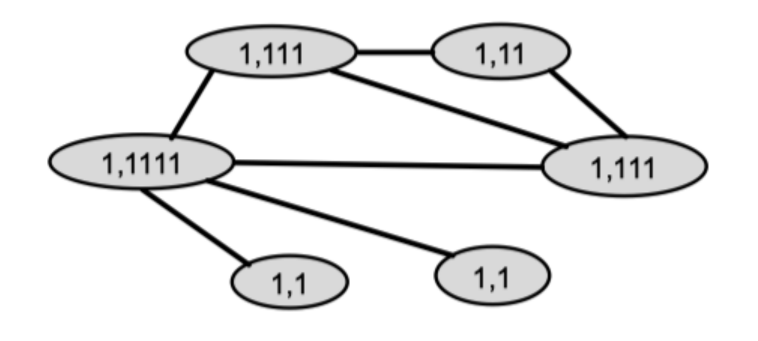
#### Example of color refinement given two graphs

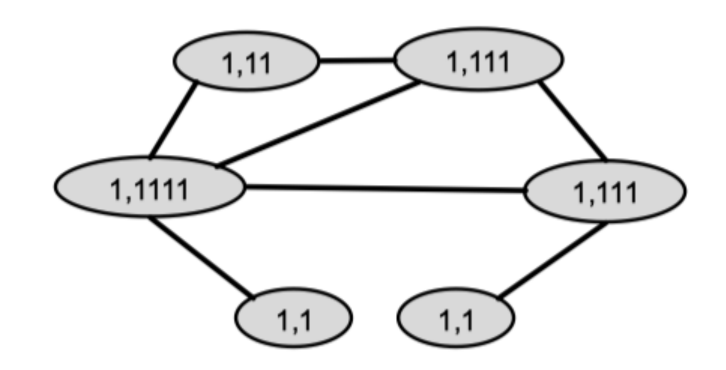
Assign initial colors





Aggregate neighboring colors

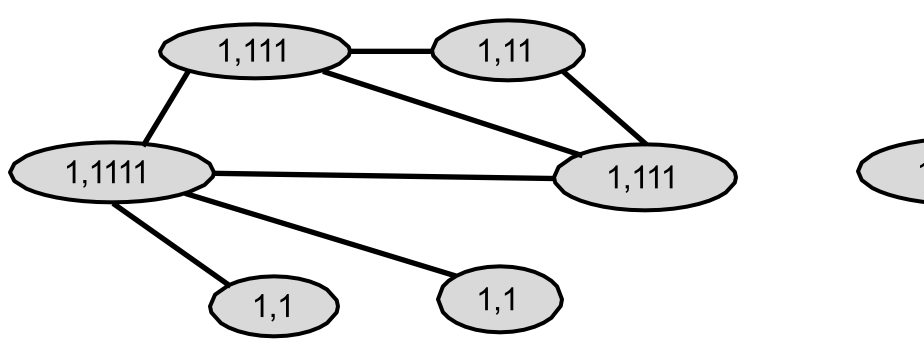


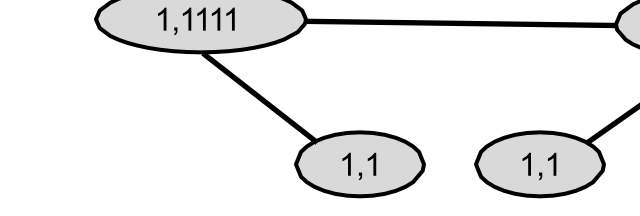


### Color Refinement (2)

#### Example of color refinement given two graphs

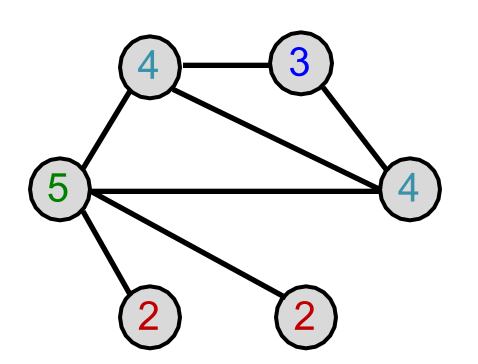
Aggregated colors

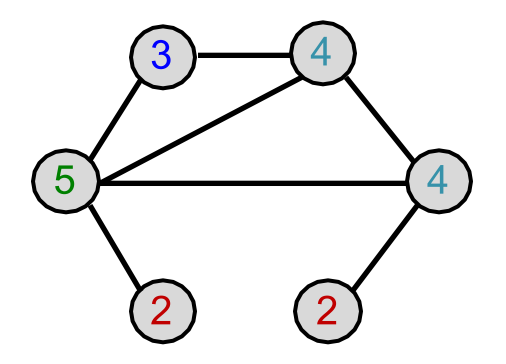




1,11

Hash aggregated colors





#### Hash table

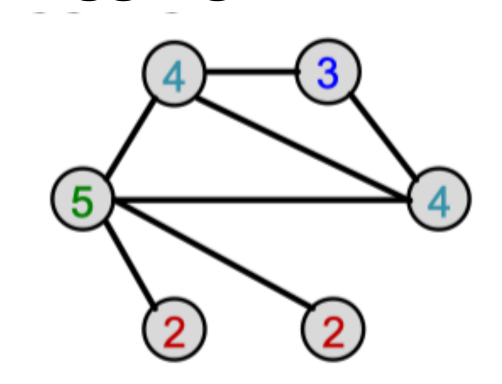
1,111

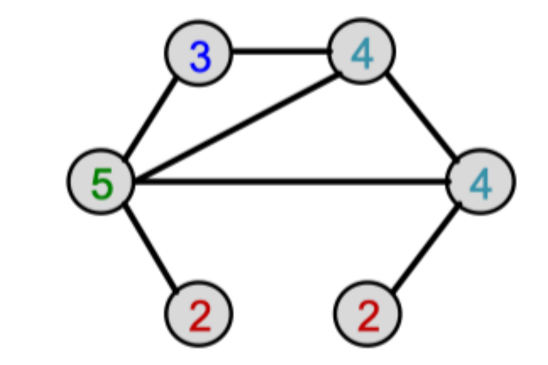
1,1	>	2	
1,11	>	3	
1,111	>	4	
1,1111	>	5	

## Color Refinement (3)

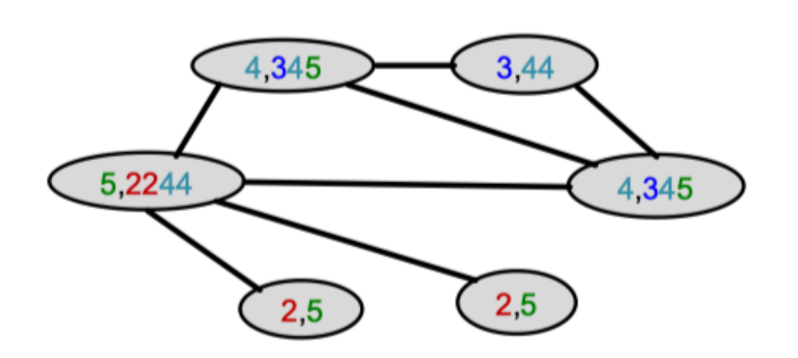
#### Example of color refinement given two graphs

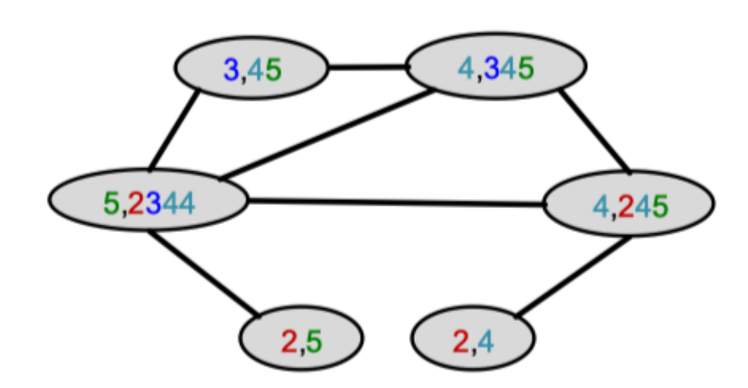
Aggregated colors





Hash aggregated colors

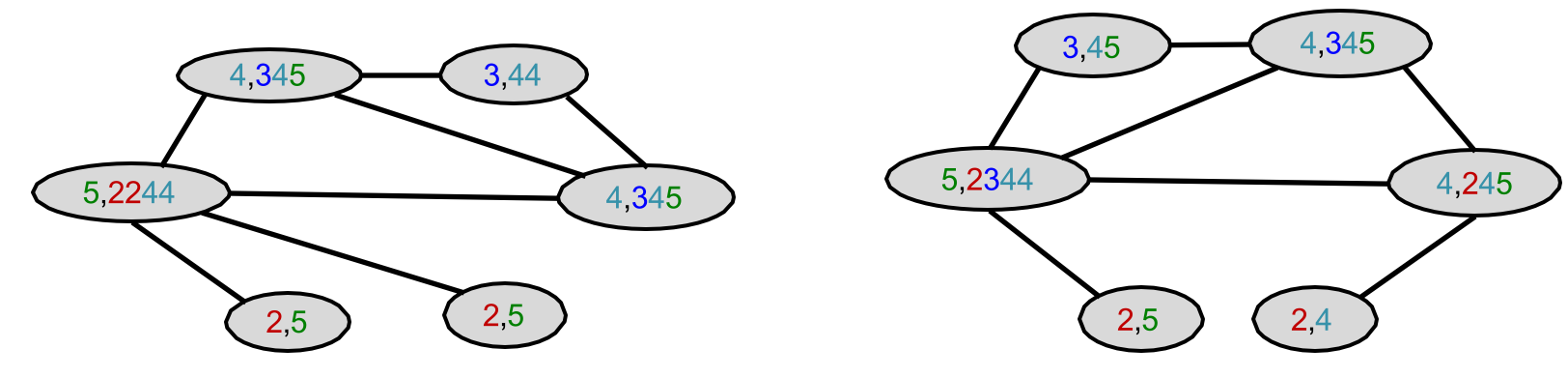




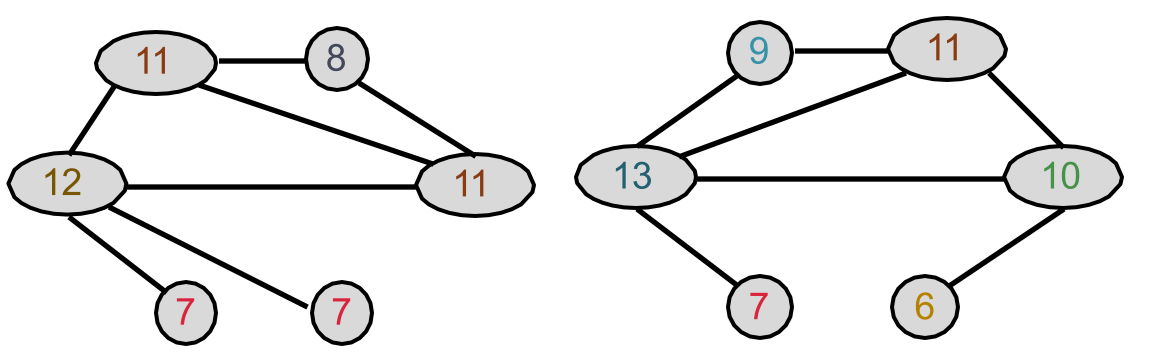
### Color Refinement (4)

#### Example of color refinement given two graphs

Aggregated colors



Hash aggregated colors



#### Hash table

2,4	<u> </u>
2,5	<b>&gt; 7</b>
3,44	<b>&gt;</b> 8
3,45	<b>&gt;</b> 9
4,245	<b>&gt;</b> 10
4,345	> 11
5,2244	<b>&gt;</b> 12
5,2344	<b>&gt;</b> 13

## Color Refinement (5)

#### **Stopping Condition:**

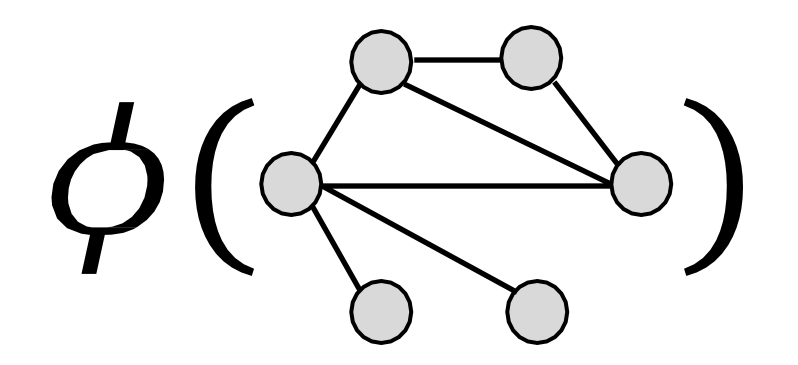
- It stops when node colors stabilize.
- Stabilization: no node's color changes in an iteration based on its current color and neighbors' colors.

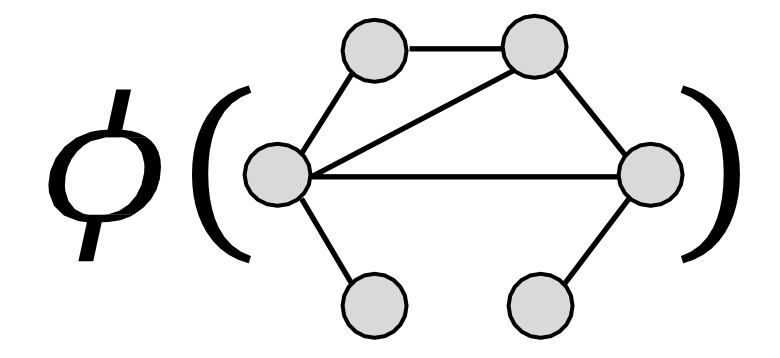
#### In Practice (for Graph Kernels):

- Often run for a predefined number of iterations (h) for consistent feature vectors.
- Guaranteed to stabilize within the number of nodes.
- Ensure that the resulting feature vectors have the same number of features.
  - NB: the list of unique colors is **global** to all the graphs in the dataset

# Weisfeiler-Lehman Graph Features

After color refinement, WL kernel counts number of nodes with a given color.





$$1,2,3,4,5,6,7,8,9,10,11,12,13$$
  
=  $[6,2,1,2,1,1,1,0,1,1,1,0,1]$ 

### Weisfeiler-Lehman Kernel

The WL kernel value is computed by the inner product of the color count vectors:

$$K(\stackrel{\checkmark}{\longleftrightarrow}, \stackrel{\checkmark}{\longleftrightarrow})$$

$$= \phi(\stackrel{\checkmark}{\longleftrightarrow})^{T}\phi(\stackrel{\checkmark}{\longleftrightarrow})$$

$$= 49$$

### Weisfeiler-Lehman Kernel

- WL kernel is computationally efficient.
  - The time complexity for color refinement at each step is linear in #(edges) since it aggregates neighboring colors.
- When computing a kernel value, only colors appearing in the two graphs must be tracked.
  - Thus, at most, #(colors) is the total number of nodes.
- Counting colors takes linear time w.r.t. #(nodes).
- In total, time complexity is linear in #(edges).

# Graph-level Features: Summary

#### Graphlet Kernel

- Graph is represented as Bag-of-graphlets
- Computationally expensive

#### Weisfeiler-Lehman Kernel

- Apply K-step color refinement algorithm to enrich node colors
  - Different colors capture different *K*-hop neighborhood structures
- Graph is represented as Bag-of-colors
- Computationally efficient
- Closely related to Graph Neural Networks (for the next course!)

# Summary

- Traditional ML Pipeline
  - Hand-crafted (structural) features + ML model
- Hand-crafted features for graph data
  - Node-level:
    - Node degree, centrality, clustering coefficient, graphlets
  - Link-level:
    - Distance-based feature
    - local/global neighborhood overlap
  - Graph-level:
    - Graphlet kernel, WL kernel
- However, we only considered featurizing the graph structure (but not the attribute of nodes and their neighbors)



