

Learning to Approximate Adaptive Kernel Convolution on Graphs

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Abstract

Various Graph Neural Networks (GNNs) have been successful in analyzing data in non-Euclidean spaces, however, they have limitations such as oversmoothing, i.e., information becomes excessively averaged as the number of hidden layers increases. The issue stems from the intrinsic formulation of conventional graph convolution where the nodal features are aggregated from a direct neighborhood per layer across the entire nodes in the graph. As setting different number of hidden layers per node is infeasible, recent works leverage a diffusion kernel to redefine the graph structure and incorporate information from farther nodes. Unfortunately, such approaches suffer from heavy diagonalization of a graph Laplacian or learning a large transform matrix. In this regards, we propose a diffusion learning framework, where the range of feature aggregation is controlled by the scale of a diffusion kernel. For efficient computation, we derive closed-form derivatives of approximations of the graph convolution with respect to the scale, so that node-wise range can be adaptively learned. With a downstream classifier, the entire framework is made trainable in an end-to-end manner. Our model is tested on various standard datasets for node-wise classification for the state-of-the-art performance, and it is also validated on a real-world brain network data for graph classifications to demonstrate its practicality for Alzheimer classification.

Introduction

Graph Neural Network (GNN) has been heavily recognized in machine learning and computer vision with various practical applications such as text classification (Huang et al. 2019), neural machine translation (Bastings et al. 2017), 3D shape analysis (Wei et al. 2020; Verma et al. 2018), semantic segmentation (Qi et al. 2017; Xie et al. 2021), social information systems (Lin et al. 2021) and speech recognition (Liu et al. 2016). At the heart of these GNN models lies the graph convolution, which develops useful representation of each node with a filtering operation on the graph. Given a graph comprised of a set of nodes/edges and signal defined on its nodes, in (Kipf et al. 2017), a convolutional layer was introduced as a linear combination of the signal within a direct neighborhood of each node using the topology of the graph, and its equivalence with spectral filtering (Hammond et al. 2011) has been shown. Stacking these convolutional layers

(together with non-linear activations) constitutes the fundamental Graph Convolutional Network (GCN) (Kipf et al. 2017), and testing a newly developed GCN on classifying node-wise labels has become a standard benchmark to validate the GCN model (Chen et al. 2018; Wu et al. 2019; Xu et al. 2020; Chen et al. 2020; Yang et al. 2021).

Within the architecture of conventional GCN and its variants, there is a fundamental issue: each convolution layer gathers information within a direct neighborhood *uniformly* across all nodes. When information aggregation from direct neighborhood is not sufficient, the convolution layers are stacked to seek for useful information from a larger neighborhood. Such an architecture with several convolution layers broadens the range of neighborhood for information aggregation *uniformly*, again, across the entire nodes in the graph. Eventually, when the same information is shared across all the nodes, it leads to “oversmoothed” representation of each node, not being able to characterize one from another. This behavior can be easily interpreted from the spectral perspective, as a filtering operation in the spectral space will uniformly affect all nodes in the graph space.

Perhaps the most intuitive solution to the oversmoothing is to use different ranges of neighborhood per node. However, it is difficult to design such a model with conventional graph convolution, as it would require different number of convolution layers for each node which is highly impractical. Several recent works tried to overcome this locality issue. Methods in (Veličković et al. 2018; Kim et al. 2022) leverage attention to capture long-range relationships among nodes, authors in (Gao et al. 2019; Wu et al. 2022b) develop pooling scheme to compress graphs, and authors in (Chen et al. 2020) improve upon the vanilla GCN with skip connection of residuals as in ResNet (He et al. 2016).

Notably, GraphHeat (Xu et al. 2020) used a diffusion kernel to redefine distances between nodes as a heat diffusion process among the nodes along the graph structure. It easily connects many local nodes within a range (i.e., scale) even though they are not directly connected. Such an approach is quite effective when the homophily condition is reasonably held even if the given edges in a graph may be noisy. But still, the scale was defined as a hyperparameter and the same range was arranged across the entire graph. Later, a framework that trains on the scale according to a target task was introduced with the gradient of a loss with re-

spect to the scale for each node. While using diffusion kernel have shown to be quite effective, such an approach is computationally inefficient with heavy diagonalization of graph Laplacian, especially when dealing with a large or a population of graphs. Other diffusion-based model such as (Zhao et al. 2021) uses diffusion on layers and channels of features instead of nodes, and Graph Neural Diffusion (GRAND) (Chamberlain et al. 2021) trains on large weight matrices for attention which can be computationally burdening.

In this regime, we propose an efficient framework that learns adaptive scales for each node of a graph with approximations, i.e., Learning Scales via APproximation (LSAP). The key idea is to train on the node-wise *range of neighborhood* instead of excessively stacking convolutional layers. For this, we first show that the formulation in (Xu et al. 2020) can be defined as a heat kernel convolution, which can be approximated with various polynomials (Huang et al. 2020). We then derive the derivatives of the expansion coefficients of these polynomials in the scale space, which can be used to define task-specific gradients to train the optimal scales *within the approximation* instead of learning exhaustive transform matrices. LSAP achieves novel node-wise representation adaptively by leveraging features from other nodes within a trained “range” defined by the diffusion kernel. The ideas above lead to the following contributions:

- We propose a GNN with an adaptive diffusion kernel whose approximations are trainable in an end-to-end manner at each node,
- We derive closed-form derivatives of various polynomial coefficients with respect to the range (i.e., scale) so that graph convolution can be efficiently trained,
- Learning on scales provides interpretable results on the semantics of each node, validated on two independent datasets with different tasks.

LSAP demonstrates superior results on the node classification task in a semi-supervised learning setting (Shchur et al. 2018), as well as on a graph classification task performed on a population of brain networks to predict diagnostic labels for Alzheimer’s Disease (AD). Especially in the AD experiment, the trained scales delineate specific regions highly responsible for the prediction of AD for interpretability.

Related Works

Graph Neural Networks. The vanilla GCN (Kipf et al. 2017) and Variant GNNs utilize graph convolution to perform feature aggregation from neighbors. Simplifying Graph Convolution (SGC) (Wu et al. 2019) captures high-order information in the graph with K-th power of the adjacency matrix, GCNII (Chen et al. 2020) extends a vanilla GCN with residual connection and identity mapping (He et al. 2016), and Graph Attention Network (GAT) (Veličković et al. 2018) introduces attention mechanism on graphs to assign relationships to different nodes. Personalized Propagation of Neural Prediction and its Approximation (APPNP) (Klicpera et al. 2019b) improved message propagation based on personalized PageRank (Page et al. 1999), Graph Random Neural Network (GRAND) (Feng

et al. 2020) developed graph data augmentation, and Deep Adaptive Graph Neural Network (DAGNN) (Liu et al. 2020) disentangled representation transform and message propagation to construct a deep model.

Also, there are recent works that discover useful graph structures from data to adaptively update the structures for message passing. DIAL-GNN (Chen et al. 2021) jointly learned graph structure and embeddings by iteratively searching for hidden graph structures, Bayesian GCNN (Zhang et al. 2019a) incorporated uncertain graph information through a parametric random graph model, and NodeFormer (Wu et al. 2022a) proposed an efficient message passing scheme for propagating layer-wise node signals.

Graph Neural Networks with Diffusion. There are several previous works that adopt diffusion on graphs for GNNs (Lee et al. 2023; Zhang et al. 2023; Huang et al. 2023). Graph Diffusion Convolution (GDC) (Klicpera et al. 2019a) introduced spatially localized graph convolution to aggregate information of indirect nodes, Adaptive Diffusion Convolution (ADC) (Zhao et al. 2021) learned a global radius applied on different layers and channels of features, GRAND (Chamberlain et al. 2021) defined diffusion PDEs on graphs and trained on weight matrices to learn attention for diffusivity, and Fast and Scalable Network Representation Learning (ProNE) (Zhang et al. 2019b) focused on effective network embedding with spectral propagation for enhancement, where they train a single global scale in the spectral propagation. Our methods differ from these methods in that it uses an adaptive parametric kernel at individual nodes, and we propose a new optimization scheme on the scales within its polynomial approximations. Also, many of methods above can be adopted for graph classification as well (Veličković et al. 2018; Klicpera et al. 2019a) with an additional layer transforming node embeddings to a graph embedding. Other literature, although not fully discussed in this section, will be introduced and used as the baselines to compare the performances with our proposed model for both node and graph classification tasks later.

Preliminaries

Graph Convolution with Heat Kernel. An undirected graph $G = \{V, E\}$ comprises a node set V with $|V| = N$ and an edge set E . A graph G is often represented as a symmetric adjacency matrix A of which individual elements a_{pq} encodes connectivity information between node p and q . A graph Laplacian is defined as $L = D - A$ where D is a degree matrix, i.e., a diagonal matrix with $D_{pp} = \sum_q A_{pq}$. Since L is positive semi-definite, it has a complete set of orthonormal basis $U = [u_1 | u_2 | \dots | u_N]$ known as Laplacian eigenvectors and corresponding real and non-negative eigenvalues $0 = \lambda_1 \leq \lambda_2 \leq \dots \leq \lambda_N$. A normalized Laplacian is defined as $\hat{L} = I_N - D^{-1/2} A D^{-1/2}$ where I_N is an identity matrix. Since \hat{L} is real symmetric, it also has a complete set of eigenvectors and eigenvalues.

In (Chung et al. 1997), the heat kernel between nodes p and q is defined in the spectral domain spanned by U as

$$h_s(p, q) = \sum_{i=1}^N e^{-s\lambda_i} u_i(p) u_i(q) \quad (1)$$

where u_i is i -th eigenvector of the graph Laplacian, and the kernel $e^{-s\lambda_i}$ captures smooth transition between the nodes as a diffusion process within the scale s . Using convolutional theorem (Oppenheim et al. 1997), graph Fourier transform, i.e., $\hat{x} = U^T x$, offers a way to define the graph convolution $*$ of a signal $x(p)$ with a filter h_s . Using Eq. (1), heat kernel convolution with h_s as a low-pass filter is defined as

$$h_s * x(p) = \sum_{i=1}^N e^{-s\lambda_i} \hat{x}(i) u_i(p) \quad (2)$$

whose band-width is controlled by the scale s .

Approximation of Convolution with Heat Kernel. The exact computation of Eq. (2) requires diagonalization of a graph Laplacian which can be computationally challenging. Existing literature uses Chebyshev polynomial as a basis to approximate the kernel convolution as a linear transform (He et al. 2022). In (Huang et al. 2020), approximation of heat kernel convolution was introduced using several orthogonal polynomials such as Chebyshev, Hermite and Laguerre. The analytic solutions to the polynomial coefficients $c_{s,n}$ for scale s were derived for Chebyshev polynomial P_n^T , Hermite polynomial P_n^H and Laguerre polynomial P_n^L , where n denotes the degree of each polynomial.

A polynomial $P_n \in \{P_n^T, P_n^H, P_n^L\}$ is often defined by a second order recurrence as

$$P_{n+1}(\lambda) = (\alpha_n \lambda + \beta_n) P_n(\lambda) + \gamma_n P_{n-1}(\lambda) \quad (3)$$

where initial conditions $P_{-1}(\lambda) = 0$ and $P_0(\lambda) = 1$ for $n \geq 0$ and parameters α_n , β_n and γ_n determine the type of polynomial. Then, the heat kernel $e^{-s\lambda}$ can be defined with polynomials P_n and expansion coefficients $c_{s,n}$ as

$$e^{-s\lambda} = \sum_{n=0}^{\infty} c_{s,n} P_n(\lambda), \quad (4)$$

Now, the solution to the heat diffusion in Eq. (2) can be expressed in terms of P_n and $c_{s,n}$ via Eq. (4) as

$$h_s * x(p) = \sum_{n=0}^{\infty} c_{s,n} \sum_{j=1}^N P_n(\lambda_j) \hat{x}(j) u_j(p). \quad (5)$$

Since $\hat{L}u_j = \lambda_j u_j$, the Eq. (5) can be further written as

$$h_s * x(p) = \sum_{n=0}^{\infty} c_{s,n} P_n(\hat{L})x(p) \quad (6)$$

where initial conditions $P_{-1}(\hat{L})x(p) = 0$ and $P_0(\hat{L})x(p) = x(p)$ from the second order recurrence. Notice that Eq. (6) represents the convolution operation as a simple linear combination of $c_{s,n}$ and P_n without u_j , and it is often approximated at the order of m for practical purposes.

Learning to Approximate Kernel Convolution

We introduce our model, i.e., LSAP, that *trains on the approximations* for the optimal range of neighborhood at individual nodes. The derivatives to train s in Eq. (6) for two separate tasks, i.e., node classification and graph classification, are derived in their closed-forms.

Model Architecture

In most of GCN frameworks (Kipf et al. 2017; Yang et al. 2021), a convolution operation at the k -th layer is given as

$$H_k = \sigma_k(\tilde{A}H_{k-1}W_k) \quad (7)$$

where \tilde{A} is a normalized adjacency matrix, W_k is a trainable weight matrix, and σ_k is a non-linear activation function. It takes an input H_{k-1} and outputs a new representation H_k . Each convolution operation takes features from direct neighbors of each node according to \tilde{A} to design the new node representation. As more convolution layers are stacked, the range of aggregation is uniformly increased for all nodes causing the infamous “oversmoothing” issue. To alleviate oversmoothing, we propose to utilize a diffusion kernel, i.e., a heat kernel, on the graph to define ranges of neighborhood for individual nodes, so that the node-wise range is adaptively defined to avoid the oversmoothing.

The architecture of LSAP is given in Fig. 1. The overall components are similar to the original GCN (Kipf et al. 2017), however, LSAP redefines the convolution with approximated heat kernel. Consider a graph G given as a Laplacian \hat{L} , feature X defined on its nodes and either node-wise or graph-wise label Y . Our framework takes \hat{L} and X as inputs, performs approximated heat kernel convolutions and outputs a prediction \hat{Y} . The \hat{Y} is then compared with the ground truth Y , and the error is backpropagated to update model parameters including the scale s .

Convolution Layer. Based on Eq. (6), Eq. (7) is reformulated by replacing the normalized adjacency matrix \tilde{A} with the heat kernel with polynomial approximation as

$$H_k = \sigma_k([\sum_{n=0}^m c_{s,n} P_n(\hat{L})]H_{k-1}W_k). \quad (8)$$

While \tilde{A} let the model combine information from nodes within 1-hop distance only, Eq. (8) let it aggregate information within a “range” of each node defined by s within $c_{s,n}$. The Eq. (8) defines a convolution layer, and we can stack K of them to achieve a better representation of the original X .

Output Layer. The output layer yields a prediction of Y , i.e., \hat{Y} , via softmax. Depending on the task, it may include a simple Multi-layer Perceptron (MLP) that can be trained. A loss L_{err} is computed at this layer which quantifies the error between \hat{Y} and Y using cross-entropy for different tasks, e.g., node classification or graph classification.

Model Update. The loss L_{err} is backpropagated to update the model parameters, i.e., W_k and s . To maintain the scale positive, an ℓ_1 -norm regularization on s is imposed. The overall objective function \mathcal{L} is given as

$$\mathcal{L} = \mathcal{L}_{\text{err}} + \alpha |\mathbf{s}| \quad (9)$$

where α is a hyperparameter. The weight W_k can be easily trained with backpropagation, and a multi-variate \mathbf{s} across all nodes can be also trained given a gradient on scale s as

$$\mathbf{s} \leftarrow \mathbf{s} - \beta_s \frac{\partial \mathcal{L}}{\partial \mathbf{s}} \quad (10)$$

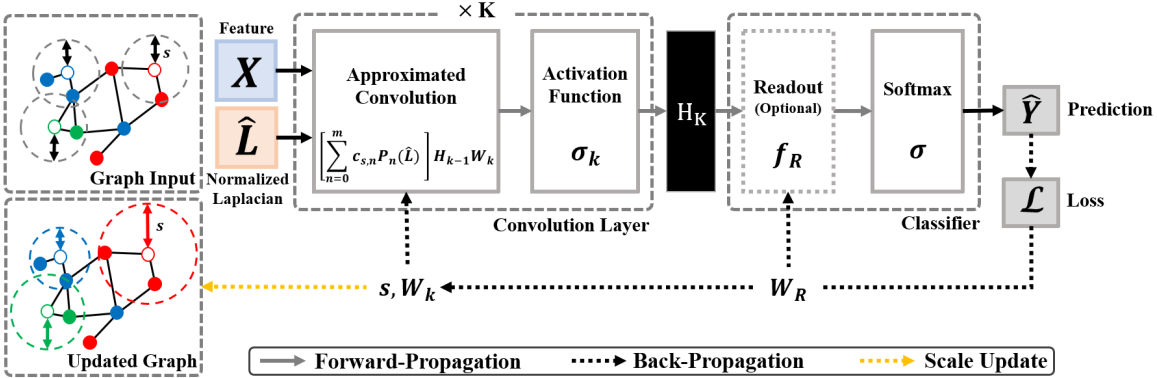


Figure 1: Illustration of LSAP. A graph (as normalized Laplacian \hat{L}) and node feature X are inputted to the convolution layer. The output H_K is inputted to a downstream classifier which yields a prediction \hat{Y} . The loss from \hat{Y} is backpropagated to update the classifier and convolution approximation with $s = [s_1, \dots, s_N]$ to adaptively adjust the scale of each node.

where β_s is a learning rate. It requires $\frac{\partial \mathcal{L}}{\partial s}$ to make the framework trainable, and we derive the $\frac{\partial \mathcal{L}}{\partial s}$ in a closed-form to train on the approximations of Eq. (8) in the following.

Gradients of Polynomial Coefficients with Scale

We denote expansion coefficients as $c_{s,n}^T$, $c_{s,n}^H$ and $c_{s,n}^L$ that correspond to P_n^T , P_n^H and P_n^L . As introduced in (Huang et al. 2020), one can obtain a solution to the heat diffusion by obtaining expansion coefficient with each polynomial in P_n . In order to design a gradient-based “learning” framework of node-wise range (i.e., scale) based on these expansions, we derived gradients of loss $\frac{\partial \mathcal{L}}{\partial s}$ in closed-forms. This is an essential component of LSAP as it let the model efficiently train without diagonalization of \hat{L} . The $\frac{\partial \mathcal{L}}{\partial s}$ can be achieved using the chain rule in a traditional way, and to obtain $\frac{\partial \mathcal{L}}{\partial s}$ in terms of the H_k , we compute $\frac{\partial c_{s,n}}{\partial s}$ for each coefficient.

Chebyshev Polynomial. The recurrence relation and expansion coefficient for Chebyshev polynomial are given as

$$P_{n+1}^T(\hat{L}) = (2 - \delta_{n0})\hat{L}P_n^T(\hat{L}) - P_{n-1}^T(\hat{L}), \\ c_{s,n}^T = (2 - \delta_{n0})(-1)^n e^{-\frac{sb}{2}} I_n\left(\frac{sb}{2}\right) \quad (11)$$

where $b > 0$ is a hyper-parameter, and I_n is the *modified Bessel function of the first kind* (Olver et al. 2010).

Hermite Polynomial. The recurrence relation and expansion coefficient for Hermite polynomial are written as

$$P_{n+1}^H(\hat{L}) = 2\hat{L}P_n^H(\hat{L}) - 2nP_{n-1}^H(\hat{L}), \\ c_{s,n}^H = \frac{1}{n!} \left(\frac{-s}{2}\right)^n e^{\frac{s^2}{4}}, \quad (12)$$

Laguerre Polynomial. For Laguerre polynomial, the recurrence relation and expansion coefficient are

$$P_{n+1}^L(\hat{L}) = \frac{(2n + 1 - \hat{L})P_n^L(\hat{L}) - nP_{n-1}^L(\hat{L})}{n + 1}, \\ c_{s,n}^L = \frac{s^n}{(s + 1)^{n+1}}. \quad (13)$$

Notice that all the expansion coefficients above are defined by s . If they are differentiable with respect to s , then we do not need to learn expensive parameters but simply train on these polynomial approximations with s directly.

Lemma 1. Consider an orthogonal polynomial P_n over interval $[a, b]$ with inner product $\int_a^b P_n(\lambda)P_k(\lambda)w(\lambda)d\lambda = \delta_{nk}$, where $w(\lambda)$ is the weight function. If P_n expands the heat kernel, the expansion coefficients $c_{s,n}$ with respect to s are differentiable and $\frac{\partial c_{s,n}}{\partial s} = -\int_a^b \lambda e^{-s\lambda} P_n(\lambda)w(\lambda)d\lambda$.

Lemma 1 (with a proof in the supplementary) says that the $c_{s,n}$ in Eq. (11), (12) and (13) have the derivatives with respect to s . These will be used in the following two sections to define gradients on loss functions for different tasks.

Semi-supervised Node Classification

The goal of node classification is to predict labels of unlabeled nodes based on the information from other nodes. The output layer (after K -convolution layers) of LSAP produces a prediction $\hat{Y} = \sigma(H_K)$, and the training should be performed to reduce the error between \hat{Y} and the true Y .

Lemma 2. Let a graph convolution be operated by Eq. (8), which approximates the convolution with P_n and $c_{s,n}$. If a loss \mathcal{L}_{err} for node-wise classification is defined as cross-entropy between a prediction $\hat{Y} = \sigma(H_K)$, where $\sigma(\cdot)$ is a softmax function, and the true Y , then

$$\frac{\partial \mathcal{L}_{err}}{\partial s} = (\hat{Y} - Y) \times \sigma'_k \left(\left[\sum_{n=0}^m c_{s,n} P_n(\hat{L}) \right] H_{k-1} W_k \right) W_k^T \\ \times \left(\sum_{n=0}^m P_n(\hat{L}) H_{k-1}^T + \left[\sum_{n=0}^m c_{s,n} P_n(\hat{L}) \right] \frac{\partial H_{k-1}}{\partial s} \right) \frac{\partial c_{s,n}}{\partial s} \quad (14)$$

where σ'_k is the derivative of σ_k .

Lemma 2 let LSAP backpropagate the error to update s to obtain the optimal scale per node for node classification.

Graph Classification

Consider a population of graphs $\{G_t\}_{t=1}^T$ with corresponding labels $\{\hat{Y}_t\}_{t=1}^T$, and learning a graph classification model

finds a function $f(G_t) = Y_t$. For this, the output layer consists of a readout layer $f_R(\cdot)$ (i.e., MLP with ReLU) and a softmax $\sigma(\cdot)$ at the end to construct pseudo-probability for each class. The $f_R(\cdot)$ with weights W_R takes the output H_K from the convolution layers as an input and returns H_R as

$$H_R = f_R(H_K; W_R), \quad (15)$$

and prediction $\hat{Y} = \sigma(H_R)$.

Lemma 3. Let H_k from Eq. (8) be a graph convolution with a heat kernel with polynomial P_n and coefficients $c_{s,n}$. If a loss \mathcal{L}_{err} for classifying graph-wise label is defined as cross-entropy between a prediction $\hat{Y} = \sigma(H_R)$, where $\sigma(\cdot)$ is a softmax function, and the true Y , then

$$\begin{aligned} \frac{\partial \mathcal{L}_{err}}{\partial s} &= (\hat{Y} - Y) \times \frac{\partial H_R}{\partial H_K} \times \sigma'_k \left(\sum_{n=0}^m c_{s,n} P_n(\hat{L}) \right) H_{k-1} W_k W_k^\top \\ &\times \left(\sum_{n=0}^m P_n(\hat{L}) H_{k-1}^\top + \left[\sum_{n=0}^m c_{s,n} P_n(\hat{L}) \right] \frac{\partial H_{k-1}}{\partial c_{s,n}} \right) \frac{\partial c_{s,n}}{\partial s} \end{aligned} \quad (16)$$

where σ'_k is the derivative of σ_k .

Lemma 3 let LSAP adaptively update the scale s across all nodes for each node using Eq. (10) towards the minimal error for predicting graph-wise label.

Experiments

In this section, we compare the performances of LSAP and baselines on node classification and graph classification. **LSAP-C**, **LSAP-H** and **LSAP-L** correspond to approximation frameworks with P_n^T , P_n^H and P_n^L , and the model with exact computation of the heat kernel convolution is referred as **Exact**. For the node classification, we used conventional benchmarks for semi-supervised learning task (Shchur et al. 2018). For the graph classification, we investigated real brain network data from Alzheimer’s Disease Neuroimaging Initiative (ADNI) to classify different diagnostic stages towards Alzheimer’s Disease (AD) for a practical application. These tasks on seven different benchmarks can demonstrate the feasibility of LSAP.

Semi-supervised Node Classification

Datasets. We conducted experiments on standard node classification datasets (in Table 1) that provide connected and undirected graphs. Cora, Citeseer and Pubmed (Sen et al. 2008) are constructed as citation networks, Amazon Computer and Amazon Photo (Shchur et al. 2018) define co-purchase networks, and Coauthor CS (Shchur et al. 2018) is a co-authorship network.

Dataset	Nodes	Edges	Classes	Features
Cora	2,708	5,429	7	1,433
Citeseer	3,327	4,732	6	3,703
Pubmed	19,717	44,338	3	500
Amazon Computers	13,752	245,861	10	767
Amazon Photo	7,650	119,081	8	745
Coauthor CS	18,333	81,894	15	6,805

Table 1: Summary of node classification datasets.

Setup. We used the accuracy as the evaluation metric. For Cora, Citeseer and Pubmed data, eighteen different baselines were used to compare the results for the node classification task as listed in Table 2. These standard benchmarks are provided with fixed split of 20 nodes per class for training, 500 nodes for validation and 1000 nodes for testing as in other literature (Kim et al. 2022; Wu et al. 2022b).

For Amazon and Coauthor datasets, seven baselines are used as in Table 3. For the MLP with 3-layers, GCN and 3ference, results are obtained from (Luo et al. 2022), and a result for DSF comes from (Guo et al. 2023). For others, the experiments were performed by randomly splitting the data as 60%/20%/20% for training/validation/testing datasets as in (Luo et al. 2022) and replicating it 10 times to obtain mean and standard deviation of the evaluation metric.

Results. Table 2 and 3 show the performance comparisons between LSAP and baseline models. As shown in Table 2, on the node classification benchmarks, learning node-wise adaptive scale performs the best in both Exact and its approximations. LSAP showed improved performance over existing models; exceeding previous best baseline performances by 2.4% (on Cora) and 1.1% (on Citeseer and Pubmed). The similar performance of LSAP with that of Exact demonstrates accurate convolution approximation. Despite slight decreases, training on adaptive scales using LSAP was much faster.

For additional datasets in Table 3, the performance of LSAP outperformed the baselines. The results for MLP, GCN and 3ference were adopted from (Luo et al. 2022), which reported the best performance out of 10 replicated experiments. We ran the same experiments for GAT, GDC, GraphHeat, Exact and LSAP, and the mean and standard deviation of metrics are given. LSAP shows significant improvements on the Amazon Computer (94.43%, LSAP-C) and Amazon Photo (96.65%, LSAP-H). On the Coauthor

Model	Cora	Citeseer	Pubmed
GCN (Kipf et al. 2017)	81.50	70.30	78.60
GAT (Veličković et al. 2018)	83.00	72.50	79.00
APPNP (Klicpera et al. 2019b)	83.30	71.80	80.10
GDC [†] (Klicpera et al. 2019a)	82.20	71.80	79.10
SGC (Wu et al. 2019)	81.70	71.30	78.90
Bayesian GCN (Zhang et al. 2019a)	81.20	72.20	-
Shoestring (Lin, Gao, and Li 2020)	81.90	69.50	79.70
GraphHeat [†] (Xu et al. 2020)	83.70	72.50	80.50
g-U-Nets (Gao et al. 2019)	84.40	73.20	79.60
GCNII (Chen et al. 2020)	85.50	73.40	80.30
GRAND (Feng et al. 2020)	85.40	75.40	82.70
DAGNN (Liu et al. 2020)	84.40	73.30	80.50
SelfSAGCN (Yang et al. 2021)	83.80	73.50	80.70
DIAL-GNN (Chen et al. 2021)	84.50	74.10	-
SuperGAT (Kim et al. 2022)	84.30	72.60	81.70
GRAND [†] (Chamberlain et al. 2021)	83.60	74.10	78.80
ADC [†] (Zhao et al. 2021)	84.50	74.50	83.00
SEP-N (Wu et al. 2022b)	84.80	72.90	80.20
LSAP-C	87.90	76.50	83.30
LSAP-H	85.00	76.10	82.60
LSAP-L	85.90	75.90	84.10
Exact	88.20	78.10	85.30

[†]: graph diffusion-based models.

Table 2: Accuracy (%) on Cora, Citeseer, and Pubmed. LSAP yields better performances over existing baselines (in bold) similar to Exact achieving the best results (underline).

Model	Amazon Computer	Amazon Photo	Coauthor CS
MLP (3-layers)	84.63	91.96	95.63
GCN	90.49	93.91	93.32
3ference	90.74	95.05	95.99
GAT	91.18 ± 0.74	94.49 ± 0.54	93.42 ± 0.31
GDC	86.03 ± 2.26	93.28 ± 1.03	92.68 ± 0.53
GraphHeat	89.59 ± 3.15	94.04 ± 0.75	92.93 ± 0.20
DSF	92.84 ± 0.10	95.73 ± 0.08	-
LSAP-C	94.43 ± 1.16	95.96 ± 1.65	94.81 ± 0.55
LSAP-H	93.64 ± 0.86	96.65 ± 0.67	93.52 ± 0.97
LSAP-L	92.76 ± 0.48	95.35 ± 0.85	93.58 ± 0.82
Exact	93.52 ± 0.65	96.41 ± 1.54	93.71 ± 1.16

Table 3: Mean node classification accuracy (%) and s.d. on Amazon Computers, Amazon Photo, and Coauthor CS. The best results are in bold, and the best results within experiments with replicates are underlined.

CS, we also achieve the highest mean accuracy (94.81%, LSAP-C) among the experiments with random replicates.

Graph Classification

Datasets. Using the magnetic resonance images (MRI) from the ADNI data, each brain was partitioned into 148 cortical regions and 12 sub-cortical regions using Destrieux atlas (Destrieux et al. 2010), and tractography on diffusion-weighted imaging (DWI) was applied to calculate the number of white matter fibers connecting the 160 brain regions to construct 160×160 structural network (i.e., graph). On the same parcellation, region-wise imaging features such as Standard Uptake Value Ratio (SUVR) of metabolism level from FDG-PET and cortical thickness from MRI were measured. For the SUVR normalization, Cerebellum was used as the reference. The dataset consists of 5 AD-specific progressive groups: Control (CN), Significant Memory Concern (SMC), Early Mild Cognitive Impairment (EMCI), Late Mild Cognitive Impairment (LMCI) and AD. The demographics of ADNI dataset are summarized in Table 4.

Setup. 5-way classification was designed to classify the different groups in Table 4. 5-fold cross validation was used to obtain unbiased results, and accuracy, precision, and recall in their mean were used as evaluation metrics. As the baseline, we adopted Linear Support Vector Machine (SVM), Multi-Layer Perceptron (MLP) with 2 layers, GCN (Kipf et al. 2017), GAT (Veličković et al. 2018), GDC (Klicpera et al. 2019a), GraphHeat (Xu et al. 2020) and ADC (Zhao et al. 2021). Each sample is given with a graph (i.e., brain network) and two node features (i.e., cortical thickness and FDG measure) which are well-known as useful biomarkers for AD diagnosis.

Biomarker	Category	CN	SMC	EMCI	LMCI	AD
Cortical Thickness	# of subjects	359	181	437	180	166
	Gender (M / F)	178 / 181	69 / 112	249 / 188	119 / 61	102 / 64
	Age (Mean±Std)	72.8±1.4	72.0±5.2	71.0±7.9	70.9±6.1	74.8±8.7
FDG	# of subjects	345	186	461	231	162
	Gender (M / F)	173 / 172	66 / 120	262 / 199	152 / 79	102 / 60
	Age (Mean±Std)	73.0±1.3	71.7±5.2	71.7±7.8	71.1±7.0	74.9±8.8

Table 4: Demographics of the ADNI dataset.

Feature	Model	Classification (ADNI)		
		Accuracy (%)	Precision	Recall
Cortical Thickness	SVM (Linear)	82.39 ± 2.73	0.822 ± 0.033	0.852 ± 0.025
	MLP (2-layers)	78.76 ± 2.21	0.792 ± 0.036	0.799 ± 0.026
	GCN	61.37 ± 3.09	0.598 ± 0.025	0.626 ± 0.044
	GAT	64.17 ± 5.46	0.627 ± 0.067	0.668 ± 0.046
	GDC	77.10 ± 4.25	0.769 ± 0.050	0.785 ± 0.044
	GraphHeat	70.90 ± 3.17	0.703 ± 0.030	0.718 ± 0.026
	ADC	82.10 ± 2.41	0.776 ± 0.019	0.728 ± 0.067
	LSAP-C	87.00 ± 2.16	0.868 ± 0.027	0.885 ± 0.027
	LSAP-H	85.41 ± 2.32	0.859 ± 0.031	0.867 ± 0.030
	LSAP-L	85.64 ± 1.86	0.859 ± 0.022	0.866 ± 0.022
FDG	Exact	86.24 ± 1.96	0.866 ± 0.017	0.867 ± 0.023
	SVM (Linear)	85.27 ± 2.09	0.857 ± 0.027	0.869 ± 0.021
	MLP (2-layers)	87.51 ± 1.62	0.882 ± 0.024	0.882 ± 0.014
	GCN	68.81 ± 1.95	0.677 ± 0.028	0.697 ± 0.025
	GAT	69.24 ± 7.13	0.670 ± 0.106	0.736 ± 0.037
	GDC	86.21 ± 3.24	0.867 ± 0.033	0.870 ± 0.029
	GraphHeat	76.97 ± 2.42	0.775 ± 0.035	0.773 ± 0.010
	ADC	88.60 ± 2.81	0.708 ± 0.062	0.753 ± 0.053
	LSAP-C	89.24 ± 2.23	0.895 ± 0.022	0.904 ± 0.023
	LSAP-H	90.11 ± 2.44	0.903 ± 0.027	0.910 ± 0.022
	LSAP-L	90.40 ± 1.38	0.909 ± 0.018	0.914 ± 0.015
	Exact	90.18 ± 2.67	0.907 ± 0.028	0.907 ± 0.028

Table 5: Classification performances on ADNI dataset.

Results. The performances including accuracy, precision, and recall between LSAP and seven baselines on the ADNI dataset are shown in Table 5. LSAP showed accuracy $\sim 86\%$ using cortical thickness and $\sim 90\%$ with FDG in classifying the 5 diagnostic stages of AD, with precision and recall ~ 0.86 and 0.91 , respectively. These numbers are approximately the same with the results from Exact and low standard deviations from LSAP demonstrate feasibility of the approximation. LSAP performed the best surpassing the second best methods by 4.61%p and 1.80%p in accuracy for cortical thickness and FDG experiments, respectively.

Model Behavior Analysis

Computation Time with Kernel Convolution. Fig. 2 compares averaged empirical time (in ms) spent for one epoch of training process of Exact and LSAP on node classification task (on Cora, Citesser and Pubmed) and graph classification task (on ADNI) with 10 replicates. The colors denote the type of methods, and as seen in Fig. 2, LSAP takes

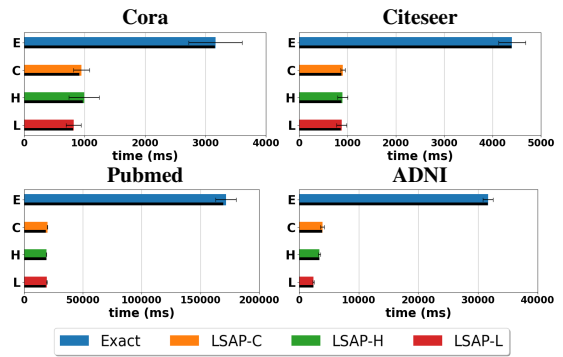


Figure 2: Comparisons of computation time (in ms) for one epoch (Forward and backpropagation). Within the epoch, time for heat kernel convolution is given in black bar. Results were obtained with 10 repetitions.

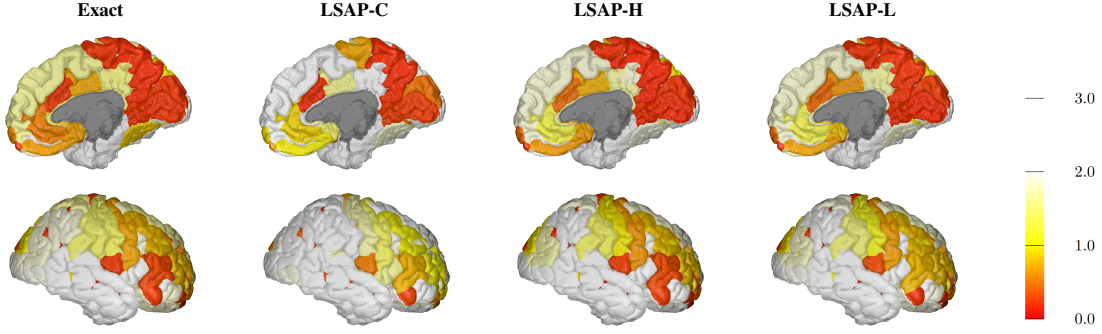


Figure 3: Visualization of the learned scales on the cortical regions of a brain. This visualization shows the scale of each ROI from the classification result using FDG feature. Top: Inner part of right hemisphere, Bottom: Outer part of right hemisphere.

ROI (Destrieux et al. 2010)	Exact	LSAP-C	LSAP-H	LSAP-L
(L) G&S.paracentral	0.034	0.052	0.049	0.066
(L) G.front.inf.Orbital	0.036	0.071	0.060	0.043
(R) G.precuneus	0.041	0.044	0.034	0.051
(R) S.ortibal.med.olfact	0.047	0.078	0.054	0.059
(R) G.cingul.Post.ventral	0.055	0.056	0.055	0.051
(R) S.oc.temp.lat	0.055	0.065	0.045	0.063
(R) G.oc.temp.med.Lingual	0.055	0.076	0.043	0.040
(L) Sub.put	0.058	0.077	0.047	0.060
(L) S.postcentral	0.061	0.069	0.060	0.018
(R) G.front.inf.Orbital	0.063	0.093	0.069	0.050

Table 6: 10 ROIs with the smallest trained scales for AD classification. (L)/(R) denote the left/right hemisphere.

far less time than Exact computation. Notice that the computation of kernel convolution takes the majority of time (in black bar), and the approximations make this process efficient. For the node classification, approximation on Pubmed showed the best efficiency as its graph had the largest number of nodes (19717) compared to Cora (2708) and Citeseer (3327). For the graph classification, approximations were even more efficient as eigendecomposition of \hat{L} from all subjects had to be performed for Exact. Comparing the computation time of a single epoch on Exact and LSAP-L, the time is saved by $\sim 93\%$.

Discussions on the Scales for Graph Classification. In AD classification, we performed graph classification to distinguish the different diagnostic labels of AD. The trained model yields node-wise optimized scale where the node corresponds to specific region of interest (ROI) in the brain. The trained scales denote the optimal ranges of neighborhood for each ROI. Therefore, if the trained scale is small for a specific node, it means that the node does not have to look far to contribute to the classification. On the other hand, the nodes with large scales need to aggregate information from far distances to constitute an effective embedding as it is not very useful on its own. The trained scales on the brain network classification with Exact and LSAP are visualized in Fig. 3 conveying two important perspectives. First, the scales delineate which of the ROIs are independently behaving to classify AD-specific labels. Second, the trained scales with LSAP are quite similar to the result from Exact meaning that the approximation is feasibly accurate for practical uses.

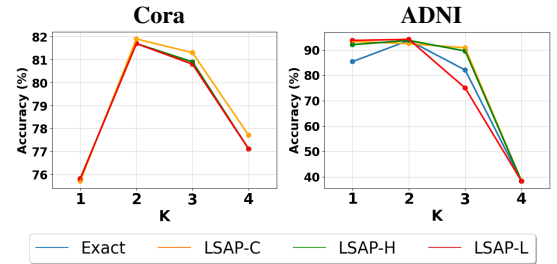


Figure 4: Effect of the number of layers K on model performance. Left: accuracy of node classification on Cora, Right: accuracy of graph classification on ADNI.

In Table 6, 10 ROIs with the smallest scales that appear in common across Exact and LSAPs are listed. *Inferior frontal orbital gyrus* on both hemispheres is captured, and several *temporal/orbital regions*, *precuneous*, and *left putamen* are shown to yield small scales. These ROIs are known as highly AD-specific by various literature (Galton et al. 2001; Bailly et al. 2015; de Jong et al. 2008; Hoesen et al. 2000).

Effect of K . We examined the performance of LSAP with respect to the number of convolution layers K on Cora and ADNI experiments. When K was varied from 1 to 4 under the same setting, $K=2$ showed the best performance in both experiments. The performance decreases when $K=3$ and 4 may be due to lack of training samples as the model sizes are drastically increased. We observed the same pattern across Exact and LSAP, which demonstrates LSAP with approximation is able to train on the scales properly.

Conclusions

In this work, we proposed efficient trainable methods to bypass exact computation of spectral kernel convolution that define adaptive ranges of neighbor for each node. We have derived closed-form derivatives on polynomial coefficients to train the scale with conventional backpropagation, and the developed framework LSAP demonstrates SOTA performance on node classification and brain network classification. The brain network analysis provides neuroscientifically interpretable results corroborated by previous AD literature.

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