

Benchmarking Graph Neural Networks

Xavier Bresson

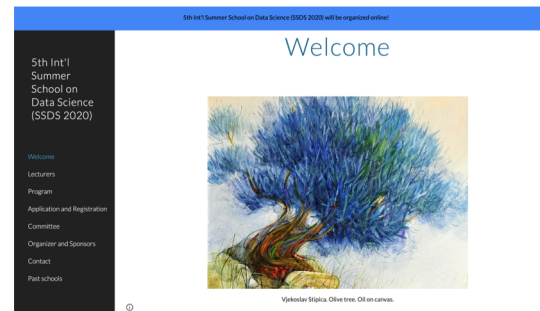


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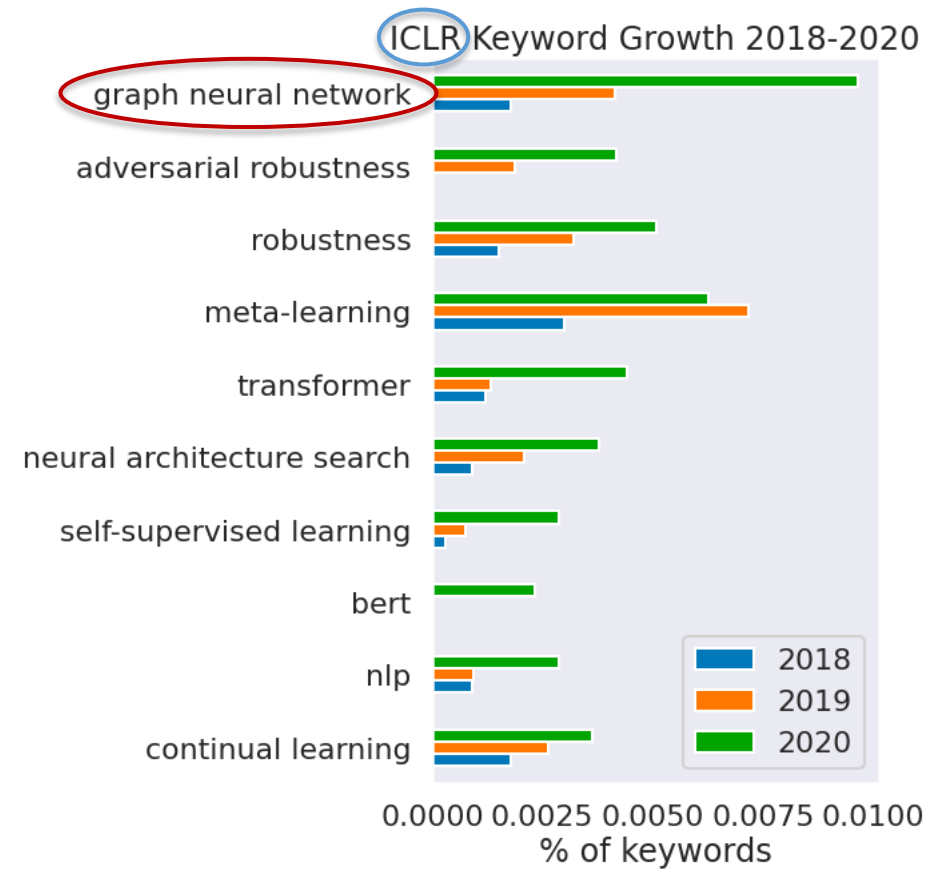
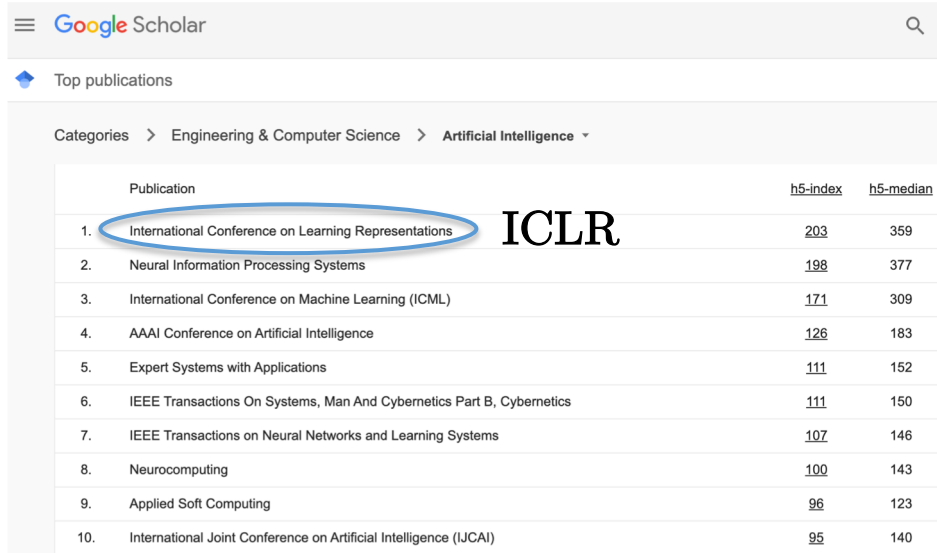
Summer School on Data Science (SSDS 2020)
Sven Loncaric, Tomislav Smuc, Vinko Zlatic, Tomislav Lipic

Sept 11th 2020



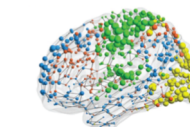
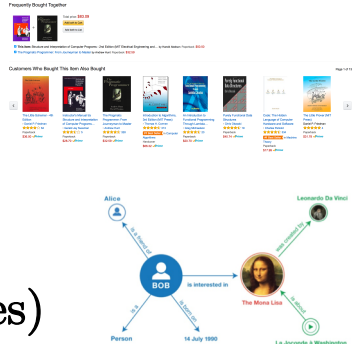
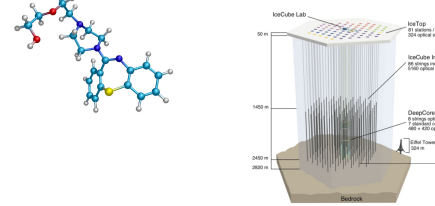
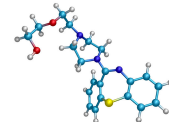
Graph Neural Networks (GNNs)

- GNNs have become the **standard toolkit** for analyzing and learning from **data on graphs**.
- **Hottest** machine learning topics in 2020.

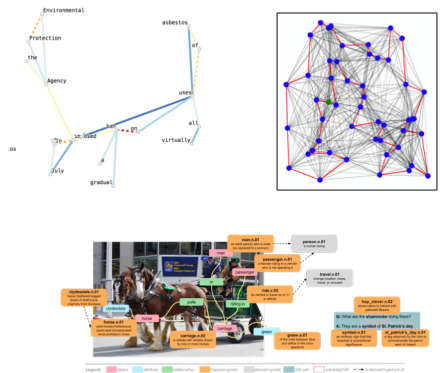


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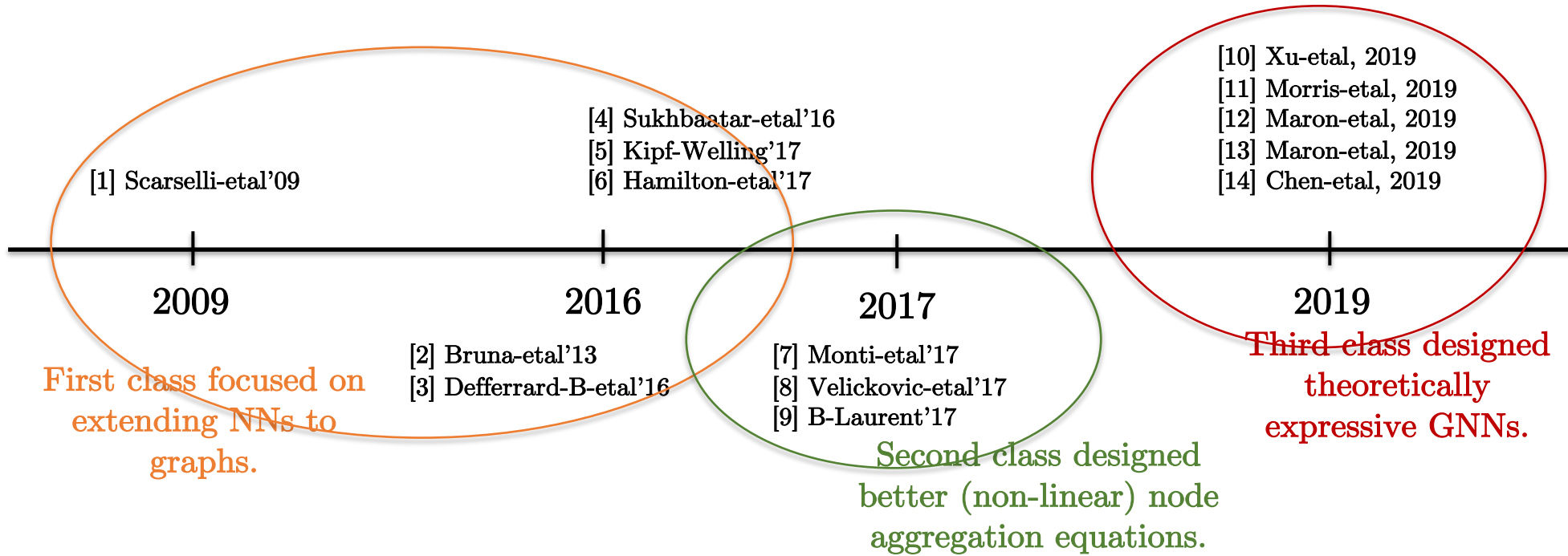
- Applicative domains :
 - Chemistry^[1,2] (generate new drugs and materials)
 - Physics^[3,4,5] (learn and accelerate physics)
 - Recommender systems^[6,7] (leverage consumer-product choices)
 - Social sciences^[8,9] (predict future collaborators, identify fake news)
 - Knowledge graphs^[10,11] (reasoning with entity-relationship)
 - Neuroscience^[12] (understanding brain mechanisms and neuro-degenerative diseases)
 - Computer Vision^[13] (scene understanding for visual reasoning)
 - Natural Language Processing^[14] (transformers)
 - Combinatorial Optimization^[15,16] (better/faster approximated solutions to NP-hard problems)



- [1] Duvenaud, Maclaurin, Iparraguirre, Bombarell, Hirzel, Aspuru-Guzik, Adams, Convolutional networks on graphs for learning molecular fingerprints, 2015
- [2] Gilmer, Schoenholz, Riley, Vinyals, Dahl, Neural message passing for quantum chemistry, 2017
- [3] Battaglia, Pascanu, Lai, Rezende, Danilo, Interaction networks for learning about objects, relations and physics, 2016
- [4] Cranmer, Xu, Battaglia, Ho, Learning symbolic physics with graph networks, 2019
- [5] Sanchez-Gonzalez, Godwin, Pfaff, Ying, Leskovec, Battaglia, Learning to simulate complex physics with graph networks, 2020
- [6] Monti, Bronstein, Bresson, Geometric matrix completion with recurrent multi-graph neural networks, 2017
- [7] Ying, He, Chen, Eksombatchai, Hamilton, Leskovec, Graph convolutional neural networks for web-scale recommender systems, 2018
- [8] Kipf, Welling, Semi-supervised classification with graph convolutional networks, 2017
- [9] Monti, Frasca, Eynard, Mannion, Bronstein, Fake news detection on social media using geometric deep learning, 2019
- [10] Schlichtkrull, Kipf, Bloem, VanDenBerg, Titov, Welling, Modeling relational data with graph convolutional networks, 2018
- [11] Chami, Wolf, Juan Sala Ravi, Re, Low-dimensional hyperbolic knowledge graph embeddings, 2020
- [12] Parisot, Ktena, Ferrante, Lee, Guerrero, Glocker, Rueckert, Disease prediction using graph networks: Application to Autism Spectrum Disorder and Alzheimer's disease, 2018
- [13] Johnson, Gupta, Fei-Fei, Image generation from scene graphs, 2018
- [14] Vaswani, Shazeer, Parmar, Uszkoreit, Jones, N Gomez, Kaiser, Polosukhin, Attention is all you need, 2017
- [15] Vinyals, Fortunato, Jaitly, Pointer, 2015
- [16] Bello, Pham, Le, Norouzi, Bengio, Neural combinatorial optimization with reinforcement learning, 2016



A Decade of GNNs



- [1] Scarselli, Gori, Tsoi, Hagenbuchner, Monfardini, The Graph Neural Network Model, 2009
- [2] Bruna, Zaremba, Szlam, LeCun, Spectral networks and locally connected networks on graphs, 2013
- [3] Defferrard, Bresson, Vandergheynst, Convolutional neural networks on graphs with fast localized spectral filtering, 2016
- [4] Sukhbaatar, Szlam, Fergus, Learning multiagent communication with backpropagation, 2016
- [5] Kipf, Welling, Semi-supervised classification with graph convolutional networks, 2017
- [6] Hamilton, Ying, Leskovec, Inductive representation learning on large graphs, 2017
- [7] Monti, Boschaini, Masci, Rodola, Svoboda, Bronstein, Geometric deep learning on graphs using mixture model cnns, 2017
- [8] Velickovic, Cucurull, Casanova, Romero, Lio, Bengio, Graph attention networks, 2017
- [9] Bresson, Laurent, Residual gated graph convnets, 2017
- [10] Xu, Hu, Leskovec, Jegelka, How powerful are graph neural networks?, 2019
- [11] Morris, Ritzert, Fey, Hamilton, Lenssen, Rattan, Grohe, Weisfeiler and leman go neural: Higher-order graph networks, 2019
- [12] Maron, Ben-Hamu, Shamir, Lipman, Invariant and equivariant graph networks, 2019
- [13] Maron, Ben-Hamu, Serviansky, Lipman, Provably powerful graph networks, 2019
- [14] Chen, Villar, Chen, Bruna, On the equivalence graph isomorphism testing and function approximation with gnnns, 2019

Developing powerful
GNNs for real-world
adoption of graph
deep learning.

Tracking Progress

- The field has grown extensively in the last three years 😊
- Unfortunately, evaluating the effectiveness of new architectures/ideas has become difficult for two major reasons :
 - We have been evaluated progress on small datasets such as Cora^[1], Citeseer^[2] and TU^[3].
 - Cora is a single graph of 2.7K nodes, TU-IMDB has 1.5K graphs with 13 nodes on average, and TU-MUTAG has 188 molecules with 18 nodes.
 - Simple or graph-agnostic architectures provide statistically same performance as complex architectures^[4,5].
 - We have not rigorously enforced standardized experimental settings for fair comparisons between models^[6].
- It has become critical to solve these issues.

[1] McCallum, Nigam, Rennie, Seymore, Automating the construction of internet portals with machine learning, 2000

[2] Getoor, Link-based classification, 2005

[3] Kersting, Kriege, Morris, Mutzel, Neumann, Benchmark data sets for graph kernels, 2020

[4] Hoang, Maehara, Revisiting graph neural networks, 2019

[5] Chen, Bian, Sun, Are powerful graph neural nets necessary? a dissection on graph classification, 2019

[6] Errica, Podda, Bacciu, Micheli, A fair comparison of graph neural networks for graph classification, 2019

Benchmarking GNNs

- Our motivation : Identify and quantify what types of architectures, first principles are universal, generalizable, and scalable to larger and more challenging datasets.
- Benchmarking has been beneficial for driving progress, identifying essential ideas, and solving domain-specific problems^[1].
 - The **2012 ImageNet challenge**^[2] has provided a benchmark dataset that has triggered the deep learning revolution.
- Challenges :
 - Developing a **rigorous experimental setting** for fair comparisons, and being reproducible,
 - Using **datasets that can statistically separate model performance**,
 - Benchmarking **distinct fundamental graph tasks**.

[1] Weber, Saelens, Cannoodt, Soneson, Hapfelmeier, Gardner, Boulesteix, Saeys, Robinson, Essential guidelines for computational method benchmarking, 2019

[2] Deng, Dong, Socher, Li, Li, Fei-Fei, ImageNet: A Large-Scale Hierarchical Image Database, 2009

Outline

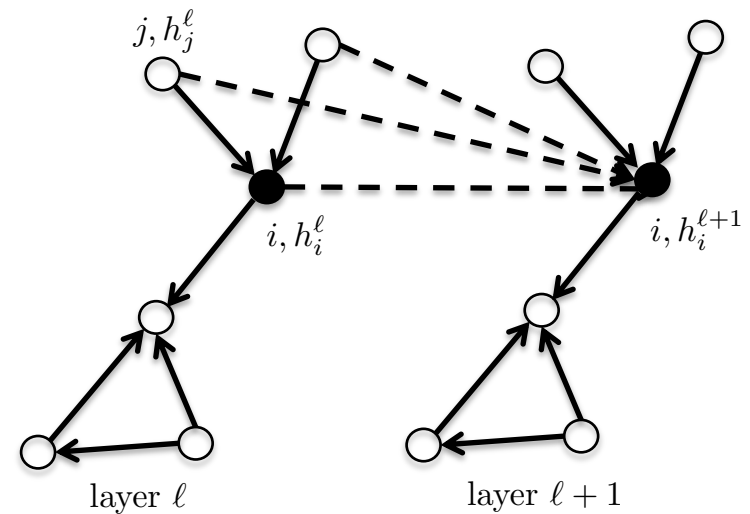
- Motivation
- Message-Passing GCNs
- Weisfeiler-Lehman GNNs
- Graph-Agnostic GNNs
- Datasets
- Infrastructure and Experimental Setting
- Benchmarking Results
- Laplacian Positional Encodings
- Link Prediction with Edge Representation
- Conclusion

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MP-GCNs

- Message-passing **Graph Convolutional Networks**
- What are the **minimal inner structures** to design a message-passing node aggregation function f_{GCN} ?
 - **Invariant by node permutation** (equivariant/invariant layers^[1]).
 - Independent of graph size n and **neighborhood size**.
 - **Locality** (local reception field - only neighbors are considered).
 - **Graph convolution operator** (weight sharing across graph).
 - Linear complexity $O(E)$, E being the number of edges (reduces to $O(n)$ for sparse graphs).



$$h_i^{\ell+1} = f_{\text{GCN}}(h_i^\ell, \{h_j^\ell : j \in \mathcal{N}_i\})$$

[1] Maron, Ben-Hamu, Shamir, Lipman, Invariant and equivariant graph networks, 2019

Isotropic GCNs

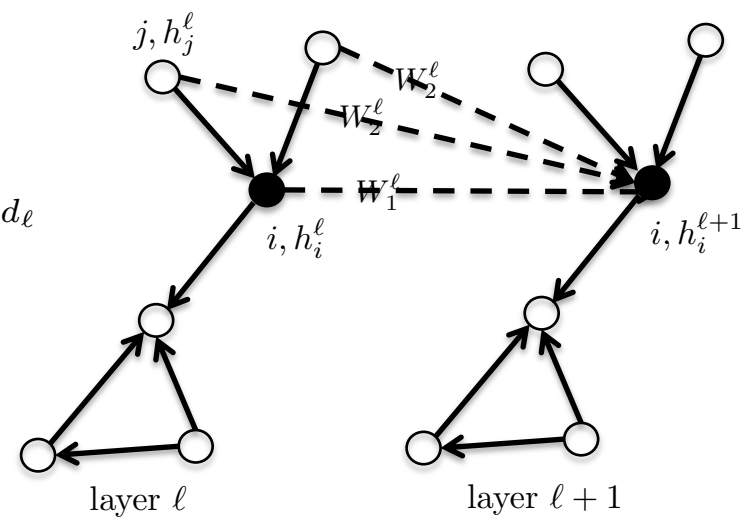
- We consider a class of **isotropic GCNs** when the node update equation **treats every “edge direction” equally**, i.e. each neighbor contributes equally to the update of the central node :

$$h_i^{\ell+1} = \sigma \left(W_1^\ell h_i^\ell + \sum_{j \in \mathcal{N}_i} W_2^\ell h_j^\ell \right), \text{ Sum/mean/max}$$

$$h^{\ell+1} \in \mathbb{R}^{n \times d_{\ell+1}}, \quad h^\ell \in \mathbb{R}^{n \times d_\ell}, \quad W_{1,2}^\ell \in \mathbb{R}^{d_{\ell+1} \times d_\ell}$$

- Models :**

- Vanilla GCNs^[1,2]
- GraphSage^[3]
- ChebNets^[4]



- Isotropic GCNs are **limited to learn** template weights for the central node h_i (weight W_1) and the neighbors h_j (weight W_2).

[1] Sukhbaatar, Szlam, Fergus, Learning multiagent communication with backpropagation, 2016

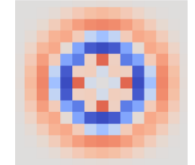
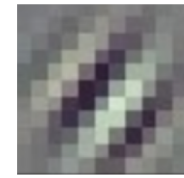
[2] Kipf, Welling, Semi-supervised classification with graph convolutional networks, 2017

[3] Hamilton, Ying, Leskovec, Inductive representation learning on large graphs, 2017

[4] Defferrard, Bresson, Vandergheynst, Convolutional neural networks on graphs with fast localized spectral filtering, 2016

Anisotropic GCNs

- Standard ConvNets^[1] produce **anisotropic filters** because Euclidean grids have **directional structure** (up, down, left, right).
- GCNs such as vanilla GCNs, GraphSage, ChebNets compute isotropic filters as there is **no notion of direction on arbitrary graphs**.
- How to **get anisotropy back** ?
 - **Natural edge features**^[2,3] if available (e.g. different bond connections between atoms in molecular graphs or distinct edge attributes in knowledge graphs).
 - **Learn anisotropy** with a mechanism invariant by index permutation and can treat neighbors differently :
 - MoNets^[4] with edge degrees
 - GatedGCNs^[5] with edge gates
 - GAT^[6] with attention mechanism^[7]



[1] LeCun Bottou Bengio, Haffner, Gradient-based learning applied to document recognition, 1998

[2] Gilmer, Schoenholz, Riley, Vinyals, Dahl, Neural message passing for quantum chemistry, 2017

[3] Bresson, Laurent, A Two-Step Graph Convolutional Decoder for Molecule Generation, 2019

[4] Monti, Boscaini, Masci, Rodolà, J. Svoboda, M. Bronstein, Geometric deep learning on graphs and manifolds using mixture model CNNs, 2016

[5] Bresson, Laurent, Residual gated graph convnets, 2017

[6] Velickovic, Cucurull, Casanova, Romero, Lio, Bengio, Graph Attention Networks, 2018

[7] Bahdanau, Cho, Bengio, Neural machine translation by jointly learning to align and translate, 2015

Anisotropic GCNs

- When the update equation **treats every edge direction differently**, we instantiate anisotropic GCNs with an aggregation equation of the form :

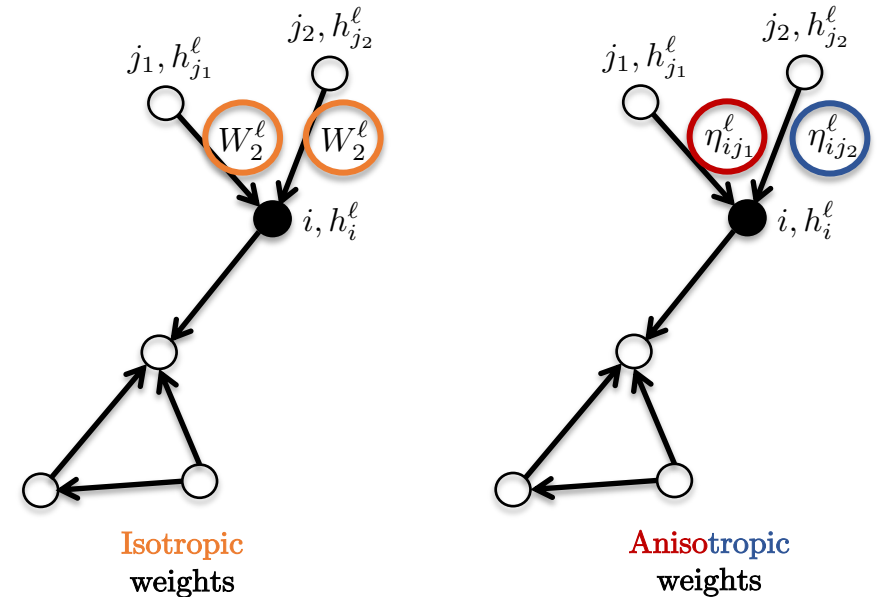
$$h_i^{\ell+1} = \sigma \left(W_1^\ell h_i^\ell + \sum_{j \in \mathcal{N}_i} \eta_{ij}^\ell W_2^\ell h_j^\ell \right), \quad h^{\ell+1} \in \mathbb{R}^{n \times d_{\ell+1}}, \quad h^\ell \in \mathbb{R}^{n \times d_\ell}, \quad W_{1,2}^\ell \in \mathbb{R}^{d_{\ell+1} \times d_\ell},$$

where $\eta_{ij}^\ell = g_W(h_i^\ell, h_j^\ell)$ is a parameterized function which weights are learned during training.

Attention mechanism
in 1-hop
neighborhood

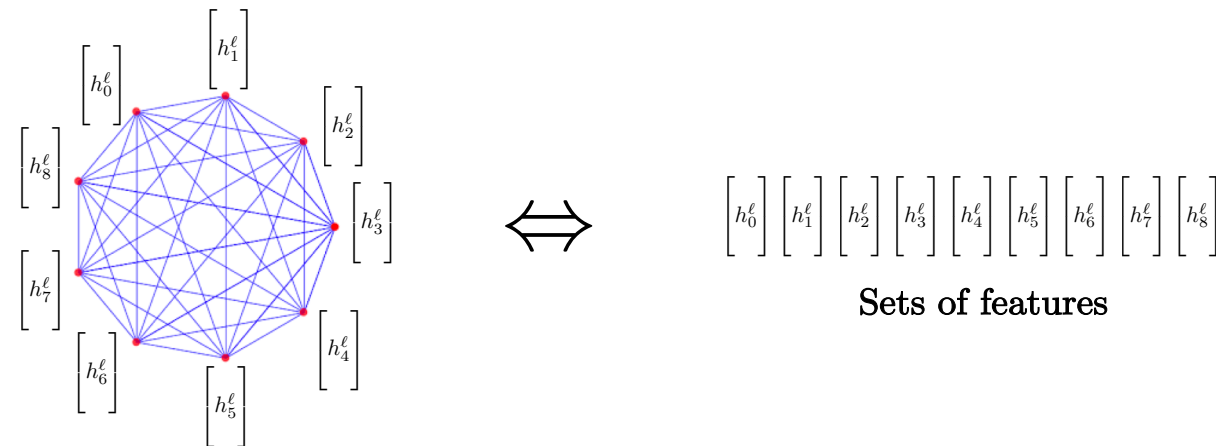
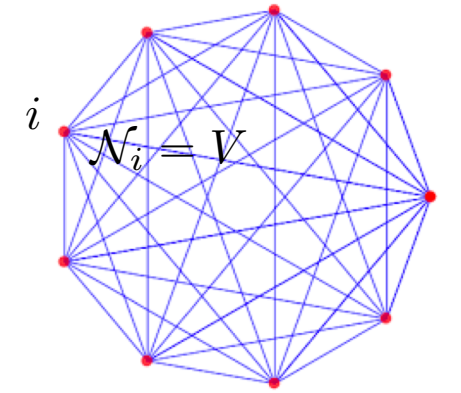
$$\begin{aligned} \eta_{ij}^\ell &= g_{W_{QK}}(h_i^\ell, h_j^\ell) \\ &= \text{Softmax}_{\mathcal{N}_i}(e_{ij}^\ell) = \frac{\exp(e_{ij}^\ell)}{\sum_{j' \in \mathcal{N}_i} \exp(e_{ij'}^\ell)} \\ &\text{with } e_{ij}^\ell = (W_Q^\ell h_i^\ell)^T (W_K^\ell h_j^\ell), \quad W_Q, W_K \in \mathbb{R}^{d_{\ell+1} \times d_\ell} \end{aligned}$$

Query Key



Transformers

- Transformers is a special case of GCNs when the graph is fully connected.
 - The neighborhood \mathcal{N}_i is the whole graph.
- What does it mean to have a fully connected graph ?
 - There is no particular graph structure that can be used. It becomes useless to talk about graphs (as each data point is connected to all other points).
 - It would be better to talk about sets.
 - Transformers are set neural networks, which analyze bags of features.



[1] A Vaswani, N Shazeer, N Parmar, J Uszkoreit, L Jones, A. Gomez, L. Kaiser, I. Polosukhin, Attention is all you need, 2017

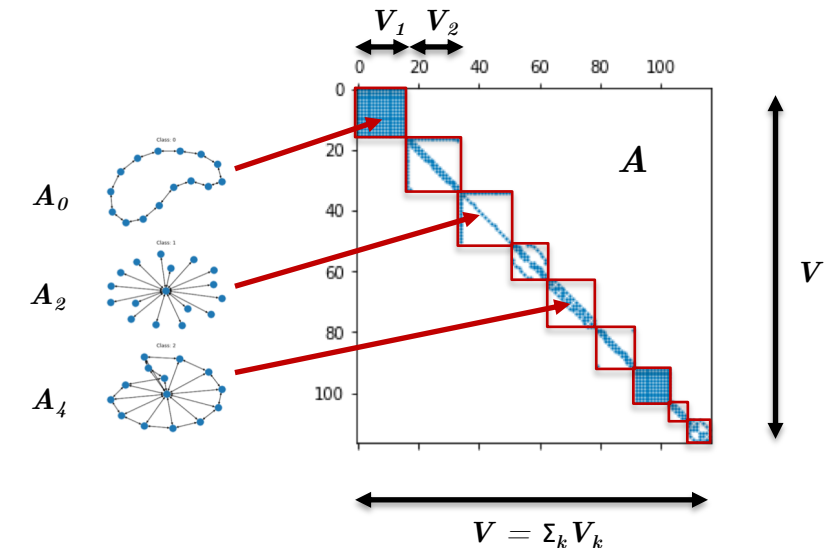
Batch Normalization and Residual Connections

- GCNs benefit from BN^[1,2] and RC^[1,3] :
 - Speed up learning process.
 - Improve performance with better generalization.

- How to batch graphs of different sizes ?
 - For images of size $B \times V_x \times V_y \times d$, BN is performed along the batch dimension, i.e. dim=0.
 - For graphs, a (big) sparse block diagonal matrix A of size $V \times d$ is first built from K matrices A_k and BN is carried out along the node direction, i.e. dim=0.

$$h_i^{\ell+1} = h_i^\ell + \sigma\left(\text{BN}\left(\hat{h}_i^{\ell+1}\right)\right)$$

$$\hat{h}_i^{\ell+1} = f_{\text{GCN}}\left(h_i^\ell, \{h_j^\ell : j \in \mathcal{N}_i\}\right)$$



[1] Bresson, Laurent, Residual gated graph convnets, 2017

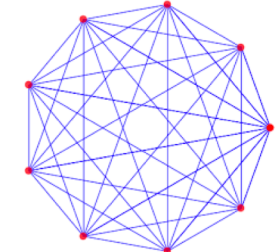
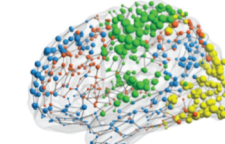
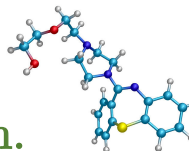
[2] Ioffe, Szegedy, Batch normalization: Accelerating deep network training by reducing internal covariate shift, 2015

[3] Li, Muller, Qian, Delgadillo, Abualshour, Thabet, Ghanem, Deepgcns: Making gcns go as deep as cnns, 2019

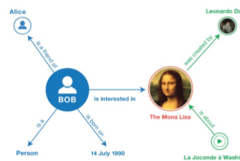
Sparsity and Local Computations

- GCNs leverage graph sparsity :

- Sparsity is a good inductive bias for generalization.
- Sparse vs full graphs.



vs.



Sparse graphs

Fully connected
graph

- Local computations :

- Node update equation is local, only depending on neighborhood \mathcal{N}_i of node i , and independent of graph size n , making the space/time complexity $O(n)$ for sparse graphs.
- GCNs are highly parallelizable on GPUs, and sparse matrix multiplications are efficiently implemented via GNN libraries s.a. DGL^[1], PyG^[2], Spektral^[3].

$$h_i^{\ell+1} = f_{\text{GCN}}(h_i^\ell, \{h_j^\ell : j \in \mathcal{N}_i\})$$

DGL



PyTorch
geometric



[1] Wang-et.al, Deep graph library: Towards efficient and scalable deep learning on graphs, 2019

[2] Fey, Lenssen, Fast graph representation learning with pytorch geometric, 2019

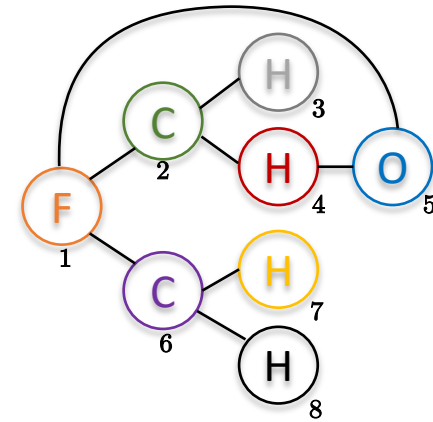
[3] Grattarola, Alippi, Graph Neural Networks in TensorFlow and Keras with Spektral, 2020

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- **Weisfeiler-Lehman GNNs**
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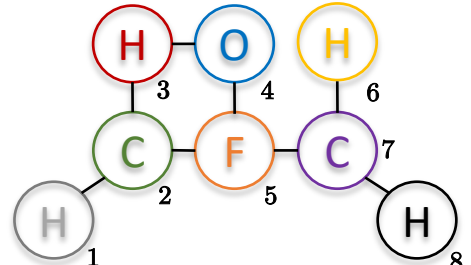
Weisfeiler-Lehman GNNs

- Motivation : Characterize expressivity power of GNNs with graph isomorphism.
- Graph isomorphism : Two graphs are isomorphic if there exists an index permutation between the nodes that preserves node adjacencies.
- Determining whether two graphs are isomorphic is NP-intermediate : It is not known if a polynomial time algorithm exists, or the problem is NP-hard.
- Weisfeiler-Lehman (WL) proposed an algorithm^[1] to test if two graphs are not isomorphic. However, the WL test is not sufficient to guarantee that two graphs are isomorphic.



Graph 1

\Updownarrow Isomorphic graphs



Graph 2

[1] B Weisfeiler, A Lehman, A reduction of a graph to a canonical form and an algebra arising during this reduction, 1968

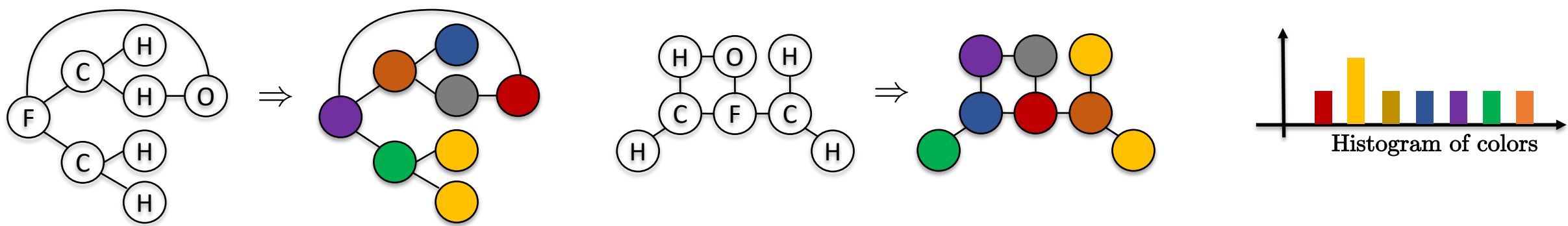
WL Test^[1]

- Idea : Design an injective “coloring” function f_{WL} :

$$f_{WL}(c_i^\ell, \{c_j^\ell\}_{j \in N_i}) = c_i^{\ell+1}$$

iteration
Multiset function Multiset (set of unordered and repetitive elements)

- WL algorithm **iteratively** applies function f_{WL} until no new colors are created.
- This produces a canonical representation of a graph as a **histogram of colors**.
- If **two graphs have different color histograms** (up to all color permutations) then the two graphs are **guaranteed to be non-isomorphic**.



[1] B Weisfeiler, A Lehman, A reduction of a graph to a canonical form and an algebra arising during this reduction, 1968

Graph Isomorphism Networks (GINs)^[1]

- Goal : Design a GNN as expressive as the WL test to distinguish non-isomorphic graphs.
- Node aggregation of the form :

$$f_{\text{NN}}(h_i^\ell, \{h_j^\ell\}_{j \in \mathcal{N}_i}) = (1 + \varepsilon)g(h_i^\ell) + \sum_{j \in \mathcal{N}_i} g(h_j^\ell)$$

irrational injective sum

- It is difficult to design an analytical injective function \Rightarrow **MLP is used to approximate g** (existence guaranteed by the universal approximation theorem) :

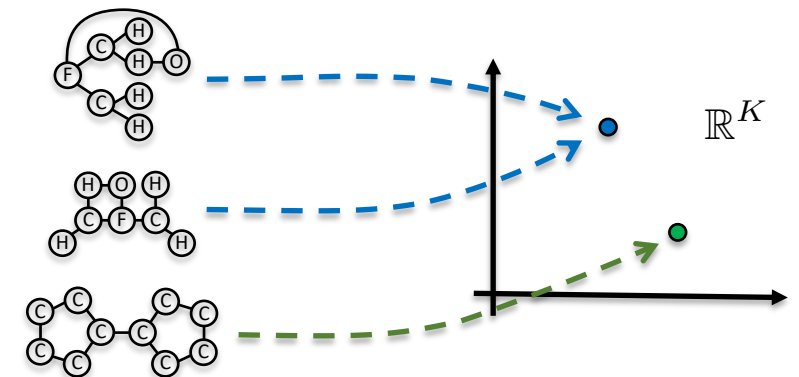
$$h_i^{\ell+1} = f_{\text{GIN}}(h_i^\ell, \{h_j^\ell\}_{j \in \mathcal{N}_i}) = \text{MLP}^\ell \left((1 + \varepsilon)h_i^\ell + \sum_{j \in \mathcal{N}_i} h_j^\ell \right)$$

- Graph **readout** function must also be **injective** :

$$h_G = \text{MLP} \left(\sum_{i \in V} h_i^L \right) \in \mathbb{R}^K$$

Sum function

- Pioneer work with^[2] on GNN expressivity.

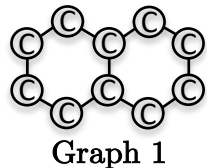


[1] Xu, Hu, Leskovec, Jegelka, How powerful are graph neural networks?, 2019

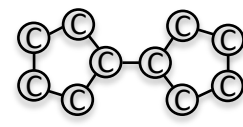
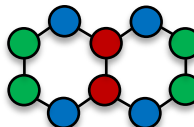
[2] Morris, Ritzert, Fey, Hamilton, Lenssen, Rattan, Grohe, Weisfeiler and leman go neural: Higher-order graph networks, 2019

Limitation

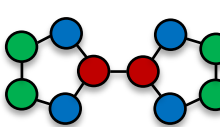
- WL test^[1] is not a sufficient condition, it can fail to differentiate non-isomorphic graphs :
 - Example of two graphs with same color signatures but not isomorphic :



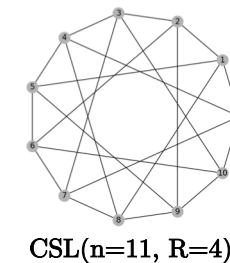
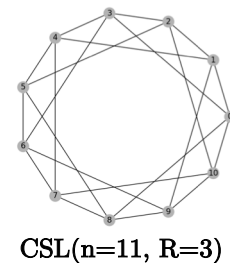
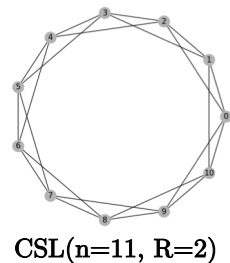
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⇒



- Another example w/ Circular Skip Length (CSL) graphs^[2] (a circular graph with skip connections for each node to R -hop nodes) :



Nodes are
isomorphic

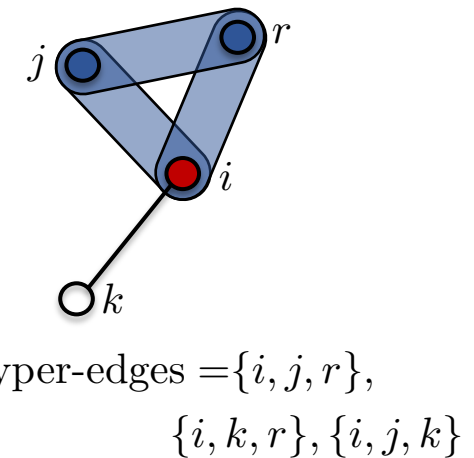
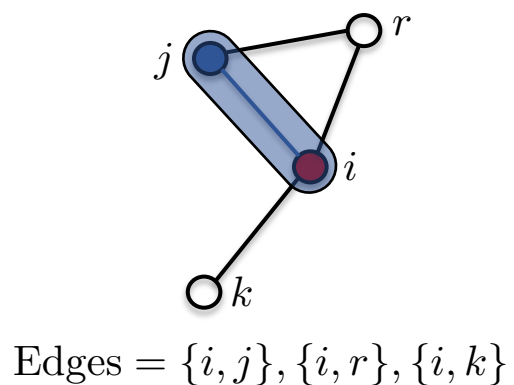
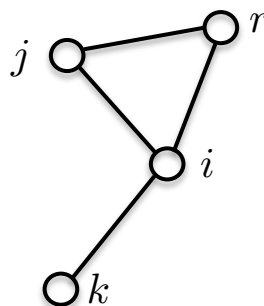
- Can we improve the expressivity of the original WL test ?

[1] B Weisfeiler, A Lehman, A reduction of a graph to a canonical form and an algebra arising during this reduction, 1968

[2] Murphy, Srinivasan, Rao, Ribeiro, Relational pooling for graph representations, 2019

k -WL Test

- Can we design better WL tests ?
 - Original WL test uses **2-tuple of nodes** to produce colors. To produce more colors and improve expressivity of WL tests, **k -tuple of nodes** with $k \geq 3$ can be used (higher-order interactions between nodes).



2-tuple of nodes can distinguish non-isomorphic graphs with the **1-WL/2-WL** tests.

3 -tuple of nodes can distinguish non-isomorphic graphs with the **3 -WL** test.

Equivariant GNNs^[1]

- How to design GNNs with the same **expressivity power** as the k -WL test ?

- Let us define a **k -order equivariant GNNs** :

$$y = m_{W^{L+1}} \circ g_{W^L} \circ \sigma \circ f_{W^{L-1}} \circ \sigma \circ f_{W^{L-2}} \circ \dots \sigma \circ f_{W^0} \circ h_0$$

with $k = \max_{\ell \in [0, L-1]} k_\ell$

Permutation
 $f_W(P \circ h) = P \circ f_W(h)$
 Equivariant linear layers
 $g_W(P \circ h) = g_W(h) \in \mathbb{R}^K, K \geq 1$
 Invariant linear layer

and $f_{W^\ell} : \mathbb{R}^{n^{k_\ell} \times d_\ell} \rightarrow \mathbb{R}^{n^{k_\ell+1} \times d_{\ell+1}}$

- Theorem^[1] : There exist **k -order E-GNNs** that can distinguish non-isomorphic graphs with the **k -WL test**.
 - However, k -order E-GNNs require **$O(n^k)$** memory/speed complexities.
 - Note that we need at least $k=3$ (hence $O(n^3)$) to be more powerful than GINs as GINs have the discriminative power of the 1-WL/2-WL tests^[2].

[1] Maron, Ben-Hamu, Shamir, Lipman, Invariant and equivariant graph networks, 2019

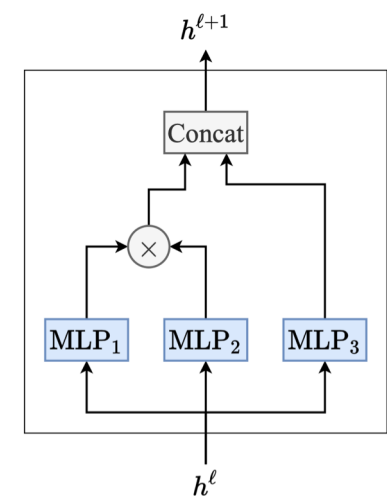
[2] Cai, Furer, Immerman, An optimal lower bound on the number of variables for graph identification, 1992

3-WL GNNs^[1]

- How to design GNNs that are **3-WL expressive** but do not require $O(n^3)$ memory/speed complexities?
- **3-WL GNNs** : To achieve **higher interactions** between nodes, it is sufficient to **multiply second-order tensors** feature-wise.
 - Theorem^[1] : There **exist** 3-WL GNNs as expressive as the 3-WL test.
 - **Memory is quadratic** $O(n^2)$ but matrix multiplication implies $O(n^3)$ speed.
 - Also, matrix multiplication densifies sparse matrix.
 - This is the simplest, most scalable and most expressive GNNs in terms of 3-WL test.
- **3-WL GNN update layer** :

$$h^{\ell+1} = \text{Concat}(m_{W_1^\ell}(h^\ell) \cdot m_{W_2^\ell}(h^\ell), m_{W_3^\ell}(h^\ell))$$

where $h^{\ell+1} \in \mathbb{R}^{n \times n \times d_{\ell+1}}$, $h^\ell \in \mathbb{R}^{n \times n \times d_\ell}$, $W_1, W_2, W_3 \in \mathbb{R}^{2 \times d_\ell \times d_{\ell+1}}$



[1] Maron, Ben-Hamu, Serviansky, Lipman, Provably powerful graph networks, 2019

RingGNNs[1]

- Related method to 3-WL GNNs to design more expressive GNNs than GINs.

- Higher-order interaction between nodes is produced by multiplying equivariant linear layers^[2].
- RingGNN update layer :

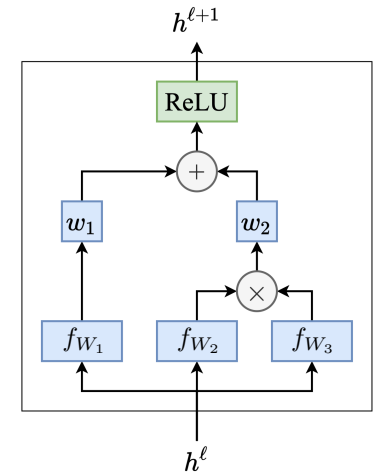
$$h^{\ell+1} = \sigma(w_1^\ell f_{W_1^\ell}(h^\ell) + w_2^\ell f_{W_2^\ell}(h^\ell) \cdot f_{W_3^\ell}(h^\ell)),$$

where $h^{\ell+1} \in \mathbb{R}^{n \times n \times d_{\ell+1}}$, $h^\ell \in \mathbb{R}^{n \times n \times d_\ell}$, $w_{1,2}^\ell \in \mathbb{R}$, $W_1, W_2, W_3 \in \mathbb{R}^{d_\ell \times d_{\ell+1} \times 17}$,

where f_W are the equivariant linear layers defined as :

$$(f_W(h))_{\cdot,\cdot,q'} = \sum_{k=1}^{15+2} \sum_{q=1}^{d_\ell} W_{k,q,q'} f_k(h_{\cdot,\cdot,q}) \in \mathbb{R}^{n \times n \times d_{\ell+1}}$$

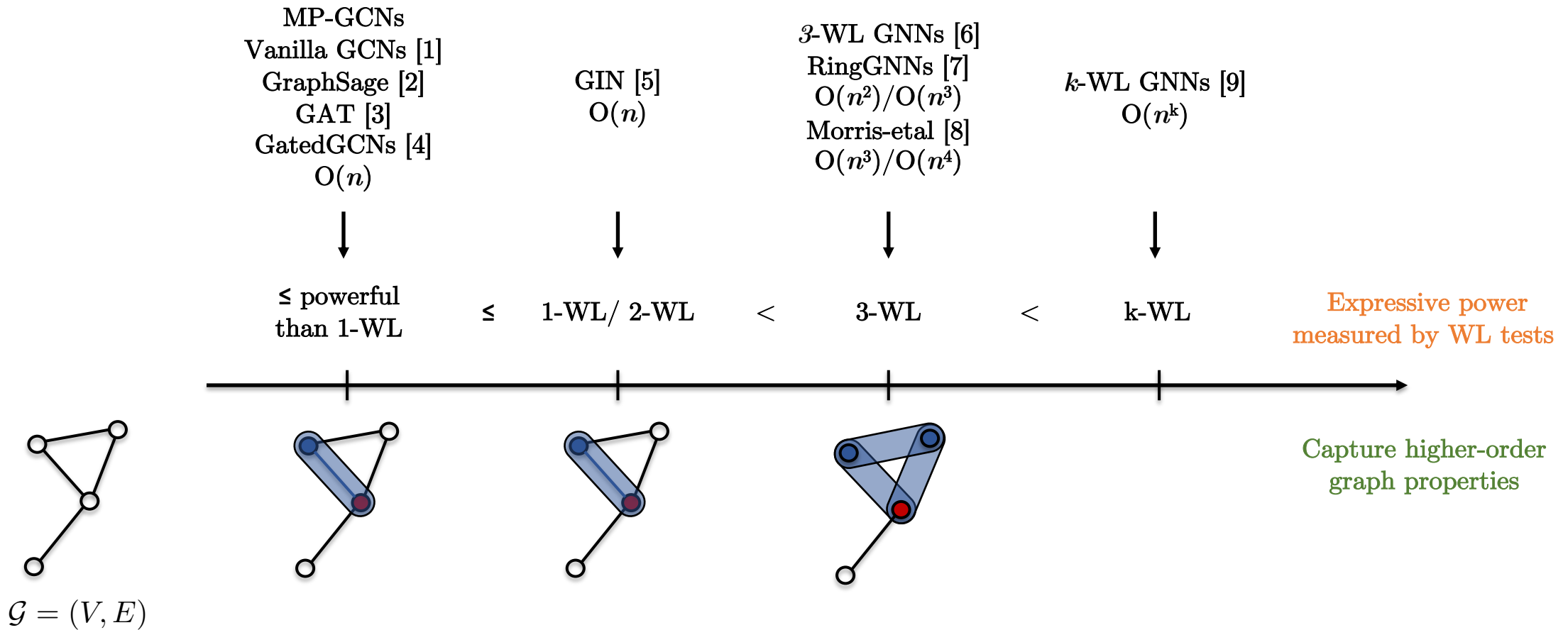
and f_k are all possible 15 equivariant linear functions $f_k : \mathbb{R}^{n \times n} \rightarrow \mathbb{R}^{n \times n}$ for a given tensor $h \in \mathbb{R}^{n \times n}$, and 2 bias functions.



[1] Chen, Villar, Chen, Bruna, On the equivalence between graph isomorphism testing and function approximation with gnns, 2019

[2] Maron, Ben-Hamu, Shamir, Lipman, Invariant and equivariant graph networks, 2019

Expressivity Hierarchy w.r.t. WL Test



- [1] Kipf, Welling, Semi-supervised classification with graph convolutional networks, 2017
- [2] Hamilton, Ying, Leskovec, Inductive representation learning on large graphs, 2017
- [3] Velickovic, Cucurull, Casanova, Romero, Lio, Bengio, Graph attention networks, 2017
- [4] Bresson, Laurent, Residual gated graph convnets, 2017
- [5] Xu, Hu, Leskovec, Jegelka, How powerful are graph neural networks?, 2019
- [6] Maron, Ben-Hamu, Serviansky, Lipman, Provably powerful graph networks, 2019
- [7] Chen, Villar, Chen, Bruna, On the equivalence between graph isomorphism testing and function approximation with gnnns, 2019
- [8] Morris, Ritzert, Fey, Hamilton, Lenssen, Rattan, Grohe, Weisfeiler and leman go neural: Higher-order graph neural networks, 2019
- [9] Maron, Ben-Hamu, Shamir, Lipman, Invariant and equivariant graph networks, 2019

Outline

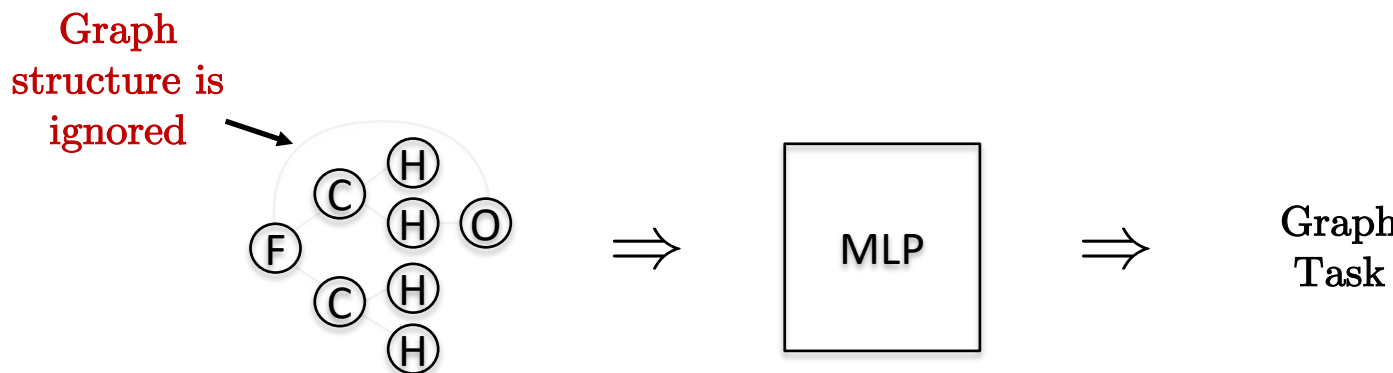
- Motivation
- Message-Passing GCNs
- Weisfeiler-Lehman GNNs
- **Graph-Agnostic GNNs**
- Datasets
- Infrastructure and Experimental Setting
- Benchmarking Results
- Laplacian Positional Encodings
- Link Prediction with Edge Representation
- Conclusion

MLP Baseline

- **Graph-agnostic NNs** : As a sanity check, we compare GNNs to a simple MLP network which updates each node independent of one-other :

$$h_i^{\ell+1} = \sigma(W^\ell h_i^\ell), \quad h^{\ell+1} \in \mathbb{R}^{n \times d_{\ell+1}}, \quad h^\ell \in \mathbb{R}^{n \times d_\ell}, \quad W^\ell \in \mathbb{R}^{d_{\ell+1} \times d_\ell}$$

and passes these features to the task-based readout layer.



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Datasets

- Main issue with prevalent datasets : **Small sizes**
- Ideal dataset : Representative, realistic, and medium/large-scale size.
- **Challenges :**
 - What theoretical tool to **define the quality of dataset** or validate its statistical representativeness for a given task ?
 - Several **arbitrary choices** when preparing graphs, such as node and edge features. For example, e-commerce product features.
 - **Appropriate size** may depend on the task complexity as well as the dimensionality and statistics of underlying data.
- The recent **Open Graph Benchmark^[1]** (OGB) project is a much needed initiative to tackle these challenges.
 - OGB offers a collection of medium/large-scale real-world graph datasets and evaluation protocols, with an emphasis on out-of-distribution generalization performance through meaningful data splits.

[1] Hu, Fey, Zitnik, Dong, Ren, Liu, Catasta, Leskovec, Open graph benchmark: Datasets for machine learning on graphs, 2020

Proposed Datasets

- Appropriate datasets :
 - Datasets that are able to statistically separate the performance of GNNs.
- Summary of the 7 medium-scale datasets :

Domain & Construction	Dataset	#Graphs	#Nodes	Total #Nodes	Task
Chemistry: Real-world molecular graphs	ZINC	12K	9-37	277,864	Graph Regression
Mathematical Modelling: Artificial graphs generated from Stochastic Block Models	PATTERN	14K	44-188	1,664,491	Node Classification
	CLUSTER	12K	41-190	1,406,436	
Computer Vision: Graphs constructed with SLIC super-pixels of images	MNIST	70K	40-75	4,939,668	Graph Classification
	CIFAR10	60K	85-150	7,058,005	
Combinatorial Optimization: Uniformly generated artificial Euclidean graphs	TSP	12K	50-500	3,309,140	Edge Classification
Social Networks: Real-world citation graph	COLLAB	1	235,868	235,868	Edge Classification
Circular Skip Links: Isomorphic graphs with same degree	CSL	150	41	6,150	Graph Classification

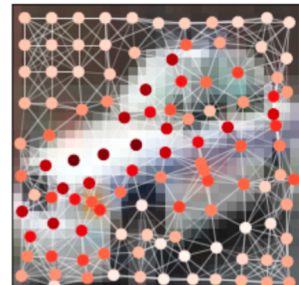
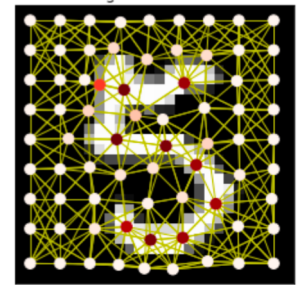
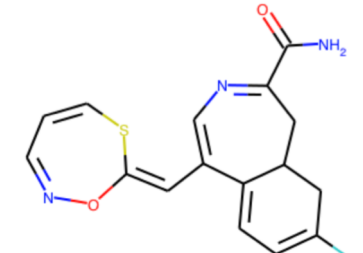
4 datasets are artificially generated, 2 datasets are semi-artificial, and 2 are real-world.

Sizes in terms of total number of nodes vary between 0.27M to 7M.

4 most fundamental graph tasks : graph regression, graph classification, node classification and link prediction.

Proposed Datasets

- **ZINC^[1]** : A popular **real-world molecular dataset** of 250K graphs, out of which we randomly select **12K for efficiency**. We consider the task of **graph property regression for constrained solubility**, an important chemical property for designing generative GNNs for molecules^[2].
 - Statistics : 10,000 train/1,000 validation/1,000 test graphs of sizes 9-37 nodes/heavy atoms.
- **MNIST^[3]/CIFAR10^[4]** : Classical **image classification** datasets converted into graphs using **super-pixels^[5]** and assigning node features as the superpixel coordinates and mean intensity. These datasets are **sanity-checks**, as we expect most GNNs to perform close to 100% for MNIST and well enough for CIFAR10.
 - Statistics : MNIST has 55,000 train/5,000 validation/10,000 test graphs of sizes 40-75 nodes and CIFAR10 has 45,000 train/5,000 validation/10,000 test graphs of sizes 85-150 nodes.



[1] Irwin, Sterling, Mysinger, Bolstad, Coleman, Zinc: a free tool to discover chemistry for biology, 2012

[2] Jia, Lin, Ying, You, Leskovec, Aiken, Redundancy-free computation graphs for graph neural networks, 2019

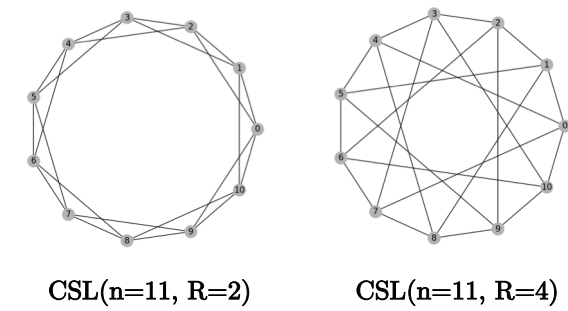
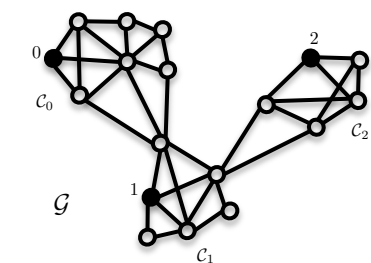
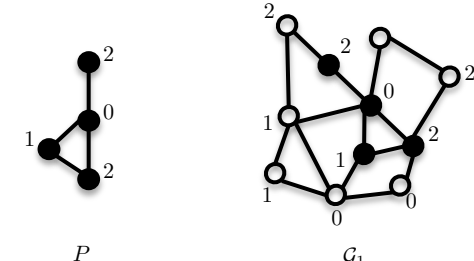
[3] LeCun, Bottou, Bengio, Haffner, Gradient-based learning applied to document recognition, 1998

[4] Alex Krizhevsky et al, Learning multiple layers of features from tiny images, 2009

[5] Achanta, Shaji, Smith, Lucchi, Fua, Süsstrunk, Slic superpixels compared to state-of-the-art superpixel methods, 2012

Proposed Datasets

- **PATTERN/CLUSTER** : Node classification tasks generated with **Stochastic Block Models**^[1], which are widely used to model communities in social networks by modulating the intra- and extra-communities connections. PATTERN tests the fundamental task of recognizing specific predetermined subgraphs^[2].
 - Statistics : PATTERN has 10,000 train/2,000 validation/2,000 test graphs of sizes 50-180 nodes. CLUSTER has 10,000 train/1,000 validation/1,000 test graphs of sizes 40-190 nodes.
- **CSL**^[3] : Synthetic to test the expressivity of GNNs. Graphs are isomorphic if they have the same degree and the task is to **classify non-isomorphic graphs**.
 - Statistics : 5-fold cross-validation split of 150 graphs of sizes 41 nodes with train-val-test ratio 3:1:1.



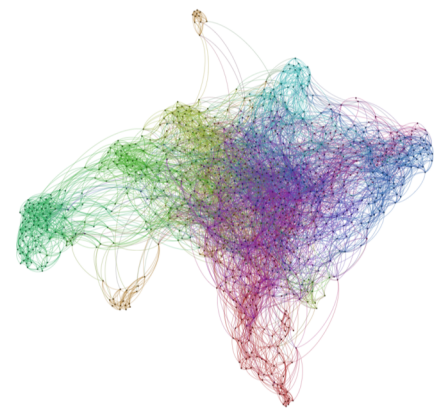
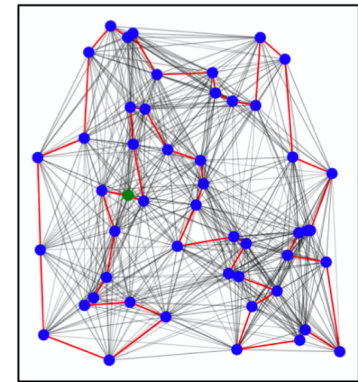
[1] Abbe, Community detection and stochastic block models: recent developments, 2017

[2] Scarselli, Gori, Tsoi, Hagenbuchner, Monfardini, The Graph Neural Network Model, 2009

[3] Murphy, Srinivasan, Rao, Ribeiro, Relational pooling for graph representations, 2019

Proposed Datasets

- **Travelling Salesman Problem (TSP)** : Link prediction on 2D Euclidean graphs to identify edges belonging to the optimal TSP solution given by Concorde^[1]. **TSP is the most studied NP-hard combinatorial problem** with a growing body of literature on leveraging GNNs to learn better solvers^[2,3].
 - Statistics : 10,000 train TSPs / 1,000 validation TSPs / 1,000 test TSPs, where the number of nodes is randomly selected in [50, 500].
- **OGB-COLLAB^[4]** : **Link prediction** dataset proposed by OGB corresponding to a collaboration network between scientists. The task is to predict future author collaboration relationships given past collaboration links.
 - Statistics : A single large temporal graph of size 235K nodes with given train/validation/test edge splits.



[1] Applegate, Bixby, Chvatal, Cook, Concorde tsp solver, 2006

[2] Vinyals, Fortunato, Jaitly, Pointer, 2015

[3] Bello, Pham, Le, Norouzi, Bengio, Neural combinatorial optimization with reinforcement learning, 2016

[4] Hu, Fey, Zitnik, Dong, Ren, Liu, Catasta, Leskovec, Open graph benchmark: Datasets for machine learning on graphs, 2020

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Benchmark Infrastructure



- GitHub Repo :
<https://github.com/graphdeeplearning/benchmarking-gnns>
- Objectives :
 - Ease-of-use and modular, enabling new users to experiment and study the building blocks of GNNs.
 - Experimentally rigorous and fair for all models being benchmarked.
 - Being comprehensive for tracking progress of new GNNs and novel dataset/task.
- Components :
 - Data pipelines
 - GNN layers and models
 - Training and evaluation functions
 - Network and hyperparameter configurations
 - Scripts for reproducibility

The screenshot shows the GitHub repository page for `graphdeeplearning/benchmarking-gnns`. The repository has 21 commits, 3 branches, and 0 tags. It includes sections for **Code**, **Issues**, **Pull requests**, **Actions**, **Projects**, **Wiki**, **Security**, **Insights**, and **Settings**. The **About** section mentions it's a repository for benchmarking graph neural networks, with links to arXiv.org and various benchmarks. The **Releases** section indicates no releases have been published. Below the repository details, there is a step-by-step guide for benchmark installation:

- 1. Benchmark Installation**
Follow these instructions to install the benchmark and setup the environment.
- 2. Download datasets**
Proceed as follows to download the benchmark datasets.
- 3. Reproducibility**
Use this page to run the codes and reproduce the published results.
- 4. Adding a new dataset**
Instructions to add a dataset to the benchmark.
- 5. Adding a Message-passing GCN**
Step-by-step directions to add a MP-GCN to the benchmark.
- 6. Adding a Weisfeiler-Lehman GNN**
Step-by-step directions to add a WL-GNN to the benchmark.

A diagram illustrating a message-passing GNN layer transition from layer ℓ to $\ell+1$. It shows nodes i, j at layer ℓ with hidden states h_i^ℓ, h_j^ℓ . They interact via a function f_{agg} to produce a new state $h_{ij}^{\ell+1}$. This state is then passed to the next layer along with node i , which has state $h_i^{\ell+1}$.

Experimental Setting

- **Data splits** : Given for ZINC, MNIST, CIFAR10, CSL, OGB-COLLAB, random for PATTERN, CLUSTER, TSP.
- **Training** :
 - **Adam** optimizer w/ **learning rate decay strategy**.
Initial learning rate is $1e-3/1e-4$, reduced by half if validation loss does not improve after 5/10 epochs.
Training is stopped if learning rate $\leq 1e-6$ or computational time ≥ 12 hours.
 - **Statistics** with 4 results using 4 different seeds are reported.
- **Parameter budgets** :
 - Our goal is not to find the optimal hyperparameters (computationally expensive) but to compare models within the same parameter budget and a maximal computational time.
 - **Two parameter budgets** :
 - 100k (w/ $L=4$) and 500k (w/ $L=16$) parameters for each GNNs for all tasks.

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Benchmarking Results

- Result #1 : MP-GCNs outperformed WL-GNNs on all datasets.**
 - Potential reasons :
 - $O(n^2)/O(n^3)$ memory/speed complexities of WL-GNNs do not scale to medium-scale datasets.
 - Best result for ZINC, which is the dataset with the smallest sizes, $n \in [9,37]$.
 - Early stage of developments of WL-GNNs.
- Result #2 : MP-GCNs benefit from batch normalization and residual connection :**
 - Boost performance
 - Accelerate training
 - WL-GNNs do not benefit from RC, BN or LN (performances degrade).
 - Batch implementation is carried out by gradient accumulation.

NODE CLASSIFICATION													
Model	L	#Param	Test Acc.±s.d.	Train Acc.±s.d.	PATTERN	#Epoch	Epoch/Total	#Param	Test Acc.±s.d.	Train Acc.±s.d.	CLUSTER	#Epoch	Epoch/Total
MLP	4	105263	50.519±0.000	50.487±0.014	42.25	8.95±0.11hr	106015	20.973±0.004	20.938±0.002	42.25	5.83±0.07hr		
GCN	4	100923	63.880±0.074	65.126±0.135	105.00	118.85±3.51hr	101655	53.445±2.029	54.041±2.197	70.00	65.72±1.30hr		
GraphSage	4	101739	50.516±0.001	50.473±0.014	41.75	93.41±0.17hr	102187	50.454±0.145	54.374±0.203	64.00	53.56±0.97hr		
	16	503842	50.492±0.001	50.487±0.014	46.50	39.11±0.09hr	502235	63.045±1.10	63.74±0.160	57.75	225.61±0.60hr		
MoNet	4	103778	85.582±0.037	85.550±0.044	89.75	87.1±0.00hr	104227	84.06±0.11	84.58±0.15	76.75	24.30±0.30hr		
	16	511487	85.582±0.038	85.720±0.068	81.75	86.40±0.58hr	511999	66.40±0.540	67.72±0.649	77.75	74.42±1.05hr		
GAT	4	109936	71.892±0.334	78.409±1.592	81.50	492.19±11.13hr	501687	68.498±0.976	71.729±2.212	79.75	270.28±6.08hr		
	16	526990	78.271±0.183	77.883±1.632	96.00	20.92±0.57hr	110700	57.732±0.323	58.331±0.342	67.25	14.17±0.27hr		
GatedGCN	4	104003	90.212±0.476	53.50	50.33±0.77hr	527874	70.587±0.447	76.074±1.362	73.50	35.94±0.75hr			
	16	502233	84.480±0.122	84.474±0.155	78.75	139.01±0.99hr	104355	60.404±0.419	61.618±0.536	94.50	79.97±2.13hr		
GatedGCN-PE	16	502457	85.568±0.085	86.007±0.123	65.25	644.71±19.11hr	502015	73.840±0.326	87.880±0.908	60.00	400.07±6.81hr		
GIN	4	100884	85.590±0.037	85.857±0.044	93.00	15.24±0.00hr	104230	58.46±0.11	59.00±0.337	74.75	10.71±0.23hr		
	16	508734	85.590±0.136	85.644±0.116	80.75	85.00±0.62hr	512779	64.71±0.153	65.97±0.116	80.75	20.71±0.77hr		
RingGNN	2	105206	86.245±0.013	86.118±0.034	75.00	573.37±12.11hr	104746	42.41±20.063	42.520±20.212	74.50	428.24±8.79hr		
	2	504766	86.244±0.025	86.105±0.021	72.00	595.97±12.15hr	524022	22.340±0.000	22.304±0.000	43.25	501.84±6.22hr		
	8	505749	Diverged	Diverged	Diverged	Diverged	514380	Diverged	Diverged	Diverged	Diverged		
3WLGNN	3	103572	85.661±0.353	85.608±0.337	95.00	304.79±7.88hr	105552	57.130±6.539	57.404±6.597	116.00	219.51±6.52hr		
	3	502872	85.341±0.207	85.270±0.198	98.15	424.23±9.56hr	105725	55.489±7.863	55.736±8.204	66.00	319.98±5.79hr		
	8	581716	Diverged	Diverged	Diverged	Diverged	586788	Diverged	Diverged	Diverged	Diverged		
GRAPH CLASSIFICATION												CIFAR10	
Model	L	#Param	Test Acc.±s.d.	Train Acc.±s.d.	#Epoch	Epoch/Total	#Param	Test Acc.±s.d.	Train Acc.±s.d.	#Epoch	Epoch/Total	CIFAR10	
MLP	4	104044	95.340±0.138	97.432±0.470	232.25	22.74±0.48hr	104380	56.340±0.181	65.113±1.685	185.25	29.48±1.53hr		
GCN	4	104337	97.312±0.097	100.000±0.000	98.25	113.12±0.13hr	101657	55.710±0.381	69.523±1.948	142.50	109.70±4.39hr		
GraphSage	4	104337	97.312±0.097	100.000±0.000	98.25	113.12±0.13hr	104517	65.767±0.308	99.719±0.062	93.50	124.61±3.39hr		
MoNet	4	104049	90.803±0.032	96.609±0.440	146.25	93.19±3.82hr	104229	54.655±0.518	65.911±2.151	141.50	97.13±3.85hr		
GAT	4	110400	95.535±0.205	99.994±0.008	104.75	42.26±0.12hr	110704	64.223±0.455	89.114±0.499	103.75	55.27±1.62hr		
GatedGCN	4	104217	97.340±0.143	100.000±0.000	96.25	128.79±0.50hr	104357	67.312±0.311	95.553±1.018	97.00	154.15±4.22hr		
GIN	4	105434	96.485±0.252	100.000±0.000	128.00	39.22±0.41hr	105654	55.255±1.527	79.412±9.700	141.50	52.12±2.07hr		
RingGNN	2	505996	11.350±0.000	11.235±0.100	14.00	11.350±0.000	105196	19.556±0.108	19.556±0.107	13.75	311.29±13.00hr		
	2	501823	91.000±0.049	92.000±0.050	14.75	257.50±9.63hr	500849	39.165±17.114	39.165±17.114	13.75	299.24±6.00hr		
	8	506357	Diverged	Diverged	Diverged	Diverged	510439	Diverged	Diverged	Diverged	Diverged		
3WLGNN	3	108092	95.075±0.061	95.830±1.338	27.75	150.92±12.40hr	108516	59.175±1.593	63.751±2.697	28.50	1506.29±12.60hr		
	3	501690	95.002±0.419	95.692±0.677	26.25	160.63±7.34±12.42hr	105270	58.043±2.512	61.574±3.575	20.00	2091.22±12.55hr		
	8	508016	Diverged	Diverged	Diverged	Diverged	501584	Diverged	Diverged	Diverged	Diverged		
LINK PREDICTION												COLLAB	
Model	L	#Param	Test F1 ±s.d.	Train F1 ±s.d.	#Epoch	Epoch/Total	#Param	Test Hits ±s.d.	Train Hits ±s.d.	#Epoch	Epoch/Total	COLLAB	
MLP	4	96956	0.544±0.001	0.544±0.000	164.25	50.15±2.31hr	39441	20.350±2.168	29.807±3.366	147.50	2.09±0.09hr		
GCN	4	95702	0.630±0.001	0.631±0.001	261.00	152.89±11.11hr	40749	50.422±1.131	92.112±0.991	122.50	351.05±12.04hr		
GraphSage	4	92963	0.665±0.003	0.669±0.003	266.00	157.26±11.68hr	39856	51.618±0.690	99.949±0.052	152.75	277.93±11.87hr		
MoNet	4	99007	0.641±0.002	0.643±0.002	282.00	84.466±6.65hr	39751	36.144±2.191	61.156±3.157	167.50	26.69±1.26hr		
GAT	4	96182	0.671±0.002	0.673±0.002	328.25	104.26±1.25hr	42637	51.501±0.962	97.851±1.114	157.00	18.12±0.80hr		
GatedGCN	4	978478	0.791±0.003	0.793±0.003	159.00	104.35±1.25hr	40966	52.635±1.168	96.103±1.874	95.00	453.47±12.09hr		
GatedGCN-E	16	500770	0.808±0.003	0.811±0.003	197.00	218.51±2.04hr	41898	52.849±1.145	96.165±0.453	94.75	45.72±12.08hr		
GatedGCN-E	16	507077	0.856±0.002	0.850±0.001	53.00	104.35±1.25hr	40965	49.212±1.560	88.747±1.058	95.00	45.12±12.03hr		
k-NN Heuristic	0	k = 2	Test F1 ±0.693	-	-	-	605461	44.206±0.452	100.000±0.000	254.33	2.66±0.21hr		
GRAPH REGRESSION - ZINC													
Model	L	#Param	Test MAE ±s.d.	Train MAE ±s.d.	#Epoch	Epoch/Total	#Param	Test MAE ±s.d.	Train MAE ±s.d.	#Epoch	Epoch/Total	ZINC	
MLP	4	108975	0.706±0.000	0.64±0.000	116.75	1.01±0.03hr	-	-	-	-	-		
GCN	4	103077	0.459±0.003	0.343±0.011	196.25	2.89±0.16hr	-	-	-	-	-		
GraphSage	4	105079	0.367±0.001	0.128±0.019	197.00	12.78±0.71hr	-	-	-	-	-		
MoNet	4	106902	0.397±0.010	0.318±0.006	183.75	1.57±0.08hr	-	-	-	-	-		
GAT	4	104013	0.292±0.004	0.093±0.014	171.75	10.82±0.52hr	-	-	-	-	-		
GatedGCN	4	102385	0.475±0.007	0.317±0.006	137.50	2.93±0.11hr	-	-	-	-	-		
GatedGCN-E	4	105135	0.384±0.007	0.067±0.006	144.00	12.98±0.53hr	-	-	-	-	-		
GatedGCN-E	16	104372	0.435±0.003	0.049±0.014	173.50	5.76±0.28hr	-	-	-	-	-		
GatedGCN-E	16	504309	0.282±0.015	0.074±0.016	166.75	20.56±0.96hr	-	-	-	-	-		
GatedGCN-E	16	505011	0.214±0.013	0.067±0.019	185.00	10.70±0.56hr	-	-	-	-	-		
GIN	4	103079	0.387±0.015	0.319±0.015	135.25	2.29±0.10hr	-	-	-	-	-		
RingGNN	2	507978	0.526±0.051	0.444±0.024	147.00	10.22±0.42hr	-	-	-	-	-		
RingGNN	2	101403	0.363±0.026	0.249±0.025	95.00	2.37±0.09hr	-	-	-	-	-		
RingGNN	2	527283	0.353±0.019	0.236±0.019	79.75	29.34±0.63hr	-	-	-	-	-		
RingGNN	8	510305	Diverged	Diverged	Diverged	Diverged	-	-	-	-	-		
3WLGNN	3	102150	0.407±0.024	0.272±0.037	111.25	28.23±0.88hr	-	-	-	-	-		
3WLGNN	3	103096	0.250±0.054	0.174±0.044	117.25	28.6±0.90hr	-	-	-	-	-		
3WLGNN	3	506003	0.249±0.008	0.173±0.008	123.25	22.98±0.80hr	-	-	-	-	-		
3WLGNN	8	582824	0.303±0.057	0.240±0.043	52.50	81.12±12.15hr	-	-	-	-	-		

Evaluation Metrics: (higher is better, except for ZINC)

- CLUSTER, PATTERN use weighted accuracy w.r.t. the class sizes.
- MNIST, CIFAR10 are multi-label classification accuracy.
- TSP uses binary F1 score for the positive edges.
- COLLAB uses Hits@50 via the evaluator provided by OGB [?].
- ZINC uses mean absolute regression error.

Notation:

- Models with the suffix -E use input edge features to initialize edge representations (ZINC: bond type, TSP: Euclidean distance, COLLAB: collaboration frequency and year).
- Models with the suffix -PE use positional and Laplacian Eigenvectors as node positional and time dimensions for ZINC, 2 for TSP, and 20 for others.
- Results denoted by Diverged indicate unstable and divergent runs across all 4 seeds and initial learning rates (10^{-3} – 10^{-1}).
- Results denoted by OOM indicate runs that throw out of memory errors on our hardware configuration.

Extended training:

- For TSP, RingGNN/3WLGNN with 100K parameters achieved 0.733±0.020 and 0.649±0.051 respectively after 48hr of training.

Benchmarking Results

- Result #3 : Anisotropic mechanism improve (isotropic) GCNs.
 - Sparse attention^[1,2] and dense attention^[3] are not injective (unlike f_{GIN}) but experiments showed that they are good at generalization. See also^[4].
- Intuitively, softmax attention is flexible to represent max/mean/weighted mean w.r.t. contextual information.
- Recent work^[5] studied effectiveness of global attention for generalization.

[1] Bahdanau, Cho, Bengio, Neural machine translation by jointly learning to align and translate, 2015

[2] Velickovic, Cucurull, Casanova, Romero, Lio, Bengio, Graph attention networks, 2017

[3] Bresson, Laurent, Residual gated graph convnets, 2017

[4] Abu-El-Haija, Perozzi, Al-Rfou, Alemi, Watch your step: Learning node embeddings via graph attention, 2018

[5] Puny, Ben-Hamu, Lipman, From Graph Low-Rank Global Attention to 2-FWL Approximation, 2020

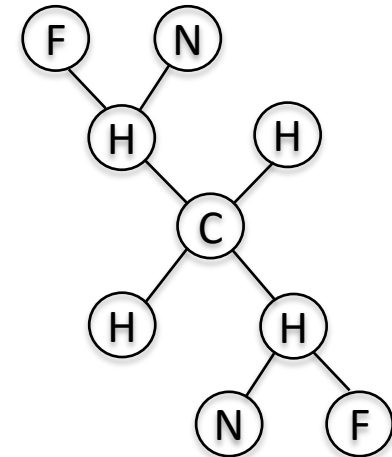
NODE CLASSIFICATION														
Model	L	#Param	Test Acc.±s.d.	Train Acc.±s.d.	PATTERN	#Epoch	Epoch/Total	#Param	Test Acc.±s.d.	Train Acc.±s.d.	CLUSTER	#Epoch	Epoch/Total	
MLP	4	105263	50.519±0.000	50.487±0.014	42.25	8.95±0.11hr	106015	20.973±0.004	20.938±0.002	42.25	5.83±0.07hr			
GCN	4	100923	63.880±0.074	65.126±0.135	105.00	11.88±3.51hr	101655	53.445±2.029	54.041±2.197	70.00	65.72±1.30hr			
GraphSage	4	101739	50.516±0.001	50.473±0.014	41.75	9.34±1.17hr	102187	50.454±0.145	54.374±0.203	64.00	53.56±0.97hr			
	16	504942	50.492±0.001	50.487±0.010	46.50	39.1±1.90hr	504253	63.664±0.110	64.710±0.167	57.75	225.61±0.50hr			
MoNet	4	103778	85.582±0.037	85.582±0.044	89.75	8.71±0.90hr	104227	84.06±0.111	84.06±0.113	76.75	24.30±0.30hr			
	16	511487	85.582±0.038	85.720±0.068	81.75	68.40±0.58hr	511999	66.40±0.540	67.72±0.649	77.75	74.42±1.05hr			
GAT	4	109936	71.892±0.334	78.409±1.592	81.50	492.19±1.13hr	501687	68.498±0.976	71.729±2.212	79.75	270.28±6.08hr			
	16	526990	78.271±0.183	77.883±1.632	96.00	20.92±0.57hr	110700	57.732±0.323	58.331±0.342	67.25	14.17±0.27hr			
GatedGCN	4	104003	84.480±0.122	84.474±0.155	78.75	84.480±0.155	104335	70.587±0.447	76.074±1.362	73.50	35.94±0.57hr			
	16	502233	85.568±0.088	86.007±0.123	65.25	64.61±1.19hr	502615	73.840±0.326	87.880±0.908	60.00	400.07±6.81hr			
GatedGCN-PE	16	502457	86.508±0.085	86.801±0.133	86.75	64.71±0.94hr	504253	76.082±0.196	88.919±0.720	57.75	399.66±6.58hr			
GIN	4	100884	85.590±0.000	85.857±0.000	93.00	15.24±0.11hr	104279	58.45±0.111	58.45±0.137	74.75	10.71±0.23hr			
	16	508734	85.590±0.136	85.844±0.116	85.75	85.0±0.62hr	512779	64.71±0.153	65.972±0.116	80.00	20.70±0.70hr			
RingGNN	2	105206	86.245±0.013	86.118±0.034	75.00	57.37±0.12hr	104746	42.41±0.206	42.520±0.212	74.50	428.24±8.79hr			
	2	504766	86.244±0.025	86.105±0.021	72.00	86.10±0.21hr	524022	22.340±0.000	22.304±0.000	43.25	501.84±6.22hr			
	8	505749	Diverged	Diverged	Diverged	Diverged	514380	Diverged	Diverged	Diverged	Diverged			
3WLNNG	3	103572	85.661±0.353	85.608±0.337	95.00	30.97±0.88hr	105552	57.130±0.539	57.404±0.597	116.00	219.51±6.52hr			
	3	502872	85.341±0.207	85.270±0.198	81.75	424.23±0.56hr	507252	55.489±0.7863	55.736±0.824	66.00	319.98±5.79hr			
	8	581716	Diverged	Diverged	Diverged	Diverged	586788	Diverged	Diverged	Diverged	Diverged			
GRAPH CLASSIFICATION														
Model	L	#Param	Test Acc.±s.d.	Train Acc.±s.d.	MNIST	#Epoch	Epoch/Total	#Param	Test Acc.±s.d.	Train Acc.±s.d.	CIFAR10	#Epoch	Epoch/Total	
MLP	4	104044	95.340±0.138	97.432±0.470	232.25	22.74±0.48hr	104380	56.340±0.181	65.113±1.685	185.25	29.48±1.53hr			
GCN	4	101365	90.705±0.218	97.196±0.223	127.50	83.41±2.99hr	101657	55.710±0.381	69.523±1.948	142.50	109.70±4.39hr			
GraphSage	4	104337	97.312±0.097	100.000±0.000	98.25	113.2±3.13hr	104517	65.767±0.308	99.719±0.062	93.50	124.61±3.29hr			
MoNet	4	104049	90.803±0.032	96.609±0.440	146.25	93.19±3.82hr	104229	54.655±0.518	65.911±2.512	141.50	97.13±3.85hr			
GAT	4	110400	95.535±0.205	99.994±0.008	104.75	42.26±1.25hr	110704	64.223±0.455	89.114±0.499	103.75	55.27±1.62hr			
GatedGCN	4	104217	97.340±0.143	100.000±0.000	96.25	128.79±0.50hr	104337	67.312±0.311	94.553±0.118	97.00	154.15±4.22hr			
GIN	4	105434	96.485±0.252	100.000±0.000	128.00	39.22±0.41hr	105654	55.255±1.527	79.412±0.700	141.50	52.12±0.20hr			
RingGNN	2	501828	91.350±0.049	92.300±0.050	14.25	257.59±0.43hr	500849	39.16±0.17114	39.16±0.17114	13.75	311.96±13.00hr			
	2	505637	Diverged	Diverged	Diverged	Diverged	510439	Diverged	Diverged	Diverged	Diverged			
3WLNNG	3	108024	95.075±0.061	95.830±1.338	27.75	152.32±0.24hr	108516	58.175±1.593	63.751±2.697	28.50	1506.29±12.60hr			
	3	501690	95.002±0.419	95.692±0.677	26.25	160.67±3.12hr	502770	58.043±2.512	61.574±3.575	20.00	2091.22±12.55hr			
	8	500816	Diverged	Diverged	Diverged	Diverged	501584	Diverged	Diverged	Diverged	Diverged			
LINK PREDICTION														
Model	L	#Param	Test F1 ± s.d.	Train F1 ± s.d.	TSP	#Epoch	Epoch/Total	#Param	Test Hits ± s.d.	Train Hits ± s.d.	COLLAB	#Epoch	Epoch/Total	
MLP	4	96956	0.544±0.001	0.544±0.000	164.25	50.15±2.31hr	39441	20.350±2.168	29.807±3.366	147.50	2.09±0.09hr			
GIN	4	95702	0.630±0.001	0.631±0.001	261.00	152.89±11.13hr	40479	50.422±1.131	92.112±0.991	122.50	351.05±12.04hr			
GraphSage	4	99263	0.665±0.003	0.669±0.003	266.00	157.26±11.68hr	39856	51.618±0.690	99.949±0.052	152.75	277.93±11.87hr			
MoNet	4	99007	0.641±0.002	0.643±0.002	282.00	84.46±6.65hr	39751	36.144±2.191	61.156±3.973	167.50	26.69±1.26hr			
GAT	4	96182	0.671±0.002	0.673±0.002	328.25	68.23±6.25hr	42637	51.501±0.962	97.851±1.114	157.00	18.12±0.80hr			
GatedGCN	4	97858	0.791±0.003	0.793±0.003	159.00	108.52±0.72hr	40965	52.635±1.168	96.103±1.181	95.00	453.74±12.09hr			
GatedGCN-E	4	97858	0.808±0.003	0.811±0.003	197.00	218.51±2.04hr	41898	52.849±1.145	96.165±0.453	94.75	45.72±12.08hr			
GatedGCN-E-PE	16	500770	0.838±0.002	0.850±0.001	53.00	160.21±1.27hr	40695	49.212±1.560	88.747±1.058	95.00	45.12±12.03hr			
GIN	4	99002	0.656±0.003	0.660±0.003	273.50	72.73±5.56hr	39544	41.730±2.284	70.555±4.444	140.25	8.66±0.34hr			
RingGNN	2	106862	0.643±0.024	0.644±0.024	2.00	1785.0±17.19hr	-	OOM						
	2	507938	0.704±0.003	0.705±0.003	3.00	1283.53±16.08hr	-	OOM						
3WLNNG	3	106366	0.694±0.073	0.699±0.073	2.00	1746.81±16.59hr	-	OOM						
	3	504681	0.288±0.311	0.290±0.312	2.00	1719.0±7.16hr	-	OOM						
8	508832	OOM	OOM	OOM	OOM	OOM	-	OOM						
k-NN Heuristic	0	k = 2	Test F1: 0.693											
Matrix Fact.	0													
								60546561	44.206±0.452	100.000±0.000	254.33	2.66±0.21hr		
GRAPH REGRESSION - ZINC														
Model	L	#Param	Test MAE ± s.d.	Train MAE ± s.d.	ZINC	#Epoch	Epoch/Total							
MLP	4	108975	0.706±0.000	0.644±0.000	116.75	1.01±0.03hr								
GCN	4	103077	0.459±0.003	0.343±0.011	196.25	2.89±0.16hr								
GraphSage	4	105079	0.367±0.011	0.128±0.019	197.00	12.78±0.71hr								
MoNet	4	106902	0.397±0.010	0.318±0.006	147.25	1.93±0.08hr								
GAT	4	104013	0.292±0.008	0.093±0.014	171.75	10.82±0.52hr								
	4	102385	0.475±0.007	0.317±0.006	137.50	2.93±0.11hr								
GatedGCN	4	103145	0.384±0.007	0.067±0.014	144.00	12.98±0.53hr								
GatedGCN-E	4	104735	0.343±0.011	0.074±0.014	173.50	5.760±0.28hr								
GatedGCN-E-PE	16	504309	0.249±0.013	0.249±0.013	102.75	10.26±0.56hr								
GatedGCN-E-PE	16	505011	0.214±0.013	0.067±0.019	185.00	10.70±0.56hr								
GIN	4	103079	0.387±0.015	0.319±0.015	135.25	2.90±0.10hr								
RingGNN	2	507938	0.526±0.051	0.444±0.039	147.00	10.22±0.42hr								
	2	106403	0.363±0.026	0.245±0.025	95.00	2.77±0.76hr								
2	527283	0.353±0.019	0.236±0.019	293.94±0.63hr										
8	510305	Diverged	Diverged	Diverged	Diverged	Diverged								
3WLNNG	3	102150	0.407±0.024	0.272±0.037	111.25	28.23±0.88hr								
3WLNNG-E	3	103096	0.250±0.054	0.147±0.044	117.25	16.69±0.90hr								
	3	504603	0.288±0.015	0.074±0.016	166.75	20.56±0.96hr								
8	508224</													

Outline

- Motivation
- Message-Passing GCNs
- Weisfeiler-Lehman GNNs
- Graph-Agnostic GNNs
- Datasets
- Infrastructure and Experimental Setting
- Benchmarking Results
- **Laplacian Positional Encodings**
- Link Prediction with Edge Representation
- Conclusion

Structural (GCNs) vs Positional Encodings

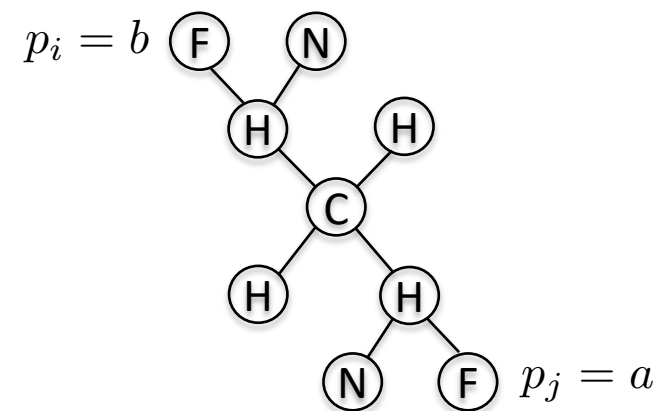
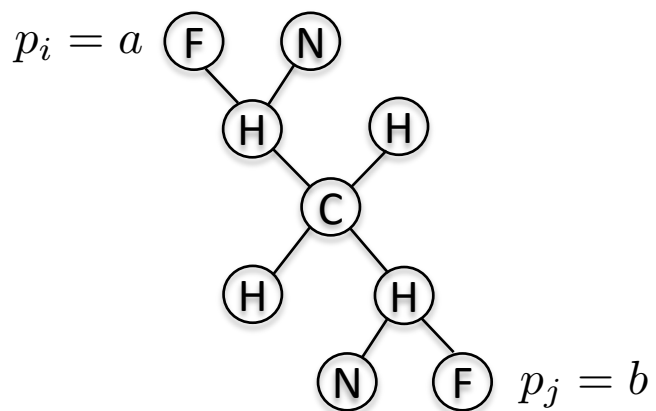
- GCNs like GINs^[1] are not able to differentiate isomorphic nodes, i.e. nodes with the same neighborhood structure :
 - All F atoms have the same representation.
 - All N atoms have the same representation.
 - Two H atoms have the same representation.
 - Two H atoms have the same representation.
- This is a limitation of the expressivity of GCNs.
 - Can we break this structural symmetry ?
 - This can be done either by
 - Higher-order WL-GNNs, but require $O(n^2)/O(n^3)$.
 - Positional encodings of nodes with $O(n)$.



[1] Xu, Hu, Leskovec, Jegelka, How powerful are graph neural networks?, 2019

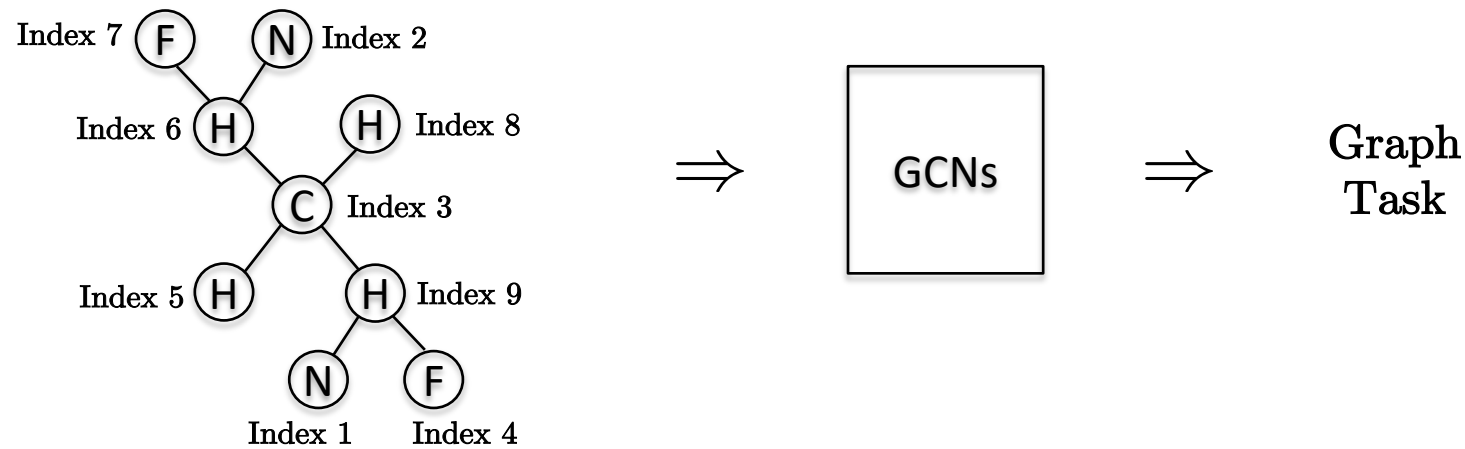
Graph Positional Encodings

- Properties of PEs :
 - Unique representation for each node.
 - Distance-sensitive : Nodes far apart on the graph should have different positional features whereas nodes nearby have similar positional features.
- Graph symmetries prevent assigning canonical representation of PEs.
 - if nodes i and j are structurally symmetric and we have PEs $p_i=a$, $p_j=b$, then it is also possible to arbitrary choose $p_i=b$, $p_j=a$.
 - PEs are always arbitrary up to the number of graph symmetries.



Index Positional Encodings

- Simplest PEs are (arbitrary) **ordering to the nodes**, among $n!$ possible orderings.
- Theorem^[1]: GCNs s.a. GINs^[2] are **more expressive** than the 1-WL test when considering (one-hot encoding of) **node indices** as PE features.
- During training, **orderings are uniformly sampled** from the $n!$ possible choices in order for the network to learn to be independent to these arbitrary choices.



[1] Murphy, Srinivasan, Rao, Ribeiro, Relational pooling for graph representations, 2019

[2] Xu, Hu, Leskovec, Jegelka, How powerful are graph neural networks?, 2019

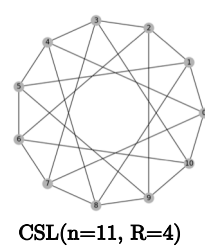
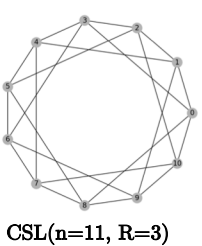
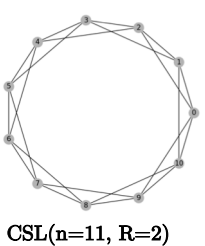
Laplacian Positional Encodings

- Laplacian PEs :
 - Eigen-decomposition : $\Delta = \mathbf{I} - D^{-1/2} A D^{-1/2} = U^T \Lambda U$
 - Hybrid positional and structural encodings, invariant by index permutation.
 - Unique and distance-sensitive : two nodes far away on a graph have large PEs distance, and inversely (two one-encoding vectors of different indices are equally distant).
- LapPEs have also natural symmetries with the arbitrary sign of eigenvectors.
 - The number of possible sign flips is 2^k , k being the number of eigenvectors.
 - In practice, we choose $k \ll n$, and therefore 2^k is much smaller than $n!$ (the number of possible ordering of the nodes).
 - During the training, sign of eigenvectors will be uniformly sampled at random between the 2^k possibilities.
- Lap PEs are graph generalizations of Transformer's PEs^[2].

[1] Vaswani, Shazeer, Parmar, Uszkoreit, Jones, Gomez, Kaiser, Polosukhin, Attention is all you need, 2017

Numerical Results

- **Results :** PEs with GatedGCN^[1].
Performance reported on the test sets of CSL, ZINC, PATTERN, CLUSTER and COLLAB (higher is better, except for ZINC).



	PE type	L	#Param	Test Acc. \pm s.d.	Train Acc. \pm s.d.	#Epochs	Epoch/Total
CSL	No PE	4	104007	10.000 \pm 0.000	10.000 \pm 0.000	54.00	0.58s/0.05hr
	EigVecs-20	4	105407	68.633 \pm 7.143	99.811 \pm 0.232	107.16	0.59s/0.09hr
	Rand sign(EigVecs)	4	105407	99.767\pm0.394	99.689 \pm 0.550	188.76	0.59s/0.16hr
	Abs(EigVecs)	4	105407	99.433 \pm 1.133	100.000 \pm 0.000	143.64	0.60s/0.12hr
	Fixed node ordering	4	106807	10.533 \pm 4.469	76.056 \pm 14.136	60.56	0.59s/0.05hr
	Rand node ordering	4	106807	11.133 \pm 2.571	10.944 \pm 2.106	91.60	0.60s/0.08hr
PATTERN	No PE	16	502223	85.605 \pm 0.105	85.999 \pm 0.145	62.00	646.03s/11.36hr
	EigVecs-2	16	505421	86.029 \pm 0.085	86.955 \pm 0.227	65.00	645.36s/11.94hr
	Rand sign(EigVecs)	16	502457	86.508\pm0.085	86.801 \pm 0.133	65.75	647.94s/12.08hr
	Abs(EigVecs)	16	505421	86.393 \pm 0.037	87.011 \pm 0.172	62.00	645.90s/11.41hr
	Fixed node ordering	16	516887	80.133 \pm 0.202	98.416 \pm 0.141	45.00	643.23s/8.27hr
	Rand node ordering	16	516887	85.767 \pm 0.044	85.998 \pm 0.063	64.50	645.09s/11.79hr
CLUSTER	No PE	16	502615	73.684 \pm 0.348	88.356 \pm 1.577	61.50	399.44s/6.97hr
	EigVecs-20	16	504253	75.520 \pm 0.395	89.332 \pm 1.297	49.75	400.50s/5.70hr
	Rand sign(EigVecs)	16	504253	76.082\pm0.196	88.919 \pm 0.720	57.75	399.66s/6.58hr
	Abs(EigVecs)	16	504253	73.796 \pm 0.234	91.125 \pm 1.248	58.75	398.97s/6.68hr
	Fixed node ordering	16	517435	69.232 \pm 0.265	92.298 \pm 0.712	51.00	400.40s/5.82hr
	Rand node ordering	16	517435	74.656 \pm 0.314	82.940 \pm 1.718	61.00	397.75s/6.88hr
COLLAB	No PE	3	40965	52.635 \pm 1.168	96.103 \pm 1.876	95.00	453.47s/12.09hr
	EigVecs-20	3	41889	52.326 \pm 0.678	96.700 \pm 1.296	95.00	452.40s/12.10hr
	Rand sign(EigVecs)	3	41889	52.849\pm1.345	96.165 \pm 0.453	94.75	452.75s/12.08hr
	Abs(EigVecs)	3	41889	51.419 \pm 1.109	95.984 \pm 1.157	95.00	451.36s/12.07hr
	PE type	L	#Param	Test MAE \pm s.d.	Train MAE \pm s.d.	#Epochs	Epoch/Total
ZINC	No PE	16	504153	0.354 \pm 0.012	0.095 \pm 0.012	165.25	10.52s/0.49hr
	EigVecs-8	16	505011	0.319 \pm 0.010	0.038 \pm 0.007	143.25	10.62s/0.43hr
	Rand sign(EigVecs)	16	505011	0.214\pm0.013	0.067 \pm 0.019	185.00	10.70s/0.56hr
	Abs(EigVecs)	16	505011	0.214\pm0.009	0.035 \pm 0.011	167.50	10.61s/0.50hr
	Fixed node ordering	16	507195	0.431 \pm 0.007	0.044 \pm 0.009	118.25	10.62s/0.35hr
	Rand node ordering	16	507195	0.321 \pm 0.015	0.177 \pm 0.015	184.75	10.55s/0.55hr

[1] Bresson, Laurent, Residual gated graph convnets, 2017

Outline

- Motivation
- Message-Passing GCNs
- Weisfeiler-Lehman GNNs
- Graph-Agnostic GNNs
- Datasets
- Infrastructure and Experimental Setting
- Benchmarking Results
- Laplacian Positional Encodings
- **Link Prediction with Edge Representation**
- Conclusion

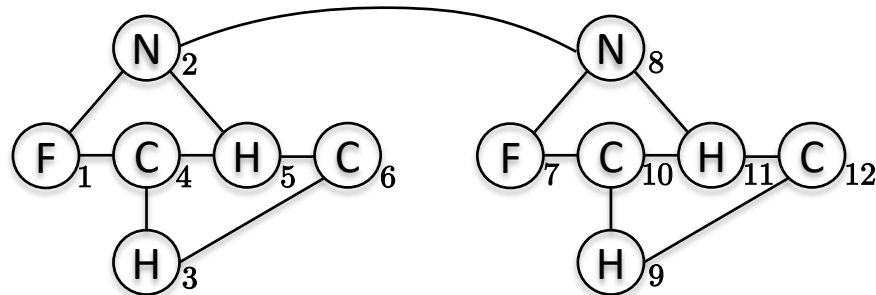
Link Prediction

- GCNs may fail the link prediction task.
 - Apply any GCN to this molecule composed of two identical compounds :

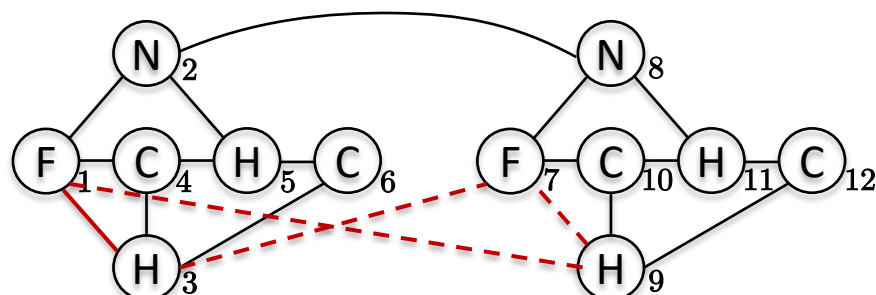
We have

$$h_1 = h_7 \in \mathbb{R}^d$$

$$h_3 = h_9 \in \mathbb{R}^d$$

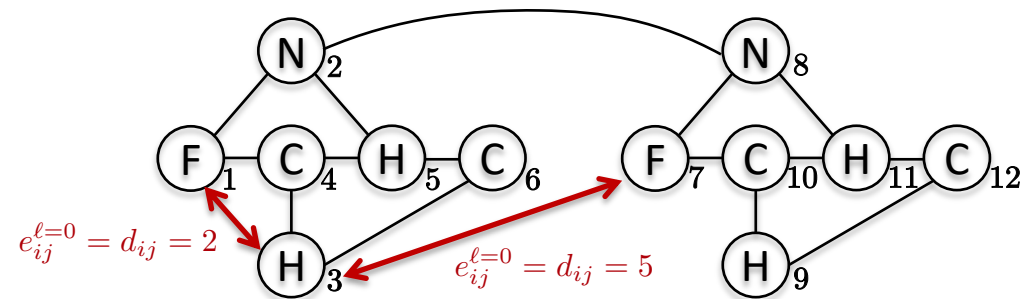


- Perform link prediction (by transfer learning) :
 - Suppose there exist a bond $(F_{(item\ 1)}, H_{(item\ 3)})$.
 - Can we predict the link $(F_{(item\ 7)}, H_{(item\ 9)})$? Yes, but the network will also predict (wrong) links between $(F_{(item\ 7)}, H_{(item\ 3)})$ and between $(F_{(item\ 1)}, H_{(item\ 9)})$.



Link Prediction with Edge Representation

- How to design expressive GCNs for the link prediction task ?
 - Theorem^[1] :
 - Link prediction is maximally expressive with a joint representation of nodes (intuitively this encodes the similarity/distance between pair of nodes).
 - This requires $O(n^2)$ edge representations.

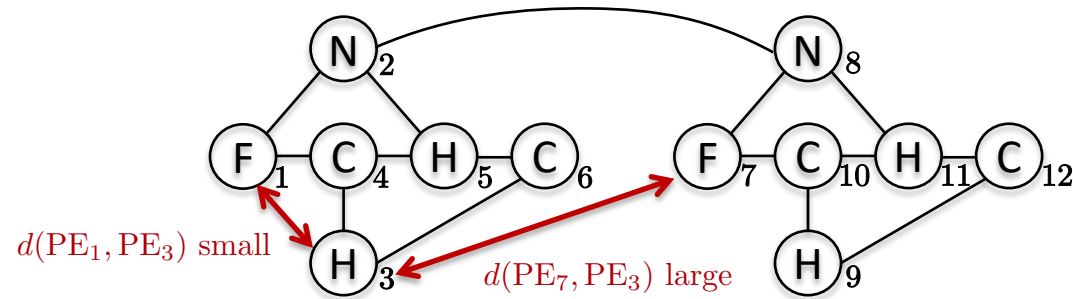


Input edge representation is $e_{ij}^{l=0} = d_{ij}$ (e.g. shortest distances).

[1] Srinivasan, Ribeiro, On the Equivalence between Positional Node Embeddings and Structural Graph Representations, 2019

Link Prediction with Edge Representation

- Alternate solution :
 - Represent links as joint representations of nodes.
 - Add node positional encodings that are unique and distance-sensitive to differentiate links, like Laplacian PEs.
 - Complexity is $O(E)$, E is the number of edges.



Numerical Results

- We study the **impact of edge representation** by instantiating three variants :

- No edge feature/representation** : Isotropic models (like vanilla GCNs^[1])

$$h_i^{\ell+1} = \sigma \left(\sum_{j \in \mathcal{N}_i} W^\ell h_j^\ell \right), \text{ identified by (E.Feat,E.Repr= } x,x)$$

- Edge feature/no edge representation** : Anisotropic models (like GAT^[2])

$$h_i^{\ell+1} = \sigma \left(\sum_{j \in \mathcal{N}_i} f_{V^\ell}(h_i^\ell, h_j^\ell) \cdot W^\ell h_j^\ell \right), \text{ with (E.Feat,E.Repr=} \checkmark, x)$$

- Edge feature and representation** : Anisotropic models (like GatedGCNs^[3])

$$h_i^{\ell+1} = \sigma \left(\sum_{j \in \mathcal{N}_i} e_{ij}^\ell \cdot W^\ell h_j^\ell \right), \quad e_{ij}^{\ell+1} = f_{V^\ell}(h_i^\ell, h_j^\ell, e_{ij}^\ell), \quad \text{with (E.Feat,E.Repr=} \checkmark, \checkmark)$$

	Model	EFeat.	E.Repr.	L	#Param	Test Acc.±s.d.	Train Acc.±s.d.	#Epochs	Epoch/Total
TSP	GatedGCN	x	x	4	99026	0.646±0.002	0.648±0.002	197.50	150.83s/8.34hr
	GatedGCN	✓	x	4	98174	0.757±0.009	0.760±0.009	218.25	197.80s/12.06hr
	GatedGCN	✓	✓	4	97858	0.791±0.003	0.793±0.003	159.00	218.20s/9.72hr
	GatedGCN-E	✓	✓	4	97858	0.808±0.003	0.811±0.003	197.00	218.51s/12.04hr
	GAT	x	x	4	95462	0.643±0.001	0.644±0.001	132.75	325.22s/12.10hr
	GAT	✓	x	4	96182	0.671±0.002	0.673±0.002	328.25	68.23s/6.25hr
	GAT	✓	✓	4	96762	0.748±0.022	0.749±0.022	93.00	462.22s/12.10hr
	GAT-E	✓	✓	4	96762	0.782±0.006	0.783±0.006	98.00	438.37s/12.11hr
	GatedGCN	x	x	3	26593	35.989±1.549	60.586±4.251	148.00	263.62s/10.90h
	GatedGCN	✓	x	3	26715	50.668±0.291	96.128±0.576	172.00	384.39s/18.44hr
COLLAB	GatedGCN	✓	✓	3	27055	51.537±1.038	96.524±1.704	188.67	376.67s/19.85hr
	GatedGCN-E	✓	✓	3	27055	47.212±2.016	85.801±0.984	156.67	377.04s/16.49hr
	GAT	x	x	3	28201	41.141±0.701	70.344±1.837	153.50	371.50s/15.97hr
	GAT	✓	x	3	28561	50.662±0.687	96.085±0.499	174.50	403.52s/19.69hr
	GAT	✓	✓	3	26676	49.674±0.105	92.665±0.719	201.00	349.19s/19.59hr
	GAT-E	✓	✓	3	26676	44.989±1.395	82.230±4.941	120.67	328.29s/11.10hr

[1] Kipf, Welling, Semi-supervised classification with graph convolutional networks, 2017

[2] Velickovic, Cucurull, Casanova, Romero, Lio, Bengio, Graph attention networks, 2017

[3] Bresson, Laurent, Residual gated graph convnets, 2017

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Conclusion

- Take-home messages^[1] :
 - MP-GCNs outperformed WL-GNNs for GINs, 3WL-GNNs, RingGNNs on the 8 datasets used in this benchmark.
 - MP-GCNs benefit from graph sparsity, and universal building blocks w/ batch normalization and residual connection.
 - Anisotropic mechanism improve (isotropic) GCNs.
 - Laplacian eigenvectors improve index positional encodings.
 - Edge representation with positional encodings enhance link prediction.
- Recent works have focused on more efficient WL-inspired techniques^[2,3,4,5,6].
- Benchmarking will bridge the gap between theory and practical performances.

[1] Dwivedi, Joshi, Laurent, Bengio, Bresson, Benchmarking graph neural networks, 2020, <https://arxiv.org/pdf/2003.00982.pdf>

[2] Bouritsas, Frasca, Zafeiriou, Bronstein, Improving Graph Neural Network Expressivity via Subgraph Isomorphism Counting, 2020

[3] Puny, Ben-Hamu, Lipman, From Graph Low-Rank Global Attention to 2-FWL Approximation, 2020

[4] Morris, Rattan, Mutzel, Weisfeiler and Leman go sparse: Towards scalable higher-order graph embeddings, 2020

[5] Corso, Cavalleri, Beaini, Lio, and Velickovic, Principal neighbourhood aggregation for graph nets, 2020

[6] Vignac, Loukas, Frossard, Building powerful and equivariant graph neural networks with message-passing, 2020

Workshop Announcement

The screenshot shows the IPAM website's workshops section. At the top, there's a navigation bar with links for Programs, Videos, News, People, Your Visit, About IPAM, Donate, and Contact Us. A search bar is also present. Below the navigation, a banner for the "Deep Learning and Combinatorial Optimization" workshop is displayed, featuring a blue background with mathematical symbols and a circular pattern of dots.

Workshops

Programs > Workshops > Deep Learning and Combinatorial Optimization

Deep Learning and Combinatorial Optimization

FEBRUARY 22 - 26, 2021

OVERVIEW LODGING APPLICATION & REGISTRATION

Overview

In recent years, deep learning has significantly improved the fields of computer vision, natural language processing and speech recognition. Beyond these traditional fields, deep learning has been extended to quantum chemistry, physics, neuroscience, and more recently to combinatorial optimization (CO). Well-known CO problems are Travelling Salesman Problem, assignment problems, routing, planning, Bayesian search, and scheduling. CO is basically used every day in finance and revenue management, transportation, manufacturing, supply chain, public policy, hardware design, computing and information technology.

Most combinatorial problems are difficult to solve, often leading to heuristic solutions which require years of research work and significant specialized knowledge. For example, the famous TSP problem has been studied for more than 80 years, and the best solver leverages 30 years of theoretical developments, data structures and heuristics from computer science. In the last few years, deep learning has developed some preliminary but promising approaches to deal with classical CO problems such as TSP, MaxCut, Minimum Vertex Cover, Knapsack, Quadratic Assignment Problem and Vehicle Routing Problems. DL is particularly attractive to address CO problems given its high flexibility, approximate

<https://www.ipam.ucla.edu/programs/workshops/deep-learning-and-combinatorial-optimization>

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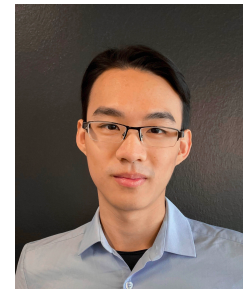
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Thank you

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