

# Deep Learning for Data Science

## DS 542

<https://dl4ds.github.io/sp2026/>

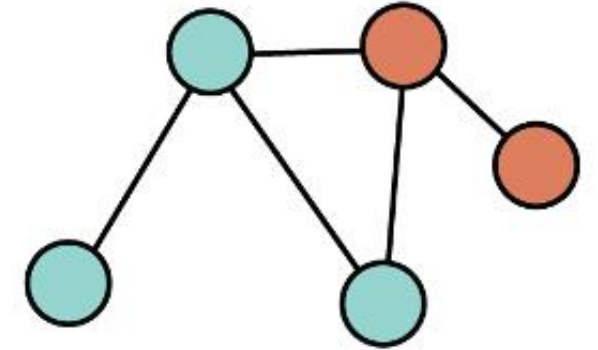
Graph Neural Networks

# Plan for Today

- Basic definition and examples
- Graph representation
- Properties of Adjacency Matrix
- Graph neural network, tasks and loss functions
- Graph convolutional network
- Graph & Node classification
- Edge graphs

# Graph Neural Networks

Neural architectures that process graphs.

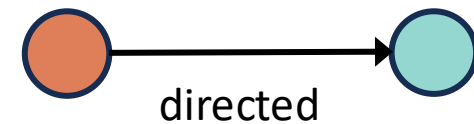
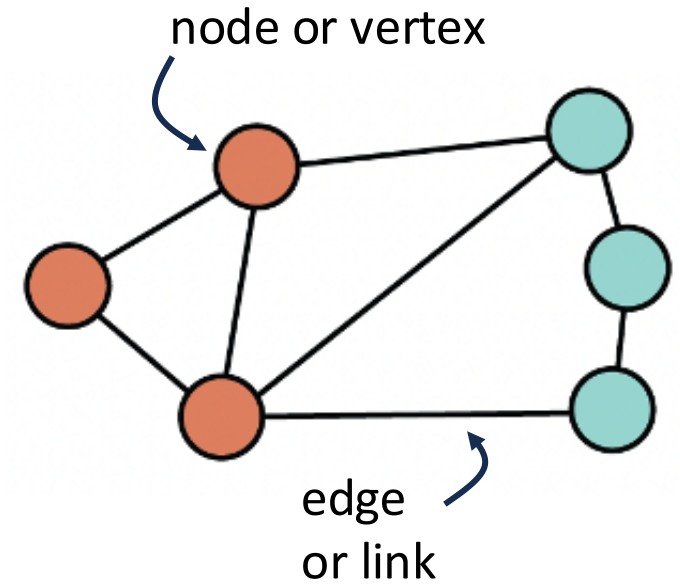


Three challenges:

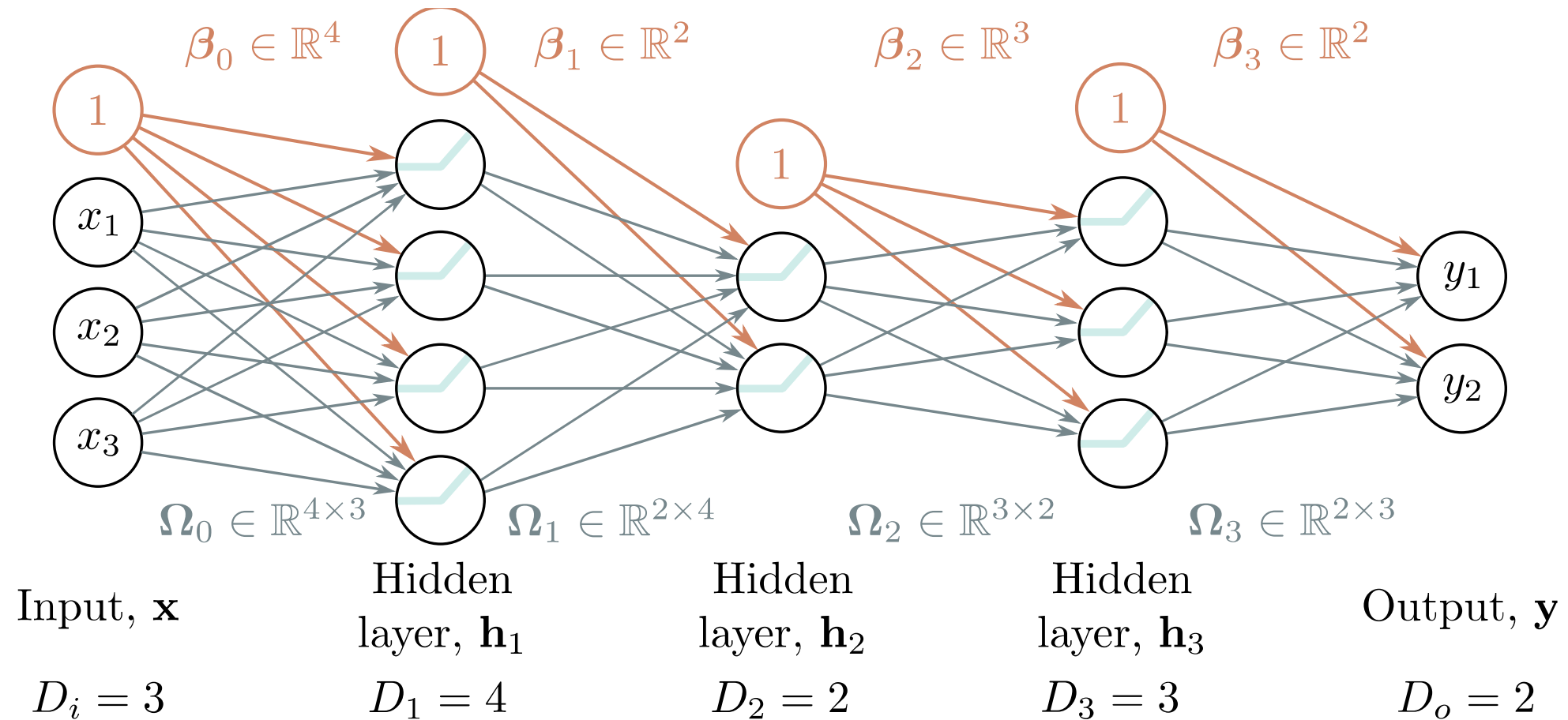
1. Variable topology
2. Size (billions of nodes)
3. Single monolithic graph

# Graph (Network)

- general structure composed of *nodes* (vertices) and *edges* (links)
- edges can be *undirected* or *directed*
- a graph with directed edges and no cycles (no loops) is called *directed acyclic graph* (DAG)

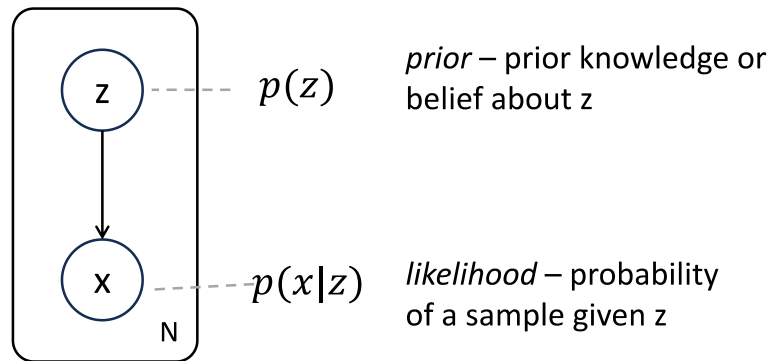


# Directed Example – Feed Forward Network

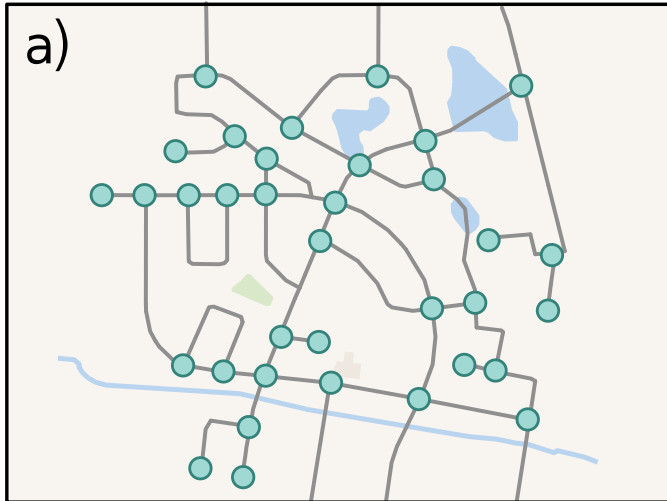


# Directed Example – Bayesian Graphical Model

## Preliminaries: Bayesian Models



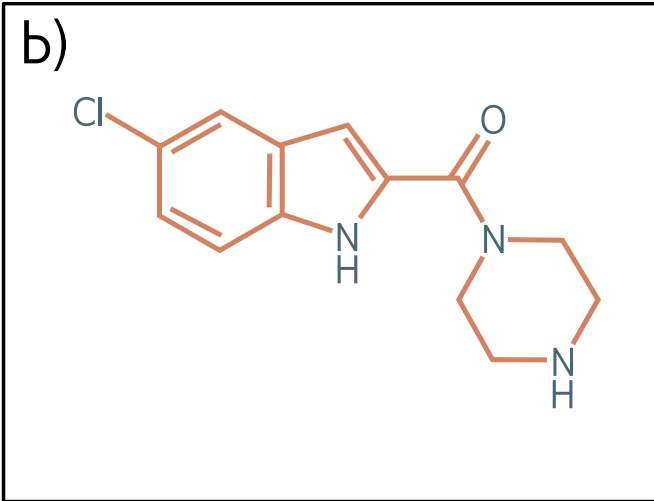
# Undirected Examples



**road networks**

**nodes:** physical locations or landmarks

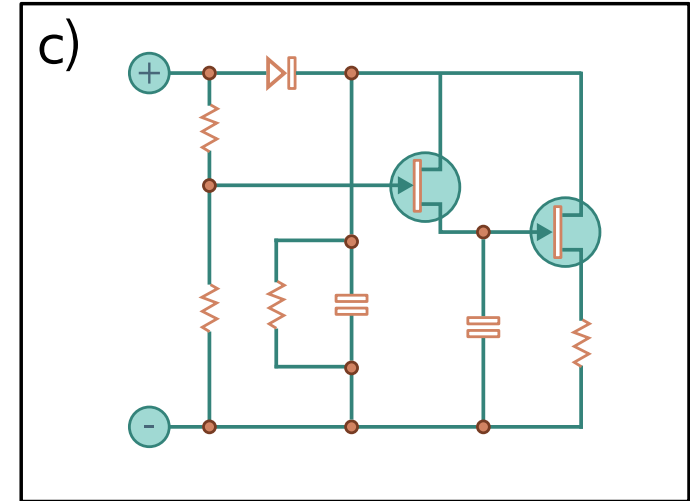
**edges:** connecting roads



**chemical molecules**

**nodes:** atoms

**edges:** chemical bonds

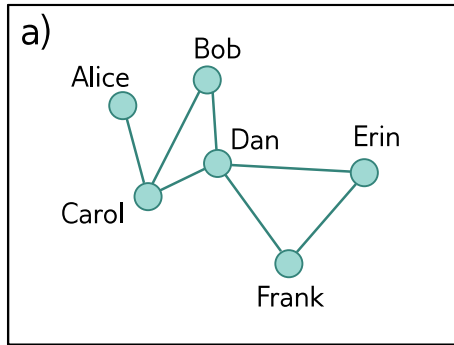


**electrical circuits**

**nodes:** components or junctions

**edges:** wires/electrical connections

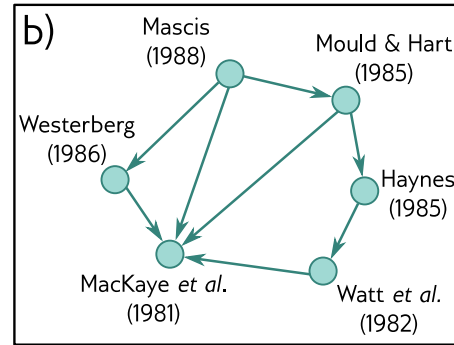
# Examples



**social networks**

**nodes:** people

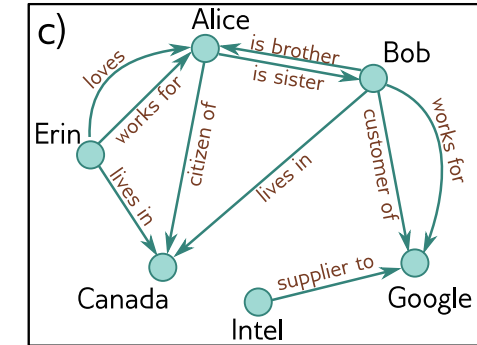
**edges:** friendships  
(undirected)



**science literature**

**nodes:** papers

**edges:** citations  
(acyclic directed)

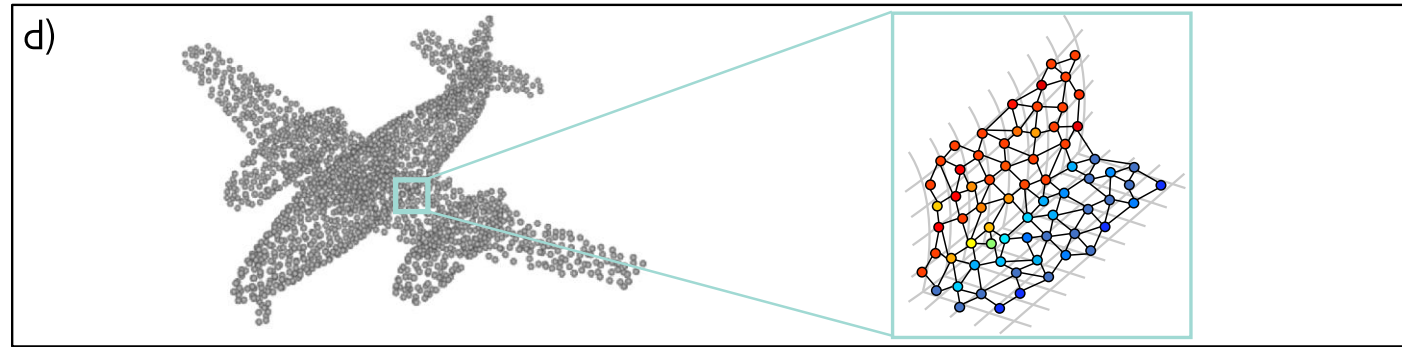


**knowledge graph**

**nodes:** objects

**edges:** named relationship  
(cyclic directed)

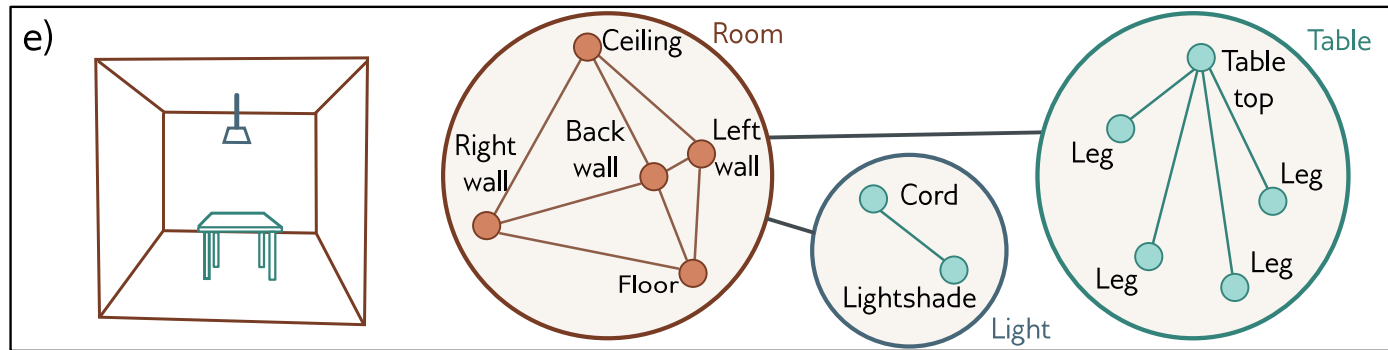
# Example – Geometric Point Cloud



**nodes:** positions in 3D space (vertex in 3D graphics)

**edges:** connections to nearby points  
(undirected)

# Example – Scene Graph



hierarchical graph showing relationship between objects in a 3D scene

**nodes:** composite graphs or objects in 3D space

**edges:** connections to nearby points  
(undirected)

# Other examples

- Wikipedia – nodes are articles, edges are hyperlinks between articles
- Computer programs – nodes are syntax tokens, edges are computation between tokens (tensor graph from Gradients lecture)
- Protein interactions – nodes are proteins, edges exist where two proteins interface
- Set or list – every element is connected to every other element
- image – each pixel is a node with edges to the eight adjacent pixels

# Any Questions?

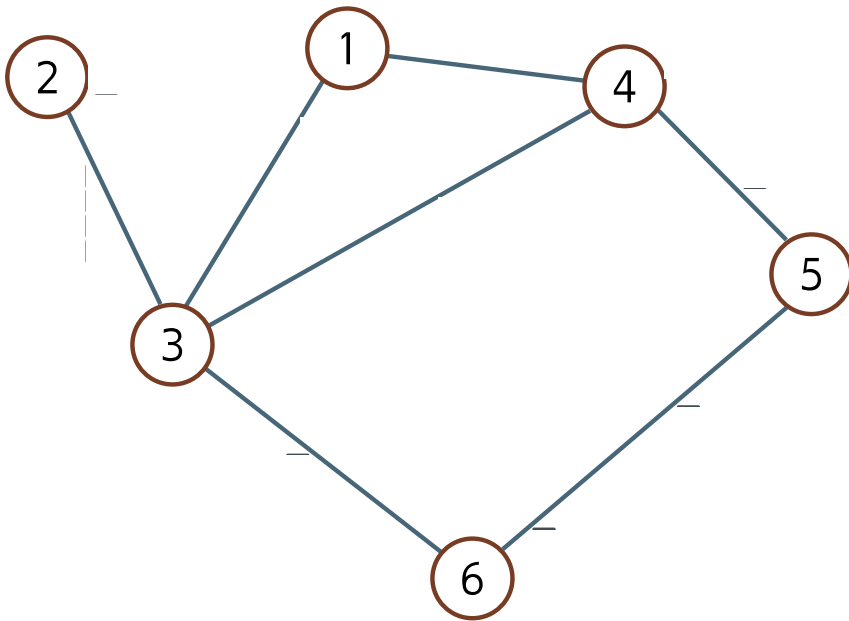


## Moving on

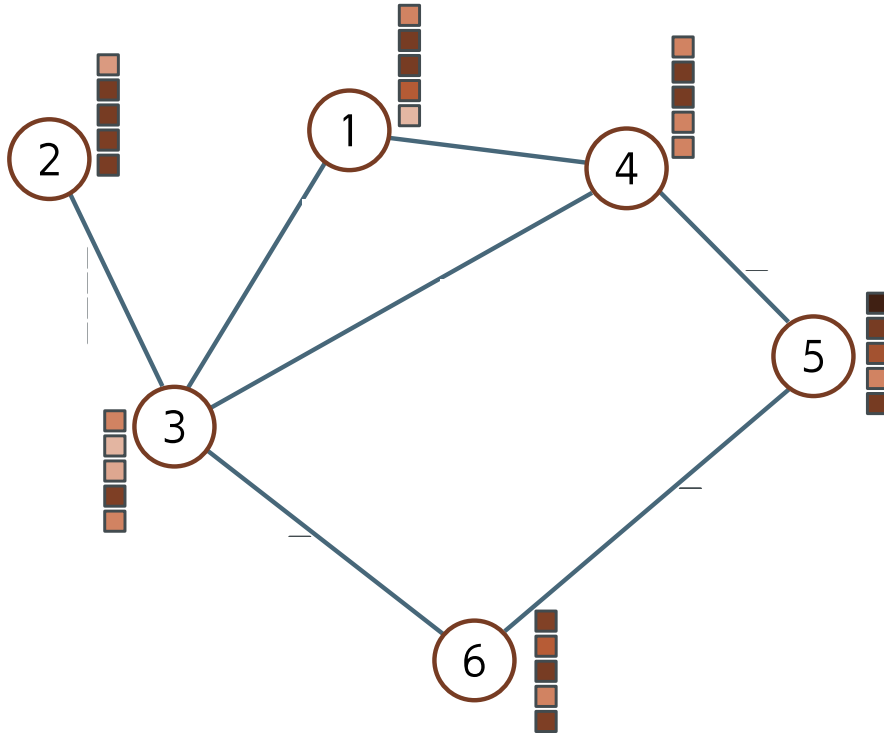
- Basic definition and examples
- **Graph representation**
- Properties of Adjacency Matrix
- Graph neural network, tasks and loss functions
- Graph convolutional network
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# Graph representation

Example undirected graph with 6 nodes



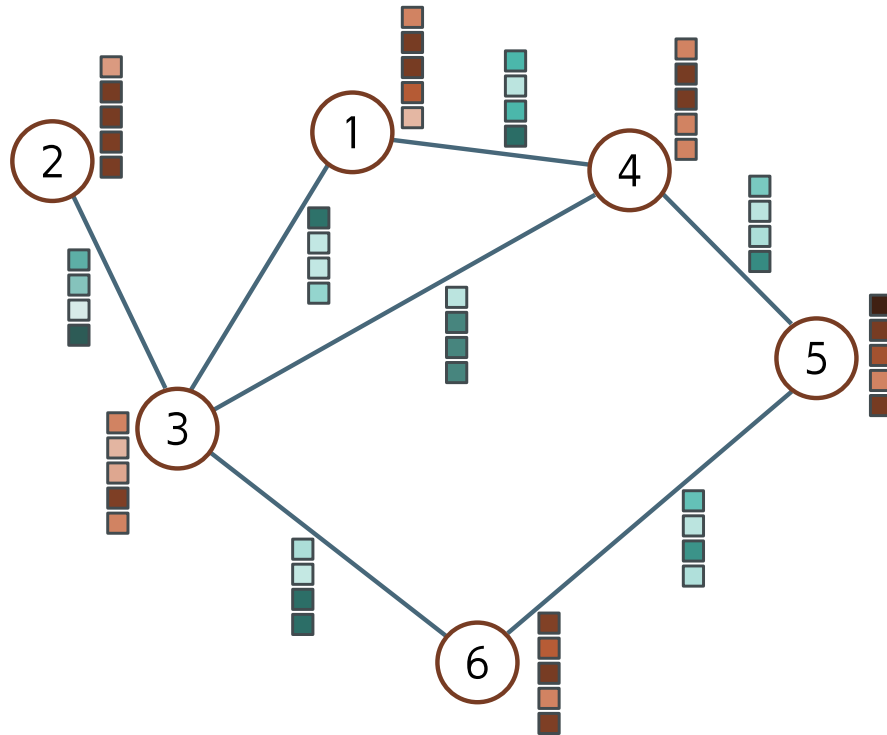
# Graph representation – node embedding



Example undirected graph with 6 nodes

Information about a node is stored in a *node embedding*

# Graph representation – edge embedding

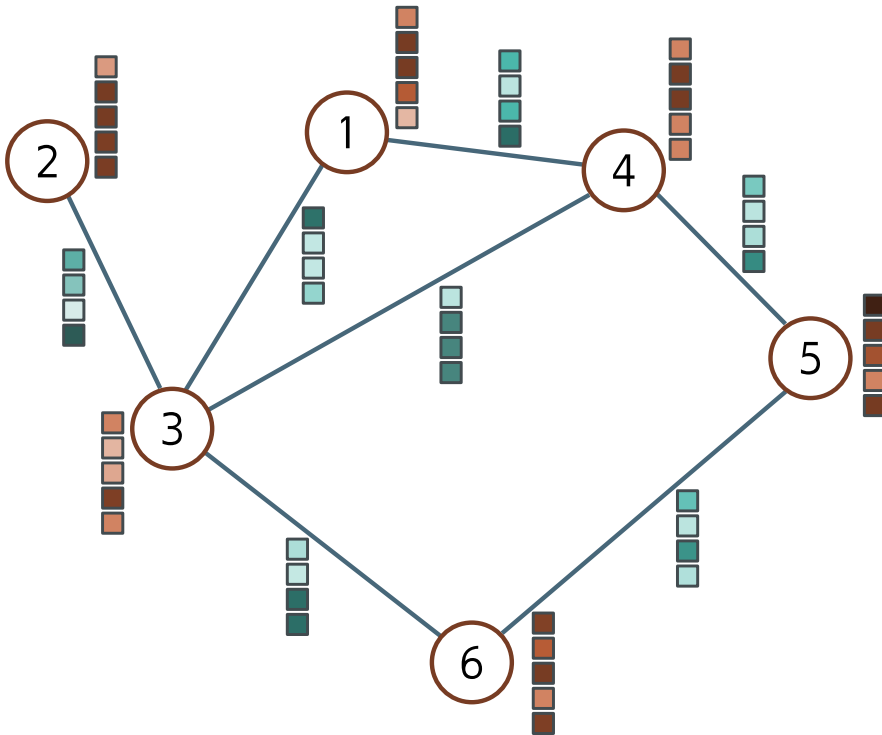


Example undirected graph with 6 nodes

Information about a node is stored in a *node embedding*

Information about an edge is stored in an *edge embedding*

# Graph representation – adjacency matrix



Adjacency  
matrix,  $A$   
 $N \times N$

	1	2	3	4	5	6
1						
2						
3						
4						
5						
6						

Assume we have  $N$  nodes

The graph connections can be represented by an *adjacency matrix*

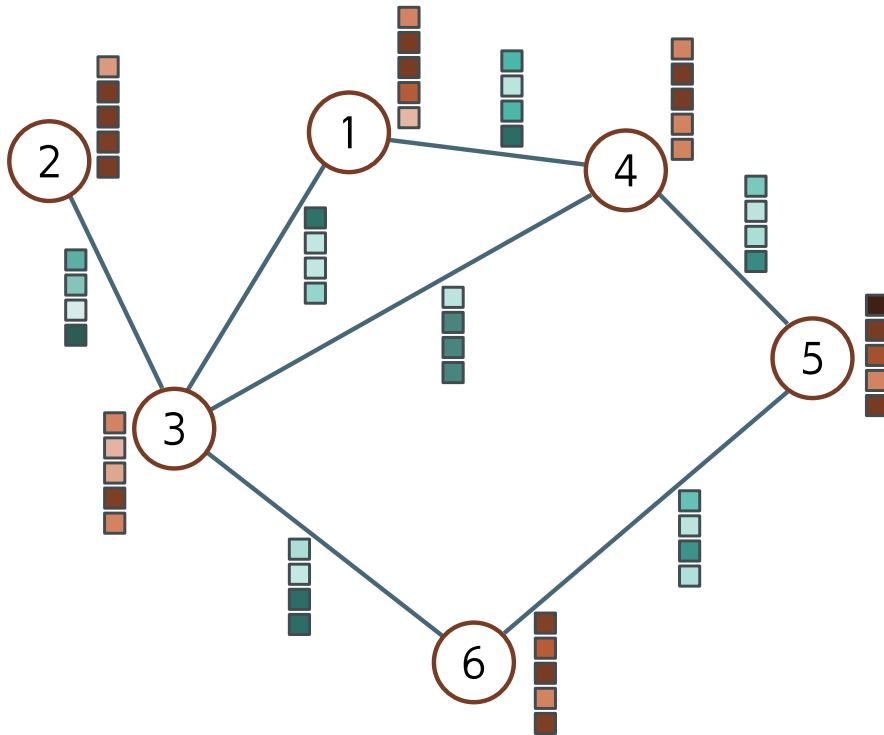
Where a value of 1 at  $(m, n)$  represents a connection between nodes  $m$  and  $n$ .

For undirected graphs the matrix is always symmetric about the diagonal

Diagonal is zero – no edge to itself

Can be very sparse

# Graph representation – node data matrix



Adjacency matrix,  $\mathbf{A}$   
 $N \times N$

	1	2	3	4	5	6
1						
2						
3						
4						
5						
6						

Node data,  $\mathbf{X}$   
 $D \times N$

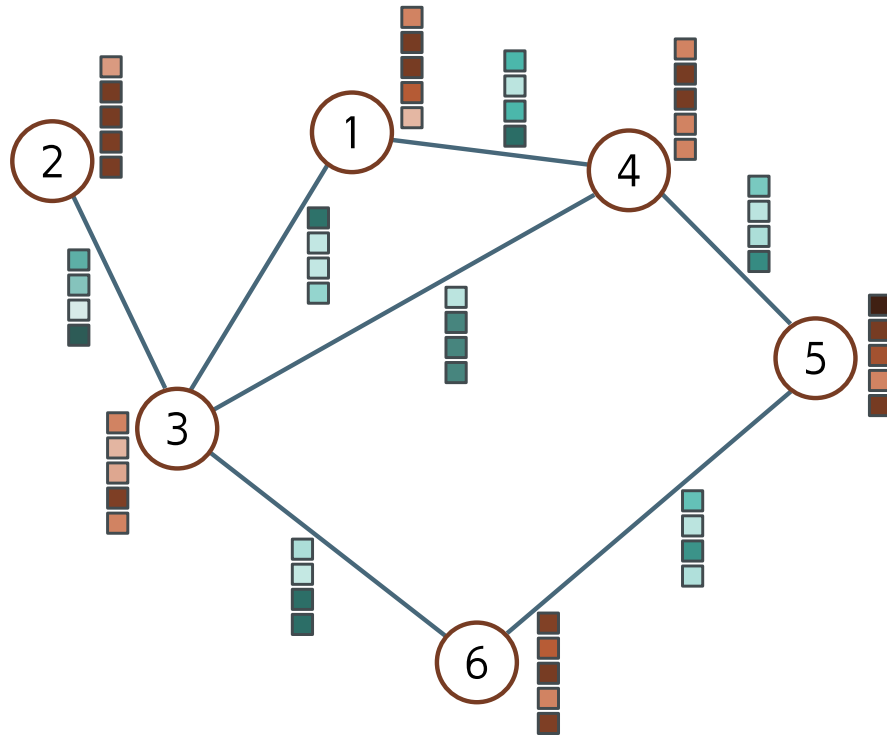
	1	2	3	4	5	6
1						
2						
3						
4						
5						
6						

All the node data in the form of node embeddings can be represented by a *Node data matrix*

Where  $D$  is the dimension of the node embedding and

$N$  is the number of nodes

# Graph representation – edge data matrix



Adjacency matrix,  $\mathbf{A}$   
 $N \times N$

	1	2	3	4	5	6
1						
2						
3						
4						
5						
6						

Node data,  $\mathbf{X}$   
 $D \times N$

	1	2	3	4	5	6
1						
2						
3						
4						
5						
6						

Edge data,  $\mathbf{E}$   
 $D_E \times E$

	1	1	2	3	3	4	5
1							
2							
3							
4							
5							
6							

Similarly, all the edge embedding information can be stored in an *Edge data matrix*, where:  
 $D_E$  is the dimension of the edge embedding vector and  
 $E$  is the number of edges

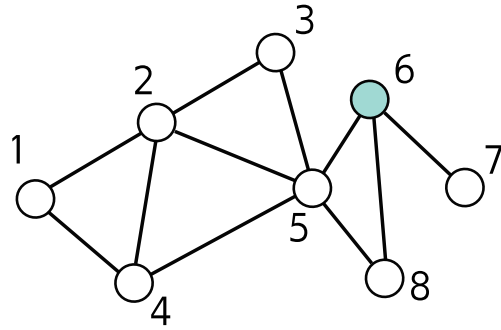
# Any Questions?



## Moving on

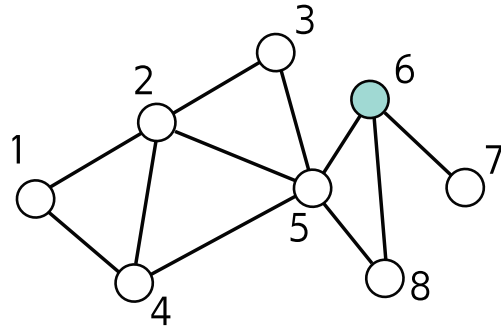
- Basic definition and examples
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- **Properties of Adjacency Matrix**
- Graph neural network, tasks and loss functions
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# Adjacency Matrix



Assume we have an 8-node undirected graph

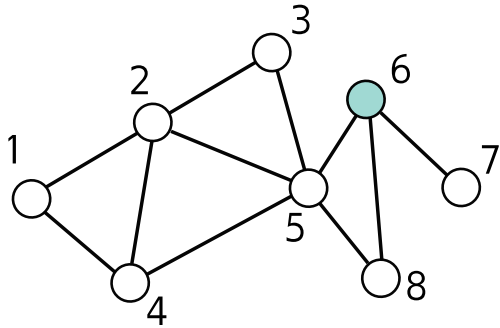
# Adjacency Matrix



$$\mathbf{A} = \begin{bmatrix} 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 \\ 1 & 0 & 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 1 & 1 & 1 & 0 & 1 & 0 & 1 \\ 0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 1 & 0 & 0 \end{bmatrix}$$

Adjacency matrix for this graph.

# Adjacency Matrix



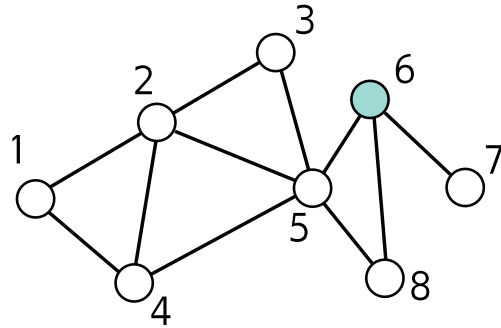
$$\mathbf{A} = \begin{bmatrix} 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 \\ 1 & 0 & 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 1 & 1 & 1 & 0 & 1 & 0 & 1 \\ 0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 1 & 0 & 0 \end{bmatrix}$$

adjacency matrix

$$\mathbf{x} = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 0 \end{bmatrix}$$

We can one hot encode  
representation of node 6

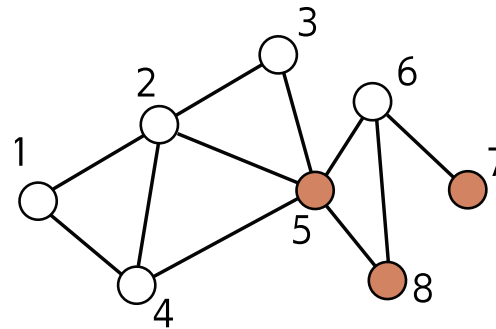
# Adjacency Matrix



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

$$x = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 0 \end{bmatrix}$$

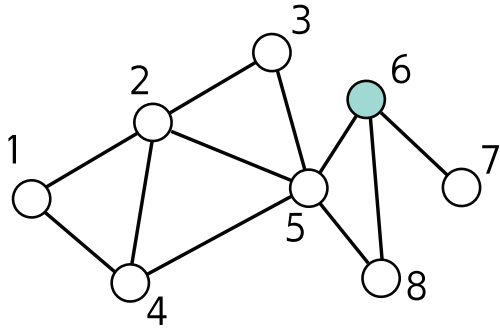
$$Ax = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 1 \end{bmatrix}$$



If we pre-multiply the one-hot encoded data node vector  $x$  by adjacency matrix  $A$  we get the 6<sup>th</sup> column of  $A$  indicating direct connections to other nodes

One-hot encoding vector of all nodes directly connected node 6

# Adjacency Matrix

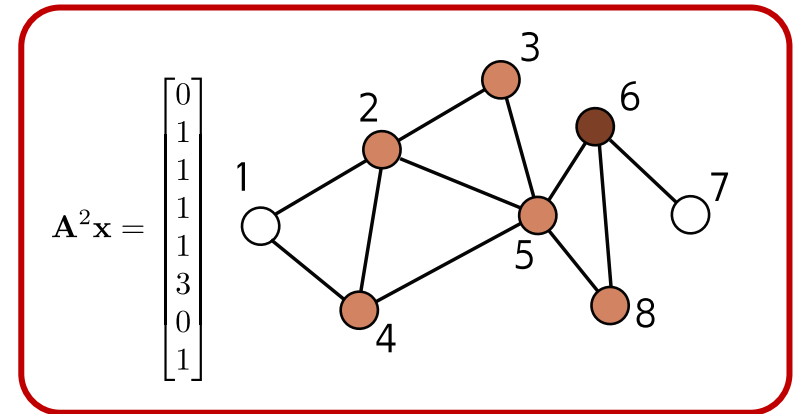
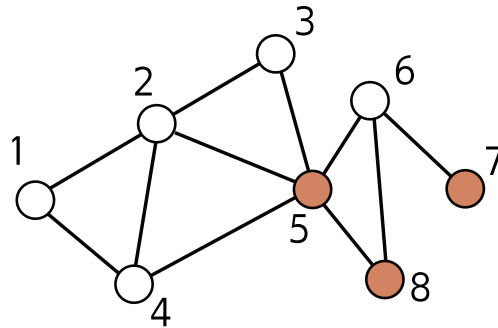


$$A = \begin{bmatrix} 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 \\ 1 & 0 & 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 1 & 1 & 1 & 0 & 1 & 0 & 1 \\ 0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 1 & 0 & 0 \end{bmatrix}$$

If we pre-multiply again by  $A$ , we get a vector showing the number of times we can get to each node in 2 steps.

$$x = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 0 \end{bmatrix}$$

$$Ax = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 1 \\ 1 \end{bmatrix}$$



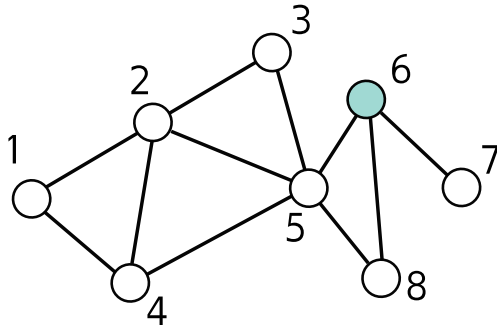
$$A^2x = \begin{bmatrix} 0 \\ 1 \\ 1 \\ 1 \\ 1 \\ 1 \\ 3 \\ 0 \\ 1 \end{bmatrix}$$

Graph showing all nodes that can be reached in *exactly* 2 steps.

# Adjacency Matrix

Pre-multiplying  $x$  by  $A$  twice is equivalent to the matrix  $A^2$

Shows how many times you can get from node  $m$  to node  $n$  in 2 steps

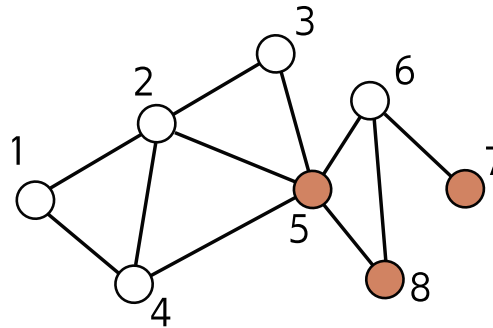


$$A = \begin{bmatrix} 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 \\ 1 & 0 & 1 & 1 & 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 1 & 1 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 1 & 1 & 1 & 0 & 1 & 0 & 1 \\ 0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\ 0 & 0 & 0 & 0 & 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 0 & 1 & 1 & 0 & 0 \end{bmatrix}$$

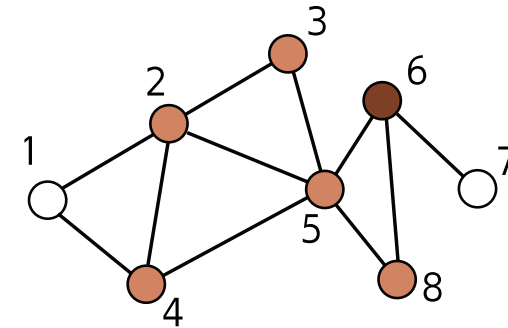
$$A^2 = \begin{bmatrix} 2 & 1 & 1 & 1 & 2 & 0 & 0 & 0 \\ 1 & 4 & 1 & 2 & 2 & 1 & 0 & 1 \\ 1 & 1 & 2 & 2 & 1 & 1 & 0 & 1 \\ 1 & 2 & 2 & 3 & 1 & 1 & 0 & 1 \\ 2 & 2 & 1 & 1 & 5 & 1 & 1 & 1 \\ 0 & 1 & 1 & 1 & 1 & 3 & 0 & 1 \\ 0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\ 0 & 1 & 1 & 1 & 1 & 1 & 1 & 2 \end{bmatrix}$$

$$x = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 0 \end{bmatrix}$$

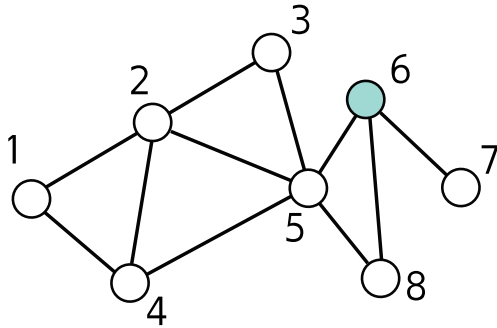
$$Ax = \begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 1 \\ 0 \\ 1 \end{bmatrix}$$



$$A^2x = \begin{bmatrix} 0 \\ 1 \\ 1 \\ 1 \\ 1 \\ 3 \\ 0 \\ 1 \end{bmatrix}$$



# Adjacency Matrix



$$\mathbf{A}^2 = \begin{bmatrix} 2 & 1 & 1 & 1 & 2 & 0 & 0 & 0 \\ 1 & 4 & 1 & 2 & 2 & 1 & 0 & 1 \\ 1 & 1 & 2 & 2 & 1 & 1 & 0 & 1 \\ 1 & 2 & 2 & 3 & 1 & 1 & 0 & 1 \\ 2 & 2 & 1 & 1 & 5 & 1 & 1 & 1 \\ 0 & 1 & 1 & 1 & 1 & 3 & 0 & 1 \\ 0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\ 0 & 1 & 1 & 1 & 1 & 1 & 1 & 2 \end{bmatrix}$$

Example for  $L = 2$

When you raise the adjacency matrix to the power of  $L$  (pre-multiply  $L-1$  times),

the entry at position  $(m, n)$  of  $\mathbf{A}^L$  contains the number of unique walks of length  $L$  from node  $n$  to node  $m$

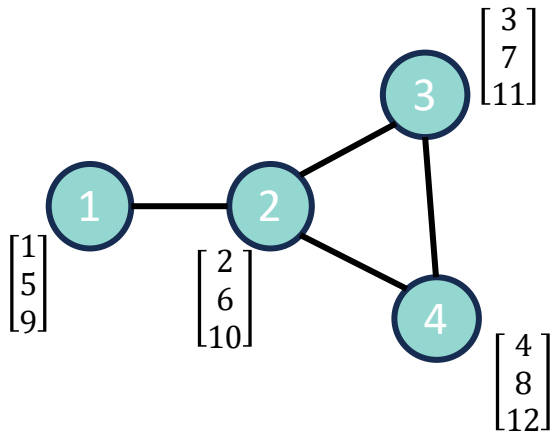
Note: this is not the same as the number of unique paths since it includes routes that visit the same node more than once.

a non-zero entry at position  $(m, n)$  indicates that the distance from  $m$  to  $n$  must be less than or equal to  $L$ .

[See Notebook 13.1 – Encoding Graphs](#)

# Permutation of node indices

Since node indexing is arbitrary, we can permute the node indices



$$\mathbf{X} = \begin{matrix} & \begin{pmatrix} 1 & 2 & 3 & 4 \end{pmatrix} \\ \begin{bmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{bmatrix} \end{matrix}$$

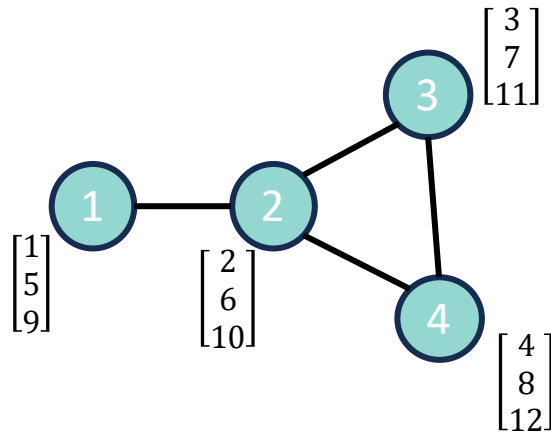
node data

$$\mathbf{A} = \begin{bmatrix} 0 & 1 & 0 & 0 \\ 1 & 0 & 1 & 1 \\ 0 & 1 & 0 & 1 \\ 0 & 1 & 1 & 0 \end{bmatrix}$$

adjacency matrix

# Permutation of node indices

Since node indexing is arbitrary, we can permute the node indices



$$\mathbf{X} = \begin{matrix} & \begin{matrix} (1 & 2 & 3 & 4) \end{matrix} \\ \begin{matrix} (1 \\ 2 \\ 3) \end{matrix} & \begin{bmatrix} 1 & 2 & 3 & 4 \\ 5 & 6 & 7 & 8 \\ 9 & 10 & 11 & 12 \end{bmatrix} \end{matrix}$$

node data

$$\mathbf{A} = \begin{bmatrix} 0 & 1 & 0 & 0 \\ 1 & 0 & 1 & 1 \\ 0 & 1 & 0 & 1 \\ 0 & 1 & 1 & 0 \end{bmatrix}$$

adjacency matrix

$$\mathbf{P} = \begin{matrix} & \begin{matrix} (3 & 4 & 2 & 1) \end{matrix} \\ \begin{matrix} (1 \\ 2 \\ 3 \\ 4) \end{matrix} & \begin{bmatrix} 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \\ 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{bmatrix} \end{matrix}$$

We can express this mathematically with a permutation matrix,  $\mathbf{P}$

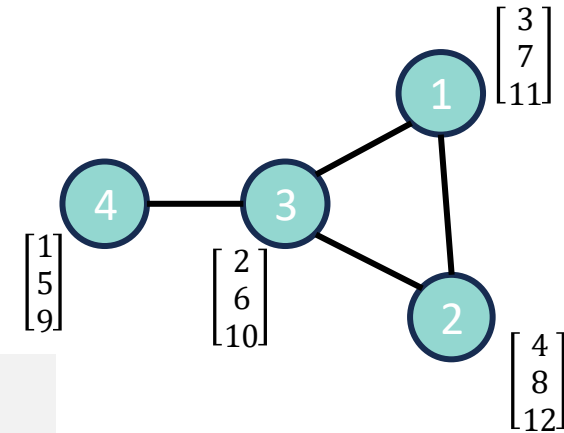
$$\begin{matrix} \text{New: } (1 & 2 & 3 & 4) \\ \text{Old: } (3 & 4 & 2 & 1) \end{matrix}$$

$$\mathbf{X}' = \mathbf{X}\mathbf{P} = \begin{bmatrix} 3 & 4 & 2 & 1 \\ 7 & 8 & 6 & 5 \\ 11 & 12 & 10 & 9 \end{bmatrix}$$

Permute the columns of the Node data matrix

$$\mathbf{A}' = \mathbf{P}^T \mathbf{A} \mathbf{P} = \begin{bmatrix} 0 & 1 & 1 & 0 \\ 1 & 0 & 1 & 0 \\ 1 & 1 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{bmatrix}$$

Permute both the rows and column of the Adjacency matrix



# Any Questions?



## Moving on

- Basic definition and examples
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- Properties of Adjacency Matrix
- Graph neural network, tasks and loss functions
- Graph convolutional network
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# Graph Neural Network

- A graph neural network is a model that takes the **node embeddings**  $\mathbf{X}$  and the **adjacency matrix**  $\mathbf{A}$  as inputs and passes them through a series of  $K$  layers.
- The node embeddings are updated at each layer to create intermediate “hidden” representations  $\mathbf{H}_K$  before finally computing **output embeddings**  $\mathbf{H}_K$ .
- At the start of this network, each column of the input node embeddings  $\mathbf{X}$  just contains information about the node itself.
- At the end, each column of the model output  $\mathbf{H}_K$  includes **information about the node and its context within the graph**.
- This is **like word embeddings passing through a transformer** network. These represent words at the start but represent the word meanings in the context of the sentence at the end.

# Graph Level Tasks

Determine

- class categories, e.g. molecule is poisonous
  - regression values, e.g. molecule boiling and freezing point
- based on graph structure and node embeddings

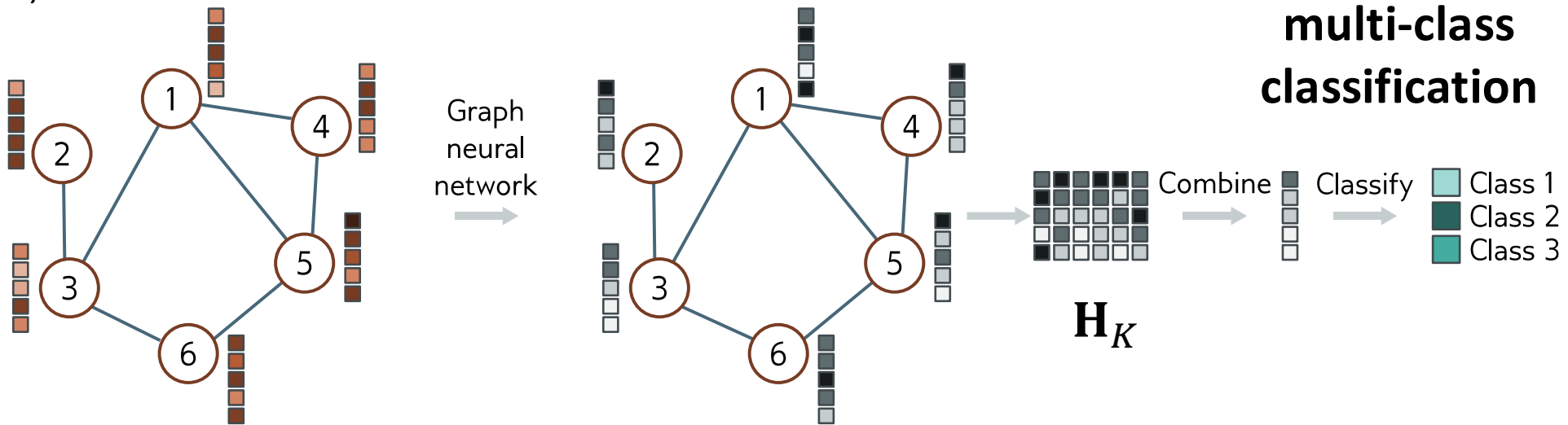
For graph-level tasks, the output node embeddings are combined (e.g., by averaging), and the resulting vector is mapped via a linear transformation or neural network to a fixed-size vector

# Typical Three Types of Models

- Graph level regression & classification
- Node level regression & classification
- Edge prediction

Look at prediction heads first.

# Graph level regression & classification



Last layer (Regression):  $\Pr(y|\mathbf{X}, \mathbf{A}) = \beta_K + \omega_K \mathbf{H}_K \mathbf{1} / N$

Last layer (Classification):  $\Pr(y = 1|\mathbf{X}, \mathbf{A}) = \text{sigmoid}[\beta_K + \omega_K \mathbf{H}_K \mathbf{1} / N]$

Regression Loss Function: Least Squares Loss  
Classification Loss Function: (Binary) Cross Entropy

$\beta_K$  is scalar

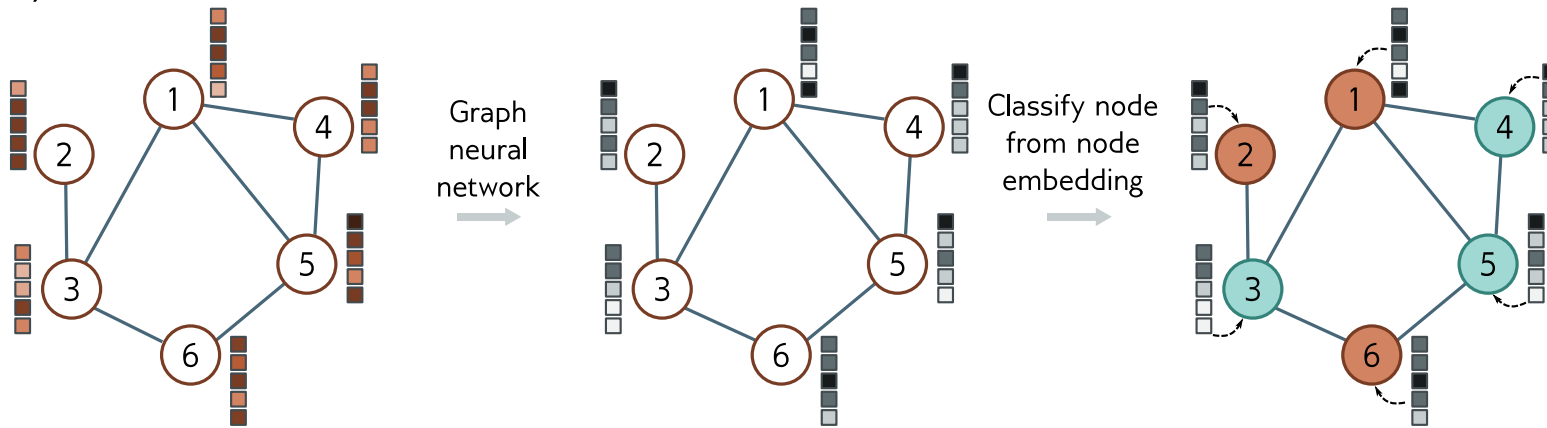
$\omega_K$  is  $1 \times D$  row vector

$\mathbf{H}_K$  is the  $D \times N$  output embedding matrix

$\mathbf{1}$  is an  $N \times 1$  column vector of 1s

Mean pooling

# Node level binary regression & classification



Last layer (Regression):  $\Pr(y^{(n)} | \mathbf{X}, \mathbf{A}) = \beta_K + \omega_K \mathbf{h}_K^{(n)}$

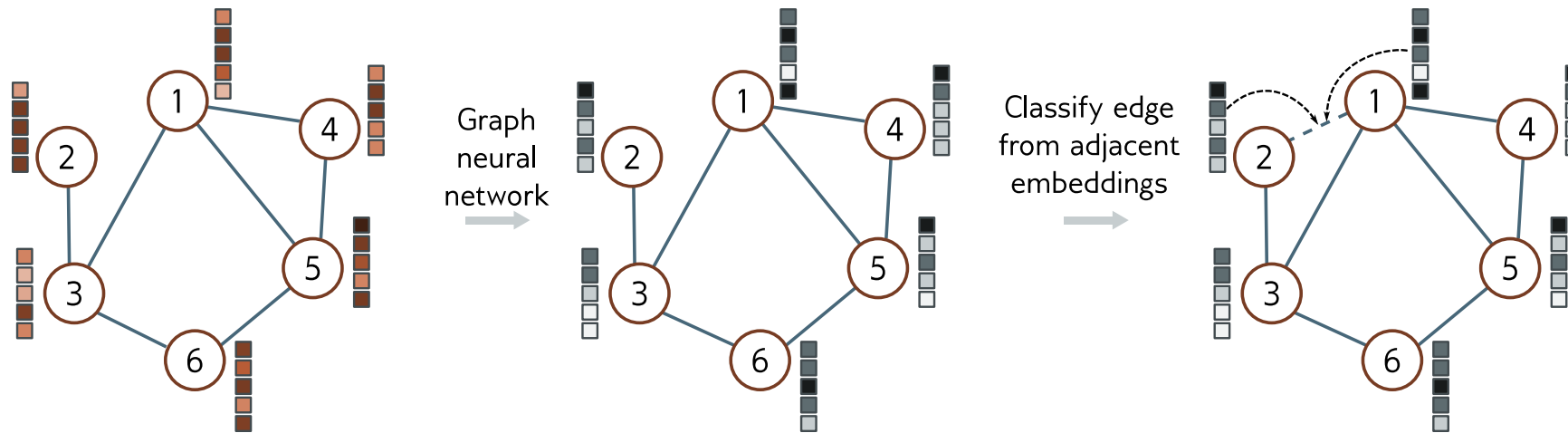
Last layer (Classification):  $\Pr(y^{(n)} = 1 | \mathbf{X}, \mathbf{A}) = \text{sigmoid}[\beta_K + \omega_K \mathbf{h}_K^{(n)}]$

$\mathbf{h}_K^{(n)}$  is the  $D \times 1$  output embedding vector node for  $n$

Regression Loss Function: Least Squares Loss  
Classification Loss Function: (Binary) Cross Entropy

# Edge prediction (classification)

Predict whether edge should exist or not.



Last layer:  $\Pr(y^{(mn)} = 1 | \mathbf{X}, \mathbf{A}) = \text{sigmoid}[\mathbf{h}_K^{(m)T} \mathbf{h}_K^{(n)}]$

$$[1 \times D][D \times 1]$$

Classification Loss Function: Binary Cross Entropy

# Any Questions?



## **Moving on**

- Basic definition and examples
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- Properties of Adjacency Matrix
- Graph neural network, tasks and loss functions
- Graph convolutional network
- Graph & Node classification
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# Graph convolutional network

These models are convolutional in that they update each node by aggregating information from nearby nodes.

As such, they induce a relational inductive bias (i.e., a bias toward prioritizing information from neighbors).

$$\begin{aligned}\mathbf{H}_1 &= \mathbf{F}[\mathbf{X}, \mathbf{A}, \phi_0] \\ \mathbf{H}_2 &= \mathbf{F}[\mathbf{H}_1, \mathbf{A}, \phi_1] \\ \mathbf{H}_3 &= \mathbf{F}[\mathbf{H}_2, \mathbf{A}, \phi_2] \\ \vdots &= \vdots \\ \mathbf{H}_K &= \mathbf{F}[\mathbf{H}_{K-1}, \mathbf{A}, \phi_{K-1}],\end{aligned}$$

A function  $F[\cdot]$  with parameters  $\phi_i$  that takes the node embeddings and adjacency matrix and outputs new node embeddings

# Equivariance and Invariance

Every layer should be *equivariant* to index permutations

$$\mathbf{H}_{k+1}\mathbf{P} = \mathbf{F}[\mathbf{H}_k\mathbf{P}, \mathbf{P}^T\mathbf{A}\mathbf{P}, \phi_k]$$

And for node classification and edge prediction the output should be *invariant* to index permutations

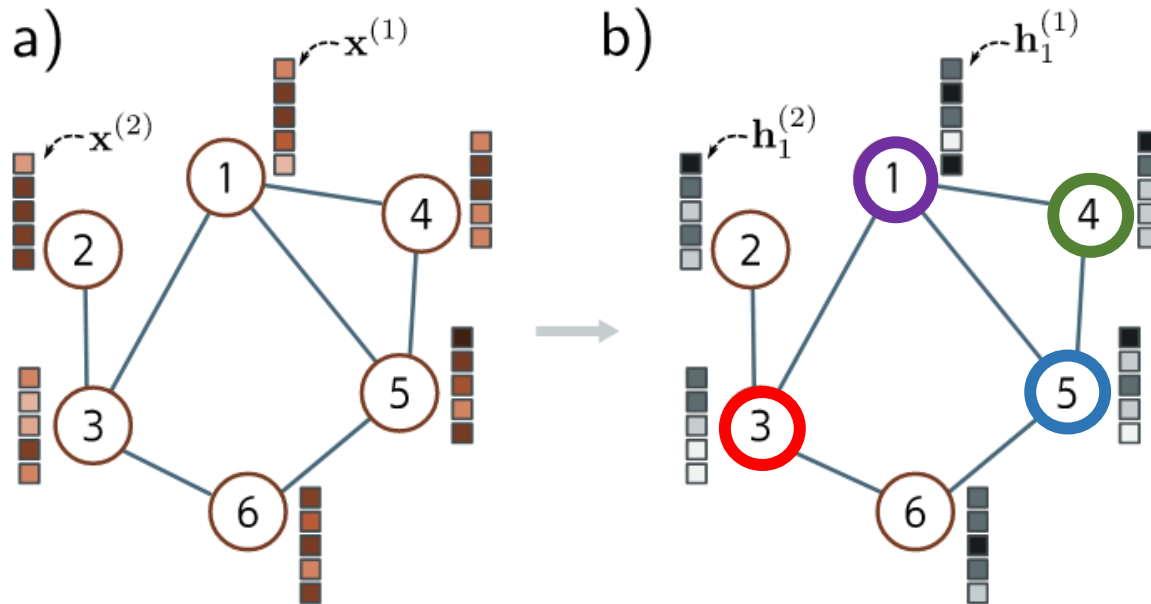
$$y = \text{sigmoid}[\beta_K + \omega_K \mathbf{H}_K \mathbf{1}/N] = \text{sigmoid}[\beta_K + \omega_K \mathbf{H}_K \mathbf{P} \mathbf{1}/N]$$

# Example Graph Convolution Network (GCN) layer

At each node  $n$  in layer  $k$ , aggregate information from neighboring nodes

$$\text{agg}[n, k] = \sum_{m \in \text{ne}[n]} \mathbf{h}_k^{(m)}$$

where  $\text{ne}[n]$  returns the set of indices of the neighbors of node  $n$ .



$$\text{ne}[1] = \{4, 5, 3\}$$

$$\text{agg}[n = 1, k = 1] = \mathbf{h}_1^{(4)} + \mathbf{h}_1^{(5)} + \mathbf{h}_1^{(3)}$$

# Example Graph Convolution Network (GCN) layer

At each node  $n$  in layer  $k$ , aggregate information from neighboring nodes

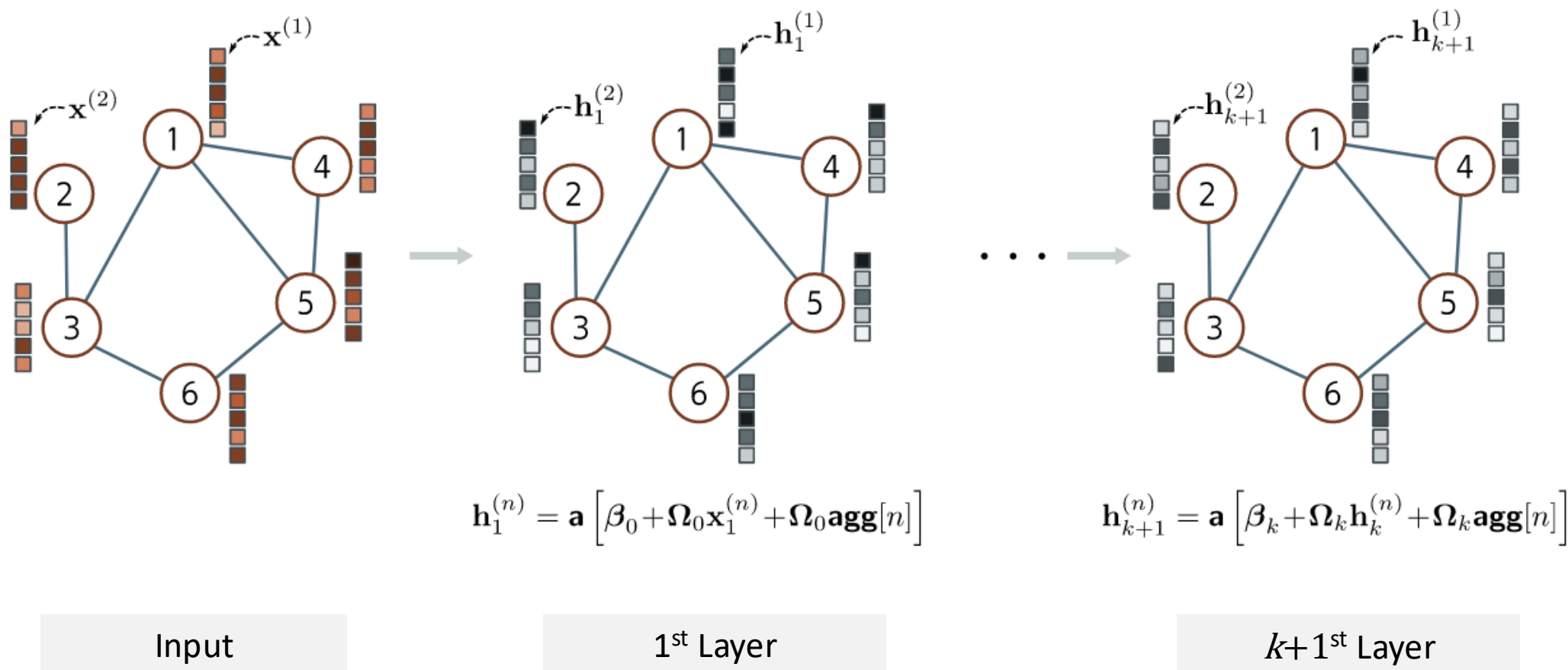
$$\text{agg}[n, k] = \sum_{m \in \text{ne}[n]} \mathbf{h}_k^{(m)}$$

where  $\text{ne}[n]$  returns the set of indices of the neighbors of node  $n$ .

Then a linear transform to the current node vector and the aggregate for the current node and add a bias.

$$\mathbf{h}_{k+1}^{(n)} = \mathbf{a} \left[ \underset{D \times 1}{\boldsymbol{\beta}_k} + \underset{D \times D}{\boldsymbol{\Omega}_k} \cdot \underset{D \times 1}{\mathbf{h}_k^{(n)}} + \underset{D \times D}{\boldsymbol{\Omega}_k} \cdot \underset{D \times 1}{\text{agg}[n, k]} \right]$$

# Graph convolution layers



# Example Graph Convolution Network (GCN) layer

We apply the following equation

$$\mathbf{h}_{k+1}^{(n)} = \mathbf{a} \left[ \beta_k + \Omega_k \cdot \mathbf{h}_k^{(n)} + \Omega_k \cdot \text{agg}[n, k] \right]$$

to the entire node hidden layers matrix,  $\mathbf{H}_k$ , by noting that  $\mathbf{H}_k \mathbf{A}$  produces a matrix where the  $n^{\text{th}}$  column is  $\text{agg}[n, k]$ .

$$\begin{aligned} \mathbf{H}_{k+1} &= \mathbf{a} \left[ \beta_k \mathbf{1}^T + \Omega_k \mathbf{H}_k + \Omega_k \mathbf{H}_k \mathbf{A} \right] \\ &= \mathbf{a} \left[ \beta_k \mathbf{1}^T + \Omega_k \mathbf{H}_k (\mathbf{A} + \mathbf{I}) \right], \end{aligned}$$

# Example Graph Convolution Network (GCN) layer

We apply the following equation

$$\mathbf{h}_{k+1}^{(n)} = \mathbf{a} \left[ \beta_k + \Omega_k \cdot \mathbf{h}_k^{(n)} + \Omega_k \cdot \text{agg}[n, k] \right]$$

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$$\begin{aligned} \mathbf{H}_{k+1} &= \mathbf{a} \left[ \beta_k \mathbf{1}^T + \Omega_k \mathbf{H}_k + \Omega_k \mathbf{H}_k \mathbf{A} \right] \\ &= \mathbf{a} \left[ \beta_k \mathbf{1}^T + \Omega_k \mathbf{H}_k (\mathbf{A} + \mathbf{I}) \right], \end{aligned}$$

Note that this is (1) equivariant to permutations, (2) handles arbitrary number of neighbors, (3) exploits graph structure and (4) share parameters

# Any Questions?



## Moving on

- Basic definition and examples
- Graph representation
- Properties of Adjacency Matrix
- Graph neural network, tasks and loss functions
- Graph convolutional network
- Graph & Node classification
- Edge graphs

# Graph classification example

We can put it all together and add a sigmoid layer

$$\begin{aligned}\mathbf{H}_1 &= \mathbf{a} [\beta_0 \mathbf{1}^T + \Omega_0 \mathbf{X}(\mathbf{A} + \mathbf{I})] \\ \mathbf{H}_2 &= \mathbf{a} [\beta_1 \mathbf{1}^T + \Omega_1 \mathbf{H}_1(\mathbf{A} + \mathbf{I})] \\ &\vdots \\ \mathbf{H}_K &= \mathbf{a} [\beta_{K-1} \mathbf{1}^T + \Omega_{K-1} \mathbf{H}_{K-1}(\mathbf{A} + \mathbf{I})] \\ f[\mathbf{X}, \mathbf{A}, \Phi] &= \text{sig} [\beta_K + \underbrace{\omega_K \mathbf{H}_K \mathbf{1} / N}_{\text{Mean pooling}}],\end{aligned}$$

For classification on molecules,

$X \in \mathbb{R}^{118 \times N}$ : one hot encoding of 118 elements

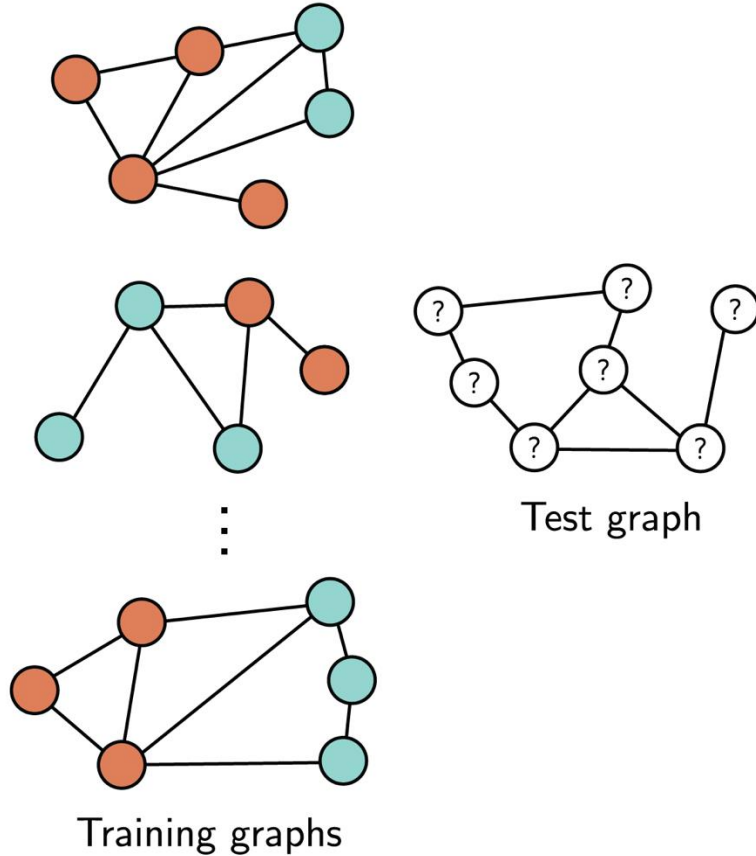
$\Omega_0 \in \mathbb{R}^{D \times 118}$ : convert to  $D$ -dimensional embeddings

$\beta_K$ : is a scalar

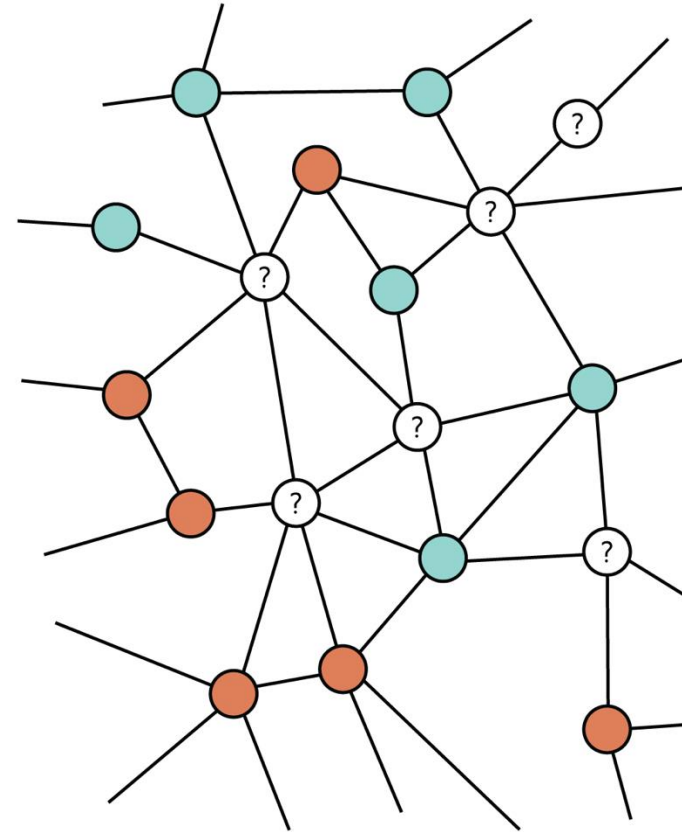
$\omega_K$ : a  $1 \times D$  parameters row vector

# Inductive

# vs. Transductive



supervised learning: train with the labeled graphs and then run inference on the unlabeled (test) graphs



semi-supervised learning: train with the labeled nodes, then run inference to determine label for unlabeled nodes

# Node classification example

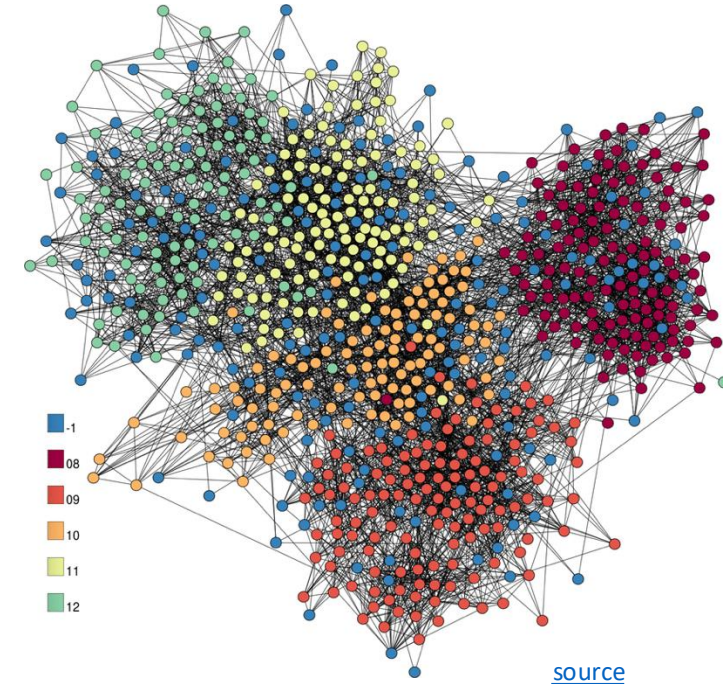
Assume *transductive* binary node *classification* with millions of nodes, *partially labeled*.

Same network body as graph classification, but different head:

$$\mathbf{f}[\mathbf{X}, \mathbf{A}, \Phi] = \text{sigmoid}[\beta_K \mathbf{1}^T + \omega_K \mathbf{H}_K]$$

No mean pooling. Output is  $1 \times N$ .

Train with binary cross-entropy loss on nodes with labels.

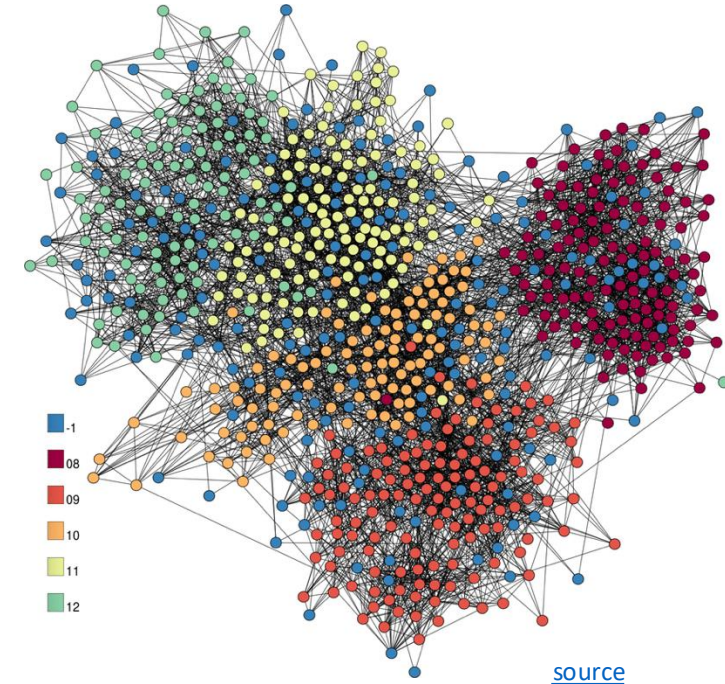


# Node classification example

Assume *transductive* binary node *classification* with millions of nodes, *partially labeled*.

Challenges:

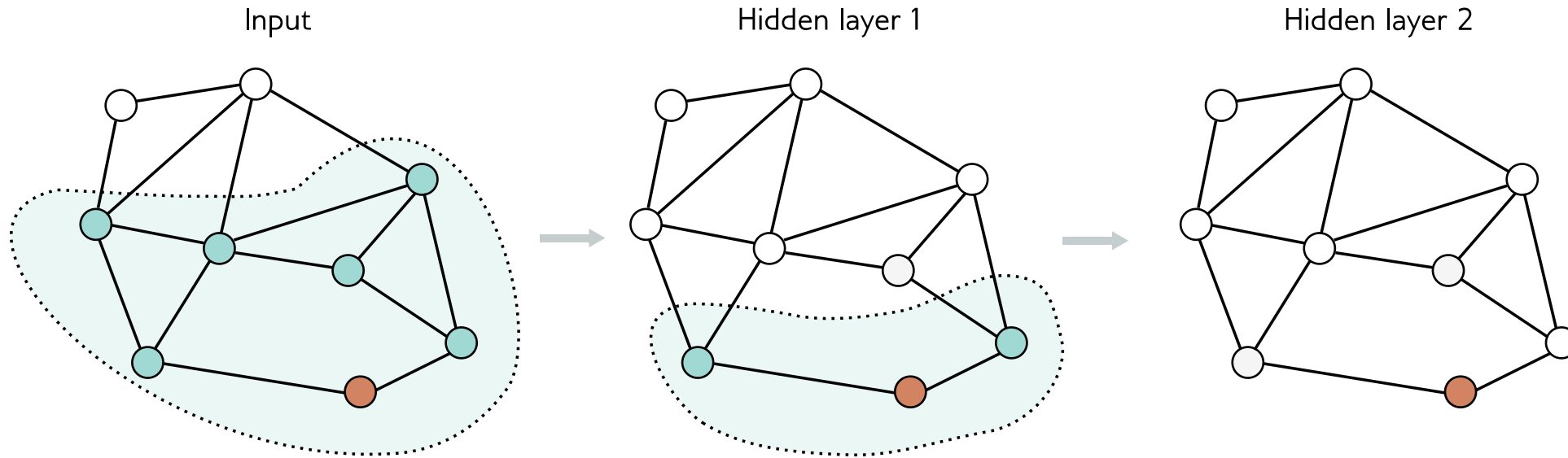
1. memory limitations: need to store every node and hidden layer embedding during training
2. how to perform SGD with basically one batch!



# Solutions: Choosing batches for graphs

1. Choose random subset of nodes
2. Neighborhood sampling
3. Graph partitioning

# Batches: Random subset



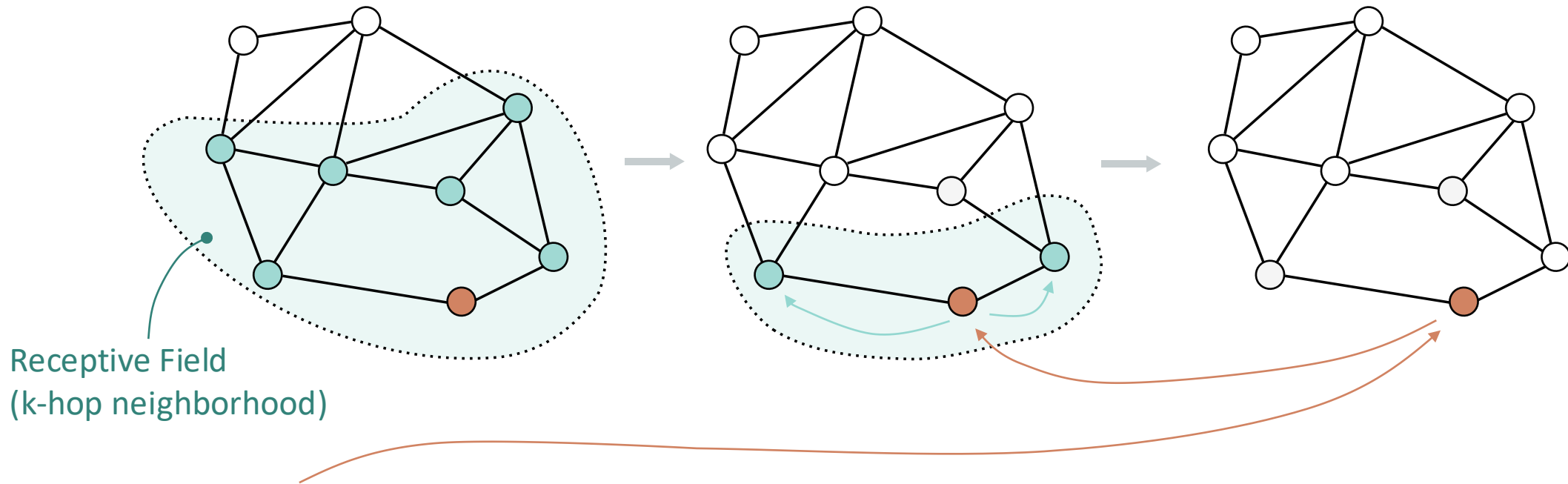
You can pick a random batch of labeled nodes at each training step,  
And only include them and their “ $k$ -hop neighborhoods”.

# Batches: Random subset

Input

Hidden layer 1

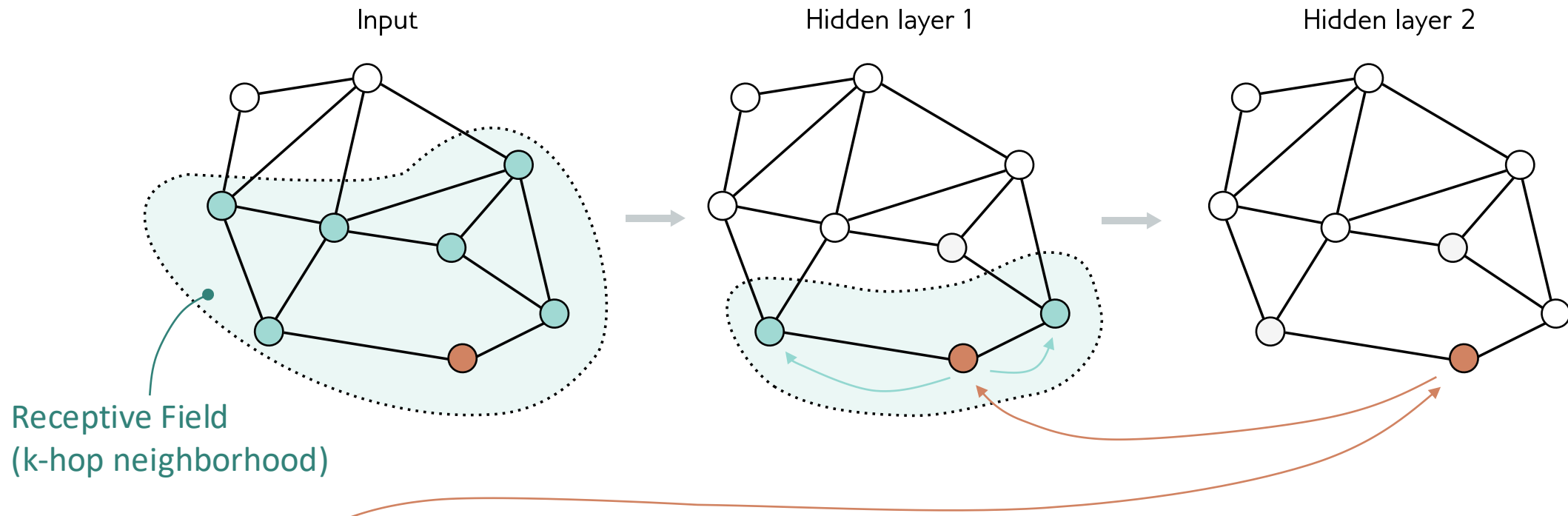
Hidden layer 2



Each **node** is dependent on the same node in the previous layer and its **neighbors** because of `agg[]`

$$\mathbf{h}_{k+1}^{(n)} = \mathbf{a} \left[ \beta_k + \Omega_k \cdot \mathbf{h}_k^{(n)} + \Omega_k \cdot \text{agg}[n, k] \right]$$

# Batches: Random subset

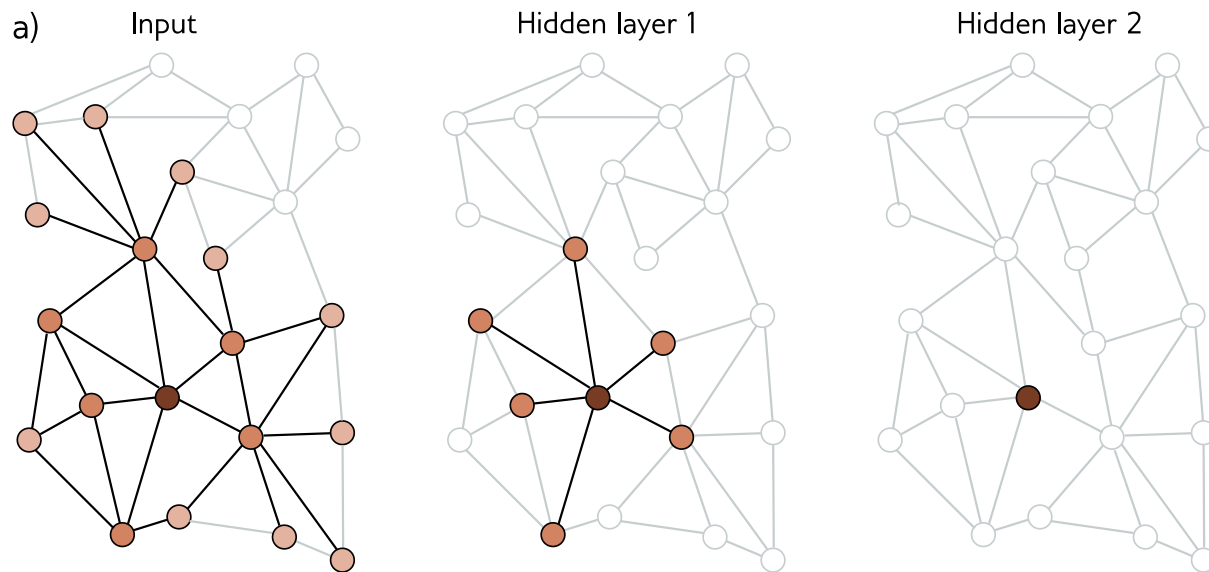


Each **node** is dependent on the same node in the previous layer and its **neighbors** because of `agg[]`.

$$\mathbf{h}_{k+1}^{(n)} = \mathbf{a} \left[ \beta_k + \Omega_k \cdot \mathbf{h}_k^{(n)} + \Omega_k \cdot \text{agg}[n, k] \right]$$

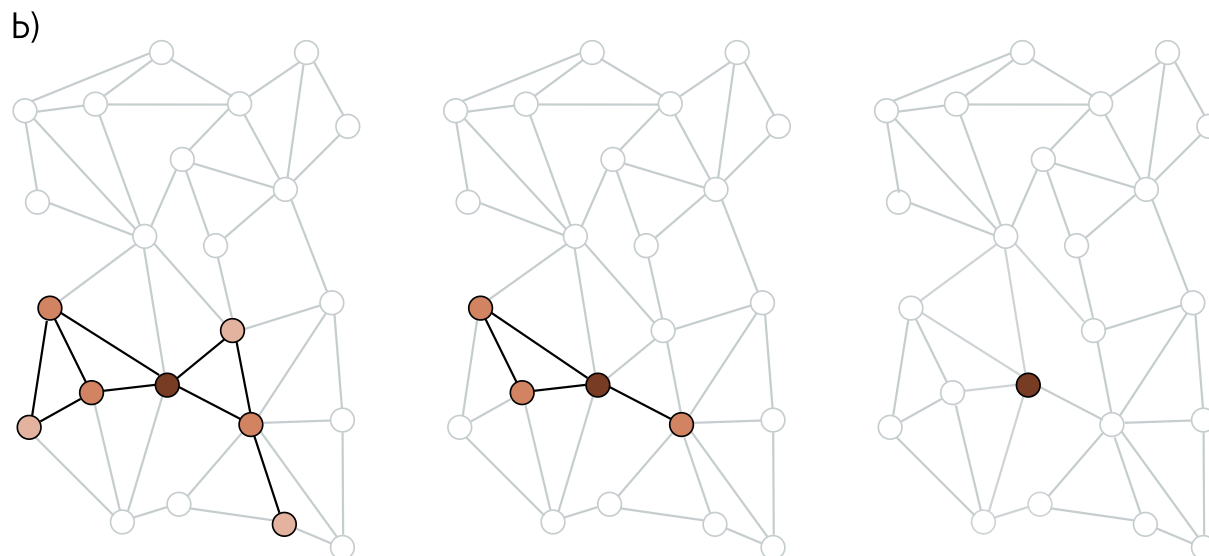
With many layers and dense connection, it can quickly expand to encompass every node.

# Neighborhood Sampling



## Random Sampling:

Use all the neighbors



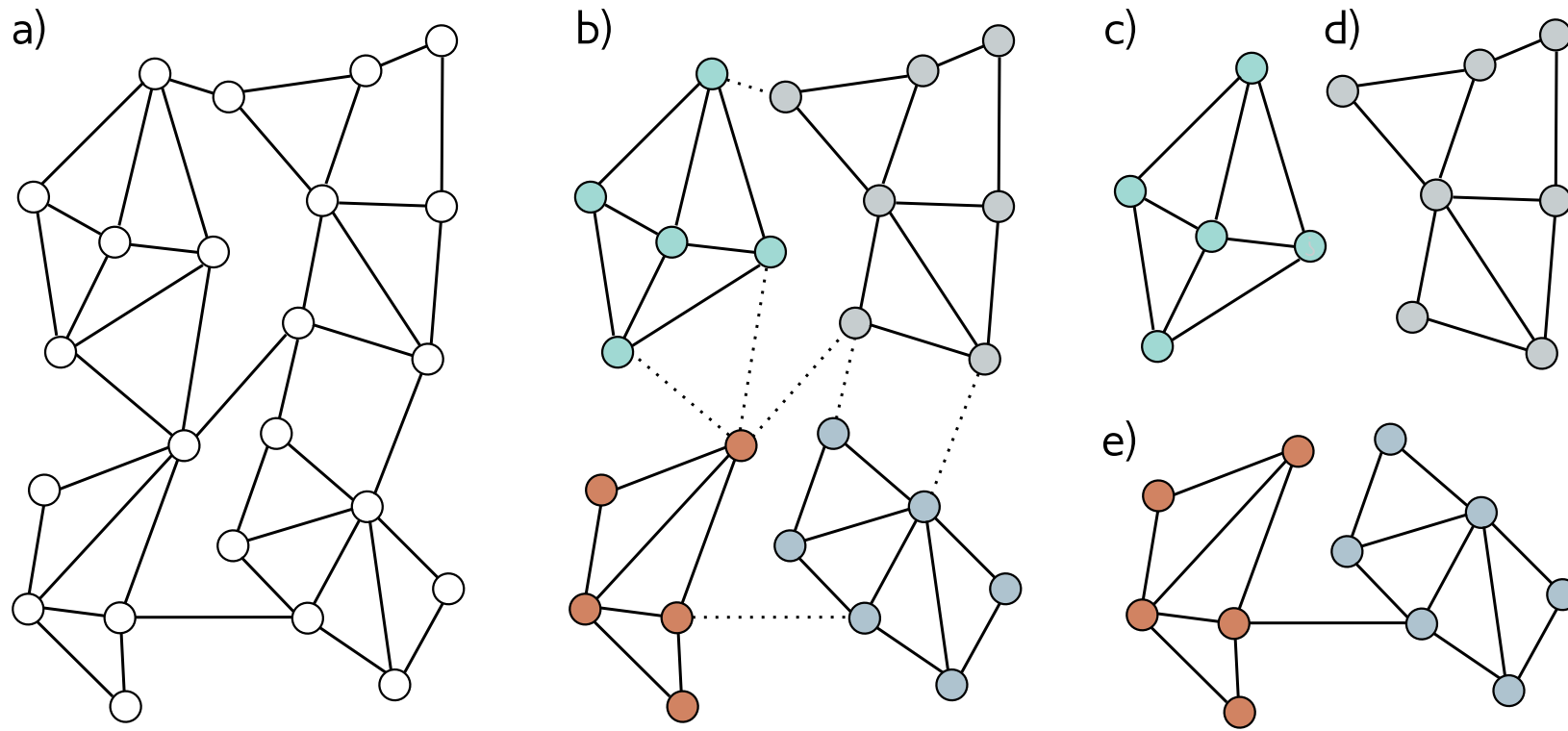
## Neighborhood Sampling:

Use max  $n$  of the neighbors.

Here  $n = 3$ .

See Notebook 13.3

# Graph Partitioning



Disconnect edges of the original to create maximally connected disjoint subsets

Split into train, test and validation sets and train just like in the inductive setting.

# Alternatives to Mean Pooling for Node Combinations

- **Diagonal enhancement:** current node is multiplied by  $(1 + \epsilon_k)$ , where  $\epsilon_k$  is a learned scalar for each layer

$$\mathbf{H}_{k+1} = \mathbf{a}[\beta_k \mathbf{1}^T + \mathbf{\Omega}_k \mathbf{H}_k (\mathbf{A} + (1 + \epsilon_k) \mathbf{I})]$$

- **Residual connections:** Include the current node in the sum

$$\mathbf{H}_{k+1} = \mathbf{a}[\beta_k \mathbf{1}^T + \mathbf{\Omega}_k \mathbf{H}_k \mathbf{A}] + \mathbf{H}_k$$

- **Mean aggregation:** take average instead of sum of neighbors

$$\text{agg}[n] = \frac{1}{|\text{ne}[n]|} \sum_{m \in \text{ne}[n]} \mathbf{h}_m$$

- **Kipf normalization:** downweight neighboring nodes with a lot of neighbors

$$\text{agg}[n] = \sum_{m \in \text{ne}[n]} \frac{h_m}{\sqrt{|\text{ne}[n]| |\text{ne}[m]|}}$$

- **Max pool aggregation:** element-wise max of all neighbors to current node

$$\text{agg}[n] = \max_{m \in \text{ne}[n]} [\mathbf{h}_m]$$

# Aggregation by Attention

Weights depend on data at the nodes.

Apply linear transform to current node:

$$\mathbf{H}'_k = \beta_k \mathbf{1}^T + \mathbf{\Omega}_k \mathbf{H}$$

Then the similarity  $s_{mn}$  of each transformed node embedding  $\mathbf{h}'_m$  to the transformed node embedding  $\mathbf{h}'_n$  is computed by concatenating the pairs, taking a dot product with a column vector  $\phi_k$  of learned parameters, and applying an activation function:

$$s_{mn} = a \left[ \phi_k^T \begin{bmatrix} \mathbf{h}'_m \\ \mathbf{h}'_n \end{bmatrix} \right]$$

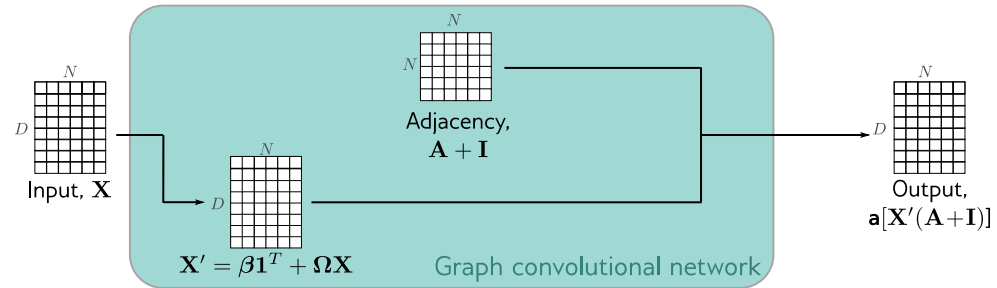
$$\mathbf{H}_{k+1} = \mathbf{a}[\mathbf{H}'_k \cdot \text{Softmask}[\mathbf{S}, \mathbf{A} + \mathbf{I}]]$$

# Softmask[S, A+I]

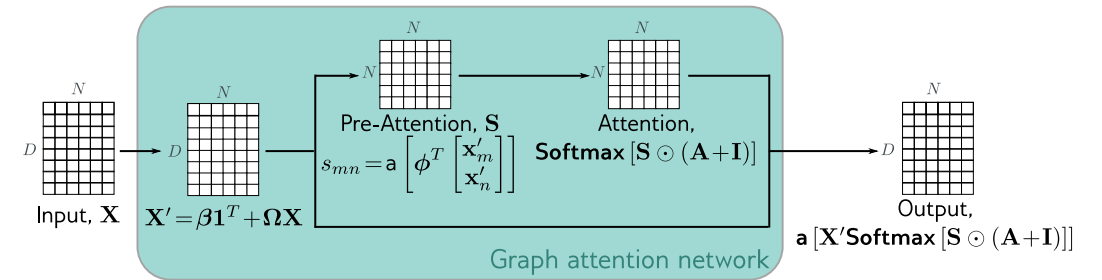
The function **Softmask[S, A+I]**

- computes the attention values by applying **softmax** operation separately to each column of its first argument S,
- but only after setting values where the second argument  $A + I$  is zero to negative infinity, so they do not contribute.
- This ensures that the attention to non-neighboring nodes is zero.

# Graph Attention

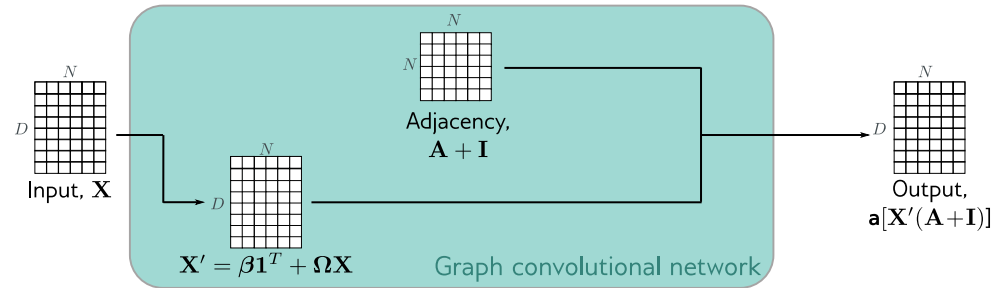


Regular graph convolution

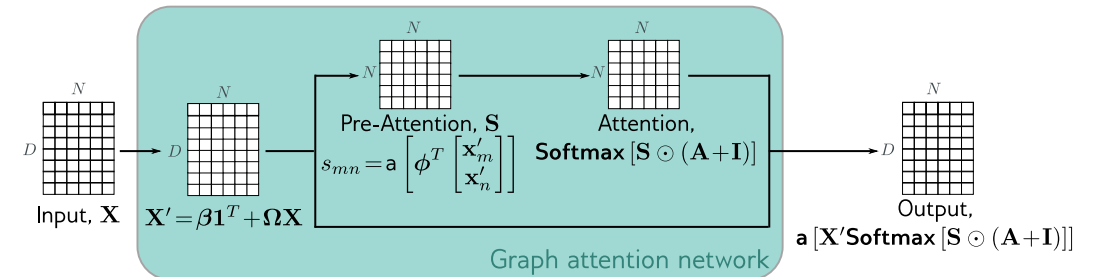


Graph attention

# Graph Attention



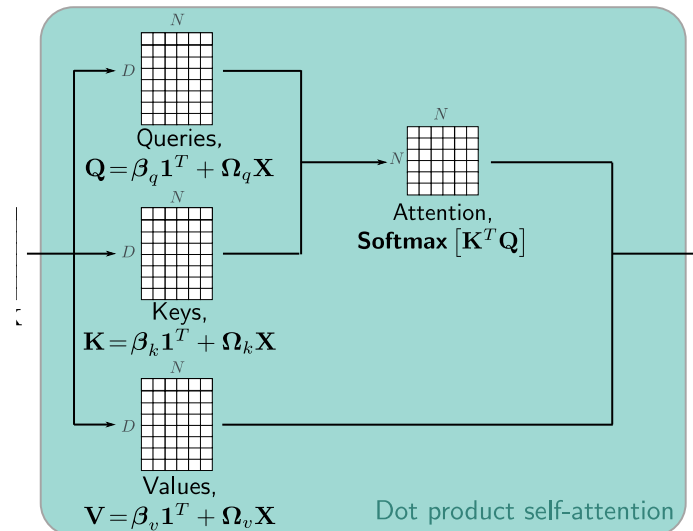
Regular graph convolution



Graph attention

*Similar to Transformer Self Attention, except*

- K, Q and V are all the same
- Different similarity measure
- Only attends to neighbors



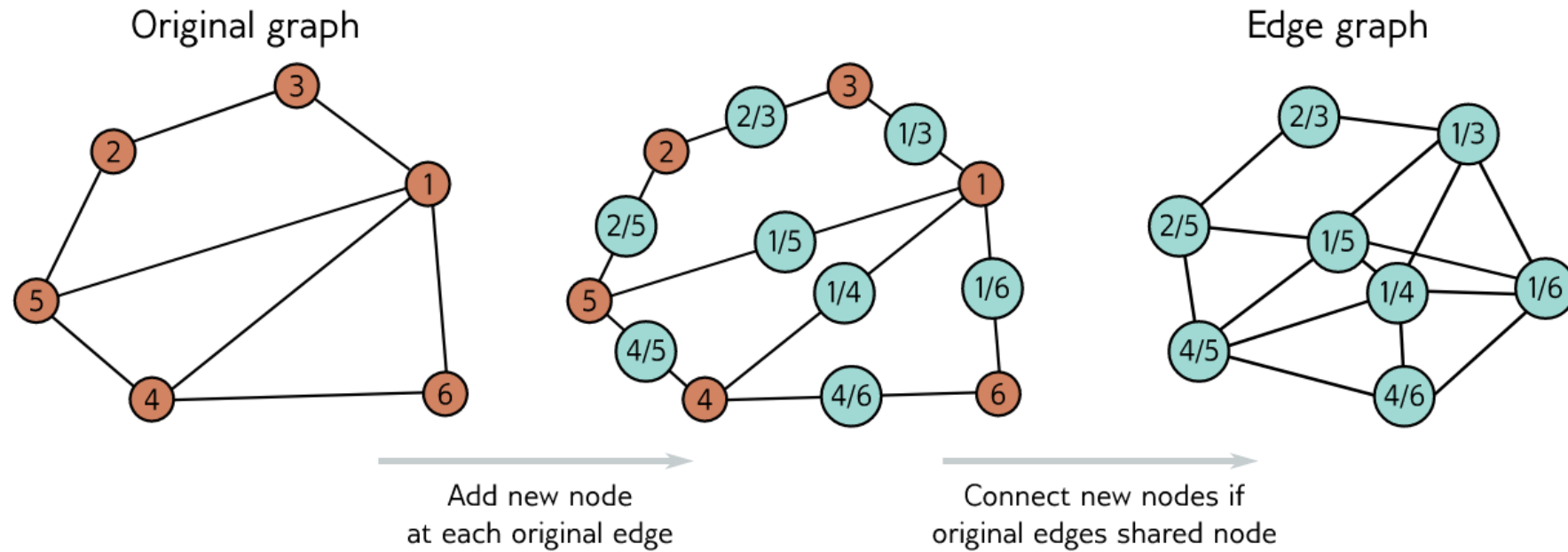
# Any Questions?



## Moving on

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# Edge Graphs



Handled by simple transformation from node graphs.

Then process as node graph.

Transform back to edge graph.

# Any Questions?



## **Moving on**

- Basic definition and examples
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