Locating Datacenter Link Faults with a Directed Graph Convolutional Neural Network

Michael P. Kenning, Jingjing Deng, Michael Edwards and Xianghua Xie
Swansea University, U.K.
csvision.swan.ac.uk

Keywords: Graph Deep Learning, Fault Detection, Datacenter Network, Directed Graph, Convolutional Neural Network.

Abstract: Datacenters alongside many domains are well represented by directed graphs, and there are many datacenter problems where deeply learned graph models may prove advantageous. Yet few applications of graph-based convolutional neural networks (GCNNs) to datacenters exist. Few of the GCNNs in the literature are explicitly designed for directed graphs, partly owed to the relative dearth of GCNNs designed specifically for directed graphs. We present therefore a convolutional operation for directed graphs, which we apply to learning to locate the faulty links in datacenters. Moreover, since the detection problem would be phrased as link-wise classification, we propose constructing a directed linegraph, where the problem is instead phrased as a vertex-wise classification. We find that our model detects more link faults than the comparison models, as measured by McNemar’s test, and outperforms the comparison models in respect of the $F_1$-score, precision and recall.

1 INTRODUCTION

The convolutional neural network’s (CNN) representational power is owed to its small kernels, permitting a small set of parameters to compose low-level features over a sequence of layers into higher-order signals. The kernel’s structure however restricts CNNs are restricted to regular structures, such as images and video, precluding their direct use on a great number of natural domains exist in non-Euclidean domains.

A great effort therefore has been expended in generalizing CNNs to irregular domains. One structure that represents irregular domains well is the graph. As with a signal over a grid of pixels, a redesigned kernel can be convolved over a graph signal both in the spectral (Bruna et al., 2014; Defferrard et al., 2016; Kipf and Welling, 2017; Levie et al., 2019) and spatial domains (Niepert et al., 2016; Gilmer et al., 2017; Monti et al., 2017; Hamilton et al., 2017). Few methods however extend the CNN to directed graphs (Ma et al., 2019; Cui et al., 2020). Fewer methods exist for learning representations on graph edges, both with and without linegraphs (Klicpera et al., 2020; Jørgensen et al., 2018; Chen et al., 2019).

Datacenters readily exhibit graph structure, especially a directed graph structure. But while there are many applications of machine learning to datacenter problems (Wang et al., 2018; Srinivasan et al., 2019; Ji et al., 2018; Xiao et al., 2019), very few use graph-based convolutional neural networks (GCNNs) (Protogerou et al., 2020; Li et al., 2020). To the best of our knowledge, none represents the datacenter as a directed graph.

In this paper we describe a GCNN that can be applied to directed graphs. The model we construct, the directed graph convolutional neural network (DGCNN), is composed of convolutional layers use edge orientation to aggregate local graph signals. We train the DGCNN to locate the link faults in a datacenter, represented by a directed linegraph, a way of representing edge signals with few antecedents Chen et al. (2019). We use in silico data generated the simulator developed by Arzani et al. (2018), the advantage of in silico data being that it permits an easy identification of the ground-truth.

To summarize, our paper contributes the following:
An approach to learning on the edges of directed graphs by constructing a directed linegraph from a directed graph.

A technique to learn feature representations on directed graphs, by altering the definition of the aggregation step of a spatial graph neural network.

A new application of graph neural networks to locating link faults in datacenters.

In Section 2 we outline previous work on graphs and machine-learning applications to datacenters. In Section 3 we present our technique for learning on a directed linegraph. In Section 4 we describe our dataset, the model’s and comparison models’ implementation, the experimental conditions, and the metrics on which we assess our model against the comparison models. The results are discussed in Section 5, and we conclude the paper in Section 6.

2 RELATED WORK

2.1 Graph Convolutional Neural Networks

As acknowledged in early work (Sperduti and Starita, 1997), fully connected neural networks do not appropriately process structured information (Bronstein et al., 2017). Models that leverage graph structure gain an analytical power: a graph can encode the relational structure of a signal, and so bestow a model with helpful inductive biases (Battaglia et al., 2018). There are two ways of defining convolution on a graph: spectrally and spatially. GCNNs are hence described as either spectral or spatial. We outline several such techniques of both kinds below.

Spectral Convolution on Graphs. Bruna et al. (2014) proposed the first graph-based convolutional neural network that uses the spectral definition of convolution. A graph may be represented as a Laplacian matrix. For undirected graphs, this matrix is symmetric and positive semidefinite, so we are able to obtain a full set of eigenvectors by eigendecomposition. These eigenvectors can then be used to project graph signals into the spectral domain, where we can apply the Hadamard product. This would yield as many parameters as vertices of the graph, so Bruna et al. proposed learning spline coefficients that are interpolated to form a filter, in turn yielding smoother filters that correspond to localized convolutional filters in the spatial domain (Shuman et al., 2013).

Owing to the $O(|V|^2)$ computational cost of computing the eigenvectors, where $V$ is the vertex set, later research focused on efficient approximations of the spectral filter. Defferrard et al. (2016) proposed an approximation using Chebyshev polynomials, reducing the computational cost of filtering to $O(k|E|)$, where $k$ is the order of the Chebyshev polynomial and $E$ is the edge set. Chebyshev polynomials are ill-suited to community detection problems, however, as they cannot produce low-order, localized, narrowband filters. Levie et al. (2019) proposed Cayley polynomials instead, the computational cost of which is $O(|V|)$.

Bruna et al.’s method and the subsequent approximations are however limited to undirected graphs, owing to the requirement that the graph Laplacian matrix be symmetric. This is not possible with directed graphs, as the adjacency matrix is not symmetric. Two methods have been proposed to overcome this problem. Ma et al. (2019) used the Perron-Frobenius theorem to extract the Perron vector from the probability matrix of the directed graph, which can then be used to construct a symmetric Laplacian matrix. To learn on directed signed graphs on the other hand, Cui et al. (2020) constructed a signed Laplacian matrix, which is used to implement convolution in a similar manner as in Bruna et al.’s work. Both these techniques require the directed graph to be strongly connected, meaning that there should be a path from every vertex to every other vertex in the graph.

The spectral methods we described above have several disadvantages. Firstly they assume a fixed graph, as the filtering is defined on the graph Laplacian, which changes as the graph’s structure changes. Consequently a model based on a Laplacian matrix cannot be applied to dynamic domains. Secondly, it is unclear whether spectral techniques work on graph localization problems where several independent signals exist. Thirdly, as mentioned above, the former techniques do not work on directed graphs, while the latter two only work on strongly connected graphs. On the contrary, the spatial techniques we discuss below can accommodate these properties.

Spatial Convolution on Graphs. Spatial convolution on graphs works locally on each vertex, like convolution on an image centers on each pixel. Each technique requires us to define a vertex’s neighborhood and the function we use to filter the neighborhood’s signals. The neighborhood is difficult as there is no immediately meaningful way to describe a locality in a graph unless defined already by the domain. The filtering function is yet more difficult. The placement of pixels in an image is fixed and a neigh-
borhood has the same structure everywhere. A CNN hence binds its kernels’ parameters to positions in a neighborhood by their offset from the center. A graph’s vertices however have no natural ordering (unless bestowed by the domain); the neighborhood sizes may also vary between each vertex. Simply binding a graph model’s parameters to a neighborhood’s vertices is hence troublesome, and must either be managed with additional procedures or circumvented. Graph models differ primarily in the way they handle the signals.

Niepert et al. (2016) were among the earliest to generalize CNNs to graphs. Their technique, PATCHY-SAN, consists of three stages: selection, aggregation and normalization. In the selection stage, a labeling procedure ranks a graph’s vertices and selects the top \( w \) vertices, mimicking striding in a CNN. Then a \( k \)-large neighborhood around each vertex is aggregated, subject to a further labeling. Finally each subgraph around the vertices is normalized. PATCHY-SAN can be extended to operate on edge attributes.

Gilmer et al. (2017) proposed the message-passing neural network. Each vertex’s state is updated by a so-called message, consisting of a summation of a message function over the one-hop neighbors’ edge and vertex attributes. The message function, being a summation, is commutative, but vulnerable to large variations in a vertex’s degree. The final layer is a readout function, commutative on the graph’s vertices.

Monti et al. (2017) developed a convolution layer that maps graph neighborhoods into a pseudo-coordinate space of a mixture of learned Gaussian distributions. The posterior probabilities thereby act as weights to each vertex in the neighborhood, the results of which are passed through an activation layer. Learning vertex weightings indirectly via Gaussian distributions means this technique copes well with variations in the degree of the vertex.

Hamilton et al. (2017) avoided separately weighting the features of vertices of a neighborhood by aggregating the one-hop neighborhood and applying a commutative function and an activation layer. With a mean aggregator it is similar to Kipf and Welling’s Graph Convolutional Network (GCN), which by contrast accepts weighted graphs, too.

A few techniques exist for learning on directed graphs and on directed linegraphs of graphs. Klicpera et al. (2020) used directional messages and second-order features on the graph to regress on molecular properties. Jørgensen et al. (2018) proposed a technique to incorporate edge information in the learning for a regression on molecular properties, while Chen et al. (2019) proposed a method to learn edge representations on a directed linegraph to inform a community detection problem on the vertices of the graph.

### 2.2 Locating Faults in Datacenters

A failure in a datacenter can exist anywhere on the multitude of machines and links that comprise it. A network administrator must collate and interrogate many sources of information to determine the location of problems (Gill et al., 2011). Manually this is time-consuming and difficult, and potentially very costly problem when responses need to be quick to reduce disruption to clients. Automating the interrogation of logs with an algorithm is quicker. For example, Zhang et al. (2005) postulated the loads on individual links as a linear transformation of unknown traffic elements that is inverted to discover the anomalies.

There are also monitoring systems available that help in diagnosing datacenter faults expeditiously. Pingmesh (Guo et al., 2015) monitors faults on distributed servers by running an agent on its every server to measure the latency of the network. 007 (Arzani et al., 2018) by contrast has at every host an agent that traces the path of detected TCP retransmission errors, on which several algorithms can be run to rank the links by likelihood of fault.

Machine-learning algorithms are particularly suitable to these problems (Wang et al., 2018). Srinivasan et al. (2019) for instance located link disconnections in an Internet-of-Things (IoT) network of up to 100 nodes using one such model, attaining a high detection rate in silico. Ji et al. (2018) used a CNN to scan log files to predict future network faults. Xiao et al. (2019) applied a CNN to detect intrusive behavior among network traffic.

But despite the power that graphs offer in explicitly representing the structure of datacenters, they are scarcely used in the literature. Protogerou et al. (2020) applied a graph network to detect denial-of-service attacks in an IoT network; whereas Li et al. (2020) applied a similar network to optimize a datacenter’s traffic-flow. The novelty of our work is that we use both a GCNN and represent the datacenter as a directed graph to locate link faults in a datacenter. To the best of our knowledge, there are no such examples of work in the literature.

### 3 METHOD

**Problem Description.** Failures in a datacenter can cost network operators and end-users time and money. There are myriad causes of failures in a datacenter.
The aim of the engineer is to identify and fix the most unreliable machines and links in the datacenter. Detecting these faults is non-trivial. Network operators must assess a plethora of machines and many more links, drawing on multiple indicators from across the network to track the health of the network. The task of the operator is to prioritize the most severe incidents (Gill et al., 2011).

One indication of a fault is packets dropping at an abnormal rate, which is picked up by host machines: When packet-drops occur in the middle of a TCP transmission, the destination machine sends a TCP retransmission error to the originating machine to request same packet again. Accordingly Arzani et al. (2018) developed the 007 system to use a datacenter’s hosts to determine the paths along which a retransmission error occurred, and assign a score of blame to each link. The system then uses the hosts’ aggregated blame scores to find the most probable locations of link faults.

But it is not so simple as finding k links assigned the greatest blame. Faults displace traffic and cause collateral faults elsewhere; genuinely faulty links are lost in a mire of overburdened links. For this reason, it is more helpful to understand the context of a link’s blame score than to look at scores in isolation. Hence understanding the structure of the datacenter is essential in locating faults.

Our objective is to design a graph-based model that can predict the faulty links in a datacenter. Graphs represent datacenters well. By incorporating the graph structure into a model, we are able to produce a model that accounts for contextual information.

**Graph-theoretical Definitions.** A graph \( G = (V, E) \) is defined by the sets of vertices or nodes \( V = V(G) \) and edges \( E = E(G) \). If two vertices \( x, y \in V \) are adjacent, they are incident to an edge \( \{x, y\} = xy = e \in E \). Every edge in a graph is therefore incident to two endvertices. For undirected graphs, \( E \) is a set of unordered pairs, meaning \( xy = yx \in E \); but for directed graphs \( E \) is a set of ordered pairs, meaning \( xy \neq yx \). Directed edges \( (x, y) \) are therefore incident to a startvertex \( x \) and endvertex \( y \). If \( xy, yx \in E \) is \( xy \) has an inverse edge \( yx \). A directed graph where every edge has an inverse is called symmetric. A directed graph where every vertex is reachable from every other vertex is strongly connected (see Fig. 1 for an example of a weakly connected graph).

The order of the graph is \( |G| = |V| = n \), the number of vertices, while the size of the graph is \( |E| \), the number of edges. In an undirected graph, a vertex’s degree \( d(x) \) is the number of adjacent vertices to \( x \). The minimum and maximum degree of a graph \( G \) are denoted respectively \( \delta(G) \) and \( \Delta(G) \). In a directed graph, the degree of a vertex \( x \) is the sum of the in-degrees \( d_i(x) = \{ (y, x) | (y, x) \in E \} \) and out-degrees \( d_o(x) = \{ (x, y) | (x, y) \in E \} \).

The neighborhood of a target vertex \( \Gamma(x) \) consists of its adjacent vertices and incident edges, the one-hop neighbors. The concept of a neighborhood can be expanded beyond the first hop to include those \( k \) hops away, which we denote \( \Gamma_k(x) \). The neighborhood of a vertex in a directed graph can be factored into neighborhoods incident to its in- and out-edges: \( \Gamma(x) = \Gamma^+(x) \cup \Gamma^-(x) \). The neighborhood can thus be conceived as a subgraph, with the order and size of its defined equivalently.

A graph can be represented in matrix form in a number of ways, requiring an indexing of the vertices. An **adjacency matrix** \( A \) is a binary matrix where each non-zero entry marks an adjacency between two vertices: \( \forall \{i, j\} \in V \), \( A_{ij} = 1 \). A degree matrix \( D \) is a diagonal matrix where the diagonal records the degree of each vertex, \( D = \text{diag}(D) \). A directed graph’s adjacency matrix is asymmetric. It has two degree matrices corresponding to the in- and out-degrees, defined as \( D^+ = \text{diag}(A^+1) \) and \( D^- = \text{diag}(A1) \) respectively.

An undirected linegraph \( L(G) \) is defined on an underlying undirected graph \( G \). It is an ordered pair \( \langle V_L, E_L \rangle \), where \( V_L = E(G) \). Consequently there is a bijective and hence invertible mapping from the edge set of \( G \) to the vertex set of \( L(G) \). Two vertices in the linegraph \( \alpha, \beta \in L(G) \) are adjacent if their edges in the underlying graph \( G \) have an endvertex in common. A directed linegraph (Aigner, 1967) is constructed in a more constrained manner. Two vertices in a directed linegraph are adjacent iff the two edges in the underlying directed graph the endvertex of one edge is identical to the startvertex of the other. (See Fig. 1 for a visual illustration of the construction of a directed and undirected linegraph.) The adjacency and degree matrices of the undirected and directed linegraphs are formed similarly to those of their underlying graphs; we denote them \( A_L \) and \( D_L \).

A \( m \)-dimensional signal on a graph or linegraph is the codomain of a mapping from its vertices to the real vector space \( f : V \to \mathbb{R}^m \). The signal may also be a mapping from a graph’s edges \( f : E \to \mathbb{R}^m \). The graph thus describes the structure of the signal. Many observed signals may share the same structure. Our dataset consists of a set of observations of a structured signal on a linegraph. Each observation is therefore a separate mapping. We often refer to a vertex’s signal simply as the vertex.
A Datacenter as a Graph. A datacenter can be easily represented as a graph if we consider each machine to be a vertex and draw edges between machines if they are connected. In our case we use an unweighted graph. As mentioned above, each connection between a pair of machines consists of an uplink and downlink, two opposing flows of traffic, which can be represented as directed edges in a symmetric directed graph.

On top of this unweighted graph, we can build a directed linegraph, where each vertex represents a link in the datacenter. The linegraph of the datacenter’s graph therefore represents the second-order structure of the datacenter, the adjacency of the links. A path of vertices in the graph represents the passage of a packet through the datacenter. We do not join inverse edges in the directed linegraph for two reasons. On the practical side, the edge’s inverse belongs to both the in- and out-neighbors: either count it doubly or exclude it. From the point of view of the domain, an edge’s inverse is irrelevant, as a packet that is routed up or down the datacenter will not traverse a link it has already traversed. In effect we are incorporating the non-backtracking operator used in the linegraph neural network used in Chen et al.’s work (2019), first proposed by Krzakala et al. (2013) designed for random walks.

As detailed in the problem description above, the features in our task sit on the links of the datacenter, on the edges of its directed graph, and therefore on the vertices of its directed linegraph. This makes it easier to apply the various techniques outlined in Section 2, as they are focused on learning representations on the vertices of the graphs rather than the edges.

A Spatial Convolution for Directed Graphs. Hamilton et al. (2017) defined graph convolution as the aggregation of signals in a vertex’s neighborhood together with the target vertex’s signal. Applying the mean aggregator, we get a local function \( g \) on graph vertex \( x \) and signal mapping \( f \),

\[
g(x, f) = \frac{1}{d(x) + 1} \sum_{y \in \Gamma^+(x)} f(y),
\]

which is identical to the convolutional layers used in the GCN (Kipf and Welling, 2017) when the graph is unweighted, as in our case. This formulation does not however account for directed graphs. Moreover, since the target vertex’s signal is summed with the neighbors’ signals, the target signal is lost.

Therefore we define our directed graph convolu-

\[
g(x, f) = \sum_{y \in \Gamma^+(x)} f(y)
\]

\[
+ f(x) \theta_1 + b,
\]

(2)

which is parameterized by the bias term \( b \) and the weights \( \theta_1 \in \mathbb{R}^c \), where \( c \) is the number of input channels.

We have partitioned the neighborhood signals into two groups according to their orientation, as we believe that the relation of the neighbors’ signals to the target’s signal changes depending on whether a neighbor is an in- or out-neighbor. The target vertex is considered separately because it does not belong to either partition, meaning it is also not lost amid the neighbors’ signals, free to inform our model independently. This is potentially very helpful in a situation where the effect of a vertex (a potentially faulty link) on its neighbors (the adjacent links) is being modeled.

4 EXPERIMENT

In this section, we compare our model, the DGCNN, empirically against several models. The task is to locate link faults in a datacenter. Our experiments were conducted in Python 3.6.6 using Tensorflow 2.2 on a computer with an NVIDIA GeForce 1080Ti graphics card, a quad-core Intel Core i7-6700k CPU at 4.00 GHz and a 32 gigabytes of RAM.

There are two primary outcomes for our experiments: (1) to establish whether separating the edge signals by their direction effects a better performance; and (2) to compare our model’s efficacy to a state-of-the-art spatial GCNN.

4.1 The Dataset

We use the flow-level datacenter simulator implemented by Arzani et al. (2018) to generate our data,
the simulator on which they developed the 007 system. The datacenters it simulates conform to the Clos topology. There are four types of machine: a host machine (Host); a top-of-rack switch (ToR) to which each Host connects; a further T1 switch to which the ToRs connect; and a top-level T2 switch. A Pod consists of a set of Hosts, ToRs and T2s. We decided on 20 T2 switches and two Pods containing 10 T1 switches, 10 Top-of-Rack (ToR) switches and 240 Host devices. Each T1 switch connects to two T2 switches, with each T2 switch connecting to one T1 switch from each Pod. The T1s and ToRs within each Pod are fully connected to one another. Each ToR switch connects to 24 Host machines. In total there are 540 machines or devices and 1440 links, as each connection consists of an up- and downlink. We use the blame scores computed by the 007 system in these simulations to locate faults. Note that faults do not occur on links joined to Hosts, for detecting such faults is simple, because they can be detected by the Host itself directly.

We ran 2,880 30-second simulations of a datacenter randomly selecting 2 to 10 links to be faulty (see Fig. 2 for an illustration of the topology and an example of a simulation). A faulty link is defined as a link with a packet-drop probability \(0.01 \leq p(F) \leq 0.1\), while healthy links had a 0% probability of dropping packets. The dataset is stored on computer as a 2880-by-1440-by-1 tensor, the one input feature being the blame scores produced by Arzani et al.’s 007 system. The labels are stored in a binary 2880-by-1440 tensor, with faults being the positive class.

Consequent on a low number of failures is a high class imbalance. The number of faults in a given experiment is uniformly random \(F \sim U(2, 10)\), meaning \(\mathbb{E}(F) = 2 + \frac{10-2}{1} = 6.5\). The expected imbalance ratio is therefore \(\rho = \mathbb{E}(F) / (1440 - \mathbb{E}(F)) = 6.5/1374 = 4.5343 \times 10^{-3}\). Unless this imbalance is addressed, it risks compromising the training, as the model could reduce the loss simply by labeling all links healthy.

### 4.2 Implementation and Comparisons

Our DGCNN consists of three of the directed convolutions described in Section 3 and a final 10-unit multi-layer perceptron (MLP), the output of which is

\[
\begin{align*}
    z &= \sigma(W(c_1 \circ c_2 \circ c_3)(V, f) + b), \\
    e_i(V, f) &= \tau(\beta g(V, f)),
\end{align*}
\]

where \(c_i(V, f) \) is the \(i\)th convolution block, consisting of a directed convolution \(g(V, f) \) supplied with a graph structure \(V\) and a signal mapping \(f\); a batch-normalization layer \(\beta\) and a rectified linear unit \(\tau\).

Three of layers are composed \((c_1 \circ c_2 \circ c_3)\) and the output is passed through a sigmoid-activated single-unit fully connected layer with weights \(W \in \mathbb{R}^{1 \times d}\) and bias \(b \in \mathbb{R}\), where \(d\) is the number of output maps from the final convolution block.

Each convolution layer \(g\) yields 10 output maps. The batch-normalization layer uses the default parameters for Tensorflow (momentum = 0.99, \(\varepsilon = 0.001, \beta = 0, \gamma = 1\); the moving mean and variance were zero- and one-initialized respectively). There are 731 trainable parameters in total. We decided on 3 convolutional layers and 10 output maps after experimentation, choosing the best trade-off between \(F\)-score and time to train (Fig. 3).

The output \(z \in \mathbb{R}^{|E|}\) is the model’s certainty of a fault at each edge, since the loss is the cross-entropy...
of the output $z$ and the binary vector of edge labels, where an entry is positive if the corresponding edge is faulty.

The architecture of the DGCNN differs from the comparison models only in its convolutional layers $g$: we endeavored to keep everything else equal to ensure that we are assessing a single factor, thereby eliminating confounding factors. Our comparison models are the following:

1. **Two Undirected Forms of the DGCNN**, (UGCNN and UGCNN Large) to determine the contribution made to the DGCNN’s performance by including edge direction. The smaller model (“undirected graph convolutional neural network (UGCNN)”) uses 10 output maps per layer, totaling 521 parameters. The larger one (“UGCNN large”) uses 12 output maps per layer, totaling 721 parameters, in order to test whether the DGCNN’s greater capacity rather than the inclusion of edge direction lends it an advantage.

$$g(x) = \theta_0 \frac{1}{d(x)} \sum_{y \in \Gamma(x)} f(y) + \theta_2 f(x) + b \quad (6)$$

2. **A GraphSAGE Network** (GraphSAGE), to compare the DGCNN to a similar spatial GCNN. It is a point of comparison with the wider literature as a well-cited model. We use the implementation in version 0.6.0 of the Spektral Tensorflow library (Grattarola and Alippi, 2020) and the mean aggregator proposed in the original paper (Hamilton et al., 2017).

3. **A Dense Model** (Dense), where the convolutional layers are replaced with 10-unit fully connected layers, to establish the utility of the structural information provided by the lineigraph.

4.3 **Experimental Conditions**

The dataset was split 3:1:1 by simulation between training (1728 simulations), validation (576) and test set (576) in five folds to study the models’ stability. In experiments, we chose batches 64 samples large because it permitted a smooth cosine decay of the learning rate while keeping the time to train minimal. By changing the magnitude of the learning rate we found that a decay from $\eta = 1 \times 10^{-2}$ to $\eta = 1 \times 10^{-7}$ resulted in the fastest optimization, which occurred by 50 epochs for all models in the experiments. We reconfirmed these observations by studying the convergences of the $F_1$-score on the training and validation sets over the training period.

The weights of the neural networks’ layers were initialized using the Glorot uniform initializer, because they work well with rectified linear units, as demonstrated in the original paper (Glorot and Bengio, 2010). The biases were initialized at zero, with the exception of the final fully connected layer, initialized to a balanced odds-ratio of the positive and negative samples (7).

$$s_+ - \frac{|L(G)|}{2} \cdot E(F) \quad s_- = \frac{|L(G)|}{2} \cdot E(\neg F) \quad (7)$$

The models are compared on statistics commonly used in binary problems. Each statistic was computed on the references from the test set and averaged over the five folds. The primary metric on which we compare the models is the $F_1$-score (8). We are however more interested in precision than recall, as we consider a set of positives as unadulterated by healthy links to be the ideal situation.

$$F_1 = 2 \frac{\text{precision} \cdot \text{recall}}{\text{precision} + \text{recall}} \quad (8)$$

We also compare our model’s predictions pairwise with the other models’ using McNemar’s test (9),

$$\chi^2 = \frac{(|N_{sf} - N_{fs}| - 1)^2}{N_{sf} + N_{fs}} \quad (9)$$

where $N_{sf}$ is the count of correct predictions by model 1 where model 2 was incorrect, and $N_{fs}$ is the opposite. When $\chi^2 \approx 0$, there is not much difference between the two models; a large value indicates a difference in performance. Model 1 in our experiments is the DGCNN, and model 2 is a comparison model. We also measure the number of parameters each model uses, and the average training and inference times on the test set.

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Table 1: The results of the experiments outlined in Sections 4.2 and 4.3. The count of parameters (Params.) is the number of trainable parameters in the model. The $F_1$-score, precision and recall scores are rounded to four decimal places. The results of McNemar’s test ($\chi^2$) are rounded to two decimal places. The time to train (“TTT”; in minutes, rounded to nearest second) and inference time (“Inf.”; rounded to nearest millisecond) are the average times over the five folds.

<table>
<thead>
<tr>
<th>Model</th>
<th>Params.</th>
<th>$F_1$</th>
<th>Precision</th>
<th>Recall</th>
<th>TTT</th>
<th>Inf.</th>
<th>$N_{i,j}$</th>
<th>$N_{f,j}$</th>
<th>$\chi^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DGCNN (ours)</td>
<td>731</td>
<td>0.8048 ± 0.0041</td>
<td>0.6736 ± 0.0056</td>
<td>0.9997 ± 0.0004</td>
<td>2.85</td>
<td>553</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>UGCNN large</td>
<td>721</td>
<td>0.7657 ± 0.0161</td>
<td>0.6211 ± 0.0213</td>
<td>0.9988 ± 0.0009</td>
<td>2.30</td>
<td>447</td>
<td>446.6</td>
<td>4.2</td>
<td>432.6</td>
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<tr>
<td>UGCNN small</td>
<td>521</td>
<td>0.7312 ± 0.0688</td>
<td>0.5803 ± 0.0822</td>
<td>0.9987 ± 0.0011</td>
<td>2.27</td>
<td>397</td>
<td>946.0</td>
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<td>927.13</td>
</tr>
<tr>
<td>GraphSAGE</td>
<td>311</td>
<td>0.3031 ± 0.0156</td>
<td>0.1790 ± 0.0109</td>
<td>0.9924 ± 0.0030</td>
<td>8.88</td>
<td>4759</td>
<td>14173.6</td>
<td>0.4</td>
<td>14170.40</td>
</tr>
<tr>
<td>Dense</td>
<td>311</td>
<td>0.3085 ± 0.0546</td>
<td>0.1839 ± 0.0389</td>
<td>0.9893 ± 0.0072</td>
<td>1.52</td>
<td>219</td>
<td>14245.0</td>
<td>0.4</td>
<td>14242.40</td>
</tr>
</tbody>
</table>

5 RESULTS AND DISCUSSION

As we can see from the results listed in Table 1, the DGCNN outperformed the comparison models on all measures. McNemar’s test shows that the DGCNN frequently classified links correctly where the other models failed. Our model was also the most stable, as measured by standard deviation between the folds—although this could be the consequence of choosing the architecture, the hyperparameters and experimental settings on the basis of the performance of the DGCNN alone.

The number of parameters in the DGCNN slightly increased the training time over its undirected forms and nearly doubled it compared to the dense network. Understandably, inference time on the DGCNN was middling, having had the greatest number of parameters; for the opposite reason, the dense network is the fastest. GraphSAGE was exceptionally slow in training and inference, although this might be owed to this particular implementation.

The performance of the DGCNN does not appear to be owed solely to the greater number of parameters: the evidence is strong that the inclusion of graph direction has helped the DGCNN. The large UGCNN’s greater capacity improved its performance, but still a distance off the DGCNN’s performance. As repeated throughout this paper and elsewhere (Bronstein et al., 2017), this experiment affords positive evidence that in knowing the structure we can learn something important about the signal. We suspect that the mean aggregator’s unification of a target vertex’s features with its neighbors’ is deleterious to performance, as it prevents the model weighting one against the other, permitting the model to compare a vertex to its neighborhood as our model can.

It is not clear how much performance would be affected in this particular case if the healthy links were initialized with small yet insignificant packet-drop probabilities, as in the original work by Arzani et al. It would be interesting to see the effect of the noise from healthy links on the models’ performance. For lack of space and time, we also did not conduct further analyses into the faults that the model missed. Such analysis could reveal any trends in the types of the faults not detected.

Having focused on spatial convolutional techniques on graphs in our research, we did not evaluate the two spectral methods outlined in Section 2 (Ma et al., 2019; Cui et al., 2020). Our suspicion is that, being spectral methods, they would not be suitable for locating multiple independent but related signals on the graph, as they inherently learn localized but global signals. We leave this analysis to future work.

6 CONCLUSION

In this paper we presented a spatial convolutional layer for directed graphs, and moreover a technique to learn a localization task on a directed linegraph. We applied these techniques to locating link faults in a datacenter, finding that the inclusion of direction found in the graph structure significantly improved the model. Future work should repeat these experiments, to establish whether similar performance gains can be yielded in other, unrelated domains, and compare spatial and spectral approaches on other vertex-focused tasks. This work has focused on simulated datacenters, because the ground-truths are easily acquired. In further and more specific future work, however, we could also apply our method to data from larger, real datacenters.

REFERENCES


Relational inductive biases, deep learning, and graph networks. ArXiv.


