Seminar Series on Graph Neural Networks 03 A graph signal processing viewpoint of graph neural networks

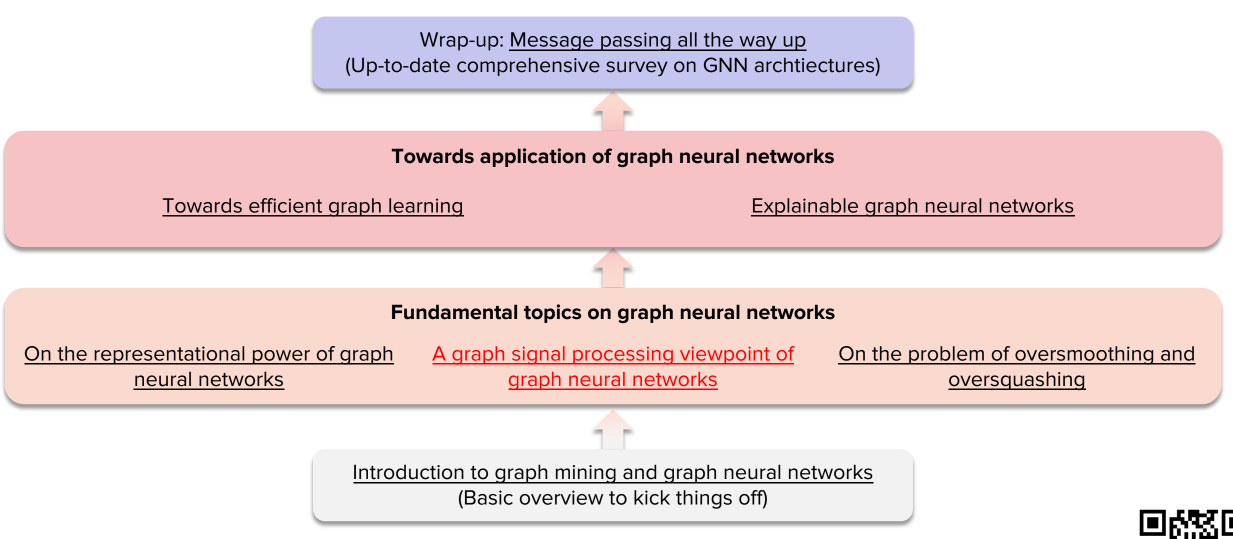
Yong-Min Shin School of Mathematics and Computing (Computational Science and Engineering) Yonsei University 2025.04.14







Before going in....





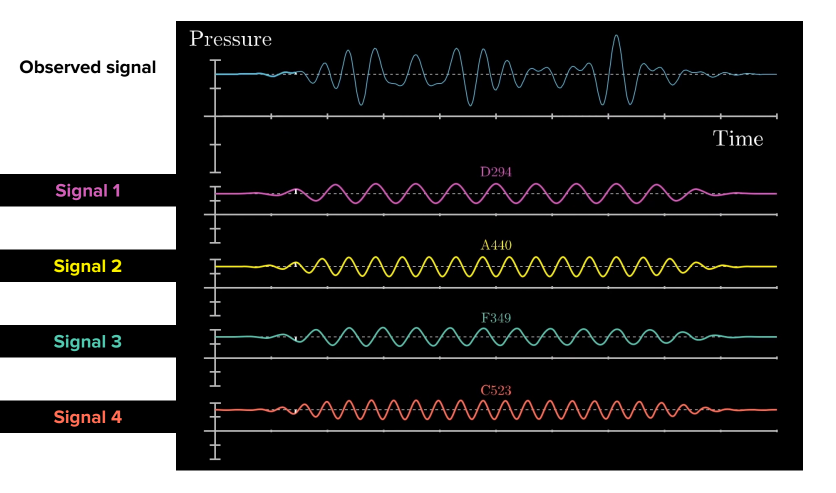
* Presentation slides are available at: (jordan7186.github.io/presentations/)

(Some of the topics may change in the future for a better alternative)

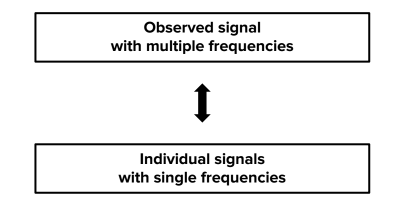
Objectives

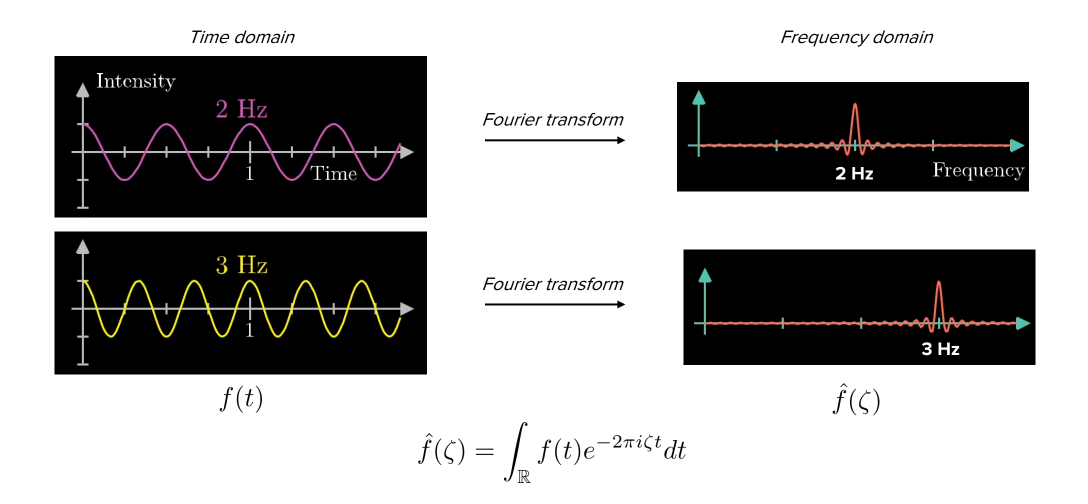
- 1. Preliminary: Singal processing (3blue1brown)
- 2. Understanding of graph signals & graph Fourier transform
- 3. Understanding the **formulation of ChebNet**
- 4. Re-reading GCN, understanding in the original author's way

Preliminary: Signal processing

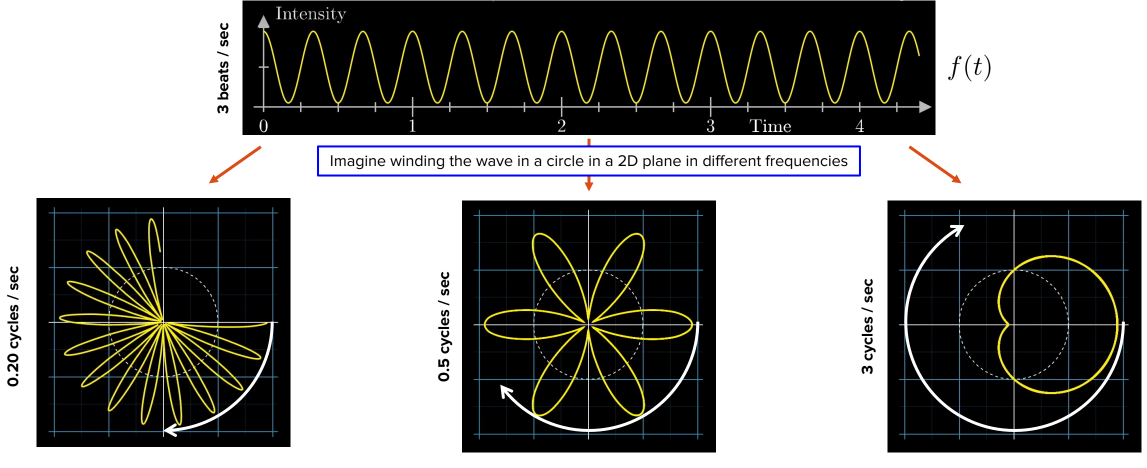


Fourier transforms can be used to analyze signals





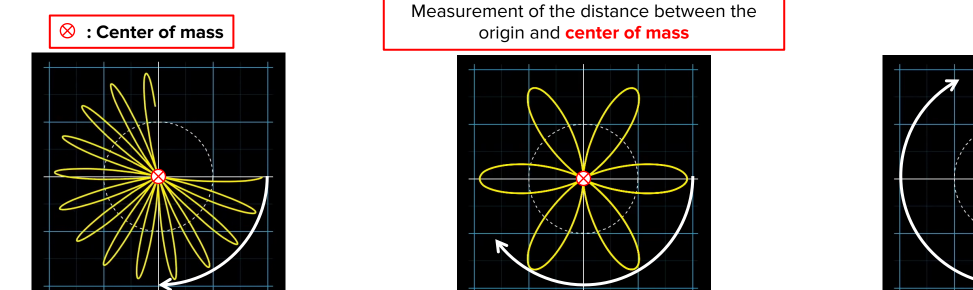
3Blue1Brown, https://www.youtube.com/watch?v=spUNpyF58BY

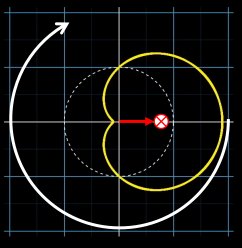


Observation

7

Something unique happens when the winding frequency *exactly* matches the signal frequency





 $\quad \Rightarrow \quad \hat{f}(\zeta) = \int_{\mathbb{R}} f(t) e^{-2\pi i \zeta t} dt$

Imagine winding the wave in a circle in a 2D plane in different frequencies

$$f(t)e^{-2\pi i\zeta t}$$

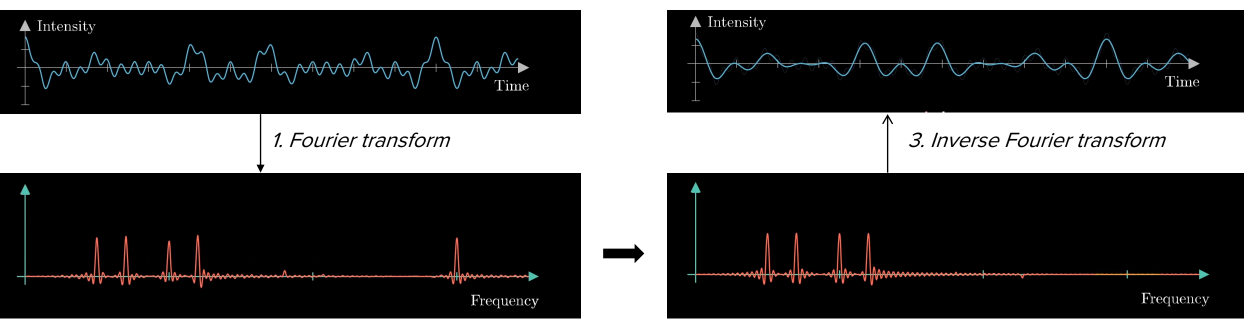
Measurement of the distance between the origin and **center of mass**

$$\frac{1}{Z} \int_{\mathbb{R}} f(t) e^{-2\pi i \zeta t} dt$$

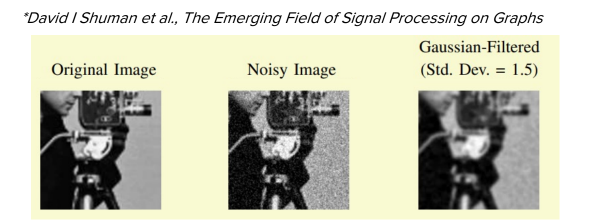
3Blue1Brown, https://www.youtube.com/watch?v=spUNpyF58BY

8

Low-pass filtering



2. Reduce <u>high-frequency</u> noise





3Blue1Brown, https://www.youtube.com/watch?v=spUNpyF58BY

Additional linear algebra: Spectral decomposition

Spectral Decomposition

The expression

$$A = PDP^7$$

is called the spectral decomposition of A. We can write it as

$$A = \begin{bmatrix} \vec{x_1} & \vec{x_2} & \cdots & \vec{x_n} \end{bmatrix} \begin{bmatrix} \lambda_1 & 0 & \cdots & \cdots & 0 \\ 0 & \lambda_2 & 0 & \cdots & 0 \\ \vdots & 0 & \ddots & \ddots & 0 \\ \vdots & \vdots & \ddots & \ddots & 0 \\ 0 & 0 & \cdots & 0 & \lambda_n \end{bmatrix} \begin{bmatrix} \vec{x_1}^T \\ \vec{x_2}^T \\ \vdots \\ \vec{x_n}^T \end{bmatrix}$$

- When decomposing the (symmetric) adjacency matrix, the eigenvalues are real.
- The eigenvalues are usually ordered.

$$\lambda_1 \ge \lambda_2 \ge \cdots \ge \lambda_n$$

We can also rearrange in an ascending order, just swap the corresponding eigenvectors accordingly.

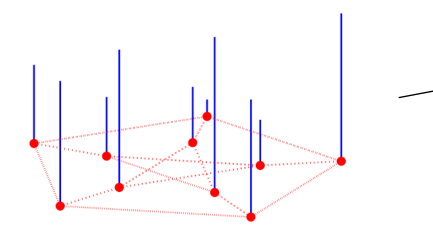
Dr. Ceni Babaoglu	cenibabaoglu.com
Linear Algebra for Machine Learning: Singular Value Decomposition and Principal Component Analysis	

Slide from: https://www.slideshare.net/CeniBabaogluPhDinMat/5-linear-algebra-for-machine-learning-singular-value-decomposition-and-principal-component-analysis-147828329

Understanding of graph signals & graph Fourier transforms

How should we think of signals on graphs?

We will consider a simple 1D node features, which can be easily extended to muti-dimensional case



We can imagine assigning a single value to each vertex Think of 1-dimensional feature matrix as a function

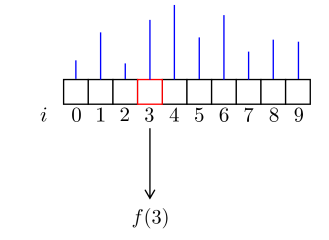
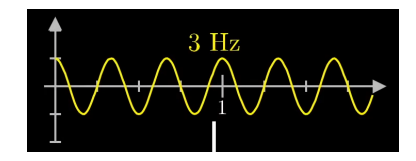


Fig. 1. A random positive graph signal on the vertices of the Petersen graph. The height of each blue bar represents the signal value at the vertex where the bar originates.



Generalization of Fourier transforms to graphs

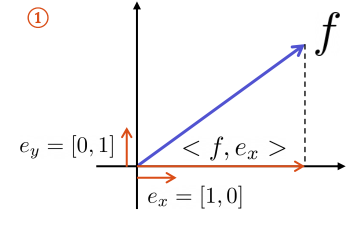
A GRAPH FOURIER TRANSFORM AND NOTION OF FREQUENCY

The classical Fourier transform

$$\hat{f}(\xi) := \langle f, e^{2\pi i \xi t} \rangle = \int_{\mathbb{R}} f(t) e^{-2\pi i \xi t} dt$$

is the expansion of a function f in terms of the complex exponentials, which are the eigenfunctions of the one-dimensional (1-D) Laplace operator

$$-\Delta \left(e^{2\pi i\xi t}\right) = -\frac{\partial^2}{\partial t^2} e^{2\pi i\xi t} = (2\pi\xi)^2 e^{2\pi i\xi t}.$$
 (2)



2

Fourier transform = Inner product with some function Some function = complex expoenetials?

3

Some function = <u>Eigenfunction</u> of the Laplace operator

$$\Delta = \frac{\partial^2}{\partial t^2} \qquad \Delta e^{2\pi i\xi t} = -(2\pi\xi)^2 e^{2\pi i\xi t}$$

Eigenfunction

Roughly, the Laplace operator measures the **local difference between the function and average**.

Generalization of Fourier transforms to graphs

Conclusion of the previous slide:

Fourier transform is the inner product between the target function and the eigenfunction of the Laplace operator.

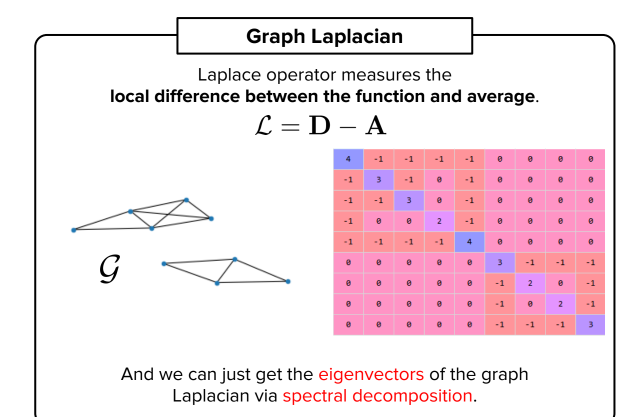
Graph signal! (# of nodes = N)

Analogously, we can define the graph Fourier transform \hat{f} of any function $f \in \mathbb{R}^N$ on the vertices of \mathcal{G} as the expansion of f in terms of the eigenvectors of the graph Laplacian:

$$\underline{\hat{f}(\lambda_{\ell}) := \langle \mathbf{f}, \mathbf{u}_{\ell} \rangle}_{i=1} = \sum_{i=1}^{N} f(i) u_{\ell}^{*}(i).$$
(3)

This part is the same from before

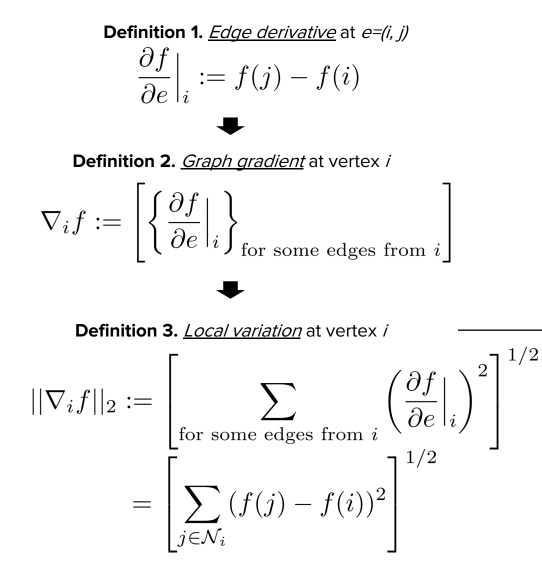
This part is different because now its discrete



Complementary note 1: On the physical intuition of the graph Laplacian

Why is Laplacian related to measuring the difference between the function and average?

Need some further generalizations of mathematical concepts... "Discrete calculus"

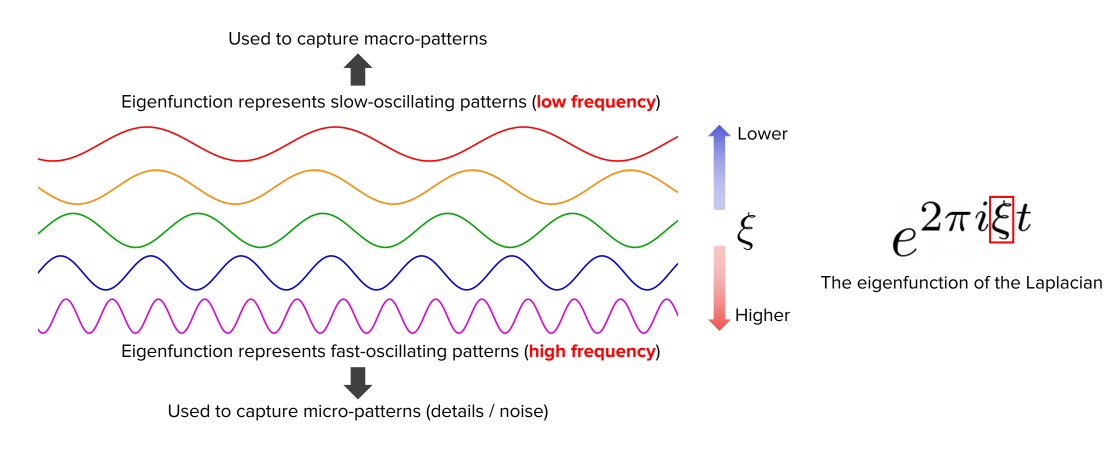


We can calculate the *total variation of the whole graph* as *the <u>sum of local variation</u> (squared) for all nodes in the graph:*

 $\frac{1}{2} \sum_{i \in V} \sum_{j \in \mathcal{N}_i} (f(j) - f(i))^2$

which is $f^T \mathcal{L} f$

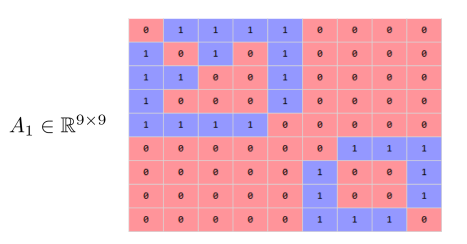
....So we can at least understand why the graph Laplacian is useful to <u>capture the patterns</u> of the graph signal. It measures how much the signal differs locally, eventually containing all information on variation of signals.

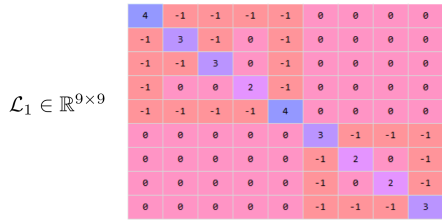


Is this also analogous to graph Fourier basis?



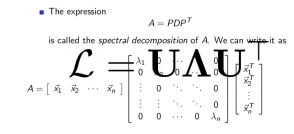






Concrete example 1

Spectral Decomposition

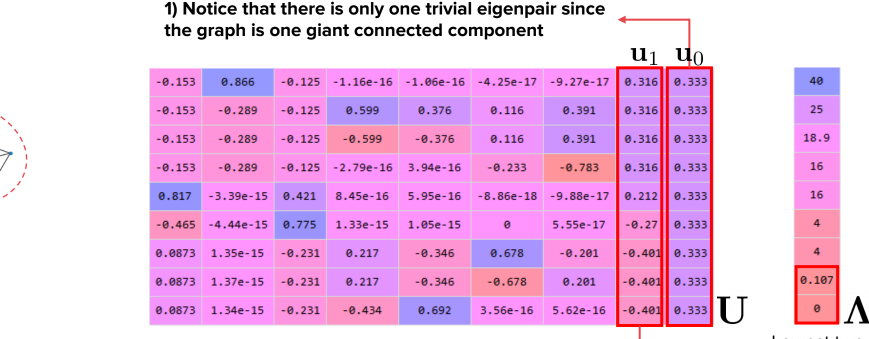


	Linear Algebra for Machine Learning: Singular Value Decomposition and Principal Component Analysis										
							\mathbf{u}_1	\mathbf{u}_0			
-2.32e-17	-0.894	-2.76e-18	0	0	0	-1.35e-16	0	0.447		25	
-0.289	0.224	0.707	0	0	0	0.408	0	0.447		25	
-0.289	0.224	-0.707	0	0	0	0.408	0	0.447		16	
-0.289	0.224	-5.67e-16	0	0	0	-0.816	0	0.447		16	
0.866	0.224	2.8e-16	0	0	0	-4.25e-16	0	0.447		16	
0	0	0	-0.866	1.02e-16	4.96e-17	0	0.5	0		4	
0	0	0	0.289	0.408	-0.707	0	0.5	0		4	
0	0	0	0.289	0.408	0.707	0	0.5	0	тт	ø	٨
0	0	0	0.289	-0.816	-1.6e-17	0	0.5	0	U	0	Δ
Matrix of eigenvectors Corresponding eigenvalues								es fre	Lowest two frequencies <i>(eigenvalues</i>		

Dr. Ceni Babaoglu

Notice that the <u>trivial</u> eigenpair reveal the most macro pattern in a graph: Number of connected compoents ("blobs")

Concrete example 2

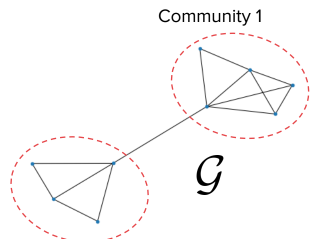


2) Still, also notice that the first non-trivial

eigenvectors returns a 'soft' community assignment,

which is the the next macro-pattern.

(HIGHLY incourage to read spectral clustering [1])

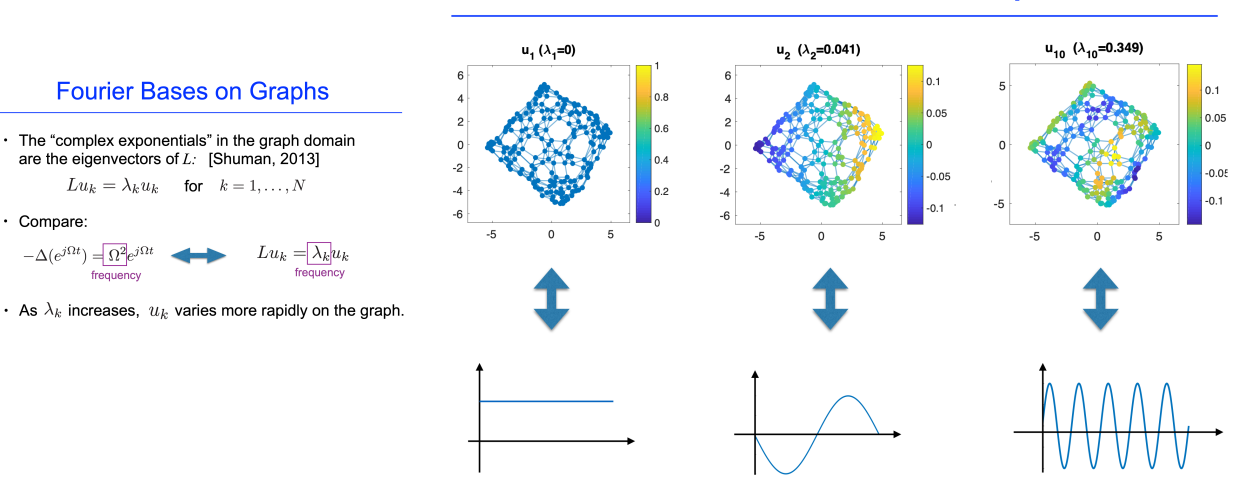


Community 2

Lowest two frequencies (eigenvalues)

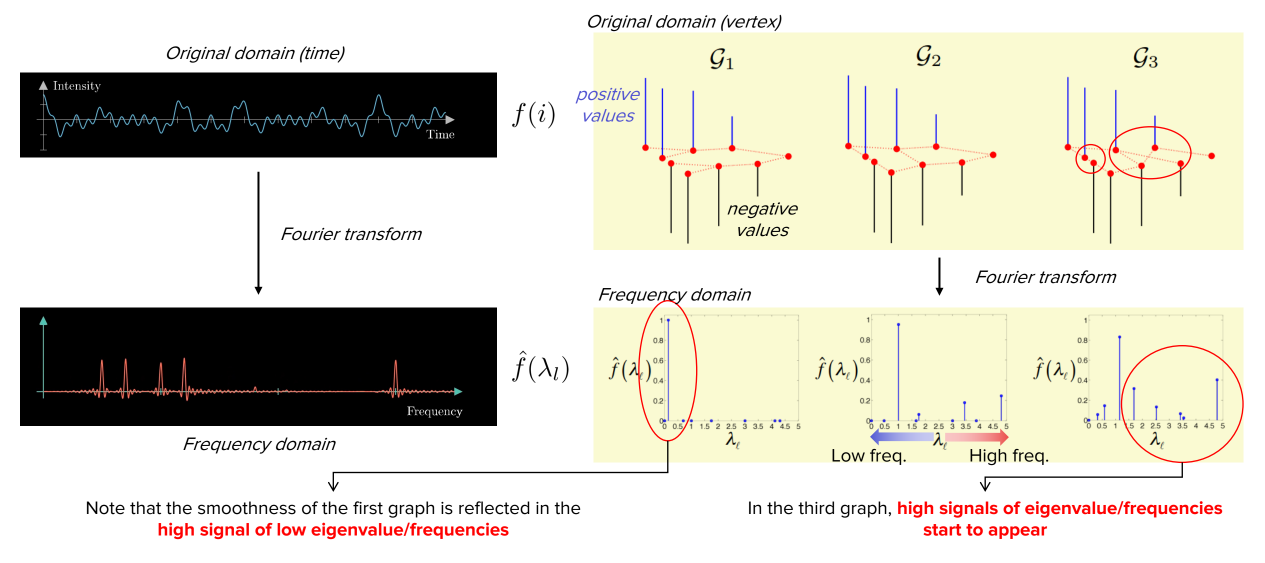
Some further examples... (external slide)

Fourier Bases on Graphs



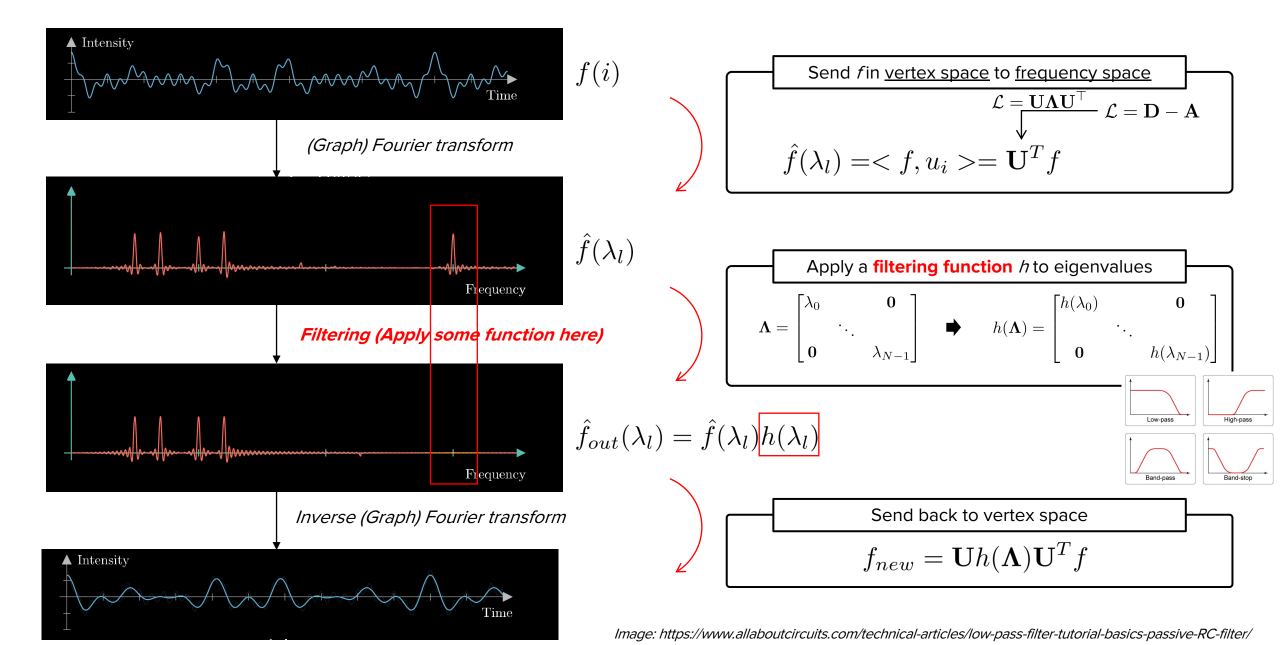
Slides from: Elif Vural, "Spectral Graph Theory and Graph Signal Processing", 2021, https://indico.truba.gov.tr/event/56/contributions/456/attachments/118/280/CizgeOkulu_ElifVural.pdf

...Coming back, completing the full graph Fourier transformation



* High frequency = rapid oscillation, high variance

...Coming back, completing the full graph Fourier transformation

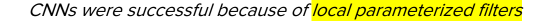


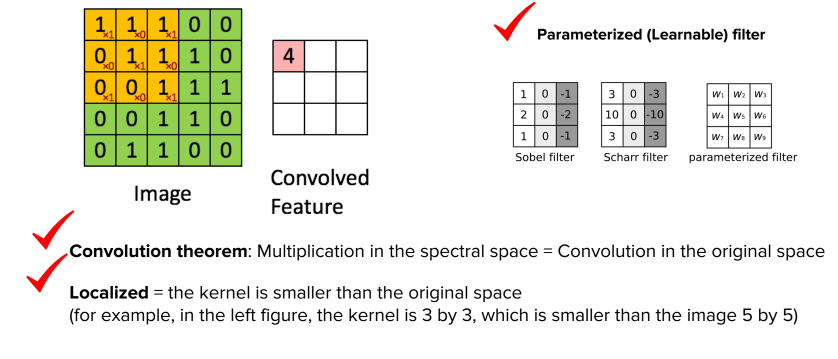
Understanding the formulation of ChebNet & GCN

Application of convolution (filters) to graphs

Defferrard et al., Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering

- 1. Spectral formulation: Extend the formulation of graph signal processing
- 2. Strictly localized filters: Design a local filter localized in K hop from the central vertex

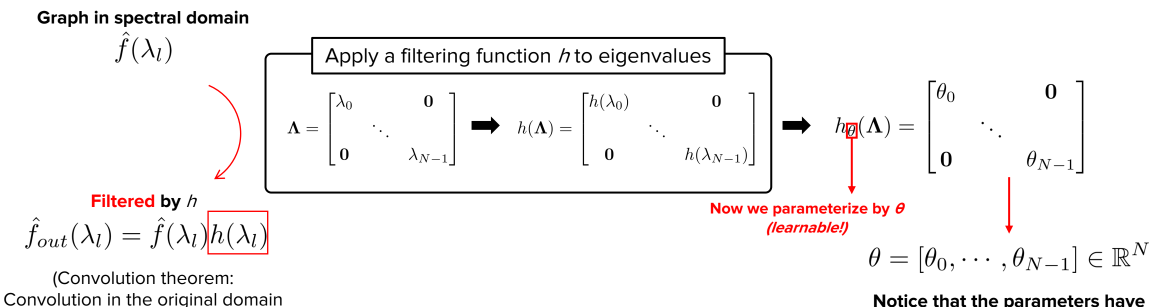




3. Low computational complexity: Expensive eigenvalue decomposition (spectral decomposition) is not needed

Localizing graph filters

Defferrard et al., Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering

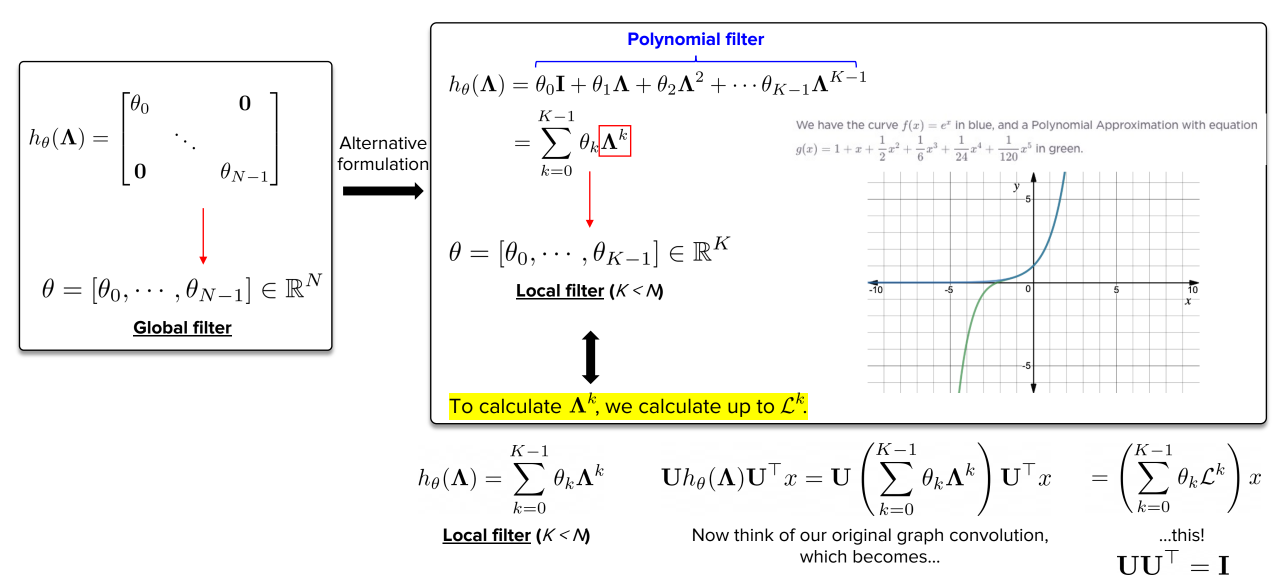


= Product in the Fourier domain)

Notice that the parameters have the same size as the input *N*

Localizing graph filters

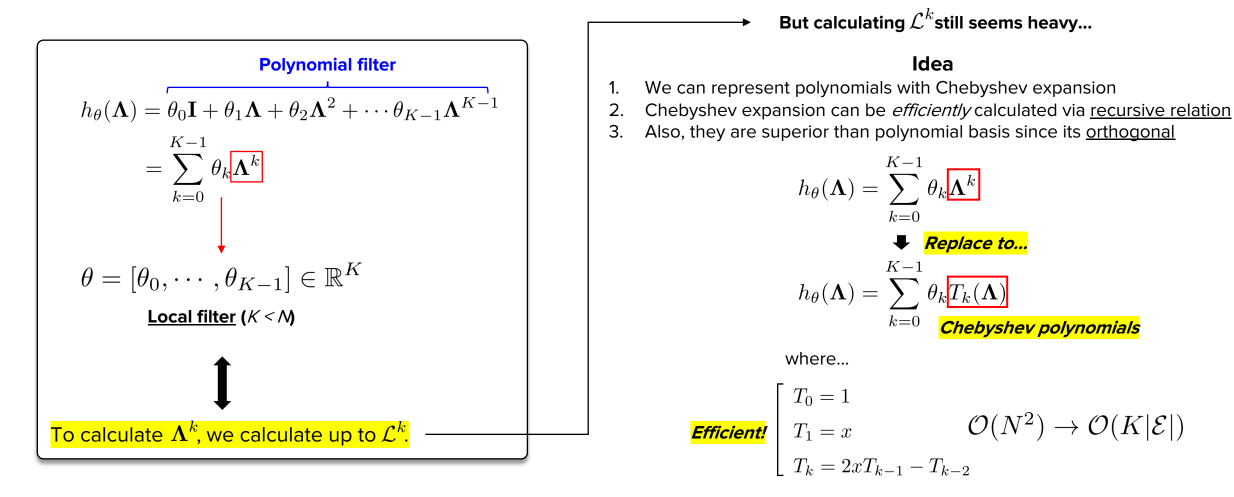
Defferrard et al., Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering



Polynomial approximation (right): https://www.expii.com/t/what-is-a-polynomial-approximation-317

Efficient implementation of local graph filters

Defferrard et al., Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering



Efficient implementation of local graph filters

Defferrard et al., Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering

conv.ChebConv

class ChebConv (in_channels: int, out_channels: int, K: int, normalization: Optional[str] = 'sym', bias: bool = True, **kwargs) [source]

Bases: MessagePassing

The chebyshev spectral graph convolutional operator from the "Convolutional Neural Networks on Graphs with Fast Localized Spectral Filtering" paper.

$$\mathbf{X}' = \sum_{k=1}^{K} \mathbf{Z}^{(k)} \cdot \mathbf{\Theta}^{(k)}$$

where $\mathbf{Z}^{(k)}$ is computed recursively by

$$egin{aligned} \mathbf{Z}^{(1)} &= \mathbf{X} \ \mathbf{Z}^{(2)} &= \mathbf{\hat{L}} \cdot \mathbf{X} \ \mathbf{Z}^{(k)} &= 2 \cdot \mathbf{\hat{L}} \cdot \mathbf{Z}^{(k-1)} - \mathbf{Z}^{(k-2)} \end{aligned}$$

and ${\bf \hat{L}}$ denotes the scaled and normalized Laplacian $\frac{2{\bf L}}{\lambda_{max}}-{\bf I}.$

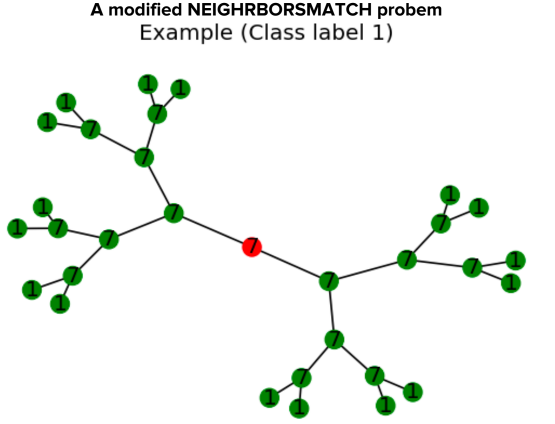
def forward(self, x: Tensor, edge_index: Tensor, edge_weight: OptTensor = None, batch: OptTensor = None, lambda_max: OptTensor = None,) -> Tensor: edge_index, norm = self.__norm__(edge_index, x.size(self.node dim), edge weight, self.normalization, lambda max, dtype=x.dtype, batch=batch, Tx 0 = xTx 1 = x # Dummy. out = self.lins[0](Tx_0) # propagate_type: (x: Tensor, norm: Tensor) if len(self.lins) > 1: Tx_1 = self.propagate(edge_index, x=x, norm=norm) out = out + self.lins[1](Tx 1) for lin in self.lins[2:]: Tx 2 = self.propagate(edge index, x=Tx 1, norm=norm) $Tx_2 = 2 * Tx_2 - Tx_0$ out = out + lin.forward(Tx 2) $Tx_0, Tx_1 = Tx_1, Tx_2$ if self.bias is not None: out = out + self.bias

[docs]

```
return out
```

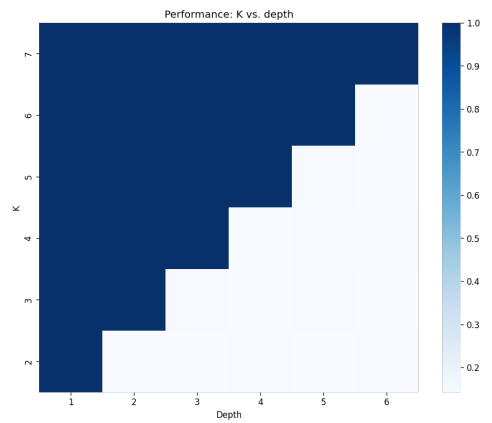
A simple example to showcase the locality





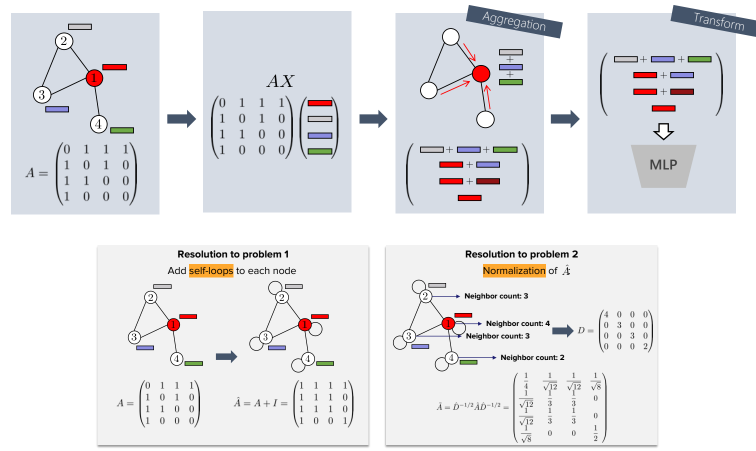
- Target: Root node (red)'s label
- The leaf node's node feature (1 in the example) is the root node's label
- The rest of the node's feature are completely irrelevant
- The model MUST be able to aggregate at least 4-hop local neighbor's information.

Alon & Yahav, On the bottleneck of graph neural networks and its practical implications, ICLR 2021 For more reading on this simple experiment, go to: https://jordan7186.github.io/blog/2022/ChebConv/



- Use a single ChebConv layer
- The model's locality (K) should at least match the minimum depth required by the NEIGHBORSMATCH problem.
- Since the problem is very easy to solve (as long as the information is properly gathered), the performance is 100% or near 1/(# of classes)%.

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks



FAST APPROXIMATE CONVOLUTIONS ON GRAPHS

In this section, we provide theoretical motivation for a specific graph-based neural network model f(X, A) that we will use in the rest of this paper. We consider a multi-layer Graph Convolutional Network (GCN) with the following layer-wise propagation rule:

$$H^{(l+1)} = \sigma \left(\tilde{D}^{-\frac{1}{2}} \tilde{A} \tilde{D}^{-\frac{1}{2}} H^{(l)} W^{(l)} \right) \,. \tag{2}$$

Here, $\tilde{A} = A + I_N$ is the adjacency matrix of the undirected graph \mathcal{G} with added self-connections. I_N is the identity matrix, $\tilde{D}_{ii} = \sum_j \tilde{A}_{ij}$ and $W^{(l)}$ is a layer-specific trainable weight matrix. $\sigma(\cdot)$ denotes an activation function, such as the ReLU(\cdot) = max(0, \cdot). $H^{(l)} \in \mathbb{R}^{N \times D}$ is the matrix of activations in the l^{th} layer; $H^{(0)} = X$. In the following, we show that the form of this propagation rule can be motivated¹ via a first-order approximation of localized spectral filters on graphs (Hammond et al., 2011; Defferrard et al., 2016).

2.1 SPECTRAL GRAPH CONVOLUTIONS

We consider spectral convolutions on graphs defined as the multiplication of a signal $x \in \mathbb{R}^N$ (a scalar for every node) with a filter $g_{\theta} = \text{diag}(\theta)$ parameterized by $\theta \in \mathbb{R}^N$ in the Fourier domain, i.e.:

$$g \star x = U g_{\theta} U^{\top} x \,, \tag{3}$$

where U is the matrix of eigenvectors of the normalized graph Laplacian $L = I_N - D^{-\frac{1}{2}}AD^{-\frac{1}{2}} = U\Lambda U^{\top}$, with a diagonal matrix of its eigenvalues Λ and $U^{\top}x$ being the graph Fourier transform of x. We can understand g_{θ} as a function of the eigenvalues of L, i.e. $g_{\theta}(\Lambda)$. Evaluating Eq. 3 is computationally expensive, as multiplication with the eigenvector matrix U is $\mathcal{O}(N^2)$. Furthermore,

We are now ready to follow the <u>original author's motivation</u> for GCN. But keep in mind that all of the previous explanations are still intact. **It is still the same model.**

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks

We now understand the first part of Section 2.1.

2.1 Spectral Graph Convolutions

(1): We are assuming a global filter with N parameters.

We consider spectral convolutions on graphs defined as the multiplication of a signal $x \in \mathbb{R}^N$ (a scalar for every node) with a filter $g_{\theta} = \text{diag}(\theta)$ parameterized by $\theta \in \mathbb{R}^N$ in the Fourier domain, i.e.:

$$g_{\theta} \star x = U g_{\theta} U^{\top} x \,, \tag{3}$$

where U is the matrix of eigenvectors of the normalized graph Laplacian $L = I_N - D^{-\frac{1}{2}}AD^{-\frac{1}{2}} = U\Lambda U^{\top}$, with a diagonal matrix of its eigenvalues Λ and $U^{\top}x$ being the graph Fourier transform of x. We can understand g_{θ} as a function of the eigenvalues of L, i.e. $g_{\theta}(\Lambda)$.

(2): Use the original eigenvalues as a basis of some approximation, then we can learn the coefficients.

 $(\mathbf{2})$

1. Initial graph signal on every node

- 2. Send the signal to the frequency domain (Fourier)
- 3. Apply the (learnable) filter function (Equalizer!)

4. Send the edited signal back to the original domain (inverse Fourier)

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks

The next part is now also familiar to us...

Evaluating Eq. 3 is

computationally expensive, as multiplication with the eigenvector matrix U is $\mathcal{O}(N^2)$. Furthermore, computing the eigendecomposition of L in the first place might be prohibitively expensive for large graphs. To circumvent this problem, it was suggested in Hammond et al. (2011) that $g_{\theta}(\Lambda)$ can be well-approximated by a truncated expansion in terms of Chebyshev polynomials $T_k(x)$ up to K^{th} order:

$$g_{\theta'}(\Lambda) \approx \sum_{k=0}^{K} \theta'_k T_k(\tilde{\Lambda}),$$
 (4)

with a rescaled $\tilde{\Lambda} = \frac{2}{\lambda_{\max}} \Lambda - I_N$. λ_{\max} denotes the largest eigenvalue of L. $\theta' \in \mathbb{R}^K$ is now a vector of Chebyshev coefficients. The Chebyshev polynomials are recursively defined as $T_k(x) = 2xT_{k-1}(x) - T_{k-2}(x)$, with $T_0(x) = 1$ and $T_1(x) = x$. The reader is referred to Hammond et al. (2011) for an in-depth discussion of this approximation.

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks

The next part is now also familiar to us...

Going back to our definition of a convolution of a signal x with a filter $g_{\theta'}$, we now have:

$$g_{\theta'} \star x \approx \sum_{k=0}^{K} \theta'_k T_k(\tilde{L}) x \,, \tag{5}$$

with $\tilde{L} = \frac{2}{\lambda_{\text{max}}}L - I_N$; as can easily be verified by noticing that $(U\Lambda U^{\top})^k = U\Lambda^k U^{\top}$. Note that this expression is now K-localized since it is a Kth-order polynomial in the Laplacian, i.e. it depends only on nodes that are at maximum K steps away from the central node (Kth-order neighborhood). The complexity of evaluating Eq. 5 is $\mathcal{O}(|\mathcal{E}|)$, i.e. linear in the number of edges. Defferrard et al. (2016) use this K-localized convolution to define a convolutional neural network on graphs.

In other words, each node requires up to K-hop local neighborhood information to capture up to K-th complex patterns.

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks

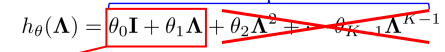
In Section 2.2, the authors start to introduce the 'deep learning' style motivations

2.2 LAYER-WISE LINEAR MODEL

Let's consider the extremely simplified case as a single layer, and let the designer choose how much layer to stack.

A neural network model based on graph convolutions can therefore be built by stacking multiple convolutional layers of the form of Eq. 5, each layer followed by a point-wise non-linearity. Now, imagine we limited the layer-wise convolution operation to K = 1 (see Eq. 5), i.e. a function that is linear w.r.t. L and therefore a linear function on the graph Laplacian spectrum.

Polynomial filter



In this linear formulation of a GCN we further approximate $\lambda_{max} \approx 2$, as we can expect that neural network parameters will adapt to this change in scale during training. Under these approximations Eq. 5 simplifies to:

$$g_{\theta'} \star x \approx \theta'_0 x + \theta'_1 \left(L - I_N \right) x = \theta'_0 x - \theta'_1 D^{-\frac{1}{2}} A D^{-\frac{1}{2}} x \,, \tag{6}$$

with two free parameters θ'_0 and θ'_1 . The filter parameters can be shared over the whole graph. Successive application of filters of this form then effectively convolve the k^{th} -order neighborhood of a node, where k is the number of successive filtering operations or convolutional layers in the neural network model.

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks

...which leads us to the final (and familiar) GCN layer.

Do we even need to differentiate θ'_0 and θ'_1 ? Let's combine them to a single parameter. In practice, it can be beneficial to constrain the number of parameters further to address overfitting and to minimize the number of operations (such as matrix multiplications) per layer. This leaves us with the following expression:

$$g_{\theta} \star x \approx \theta \left(I_N + D^{-\frac{1}{2}} A D^{-\frac{1}{2}} \right) x \,, \tag{7}$$

with a single parameter $\theta = \theta'_0 = -\theta'_1$. Note that $I_N + D^{-\frac{1}{2}}AD^{-\frac{1}{2}}$ now has eigenvalues in the range [0, 2]. Repeated application of this operator can therefore lead to numerical instabilities and exploding/vanishing gradients when used in a deep neural network model. To alleviate this problem, we introduce the following *renormalization trick*: $I_N + D^{-\frac{1}{2}}AD^{-\frac{1}{2}} \rightarrow \tilde{D}^{-\frac{1}{2}}\tilde{A}\tilde{D}^{-\frac{1}{2}}$, with $\tilde{A} = A + I_N$ and $\tilde{D}_{ii} = \sum_j \tilde{A}_{ij}$.

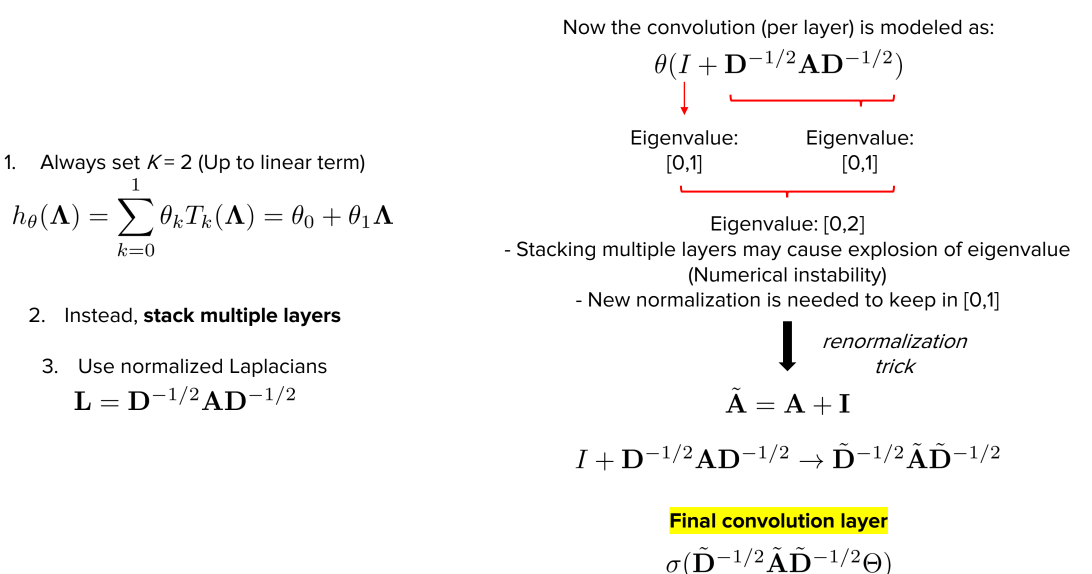
We can generalize this definition to a signal $X \in \mathbb{R}^{N \times C}$ with C input channels (i.e. a C-dimensional feature vector for every node) and F filters or feature maps as follows:

$$Z = \tilde{D}^{-\frac{1}{2}} \tilde{A} \tilde{D}^{-\frac{1}{2}} X \Theta, \qquad (8)$$

GCNs: Let's be more efficient then ChebConv

Kipf & Welling, Semi-supervised Classification with Graph Convolutional Networks

2.



Takeaways

- 1. Graph Fourier Transform: Start with the generalized concept of Fourier transform, everything else is the same.
- 2. ChebNet / ChebConv: A learnable, localized filter for graphs
- 3. GCN: Push the simplification of graph filters to the extreme, compensate by stacking multiple layers.

*If you are interested in the actual efficiency between ChebNet vs. GCN, check out https://jordan7186.github.io/blog/2022/Efficiency_Comparison/

*Highly recommended reading (for follow-up work and great summary): Wang & Zhang, How powerful are spectral graph neural networks, ICML 2022

Thank you!

Please feel free to ask any questions :) *jordan7186.github.io*