Intelligent Performance Inference: a Graph Neural Network Approach to Modelling Maximum Achievable Throughput in Optical Networks

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One of the key performance metrics for optical networks is the maximum achievable throughput for a given network. Determining it, however, is an NP-hard optimisation problem, often solved via computationally expensive integer linear programming (ILP) formulations, infeasible to implement as objectives, even on very small node scales of a few tens of nodes. Alternatively heuristics are used, although these too require considerable computation time for large numbers of networks. There is, thus, a need for ultra-fast and accurate performance evaluation of optical networks. For the first time, we propose the use of a geometric deep learning model, message passing neural networks (MPNN), to learn the relationship between node and edge features, the structure and the maximum achievable throughput of networks. We demonstrate that MPNNs can accurately predict the maximum achievable throughput while reducing the computational time by up to 5-orders of magnitude compared to the ILP for small networks (10-15 nodes) and compared to the heuristic for large networks (25-100 nodes) - proving their suitability for the design and optimisation of optical networks on different time- and distance- scales.

I. INTRODUCTION

Multiwavelength optical networks underpin the global data communication network infrastructure and use the wavelength domain both for routing and to increase point-to-point data transmission. Increasingly, optical networks are being used for intra- and inter-data centre communications and in high performance computing.¹ To enhance the throughput of these networks, data is using many wavelength channels on a single fibre, using wavelength division multiplexing (WDM). This has greatly improved the capacity of optical networks, but requires the solution of the routing and wavelength assignment (RWA), shown to be NP-hard and, therefore, computationally difficult to solve optimally.²³

The overarching goal of physical network design is to maximise the performance, measured by throughput, latency and resilience, whilst minimising the cost and/or resource use, making them intelligent and adaptive.⁵⁶ This is an evolution from the previous goals of minimising the number of wavelengths needed to optically route data within the network, which quantified the relationship between wavelength requirements and the physical topology.³⁹ However, due to growing number of wavelengths in fibres and the growing understanding of the linear and nonlinear physical layer impairments, have highlighted the importance of including the optical fibre physical properties in network analysis and design, since these play a significant role in determining both routing and throughput, and must be taken into account.⁶

The maximum achievable throughput is defined here as the throughput when the resources (wavelengths) are fully exploited, at zero blocking for a given demand distribution (measured in bits per second). To quantify the maximum achievable throughput in a network, requires an optimal solution to the RWA problem. Integer linear programming (ILP) formulations have been shown to solve the RWA problem optimally, however are infeasible for networks larger than ~ 30 nodes. Problem-agnostic optimisation frameworks, that aim at efficiently exploring the solution space, i.e. meta-heuristics, have been shown to give good solutions in this area, however have no guarantee of achieving a global optimum, whilst often still taking a long time to solve.⁴¹⁶ Heuristics are handcrafted algorithms for specific purposes, based on automated rules-of-thumb, which are highly-scalable, but have shown limited success in terms of reaching optimum values, compared to ILP solutions.¹⁴ In addition, graph cuts have been analysed to estimate the maximum achievable throughput, however also succumb to computational complexity for larger graphs and have been shown to be non-exact for non-uniform traffic distribution.⁷¹⁰ The efficient and accurate measurement of the maximum achievable throughput of optical networks remains a considerable challenge.

To make this task more computationally efficient, machine learning has been introduced to learn the relationship between the topology and performance parameters, based on previously labelled datasets.

More specifically, supervised deep learning has been used, where large labelled datasets are used to train learnable functions. There are two broad approaches to deep learning: (i) ar-

FIG. 1: An example of message passing on a 6 node topology
tificial neural networks (ANN) based on euclidian data, or (ii) geometric deep learning using graph neural networks (GNN), based on graph structured data to perform classification or regression tasks. If[19] an ANN was used to estimate the blocking probability of a network and if[20] it was used to estimate the fast optical packet loss rate for bufferless optical packet-switched networks. The problem however with ANNs, is that they operate on grid-style data, i.e. vector/matrix inputs. Therefore, they are not able to use relational data, i.e. the relation of one node to other nodes, nor fully represent the structural features of the graphs, such as node degree variance or algebraic connectivity. This makes it difficult to learn a general relationship between these features and the performance parameters as the structure and physical properties of graphs have a significant impact on overall network performance. More recently there have been many works that look at applying reinforcement learning to solve the RWA problem. These works are promising works in applying novel combinatorial optimisation techniques to the RWA problem, however still do not achieve performance comparable to ILP solutions.

Geometric deep learning applies deep learning to graph structured data incorporating the graph structure within the learning process. By including node features, edge features, and learnable functions that operate in a graph-structured manner, geometric deep learning learns relationships on graphs better than traditional deep learning. Graph neural networks (GNN) are a collection of supervised learning frameworks for geometric deep learning, able to aggregate information in graphs. Message passing neural networks (MPNN) are a specific formulation of GNN, shown to perform well for regression tasks, one of the fundamental goals of our work. The benefit of this type of model, is that it has been shown to generalise well to graphs of different sizes and structures, as well as being computationally efficient. The downside, however, is that it requires large datasets to learn and predict graph properties. More often than not these are far and few between, making it difficult to apply these frameworks. Previously, this approach has been successfully applied in quantum chemistry and non-optical communication networks. However, in optical communications, estimating the maximum achievable throughput of a network is an infamously NP-hard problem, so that this regression problem remains unsolved.

In this paper we apply a MPNN architecture to model the performance of optical networks, by learning the relationship between the topology and the maximum achievable throughput. We generated three large datasets (~30000 graphs per dataset), with the number of nodes in the range 10-100, expanding previous work. The datasets were used to train an accurate MPNN model to learn the relationship between the topology and the performance of optical networks. We demonstrate the significant computational time reduction with this model to estimate graph performance. This in turn reduces the complexity of topology design problems for optical networks, by replacing computationally complex objectives with surrogate models, such as the one demonstrated here. Thus, enabling future topology optimisation to maximise the achievable throughput, making topology design more intelligent and computationally efficient.

The rest of the paper is structured as follows. Section II describes the MPNN model and its operation. Section III details the methodology used to generate the three graph datasets. Section IV explains the training procedure and the different parameter settings for the MPNN model. Finally, using the generated datasets, Section V analyses the model’s capability to predict maximum achievable throughput, in terms of accuracy, computation time and generalisation capability.

II. MESSAGE PASSING NEURAL NETWORKS (MPNN)

Optical networks are a set of nodes (for example major cities or data centre or servers within data centres) interconnected, with arbitrary connectivity, by optical fibres. Optical fibres carry multiple discrete wavelength channels, each carrying individually modulated data. To transfer data between source and destination nodes, for each source – destination demand, a lightpath is setup, consisting of a route (a series of edges) and a wavelength designated for transmission. The process of selection of wavelength (colour) and route is determined via the RWA algorithm. Optical networks are usually modelled as graphs and so ideally lend themselves as inputs to graph neural networks. Recently there have been works that apply graph neural networks in the context of reinforcement learning (RL) to the RWA problem in optical networks. These works however are only compared to heuristics, where sometimes they perform better and sometimes worse. We apply MPNNs to learn from the solutions directly from integer linear programming formulations and heuristics to consistently predict performance properties of optical networks such as the maximum achievable throughput.

We start with a digraph denoted as \(G(N,E)\), where \(N\) and \(E\) are its set of nodes and edges respectively. All nodes and edges have a pre-determined set of node and edge features, \(x_n\) and \(e_u\), respectively, where \(n \in N\) and \((n,u) \in E\). In this work, the degree \(\delta_n\) and the total traffic originating from a node is used to describe node features, and the worst case noise-to-signal (NSR) of a fully populated link is used to describe the corresponding edge feature. These node and edge features, are vectors with information related to either nodes: degree and traffic, or for the edges: distance and signal-to-noise (SNR) ratio. MPNNs use abstract vectors, referred to as a node or edge hidden state. In this work, for simplicity, we confine the hidden states to nodes, represented as \(h_n^t\), where \(t\) represents the message passing iteration. These hidden states, are vectors that hold embeddings for nodes, i.e. for a specific node, they capture structural information from the rest of the graph. We define the set of node features/edge features as \(X^N/X^E\) respectively, and the set of hidden node states as \(H^N\). GNNs can be used for either regression or classification tasks, and in this work we focus on the regression of graph properties.

The MPNN framework centres around three functions: message function \(M_t(h_n^t, h_n^t, e_{nu})\), update function \(U_t(h_n^t, m_n^{t+1})\) and readout function \(R(H^N, X^N)\), where \(n\) is a node in the node set \(N\), \(t\) is the message passing iteration out of \(T\) iterations and \(u\) is a node in the neighbourhood of \(n\) \((u \in \mathcal{N}(n))\). \(T\) generally is chosen to be in the order of ei-
ther the average shortest path length or diameter of the graph. There have been several different formulations of these functions in the past\cite{29,30}, however, they all follow the same general algorithm.

**Algorithm 1:** Message Passing Neural Network

*Input:* $G, X_N, X_E$

*Output:* $y$

\begin{verbatim}
\textbf{begin}
  \textbf{for} $n \text{ to } N$ \textbf{do}
  \hspace{1em} $h_n^1 \leftarrow [x_n, 0, 0, \ldots, 0]$;
  \textbf{end}
  \textbf{for} $t = 1 \text{ to } T$ \textbf{do}
  \hspace{1em} $m_n^{t+1} \leftarrow \sum_{\omega \in \mathcal{N}(n)} M_t(h_{\omega}^t, h_n^t, e_{\nu});$
  \hspace{1em} $h_n^{t+1} \leftarrow U_t(h_n^t, m_n^{t+1});$
  \textbf{end}
  $y \leftarrow R(H_N, X_N);$
\textbf{end}
\end{verbatim}

The MPNN - outlined in algorithm\cite{31} - is made up of three stages: (i) message passing (line 7) (ii) update (line 8) (iii) readout (line 11). In (i) each node in the graph, requests information (messages) from its neighbourhood ($\mathcal{N}(n)$). This information (messages) is given by feeding in node and edge information into the message function ($M_t(h_{\omega}^t, h_n^t, e_{\nu})$). To form the nodal message ($m_n^{t+1}$) of node $n$, we have to aggregate the information (messages) given by the neighbourhood of $n$. This can be done via different operations, i.e. averaging, sampling or summing. We obtain the message ($m_n^{t+1}$) of node $n$, by summing the messages from the neighbourhood of $n$. This is then fed through an update function (line 8) defined as $U_t(h_n^t, m_n^{t+1})$, which updates the state ($h_n^{t+1}$) of each node. These two steps are illustrated in the inner block of the diagram shown in figure\cite{2}, where one can see that the process is repeated for each node $n \in N$. This procedure iteratively distributes the information of the graph to every node by collecting local messages and using these to update the new hidden vectors. Figure\cite{1} demonstrates this process of message passing for the computation of the node state of node 1, on an example 6-node topology. Here the process is shown for two message passing iterations, $t=1, 2$. Working backwards from $t=2$ and node 1, we can see that we use the neighbours of node 1, which feed their messages into the aggregation that then are updated. Before this, their neighbours do the same for their states. This demonstrates how the information about the graph is distributed across the graph during message passing.

After $T$ message passing rounds and update layers, shown by the outer blocks in figure\cite{2} the hidden states are aggregated and used to create a graph level prediction $y$, seen in line 11 of algorithm\cite{1}. This process of aggregation and graph-level readout is summarised by the readout function $R(H_N, X_N)$, where $H_N$ denotes the set of hidden states and $X_N$ the set of node features. The readout function outputs a scalar value used for prediction. This architecture has been shown to provide good learning capability for general graph structured data\cite{22}, where one can incorporate individual node and edge features. Another advantage of such a model is that the architecture is size agnostic, meaning that the model can generalise to graphs of different sizes.

### A. Message Function

The message function is used to extract information from both hidden states of neighbouring nodes and adjacent edge features, thus producing messages (line 7). The functions that are used to formulate these messages are generally learnable functions. Within optical networking, the fibre lengths of edges significantly impact the transmission performance over these edges, therefore it is essential to include this information in the formulation of messages.

$$M_t(h_{\omega}^t, h_n^t, e_{\nu}) = \Lambda(e_{\nu}) \times h_{\omega}^t + b(e_{\nu})$$

(1)

As shown in Eq.\cite{1}, we use a matrix ANN ($\Lambda$) and vector valued ANN ($b$) to extract features from the edge feature vectors $e_{\nu}$ in the constructed messages. These messages are then aggregated via a sum operation, as seen in line 7 of algorithm\cite{1} and the inner block of figure\cite{2} to give the future message of node $n$, $m_n^{t+1}$. The sum operator is chosen due to it being permutation agnostic and for its simplicity.

### B. Update Function

The update function is used to take the information (messages) aggregated from the neighbourhood of node $n$ and learn a new hidden state vector $h_n^{t+1}$ to incorporate this new information in the state (line 8). As this is a sequential process, we use a recurrent neural network (RNN) architecture, which can take previous states into account and learn how much of the previous states to use in the next state, whilst producing new abstract representations of the data. RNNs have been shown to struggle with vanishing gradients during training, therefore
we chose to use a gated recurrent unit (GRU). GRUs use reset and update gates to learn how much of the state and input to use or forget depending on the target. They exhibit improved performance over standard RNNs, however have been shown to have reasonable computational complexity.\footnote{\cite{8}}

\begin{equation}
U_t(h^t_n, m^t_n) = \text{GRU}(h^t_n, m^t_n) \tag{2}
\end{equation}

As shown in Eq. (2), the update function consists solely of a GRU function. The current state, \(h^t_n\), and the aggregated messages, \(m^t_n\), are fed in, to produce an updated representation, \(h^{t+1}_n\).

### C. Readout Function

The final graph-level aggregation of the states and features is carried out via the readout function (line 11). The aim is to aggregate all the relevant inter-dependencies between the nodes, via the hidden states, \(h^T_n\), and represent it as a single vector, on which one can regress to the target outputs (throughput, in our case). The readout function is shown in Eq. (3).

\begin{equation}
R(H_N, X_N) = b(\sum_n \sigma[i(h^T_n, x_n)] \odot j(h^T_n)) \tag{3}
\end{equation}

To learn which parts of the hidden vector, \(h^T_n\), are important for the prediction of the target, an attention mechanism was used. The attention mechanism learns weights in the range of \([0, 1]\), that are used to weight a vector, with the goal of learning which parts of the vector are important for the learning task. This is achieved, by feeding the concatenated hidden vector, \(h^T_n\), and node features, \(x_n\), through an ANN and passing the output through a sigmoid function \(\sigma\), which normalises the output between \([0, 1]\). Using an element-wise multiplication (Hadamard product), this vector acts as attention scores to the original hidden vector, \(h^T_n\), which is fed through another ANN \footnote{\cite{5}}. Summing these operations over all nodes gives us the final vector used for the regression layer. The regression layer, \(b\) consists of a single ANN layer, which reduces the output to a scalar value.

Having defined the MPNN model architecture, it now needs to be trained on graph labels, i.e. maximum achievable throughput, to predict this on all unseen graphs. The next section details the methodology for generating the training and testing datasets.

### III. DATASET GENERATION

To train the MPNN, three large sets of graphs, with different numbers of nodes, were generated. To calculate the maximum achievable throughput for nodes sizes below 25, an ILP was used to find optimal RWAs, although for the two larger sets of graphs (\(N = 25 - 100\)), a heuristic algorithm was used. Using these RWAs, the throughput was calculated and then stored as a graph label to be used for training and testing. The following section details the methodology used for generating the topologies, the ILP formulation, the heuristic algorithms, the physical layer impairments (PLI) model and the final calculation of the maximum achievable throughput.

#### A. Topology Generation

A single generative graph model was chosen to create the graph structures for training and testing. These were created via the SNR-BA model, which has been shown to reflect optical core network structures and physical properties\footnote{\cite{6}}. In this generative graph model, distances between nodes are incorporated in the process of edge selection, creating localised hubs within graphs. Nodes are chosen uniformly over a grid, representing the size of the north-American continent, resulting in unique node locations for each graph and therefore giving a greater range of graph structures and physical properties than \(m^3\), where the same real network node locations were used in all generated graphs. The SNR-BA generative graph model then uses these unique node locations to generate the graphs. Inter-node distances were constrained to be at least 100km.

\begin{equation}
D_{\text{fibre}} = \begin{cases} 
1.5 \cdot D_{\text{hav}} & \text{if } D_{\text{hav}} < 1000 \text{ km} \\
1500 \text{ km} & \text{if } 1000 \text{ km} \leq D_{\text{hav}} \leq 1200 \text{ km} \\
1.25 \cdot D_{\text{hav}} & \text{if } D_{\text{hav}} > 1200 \text{ km}
\end{cases} \tag{4}
\end{equation}

To model distance, \(D_{\text{hav}}\), in the generated graphs, their geographical coordinates were used in conjunction with the haversine formula\footnote{\cite{39}}. The haversine formula takes into account the curvature of the earth and calculates distances over a sphere rather than a plane. We also account for realistic fibre distances as shown in Eq. (4). The fibre distances are estimated according to the European Telecommunications Standards Institute (ETSI) guidelines for estimating distance overheads in communication networks\footnote{\cite{39}}.

To demonstrate the model operation on different size topologies, three separate datasets were generated: (i) alpha set with \(10 \leq |N| \leq 15\) (\(\sim 75000\) graphs), beta set with \(25 \leq |N| \leq 45\) (\(\sim 95000\) graphs) and gamma set with \(55 \leq |N| \leq 100\) (\(\sim 75000\) graphs). The nodes increased by 1 and 5 for the alpha and beta, gamma datasets respectively, giving 6, 5 and 10 node scales for the alpha, beta and gamma datasets respectively. To make sure that the model learns performance trends over a variety of edge numbers, edge numbers were chosen by adding an empirically determined percentage of the nodes for a given graph, multiple times, as seen in Eq. (5). Here \(d_n\) was chosen to be 0.2, typical of the relatively sparse core networks and \(i\) varied between 1 and 10 - so that the graph, empirically, has approximately 20% more edges than nodes, as the sparsest core networks have about 20% more edges than nodes, we have used this as the minimum value.\footnote{\cite{8}}

\begin{equation}
|E| = |N| + d_n \cdot |N| \cdot i \tag{5}
\end{equation}

To generate the labels for the maximum achievable throughput values, an optimum RWA that maximises the number of allocated lightpaths (simply referred to as the optimum RWA)
is required for each network considered. To find these optimum RWAs, an ILP formulation was used for $|N| < 25$, and for $|N| = 25 – 100$, a heuristic, as described in the next two sections.

B. Integer Linear Program

To maximise the maximum achievable throughput of the network, an ILP is developed that maximises the number of lightpaths allocated, maximised using the objective function (6). The ILP considered here does not directly optimise the maximum achievable throughput, however assumes that maximising the number of lightpaths allocated is highly correlated to maximising throughput within the network. Maximising the number of lightpaths allows the problem to be specified in terms of an integer objective making the problem less computationally complex. Maximising throughput directly by solving the routing and wavelength allocation problem, a more complex problem to be solved, because of nonlinear interactions between lightpaths. A decision variable $\delta_{w,k,z}$, is used to define a lightpath, where $w \in W$, $k \in K$ and $z \in Z$, are the set of wavelengths, k-shortest paths and node-pairs, respectively. It is defined as in Eq.(7), and is constrained to assigning a lightpath, subject to the normalised traffic matrix ($T^c$) and the objective $M$, defined in Eq.(8). For this work all the training data was generated with uniform traffic, meaning that uniform bandwidth was assumed to be routed between all node-pairs. Here the objective is to maximise $M$, as summarised in Eq.(6).

Using this ILP formulation, optimum RWAs were found for each graph in the alpha dataset, using 156 wavelengths and 20 k-shortest paths. ILP formulations, however, do not scale for larger graphs, and heuristics in conjunction with sequential loading were used for the beta and gamma datasets, as described in the following section.

C. Heuristics

Previously, it has been argued, that heuristics, and specifically, first-fit k-shortest-paths (FF-kSP), can achieve similar performance for estimating capacity in optical networks. These heuristics are used in combination with sequential loading, to load the network until the maximum achievable throughput can be found and FF-kSP heuristic has been shown to give very good performance for this type of task, with low computational complexity. The heuristics here are used for two purposes: (i) to benchmark the performance of the MPNN ($N \leq 25$) (ii) to generate training labels for larger graphs ($N \geq 25$), for which the ILP is infeasible.

Algorithm 2: Sequential loading algorithm

Input: $T_r$, $G$
Output: $RWA_t$

\begin{algorithm}
 ibegin
  1 $i = 0$
  2 $M = 0$
  3 $m_i = 100$
  4 for $i \leq m_i$ do
  5     while $RWA_i \neq blocked$ do
  6         $M \leftarrow M + m_i$
  7         $T_r \leftarrow \lfloor M \cdot T^c \rfloor$
  8         $RWA_i \leftarrow R_F(G, T_r, RWA_{i-1})$
  9     end
  10     $M \leftarrow M - m_i$
  11     $m_i \leftarrow \lfloor m_i/2 \rfloor$
  12 end
 iend
\end{algorithm}

The sequential loading applied is described by algorithm 2. Here the same objective ($M$) is maximised as in the ILP, where $T_r$ and $T_c$ are $N \times N$ matrices representing the connections to be routed and the normalised traffic matrix respectively. $F_R$ is the routing function, which takes three inputs: a graph ($G$), a connections matrix ($T_r$) and the previous RWA. Once blocking is achieved, the iteration ($m_i$) is halved and the network is loaded until blocking is achieved. This iterates $m_i$ times, after which the final RWA is returned, where $m_i$ was chosen to be 6, as more would be adding negligible amounts to the objective. For this work kSP-FF and FF-kSP were used, both of which are common heuristics for solving the RWA problem. FF-kSP was used to generate optimum RWA configurations for the graphs in the beta and gamma datasets, due to its high linear correlation to the ILP-determined performance data, as seen later in Section V A.
Having optimised the RWA of a given network, we need to use these optimum RWA values to calculate the maximum achievable throughput. For this, the physical layer impairments (PLI) over the fibre in which the data is routed need to be taken into account to calculate the SNR values of the individual lightpaths.

D. Physical Layer Impairments Model

To calculate the resultant throughput for a given RWA, we calculated the accumulated SNR for each lightpath assignment. A lightpath \( i = \{ p_i, \lambda_i \} \in R \) has a path \( p_i \) and a wavelength \( \lambda_i \) associated with it and is part of the set of lightpaths for a routing \( R \). To calculate the capacity for this lightpath, one first needs to take into account the edges along which the lightpath travels and their corresponding transmission SNR values. Using \( \text{Eq.} (10) \) and the state of the edge (which wavelengths are occupied), one can calculate the \( \text{SNR}_{i,e} \) value on each of the links \( e \in p_i \). Here \( P_i \) is the launch power, \( \eta_n \) is the nonlinear coefficient, which can be calculated by Eq. (5) in [2] and \( P_{\text{ASE}} \) is the power of the amplified spontaneous emission. In this work, just by way of an example, we assumed multiples of 80km standard single mode fibre spans, with \( \alpha = 0.2 \text{dB/km}, D = 18 \text{ps}/\mu\text{m} \cdot \text{km} \) and \( \gamma = 1.2 \text{W}^{-1} \text{km}^{-1} \), amplified with erbium doped fibre amplifiers (EDFA) with a noise figure of 4dB. Nyquist spaced 32Gbd channels were interfaced with colourless, directionless, and contentionless, reconfigurable optical add-drop multiplexers (CDC-ROADMs), over a constrained C-band (1530–1570 nm) optical bandwidth. The total SNR of that path is then calculated by taking the inverse sum of the noise-to-signal ratio (NSR) values of each link traversed by the path \( p_i \), shown in Eq. (11):

\[
\text{SNR}(i,e) \approx \frac{P_i}{P_{\text{ASE}} + \eta_n P_i^3} \quad (10)
\]

\[
\text{SNR}_i = \left( \sum_{e \in p_i} \frac{1}{\text{SNR}_{i,e}} \right)^{-1} \quad (11)
\]

This SNR is used to calculate the maximum achievable data rate over this lightpath using Eq. (12) in [2]. It can be seen that the capacity of a lightpath mainly depends on the SNR of that path, which, in turn, depends on the length and congestion along the path. \( B_{CH} \) represents the channel bandwidth used, which was kept constant at 32 GHz for all channels.

\[
T_i = 2B_{CH} \log_2(1 + \text{SNR}_i) \quad (12)
\]

\[
T_A = \sum_{i \in R} T_i \quad (13)
\]

The throughputs for all the lightpaths were calculated and summed, as in Eq. (13) to give the total achievable throughput for a particular RWA and network. For the routing of larger networks, multiple fibres needed to be incorporated to allow for all-to-all connectivity, where 4 fibres per edge for \( 25 \leq |N| \leq 45 \) and 16 fibres per edge for \( 55 \leq |N| \leq 100 \) were used. Multiple fibres generally increase the throughput of a network by at least \( \xi \) times, where \( \xi \) is the number of fibres per edge[2]. The hierarchy of graphs however does not change when adding more fibres, i.e. which network is better does not change, therefore not changing the overall distribution of the generated graphs.

Using this methodology, the three datasets, alpha, beta and gamma, were generated via the SNR-BA model, their RWAs optimised either via ILP or FF-kSP (depending on their node scale) and finally their maximum achievable throughputs \( (T_A) \) calculated. The whole process is illustrated in figure [3]. Using \( T_A \) as a training label (target), the MPNN model, defined in Section III was trained in a supervised manner, described in the following section.

IV. TRAINING

Using the datasets created as described in Section III, the three MPNN models were trained to predict the maximum achievable throughput, \( T_A \). The node features \( (x_n) \), chosen to be the degree of each node \((\delta_n)\) and its normalised traffic:

\[
x_n = \delta_n \sum_{u \in N} T_c^{(n,u)}
\]

were used to initialise the node hidden vectors \((h_n^0)\). As throughput depends on the physical properties of the fibre links, the overall transmission quality metric is a critical feature. The number of lightpaths carried over an edge or set of edges, as well as the length of paths taken, influences the overall system performance via the achievable signal-to-noise ratio in the nonlinear optical regime. Throughout testing it was seen that the inverse of the SNR (NSR) was a better feature to use, as it is an additive quantity over a set of edges. In an ideal case, optimal launch powers would be found for each wavelength in the network, however due to the computational complexity associated with carrying this out for many networks (240000 topologies), the assumption of uniform launch powers, i.e. equal input power/lightpath for all lightpaths, was used, similarly to other work[3].

A hidden vector size of 16, with 8 message passing rounds was used for all model training. Larger hidden vector sizes did not seem to provide more accuracy, however added higher computational complexity. The message passing rounds were chosen to work well over the whole spectrum of node scales, where \( T = 8 \) gave good performance for all. As overfitting was initially a problem, a dropout rate of 0.65 with L2 regularisation at a rate of 0.03 was used, where higher values started reducing the accuracy of the model. The learning rate was initialised with a value of 0.001 and decayed exponentially using 10000 steps at a rate of 0.95. The Adam optimisation algorithm was used to train all models, with the graphs in batches of 50 and a single fully-connected regression layer consisting of 256 neurons was used for the final regression output, where larger values did not provide further accuracy without overfitting. To monitor whether the model was tending to overfit, a
validation set of 500 graphs was used to evaluate the performance at each epoch. The training was conducted on a Nvidia V100 16GB GPU, with training generally taking around 72 hours, covering 2000 epochs.

Having trained the three separate models on the different datasets, we used three separate test sets to gauge the performance of the trained inference models. These were generated identically to the methodology laid out in Section III, however were unseen (not used for training) by the model. The following section analyses the accuracy, computation time and generalisation capabilities of the MPNN model using these test sets.

V. RESULTS

To understand how well the MPNN model performs in comparison to other methodologies for estimating the maximum achievable throughput of an optical network, there are three aspects we evaluate: (i) model accuracy (ii) time complexity (iii) generalisation capability to unseen graphs.

To evaluate the accuracy of the MPNN, the three test sets of 6000, 5000 and 10000 graphs, were used to test the alpha, beta and gamma models, respectively. Here 1000 graphs for each node scale were generated. Labels for the graphs in the alpha test set were generated via the ILP formulation, whilst for the beta and gamma test sets, FF-kSP was used. Again, uniform traffic was generated and used for all these results. To determine $R^2$, the coefficient of determination ($R^2$) was used, which measures how much of the variance in the data is explained by the model, by comparing the prediction variance to that of the original data.

$$R^2 = 1 - \frac{\sum_n (y_n - p_n)^2}{\sum_n (y_n - \bar{y})^2}$$

(14)

Where $y_n$ refers to the true label and $p_n$ the predicted label and $\bar{y}$ the mean.

The Pearson’s correlation coefficient $\rho$ was used as an alternative metric and determines how linearly the labels and predictions are related, with a value of 1 signifying a perfect linear correlation. This coefficient is important in the context of surrogate models or cost functions, as a model might be inaccurate (low $R^2$), however have a high linear correlation (high $\rho$), which means that although inaccurate it can predict the relative performance of a network well - vital for optimisation. The predictive accuracy for all three models is analysed and discussed in the following section.

A. Maximum Achievable Throughput

The label that was evaluated is that of maximum achievable throughput ($T_A$). For each of the 6000 graphs in the alpha test set, the performance was evaluated via ILP (used for the labels), FF-kSP, kSP-FF and MPNN and plotted in figure 4(a). It can be seen that the kSP-FF and FF-kSP heuristics underperform compared to the ILP, giving lower $R^2$ values in both, although kSP-FF is worse as it has even lower values for $R^2$ and $\rho$, as seen in table I. This signifies poor predictive accuracy of the actual labels ($T_A$). The performance of kSP-FF is worse than that of FF-kSP, as it prioritises shorter paths over optimising wavelength selection, this is an expected result seen in [23]. FF-kSP generally spreads the usage more evenly across all network links compared to kSP-FF. This spreading of resources uses slightly more spectrum, however achieves much less network congestion. However, the MPNN has learnt on a variety of graphs of these sizes and can accurately predict the throughput trend here, as indicated by its high $R^2$ value of 0.951. Furthermore, one can see the difference in throughput distributions, where the CDF of the different methodologies are plotted in figure 4(b). Here it is clear that the kSP-FF and FF-kSP heuristics give different distributions compared to the ILP. The MPNN however, is able to replicate the original distribution of the ILP well, and therefore it can predict the throughput of the networks better. In addition to the MPNN, a linear and nonlinear ML regression method were evaluated to compare other ML frameworks. Here the degree, variance, connectivity, algebraic connectivity, communicability distance and communicability traffic index were used as input features for the graph. ElasticNet and the artificial neural network (ANN) were trained and tested with the same data, both scoring lower with $R^2$ values of 0.763 and 0.768, respectively.

Another metric evaluated was the Pearson’s correlation coefficient ($\rho$). As for the optimisation, there is significant linear correlation between the real performance and the predicted performance properties. It can be seen that the heuristics generally perform well, and the FF-kSP has a high linear correlation ($\rho = 0.969$) between the estimated throughput values and those calculated via the ILP, as seen in table I. The MPNN, has a similar correlation, with $\rho = 0.975$, meaning it predicts the relative throughput performance of networks well. ElasticNet and the ANN both score $\rho$ values lower than the MPNN and FF-kSP. The high linear correlation of FF-kSP compared to the ILP, makes it a good candidate to evaluate the maximum achievable throughput for the larger graphs, even though the real maximum achievable throughput might be larger.

To further evaluate the performance of the model, the model was applied to the beta and gamma testing datasets that included larger graphs. Here the labels were generated via the FF-kSP heuristic, as the ILP is not able to scale to these

<table>
<thead>
<tr>
<th>Node Scale</th>
<th>Method</th>
<th>$R^2$</th>
<th>$\rho$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10 \leq</td>
<td>N</td>
<td>\leq 15$</td>
<td>MPNN</td>
</tr>
<tr>
<td>$10 \leq</td>
<td>N</td>
<td>\leq 15$</td>
<td>FF-kSP</td>
</tr>
<tr>
<td>$10 \leq</td>
<td>N</td>
<td>\leq 15$</td>
<td>kSP-FF</td>
</tr>
<tr>
<td>$10 \leq</td>
<td>N</td>
<td>\leq 15$</td>
<td>ElasticNet</td>
</tr>
<tr>
<td>$10 \leq</td>
<td>N</td>
<td>\leq 15$</td>
<td>ANN</td>
</tr>
<tr>
<td>$25 \leq</td>
<td>N</td>
<td>\leq 45$</td>
<td>MPNN</td>
</tr>
<tr>
<td>$55 \leq</td>
<td>N</td>
<td>\leq 100$</td>
<td>MPNN</td>
</tr>
</tbody>
</table>

TABLE I: Accuracy of the MPNN model and other capacity estimation methods, measured by the coefficient of determination ($R^2$) and the Pearson’s correlation coefficient ($\rho$).
larger topologies. The MPNN was used to infer the maximum achievable throughput of the graphs in the respective test sets and plotted in figures 5(a) and (b). It is clear that for the beta model, the MPNN performs better than the alpha and gamma models, with a $R^2$ value of 0.973. This is because the beta training set had the most training graphs per node scale. This was possible for the beta model, as the heuristic runs faster on the smaller beta graphs than the larger gamma set graphs, and also faster than the ILP. This can be seen in figures 6(a) and (b). Although the gamma model still achieves a high $R^2$ of 0.948, one can see that the larger throughput values have more variance in them. To remedy this, more samples are needed in the training set for larger node scales. Both the beta and gamma models achieved high linear correlation values ($\rho$) of 0.986 and 0.974 respectively.

One can conclude that heuristics consistently underperform in estimating the throughput in optical networks compared to the ILP solutions. However, FF-kSP is able to predict the trend (linear correlation - $\rho$) well, making it suitable for optimisation tasks and training data generation. The MPNN on the other hand is able to accurately predict throughput values of unseen data, based on similar structure and sizes seen during training. It also has a high linear correlation between the labels and its own predictions, making it an excellent candidate for modelling the throughput of these optical networks. Therefore, the MPNN is able to predict maximum achievable
throughput accurately, although it does not provide a solution, for example for the RWA, of how to reach it. The model could potentially be adjusted to learn the RWA, this is outside the scope of the current work and is left for future research. However, given the extensive training and data needs of this model, what is the real benefit of using it in place of heuristics? This question is addressed in the following section.

B. Computational Time Comparison

The key advantage of using machine learning to model relationships in data, is very low inference times compared to ILP and, even, heuristic methods.

The ILP has a worst-case computational complexity as described in Eq. (15), where \( D \) is the number of demands or connection requests (\( D = \sum_{c \in Z} [T^C \cdot M] \)), \( E \) the number of edges and \( W \) the number of wavelengths used:

\[
O(2^D \cdot E^W)
\]  
(15)

For the heuristic algorithms used, the complexity generally scales as in Eq. (16) but must be modified to include the term \( R \cdot n \) number of sequential loading iterations as the heuristic needs to run many times before finding the optimum RWA:

\[
O(Rn^3(E + Nlog(N)))
\]  
(16)

The advantage of the MPNN, thus, is that it can directly evaluate the network properties, learnt from previous data. For the inference of the MPNN, the complexity is defined in Eq. (17), where \( T \) denotes the number of message passing rounds, defined previously and \( d \) is the length of hidden dimensional vector used, 16 in our case.

\[
O(T \cdot N^2 \cdot d^2)
\]  
(17)

Comparing the equations Eq. (15), Eq. (16) and Eq. (17), it can be seen that the MPNN scales the best, computationally, with number of nodes. To quantify the computational benefits of using MPNNs to model optical networks, graphs with nodes varying from 10 to 20, were used to evaluate the ILP, FF-kSP, kSP-FF and MPNN time performance. Here the node size has been expanded to 20 nodes for the test set to see the computational time scaling better with smaller graphs. This, however, was not possible in the case of the training set, since the ILP computation is too complex. For each graph, the respective methodologies were used to calculate the maximum achievable throughput and their computation times measured and plotted in figure 6(a). The reduction in computation time through using MPNN model can be clearly seen, where it takes approximately 10s of ms, compared to 10s, 100s and 1000s of seconds for kSP-FF, FF-kSP and ILP respectively. The same trend, between FF-kSP and the MPNN, was analysed for the larger networks using the beta and gamma test sets, with the results shown in figure 6(b). The MPNN again shows minimum computation time increase for these node ranges, compared to the heuristic method.

Therefore, the MPNN model can be seen as an accurate and fast method for evaluating performance metrics of graphs, which are generally computationally difficult to evaluate. This would allow for the fast evaluation of a large number of graphs within any topology design process. However, how well does this model generalise when varying the input structure and traffic distributions that affect the throughput of the network? This is the question explored in the next section.

C. Generalisation Capability

The generalisation capability of an ML model refers to the ability of the model to operate over distributions not seen during the training process. In the context of this work, this could encompass different graph structures, different traffic distributions or distance scales. Here we choose to evaluate the model against two different graph structures and a change in input traffic distribution.

To test different graph structures, two common generative models were used to vary the structures tested, the Erdos-Renyi (ER) and Barabasi-Albert (BA) models. Using these models 5000 graphs with \( 25 \leq |N| \leq 45 \) were generated per generative model. After calculating the throughputs of these graphs using the FF-kSP heuristic and the Gaussian Noise (GN) model, we feed the graphs through the MPNN model to predict the throughput values.

When comparing the accuracy of the model over these varied graph structures, one can see that the accuracy, in terms of coefficient of determination \( R^2 \), drops significantly, as seen in table II.

<table>
<thead>
<tr>
<th>Graph Type</th>
<th>( \gamma )</th>
<th>( R^2 )</th>
<th>( \rho )</th>
<th>( D )</th>
<th>( F_{WSD} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>ER</td>
<td>0.0</td>
<td>0.776</td>
<td>0.928</td>
<td>0.096</td>
<td>0.000286</td>
</tr>
<tr>
<td>BA</td>
<td>0.0</td>
<td>0.830</td>
<td>0.938</td>
<td>0.125</td>
<td>0.000201</td>
</tr>
<tr>
<td>SNR-BA</td>
<td>0.0</td>
<td>0.973</td>
<td>0.986</td>
<td>0.000</td>
<td>0.000000</td>
</tr>
<tr>
<td>SNR-BA</td>
<td>0.2</td>
<td>0.901</td>
<td>0.983</td>
<td>0.079</td>
<td>2.13\times10^{-6}</td>
</tr>
<tr>
<td>SNR-BA</td>
<td>0.4</td>
<td>0.907</td>
<td>0.983</td>
<td>0.075</td>
<td>2.13\times10^{-6}</td>
</tr>
<tr>
<td>SNR-BA</td>
<td>0.6</td>
<td>0.906</td>
<td>0.983</td>
<td>0.060</td>
<td>2.13\times10^{-6}</td>
</tr>
<tr>
<td>SNR-BA</td>
<td>0.8</td>
<td>0.911</td>
<td>0.984</td>
<td>0.058</td>
<td>2.13\times10^{-6}</td>
</tr>
<tr>
<td>SNR-BA</td>
<td>1.0</td>
<td>0.907</td>
<td>0.982</td>
<td>0.062</td>
<td>2.13\times10^{-6}</td>
</tr>
</tbody>
</table>

| TABLE II: Accuracy for generalisation capability, measured by the coefficient of determination \( R^2 \) and \( \rho \). The variable \( \gamma \) determines how locally skewed the traffic is and \( D \) is the absolute distance between the test throughput and the original training throughput (ks-2s test). |
FIG. 6: (a) Comparison between computation times for throughput of networks over different node scales, using ILP, FF-kSP, kSP-FF and MPNN. (b) Comparison between computation times for throughput of networks, using FF-kSP and MPNN for 25 ≤ 100.

BA graphs. We can see that the ER and BA graph’s WSD is larger, where the ER structures are the most different from the original test graphs. In addition, due to the difference in graph structures, there are large differences in the throughput distributions. This is measured by the Kolmogorov-Smirnov two sample test, which returns a distance $D$, which signifies the largest absolute difference between the CDF of two distributions. In Table II, the large $D$ values for ER and BA test sets, signify the difference in throughput distributions to those seen during training. These $D$ values are larger than those of the SNR-BA test distributions, meaning their throughput distributions are further from those arising from SNR-BA distributions. The Pearson’s correlation coefficient ($\rho$) however remains high, which shows that it still indicates the throughput trends of the networks well. This is important for optimisation, as the cost function might not need to accurately describe throughput exactly, however it needs to describe one graph being better than another.

To investigate how a change in the traffic demand matrix affects the accuracy of the model, we generated localised skewed traffic matrices for 5000 graphs. By defining the traffic as in Eq. (18), we generated 5 different traffic skews shown in Table II.

$$T_{ij}^c = \frac{1}{\sum_{k \in N} D_{i,j}}$$

(18)

For each of these skewed traffic matrices, we tested the MPNN accuracy in terms of $R^2$ and $\rho$. The $R^2$ values drop by about 7%, and remain constant for the different skew values ($\gamma > 0$). The Pearson’s correlation coefficient ($\rho$) value remain high at around 0.98, unchanged from the uniform traffic distribution results, meaning that it still predicts the throughput trend well.

The large variation in performance, in terms of $R^2$ for ER and BA graphs, indicates that the trained MPNN model does not generalise well to largely different graph structures. This highlights the importance of using a variety of different structures within the training dataset and that an expansion of the training data is necessary here to represent different graph structures. Once the training set is more representative, by generating a variety of graph structures, it accurately evaluates the vast solution space of the topology design problem.

VI. CONCLUSION

In this paper the well-known NP-hard problem of estimating optimal performance of an optical network, was investigated, focusing, in particular, on the maximum achievable throughput. Understanding and being able to predict it, is vital for optimising optical networks for the future, as to maximise the longevity of investments made in infrastructure for communications or for improving existing topologies. We have proposed the use of geometric deep learning framework to learn this network property from a previously generated training dataset of graphs.

By applying this methodology, we showed that the model can accurately model the maximum achievable throughput of previously unseen graphs, proven by high coefficient of determination accuracy ($R^2$) and Pearson’s correlation coefficient ($\rho$) values. This accuracy was achieved with significant time savings of multiple orders of magnitude, enabling rapid prediction (~ 30ms for 100 node graphs) of maximal performance of a large number of graphs.

Furthermore, both the importance and weaknesses of the training datasets were highlighted by applying the model to other graph structures, such as ER and BA graphs, where the model struggled to give accurate predictions, showing that the training set of graphs needed to be expanded. The generalisation of different traffic input distributions generally performed well and showed suitability for future optimisation purposes.
This type of modelling is invaluable to further optimise the physical topologies of optical networks, by using this as a surrogate model in future optimisation. It directly enables using the maximum achievable throughput of a topology in the cost function of any number of optimisation algorithms, as well as giving huge time savings and therefore enhancing the optimisation. In addition, networks are ever-evolving and are not necessarily purely fixed grid. They might have a flexible grid, regeneration devices or intermediate grooming at routers or switches. These changes in network configuration could be included in future training sets by using them as node or edge features and generating more training data using a range of these configurations. Further investigations will include work on analysing the resilience of networks, accurate traffic modelling and generalisation to multiple fibre systems using MPNNs.

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