Graph Convolution Networks (Deep Learning After You Drop The Camera)

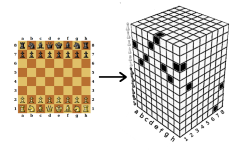
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Convolutional Neural Networks

CNNs, as you've seen are

- 1. Flexible
- 2. Computationally Efficient
- 3. Expressive
- 4. Give great results.



THE DARK SIDE OF CNNs

But what is assumed when you use a CNN?

The data should be

- 1. Regular
- 2. Dense (or non–sparse)
- 3. Have some local relational properties
- Consistent
- 5. Suitable for taking layers of filtering

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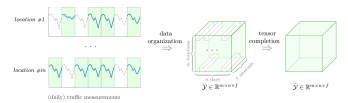
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Breaking these assumptions?

What happens if these assumptions don't hold?

You can always try to map data into a more regular structure, or downsample your data—Uber does this for demand estimation.

Imputation is another option too, where we try and fill in missing data to make a regular structure



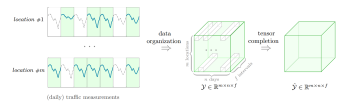
And even if the assumptions hold, it doesn't mean that there's not a better way.

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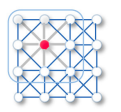
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MOTIVATING QUESTION





What happens if the data isn't nicely structured and suitable for a CNN? How do we manage real world data sets?

Real World Data

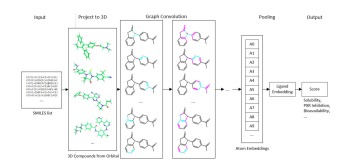
Real world data doesn't often exist on nice grids!

- 1. Traffic Graphs
- 2. Sensor network data
- 3. The relationship between your limbs as you move
- 4. Taxi demand
- 5. Social networks

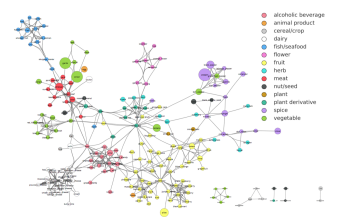
We need a general language to describe the relationships between entities.

One potential solution is to use *graphs*.

Real World Data



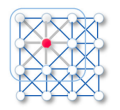
REAL WORLD DATA



It is possible to represent this as a matrix—specifically an adjacency matrix corresponding to all the links. But the matrix would inherently be incredibly sparse, which would make it inappropriate for CNNs.

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GRAPHS





- ► Graphs are a collection of Vertices (or Nodes) V and Edges E. For deep learning, we presume that the Graph Nodes have attributes $\mathbf{X} \in \mathbb{R}^{n \times d}$.
- ▶ It is also possible that the edges have attributes $\mathbf{X}^e \in \mathcal{R}^{m \times c}$. These properties can be constant or vary with time.
- ▶ With a Graph Convolution Network, we want to aggregate information from the neighbours of a node.
- ► A neighbourhood can be more than just the nodes immediate neighbours.

PREDICTIONS ON GRAPHS

The big question is: How do we take advantage of this relational structure to make better predictions?

Rather than abstracting them to make the data fit modelling frameworks (like CNNs), we instead want to explicitly model these relationships to improve predictive performance.

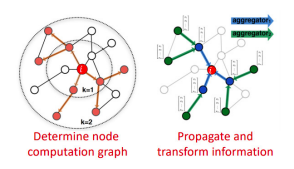
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GRAPH NEURAL NETWORKS

With a graph neural network, we want to learn how to aggregate and propagate information across the graph, in a way that helps us extra **local** (node specific) or **global** (graph specific) features.

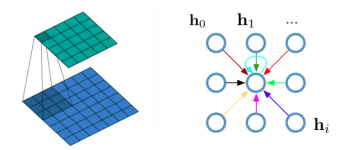


What exactly is Convolution again?

$$(f * g)(x) = \int_{\mathcal{R}^d} f(y)g(x - y)dy = \int_{\mathcal{R}^d} f(x - y)g(y)dy.$$

- ► In general, a convolution is the distortion of one function by another, so one takes the properties of the other.
- ► In a CNN, we project the data onto the convolution kernel, and extract properties about the local neighbourhood within the matrix representation.

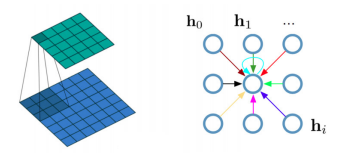
CNN Networks



 $\mathbf{h}_i \in \mathcal{R}^F$ are the hidden layer activations of each pixel.

Update the hidden layer by $\mathbf{h}_{i}^{(l+1)} = \sigma\left(\sum_{\forall i} \mathbf{W}_{i}^{(l)} h_{i}^{(l)}\right)$

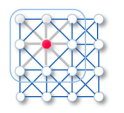
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GRAPH CONVOLUTIONAL NETWORKS



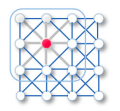


To update:

$$\mathbf{h}_0^{(l+1)} = \sigma \left(\mathbf{W}_0^{(l)} \mathbf{h}_0^{(l)} + \mathbf{W}_1^{(l)} \mathbf{h}_1^{(l)} + \mathbf{W}_2^{(l)} \mathbf{h}_2^{(l)} + \mathbf{W}_3^{(l)} \mathbf{h}_3^{(l)} + \mathbf{W}_4^{(l)} \mathbf{h}_4^{(l)} \right)$$

Note how **each node** has a **single weight** W_i , rather than a unique weight for each link.

GRAPH CONVOLUTIONAL NETWORKS



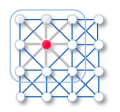


But how do we account for the fact that some nodes have fewer connections?

Weight the update, so that:

$$\mathbf{h}_{i}^{(l+1)} = \sigma \left(\mathbf{W}_{i} \mathbf{h}_{0}^{(l)} + \sum_{\forall j \in \mathcal{N}_{i}} f(i, |\mathcal{N}_{j}|) \mathbf{h}_{j}^{(l)} W_{j}^{(l)} \right)$$
One possible weighting is $f(i, |\mathcal{N}_{i}|) = \frac{1}{|\mathcal{N}_{i}|}$

GRAPH CONVOLUTIONAL NETWORKS





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GRAPH NETWORK EXTENSIONS

$$\mathbf{H}^{(l+1)} = \sigma \left(\mathbf{H}^{(l)} \mathbf{W}_0^{(l)} + \tilde{\mathbf{A}} \mathbf{H}^{(l)} \mathbf{W}_1^{(l)} \right)$$

Can be generalised as

$$H^{(l+1)} = \sigma \left(H^{(l)} W_0^{(l)} + Agg \left(\{ h_j^{(l)}, \forall j \in \mathcal{N}_i \} \right) \right)$$

Where the *Agg* function can be nearly anything. Examples include max–pooling

$$Agg = \gamma\left(\{\mathbf{Qh}\}\right)$$

where γ is a mean, max, or min function. We can also apply LSTMs to the output (or even, in some cases, on the hidden layers too)

$$Agg = LSTM(\mathbf{h})$$

EFFICIENT GRAPH UPDATES

Just summing over all the connecting nodes is neither efficient, nor tensor–like. The update procedure can instead by framed as

$$\mathbf{H}^{(l+1)} = \sigma \left(\mathbf{H}^{(l)} \mathbf{W}_0^{(l)} + \tilde{\mathbf{A}} \mathbf{H}^{(l)} \mathbf{W}_1^{(l)} \right)$$

where $\tilde{\mathbf{A}} = \mathbf{D}^{-1/2} \mathbf{A} \mathbf{D}^{1/2}$ is the Laplacian operator. $\mathbf{H}^{(l)} = [\mathbf{h}_1^{(l)}, \mathbf{h}_1^{(l)}, \dots, \mathbf{h}_N^{(l)}]$. In this \mathbf{A} is the graph adjacency

matrix, for which $\mathbf{A}_{i,j} = 1$ if there's a link from node i to j, \mathbf{D} is the diagonal degree matrix of \mathbf{A} , where

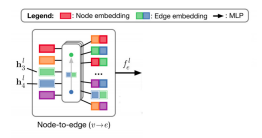
$$\mathbf{D}_{ii} = \sum_{\forall i} \mathbf{A}_{ij}$$

Instead of using $\tilde{\mathbf{A}} = \mathbf{D}^{-1/2} \mathbf{A} \mathbf{D}^{1/2}$, can use the modified Laplacian $\mathbf{L} = \mathbf{I}_N + \mathbf{D}^{-1/2} \mathbf{A} \mathbf{D}^{1/2}$, so that

$$\mathbf{H}^{(l+1)} = \sigma \left(\mathbf{L} \mathbf{H}^{(l)} \mathbf{W}^{(l)} + \mathbf{b}^{(l)} \right)$$

Graph Edge Networks

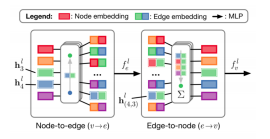
But real data of interest might exist on the nodes and the edges.



Edge hidden weights can be assigned as

$$\mathbf{h}_{(i,j)}^{(l)} = f_e^{(l)}\left(\mathbf{h}_i^{(l)}, \mathbf{h}_j^{(l)}; \mathbf{x}_{(i,j)}\right)$$

GRAPH EDGE NETWORKS



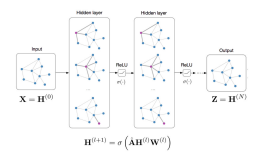
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Vertex hidden weights are then

$$\mathbf{h}_{j}^{(l+1)} = f_{v}^{(l)} \left(\sum_{\forall i \in \mathcal{N}_{j}} \mathbf{h}_{(i,j)}^{(l)}; \mathbf{x}_{i} \right)$$

THE FINAL LAYER



The final layer can be processed in a number of ways. softmax(\mathbf{z}_i) can be used for node classification, and softmax($\sum \mathbf{z}_i$) for graph classification. The importance of links can be predicted by $\sigma(\mathbf{z}_i^T\mathbf{z}_j)$. Other activation functions can be used for regression.

Summary so far

- ► Graph networks discard the structure of tensor/matrix blocks in exchange for flexible discretisations.
- ► Information can be encoded at the graph nodes and edges. These properties can even be time varying subject to local or global conditions.
- ► Care needs to be taken with how the networks weigh varying numbers of connections into a node.

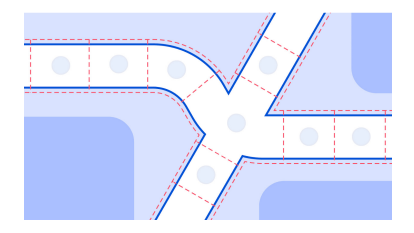
Training

- ▶ With a CNN, all information is loaded into memory. If our data is an image, then we load the entire image at once.
- ▶ In a GCN, we can do the same thing. But a Graph G = (E, V) has a lot more information to load in, and there's no implicit structure.
- ▶ But with a GCN, we don't need global information to train or run the model. So can randomly select a node, expand over its neighbours, and train over a subset of the graph. Changing this subset over training allows global behaviour to be learned.

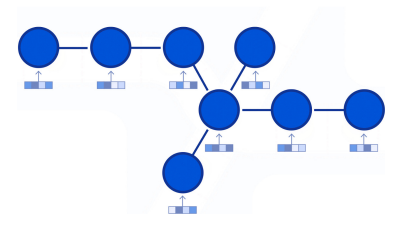
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CASE STUDY: GOOGLE MAPS

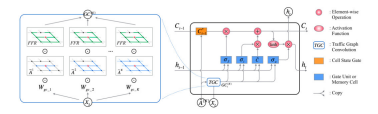


CASE STUDY: GOOGLE MAPS



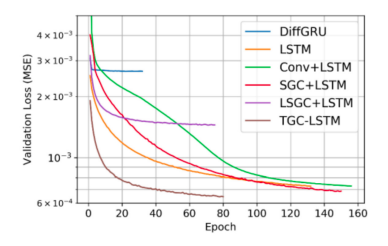
Create the graph network, and then at each node enter a sequence of time series data.

Case Study: Google Maps

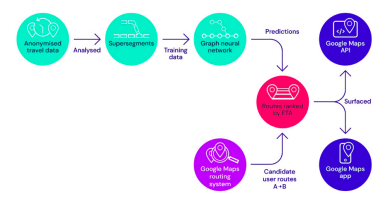


Behaviour across the whole network can be described by a Graph Convolution Network, embedded within an LSTM-like structure.

CASE STUDY: GOOGLE MAPS



CASE STUDY: GOOGLE MAPS



Case Study: Point Clouds

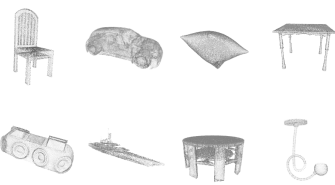


Figure: ShapeNet point clouds

These can be represented as a density matrix, but this approach may fail in the case of complex geometries, noisy data, or areas with holes. Would require significant preprocessing, as point clouds are irregular and unordered.

CASE STUDY: POINT CLOUDS

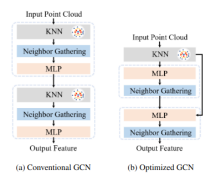


Figure: Architecture from "Towards Efficient Graph Convolutional Networks for Point Cloud Handling" Li et. al, 2021

Case Study: Point Clouds

Network	Method	Points	mIoU	Runtime [ms]	GPU mem. [GB]	FLOPs [G]	#GPU
PointCNN [21]	Baseline	2048	83.34	123.0 / 100.%	3.3 / 100.%	9.7 / 100.%	1
	Accel.	2048	83.21	111.9 / 91.0%	2.7 / 82.7%	7.6 / 78.8%	1
DGCNN [45]	Baseline	2048	84.95	116.1 / 100.%	17.2 / 100.%	158.8 / 100.%	2
	Accel.	2048	84.78	81.8 / 70.5%	4.1 / 23.8%	71.2 / 44.8%	1
[50]	Baseline	2048	84.13	365.3 / 100.%	9.7/ 100.%	202.9 / 100.%	2
	Accel.	2048	84.02	46.2 / 12.6 %	1.9 / 19.5%	52.7 / 26.0%	1

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While GNNs are less efficient than CNNs, as a relatively new and emerging architecture there's still plenty of scope for improvements.

Overview

The biggest feature of Graph Neural Networks is their flexibility. A GNN can perform almost any operation that you could see with a traditional CNN, but with extra flexibility in how you design the network.

This flexibility in turn allows ML practitioners to both approach a greater range of problems, and to tackle traditional problems with the potential for greater accuracy.

With careful design, even with the additional overhead of managing the graph, they also hold the potential to be more scalable to large data sets than other NN approaches.