Learning Local Receptive Fields and their Weight Sharing Scheme on Graphs

Jean-Charles Vialatte\textsuperscript{1,2}, Vincent Gripon\textsuperscript{2}, Gilles Coppin\textsuperscript{2}
\textsuperscript{1}City Zen Data
55 rue Charles Nungesser
29490 Guipavas, France
jean-charles.vialatte@cityzendata.com
\textsuperscript{2}IMT Atlantique / CNRS Lab-STICC
Technopole Brest Iroise
29238 Brest, France
name.surname@imt-atlantique.fr

\textbf{Abstract}—We propose a generic layer formulation that extends the properties of convolutional layers to any domain that can be described by a graph topology. Namely, we use the support of its adjacency matrix to design learnable weight sharing filters able to exploit the underlying structure of signals in the same fashion as for images. The proposed formulation makes it possible to learn the weights of the filter as well as a scheme that controls how they are shared across the graph. We perform validation experiments with image datasets and show that these filters offer performances comparable with convolutional ones.

\textbf{Index Terms}—deep learning, convolutional neural networks, local receptive fields, graph signal processing

\section{I. INTRODUCTION}

Convolutional Neural Networks (CNNs) have achieved state-of-the-art accuracy in many supervised learning challenges \cite{1,2,3,4,5,6}. For their ability to absorb huge amounts of data with lesser overfitting, deep learning \cite{7} models are the golden standard when a lot of data is available. CNNs benefit from the ability to create stationary and multi-resolution low-level features from raw data, independently from their location in the training images. Some authors draw a parallel between these features and scattering transforms \cite{8}.

Obviously CNNs rely on the ability to define a convolution operator (or a translation) on signals. On images, this amounts to learn local receptive fields \cite{9} that are convolved with training images. Considering images to be defined on a grid graph, we point out that the receptive fields of vertices are included in their neighbors – or, more generally, a neighborhood.

Reciprocally, convolution requires more than the neighborhoods of vertices in the underlying graph, as the operator is able to match specific neighbors of distinct vertices together. For instance, performing convolution on images requires the knowledge of coordinates of pixels, that is not directly accessible when considering a grid graph (c.f. \cite{10}, \cite{11}). In this paper we are interested in demonstrating that the underlying graph is nevertheless enough to achieve comparable results.

The convolution of a signal can be formalized as its multiplication with a convolution matrix. In the case of images and for small convolution kernels, it is interesting to note that this convolution matrix has the same support as a lattice graph. Using this idea, we propose to introduce a type of layer based on a graph that connects neurons to their neighbors. Moreover, convolution matrices are entirely determined by a single row, since the same weights appear on each one. To imitate this process, we introduce a weight sharing learning procedure, that consists in using a limited pool of weights that each row of the obtained operator can make use of.

Section II presents related work. Section III describes our methodology and the links with existing architectures. Section IV contains experimental results. Section V is a conclusion.

\section{II. RELATED WORK}

Due to the effectiveness of CNNs on image datasets, models have been proposed to adapt them to other kind of data, e.g. for shapes and manifolds \cite{12,13}, molecular datasets \cite{14}, or graphs \cite{15,16,13}. A review is done in \cite{17}. In particular, CNNs have also been adapted to graph signals, such as in \cite{18,19} where the convolution is formalized in the spectral domain of the graph defined by its Laplacian \cite{20}. This approach have been improved in \cite{21}, with a localized and fast approximated formulation, and has been used back in vision to breed isometry invariant representations \cite{22}.

For non-spectral approaches, feature correspondences in the input domain allow to define how the weights are tied across the layer, such as for images or manifolds. For graphs and graph signals, such correspondences doesn’t necessarily exist. For example, in \cite{16} (where the convolution is based on multiplications with powers of the probability transition matrix) weights are tied according to the power to which they are attached, in \cite{15} an ordering of the nodes is used, in \cite{13} an embedding is learned from the degrees of the nodes. These choices are arbitrary and unsimilar to what is done by regular convolutions. On the contrary, we propose a generic layer formulation that allows to also learn how the weights are linearly distributed over the local receptive field.

Our model is first designed for the task of graph signal classification, but another common task is the problem of node classification such as in \cite{16,23,24,13}. Models learning part of their structures have also been proposed, such as in \cite{25,26}. Moreover, because our model strongly resembles regular convolutions, it can be related to variants derived of those, such as group equivariant convolutions \cite{27}.
III. METHODOLOGY

We first recall the basic principles of Deep Neural Networks (DNNs) and CNNs, then introduce our proposed graph layer.

A. Background

DNNs [28] consist of a composition of layers, each one parametrized by a learnable weight kernel \(W\) and a nonlinear function \(f: \mathbb{R} \rightarrow \mathbb{R}\). Providing the input of such a layer is \(\mathbf{x}\), the corresponding output is then:

\[
y = f(W \cdot \mathbf{x} + \mathbf{b}),
\]

where \(\cdot\) is the matrix product operator, \(f\) is applied component-wise and \(\mathbf{b}\) is a learnable bias vector.

The weight kernels are learned using an optimization routine usually based on gradient descent, so that the DNN is able to approximate an objective function. A DNN containing only this type of layer is called Multi-Layer Perceptron (MLP).

In the case of CNNs [29], some of the layers have the particular form of convolution filters. In this case, the convolutional operation can also be written as the product of the input signal with a matrix \(W\), where \(W\) is a Toeplitz matrix. Previous works [30], [31], [32], [33] have shown that to obtain the best accuracy in vision challenges, it is usually better to use very small kernels, resulting in a sparse \(W\). Figure 1 depicts a convolutional layer.

![Figure 1. Depiction of a 1D-convolutional layer and its associated matrix \(W\).](image)

B. Proposed Method

We propose to introduce another type of layer, that we call receptive graph layer. It is based on an adjacency matrix and aims at extending the principle of convolutional layers to any domain that can be described using a graph.

Consider an adjacency matrix \(A\) that is well fitted to the signals to be learned, in the sense that it describes an underlying graph structure between the input features. We define the receptive graph layer associated with \(A\) using the product between a third rank tensor \(S\) and a weight kernel \(W\). For now, the tensor \(W\) would be one-rank containing the weights of the layer and \(S\) is of shape \(n \times n \times \omega\), where \(n \times n\) is the shape of the adjacency matrix and \(\omega\) is the shape of \(W\).

On the first two ranks, the support of \(S\) must not exceed that of \(A\), such that \(A_{ij} = 0 \Rightarrow \forall k, S_{ijk} = 0\).

Overall, we obtain:

\[
y = f(W \cdot S \cdot \mathbf{x} + \mathbf{b}),
\]

where here \(\cdot\) denotes the tensor product.

Intuitively, the values of the weight kernel \(W\) are linearly distributed to pairs of neighbours in \(A\) with respect to the values of \(S\). For this reason, we call \(S\) the scheme (or weight sharing scheme), of the receptive graph. In a sense, this scheme tensor is to the receptive graph what the adjacency matrix is to the graph. An example is depicted in Figure 2.

![Figure 2. Depiction of a graph, the corresponding receptive graph and its associated weight sharing scheme \(S\).](image)

Alike convolution on images, \(W\) can be extended as a third-rank tensor to include multiple input and output channels (also known as feature maps). It is worth mentioning that an implementation must be memory efficient to take care of a possibly large sparse \(S\).

C. Training

The training procedure requires learning both \(S\) and \(W\). We perform the two jointly. Learning \(W\) amounts to learning weights as in regular CNNs, whereas learning \(S\) amounts to learning how these weights are tied over the receptive fields. We also experiment a fine-tuning step, which consists in freezing \(S\) in the last epochs.

Because of our inspiration from CNNs, we propose constraints on the parameters of \(S\). Namely, we impose them to be between 0 and 1, and to sum to 1 along the third dimension. Therefore, the vectors on the third rank of \(S\) can be interpreted as performing a weighted average of the parameters in \(W\).

We test two types of initialization for \(S\). The first one consists in distributing one-hot-bit vectors along the third rank. We impose that for each receptive field, a particular one-hot-bit vector can be distributed at most once more than any other. We refer to it as one-hot-bit initialization. The second one consists in using a uniform random distribution with limits as described in [34].

D. Extension and Discussion

For simplicity we restricted our explanation to square adjacency matrices. In the case of oriented graphs, one could remove the rows and columns of zeros and obtain a receptive graph with a distinct number of neurons in the input \((n)\) than
in the output \((m)\). As a result, receptive graph layers extend usual ones, as explained here:

1) To obtain a fully connected layer, one can choose \(\omega\) to be \(nm\) and \(S\) the matrix of vectors that contains all possible one-hot-bit vectors.

2) To obtain a convolutional layer, one can choose \(\omega\) to be the size of the kernel. \(S\) would be one-hot-bit encoded along its third rank and circulant along the first two ranks. A stride \(>1\) can be obtained by removing the corresponding rows.

In our case, \(S\) is more similar to that obtained when considering convolutional layers, with the noticeable differences that we do not force which weight to allocate for which neighbor along its third rank and it is not necessarily circulant along the first two ranks.

Note that without the constraint that the support of \(S\) must not exceed that of \(A\) (or if the used graph is complete), the proposed formulation could also be applied to structure learning for the input features \([35],[36]\). For instance, summing such learned \(S\) along the third rank gives hints about how they are correlated to each others. Also, a regularization along the third rank could be used to drop connections during training \([37]\).

In Figure 3, we plot the test error rate on MNIST for various receptive graph layers with convolutional ones. For this purpose, we use image datasets, but restrain priors about the underlying structure.

We observe that even without knowledge of the underlying euclidean structure, receptive grid graph layers obtain comparable performances as convolutional ones. When the ordering is known, they match convolutions.

To compare with convolutions, we also plot in Figure 3 the obtained product \(W \cdot S\) for one feature map, when using the square of the grid graph. For a convolutional layer, each diamond of this plot would be identical.

We consider a grid graph that connects each pixel to itself and its 4 nearest neighbors (or less on the borders). We also use the square of this graph (pixels are connected to their 13 nearest neighbors, including themselves), the cube of this graph (25 nearest neighbors), up to 10 powers (211 nearest neighbors). In this subsection, we use one-hot-bit initialization.

We test the model under two setups: either the ordering of the node is unknown, and then the one-hot-bit vectors are distributed randomly and modified upon training; either an ordering of the node is known, and then the one-hot-bit vectors are distributed in a circulant fashion in the third rank of \(S\) which is freeze in this state. We use the number of nearest neighbors as for the dimension of the third rank of \(S\). We also compare with a convolutional layer of size 5x5, thus containing as many weights as the cube of the grid graph.

Table I summarizes the obtained results. The ordering is unknown for the first result given, and known for the second result between parenthesis.

We observe that even without knowledge of the underlying structure, receptive grid graph layers obtain comparable performances as convolutional ones. When the ordering is known, they match convolutions.

A. Description

In the following experiments, we are interested in comparing various receptive graph layers with convolutional ones. For this purpose, we use image datasets, but restrain priors about the underlying structure.

We first present experiments on MNIST \([38]\). It contains 10 classes of gray levels images (28x28 pixels) with 60'000 examples for training, 10'000 for testing. We also do experiments on a scrambled version to hide the underlying structure, as done in previous work \([39]\). Then we present experiments on Cifar10 \([40]\). It contains 10 classes of RGB images (32x32 pixels) with 50'000 examples for training, 10'000 for testing.

B. Experiments with MNIST

On MNIST, we use models composed of a single receptive graph (or convolutional) layer without pooling. For each of those models, the first layer consists of 50 feature maps, followed by a fully connected layer of 300 neurons, and terminated by a softmax layer of 10 neurons. Rectified Linear Units \([41]\) are used for the activations and a dropout \([42]\) of 0.5 is applied on the fully-connected layer. Unless stated otherwise, weights are initialized random uniformly \([34]\). A regularization with weight \(10^{-5}\) is applied to input layers \([43]\).

Optimizations are done with ADAM \([44]\) up to 100 epochs and a fine-tuning step while \(S\) is frozen is done for up to 50 additional epochs.

In Figure 4, we plot the test error rate on MNIST for various normalizations when using the square of the grid graph, as a function of the number of epochs of training. We observe that they have little influence on the performance and sometimes improve it a bit. Thus, their usage is validated as optional hyperparameters.
Convolutional layers of 64 feature maps, with max pooling and local response normalization, followed by two fully connected layers of 384 and 192 neurons. We switched each convolutional layer with receptive graph layers, but kept the pooling ones. We compare two different supports: the one obtained by using the underlying graph of a regular 5x5 convolution, and the support of the square of the grid graph. Optimization is done with stochastic gradient descent on 375 epochs where $S$ is freezeed on the 125 last ones. $S$ is initialized uniform randomly (except for the baseline where $S$ is not learned). Hence, the ordering of the nodes is unknown, but the multiresolution structure information is kept through pooling layers. Results are summarized in table III.

Despite the fact that $S$ have been initialized without the knowledge of the ordering of the nodes, both receptive graph layers attain accuracies comparable with the baseline. It is even surpassed by a little amount. This is quite remarkable because it means that learning the weight sharing scheme $S$ have made the classifier understand the ordering structure of the nodes.

V. Conclusion

We introduced a new class of layers for deep neural networks which consists in using the support of a graph operator and linearly distributing a pool of weights over the defined edges. The linear distribution is learned jointly with the pool of weights. Thanks to these structural dependencies, we showed it is possible to share weights in a fashion similar to Convolutional Neural Networks (CNNs).

We performed experiments on vision datasets where the receptive graph layer obtains similar performance as convolutional ones, even when the underlying image structure is hidden. We believe that with further work, the proposed layer could fully extend the performance of CNNs to many other domains described by a graph.

Future works will also include exploration of more advanced graph inference techniques. One example is using gradient descent from the supervised task at hand [19]. We can also notice that in our case, this amounts to select receptive fields, breeding another avenue [46].

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