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On

Convolution of Graph Signals

And

Deep Learning on Graph Domains

Abstract

This manuscript is a thesis submitted to apply for a doctorate. It is devoted to two subjects. The first one is about extending the discrete convolution to graph signals. The second one is about extending neural networks to graph domains. Both subjects are related since neural networks can make use of convolutions to leverage the underlying structure of their input domain.

Résumé

Ce manuscrit est une thèse soumise pour candidater au grade de docteur. Il est dévolu à deux sujets. Le premier traite d'extensions de la convolution discrète aux signaux sur graphe. Le second traite d'extensions de l'ensemble de définition des réseaux de neurones à des graphes. Les deux sujets sont reliés car les réseaux de neurones peuvent tirer profit de la structure sous-jacente de leur ensemble de définition à l'aide de convolutions.

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Introduction

One of the first appearances of the *convolution* operation was in the eighteenth century (D'Alembert, 1754, according to Dominguez-Torres, 2010). Since then it has been used in a wide range of domains, including applied mathematics, physics, engineering, computer science and real world problems. In today's era of computerized industry and big data, the convolution has never been more useful. A famous example is its usage in deep learning algorithms for image processing (LeCun et al., 2015). But this is just the tip of the iceberg. Years after years, IT companies acquire more and more data. With hashing algorithms, accessing single points or moderately sized batches of data is relatively easy. However, processing the entire database's knowledge is another story. Algorithms that do not scale well are too time consuming, so only those that traverse the entire database only a few times remain feasible. And that is the point: traversing the database (done in a paralleled manner) to process it can be modeled with a convolution. Therefore this little mathematical tool is bound to play a great role! An example can be seen in the company that funded this Ph.D.: one of its technologies for processing large quantities of data is a programming language centered around a few frameworks. One of them is nothing more than a reimplementation of the convolution by various complicated functions, which helps tremendously to produce simplified scripts.

The subject of this thesis is a timely one, since deep learning have never received as much spotlight as in the last decade. However, deep learning

models are often very specialized to their use cases. Standard Convolutional Neural Networks (CNNs) can only be applied to datasets for which each object can be modeled by a signal defined on an Euclidean domain, like images or sounds. This is because the discrete convolution takes into account the structure of the domain of its inputs, in addition to raw data. Our goal is to study and find ways to extend deep learning models to signals defined on a broader range of domains. To this end we build an algebraic theory of convolutions of graph signals, with the hope to characterize what a more generic definition of convolution should be and what properties it should preserve to keep its usefulness in deep learning models. We choose to model the domain of signals under study with a graph, since this structure can represent a wide range of domains. Our subject fits the idea that efforts in the Artificial Intelligence (AI) field should tend toward a general-purpose AI, and in a lesser extent, that AI-based algorithm, like deep learning, should seek genericity.

The ultimate aim is to use these general-purpose technologies and apply them to all sorts of important real world problems.

— Demis Hassabis

This manuscript is broken down into three chapters. Each chapter is preceded by a short overview so that the reader can grasp its essential contents at a glance. In Chapter 1, we present our domains of interest with a selected literature review. Then, in Chapter 2, we theorize an algebraic understanding of convolutions of graph signals. Finally, in Chapter 3, we study neural networks intended for graph domains. We summarize our contributions in the conclusion and dissuss limitations and perspectives.

Chapter 1

Presentation of the field

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Chapter overview

In this chapter, we present notions related to our domains of interest. One of them, deep learning, is the field of research that focuses on a particular class of functions: neural networks. Since we try to employ a rigorous approach, we first define properly their input domain and their codomain, which can be modeled as tensor spaces. In particular, we give original definitions of tensors in Section 1.1 that are appropriate for the study of neural networks. We also explain how data is handled and manipulated. We give definitions of some binary operations that are important for our study: tensor contraction, and convolution. In Section 1.2, we define neural networks, discuss their biological interpretation, present how they learn, and relate some historical advances. Then we introduce common layers, especially convolutional ones for which we demonstrate a little result that helps toward our study. In the last section, Section 1.3, we present the field of deep learning on graphs. We start with definitions about graphs and signals, and then we describe use cases. Finally, we give a review of stateof-the-art models in two separate subsections, one on spectral methods, and the other one on vertex-domain methods.

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1.1 Tensors

Intuitively, tensors in the field of deep learning are defined as a generalization of vectors and matrices, as if vectors were tensors of rank 1 and matrices were tensors of rank 2. That is, they are objects in a vector space and their dimensions are indexed using as many indices as their rank, so that they can be represented by multidimensional arrays. In mathematics, a tensor can be defined as a special type of multilinear function (Bass, 1968; Marcus, 1975; Williamson, 2015) which can be represented by a multidimensional array. Alternatively, Hackbush proposes a mathematical construction of a tensor space as a quotient set of the span of an appropriately defined tensor product (Hackbusch, 2012), which coordinates in a basis can also be represented by a multidimensional array. In particular in the field of mathematics, tensors enjoy an intrinsic definition that neither depends on a representation nor would change the underlying object after a change of basis, whereas in our domain, tensors are confounded with their representation.

1.1.1 Definition

Our definition of tensors is such that they are a bit more than multidimensional arrays but not as much as mathematical tensors. They are embedded in a vector space, called tensor space, so that deep learning objects can be later defined rigorously.

Given canonical bases, we first define a tensor space, then we relate it to the definition of the tensor product of vector spaces.

Definition 1. Tensor space

We define a *tensor space* \mathbb{T} of rank r as a vector space such that its canonical basis is a Cartesian product of the canonical bases of r finite-dimensional vector spaces.

Its shape is denoted $n_1 \times n_2 \times \cdots \times n_r$, where the $\{n_k\}$ are the dimensions of the vector spaces.

Remark. Unless stated otherwise, vector spaces are assumed to be over the field of real numbers \mathbb{R} .

Definition 2. Tensor product of vector spaces

Given r vector spaces V_1, V_2, \dots, V_r , their *tensor product* is the tensor space \mathbb{T} spanned by the Cartesian product of their canonical bases under coordinatewise sum and outer product.

We use the notation
$$\mathbb{T} = \bigotimes_{k=1}^{r} \mathbb{V}_k$$
.

Remark. This simpler definition is indeed equivalent with the definition of the tensor product given in (Hackbusch, 2012, p. 51). The drawback of our definition is that it depends on the canonical bases, which at first can seem limiting as being canonical implies that they are bounded to a certain system of coordinates. However this is not a concern in our domain as we need not distinguish tensors from their representation.

Naming convention

We will also call *vector space* a tensor space of rank 1. In case there is a vector space that we do not need to see as a tensor space of rank 1, we may use the term *linear space* instead. We also make a clear distinction between the terms *dimension* (that is, for a tensor space it is equal to $\prod_{k=1}^{r} n_k$) and the term *rank* (equal to r). Note that some authors use the term *order* or *mode* instead of *rank* as the latter is also affected to another notion.

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Definition 3. Tensor

A *tensor* t is an object of a tensor space. The *shape* of t, which is the same as the shape of the tensor space it belongs to, is denoted $n_1^{(t)} \times n_2^{(t)} \times \cdots \times n_r^{(t)}$.

1.1.2 Manipulation

In this subsection, we describe notations and operators used to manipulate data stored in tensors. The formalism we present here should be familiar to the researcher in the domain, since it is similar to the notations used by NumPy (Oliphant, 2006) and most deep learning libraries *e.g.* TensorFlow (Abadi et al., 2015), PyTorch (Paszke et al., 2017), Mxnet (Chen et al., 2015), Keras (Chollet, 2015).

Definition 4. Indexing

An *entry* of a tensor $t \in \mathbb{T}$ is one of its scalar coordinates in the canonical basis, denoted $t[i_1, i_2, \dots, i_r]$.

More precisely, if $\mathbb{T}=\bigotimes_{k=1}^r\mathbb{V}_k$, with bases $((e_k^i)_{i=1,\dots,n_k})_{k=1,\dots,r}$, then we have

$$t = \sum_{i_1=1}^{n_1} \cdots \sum_{i_r=1}^{n_r} t[i_1, i_2, \dots, i_r](e_1^{i_1}, \dots, e_r^{i_r})$$

We call the *index space* of $\mathbb T$ the Cartesian product of integer intervals $\mathcal I=\prod_{k=1}^r [\![1,n_k]\!].$

Remark. When using an index i_k for an entry of a tensor t, we implicitly assume that $i_k \in [1, n_k^{(t)}]$ unless otherwise specified.

Definition 5. Subtensor

A *subtensor* t' is a tensor of same rank composed of entries of t. We denote

$$\begin{cases} t' = t[[i_1^1, \dots, i_{n_1(t')}^1], \dots, [i_1^r, \dots, i_{n_r(t')}^r]] \\ \text{where } t'[j_1, \dots, j_r] = t[i_{j_1}^1, \dots, i_{j_r}^r] \end{cases}$$

Given p, if $[i_1^p,\ldots,i_{n_p(t')}^p]$ is a singleton, we drop the brackets. If it is a strictly increasing contiguous sequence, we write instead $i_1^p:i_{n_p(t')}^p$. We drop the left (or right) of: if it is the possible lower (or upper) bound.

Remark. This notation for indexing subtensors is not the same as the one used by NumPy. However the use of : is similar (except that we also include the right of :).

Definition 6. Slicing

A *slice* operation, along the last ranks $\{r_1, r_2, \dots, r_s\}$, and indexed by $(i_{r_1}, i_{r_2}, \dots, i_{r_s})$, is a morphism $s : \mathbb{T} = \bigotimes_{k=1}^r \mathbb{V}_k \to \bigotimes_{k=1}^{r-s} \mathbb{V}_k$, such that:

$$s(t)[i'_1, i'_2, \dots, i'_{r-s}] = t[i'_1, i'_2, \dots, i'_{r-s}, i_{r_1}, i_{r_2}, \dots, i_{r_s}]$$

 $i.e.$ $s(t) := t[:, :, \dots, :, i_{r_1}, i_{r_2}, \dots, i_{r_s}]$

where := means that entries of the right operand are assigned to the left operand. We denote $t_{i_{r_1},i_{r_2},...i_{r_s}}$ and call it the *slice* of t. Slicing along a subset of ranks that are not the lasts is defined similarly. $s(\mathbb{T})$ is called a *slice subspace*.

Definition 7. Flattening

A *flatten* operation is an isomorphism $f : \mathbb{T} \to \mathbb{V}$, between a tensor space \mathbb{T} of rank r and an n-dimensional vector space \mathbb{V} , where $n = \prod_{k=1}^{r} n_k$. It is

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characterized by a bijection in the index spaces $g:\prod_{k=1}^r [\![1,n_k]\!] \to [\![1,n]\!]$ such that

$$\forall t \in \mathbb{T}, f(t)[g(i_1, i_2, \dots, i_r)] = f(t[i_1, i_2, \dots, i_r])$$

We call an inverse operation a *de-flatten* operation.

Row major ordering

The choice of g determines in which order the indexing is made. g is reminiscent of how data of multidimensional arrays or tensors are stored internally by programming languages. In most tensor manipulation languages, incrementing the memory address (*i.e.* the output of g) will first increment the last index i_r if $i_r < n_r$ (and if else $i_r = n_r$, then $i_r := 1$ and ranks are ordered in reverse lexicographic order to decide what indices are incremented). This is called *row major ordering*, as opposed to *column major ordering*. That is, in row major, g is defined as

$$g(i_1, i_2, \dots, i_r) = \sum_{p=1}^r \left(\prod_{k=p+1}^r n_k\right) i_p$$
 (1)

Definition 8. Reshaping

A *reshape* operation is an isomorphism defined on a tensor space $\mathbb{T} = \bigotimes_{k=1}^r \mathbb{V}_k$ such that some of its basis vector spaces $\{\mathbb{V}_k\}$ are de-flattened and some of its slice subspaces are flattened.

1.1.3 Binary operations

We define binary operations on tensors that we'll later have use for. In particular, we define *tensor contraction* which is sometimes called *tensor multiplication*, *tensor product* or *tensor dotproduct* by other sources. We also define *convolution* and *pooling* which serve as the common building blocks of convolution neural network architectures.

Definition 9. Contraction

A *tensor contraction* between two tensors, along ranks of same dimensions, is defined by natural extension of the dot product operation to tensors. More precisely, let \mathbb{T}_1 a tensor space of shape $n_1^{(1)} \times n_2^{(1)} \times \cdots \times n_{r_1}^{(1)}$, and \mathbb{T}_2 a tensor space of shape $n_1^{(2)} \times n_2^{(2)} \times \cdots \times n_{r_2}^{(2)}$, such that $\forall k \in [\![1,s]\!], n_{r_1-(s-k)}^{(1)} = n_k^{(2)}$, then the tensor contraction between $t_1 \in \mathbb{T}_1$ and $t_2 \in \mathbb{T}_2$ is defined as:

$$\begin{cases} t_1 \otimes t_2 = t_3 \in \mathbb{T}_3 \text{ of shape } n_1^{(1)} \times \dots \times n_{r_1-s}^{(1)} \times n_{s+1}^{(2)} \times \dots \times n_{r_2}^{(2)} \text{ where } \\ t_3[i_1^{(1)}, \dots, i_{r_1-s}^{(1)}, i_{s+1}^{(2)}, \dots, i_{r_2}^{(2)}] = \\ \sum_{k_1=1}^{n_1^{(2)}} \dots \sum_{k_s=1}^{n_s^{(2)}} t_1[i_1^{(1)}, \dots, i_{r_1-s}^{(1)}, k_1, \dots, k_s] \, t_2[k_1, \dots, k_s, i_{s+1}^{(2)}, \dots, i_{r_2}^{(2)}] \end{cases}$$

For the sake of simplicity, we omit the case where the contracted ranks are not the last ones for t_1 and the first ones for t_2 . But this definition still holds in the general case subject to a permutation of the indices.

Definition 10. Covariant and contravariant indices

Given a tensor contraction $t_1 \otimes t_2$, indices of the left hand operand t_1 that are not contracted are called *covariant* indices. Those that are contracted are called *contravariant* indices. For the right operand t_2 , the naming convention is the opposite. The set of covariant and contravariant indices of both operands are called the *transformation laws* of the tensor contraction.

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Remark. Contrary to most mathematical definitions, tensors in deep learning are independent of any transformation law, so that they must be specified for tensor contractions.

Einstein summation convention

The Einstein summation convention is a notational convention to write a sum-product expression as a product expression. The summation indices are those that appear simultaneously in the superscript of the left operand and in the subscript of the right one, if subscripts precede superscripts in the notation, or else vice-versa. For example, a dot product is written $u_k v^k = \lambda$ and a matrix product is written $A_i{}^k B_k{}^j = C_i{}^j$.

The tensor contraction of Definition 9 can be rewritten using this convention:

$$t_{1i_{1}^{(1)}\cdots i_{r_{1}-s}^{(1)}}{}^{k_{1}\cdots k_{s}}t_{2k_{1}\cdots k_{s}}{}^{i_{s+1}^{(2)}\cdots i_{r_{2}}^{(2)}}=t_{3i_{1}^{(1)}\cdots i_{r_{1}-s}^{(1)}}{}^{i_{s+1}^{(2)}\cdots i_{r_{2}}^{(2)}}\tag{2}$$

Proposition 11. A contraction can be rewritten as a matrix product.

Proof. Using notation of (2), with the reshapings $t_1 \mapsto T_1$, $t_2 \mapsto T_2$ and $t_3 \mapsto T_3$ defined by grouping all covariant indices into a single index and all contravariant indices into another single index, we can rewrite

$$T_{1_{g_{i}(i_{1}^{(1)},\ldots,i_{r_{1}-s}^{(1)})}g_{k}(k_{1},\ldots,k_{s})}T_{2_{g_{k}(k_{1},\ldots,k_{s})}}g_{j}(i_{s+1}^{(2)},\ldots,i_{r_{2}}^{(2)})=T_{3_{g_{i}(i_{1}^{(1)},\ldots,i_{r_{1}-s}^{(1)})}}g_{j}(i_{s+1}^{(2)},\ldots,i_{r_{2}}^{(2)})$$

where g_i , g_k and g_j are bijections defined similarly as in (1).

Definition 12. Convolution

The *n*-dimensional convolution, denoted $*^n$, between $t_1 \in \mathbb{T}_1$ and $t_2 \in \mathbb{T}_2$, where \mathbb{T}_1 and \mathbb{T}_2 are of the same rank n such that $\forall p \in [\![1,n]\!], n_p^{(1)} \geq n_p^{(2)}$,

is defined as:

$$\begin{cases} t_1 *^n t_2 = t_3 \in \mathbb{T}_3 \text{ of shape } n_1^{(3)} \times \dots \times n_n^{(3)} \text{ where} \\ \forall p \in [1, n], n_p^{(3)} = n_p^{(1)} - n_p^{(2)} + 1 \\ t_3[i_1, \dots, i_n] = \sum_{k_1=1}^{n_1^{(2)}} \dots \sum_{k_n=1}^{n_n^{(2)}} t_1[i_1 + n_1^{(2)} - k_1, \dots, i_n + n_n^{(2)} - k_n] t_2[k_1, \dots, k_n] \end{cases}$$

Proposition 13. A convolution can be rewritten as a matrix product.

Proof. Let $t_1 *^n t_2 = t_3$ defined as previously with $\mathbb{T}_1 = \bigotimes_{k=1}^r \mathbb{V}_k^{(1)}$, $\mathbb{T}_2 =$

$$\bigotimes_{k=1}^{r} \mathbb{V}_{k}^{(2)}. \text{ Let } t_{1}' \in \bigotimes_{k=1}^{r} \mathbb{V}_{k}^{(1)} \otimes \bigotimes_{k=1}^{r} \mathbb{V}_{k}^{(2)} \text{ such that } t_{1}'[i_{1}, \ldots, i_{n}, k_{1}, \ldots, k_{n}] = t_{1}[i_{1} + n_{1}^{(2)} - k_{1}, \ldots, i_{n} + n_{n}^{(2)} - k_{n}], \text{ then}$$

$$t_3[i_1,\ldots,i_n] = \sum_{k_1=1}^{n_1^{(2)}} \cdots \sum_{k_n=1}^{n_n^{(2)}} t_1'[i_1,\ldots,i_n,k_1,\ldots,k_n] t_2[k_1,\ldots,k_n]$$

where we recognize a tensor contraction. Proposition 11 concludes. \Box

The two following operations are meant to further decrease the shape of the resulting output.

Definition 14. Strided convolution

The *n*-dimensional *strided* convolution, with strides $s = (s_1, s_2, \ldots, s_n)$, denoted $*_s^n$, between $t_1 \in \mathbb{T}_1$ and $t_2 \in \mathbb{T}_2$, where \mathbb{T}_1 and \mathbb{T}_2 are of the same rank n such that $\forall p \in [1, n], n_p^{(1)} \geq n_p^{(2)}$, is defined as:

$$\begin{cases} t_1 *_s^n t_2 = t_4 \in \mathbb{T}_4 \text{ of shape } n_1^{(4)} \times \dots \times n_n^{(4)} \text{ where} \\ \forall p \in [1, n], n_p^{(4)} = \lfloor \frac{n_p^{(1)} - n_p^{(2)} + 1}{s_p} \rfloor \\ t_4[i_1, \dots, i_n] = (t_1 *^n t_2)[(i_1 - 1)s_n + 1, \dots, (i_n - 1)s_n + 1] \end{cases}$$

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Remark. Informally, a strided convolution is defined as if it were a regular subsampling of a convolution. They match if s = (1, 1, ..., 1).

Definition 15. Pooling

Let a real-valued function f defined on all tensor spaces of any shape, e.g. the max or average function. An f-pooling operation is a mapping $t \mapsto t'$ such that each entry of t' is an image by f of a subtensor of t.

Remark. Usually, the set of subtensors that are reduced by f into entries of t' are defined by a regular partition of the entries of t.

1.2 Deep learning

In this manuscript, we adopt the point of view that a neural network is first a mathematical function, even though it derives its name from biological inspiration. That is, we won't discuss whether any of our works are biologically plausible or not, but we may provide biological interpretation when it happens.

In this section, we present a mathematical formalization and its biological interpretation. Then, we review a few important advances in the field before we finally present the most commonly used layers.

1.2.1 Neural networks

A feed-forward neural network could originally be formalized as a composite function chaining linear and non-linear functions (Rumelhart et al., 1985; LeCun et al., 1989; LeCun and Bengio, 1995). That was still the case in 2012 when important breakthroughs regenerated a surge of interest in the field (Hinton et al., 2012; Krizhevsky et al., 2012; Simonyan and Zisserman, 2014). However, in more recent years, more complex architectures have emerged (Szegedy et al., 2015; He et al., 2016a; Zoph and Le, 2016; Huang et al., 2017), such that the former formalization does not suffice. We provide a definition for the first kind of neural networks (Definition 16) and use it to present its related concepts. Then we give a more generic definition (Definition 20).

Note that in this manuscript, we only consider neural networks that are *feed-forward* (Zell, 1994; Wikipedia, 2018a), as opposed to *recurrent*.

We denote by I_f the domain of definition of a function f ("I" stands for "input") and by $O_f = f(I_f)$ its image ("O" stands for "output"), and we represent it as $I_f \xrightarrow{f} O_f$ or $f: I_f \to O_f$.

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Definition 16. Neural network (simply connected)

Let f be a function such that I_f and O_f are vector or tensor spaces.

f is a (simply connected) neural network function if there are a series of affine functions $(g_k)_{k=1,2,..,L}$ and a series of non-linear derivable univariate functions $(h_k)_{k=1,2,..,L}$ such that:

$$\begin{cases}
\forall k \in [1, L], f_k = h_k \circ g_k, \\
I_f = I_{f_1} \xrightarrow{f_1} O_{f_1} \cong I_{f_2} \xrightarrow{f_2} \dots \xrightarrow{f_L} O_{f_L} = O_f, \\
f = f_L \circ \dots \circ f_2 \circ f_1
\end{cases}$$

The couple (g_k, h_k) is called the k-th layer of the neural network. L is its depth. For $x \in I_f$, we denote by $x_k = f_k \circ ... \circ f_2 \circ f_1(x)$ the activations of the k-th layer. We denote by $\mathcal N$ the set of neural network functions.

Definition 17. Activation function

An *activation function* h is a real-valued univariate function that is non-linear and derivable, that is also defined by extension with the functional notation h(v)[i] = h(v[i]).

Definition 18. Layer

A layer is a couple $\mathcal{L} = (g, h) : I \to O$, where $g : I \to O$ is a linear function, and $h : O \to O$ is an activation function. It computes the function

$$y = h(g(x) + b)$$

where b is a constant called *bias*.

That is, in the simple formalization, a neural network is just a sequence of layers.

Remark. The bias augments the expressivity of the layers. For notational convenience, we may sometimes omit to write it down.

The most common activation function is the *rectified linear unit* (ReLU) (Glorot et al., 2011), used for its better practical performances and faster computation times. It implements the *rectifier* function $h: x \mapsto max(0, x)$ (with convention h'(0) = 0), as depicted on Figure 1.

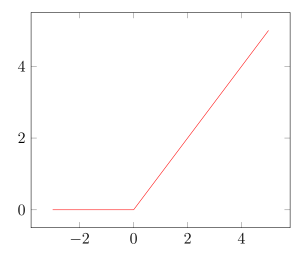


Figure 1: ReLU activation function

Examples

Let $f: x \to y$ be a neural network. For example, if f is used to classify its input x in one of c classes, then its output y would be a vector of dimension c, and each dimension corresponds to a class. The prediction of f for the class of x is the dimension of y where it has the bigger value. Typically, f is terminated by a softmax activation (Wikipedia, 2018b), so that values of the output y fall in the range [0,1], and so that y tends to have a dimension with a much bigger weight as to facilitates discrimination.

A neural network that comprises convolutional layers, *i.e.* layers s.t. g is expressed with a convolution, is called a Convolutional Neural Network (CNN). An old example is the LeNet-5 architecture (LeCun et al., 1989) as

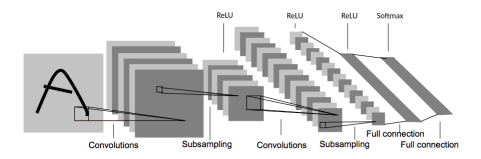


Figure 2: LeNet-5 (LeCun et al., 1989)

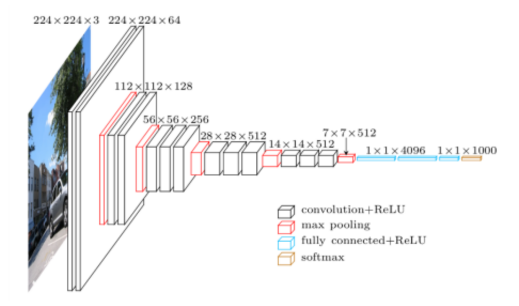


Figure 3: VGG-16 (Simonyan and Zisserman, 2014, figure from Cord, 2016)

depicted in Figure 2. It implements a function

$$f = h_4 \circ g_4 \circ \cdots \circ h_1 \circ g_1$$

where g_1 and g_2 are linear functions that applies 5x5 convolutions followed by subsampling, h_1 , h_2 and h_3 are ReLU activations, and h_4 is a softmax activation. It was originally applied to the task of handwritten digit classifications (for example for automatically reading postal ZIP codes).

Another example is the VGG architecture, a very deep CNN, and was state-of-the-art in image classification in 2014 (Simonyan and Zisserman). It is depicted on Figure 3. In more recent years, state-of-the-art architectures can no longer be described with a simple formalization.

The former neural networks are said to be *simply connected* because each layer only takes as input the output of the previous one. We give a more general definition after first defining branching operations.

Definition 19. Branching

A binary branching operation between two tensors, $x_{k_1} \bowtie x_{k_2}$, outputs, subject to shape compatibility, either their addition, either their concatenation along a rank, or their concatenation as a list.

Definition 20. Neural network (generic definition)

The set of *neural network* functions N is defined inductively as follows

- 1. $Id \in \mathcal{N}$
- 2. $f \in \mathcal{N} \land (g, h)$ is a layer $\land O_f \subset I_g \Rightarrow h \circ g \circ f \in \mathcal{N}$
- 3. for all shape compatible branching operations:

$$f_1, f_2, \dots, f_n \in \mathcal{N} \Rightarrow f_1 \bowtie f_2 \bowtie \dots \bowtie f_n \in \mathcal{N}$$

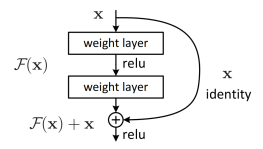


Figure 4: Module with a residual connection (He et al., 2016a)

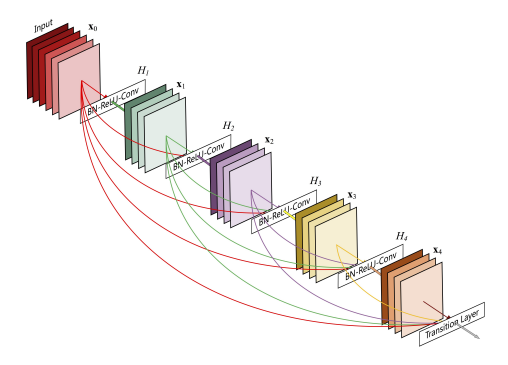


Figure 5: DenseNet (Huang et al., 2017)

Examples

The neural network proposed in (Szegedy et al., 2015), called *Inception*, use depth-wise concatenation of feature maps. Residual networks (ResNets, He et al., 2016a) make use of *residual connections*, also called *skip connections*, *i.e.* an activation that is used as input in a lower level is added to another activation at an upper level, as depicted on Figure 4. Densely connected networks (DenseNets, Huang et al., 2017) have their activations concatenated with all lower level activations. These neural networks had demonstrated state of the art performances on the imagenet classification challenge (Deng et al., 2009), outperforming simply connected neural networks. For example, DenseNet is depicted on Figure 5.

Remark. For layer indexing convenience, we still use the simple formalization in the subsequent subsections, even though the presentation would be similar with the generic formalization.

1.2.2 Interpretation

Until now, we have formally introduced a neural network as a mathematical function. As its name suggests, such function can be indeed interpreted from a connectivity perspective (LeCun, 1987).

Definition 21. Connectivity matrix

Let g a linear function. Without loss of generality subject to a flattening, let's suppose I_g and O_g are vector spaces. Then there exists a *connectivity matrix* W_g , such that:

$$\forall x \in I_g, g(x) = W_g x$$

We denote W_k the connectivity matrix of the k-th layer.

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Biological inspiration

A *neuron* is defined as a computational unit that is biologically inspired (McCulloch and Pitts, 1943). Each neuron is capable of:

- 1. receiving modulated signals from other neurons and aggregate them,
- 2. applying to the result an activation function,
- 3. passing the signal to other neurons.

That is to say, each domain $\{I_{f_k}\}$ and O_f can be interpreted as a layer of neurons, with one neuron for each dimension. The connectivity matrices $\{W_k\}$ describe the connections between each successive layers. A neuron is illustrated on Figure 6.

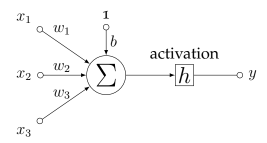


Figure 6: The McCulloch-Pitts model of a neuron

1.2.3 Training

Given an objective function F, training is the process of incrementally modifying a neural network f upon obtaining a better approximation of F. The most used training algorithms are based on gradient descent, as proposed in (Widrow and Hoff, 1960). These algorithms became popular since (Rumelhart et al., 1985). Informally, f is parameterized with initial weights that characterize its linear parts. These weights are modified step by step. At each step, a batch of samples are fed to the network, and their approximation errors sum to a loss. The weights of the network are updated in the opposite direction to their gradient with respect to that

loss. If the samples are shuffled and grouped in batches, this is called *Stochastic* gradient descent (SGD). Stochastic approximation (Robbins and Monro, 1985) tends to minimize effects of outliers on the training and is agnostic of the order in which the samples are fed.

Definition 22. Weights

Let consider the k-th layer of a neural network f. We define its weights as coordinates of a vector θ_k , called the *weight kernel*, such that:

$$\forall (i, j), \begin{cases} \exists p, W_k[i, j] := \theta_k[p] \\ \text{or } W_k[i, j] = 0 \end{cases}$$

A weight p that appears multiple times in W_k is said to be *shared*. Two parameters of W_k that share a same weight p are said to be *tied*. The number of weights of the k-th layer is $n_1^{(\theta_k)}$.

Learning

A loss function \mathcal{L} penalizes the output $x_L = f(x)$ relatively to the approximation error |f(x) - F(x)|. Gradient w.r.t. θ_k , denoted ∇_{θ_k} , is used to update the weights via an optimization algorithm based on gradient descent and a learning rate α , that is:

$$\theta_k^{(\text{new})} = \theta_k^{(\text{old})} - \alpha \cdot \vec{\nabla}_{\theta_k} \left(\mathcal{L} \left(x_L, \theta_k^{(\text{old})} \right) + \mathcal{R} \left(\theta_k^{(\text{old})} \right) \right)$$
(3)

where \mathcal{R} is a regularizer, and where α can be a scalar or a vector and \cdot can denote outer or coordinate-wise product, depending on the optimization algorithm that is used.

Linear complexity

Without loss of generality, we assume that the neural network is simply connected. Thanks to the chain rule, $\vec{\nabla}_{\theta_k}$ can be computed using gradients that are w.r.t. x_k , denoted $\vec{\nabla}_{x_k}$, which in turn can be computed using

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gradients w.r.t. outputs of the next layer k + 1, up to the gradients given on the output layer.

That is:

$$\vec{\nabla}_{\theta_k} = J_{\theta_k}(x_k) \vec{\nabla}_{x_k} \tag{4}$$

$$\vec{\nabla}_{x_k} = J_{x_k}(x_{k+1}) \vec{\nabla}_{x_{k+1}}$$

$$\vec{\nabla}_{x_{k+1}} = J_{x_{k+1}}(x_{k+2}) \vec{\nabla}_{x_{k+2}}$$

$$\vec{\nabla}_{x_{k+1}} = J_{x_{k-1}}(x_k) \vec{\nabla}_{x_k}$$

Obtaining,

$$\vec{\nabla}_{\theta_k} = J_{\theta_k}(x_k) (\prod_{p=k}^{L-1} J_{x_p}(x_{p+1})) \vec{\nabla}_{x_L}$$
(6)

where $J_{\text{wrt}}(.)$ are the respective Jacobians which can be determined with the layer's expressions and the $\{x_k\}$; and $\vec{\nabla}_{x_L}$ can be determined using \mathcal{L} , \mathcal{R} and x_L . This allows to compute the gradients with a complexity that is linear with the number of weights (only one computation of the activations), instead of being quadratic if it were done with the difference quotient expression of the derivatives (one more computation of the activations for each weight).

Backpropagation

We can remark that (5) rewrites as

$$\vec{\nabla}_{x_k} = J_{x_k}(x_{k+1}) \vec{\nabla}_{x_{k+1}}
= J_{x_k'}(h(x_k')) J_{x_k}(W_k x_k) \vec{\nabla}_{x_{k+1}}$$
(7)

where $x'_k = W_k x_k$, and these Jacobians can be expressed as:

$$J_{x'_{k}}(h(x'_{k}))[i,j] = \delta_{i}^{j}h'(x'_{k}[i])$$

$$J_{x'_{k}}(h(x'_{k})) = I h'(x'_{k})$$
(8)

$$J_{x_k}(W_k x_k) = W_k^T (9)$$

That means that we can write $\overrightarrow{\nabla}_{x_k} = (\widetilde{h}_k \circ \widetilde{g}_k)(\overrightarrow{\nabla}_{x_{k+1}})$ such that the connectivity matrix \widetilde{W}_k is obtained by transposition. This can be interpreted as gradient calculation being a *back-propagation* on the same neural network, in opposition of the *forward-propagation* done to compute the output.

1.2.4 Some historical advances

Universal approximation

Early researches have shown that neural networks with one level of depth can approximate any real-valued function defined on a compact subset of \mathbb{R}^n . This result was first proved for sigmoidal activations (Cybenko, 1989), and then it was shown it did not depend on the sigmoidal activations (Hornik et al., 1989; Hornik, 1991).

For example, this result brings theoretical justification that objective functions exists (even though it does not inform whether an algorithm to approach it exists or is efficient).

Computational difficulty

However, reaching such objective is a computationally difficult problem, which drove back interest from the field. Thanks to better hardware and to using better initialization schemes that speed up learning, researchers started to report more successes with deep neural networks (Hinton et al., 2006; Glorot and Bengio, 2010); see (Bengio, 2009) for a review of this period. It ultimately came to a surge of interest in the field after a significant breakthrough on the imagenet dataset (Deng et al., 2009) with deep

CNNs (Krizhevsky et al., 2012). The use of the fast ReLU activation function (Glorot et al., 2011) as well as leveraging graphical processing units with CUDA (Nickolls et al., 2008) were also key factors in overcoming this computational difficulty.

Adoption of ReLU activations

Historically, sigmoidal and tanh activations were mostly used (Cybenko, 1989; LeCun et al., 1989). However in recent practice, the ReLU activation (first introduced as the *positive part*, Jarrett et al., 2009), become the most used activation, as it was demonstrated to be faster and to obtain better results (Glorot et al., 2011). ReLU originated numerous variants *e.g. leaky rectified linear unit* (Maas et al., 2013), *parametric rectified linear unit* (PReLU, He et al., 2015), *exponential linear unit* (ELU, Clevert et al., 2015), *scaled exponential linear unit* (SELU, Klambauer et al., 2017), each one having particular advantages in some applications.

Avoiding overfitting

Neural networks, like any other machine learning technique, may overfit. That is, a model may behave well on the training set but fails to generalize well on unseen examples. The introduction of dropout (Srivastava et al., 2014) have helped models with more parameters to be less prone to overfitting, as dropout consists in hiding some parts of the training samples and their intermediate activations. Another good practice is to normalize per batch (Ioffe and Szegedy, 2015).

Expressivity and expressive efficiency

The study of the *expressivity* (also called *representational power*) of families of neural networks is the field that is interested in the range of functions that can be realized or approximated by this family (Håstad and Goldmann, 1991; Pascanu et al., 2013). In general, given a maximal error ϵ and an objective F, the more expressive is a family $N \subset \mathcal{N}$, the more likely it

is to contain an approximation $f \in N$ such that $d(f, F) < \epsilon$. However, if we consider the approximation $f_{min} \in N$ that have the lowest number of neurons, it is possible that f_{min} is still too large and may be unpractical. For this reason, expressivity is often studied along the related notion of expressive efficiency (Delalleau and Bengio, 2011; Cohen et al., 2018).

Rectifier neural networks

Of particular interest for the intuition is a result stating that a simply connected neural networks with only ReLU activations (a rectifier neural network) is a piecewise linear function (Pascanu et al., 2013; Montufar et al., 2014), and that conversely any piecewise linear function is also a rectifier neural network such that an upper bound of its depth is logarithmically related to the input dimension (Arora et al., 2018, th. 2.1.). Their expressive efficiency have also been demonstrated compared to neural networks using threshold or sigmoid activations (Pan and Srikumar, 2016).

Benefits of depth

Expressive efficiency analysis have demonstrated the benefits of depth, *i.e.* a shallow neural network would need an unfeasible large number of neurons to approximate the function of a deep neural network (*e.g.* Delalleau and Bengio, 2011; Bianchini and Scarselli, 2014; Poggio et al., 2015; Eldan and Shamir, 2016; Poole et al., 2016; Raghu et al., 2016; Cohen and Shashua, 2016; Mhaskar et al., 2016; Lin et al., 2017; Arora et al., 2018). This field seeks to give theoretical grounds to the practical observation that state-of-the-art architectures are getting deeper.

Benefits of branching operations

Recent works have provided rationales supporting benefits of using branching operations, thus giving justifications for architectures obtained with the generic formalization. In particular, (Cohen et al., 2018) have analyzed the impact of residual connections used in Wavenet-like architec-

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tures (Van Den Oord et al., 2016) in terms of expressive efficiency, using tools from the field of tensor analysis; (Orhan and Pitkow, 2018) have empirically demonstrated that residual connections can resolve some inefficiency problems inherent of fully-connected networks (dead activations, activations that are always equal, linearly dependent sets of activations).

1.2.5 Common layers

Definition 23. Connections

The set of *connections* of a layer (g, h), denoted C_g , is defined as:

$$C_g = \{(i, j), \exists p, W_g[i, j] := \theta_g[p]\}$$

We have $0 \le |C_g| \le n_1^{(W_g)} n_2^{(W_g)}$.

Definition 24. Dense layer

A dense layer (g,h) is a layer such that $|C_g| = n_1^{(W_g)} n_2^{(W_g)}$, i.e. all possible connections exist. The map $(i,j) \mapsto p$ is usually a bijection, meaning that there is no weight sharing.

A neural network made only of dense layers is called a Multi-Layer Perceptron (MLP, Hornik et al., 1989).

Definition 25. Partially connected layer

A partially connected layer (g,h) is a layer such that $|C_g| < n_1^{(W_g)} n_2^{(W_g)}$. A sparsely connected layer (g,h) is a layer such that $|C_g| \ll n_1^{(W_g)} n_2^{(W_g)}$.

Definition 26. Convolutional layer

A n-dimensional convolutional layer (g,h) is such that the weight kernel θ_g can be reshaped into a tensor w of rank n+2, and such that

$$\begin{cases} I_g \text{ and } O_g \text{ are tensor spaces of rank } n+1 \\ \forall x \in I_g, g(x) = (g(x)_q = \sum\limits_p x_p *^n w_{p,q})_{\forall q} \end{cases}$$

where p and q index slices along the last ranks.

A neural network that contains convolutional layers is called convolutional neural network (CNN).

Definition 27. Feature maps and input channels

The slices $g(x)_q$ are typically called *feature maps*, and the slices x_p are called *input channels*. Let's denote by $n_o = n_{n+1}^{(O_g)}$ and $n_i = n_{n+1}^{(I_g)}$ the number of feature maps and input channels. In other words, Definition 26 means that for each feature maps, a convolution layer computes n_i convolutions and sums them, computing a total if $n_i \times n_o$ convolutions.

Remark. Note that because they are simply summed, entries of two different input channels that have the same coordinates are assumed to share some sort of relationship. For instance on images, entries of each input channel (typically corresponding to Red, Green and Blue channels) that have the same coordinates share the same pixel location.

Benefits of convolutional layers

Comparatively with dense layers, convolution layers enjoy a significant decrease in the number of weights. For example, an input 2×2 convolution on images with 3-color input channels, would breed only 12 weights per feature maps, independently of the numbers of input neurons. On image datasets, their usage also breeds a significant boost in performance compared with dense layers (Krizhevsky et al., 2012), for they allow to take advantage of the topology of the inputs while dense layers do not (LeCun and Bengio, 1995). A more thorough comparison and explanation of their assets will be discussed in Section 2.1.3.

Decrease of spatial dimensions

Given a tensor input x, the n-dimensional convolutions between the inputs channels x_p and slices of a weight tensor $w_{p,q}$ would result in outputs

 y_q of shape $n_1^{(x)}-n_1^{(w)}+1\times\ldots\times n_n^{(x)}-n_n^{(w)}+1$. So, in order to preserve shapes, a padding operation must pad x with $n_1^{(w)}-1\times\ldots\times n_n^{(w)}-1$ zeros beforehand. For example, the padding function of the library *tensor-flow* (Abadi et al., 2015) pads each rank with a balanced number of zeros on the left and right indices (except if $n_t^{(w)}-1$ is odd then there is one more zero on the left).

Definition 28. Padding

A convolutional layer with *padding* (g,h) is such that g can be decomposed as $g=g_{\rm pad}\circ g'$, where g' is the linear part of a convolution layer as in Definition 26, and $g_{\rm pad}$ is an operation that pads zeros to its inputs such that g preserves tensor shapes.

Remark. One asset of padding operations is that they limit the possible loss of information on the borders of the subsequent convolutions, as well as preventing a decrease in size. Moreover, preserving shape is needed to build some neural network architectures, especially for ones with branching operations *e.g.* examples in Section 1.2.1. On the other hand, they increase memory and computational footprints.

Definition 29. Stride

A convolutional layer with *stride* is a convolutional layer that computes strided convolutions (with stride > 1) instead of convolutions.

Definition 30. Pooling

A layer with *pooling* (g,h) is such that g can be decomposed as $g=g'\circ g_{\text{pool}}$, where g_{pool} is a pooling operation.

Layers with stride or pooling downscale the signals that passes through the layer. These types of layers allows to compute features at a coarser level, giving the intuition that the deeper a layer is in the network, the more abstract is the information captured by the weights of the layer. From a practical point of view, they contribute to lower the computational complexity at deeper levels.

A simple result

In two dimensions, convolutional operations can be rewritten as a matrix-vector multiplication where the matrix is Toeplitz. We show below that it is still the case in n dimensions.

Proposition 31. Connectivity matrix of a convolution with padding

A convolutional layer with padding (g,h) is equivalently defined as its connectivity matrix W_g being a $n_i \times n_o$ block matrix such that its blocks are Toeplitz matrices, and where each block corresponds to a couple (p,q) of input channel p and feature map q.

Proof. Let's consider the slices indexed by p and q, and to simplify the notations, let's drop the subscripts p,q. We recall from Definition 12 that

$$\begin{split} y &= (x *^n w)[j_1, \dots, j_n] \\ &= \sum_{k_1 = 1}^{n_1^{(w)}} \dots \sum_{k_n = 1}^{n_n^{(w)}} x[j_1 + n_1^{(w)} - k_1, \dots, j_n + n_n^{(w)} - k_n] \, w[k_1, \dots, k_n] \\ &= \sum_{j_1 + n_1^{(w)} - 1} \dots \sum_{j_n + n_n^{(w)} - 1} x[i_1, \dots, i_n] \, w[j_1 + n_1^{(w)} - i_1, \dots, j_n + n_n^{(w)} - i_n] \\ &= \sum_{i_1 = j_1}^{n_1^{(x)}} \dots \sum_{i_n = 1}^{n_n^{(x)}} x[i_1, \dots, i_n] \, \widetilde{w}[i_1, j_1, \dots, i_n, j_n] \\ &= \sum_{i_1 = 1}^{w[j_1 + n_1^{(w)} - i_1, \dots, i_n, j_n] = \\ &= \begin{cases} w[j_1 + n_1^{(w)} - i_1, \dots, j_n + n_n^{(w)} - i_n] & \text{if } \forall t, 0 \leq i_t - j_t \leq n_t^{(w)} - 1 \\ 0 & \text{otherwise} \end{cases} \end{split}$$

Using Einstein summation convention as in (2) and permuting indices, we recognize the following tensor contraction

$$y_{j_1\cdots j_n} = x_{i_1\cdots i_n} \widetilde{w}^{i_1\cdots i_n}{}_{j_1\cdots j_n} \tag{10}$$

Following Proposition 11, we reshape (10) as a matrix product. To reshape $y \mapsto Y$, we use the row major order bijections g_j as in (1) defined onto $\{(j_1,\ldots,j_n), \forall t,1\leq j_t\leq n_t^{(y)}\}$. To reshape $x\mapsto X$, we use the same row major order bijection g_j , however defined on the indices that support non zero-padded values, so that zero-padded values are lost after reshaping. That is, we use a bijection g_i such that $g_i(i_1,i_2,\ldots,i_n)=g_j(i_1-o_1,i_2-o_2,\ldots,i_n-o_n)$ defined if and only if $\forall t,1+o_t\leq i_t\leq n_t^{(y)}$, where the $\{o_t\}$ are the starting offsets of the non zero-padded values. $\widetilde{w}\mapsto W$ is reshaped by using g_j for its covariant indices, and g_i for its contravariant indices. The entries lost by using g_i do not matter because they would have been nullified by the resulting matrix product. We remark that W is exactly the block (p,q) of W_g (and not of $W_{g'}$). Now let's prove that it is a Toeplitz matrix.

Thanks to the linearity of the expression (1) of g_j , by denoting $i'_t = i_t - o_t$, we obtain

$$g_i(i_1, i_2, \dots, i_n) - g_j(j_1, j_2, \dots, j_n) = g_j(i'_1 - j_1, i'_2 - j_2, \dots, i'_n - j_n)$$
 (11)

To simplify the notations, let's drop the arguments of g_i and g_j . By bijectivity of g_j , (11) tells us that $g_i - g_j$ remains constant if and only if $i'_t - j_t$ remains constant for all t. Recall that

$$W[g_i, g_j] = \begin{cases} w[j_1 + n_1^{(w)} - i_1', \dots, j_n + n_n^{(w)} - i_n'] & \text{if } \forall t, 0 \le i_t' - j_t \le n_t^{(w)} - 1\\ 0 & \text{otherwise} \end{cases}$$
(12)

Hence, on a diagonal of W, $g_i - g_j$ remaining constant means that $W[g_i, g_j]$ also remains constants. So W is a Toeplitz matrix.

The converse is also true as we used invertible functions in the index spaces through the proof. \Box

Remark. Note that the proof does not hold in case there is no padding. This is due to border effects when the index of the n^{th} rank resets in the definition of the row-major ordering function g_j that would be used. Indeed, under appropriate definitions, the matrices could be seen as almost Toeplitz.

This proposition provides an equivalent-characterization of convolutional layers by their connectivity matrix. Therefore, a first avenue to define convolutions on graph signals could be to define them with the connectivity matrix being as in this characterization. However, the Toeplitz property implies that the dimensions have a specific order, which is not possible when dimensions correspond to vertices of a graph. This is because permuting the order of the vertices wouldn't change the graph, but would change the connectivity matrix (which cannot be Toeplitz for every ordering).

1.3 Deep learning on graphs

Deep learning algorithms have been particularly successful for datasets of signals defined on regular domains such as images or time series. One key ingredient of their success is the use of convolutions. However, classical convolutions are only defined on Euclidean domains, so that extending deep learning on non-Euclidean domains is not straightforward. One way to represent a non-Euclidean domain is through a graph *i.e.* as points (the vertices) and relations between them (the edges). This field also comprises the study of deep learning on manifolds (*i.e.* locally Euclidean domains). The field of deep learning on graphs or manifolds have been called recently Geometric Deep Learning (Bronstein et al., 2017). In this manuscript, we are only interested in deep learning on graphs.

1.3.1 Graph and signals

We present the vocabulary, notation and conventions we will employ for graphs and signals.

Definition 32. Graph

A graph G is a couple of countable vertex and edge sets $\langle V, E \rangle$ s.t. $E \subset V^2$.

The terms *vertex* and *node* are used interchangeably. Additionally, we consider that a graph is always *simple i.e.* no two edges share the same set of vertices. Unless stated otherwise, a graph is undirected, *i.e.* (u,v) and (v,u) refer to the same edge. When it is not the case, it is called a *digraph*. We define the relation $u \sim v \Leftrightarrow (u,v) \in E$. We precise the graph if needed over the symbol $\stackrel{G}{\sim}$. For a digraph we use the symbol \rightarrow instead of \sim . A *walk* is a sequence $v_1 \sim \cdots \sim v_r$. It is said to be *simple* if its vertices are distinct, except possibly for the first and last. A graph is said to be *connected* if there exists a walk from any vertex to any other vertex. We define the *neighborhood* of a vertex as $\mathcal{N}_u = \{v \in V, u \sim v\}$. For digraphs, it is equal

to the union of the *in*- and *out*-neighborhoods. We only consider graphs without isolated vertex (a vertex with an empty neighborhood). We also only consider *weighted* graphs. That is, a graph $G = \langle V, E \rangle$ is associated with a weight mapping $w: V^2 \to \mathbb{R} + s.t.$ $w(u,v) = 0 \Leftrightarrow u \nsim v.$ If G is finite, its *adjacency matrix* $A \in \mathbb{R}^{V \times V}$ is defined *w.r.t.* to a vertex ordering $V = \{v_1, \ldots, v_n\}$ as $A[i,j] = w(v_i,v_j)$. Figure 7 illustrates an example of a graph and its adjacency matrix.

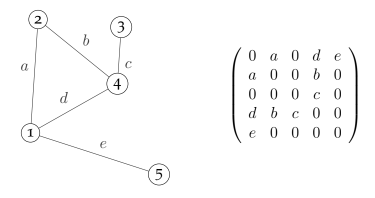


Figure 7: Example of a graph and its adjacency matrix

The *order* of G is equal to its number of vertices, possibly infinite. The *degree* of a vertex v is equal to the number of edges it is attached to. For digraphs the degree is the sum of the *in*- and *out*-degrees. The *degree* of G refers to its max degree. G is said to be *degree-regular* if all its vertices have the same degree. If it is finite, its *degree matrix* D (w.r.t. to a vertex ordering $V = \{v_1, \ldots, v_n\}$) is the diagonal matrix for which the diagonal entry corresponding to a vertex is the sum of the weights of the edges it is part of. Its *Laplacian matrix* L is the substraction L = D - A, which can be *normalized* $L = I - D^{-\frac{1}{2}}AD^{-\frac{1}{2}}$, *left-normalized* $L = I - D^{-1}A$, or *right-normalized* $L = I - AD^{-1}$. The same naming convention is used for normalized version of the adjacency matrix A. A subgraph of G induced by a subset $U \subset V$ is the graph with vertex and edge set restricted by G. The *complement* graph G shares the same vertex set but G vertex and G vertex G vertex

complete graph is such that there exists an edge between any two vertices.

Definition 33. Grid graph

Let a graph $G = \langle V, E \rangle$ such that the expression $u \sim v \Leftrightarrow ||u - v||_1 = 1$ makes sense. G can be called:

- a grid graph if $V = \mathbb{Z}^2$
- a finite grid graph if $\exists (n,m) \in \mathbb{Z}^2, V = [1,n] \times [1,m]$
- a circulant grid graph if $\exists (n,m) \in \mathbb{Z}^2, V = \mathbb{Z}/n\mathbb{Z} \times \mathbb{Z}/m\mathbb{Z}$

Definition 34. Bipartite graph

A graph is called *bipartite* if its vertex set is a disjoint union $V = V_1 \cup V_2$ s.t.

$$u \sim v \Rightarrow (u, v) \in V_1 \times V_2 \lor (u, v) \in V_2 \times V_1$$

If it is finite, its *bipartite-adjacency* matrix $A \in \mathbb{R}^{V_1 \times V_2}$ is a rectangular matrix defined w.r.t. to a vertex ordering $V_1 = \{u_1, \dots, u_n\}$, $V_2 = \{v_1, \dots, v_n\}$ and weight mapping w as $A[i, j] = w(u_i, v_j)$.

Definition 35. Signal

A *signal* on V, $s \in \mathcal{S}(V)$, is a function $s : V \to \mathbb{R}$. The *signal space* $\mathcal{S}(V)$ is the linear space of signals on V.

Remark. In particular, a vector space, and more generally a tensor space, are finite-dimensional signal spaces on any of their bases. Reciprocally, a signal space is a linear space which canonical basis is the signal space domain. Therefore, signals can be represented as vectors (or tensors) and then fed to neural networks.

A graph signal on a graph $G = \langle V, E \rangle$ is a signal on its vertex set V. We denote by S(G) or S(V) the graph signal space. G can be referred as the underlying structure of S(V). An entry of a signal s is an image by s of some $v \in V$ and we denote s[v]. If v is represented by an n-tuple, we can also write $s[v_1, v_2, \ldots, v_n]$. The support of a signal $s \in S(V)$ is the subset

 $\operatorname{supp}(s) \subset V$ on which $s \neq 0$. For spaces of signals that aren't real-valued, their codomain \mathbb{E} is precised in the subscript $\mathcal{S}_{\mathbb{E}}(V)$. The signal Id is the identity function. Given $v \in V$, the Dirac signal $\delta_v \in \mathcal{S}(V)$ is the signal valued as 1 on v and 0 everywhere else. The family of all Dirac signals spans $\mathcal{S}(V)$.

1.3.2 Learning tasks

There are two main tasks related to deep learning and graph signals.

Supervised classification of graph signals

This is the classical application of deep learning transposed to graph signals, rather than image or audio signals. It is the main task we will have in mind in the course of this manuscript. Given a graph $G = \langle V, E \rangle$ and an input signal $x \in \mathcal{S}(G)$ the goal is to classify x. If there are c possible classes, a neural network f outputs a vector y = f(x) of dimension c, and its dimension with the biggest weight determines the predicted class. Indeed, a standard MLP can be trained on a dataset of graph signals. However, an MLP wouldn't take the graph structure G into consideration. By similarity with CNNs that leverage the grid structure of images to achieve better performances than MLPs, a challenge is to define a neural network on graph signals that can leverage G. We review some models from the literature in Section 1.3.4 and in Section 1.3.5. We develop an algebraic understanding in Chapter 2 of why and how they should work, and also propose our own models and point of view in Chapter 3.

Semi-supervised classification of nodes

This task is in some way obtained from a transposed perspective of the previous one. Given a dataset of graph signals, represented as a matrix $X \in \mathbb{R}^{n \times N}$, where the rows represent the nodes, and the columns represent the signals, the goal is to classify the nodes. This amounts to classify

the rows, whereas the previous task amounts to classify the columns. As opposed to the previous one, this task is *transductive i.e.* every node data is available during training, including those from the validation and test set (but their labels are not), and it is *semi-supervised i.e.* some nodes have no label. This allows to learn on much more data than if we were restricted to labeled data. In this task, the edges connect learning samples, however in the previous one, the edges were connecting features of learning samples. This is this edge relationship between learning samples that renders the semi-supervised approach possible. This task have received much more attention than the previous one in the recent literature. We explain why in Section 1.3.4.

Other learning tasks

In this manuscript, we are less interested in other deep learning tasks related to graphs, so we briefly discuss them here. One is supervised classification of graphs, which is different than classifying graph signals. Examples include (Niepert et al., 2016; Tixier et al., 2017; Nikolentzos et al., 2017; Bai et al., 2018). Another related interesting task is called representation learning of nodes, which tackles the challenge to learn a linear representation of nodes. A common approach, derived from word2vec (Mikolov et al., 2013b; Mikolov et al., 2013a), is called node2vec (Grover and Leskovec, 2016), and was later improved in graphSAGE (Hamilton et al., 2017a). A review on this subject is done by Hamilton et al., 2017b. A thorough survey on representation learning for networks is given in (Zhang et al., 2017).

1.3.3 Datasets

We present here the standard datasets that we will use in this manuscript.

Images

Images are signals on grid graphs. Therefore they constitute a first test for models that are designed to be able to classify graph signals. A second step is to test on scrambled versions of image datasets, *i.e.* a random permutation of the pixels is fixed and the tested models are fed with images whose pixels have been shuffled according to this permutation. Therefore the domains of the scrambled input signals are not grid graphs.

- MNIST (LeCun et al., 1998) is a dataset of handwritten digits of size 28x28. It contains 10 classes and is split between 50'000 samples for training and 10'000 samples for testing.
- CIFAR-10 (Krizhevsky, 2009) is a dataset of tiny pictures of size 32x32. It contains 10 classes and is split between 50'000 samples for training and 10'000 samples for testing.
- Scrambled MNIST is the scrambled version of MNIST. A graph based on nearest neighbors from the pixel covariances is used to represent each scrambled sample as a graph signal.
- Scrambled CIFAR-10 is analogous.

Functional magnetic resonance imaging (fMRI)

fMRI samples can be represented by graph signals. The graphs are resembling grid graphs to some extent since they are embedded in an Euclidean space.

 The PINES dataset consists of fMRI scans on 182 subjects, during an emotional picture rating task (Chang et al., 2015). In (Lassance et al., 2018), we fetched individual first-level statistical maps (beta images) for the minimal and maximal ratings from https://neurovault.org/collections/1964/, to generate the dataset. Full brain data was masked on the MNI template and resampled to a 16mm cubic grid, in order to reduce dimensionality of the dataset while keeping a regular geometrical structure to infer the graph. Final volumes used for classification contain 369 signals for each subject and rating.

Text documents

Text documents can be represented as graph signals. Words are the vertices, and the value of a signal at a given vertex corresponds to a normalized occurrence of the corresponding word. Edges connect nearest neighbors in some metric space.

• 20NEWS (Joachims, 1996) is a dataset of text documents. It contains 20 classes and is split between 11'314 samples for training and 7'532 samples for testing. Each document is represented by a bag-of-word. In the version used by Defferrard et al., 2016, that we consider, documents are treated as signals on a graph of 10'000 vertices which represent the 10'000 most common words (the other words are not used). Edges are drawn from each vertex to their 16 nearest neighbors in the cosine similarity metric space.

Citation networks

In a citation network (*i.e.* a graph of citations), nodes are bag-of-word documents and edges represent citations. Models are tested on a citation network to the task of semi-supervised classification of the nodes. We use three standard datasets: Cora, Citeseer and Pubmed (Sen et al., 2008), for which we follow the experimental settings of Yang et al., 2016, and the dataset split from Kipf and Welling, 2016. In particular, there are only 20 training samples per class, but they are connected to other samples.

- Cora is a dataset of 2'708 nodes of dimension 1'433 with 5'429 edges. It contains 7 classes and consists of 140 samples for training, 500 samples for validation, 1'000 samples for testing, and 1'068 unlabeled samples.
- Citeseer is a dataset of 3'327 nodes of dimension 3'703 with 4'732 edges. It contains 6 classes and consists of 120 samples for training, 500 samples for validation, 1000 samples for testing, and 1'707 unlabeled samples.
- Pubmed is a dataset of 19'717 nodes of dimension 500 with 44'338 edges. It contains 3 classes and consists of 60 samples for training, 500 samples for validation, 1000 samples for testing, and 18'157 unlabeled samples.

1.3.4 Spectral methods

Spectral methods are based on spectral graph theory (Chung, 1996) which aims at characterizing structural properties of a graph $G = \langle V, E \rangle$ through the eigenvalues of the Laplacian matrix L. In particular, since it is Hermitian, it admits a complete set of normalized eigenvectors. By fixing a normalized eigenvector basis ordered in the rows of U (by ascending eigenvalues), U is used to define the *Graph Fourier Transform* (GFT) of a signal $s \in \mathcal{S}(G)$ (Shuman et al., 2013), and the conjugate-transpose U^* defines the inverse GFT. We write

$$\widehat{s} = Us \tag{13}$$

$$\widetilde{s} = U^* s \tag{14}$$

Remark. The GFT extends the notion of Discrete Fourier Transform (DFT) to general graphs, since that for circulant grid graphs U can be the DFT matrix.

By analogy with the convolution theorem, a convolution can be defined as pointwise multiplication, denoted \cdot , in the spectral domain of the graph (Hammond et al., 2011). For $s, g \in \mathcal{S}(G)$, we have:

$$s * g = \widetilde{\widehat{s} \cdot \widehat{g}} \tag{15}$$

This expression can be used to define convolutional layers and spectral CNNs on graphs. However, Bruna et al., 2013 pointed out that (15) would generate filters with $\mathcal{O}(n)$ weights, where n is the order of G. So they proposed to learn filters θ with only $\mathcal{O}(1)$ weights and then to smoothly interpolate the remaining weights as $g = K\theta$, where K is a linear smoother matrix. They motivate their construction by the fact that smooth multipliers in the spectral domain should simulate local operations in the vertex domain. To elaborate a bit on this, note that we have:

$$Ls[u] = \sum_{v \in V} w(u, v)(s[u] - s[v])$$
(16)

And so,

$$s^{T}Ls = \sum_{u \in V} \sum_{v \in V} w(u, v)s[u](s[u] - s[v])$$

$$= \frac{1}{2} \sum_{u \in V} \sum_{v \in V} w(u, v)s[u](s[u] - s[v]) + \frac{1}{2} \sum_{v \in V} \sum_{u \in V} w(v, u)s[v](s[v] - s[u])$$

$$= \sum_{u \in V} \sum_{v \in V} \frac{w(u, v)}{2} (s[u] - s[v])^{2}$$
(17)

That is, $s^T L s$ is some sort of measure of *smoothness* of the signal s, penalized by the weights w. The bigger is w(u, v), the closest s(u) and s(v) must

be to lower the smoothness (17). Since L is symmetric, its eigenvalues are non-negative real numbers, and U diagonalizes L as $\Lambda = ULU^*$. Denote $(\lambda_i)_i$ the eigenvalues, the smoothness measure rewrites:

$$s^{T}Ls = \hat{s}^{*}\Lambda \hat{s} = \sum_{i=1}^{n} \lambda_{i} \hat{s}[i]^{2}$$
(18)

Therefore, as they pointed out, smoothness of s can be read off the coordinates of \hat{s} , like for the DFT. Moreover, spectral multipliers modulate its smoothness, and decay in the spectral domain is related to smoothness in the vertex domain. But contrary to their conjecture, smoothness in the spectral domain is not necessary related to decay is the vertex domain (and so to some form of locality). For instance, since the Laplacian L^C of the complement graph G^C commutes with L, it can share the same eigenvector basis U, and thus define the same GFT, but their notion of locality in the vertex domain are opposed. Another drawback is that this method requires computing the GFT which complexity is at least $\mathcal{O}(n^2)$ as there is no equivalent of the Fast Fourier Transform (FFT) on graphs, so the authors suggest to use a lower number of eigenvectors d < n from the Laplacian eigenbasis.

Then, Defferrard et al., 2016 remedy to these issues by proposing an approximate formulation based on the Chebychev polynomials, denoted by $(T_i)_i$, where i is the polynomial order. That is, their proposed approximate filters are in the form

$$g_{\theta}(L) = \sum_{i=0}^{k} \theta[i] T_i(\widetilde{L})$$
(19)

where $\widetilde{L} = \frac{\lambda_{\max}}{2} L - I_n$ is the scaled normalized Laplacian with eigenvalues lying in the range [-1,1]. $g_{\theta}(L)$ are spectral multipliers since we have:

$$g_{\theta}(L)s = g_{\theta}(U^*\Lambda U)s = U^*g_{\theta}(\Lambda)Us$$

$$= \widetilde{g_{\theta}(\Lambda)} \mathbf{1} * s \tag{20}$$

These filters enjoy locality properties, they contain $\mathcal{O}(1)$ weights, and their complexity is $\mathcal{O}(n)$ when rows of L are sparse. The use of truncated Chebychev expansion (Hammond et al., 2011) ensures that in theory any set of spectral multipliers can be approximated. Also, since they are Laplacian polynomials, some authors would argue that these filters are transferable from one graph to another. From a combinatorial point of view this is true. However there is no reason that spectral multipliers from a spectral domain make sense in another one, and there are no experiment in the literature to support the hypothesis. On the other hand, (Yi et al., 2016) (who do not use polynomial filters) fix a canonical spectral base in order to synchronize every spectral domains. Their idea is to learn a warping from any eigenbasis to the canonical one, prior to performing spectral multiplication, in the manner of spatial transformer networks (STN, Jaderberg et al., 2015).

However, it is hard to evaluate if a model performs well on the task of supervised classification of graph signals, because there are not much known datasets in the literature for which the given graph domain holds enough information.

For example, Defferrard et al. built a graph signal dataset from the text categorization dataset 20NEWS (Joachims, 1996, see Section 1.3.3). However, their model (ChebNet32) fails to surpass Multinomial Naive Bayes (MNB). Moreover, even though they report that their model beats MLPs, but through experiments we noticed the contrary. In results we report in Table 1, we see that a lighter MLP, composed of a single Fully-Connected (FC) layer with ReLU and 20% dropout outperforms ChebNet32. We replicated their preprocessing phase from the code on their official repository and averaged our results on 100 runs of 10 epochs¹.

¹A few epochs are enough since models seem to overfit fast on this dataset.

MNB	FC2500	FC2500-FC500	ChebNet32	FC500	
68.51^{a}	64.64^{a}	65.76^{a}	68.26^{a}	71.96 ± 0.15^b	

^a As reported in Defferrard et al., 2016

Table 1: Accuracies (in %) on 20NEWS

Despite the significant theoretical contribution, this negative result stresses out the importance of the graph used in practice to support the convolution, a point that they also discussed. Henaff et al., 2015, proposed supervised graph estimation techniques, but a better graph signal dataset would be one that come with an already suitable graph, that of current literature is still lacking.

On the other hand, attention in the domain has shifted toward the task of semi-supervised classification of nodes, where good datasets are not lacking. For example, Levie et al., 2017, mainly demonstrate the usefulness of their model on these type of tasks. They define polynomial filters, for which Chebychev filters are a special case, that are capable to specialize in narrow bands of frequency in the spectral domain.

Another spectral avenue consists in using wavelets defined in the graph spectral domain (Hammond et al., 2011), in order to build a scattering network (Bruna and Mallat, 2013; Chen et al., 2014). This idea have been exploited recently by Zou and Lerman, 2018, then by Gama et al., 2018.

^b From our experiments.

1.3.5 Vertex-domain methods

As their name suggests, vertex-domain methods operates directly on the vertices of the graph. Convolution can be modeled as a function f of the kernel weights θ and neighboring vertices (contained in the local receptive field $\mathcal{R}(v)$), usually based on dot products. That is

$$y[v] = f_{\theta}\left(\left\{u \in \mathcal{R}(v)\right\}\right) \tag{21}$$

As such, it retains the property of being localized and of sharing weights in some way. But there remains the need to specify how the shared weights are allocated in this local receptive field (Vialatte et al., 2016). This allocation can depend on e.g. an arbitrary order (Niepert et al., 2016), on a diffusion process (Atwood and Towsley, 2016), on a function of both vertices and their neighbors (Monti et al., 2016; Simonovsky and Komodakis, 2017), on a random walk (Hechtlinger et al., 2017), on another learned kernel (Vialatte et al., 2017), on an attention mechanism (Velickovic et al., 2017; Lee et al., 2018), on pattern identification (Sankar et al., 2017), or on translation identification (Pasdeloup et al., 2017a). All these methods differ in the function f, but in the end, their definition highly overlap. That is why some authors have proposed unified frameworks (Gilmer et al., 2017). The representation we present in Chapter 3 is also unifying in that sense (see Section 3.2.1).

In particular, Kipf and Welling, 2016, were first to transpose ChebNet to the task of semi-supervised node classification. Chebychev filters (19) then take a form that is interpretable in the vertex domain, which is

$$Y = \sum_{i=0}^{k} T_i(\widetilde{L}) X \Theta \tag{22}$$

where $X \in \mathbb{R}^{n \times N}$, $\Theta \in \mathbb{R}^{N \times M}$, n is the number of nodes, N is the number

of input channels (features per node), and M is the number of output feature maps. On the left, powers of \widetilde{L} diffuse the graph signal X to share node information. On the right, Θ maps the diffused signals to another representation. So in essence, this formulation is more a vertex-domain method. They found that the best performing filters were expressed in a simplified form

$$Y = \widetilde{A}X\Theta \tag{23}$$

where \widetilde{A} is the normalized adjacency matrix of the graph to which self-loops are added. They call the architecture composed with these filters a Graph Convolution Network (GCN). Similarly, $\widetilde{A}X$ shares node information via the edges and Θ makes the model learns. This formulation attracted a lot of research attention and was, in particular, extended with attention mechanism (no pun intended), inspired from the field of neural machine translation (Bahdanau et al., 2014). A review is done by Lee et al., 2018.

For example, Velickovic et al., 2017, propose a model that learns attention in a local receptive field. They call it Graph ATtention network (GAT). The attention mechanism is parameterized by a neural network a, containing a single FC layer (g, LeakyReLU), which takes as input a couple of neighboring nodes (i,j) and outputs a scalar $\alpha_{i,j}$. The attention that i deserves to j is:

$$\alpha_{i,j} = \operatorname{softmax}_{j} \left(a(X[i,:] \Theta \parallel X[j,:] \Theta) \right) \tag{24}$$

where \parallel denotes concatenation. In a sense, a learns which input feature maps are most useful to describe the attention a node should deserve to another. The forward propagation is done similarly than (23), but with a matrix A_k filled with the attention coefficients $\alpha_{i,j}$ instead of \widetilde{A} . They also propose that the model learns multiple attention heads, so that a GAT

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layer amounts to:

$$Y = \prod_{k=1}^{K} A_k X \Theta_k \tag{25}$$

or
$$Y = \frac{1}{K} \sum_{k=1}^{K} A_k X \Theta_k$$
 (26)

Another method called Topology Adaptive GCN (TAGCN, Du et al., 2017) uses a convolution filter borrowed from graph signal processing literature (Sandryhaila and Moura, 2013), which is defined in the vertex domain as:

$$Y[:,f] = \sum_{c=1}^{N} \left(\sum_{k=1}^{K} \Theta_k[c,f] \, \widetilde{A}^k \right) X[:,c]$$
 (27)

It can be rewritten as:

$$Y = \sum_{k=1}^{K} \widetilde{A}^k X \Theta_k \tag{28}$$

Each successive powers of the normalized adjacency matrix \widetilde{A} allows for considering wider neighborhoods.

Other works extending or resembling GCN are numerous in recent days (e.g. Niepert and Garcia-Duran, 2018). We do not cover them since that their novelty compared to GCN is limited.

In the next chapter, we study how to characterize a convolution of graph signals in the vertex domain.

Chapter 2

Convolution of graph signals

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Chapter overview

Defining a convolution of signals over graph domains is a challenging problem. If the graph is not a grid graph, there exists no natural extension of the Euclidean convolution. In Section 2.1, we analyze the reasons why the Euclidean convolution operator is useful in deep learning. In particular, we recall a classical characterization: that convolution operators are exactly the class of linear functions that are equivariant to translations (Theorem 41). Therefore, we then search for domains onto which a convolution with these properties can be naturally obtained. This leads us to put our interest on representation theory and convolutions defined on groups. Since the Euclidean convolution is just a particular case of the group convolution, it makes perfect sense to steer our construction in this direction. In Section 2.2, we seek to transfer the definition of the group convolution onto the vertex domain, through its symmetric group. To obtain the wanted characterization, we will see that we need to base our construction on actions of groups, rather than on their elements. We manage to obtain it should we fix an equivariant mapping between the active group and the vertex domain (Theorem 54). Then, we propose a mixed formulation of this convolution as a binary operation between a signal defined on the vertex domain and a signal defined on the corresponding group, for which we demonstrate that the characterization also holds under abelianity (Corrolary 61). In Section 2.3, we introduce the role of the edge set and see how it influences the construction. In particular, we define a notion of edge constraint and a notion of locality preservation. For both, we obtain a characterization of graphs that admit a natural construction of convolutions with this property (Theorem 65 and Theorem 73). We analyze the notions of locality and weight sharing in this construction, and give a formulation for small kernels. At this point, with the obtained theorems we are able to describe convolutions on any

graphs, as convolutions on appropriate subgraphs. Then, in Section 2.4, we relax some aspect of the construction to better adapt it to general graphs. We explain why a construction based on groups is less interesting for some graphs, and introduce the notion of groupoid. We extend the previous construction with groupoids of partial transformations, and prove that under a mild condition the characterization by equivariance is preserved (Theorem 83). Finally, we extend it another time with another type of groupoid, that we call path groupoids. Path groupoids allow to tackle the more general case, and for them we obtain the characterization should we fix a way to traverse the vertex set using a subset of the edge set (Theorem 90), but at the price of allowing more degenerated cases. We summarize our constructions in a conclusive Section 2.5. The result of this chapter is the obtention of a set of general expressions and theorems that describe convolutions of graph signals.

2.1 Analysis of the classical convolution

In this section, we are exposing a few properties of the classical convolution that a generalization to graphs would likely try to preserve. For now let's consider a graph G agnostically of its edges i.e. $G \cong V$ is just the set of its vertices.

2.1.1 Properties of the convolution

Consider an edge-less grid graph *i.e.* $G \cong \mathbb{Z}^2$. By restriction to compactly supported signals, this case encompass the case of images.

Definition 36. Convolution on $\mathcal{S}(\mathbb{Z}^2)$

Recall that the (discrete) convolution between two signals s_1 and s_2 over \mathbb{Z}^2 is a binary operation in $\mathcal{S}(\mathbb{Z}^2)$ defined as:

$$\forall (a,b) \in \mathbb{Z}^2, (s_1 * s_2)[a,b] = \sum_{i} \sum_{j} s_1[i,j] \, s_2[a-i,b-j]$$

Definition 37. Convolution operator

A *convolution operator* is a function of the form $f_w: x \mapsto x*w$, where x and w are signals of domains for which the convolution * is defined. When * is not commutative, we differentiate the *right* operator $x \mapsto x*w$ from the *left* one $x \mapsto w*x$.

The following properties of the convolution on \mathbb{Z}^2 are of particular interest for our study.

Linearity

Operators produced by the convolution are linear. So they can be used as linear parts of layers of neural networks.

Locality and weight sharing

When w is compactly supported on K, an impulse response $f_w(x)[a,b]$ amounts to a weighted aggregation of entries of x in a neighborhood of (a,b), called the *Local Receptive Field* (LRF). The weight kernel w used for the aggregation is fixed w.r.t. (a,b), so that we say that the weights are shared.

Commutativity

This convolution is commutative. However, it won't necessarily be the case on other domains.

Equivariance to translations

Convolution operators are equivariant to translations. Below, we show that the converse of this result also holds with Theorem 41.

2.1.2 Characterization on grid graphs

We first define what a transformation of a domain is, and how it can be extended to signals defined on this domain.

Definition 38. Transformation

A transformation $f:V\to V$ is a function with same domain and codomain. The set of transformations is denoted $\Phi(V)$. The set of invertible transformations is denoted $\Phi^*(V)\subset\Phi(V)$. If V is a linear space, the set of its linear transformation is denoted $\mathcal{L}(V)$.

Remark. Note that $\Phi^*(V)$ forms what is usually called the symmetric group of V.

 $\Phi^*(V)$ can *move* signals of $\mathcal{S}(V)$ by linear extension of its group action, as we explain with the lemma that follows.

Lemma 39. Extension of $\Phi^*(V)$ to $\mathcal{L}(\mathcal{S}(V))$

An invertible transformation $f \in \Phi^*(V)$ can be extended linearly to the signal space S(V), and we have:

$$\forall s \in \mathcal{S}(V), \forall v \in V, f(s)[v] = s[f^{-1}(v)]$$

Proof. Let $s \in \mathcal{S}(V)$, $f \in \Phi^*(V)$, $L_f \in \mathcal{L}(\mathcal{S}(V))$ s.t. $\forall v \in V, L_f(\delta_v) = \delta_{f(v)}$. Then, by linear decomposition of s on the family of Dirac signals, we have:

$$L_f(s) = \sum_{v \in V} s[v] L_f(\delta_v)$$

$$= \sum_{v \in V} s[v] \delta_{f(v)}$$
 So, $\forall v \in V, L_f(s)[v] = s[f^{-1}(v)]$

 L_f extends f to S(V). When there is no ambiguity, we use the same symbol for f(.) and $L_f(.)$.

For translations, we use the following formalism.

Definition 40. Translation on $\mathcal{S}(\mathbb{Z}^2)$

A translation on \mathbb{Z}^2 is defined as a transformation $t \in \Phi^*(\mathbb{Z}^2)$ such that

$$\exists (a,b) \in \mathbb{Z}^2, \forall (x,y) \in \mathbb{Z}^2, t(x,y) = (x+a,y+b)$$

By Lemma 39, it also acts on $\mathcal{S}(\mathbb{Z}^2)$ with the notation $t_{a,b}$ *i.e.*

$$\forall s \in \mathcal{S}(\mathbb{Z}^2), \forall (x, y) \in \mathbb{Z}^2, t_{a,b}(s)[x, y] = s[x - a, y - b]$$

The next theorem fully characterizes convolution operators with their translational equivariance property. This can be seen as a discretization

of a classic result from the theory of distributions (Schwartz, 1957).

Theorem 41. Characterization of convolution operators on $\mathcal{S}(\mathbb{Z}^2)$

On real-valued signals over \mathbb{Z}^2 , the class of linear transformations that are equivariant to translations is exactly the class of convolutive operations *i.e.*

$$\exists w \in \mathcal{S}(\mathbb{Z}^2), f = . * w \Leftrightarrow \begin{cases} f \in \mathcal{L}(\mathcal{S}(\mathbb{Z}^2)) \\ \forall t \in \mathcal{T}(\mathcal{S}(\mathbb{Z}^2)), f \circ t = t \circ f \end{cases}$$

Proof. The result from left to right is a direct consequence of the definitions:

$$\forall s \in \mathcal{S}(\mathbb{Z}^2), \forall s' \in \mathcal{S}(\mathbb{Z}^2), \forall (\alpha, \beta) \in \mathbb{R}^2, \forall (a, b) \in \mathbb{Z}^2,$$

$$f_w(\alpha s + \beta s')[a, b] = \sum_i \sum_j (\alpha s + \beta s')[i, j] w[a - i, b - j]$$

$$= \alpha f_w(s)[a, b] + \beta f_w(s')[a, b] \qquad \text{(linearity)}$$

$$\forall s \in \mathcal{S}(\mathbb{Z}^2), \forall (x, y) \in \mathbb{Z}^2, \forall (a, b) \in \mathbb{Z}^2,$$

$$f_w \circ t_{x,y}(s)[a, b] = \sum_i \sum_j t_{x,y}(s)[i, j] w[a - i, b - j]$$

$$= \sum_i \sum_j s[i - x, j - y] w[a - i, b - j]$$

$$= \sum_{i'} \sum_{j'} s[i', j'] w[a - i' - x, b - j' - y] \qquad (29)$$

$$= f_w(s)[a - x, b - y]$$

$$= t_{x,y} \circ f_w(s)[a, b] \qquad \text{(equivariance)}$$

Now let's prove the result from right to left.

Let $f \in \mathcal{L}(\mathcal{S}(\mathbb{Z}^2))$, $s \in \mathcal{S}(\mathbb{Z}^2)$ and we suppose that f commutes with translations. By linear decomposition of s on the family of Dirac signals,

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we have:

$$f(s) = \sum_{i} \sum_{j} s[i, j] f(\delta_{i,j})$$

$$= \sum_{i} \sum_{j} s[i, j] f \circ t_{i,j}(\delta_{0,0})$$

$$= \sum_{i} \sum_{j} s[i, j] t_{i,j} \circ f(\delta_{0,0})$$

By denoting $w = f(\delta_{0,0}) \in \mathcal{S}(\mathbb{Z}^2)$, we obtain:

$$\forall (a,b) \in \mathbb{Z}^2, f(s)[a,b] = \sum_{i} \sum_{j} s[i,j] t_{i,j}(w)[a,b]$$

$$= \sum_{i} \sum_{j} s[i,j] w[a-i,b-j]$$
i.e. $f(s) = s * w$ (30)

2.1.3 Usefulness of convolutions in deep learning

Equivariance property of CNNs

In deep learning, an important argument in favor of CNNs is that convolutional layers are equivariant to translations. Intuitively, that means that a detail of an object in an image should produce the same features independently of its position in the image.

Lossless superiority of CNNs over MLPs

The converse result, as a consequence of Theorem 41, is never mentioned in deep learning literature. However it is also a strong one. For example, let's consider a linear function that is equivariant to translations. Thanks to the converse result, we know that this function is a convolution operator parameterized by a weight vector w, f_w : . * w. If the domain is

compactly supported, as in the case of images, we can break down the information of w in a finite number n_q of kernels w_q with small compact supports of same size (for instance of size 2×2), such that we have $f_w = \sum_{q \in \{1,2,\dots,n_q\}} f_{w_q}$. The convolution operators f_{w_q} are all in the search space of 2×2 convolutional layers. In other words, every translational equivariant linear function can have its information parameterized by these layers. So that means that the reduction of parameters from an MLP to a CNN is done without loss of expressivity (provided the objective function is known to bear this property). Besides, it also helps the training to search in a much more confined space. For example, on CIFAR-10 (see description in Section 1.3.3), CNNs reportedly attain up to 2.31% error on classification (Yamada et al., 2018), while MLPs plateaued at 21.38% (Lin et al., 2015). Intuitively, the reason for this success is *simplification by* symmetry: the supposed translational equivariance of the objective function is a symmetry that is exploited by the convolution layer to simplify its input.

Methodology for extending to general graphs

Hence, in our construction, we will try to preserve the characterization from Theorem 41 since it is mostly the reason why they are successful in deep learning. Note that other useful properties are also a consequence of this characterization. For example, the fact that convolutional layers have less parameters than a dense layer is also a consequence of this characterization.

2.2 Construction on the vertex set

As Theorem 41 is a complete characterization of convolutions, it can be used to define them i.e. convolution operators can be constructed as the set of linear transformations that are equivariant to translations. However, in the general case where G is not a grid graph, translations are not defined, so that construction needs to be generalized beyond translational equivariances.

In mathematics, convolutions are more generally defined for functions defined over a group structure. The classical convolution that is used in deep learning is just a narrow case where the domain group is an Euclidean space. Therefore, constructing a convolution on graphs should start from the more general definition of convolution on groups rather than convolution on Euclidean domains.

Our construction is motivated by the following questions:

- Does the equivariance property holds? Does the characterization from Theorem 41 still holds?
- Is it possible to extend the construction on non-group domains, or at least on mixed domains? (*i.e.* one signal is defined over a set, and the other is defined over its transformations).
- Can a group domain draw an underlying graph structure? Is the group convolution naturally defined on this class of graphs?
- Can we characterize graphs accepting our construction?

In this section, we first aim at transferring the group convolution onto the vertex set. Then, in Section 2.3, we will see the implications of considering the edge set in the process.

2.2.1 Preliminaries

We first recall preliminary notions about groups and group convolutions.

Definition 42. Group

A group Γ is a set equipped with a closed, associative and invertible composition law that admits a unique left-right identity element e.

When the law is commutative, the group is said to be *abelian*. The group convolution extends the notion of the classical discrete convolution and is defined as follows.

Definition 43. Group convolution

Let a group Γ , the *group convolution* between two signals s_1 and $s_2 \in \mathcal{S}(\Gamma)$ is defined as:

$$\forall h \in \Gamma, (s_1 *_{\Gamma} s_2)[h] = \sum_{g \in \Gamma} s_1[g] s_2[g^{-1}h]$$

provided at least one of the signals has finite support if Γ is not finite.

We call *left multiplication* the transformation $L_g: h \mapsto gh$, and *right multiplication* the transformation $R_g: h \mapsto hg$. Then, thanks to Lemma 39, the group convolution can be rewritten with the functional formulation:

$$s_1 *_{\Gamma} s_2 = \sum_{g \in \Gamma} s_1[g] L_g(s_2)$$
 (31)

$$= \sum_{g \in \Gamma} s_2[g] R_g(s_1) \tag{32}$$

which would then be commutative if, and only if, Γ were abelian. The group convolution preserves the characterization from Theorem 41.

Theorem 44. Characterization of group convolution operators Let a group Γ , let $f \in \mathcal{L}(\mathcal{S}(\Gamma))$,

- (i) f is a group convolution right operator $\Leftrightarrow f$ is equivariant to left multiplications,
- (ii) f is a group convolution left operator $\Leftrightarrow f$ is equivariant to right multiplications,
- (iii) f is a group convolution commutative operator $\Leftrightarrow f$ is equivariant to multiplications.

Proof. The proof is similar than the proof of Theorem 41.

 \Rightarrow : Given $w \in \mathcal{S}(\Gamma)$, let $f_R = .*_{\Gamma} w$, $f_L = w *_{\Gamma} ., s \in \mathcal{S}(\Gamma)$, and $p \in \Gamma$. We have:

$$(L_{p} \circ f_{R})(s) = L_{p}(s *_{\Gamma} w) \qquad (R_{p} \circ f_{L})(s) = R_{p}(w *_{\Gamma} s)$$

$$= L_{p}\left(\sum_{g \in \Gamma} s[g] L_{g}(w)\right) \qquad = R_{p}\left(\sum_{g \in \Gamma} s[g] R_{g}(w)\right)$$

$$= \sum_{g \in \Gamma} s[g] (L_{p} \circ L_{g})(w) \qquad = \sum_{g \in \Gamma} s[g] (R_{p} \circ R_{g})(w)$$

$$= \sum_{pg \in \Gamma} s[p^{-1}pg] L_{pg}(w) \qquad = \sum_{g \in \Gamma} s[gpp^{-1}] R_{gp}(w)$$

$$= \sum_{g \in \Gamma} L_{p}(s)[g] L_{g}(w) \qquad = \sum_{g \in \Gamma} R_{p}(s)[g] R_{g}(w)$$

$$= L_{p}(s) *_{\Gamma} w \qquad = w *_{\Gamma} R_{p}(s)$$

$$= (f_{R} \circ L_{p})(s) \qquad = (f_{L} \circ R_{p})(s)$$

 \Leftarrow : We show the converse only for (i) since it is similar for (ii). Let $f \in \mathcal{L}(\mathcal{S}(\Gamma))$. We suppose $\forall p \in \Gamma, L_p \circ f = f \circ L_p$. First, note that

$$\forall h \in \Gamma, L_g(\delta_g)[h] = 1 \Leftrightarrow \delta_g[g^{-1}h] = 1$$

$$\Leftrightarrow g^{-1}h = g$$

$$\Leftrightarrow h = e$$

$$i.e. \ L_g(\delta_g) = \delta_e$$

Then, we define $w = f(\delta_e)$. Let $s \in \mathcal{S}(\Gamma)$, we have:

$$f(s) = \sum_{g \in \Gamma} s[g] f(\delta_g)$$

$$= \sum_{g \in \Gamma} s[g] (f \circ L_g)(\delta_e)$$

$$= \sum_{g \in \Gamma} s[g] (L_g \circ f)(\delta_e)$$

$$= \sum_{g \in \Gamma} s[g] L_g(w)$$

$$= s *_{\Gamma} w$$

(iii) derives from (i) and (ii).

2.2.2 Steered construction from groups

For a graph $G = \langle V, E \rangle$ and a subgroup $\Gamma \subset \Phi^*(V)$ of its invertible transformations, Definition 43 is applicable for $\mathcal{S}(\Gamma)$, but not for $\mathcal{S}(V)$ as V is not a group. Nonetheless, we will try to use the group convolution on $\mathcal{S}(\Gamma)$ to construct the convolutions on $\mathcal{S}(V)$. Our goal is to construct a formulation like (31) while preserving Theorem 44 at the same time.

For now, let us assume that there is a subgroup $\Gamma \subset \Phi^*(V)$, which we associate through a one-to-one correspondence φ with V. We denote $\Gamma \stackrel{\varphi}{\cong} V$ and $g_v \stackrel{\varphi}{\mapsto} v$. Then, the linear morphism $\widetilde{\varphi}$ from $\mathcal{S}(\Gamma)$ to $\mathcal{S}(V)$ defined on the Dirac bases by $\widetilde{\varphi}(\delta_{g_v}) = \delta_{\varphi(g_v)}$ is a linear isomorphism. Hence $\mathcal{S}(V)$ inherits the same structural properties as $\mathcal{S}(\Gamma)$. For the sake of notational simplicity, we will use the same symbol φ for both φ and $\widetilde{\varphi}$ (as done between f and L_f). A commutative diagram between the sets is depicted on Figure 8.

$$\begin{array}{ccc} \Gamma & \stackrel{\varphi}{\longrightarrow} V \\ s \downarrow & & \downarrow s \\ \mathcal{S}(\Gamma) & \stackrel{\widetilde{\varphi}}{\longrightarrow} \mathcal{S}(V) \end{array}$$

Figure 8: Commutative diagram between sets

We naturally obtain the following relation, which put in simpler words means that signals on $S(\Gamma)$ are mapped to S(V) when φ is simultaneously applied on both the signal space and its domain. At the same time, this relation is reminiscent of Lemma 39.

Lemma 45. Relation between
$$S(\Gamma)$$
 and $S(V)$
 $\forall s \in S(\Gamma), \forall u \in V, \varphi(s)[u] = s[\varphi^{-1}(u)] = s[g_u]$

Proof.

$$\forall s \in \mathcal{S}(\Gamma), \varphi(s) = \varphi(\sum_{g \in \Gamma} s[g] \, \delta_g) = \sum_{g \in \Gamma} s[g] \, \varphi(\delta_g) = \sum_{g \in \Gamma} s[g] \, \delta_{\varphi(g)}$$
$$= \sum_{v \in V} s[g_v] \, \delta_v$$

So $\forall u \in V, \varphi(s)[u] = s[g_u].$

With this lemma, we can steer the definition of the group convolution from $S(\Gamma)$ to S(V) as in the following definition.

Remark. Note that the following definition is just a step in our construction since we will see that it is not enough to obtain the wanted counterpart of Theorem 41.

Definition 46. Vertex-group convolution

Let a subgroup $\Gamma \subset \Phi^*(V)$ such that $\Gamma \stackrel{\varphi}{\cong} V$. The vertex-group convolution between two signals s_1 and $s_2 \in \mathcal{S}(V)$ is defined as:

$$\forall u \in V, (s_1 *_{\Gamma} s_2)[u] = \sum_{v \in V} s_1[v] \, s_2[\varphi(g_v^{-1} g_u)]$$

We use the same symbol $*_{\Gamma}$ since the group and vertex-group convolution are inherently the same operation but applied to different domains.

Lemma 47. Relation between group and vertex-group convolutions

Let a subgroup $\Gamma \subset \Phi^*(V)$ such that $\Gamma \stackrel{\varphi}{\cong} V$,

$$\forall s_1, s_2 \in \mathcal{S}(\Gamma), \forall u \in V, (\varphi(s_1) *_{\Gamma} \varphi(s_2))[u] = (s_1 *_{\Gamma} s_2)[g_u]$$

Proof. Using Lemma 45,

$$(\varphi(s_1) *_{\Gamma} \varphi(s_2))[u] = \sum_{v \in V} \varphi(s_1)[v] \varphi(s_2)[\varphi(g_v^{-1}g_u)]$$

$$= \sum_{v \in V} s_1[g_v] s_2[g_v^{-1}g_u]$$

$$= \sum_{g \in \Gamma} s_1[g] s_2[g^{-1}g_u]$$

$$= (s_1 *_{\Gamma} s_2)[g_u]$$

Let f be a convolution right operator on S(V), s.t. $f = . *_{\Gamma} w$, and \widetilde{f} be its

counterpart on $\mathcal{S}(\Gamma)$, *i.e.* $\widetilde{f}= .*_{\Gamma} \varphi^{-1}(w).$ Then, we have

$$\forall s \in \mathcal{S}(V), f(s)[u] = \widetilde{f}(\varphi^{-1}(s))[g_u] \qquad \text{(Lemma 47)}$$

$$= \varphi(\widetilde{f}(\varphi^{-1}(s)))[u] \qquad \text{(Lemma 45)}$$

$$i.e. \ f = \varphi \circ \widetilde{f} \circ \varphi^{-1} \qquad \qquad (33)$$

This is also better depicted on a commutative diagram, see Figure 9.

$$\mathcal{S}(\Gamma) \xleftarrow{\varphi^{-1}} \mathcal{S}(V)
\widetilde{f} \downarrow \qquad \qquad \downarrow^{f}
\mathcal{S}(\Gamma) \xrightarrow{\varphi} \mathcal{S}(V)$$

Figure 9: Commutative diagram of (33)

And so, given $p \in \Gamma$, the equivariance of \widetilde{f} rewrites for f:

$$\varphi \circ L_p \circ \widetilde{f} \circ \varphi^{-1} = \varphi \circ \widetilde{f} \circ L_p \circ \varphi^{-1} \tag{34}$$

i.e.
$$(\varphi \circ L_p \circ \varphi^{-1}) \circ f = f \circ (\varphi \circ L_p \circ \varphi^{-1})$$
 (35)

That is, if we paraphrase Theorem 41, the class of vertex-group convolution operators is the class of linear operators that are equivariant to $\{\varphi \circ L_p \circ \varphi^{-1}, p \in \Gamma\}$. However, our goal is to obtain equivariance to Γ . Since L_p is not exactly an object of Γ , but a realization of a group action, we are going to consider group actions next.

2.2.3 Construction under group actions

We have been previously using the notion of group action implicitly, let us define it here.

Definition 48. Group action

An *action* of a group Γ on a set V is a homomorphism $L: \Gamma \to \Phi^*(V)$.

In particular, $L(\Gamma)$ is a subgroup of the symmetric group $\Phi^*(V)$.

Remark. Given $g \in \Gamma$, we denote $L_g = L(g)$, or just g(.) when there is no ambiguity. L_g is called the action of g by L on V. When a group Γ is in one-to-one correspondence with a set V, we will abusively attribute actions of the objects of the group to their corresponding objects in the set.

Hence, note that an group element $g \in \Gamma$ can act on both Γ through the left multiplication L and on V. This ambivalence can be seen on a commutative diagram, see Figure 10.

$$g_u \xrightarrow{L_{g_v}} g_v g_u$$

$$\varphi \downarrow \qquad \qquad \downarrow \varphi$$

$$u \xrightarrow{-\frac{(36)}{g_v}} \varphi(g_v g_u)$$

Figure 10: Commutative diagram. All arrows except for the one labeled with (36) are always true.

For (36) to be true means that φ is an equivariant map *i.e.* whether the mapping is done before or after the action of Γ has no impact on the result. When such φ exists, Γ and V are said to be equivalent and we denote $\Gamma \equiv V$.

Definition 49. Equivariant map

A map φ from a group Γ acting on the destination set V and itself, is said to be an *equivariant map* if

$$\forall g, h \in \Gamma, g(\varphi(h)) = \varphi(g(h))$$

Remark. For example, if g acts on Γ through left multiplication then $g(h) = L_g(h) = gh$.

Suppose we have $\Gamma \stackrel{\varphi}{\cong} V$. If we also have that $\Gamma \equiv V$, we are interested to know if then φ exhibits the equivalence and under which auto-action.

Definition 50. φ -Equivalence

A group Γ acting on V such that $\Gamma \stackrel{\varphi}{\cong} V$, is said to be (*left*) φ -equivalent if φ is a bijective equivariant map under left multiplication *i.e.*:

$$\forall v, u \in V, g_v(u) = \varphi(g_v g_u) \tag{36}$$

i.e.
$$\forall g \in \Gamma, g(.) = \varphi \circ L_g \circ \varphi^{-1}$$
 (37)

It is said to be *right* φ -equivalent if φ is a bijective equivariant map under right multiplication *i.e.* :

$$\forall v, u \in V, g_v(u) = \varphi(g_u g_v) \tag{38}$$

i.e.
$$\forall g \in \Gamma, g(.) = \varphi \circ R_g \circ \varphi^{-1}$$
 (39)

We denote $\Gamma \stackrel{\varphi}{\equiv} V$. Unless stated otherwise, we will implicitly consider it is under left multiplication.

Remark. For example, translations on the grid graph, with $\varphi(t_{i,j})=(i,j)$, are φ -equivalent since $t_{i,j}(a,b)=\varphi(t_{i,j}\circ t_{a,b})$. However, with $\varphi(t_{i,j})=(-i,-j)$, they would not be φ -equivalent since in general $t_{i,j}(a,b)\neq t_{i,j}(-a,-b)$. We are now equipped to define the vertex-action convolution.

Definition 51. Vertex-action convolution

Let a group Γ acting on V such that $\Gamma \stackrel{\varphi}{\cong} V$. The vertex-action convolution between two signals s_1 and $s_2 \in \mathcal{S}(V)$ is defined as:

$$s_1 *_V s_2 = \sum_{v \in V} s_1[v] g_v(s_2)$$
 (40)

$$= \sum_{g \in \Gamma} s_1[\varphi(g)] g(s_2) \tag{41}$$

The two expressions differ on the domain upon which the summation is done. Expression (40) puts the emphasis on each vertex and its corresponding action, whereas expression (41) puts the emphasis on the action of Γ .

Lemma 52. Relation with vertex-group convolution

There is φ -equivalence if, and only if, vertex-group and vertex-action convolutions are equal, *i.e.* $\Gamma \stackrel{\varphi}{\equiv} V \Leftrightarrow *_{\Gamma} = *_{V}$

Proof.

$$\forall s_1, s_2 \in \mathcal{S}(V),$$

$$s_1 *_{\Gamma} s_2 = s_1 *_{V} s_2$$

$$\Leftrightarrow \forall u \in V, \sum_{v \in V} s_1[v] \, s_2[\varphi(g_v^{-1}g_u)] = \sum_{v \in V} s_1[v] \, s_2[g_v^{-1}(u)] \qquad (42)$$

Hence, the direct sense is obtained by applying (36).

For the converse, given $u, v \in V$, we first realize (42) for $s_1 := \delta_v$, obtaining $s_2[\varphi(g_v^{-1}g_u)] = s_2[g_v^{-1}(u)]$, which we then realize for a real signal s_2 having

no two equal entries, obtaining $\varphi(g_v^{-1}g_u)=g_v^{-1}(u)$. From the latter we finally obtain (36) with the one-to-one correspondence $g_{v'}:=g_v^{-1}$.

Remark. Note that we used the fact that the φ -equivalence is implicitly left. If it were right, we would have instead $s_1 *_{\Gamma} s_2 = s_2 *_{V} s_1$.

We now coin the term φ -convolution, for when there is φ -equivalence.

Definition 53. φ -convolution

The φ -convolution is defined as the vertex-action convolution such that $\Gamma \stackrel{\varphi}{=} V$, *i.e.* given s_1 and $s_2 \in \mathcal{S}(V)$, we have:

$$s_1 *_{\varphi} s_2 = s_1 *_V s_2$$
$$= s_1 *_{\Gamma} s_2$$

This time we do obtain equivariance to Γ as expected, and the full characterization as well.

Theorem 54. Characterization of φ -convolution right operators

Let a group Γ acting on V *s.t.* $\Gamma \stackrel{\varphi}{\equiv} V$, let $f \in \mathcal{L}(\mathcal{S}(\Gamma))$, then:

f is a $\varphi\text{-convolution}$ right operator $\Leftrightarrow f$ is equivariant to Γ

Proof. As a consequence of (35), (37), Lemmas 47, and 52, we can use bullet (i) of Theorem 44 to obtain the proof.

Corollary 55. Characterization of φ -convolution operators

Let a group Γ acting on V *s.t.* $\Gamma \stackrel{\varphi}{=} V$, let $f \in \mathcal{L}(\mathcal{S}(\Gamma))$, then:

f is equivariant to $\Gamma\Leftrightarrow f$ is a φ -convolution operator s.t. its laterality is opposed to the laterality of the φ -equivalence

Proof. As per Theorem 54 and its counterpart for bullet (ii) of Theorem 44.

Remark. The counterpart of bullet (iii) of Theorem 44 would just inform that it is equivalent to say that either the φ -equivalence is commutative, or the φ -convolution is commutative, or Γ is abelian.

Theorem 54 tells us that in order to define a convolution on the vertex domain of a graph $G = \langle V, E \rangle$, all we need is a group Γ acting on the vertex set V (or on a subset of V), that is equivalent to it. The choice of Γ can be done with respect to the edge set E. This is discussed in more details in Section 2.3, where we will see that in fact, we only need a generating set of Γ .

The previous construction relies on exposing a bijective equivariant map φ between Γ and V. In the next subsection, we show that in case Γ is abelian, we even need not expose φ and the characterization still holds.

2.2.4 Mixed domain formulation

From (41), we can define a mixed domain convolution *i.e.* that is defined for $r \in \mathcal{S}(\Gamma)$ and $s \in \mathcal{S}(V)$, without the need of expliciting φ .

Definition 56. Mixed domain convolution

Let a group Γ acting on V. The *mixed domain convolution* between two signals $r \in \mathcal{S}(\Gamma)$ and $s \in \mathcal{S}(V)$ results in a signal $r *_{\mu} s \in \mathcal{S}(V)$ and is defined as:

$$r *_{\mu} s = \sum_{g \in \Gamma} r[g] g(s)$$

We call it μ -convolution. From a practical point of view, this expression of the convolution is useful because it relegates φ as an underpinning object.

Lemma 57. Relation with vertex-action convolution

$$\forall \varphi \in \mathrm{Bij}(\Gamma, V), \forall (r, s) \in \mathcal{S}(\Gamma) \times \mathcal{S}(V),$$
$$r *_{u} s = \varphi(r) *_{V} s$$

Proof. Let $\varphi \in BIJ(\Gamma, V), (r, s) \in \mathcal{S}(\Gamma) \times \mathcal{S}(V)$,

$$r *_{\mu} s = \sum_{g \in \Gamma} r[g] g(s) = \sum_{v \in V} r[g_v] g_v(s) \stackrel{(\diamond)}{=} \sum_{v \in V} \varphi(r)[v] g_v(s)$$
$$= \varphi(r) *_{V} s$$

Where $\stackrel{(\diamond)}{=}$ comes from Lemma 45.

In other words, $*_{\mu}$ is a convenient reformulation of $*_{V}$.

Lemma 58. Relation with group, vertex-group and φ -convolutions

Let $\varphi \in \text{BIJ}(\Gamma, V), (r, s) \in \mathcal{S}(\Gamma) \times \mathcal{S}(V)$, we have:

$$\Gamma \stackrel{\varphi}{\equiv} V \Leftrightarrow \forall v \in V, (r *_{\mu} s)[v] = (r *_{\Gamma} \varphi^{-1}(s))[g_v]$$
$$\Leftrightarrow r *_{\mu} s = \varphi(r) *_{\Gamma} s$$
$$\Leftrightarrow r *_{\mu} s = \varphi(r) *_{\varphi} s$$

Proof. On one hand, Lemma 57 gives $r *_{\mu} s = \varphi(r) *_{V} s$. On the other hand, Lemma 47 gives $\forall v \in V, (r *_{\Gamma} \varphi^{-1}(s))[g_{v}] = (\varphi(r) *_{\Gamma} s)[v]$. Then Lemma 52 concludes.

From μ -convolution, a left operator can be used as a layer of a neural network since it is defined over S(V), whereas right operators are not.

Proposition 59. Condition of equivariance for $*_{\mu}$

Let a group Γ acting on V. Γ is abelian, if and only if, μ -convolution left operators are equivariant to Γ .

Proof. Let $w, g \in \Gamma$, and define $f_w : s \mapsto w *_{\mu} s$. In the following expressions, Γ is abelian if and only if (43) and (44) are equal (the converse is obtained by particularizing on well chosen signals):

$$(f_w \circ g)(s) = \sum_{h \in \Gamma} w[h] hg(s)$$

$$= \sum_{h \in \Gamma} w[h] gh(s)$$

$$= g\left(\sum_{h \in \Gamma} w[h] h(s)\right)$$

$$= g(w *_{\mu} s)$$

$$= (g \circ f_w)(s)$$

$$(43)$$

Remark. We used this proof because it is simpler and does not require one-to-one correspondence between Γ and V. However, the reason behind the abelian condition is that mixed domain convolutions are expressed as if the underlying φ -equivalence were left. Therefore it is a consequence of Corrolary 55. Note that if we used a right expression for $*_{\mu}$ (which amounts to commute the operands), we would be interested in right operators instead of left ones, which would just change the problem to "right-right" instead of "left-left" which would also require abelianity for the equivariance.

We then coin the term M-convolution, for which Γ is abelian.

Definition 60. M-convolution

Let a group Γ acting on V. The M-convolution is defined as a mixed domain convolution such that Γ is an abelian subgroup of $\Phi^*(V)$.

Corollary 61. Characterization of M-convolution left operators

Let a group Γ acting on V *s.t.* $\Gamma \cong V$, and Γ is abelian. Let $f \in \mathcal{L}(\mathcal{S}(\Gamma))$, then:

f is a M-convolution left operator $\Leftrightarrow f$ is equivariant to Γ

Proof. As a corollary of Corrolary 55 and Proposition 59.

Remark. Note that despite abelianity, we do not call them "commutative" operators but still call "left" operators since the domain is mixed. Also, note that unlike for Proposition 59, the converse of Corrolary 61 requires Γ and V to be in one-to-one correspondence.

When the group domain is abelian, it is preferable to use $*_{\mathsf{M}}$ rather than $*_{\varphi}$ since φ is not needed. When Γ and V are not in one-to-one correspondence, the equivariance is preserved but not the characterization. This is not a limitation on the size of the kernel w of corresponding convolution operators since small kernels can be obtained by limiting the support of w.

The obtained constructions φ - and M-convolutions are independent of the edge set. In fact, they characterize more generally convolutions for signals defined on sets. Next, we will see what happens should we also consider the edge set.

2.3 Inclusion of the edge set in the construction

The constructions from the previous section involve the vertex set V and a group Γ acting on it. Therefore, it looks natural to try to relate the edge set and Γ .

There are two point of views. Either Γ describes an underlying graph structure $G = \langle V, E \rangle$, either G can be used to define a relevant subgroup Γ to which the produced convolutive operators will be equivariant. Both approaches will help characterize classes of graphs that can support natural definitions of convolutions.

2.3.1 Edge-constrained convolutions

Definition 62. Edge-constrained transformation

An *edge-constrained* (EC) transformation on a graph $G = \langle V, E \rangle$ is a transformation $f: V \mapsto V$ such that

$$\forall u, v \in V, f(u) = v \Rightarrow u \stackrel{E}{\sim} v$$

We denote $\Phi_{\text{EC}}(G)$ and $\Phi_{\text{EC}}^*(G)$ the sets of EC and invertible EC transformations. Note that $\Phi_{\text{EC}}^*(G)$ is not a group. Therefore, we are interested in generating sets of groups.

Definition 63. Edge-constrained convolution

An edge-constrained (EC) convolution on a graph $G = \langle V, E \rangle$ is a φ - or m-convolution of group Γ , such that Γ is generated by a set \mathcal{U} which actions on V are EC transformations.

This leads us to consider Cayley graphs (Cayley, 1878).

Definition 64. Cayley graph

Let a group Γ and one of its generating set \mathcal{U} . The *Cayley graph* generated by \mathcal{U} , is the digraph $\vec{G} = \langle V, E \rangle$ such that $V = \Gamma$ and E is such that, either:

- (i) $\forall a, b \in \Gamma, a \to b \Leftrightarrow \exists g \in \mathcal{U}, ga = b$ (left Cayley graph)
- (ii) $\forall a, b \in \Gamma, a \to b \Leftrightarrow \exists g \in \mathcal{U}, ag = b$ (right Cayley graph)

We denote $\vec{G} = \langle \mathcal{U}, \Gamma, E \rangle$. If Γ is abelian, we call it an *abelian Cayley graph*. Also, we call *Cayley subgraph* a directed subgraph that is isomorphic to a Cayley graph.

Remark. Note that in the literature, a Cayley graph is usually what we defined as a right one.

In the case of left Cayley graphs of vertex set Γ , it is clear that the φ -convolution based on Γ is EC (since the φ -equivalence is implicitly under left multiplication). More precisely, we obtain the following characterization:

Theorem 65. EC characterization by Cayley subgraphs

Let a graph $G = \langle V, E \rangle$,

- (i) its left Cayley subgraphs characterize its EC φ -convolutions,
- (ii) its abelian Cayley subgraphs characterize its EC м-convolutions.

Proof. We show the result only in the first case since the proof in the abelian case is similar.

1. Let an EC φ -convolution of group Γ . Since $*_{\varphi}$ is EC, there is a generating set \mathcal{U} such that its actions are EC. So the graph \vec{G}_s defined such that $u \to v \Leftrightarrow \exists g \in \mathcal{U}, g(u) = v$ is a subgraph of G, and we have:

$$u \to v \Leftrightarrow \exists g \in \mathcal{U}, g(u) = v$$

$$\Leftrightarrow \exists g \in \mathcal{U}, g(\varphi(g_u)) = \varphi(g_v)$$

$$\Leftrightarrow \exists g \in \mathcal{U}, \varphi(gg_u) = \varphi(g_v)$$

$$\Leftrightarrow \exists g \in \mathcal{U}, gg_u = g_v$$
(since (36))

Therefore the subgraph \vec{G}_s and the Cayley graph $\langle \mathcal{U}, \Gamma, E_c \rangle$ are isomorphic via φ .

2. Let a subgraph $\vec{G}_s = \langle V_s, E_s \rangle$ that is isomorphic to a left Cayley graph $\vec{G}_c = \langle \mathcal{U}, \Gamma, E_c \rangle$. Let ψ be a graph isomorphism from G_s to G_c .

Let us define the group action $L: \Gamma \to \Phi^*(V_s)$ inductively as follows:

(a)
$$\forall g \in \mathcal{U}, L_g(u) = v \Leftrightarrow g\psi(u) = \psi(v)$$

- (b) Whenever L_g and L_h are defined, the action of gh is defined by homomorphism as $L_{gh}=L_g\circ L_h$
- (c) Whenever L_g is defined, the action of g^{-1} is defined by homomorphism as $L_{g^{-1}} = L_g^{-1}$ i.e. $L_{g^{-1}}(u) = v \Leftrightarrow \psi(u) = g\psi(v)$

Note that the induction transfers the property (a) to all $g \in \Gamma$ in a transitive manner because

$$L_{gh}(u) = L_g(L_h(u)) = w \Leftrightarrow \exists v \in V_s \begin{cases} L_h(u) = v \\ L_g(v) = w \end{cases}$$

and

$$\exists v \in V_s \begin{cases} h\psi(u) = \psi(v) \\ g\psi(v) = \psi(w) \end{cases} \Leftrightarrow gh\psi(u) = \psi(w)$$

We must also verify that this construction is well-defined, *i.e.* whenever we define an action with (b) or (c), if the action was already defined, then they must be equal. This is the case because the homo-

morphism $g\mapsto L_g$ on Γ is in fact an isomorphism as

$$L_g = L_h \Leftrightarrow \forall u \in V, L_g(u) = L_h(u)$$

 $\Leftrightarrow \forall u \in V, g\psi(u) = h\psi(u)$
 $\Leftrightarrow g = h$

Also note that (c) is needed only in case that Γ is infinite.

Denote $\varphi = \psi^{-1} : g_u \mapsto u$ the inverse graph isomorphism. Since (a) has been transferred from \mathcal{U} onto Γ , we have, for all $g, h \in \Gamma$:

$$L_g(u) = v \Leftrightarrow g\psi(u) = \psi(v)$$

 $\Leftrightarrow \varphi(g\psi(u)) = v$
i.e. $L_g(u) = \varphi(gg_u)$

So $\Gamma \stackrel{\varphi}{=} V$. Furthermore, the corresponding φ -convolution is EC because of (a).

Remark. Note that right φ -equivalences are instead characterized by right Cayley graphs.

Through the former proof, we also obtain the following corollary. Note that we call an orientation $\vec{G}_s = \langle V_s, E_s \rangle$ of a graph $G = \langle V, E \rangle$, a directed subgraph with same vertex set $V_s = V$ and such that $E_s \subset E$.

Corollary 66. φ is a graph isomorphism from a Cayley graph

Let a graph $G = \langle V, E \rangle$, a group Γ of generating set \mathcal{U} such that $\Gamma \stackrel{\varphi}{\equiv} V$. Then φ is also a graph isomorphism between the left Cayley graph $\langle \mathcal{U}, \Gamma, E_c \rangle$ and an orientation of G.

2.3.2 On properties of the corresponding operators

Until now we have described operators with kernel w of any size. However, in practice in deep learning we rather use convolution operators with small kernels. This case is indeed encompassed by kernel of any size by considering a smaller support. This is convenient with edge-constrained convolutions, since we can choose a support not larger than the span of the generating set \mathcal{U} , so that an EC convolution operator can be expressed with only actions of \mathcal{U} . We formalize this discussion as follows.

Definition 67. Supporting set

Let f_w^{φ} be an EC φ -convolution right operator of group Γ . The supporting set of f_w^{φ} is the largest subset $\mathcal{M} \subset \Gamma$ such that $\varphi(\mathcal{M}) \subset \operatorname{supp}(w)$. Let f_w^{M} be an EC M -convolution left operator of group Γ . The supporting set of f_w^{M} is the largest subset $\mathcal{M} \subset \Gamma$ such that $\mathcal{M} \subset \operatorname{supp}(w)$.

Definition 68. Strictly edge-constrained convolution operator

We say that an EC φ or M-convolution operator, of group Γ and generating set \mathcal{U} , is strictly edge-constrained (EC*) if its supporting set $\mathcal{M} \subset \mathcal{U}$ and \mathcal{M} is symmetric (i.e. $g \in \mathcal{M} \Leftrightarrow g^{-1} \in \mathcal{M}$).

Remark. EC* convolution operators are simpler to obtain as we can construct them just with the actions $\mathcal{U} \to \Phi_{\text{EC}}^*(G)$ without composing the transformations. Also, their complexity is $\mathcal{O}(kn)$, where n = |V| is the order of the graph and $k = |\mathcal{M}|$ is the size of the kernel. In comparison, EC convolutions have complexity up to $\mathcal{O}(n^2)$.

Regardless of being EC, EC*, or not EC, the supporting set \mathcal{M} allows for writing the operators as a sum with less elements. In the abelian case, the formulation of $f_w^{\mathbf{M}}$ is simplified as:

$$f_w^{\mathbf{M}}(s) = \sum_{g \in \mathcal{M}} w[g] g(s) \tag{45}$$

In the non-abelian case, the formulation is less explicit and depends on the vertex u on which the convolution operator is realized. Since we have

$$w[g_v^{-1}(u)] \neq 0 \Leftrightarrow g_v^{-1}(u) \in \varphi(\mathcal{M})$$
$$\Leftrightarrow g_v^{-1}g_u \in \mathcal{M}$$
$$\Leftrightarrow g_v^{-1} \in \mathcal{M}g_u^{-1}$$
$$\Leftrightarrow v \in g_u(\varphi(\mathcal{M}^{-1}))$$

where $\mathcal{M}^{-1} = \{g^{-1}, g \in \mathcal{M}\}$ (which equals \mathcal{M} in the usual case if it is symmetric). Denote $\mathcal{K}_u = g_u(\varphi(\mathcal{M}^{-1}))$, f_w^{φ} can be rewritten as:

$$\forall u \in V, f_w^{\varphi}(s)[u] = \sum_{v \in \mathcal{K}_u} s[v] w[g_v^{-1}(u)]$$
(46)

Let us call r the *root vertex* defined as the φ -fiber of the identity element e (i.e. $g_r = e$). We can interpret (46) as if the convolution was first realized on r with a sum over a vertex patch initially localized on $\mathcal{K}_e = \varphi(\mathcal{M}^{-1})$, and then its realization on another vertex u would amount to move this patch under the action of g_u , exactly like with convolution operators on Euclidean domains.

From this expression, we can observe the following proposition.

Proposition 69. Weight sharing

Let a φ - or M-convolution operator f_w of supporting set $\mathcal{M} \subset \Gamma$. Then the entries of w are shared between each realization of f_w .

Proof. By reading (45), this result is obvious for M-convolution operators. So let us assume that f_w is a φ -convolution operator. Given a realization vertex $a \in V$, the non-null entries of w in expression (46) are characterized by vertices $\alpha \in \mathcal{K}_a$. Given another realization vertex b, its non-nil entries

match those of a if, and only if,

$$g_{\mathcal{K}_b}^{-1}(b) = g_{\mathcal{K}_a}^{-1}(a) \Leftrightarrow g_b^{-1}g_{\mathcal{K}_b} = g_a^{-1}g_{\mathcal{K}_a}$$
$$\Leftrightarrow \mathcal{K}_b = g_b g_a^{-1}(\mathcal{K}_a) = g_b g_a^{-1}g_a(\varphi(\mathcal{M}^{-1}))$$
$$\Leftrightarrow \mathcal{K}_b = g_b(\varphi(\mathcal{M}^{-1}))$$

which is true by definition.

Although the weight-sharing property is preserved and the φ -convolution can be seen as moving a patch \mathcal{K}_u around V, the latter patch can have a different shape for each vertex u. We discuss in the next subsection what characterizes convolutions for which this shape is stationary.

2.3.3 Locality-preserving convolutions

Euclidean convolution operators have the property of preserving the local shape, in the sense that the shape of the kernel w is preserved while being moved over the domain. In the case of graphs, what we call shape is more formally a subgraph. Therefore, the notion of locality preservation we aim to define is associated with graph automorphisms.

Definition 70. Locality-preserving convolutions

Let a graph $G = \langle V, E \rangle$, we say that a φ - or M-convolution, of group Γ , is locality-preserving (LP) if the actions of Γ are graph automorphisms.

Similarly than with EC and EC*, we are also interested in this property for operators with smaller kernels.

Definition 71. Strictly locality-preserving convolution operator

We say that a φ -convolution operator, of supporting set $\mathcal{M} \subset \Gamma$, is strictly locality-preserving (LP*) if every subgraph $\mathcal{K}_u = g_u(\mathcal{K}_e)$ are isomorphic.

Remark. We assume that M-convolution operator are always LP* in the sense that the sum is always over \mathcal{M} . Also note that LP is a stronger property than LP* (*i.e.* LP implies LP*) and that LP* and LP are not related in the same way than EC* and EC.

Similarly than for EC convolutions, LP convolutions can be characterized with Cayley subgraphs.

Lemma 72. On left Cayley graphs, right multiplicative auto-actions are automorphisms (and vice-versa).

Proof. Since
$$a \to b \Leftrightarrow \exists g, ga = b \Leftrightarrow \exists g, gah = bh \Leftrightarrow ah \to bh$$
.

Theorem 73. LP characterization by Cayley subgraphs Let a graph $G = \langle V, E \rangle$,

- (i) its right Cayley subgraphs characterize its LP φ -convolutions
- (ii) its abelian Cayley subgraphs characterize its LP м-convolutions

Proof. Let a group Γ of generating set \mathcal{U} . Denote $\vec{G^R}$ the right Cayley graph and $\vec{G^L}$ the left one. Let us assume that $\Gamma \stackrel{\varphi}{\equiv} V$. We denote the action corresponding to the usual left equivalence by $g^L(.) = \varphi \circ L_g \circ \varphi^{-1}$, and the action corresponding to the right equivalence by $g^R(.) = \varphi \circ R_g \circ \varphi^{-1}$, where L and R are the left and right multiplicative auto-actions by Γ on itself, as in (37) and (39). By Lemma 72, $g^L(.)$, that is EC on $\vec{G^L}$, is LP on $\vec{G^R}$, and vice-versa. By Theorem 65, $\vec{G^L}$ characterizes $g^L(.)$, while naturally also characterizing $\vec{G^R}$. Therefore, $\vec{G^R}$ characterizes $g^L(.)$, onto which it is LP, so is the corresponding φ -convolution.

Corollary 74. Characterization of convolutions that are EC and LP Let a graph $G = \langle V, E \rangle$,

- (i) if a φ -convolution of group Γ is EC and LP then Γ is abelian,
- (ii) an M-convolution is EC if, and only if, it is also LP.

Proof. As a consequence of Theorem 65 and Theorem 73. □

2.3.4 Checkpoint summary

In Section 2.2, we obtained two formulations of convolutions of signals on a graph $G = \langle V, E \rangle$, $*_{\varphi}$ and $*_{\mathsf{M}}$, depending on a group Γ , that preserve fully the characterization by equivariance to actions of Γ . The φ -convolution can be used given a bijective equivariant map φ between Γ and a subset $V_s \subset V$, whereas the M-convolution can be used under the condition that Γ is abelian. Then in Section 2.3, we introduced conditions related to the edge set E, which leads us to define EC and LP convolutions, that we were able to characterize by Cayley subgraph isomorphism, and for which we were capable to understand the impact of abelianity. We also defined EC* and LP* convolution operators, that are closer to what can be used in deep learning practice. In particular, to define an EC* convolution operator, we only need to search for a set of actions that is a generating set of Γ . A practical example will be seen in Section 3.4.

The sets of theorems we obtained are already satisfactory enough to understand how to convolute signals on any graph, using symmetries defined by a group acting on the vertex domain (or on a subset of the vertex domain). Therefore we can consider that the main contributions of this chapter have already been presented.

Nonetheless, symmetries defined by groups can have limitations, since they may be too perfect for some graphs, as discussed in the next section. Hence, we can try to extend further the construction with partial symmetries defined by groupoids. That is the research avenue that we explore in the remainder of this chapter.

2.4 From groups to groupoids

2.4.1 Motivation

One possible limitation coming from searching for Cayley subgraphs is that they are degree-regular *i.e.* the in- and the out-degree $d = |\mathcal{U}|$ of each vertex is the same. That is, for a general graph G, the size of the weight kernel w of an EC* convolution operator f_w supported on \mathcal{U} is bounded by d, which in turn is bounded by twice the minimal degree of G (twice because G is undirected and \mathcal{U} can contain every inverse).

There are a lot of possible strategies to overcome this limitation. For example:

- 1. connecting each vertex with its k-hop neighbors, with k > 1,
- 2. artificially creating new connections for less connected vertices,
- 3. ignoring less connected vertices,
- 4. allowing the supporting set \mathcal{M} to exceed \mathcal{U} *i.e.* dropping * in EC*.

These strategies require to concede that the topological structure supported by G is not the best one to support an EC* convolution on it, which breeds the following question:

 What can we relax in the previous EC* construction in order to unbound the supporting set, and still preserve the equivariance characterization?

The latter constraint is a consequence that every vertex of the Cayley subgraph \vec{G} must be composable with every generator from \mathcal{U} . Therefore, an answer consists in considering groupoids (Brandt, 1927) instead of groups. Roughly speaking, a groupoid is almost a group except that its composition law needs not be defined everywhere. Weinstein, 1996, unveiled the

benefits to base convolutions on groupoids instead of groups in order to exploit partial symmetries.

2.4.2 Definition of notions related to groupoids

Definition 75. Groupoid

A *groupoid* Υ is a set equipped with a partial composition law with domain $\mathcal{D} \subset \Upsilon \times \Upsilon$, called *composition rule*, that is

1. closed into Υ *i.e.* $\forall (g,h) \in \mathcal{D}, gh \in \Upsilon$

$$\textbf{2. associative } \textit{i.e.} \ \forall f,g,h \in \Upsilon, \begin{cases} (f,g),(g,h) \in \mathcal{D} \Leftrightarrow (fg,h),(f,gh) \in \mathcal{D} \\ (f,g),(fg,h) \in \mathcal{D} \Leftrightarrow (g,h),(f,gh) \in \mathcal{D} \\ \text{when defined, } (fg)h = f(gh) \end{cases}$$

3. invertible i.e.
$$\forall g \in \Upsilon, \exists ! g^{-1} \in \Upsilon$$
 s.t.
$$\begin{cases} (g, g^{-1}), (g^{-1}, g) \in \mathcal{D} \\ (g, h) \in \mathcal{D} \Rightarrow g^{-1}gh = h \\ (h, g) \in \mathcal{D} \Rightarrow hgg^{-1} = h \end{cases}$$

Optionally, it can be *domain-symmetric i.e.* $(g,h) \in \mathcal{D} \Leftrightarrow (h,g) \in \mathcal{D}$, and *abelian i.e.* domain-symmetric with gh = hg.

Remark. Note that left and right inverses are necessarily equal (because $(gg^{-1})g = g(g^{-1}g)$). Also note we can define a right identity element $e_g^r = g^{-1}g$, and a left one $e_g^l = gg^{-1}$, but they are not necessarily equal and depend on g.

Most definitions related to groups can be adapted to groupoids. In particular, let's adapt a few notions.

Definition 76. Partial transformations groupoid

The partial transformations groupoid $\Psi^*(V)$, is the set of invertible partial transformations, equipped with the functional composition law with domain \mathcal{D} such that

$$\begin{cases} \mathcal{D}_{gh} = h(\mathcal{D}_h) \cap \mathcal{D}_g \\ (g, h) \in \mathcal{D} \Leftrightarrow \mathcal{D}_{gh} \neq \emptyset \end{cases}$$

Remark. Note that a subgroupoid $\Upsilon \subset \Psi^*(V)$ is domain-symmetric when $\exists v \in V, g(v) \in \mathcal{D}_h \Leftrightarrow \exists u \in V, h(u) \in \mathcal{D}_g$

Definition 77. Groupoid partial action

A partial *action* of a groupoid Υ on a set V is a groupoid homomorphism $L: \Upsilon \to \Psi^*(V)$.

Remark. As usual, we will confound L_g and g(.) when there is no possible confusion, and we denote $\mathcal{D}_{L_g} = \mathcal{D}_g = \{v \in V, (g, v) \in \mathcal{D}_L\}.$

Definition 78. Partial equivariant map

A map φ from a groupoid Υ partially acting on the destination set V is said to be a *partial equivariant map* if

$$\forall g, h \in \Upsilon, \begin{cases} \varphi(h) \in \mathcal{D}_g \Leftrightarrow (g, h) \in \mathcal{D} \\ g(\varphi(h)) = \varphi(g(h)) \end{cases}$$

Also, φ -equivalence between a groupoid and a set is defined similarly than for groups, with also left multiplicative partial action being implicit.

2.4.3 Construction of partial convolutions

The expression of the convolution we constructed in the previous section cannot be applied as is. We first need to extend the algebraic objects we work with. Extending a partial transformation g on the signal space $\mathcal{S}(V)$ (and thus the convolutions) is a bit tricky, because only the signal entries corresponding to \mathcal{D}_g are moved. A convenient way to do this is to consider the groupoid closure obtained with the addition of an absorbing element.

Definition 79. Zero-closure

The *zero-closure* of a groupoid Υ , denoted Υ^0 , is the set obtained by addition of an absorbing element, denoted 0, such that the groupoid axioms 1, 2 and 3, and the domain \mathcal{D} are left unchanged, and

4. the composition law is extended to $\Upsilon^0 \times \Upsilon^0$ with $\forall (g,h) \notin \mathcal{D}, gh = 0$

Remark. Note that this is coherent as the properties 2 and 3 are still partially defined on the original domain \mathcal{D} .

Now, we will also extend every other algebraic object used in the expression of the φ -convolution and the M-convolution, so that we can directly apply our previous constructions.

Lemma 80. Zero-extension of φ on V^0

Let a partial equivariant map $\varphi: \Upsilon \to V$. It can be extended to a (total) equivariant map $\varphi: \Upsilon^0 \to V^0 = V \cup \varphi(0)$, such that $\varphi(0) \notin V$, that we denote $0_V = \varphi(0)$, and such that

$$\forall g \in \Upsilon^0, \forall v \in V^0, g(v) = \begin{cases} \varphi(gg_v) & \text{if } g_v \in \mathcal{D}_g \\ 0_V & \text{else} \end{cases}$$

Proof. We have $\varphi(0) \notin V$ because φ is bijective. Additionally, we must have $\forall (g,h) \notin \mathcal{D}, g(\varphi(h)) = \varphi(gh) = \varphi(0) = 0_V$.

Remark. Note that for notational convenience, we may use the same symbol 0 for 0_{Υ} , 0_V and $0_{\mathbb{R}}$.

Similarly to $\Phi^*(V)$, $\Psi^*(V)$ can also move signals of $\mathcal{S}(V)$.

Lemma 81. Zero-extension of partial actions to S(V)

Let $g \in \Psi^*(V)$. Its extension is done in two steps:

- 1. g is extended to $V^0 = V \cup \{0_V\}$ as $g(v) = 0_V \Leftrightarrow v \notin \mathcal{D}_g$.
- 2. Under the convention $\forall s \in \mathcal{S}(V), s[0_V] = 0_{\mathbb{R}}, g$ is extended via linear extension to $\mathcal{S}(V)$, and we have

$$\forall s \in \mathcal{S}(V), \forall v \in V, g(s)[v] = s[g^{-1}(v)]$$

Proof. Straightforward.

With these extensions, we can obtain the partial φ - and M-convolutions related to Υ almost by substituting Υ^0 to Γ in Definition 53 and Definition 56.

Definition 82. Partial convolutions

Let a groupoid Υ acting on V^0 . The partial φ - and M-convolutions extend their definition on groups to groupoids using the appropriate zero-extensions *i.e.*

(i)
$$\forall s, w \in \mathcal{S}(V), s *_{\varphi} w = \sum_{v \in V} s[v] g_v(w) = \sum_{g \in \Upsilon} s[\varphi(g)] g(w) s.t. \Upsilon \stackrel{\varphi}{\equiv} V$$

(ii)
$$\forall (w,s) \in \mathcal{S}(\Upsilon) \times \mathcal{S}(V), w *_{\mathbf{M}} s = \sum_{g \in \Upsilon} w[g] \, g(s) \, \textit{s.t.} \, \Upsilon \text{ is abelian}$$

Symmetrical expressions

Note that, as $\forall r, r[0] = 0$, the partial convolutions can also be expressed on the domain \mathcal{D} with a convenient symmetrical expression:

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(i)
$$\forall u \in V, (s *_{\varphi} w)[u] = \sum_{\substack{(g_a, g_b) \in \mathcal{D} \\ s.t. \ g_a g_b = g_u}} s[a] w[b]$$

(ii)
$$\forall u \in V, (w *_{\mathbf{M}} s)[u] = \sum_{\substack{v \in \mathcal{D}_g \\ s.t. \ g(v) = u}} w[g] \, s[v]$$

We obtain an equivariance characterization similar to Theorem 54 and Theorem 61, under the mild condition that Υ is at least domain-symmetric if it is not abelian.

Theorem 83. Characterization by equivariance to Υ

Let a groupoid Υ acting on V^0 . Supposedly for (i), $\Upsilon \stackrel{\varphi}{=} V$, and for (ii), Υ is abelian.

- 1. Then,
 - (i) partial φ -convolution right operators are equivariant to Υ ,
 - (ii) partial M-convolution left operators are equiv to Υ .
- 2. Conversely,
 - (i) if Υ is domain-symmetric, linear transformations of $\mathcal{S}(V)$ that are equivariant to Υ are partial φ -convolution right operators,
 - (ii) linear transformations of $\mathcal{S}(V)$ that are equivariant to Υ are partial M-convolution left operators.

Proof. (i) (a) Direct sense:

Using the symmetrical expressions, and the fact that $\forall r, r[0] = 0$, we have

$$(f_w \circ g(s))[u] = \sum_{\substack{(g_a, g_b) \in \mathcal{D} \\ s.t. \ g_a g_b = g_u}} g(s)[a] w[b]$$
$$= \sum_{\substack{(g_a, g_b) \in \mathcal{D} \\ s.t. \ g_a g_b = g_u}} s[g^{-1}(a)] w[b]$$

$$= \sum_{\substack{(g_a,g_b) \in \mathcal{D} \\ s.t. \ (g,g_a) \in \mathcal{D} \\ s.t. \ gg_ag_b = g_u}} s[a] w[b]$$

$$= \sum_{\substack{(g_a,g_b) \in \mathcal{D} \\ s.t. \ (g,g_a) \in \mathcal{D} \\ s.t. \ (g,g_a) \in \mathcal{D} \\ s.t. \ gag_b = g^{-1}g_u = g_{\varphi(g^{-1}g_u)} = g_{g^{-1}(u)}}} s[a] w[b]$$

$$= f_w(s)[g^{-1}(u)]$$

$$= (g \circ f_w(s))[u]$$

(b) Converse:

Let $v \in V$. Denote $e^r_{g_v} = g^{-1}_v g_v$ the right identity element of g_v , and $e^r_v = \varphi(e^r_{g_v})$. We have that

$$g_v(e_v^r) = v$$

So, $\delta_v = g_v(\delta_{e_v^r})$

Let $f \in \mathcal{L}(\mathcal{S}(V))$ that is equivariant to Υ , and $s \in \mathcal{S}(V)$. Thanks to the previous remark we obtain that

$$f(s) = \sum_{v \in V} s[v] f(\delta_v)$$

$$= \sum_{v \in V} s[v] f(g_v(\delta_{e_v^r}))$$

$$= \sum_{v \in V} s[v] g_v(f(\delta_{e_v^r}))$$

$$= \sum_{v \in V} s[v] g_v(w_v)$$
(47)

where $w_v = f(\delta_{e_v^r})$. In order to finish the proof, we need to find w such that $\forall v \in V, g_v(w) = g_v(w_v)$.

Let's consider the equivalence relation \mathcal{R} defined on $V \times V$ such

that:

$$a\mathcal{R}b \Leftrightarrow w_a = w_b$$

$$\Leftrightarrow e_a^r = e_b^r$$

$$\Leftrightarrow g_a^{-1}g_a = g_b^{-1}g_b$$

$$\Leftrightarrow (g_b, g_a^{-1}) \in \mathcal{D}$$

$$\Leftrightarrow (g_a^{-1}, g_b) \in \mathcal{D}$$
(48)

with (48) owing to the fact that Υ is domain-symmetric.

Given $x \in V$, denote its equivalence class $\mathcal{R}(x)$. Under the hypothesis of the axiom of choice (Zermelo, 1904) (if V is infinite), define the set \aleph that contains exactly one representative per equivalence class. Let $w = \sum_{n \in \aleph} w_n$. Then V is the disjoint union $V = \bigcup_{n \in \aleph} \mathcal{R}(n)$ and (47) rewrites:

$$\forall u \in V, f(s)[u] = \sum_{n \in \aleph} \sum_{v \in \mathcal{R}(n)} s[v] g_v(w_n)[u]$$

$$= \sum_{n \in \aleph} \sum_{v \in \mathcal{R}(n)} s[v] w_n[g_v^{-1}(u)]$$

$$= \sum_{n \in \aleph} \sum_{v \in \mathcal{R}(n)} s[v] w[g_v^{-1}(u)]$$

$$= (s *_{\varphi} w)[u]$$

$$(49)$$

where (49) is obtained thanks to (48).

(ii) With symmetrical expressions, it is clear that the convolution is abelian, if and only if, Υ is abelian. Then (i) concludes.

Inclusion of EC and LP

Similarly to the construction in Section 2.3, partial convolutions can define EC, EC*, LP and LP* counterparts with the same characterizations by Cayley subgraphs whose vertex sets are zero-closure of groupoids, and other similar results.

Limitation of partial convolutions

However, because of the groupoid associativity, if $g \in \Psi_{\text{EC}}^*(G)$, then, any $v \in V$ s.t. g(u) = v would be constrained to allow to be acted by every h s.t. $(h,g) \in \mathcal{D}$. That is, partial convolutions alleviates the limitations of convolutions based on groups by partitioning the graph into parts onto which it behaves like if it was based on local groups. This can be seen as a limitation since we might want the convolution to behave similarly on the whole graph.

2.4.4 Construction of path convolutions

To answer the limitation of partial convolutions, given $\mathcal{U} \subset \Psi_{\text{EC}}^*(G)$, we explore the idea of proceeding with a foliation of each $g \in \mathcal{U}$ into pieces, each piece corresponding to an edge $e \in E$, and together generating another groupoid with a different associativity law, as follows.

Definition 84. Path groupoid

Let $\mathcal{U} \subset \Psi_{\text{EC}}^*(G)$. The *path groupoid* generated from \mathcal{U} , denoted $\mathcal{U} \ltimes G$, with composition rule \mathcal{D}_{\ltimes} , is the groupoid obtained inductively with:

1.
$$\mathcal{U} \ltimes_1 G = \{(g, v) \in \mathcal{U} \times V, v \in \mathcal{D}_g\} \subset \mathcal{U} \ltimes G$$

2.
$$((g_n, v_n) \cdots (g_1, v_1), (h_m, u_m) \cdots (h_1, u_1)) \in \mathcal{D}_{\kappa} \Leftrightarrow h_m(u_m) = v_1$$

3.
$$((g_n, v_n) \cdots (g_1, v_1))^{-1} = (g_1^{-1}, g_1(v_1)) \cdots (g_n^{-1}, g_n(v_n))$$

We call path its objects. Given a length $l \in \mathbb{N}$, we denote $\mathcal{U} \ltimes_l G$ the subset composed of the paths that are the composition of exactly l paths of $\mathcal{U} \ltimes_1 G$.

Remark. This groupoid construction is inspired from the field of operator algebra where partial action groupoids have been extensively studied, *e.g.* Nica, 1994; Exel, 1998; Li, 2016.

Such groupoids usually come equipped with source and target maps. We also define the path map.

Definition 85. Source, target and path maps

Let a path groupoid $\mathcal{U} \ltimes G$. We define on it the *source map* α the *target map* β and the *path map* γ as:

$$\begin{cases} \alpha : (g_n, v_n) \cdots (g_1, v_1) \mapsto v_1 \in V \\ \beta : (g_n, v_n) \cdots (g_1, v_1) \mapsto g_n(v_n) \in V \\ \gamma : (g_n, v_n) \cdots (g_1, v_1) \mapsto g_n g_{n-1} \dots g_1 \in \Psi^*(V^0) \end{cases}$$

Remark. Note that the path groupoid can be seen as the local structure of the partial transformation groupoid.

Lemma 86.

Note the following properties:

1.
$$(p,q) \in \mathcal{D}_{\bowtie} \Leftrightarrow \alpha(p) = \beta(q)$$

2.
$$\alpha(p) = \beta(p^{-1})$$

3.
$$e_p^l=pp^{-1}=(\operatorname{Id},\beta(p))$$
 and $e_p^r=p^{-1}p=(\operatorname{Id},\alpha(p))$

4. γ is a groupoid partial action. We will denote γ_p instead of $\gamma(p)$.

Remark. Note that this time we won't use the notation p(v) for $\gamma_p(v)$ for clarity.

One of the key object of our construction is the use of φ -equivalence in order to transform a sum over a group(oid) of (partial) transformations, into a sum over the vertex set. With the current notion of path groupoid, searching for something similar amounts to searching for a graph traversal.

Definition 87. Traversal set

Let a graph $G = \langle V, E \rangle$ that is connected. A *traversal set* is a pair $(\mathcal{U}, \mathcal{T})$ of EC partial transformations subsets $\subset \Psi_{\text{EC}}^*(G)$, such that

- 1. \mathcal{U} is *edge-deterministic*, in the sense that an edge can only correspond to a unique g, *i.e.* $\forall g, h \in \mathcal{U} : \exists v \in V, g(v) = h(v) \Rightarrow g = h$
- 2. The EC partial transformations of \mathcal{T} are restrictions of those of \mathcal{U} , i.e. $\forall g \in \mathcal{U}, \exists ! h \in \mathcal{T}, \begin{cases} \mathcal{D}_h \subset \mathcal{D}_g \\ \forall v \in \mathcal{D}_h, h(v) = g(v) \end{cases}$ (equivalently, $\mathcal{T} \ltimes G$ is a path subgroupoid of $\mathcal{U} \ltimes G$ s.t. $|\mathcal{T}| = |\mathcal{U}|$)
- 3. The subgraph $G_{\mathcal{T}} = \langle V, \mathcal{T} \ltimes_1 G \rangle$ is a spanning tree of G.

We denote $(\mathcal{U}, \mathcal{T}) \in \text{trav}(G)$, and denote by r the root of $G_{\mathcal{T}}$. For $p \in \mathcal{T} \ltimes G \subset \mathcal{U} \ltimes G$, we denote $\gamma_p^{\mathcal{T} \ltimes G}$ and $\gamma_p^{\mathcal{U} \ltimes G}$ its path maps.

Remark. The assumption that the graph G is connected does not lose generality as the construction can be replicated to each connected component in the general case.

A traversal set $(\mathcal{U}, \mathcal{T})$ defines a φ -equivalence between the α -fiber of the root r and the vertex set V as follows.

Lemma 88. Path φ -Equivalence

Let $(\mathcal{U}, \mathcal{T}) \in \text{trav}(G)$. Given $v \in V$, there exists a unique $p_v \in \mathcal{T} \ltimes G$ such that $\alpha(p_v) = r$ and $\beta(p_v) = v$. Denote $\mathcal{T} \ltimes^r G = \alpha_{\mathcal{T} \ltimes G}^{-1}\{r\}$. We can do the following construction:

- 1. Define $\varphi: p_v \mapsto v$.
- 2. Define $(p_v, p_u) \mapsto p_v^u \in \mathcal{U}^0 \ltimes^r G$ such that the sequence of partial transformations of p_v^u and p_v are the same (i.e. $\gamma_{p_v^u}^{\mathcal{U}^0 \ltimes G} = \gamma_{p_v}^{\mathcal{U} \ltimes G}$), and the source of p_v^u is the target of p_u (i.e. $\alpha(p_v^u) = \beta(p_u) = u$)
- 3. Define the external composition $p_v p_u = p_v^u p_u \in \mathcal{U}^0 \ltimes^r G$.

Then $\varphi: \alpha_{\mathcal{T} \ltimes G}^{-1}\{r\} \to V$ is a bijective partial equivariant map.

Proof. Bijectivity is a consequence of the spanning tree structure of \mathcal{T} . Equivariance because $\gamma_{p_v}(u) = \gamma_{p_v}\gamma_{p_u}(r) = \gamma_{p_vp_u}(r) = \varphi(p_vp_u)$.

We can now define the convolution that is based on a path groupoid.

Definition 89. Path convolution

Let $(\mathcal{U}, \mathcal{T}) \in \operatorname{trav}(G)$. The *path convolution* is the partial convolution based on the path subgroupoid $\mathcal{T} \ltimes G$, which uses the groupoid partial action $\gamma := \gamma^{\mathcal{U}^0 \ltimes G}$ of the embedding groupoid zero-closure $\mathcal{U}^0 \ltimes G$.

(i) In what follows are the three expressions of the path φ -convolution for signals $s_1, s_2 \in \mathcal{S}(V)$, and $u \in V$:

$$(s *_{\varphi} w) = \sum_{v \in V} s[v] \gamma_{p_v}(w)$$

$$= \sum_{\substack{p \in \mathcal{T} \ltimes G \\ s.t. \ \alpha(p) = r}} s[\varphi(p)] \gamma_p(w)$$

$$(s *_{\varphi} w)[u] = \sum_{\substack{(a,b) \in V \\ s.t. \ \gamma_{p_a}(b) = u}} s[a] w[b]$$

(ii) The mixed formulations with $w \in \mathcal{S}(\mathcal{T} \ltimes G)$ are:

$$(w *_{\mathbf{M}} s) = \sum_{\substack{p \in \mathcal{T} \ltimes G \\ s.t. \ \alpha(p) = r}} w[p] \gamma_p(s)$$
$$(w *_{\mathbf{M}} s)[u] = \sum_{\substack{(p,v) \in \mathcal{T} \ltimes G \times V \\ s.t. \ \alpha(p) = r \\ s.t. \ \gamma_p(v) = u}} w[p] s[v]$$

Remark. The role of \mathcal{T} is to provide a φ -equivalence. The role of \mathcal{U} is to extend every partial transformation $\gamma_g^{\mathcal{T} \ltimes G}$ to the domain of its unrestricted counterpart $\gamma_q^{\mathcal{U} \ltimes G}$.

Theorem 83 also holds for path groupoids, except that the domain-symmetric condition of 2.(i) is not needed.

Theorem 90. Characterization by equivariance to $\mathcal{U} \ltimes G'$ s action Let $(\mathcal{U}, \mathcal{T}) \in \operatorname{trav}(G)$.

- (i) The class of linear transformations of S(V) that are equivariant to the path actions of $U \ltimes G$ is exactly the path φ -convolution right-operators;
- (ii) in the abelian case, they are also exactly the M-convolution leftoperators.

Proof. Instead of the domain-symmetric condition that was used in the proof of the converse of Theorem 83 (2.(i)), we use the fact that any vertex can be reached with an action from the root of the spanning tree of the traversal set. Indeed, given $v \in V$, as we have $\gamma_{p_v}(r) = v$, then $\gamma_{p_v}(\delta_r) = \delta_v$. Therefore, by developing a linear transformation f(s) on the Dirac family, and commuting f with γ_{p_v} , we obtain that $f(s) = s *_{\varphi} w$, where $w = f(\delta_r)$. The rest of the proof is similar to that of Theorem 83.

Remark. Note that $\mathcal{U} \ltimes V'$ s action is almost the same as the groupoid partial action of $\Upsilon = \langle \mathcal{U} \rangle$ (only "almost" because not all combinations of partial transformations might exist in the paths).

Even though $\mathcal{U} \ltimes V$ alleviates the limitations of Υ we discussed earlier, it introduces another one: the zero-extension of the partial actions p_v^u can be harsh for some graphs with p_v^u often being equal to the zero of the groupoid (since the edge sequence of p_u must be walkable from v). However, there would be less degenerated scenarios for EC* operators counterparts supported on paths of small length. On the other hand, it can be preferable to avoid the degenerated cases and stick to partial actions or strategies discussed in Section 2.4.1.

2.5 Conclusion

In this chapter, we constructed convolutions on graph domains. We first saw that classical convolutions are in fact the class of linear transformations of the signal space that are equivariant to translations. For signals defined on graph domains, there is no natural definition of translations. Therefore, we adopted a more abstract standpoint and considered in the first place any kind of transformation of the vertex set V. Hence, given a group Γ acting on V, we constructed the class of linear transformations of the signal space that are equivariant to it. This provided us with an expression of a convolution based on this subgroup, and a bijective equivariant map between Γ and V, in order to transport a sum over Γ into a sum over V. We also proposed a simpler expression in the abelian case. Then, we introduced the role of the edge set E, and we constrained Γ by it. This leads us to define two types of properties. One is about constraining the transformations to follow the edges, and the other one is about preserving locality through them. For each property, we obtain characterizations of graphs that admit convolutions that bear them. We analyzed intrinsic properties of the constructed convolution operator, namely locality and weight sharing. We also discussed operators with a smaller kernel, as they are more practical and simpler to construct. Finally, we explored avenues to overcome the limitation that groups aren't well representative of symmetries on some graphs. So we extended the previous construction with two types of groupoids, and we obtained constructions for which the characterization by equivariance also holds. We saw that they can handle degenerated cases, but the expressions of the obtained convolutions can be degenerated as well.

Chapter 3

Deep learning on graph domains

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Chapter overview

Our goal in this chapter is to understand how neural networks can be extended to other domains than what they were intended for. To this end, in Section 3.1, we make an effort to interpret the linear algebra that underpins a layer to build our intuition. First we state the obvious by explaining further the interpretation of tensor space as a neural space, as well as how to juggle between tensors and signals. Then, we propose a representation based on graphs. Between two layers, a propagation graph explains how the propagation is done. On the input layer, the neurons can have an underlying graph structure. We show a relation between these two graphs, obtained if and only if the local receptive fields of the neurons are intertwined. By introducing the notion of weight sharing in our analysis, we discover that a layer on any domain can be expressed by a linear ternary operation, that we call neural contraction. Its operands are the input signal X, the weight kernel Θ , and the weight sharing scheme S. We denote $\Theta S \hat{X}$. We study it in Section 3.2. We see that it is generic in comparison with related works, and propose an efficient implementation. With an experiment based on it, we see how exploiting symmetries is beneficial, which justifies the use of convolutions. Through other experiments, we explore ideas based on randomizations to apply it on general graphs. In Section 3.3, we study the effect of learning how the weights are shared, which amounts to learn both S and Θ . We explore this avenue for graph domains, with experiments on grids, covariance graphs and on citation networks. Finally, in Section 3.4, we investigate an example of a CNN architecture used for graph signals. The convolution is based on translations on graphs which define the weight sharing scheme S of the convolutional layer. We present the model of translations and the approximation we use, the subsampling layer, the data augmentation, and experiments on grid graphs or graphs resembling to grids.

3.1 Layer representations

Let $\mathcal{L}=(g,h)$ be a neural network layer, where $g:I\to O$ is its linear part, $h:O\to O$ is its activation function, I and O are its input and output spaces, which are tensor spaces.

3.1.1 Neural interpretation of tensor spaces

Recall from Definition 1 that a tensor space has been defined such that its canonical basis is a Cartesian product of canonical bases of vector spaces. Let $I = \bigotimes_{k=1}^p \mathbb{V}_k$ and $O = \bigotimes_{l=1}^q \mathbb{U}_l$. Their canonical bases are denoted $\mathbb{V}_k = (\mathbb{V}_k^1, \dots, \mathbb{V}_k^{n_k})$ and $\mathbb{U}_l = (\mathbb{U}_l^1, \dots, \mathbb{U}_l^{n_l})$.

Remark. Note that a tensor space is isomorphic to the signal space defined over its canonical basis.

More precisely, we have the following relation.

Lemma 91. Relation between tensor and signal spaces

Let V, U be vector spaces, and v, u be their canonical bases. Let T be a tensor space. \otimes and \times denote tensor and Cartesian products. Then,

- (i) $\mathbb{V} \cong \mathcal{S}(\mathbb{V})$
- (ii) $\mathbb{V} \otimes \mathbb{U} \cong \mathcal{S}(\mathbb{V} \times \mathbb{U})$
- (iii) $\mathbb{V} \otimes \mathbb{T} \cong \mathcal{S}_{\mathbb{T}}(\mathbb{V})$

where $\mathcal{S}_{\mathbb{T}}$ are signals taking values in \mathbb{T} (and \mathcal{S} are real-valued signals).

Proof. (i) Given $x \in V$, define $\widetilde{x} \in \mathcal{S}(v)$ such that $\forall i, \widetilde{x}[v^i] = x[i]$. The mapping $x \mapsto \widetilde{x}$ is a linear isomorphism.

(ii)
$$\widetilde{x}[\mathbf{v}^i, \mathbf{u}^j] = x[i, j]$$

(iii)
$$\widetilde{x}[v^i] = x[i,:,\ldots,:]$$

Let $d \leq n_k$ and $e \leq n_l$. Define V and U as the Cartesian products $V = \times_{k=1}^d \mathbb{V}_k$ and $U = \times_{l=1}^e \mathbb{U}_l$. Thanks to Lemma 91, we can identify the input and output spaces as $I = \mathcal{S}(V) \otimes \bigotimes_{k=d+1}^p \mathbb{V}_k$ and $O = \mathcal{S}(U) \otimes \bigotimes_{l=e+1}^q \mathbb{U}_l$. As $\mathcal{S}(\mathbb{V}) \otimes \mathbb{T} = \mathcal{S}_{\mathbb{T}}(\mathbb{V})$, an object of V or U can be interpreted as the representation of a *neuron* which can take multiple values.

In what follows, without loss of generality, we will make the simplification that a neuron can only take a single value (we do not consider input channels and feature maps yet). We'll thus consider that $I = \mathcal{S}(V)$ and $O = \mathcal{S}(U)$, where V is the set of *input neurons*, and U is the set of *output neurons*.

3.1.2 Propagational interpretation

Let $\mathcal{L} = (g, h)$, recall that $g : \mathcal{S}(V) \to \mathcal{S}(U)$ is characterized by a connectivity matrix W such that, g(x) = Wx.

Remark. Using the mapping defined in the proof of Lemma 91, for notational convenience, we'll abusively consider x as a vector (eventually reshaped from a tensor), and W as an object of a binary tensor product for its indexing (i.e. W[u, v] := W[i, j] where $u = u^i$ and $v = v^j$).

Definition 92. Propagation graph

The propagation graph $P = \langle (V, U), E_P \rangle$ of a layer $\mathcal{L} = (W, h)$ is the bipartite graph that has the connectivity matrix W for bipartite adjacency matrix.

The propagation graph defines an input topological space \mathcal{T}_V , and an output topological space \mathcal{T}_U .

Definition 93. Topological space

A *topological space* is a pair $\mathcal{T} = (X, \mathcal{O})$, where X is a set of points, \mathcal{O} is a set of sets that is closed under intersection (the *open sets*), and such that every point $x \in X$ is associated with a set $\mathcal{N}(x) \in \mathcal{O}$, called its *neighborhood*.

Hence, the *neural topologies* \mathcal{T}_V and \mathcal{T}_U are defined as

1.
$$\mathcal{T}_V = (V, \mathcal{O}(U))$$
, with $\forall v \in V, \mathcal{N}(v) = \{u \in U, v \stackrel{P}{\sim} u\}$

2.
$$\mathcal{T}_U = (U, \mathcal{O}(V))$$
, with $\forall u \in U, \mathcal{N}(u) = \{v \in V, v \stackrel{P}{\sim} u\}$

In particular, given an output neuron $u \in U$, a neighborhood $\mathcal{N}(u)$ is also called a *local receptive field* (LRF), that we denote $\mathcal{R}(u)$.

3.1.3 Graph representation of the input space

Let's consider that the input neurons V have a (possibly edge-less) graph structure $G = \langle V, E \rangle$.

Definition 94. Underlying graph structure

An underlying graph structure of an input space V is simply a graph such that V is its vertex set.

For our definition of edge-constrained layer, we want that the LRF are constrained by the edges of the underlying graph. So we define it as follows.

Definition 95. Edge-constrained layer

A layer $\mathcal{L}: G = \langle V, E \rangle \to U$, is said to be *edge-constrained* (EC) if:

- 1. There is a one-to-one correspondence $\pi: V_{\pi} \to U$, where $V_{\pi} \subset V$.
- 2. Given an output neuron u, an input neuron v is in its receptive field, if and only if, v and the π -fiber of u are connected in G,

i.e.
$$\forall u \in U, v \in \mathcal{R}(u) \Leftrightarrow v \stackrel{E}{\sim} \pi^{-1}(u)$$

Figure 11 illustrate an example of a local receptive field of an edge-constrained layer.



Figure 11: An LRF of a vertex (in red) of an EC layer

Remark. Note that EC convolutions from Chapter 2 are indeed EC layers. Also, the following property is important.

Proposition 96. Let W the connectivity matrix of an EC layer w.r.t. a graph $G = \langle V, E \rangle$, which adjacency matrix is A. Then W is masked by A, i.e.

$$A[i,j] = 0 \Rightarrow W[i,j] = 0$$

Proof. Suppose A[i,j]=0, then $v_i\nsim v_j$, hence $v_i\notin \mathcal{R}(\pi(v_j))$ so W[i,j]=0.

We have the following characterization to determine which layers admit an underlying graph for which they are EC layers.

Proposition 97. EC Characterization with receptive fields

Let a layer $\mathcal{L}: V \to U$, $V_{\pi} \subset V$, and a one-to-one correspondence $\pi: V_{\pi} \to U$. There exists a graph $G = \langle V, E \rangle$ for which \mathcal{L} is EC, if and only if, the receptive fields are *intertwined* (i.e. $\forall a, b \in V_{\pi}, a \in \mathcal{R}(\pi(b)) \Leftrightarrow b \in \mathcal{R}(\pi(a))$).

Proof. (
$$\Rightarrow$$
) Thanks to $a \in \mathcal{R}(\pi(b)) \Leftrightarrow a \stackrel{E}{\sim} b \Leftrightarrow b \in \mathcal{R}(\pi(a))$

(\Leftarrow) If the receptive fields are intertwined, then the relation defined as $a \sim b \Leftrightarrow a \in \mathcal{R}(\pi(b))$ is symmetric, and thus can define an edge set.

Therefore, any layer that has its receptive fields intertwined, admits an *underlying* graph structure. For example, a 2-d convolution operator can be rewritten as an EC* convolution on a lattice graph, and as an EC convolution on a grid graph.

3.1.4 Novel ternary representation with weight sharing

Weight sharing refers to the fact that some parameters of the connectivity matrix W are equal, and stay equal after each learning iteration. In other words they are tied together. From a propagational point of view, this amounts to label the edges of the propagation graph P with weights, where weights can be used multiple times to label these edges. Supposed W is of shape $m \times n$ and there are ω weights in the kernel used to label the edges. Given a input neuron i and an output neuron j, the edge labeling can be expressed as:

$$W[j,i] = \theta[h] = \theta^T a \tag{50}$$

where θ , the weight kernel, is a vector of size ω and a is a one-hot vector (full of 0s except for one 1) of size ω with a 1 at the index h corresponding

to the weight that labels the edge $i \sim j$; or a is the zero vector in case $i \nsim j$.

This equation (50) can be rewritten as a tensor contraction under Einstein summation convention, by noticing that a depends on i, j and by defining a tensor S such that a = S[:, i, j], as follows:

$$W_{ij} = \theta_k S^k_{ij} \tag{51}$$

Therefore, the linear part of \mathcal{L} can be rewritten as:

$$g(x)_j = \theta_k S^k_{\ j}{}^i x_i \tag{52}$$

If we consider that the layer \mathcal{L} is duplicated with input channels and feature maps, then θ , x and g(x) become tensors, denoted Θ , X and g(X). Usually, for stochastic gradient descent, X and g(X) are also expanded with a further rank corresponding to the batch size and we obtain:

$$g(X) = \Theta SX \text{ where } \begin{cases} W_{pq}^{ij} = \Theta_{pq}^{\ k} S_k^{ij} \\ g(X)_{jq}^{\ b} = W_{jq}^{ip} X_{ip}^{\ b} \end{cases}$$
(53)

index	size	description
$\overline{}$	n	input neuron
j	m	output neuron
p	N	input channel
q	M	feature map
k	ω	kernel weight
b	B	batch instance

Table 2: Table of indices

Since the multiplicative expression ΘSX is written regardless of the ordering of the tensors ranks and is defined by index juggling, we will write instead $\widehat{\Theta SX}$ to avoid confusion. $\widehat{\cdots}$ is a ternary operator which

we will call *neural contraction*. Note that it is associative and commutative. This can be seen by the index symmetry of (54), which rewrites (53), and where the sum symbols Σ and scalar values commute:

$$\widehat{\Theta SX}[j,q,b] = \sum_{k=1}^{\omega} \sum_{p=1}^{P} \sum_{i=1}^{n} \Theta[k,p,q] S[k,i,j] X[i,p,b]$$
 (54)

Definition 98. Ternary representation, neural contraction

The ternary representation of a layer $\mathcal{L}: X \mapsto Y$, with activation function h, is the equation $Y = h\left(\widehat{\Theta SX}\right)$, where the neural contraction $\widehat{\cdots}$ is defined by (54), Θ is the weight kernel, and S is called the weight sharing scheme.

Remark. In (50), we defined a = S[:, i, j] as a one-hot vector when $i \sim j$, as its role is to select a weight in $\theta = \Theta[p, q, :]$. However, a can also do this selection *linearly*, so in fact it is not necessarily a one-hot vector.

Figure 12 depicts an example of how the equation (53) labels the edges of P.

The ternary representation uncouples the roles of Θ and S in W, and is the most general way of representing any kind of partially connected layer with weight sharing.

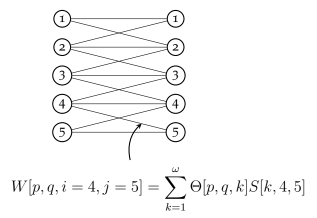


Figure 12: Example of a propagation graph P for a given input channel p and feature map q. The edge $4 \sim 5$ is labelled with a linear combination of kernel weights from $\Theta[p,q,:]$. In the usual case, S[k,4,5] is a one-hot vector that selects a single kernel weight: $\exists h, W[p,q,4,5] = \Theta[p,q,h]$.

3.2 Study of the ternary representation

In this section, we study the ternary representation, which is the general representation with weight sharing we obtained above. We already saw that it is linear, associative and commutative.

3.2.1 Genericity

The ternary representation can represent any kind of layer. We explain below how to obtain standard ones. See Table 2 and Table 3 for a description of the indices, notations and shapes.

- ightharpoonup To obtain a fully connected layer, one can choose $\omega=nm$ such that the slices S[:,i,j] constitute every possible one-hot vectors.
- \gt To obtain a convolutional layer, one can choose ω to be the size of the kernel. S would contain one-hot vectors. A stride \gt 1 can be

obtained by removing the corresponding dimensions. If the convolution is a classical convolution, or more generally, is characterized by a Cayley subgraph (see Chapter 2), then S would be circulant along the input neurons rank in the canonical basis.

- To obtain a graph convolutional layer (Kipf and Welling, 2016, see description in Section 1.3.5), one can choose B=1, $\omega=1$, and S (viewed as a matrix) to be the normalized adjacency matrix \widetilde{A} .
- \triangleright Other variants of GCN can also be obtained, for instance to obtain Chebychev filters (Defferrard et al., 2016), S would be $\sum_{i=0}^{k} T_i(\widetilde{L})$.
- To obtain a graph attention layer (Velickovic et al., 2017, see description in Section 1.3.5), one can concatenate K graph convolutional layers, where K is the number of attention heads, and with \widetilde{A} filled with the learned attention coefficients. Instead of concatenation, one could also choose $\omega = K$, and given an attention head k, the slices S[k,:,:] would be matrices containing the coefficients.
- > A topology-adaptive graph convolution layer (Du et al., 2017, see description in Section 1.3.5) is a neural contraction layer for which S contains the powers of \widetilde{A} along the first rank.
- A mixture model convolutional layer (Monti et al., 2016, equations (9) and (10)) is a neural contraction layer for which S contains the values of the weighting functions *i.e.* $S[k, i, j] = w_k(u[i, j])$.
- > A generalized convolution (Vialatte et al., 2016, equations (1) and (10)) is a neural contraction layer for which S is the allocation tensor A^e which has the sparse priors mentioned in the next subsection.
- ➤ Any partially connected layer with (or without) weight sharing can be obtained with appropriate construction of *S*.

3.2.2 Sparse priors for the classification of signals

In the more general case, the scheme S is a real-valued tensor, which may introduce more weights in the layer. However, for Euclidean convolutions, S is already determined by the Euclidean convolution operator and does not count for more memory. To imitate this property, we can focus on the class of layer with these sparse priors:

- 1. Given a couple of neuron indices (i, j), S[:, i, j] is either a one-hot vector, or the zero vector, as in (50).
- 2. Given an output neuron index j, a weight $\theta[h]$ can appear only once in its LRF.

Remark. For these sparse priors we consider the case of classification of signals. In the case of semi-supervised node classification, S would only be sparse between the rank indexed by i and the one indexed by j, but not on the other rank.

Therefore, we can describe S with a weight assignment table T, which is a matrix such that

$$T[i,j] = \begin{cases} h \text{ if } \exists! h, \theta^T S[:,i,j] = \theta[h] \\ 0 \text{ else} \end{cases}$$
 (55)

In turn, T can be described by the set of transformations (or partial transformations), defined as

$$g_h(j) = i \Leftrightarrow T[i, j] = h \tag{56}$$

where each (partial) transformation g_h corresponds to the weight indexed by h.

Without loss of generality, let us reduce to the simple case where B=P=Q=1. The neural contraction (54) rewrites as:

$$\widehat{\theta Sx}[j] = \sum_{k=1}^{\omega} \sum_{i=1}^{n} \theta[k] S[k, i, j] x[i]$$
(57)

$$=\sum_{h=1}^{\omega}\theta[h]\,x[g_h(j)]\tag{58}$$

which for EC layers, can be rewritten as an EC* convolution of graph signals as studied in Chapter 2, since the (partial) transformations are invertible thanks to the sparse priors. Conversely, given a convolution of graph signals (formulated as $*_{\varphi}$ or $*_{\mathtt{M}}$), we can derive an assignment table T and its corresponding scheme S to obtain a neural contraction.

3.2.3 Efficient implementation under sparse priors

Remark. This section applies in supervised settings but not in semi-supervised setting for which it is best to use sparse functions of common deep learning libraries.

What is the fastest way to compute $\widehat{\Theta SX}$?

As the equation (53) is associative and commutative, there are three ways to start to calculate it: with ΘS , SX, or ΘX , which we will call *middle-stage tensors*. The computation of a middle-stage tensor is the bottleneck to compute (53) as it imposes to compute significantly more entries than for the second tensor contraction. In Table 3, we compare their shapes. We refer the reader to Table 2 for the denomination of the indices.

tensor	shape
Θ	$\omega \times N \times M$
S	$\omega \times n \times m$
X	$n \times N \times B$
ΘS	$n \times m \times N \times M$
SX	$\omega \times m \times N \times B$
ΘX	$\omega \times n \times M \times B$
$\widehat{\Theta SX}$	$m \times M \times B$

Table 3: Table of shapes

We usually want to have $\omega \ll n$ and $\omega \ll m$, which means that we have weight kernels of small sizes (for example in the case of images, convolutional kernel are of size significantly smaller than that of the images). Also, the number of input channels N and of feature maps M are roughly in the same order, with N < M more often than the contrary. So in practice, the size of ΘS is significantly bigger than the size of S X and of ΘX , and the size of S X is usually the smallest.

How to exploit S sparsity?

Also, in usual supervised settings, S is sparse as S[:,i,j] are one-hot vectors. So computing SX should be faster than computing SX, provided we exploit the sparsity. Although S is very sparse as it contains at most a fraction $\frac{1}{w}$ -th of non-zero values, it is only sparse along the first rank, which makes implementation with sparse classes of common deep learning libraries use less parallelization than it can. However, we can benefit from the sparse priors mentioned in Section 3.2.2.

So we proceed differently. The idea is to use a non-sparse tensor $X_{\rm LRF}$ that has a rank that indexes the LRF, and another rank that indexes elements of these LRF, in order to lower the computation to a dense matrix multiplication (or a dense tensor contraction) which is already well optimized. This approach is similar to that proposed in Chellapilla et al., 2006, which

is also exploited in the cudnn primitives (Chetlur et al., 2014) to efficiently implement the classical convolution.

The LRF representation

In our case, it turns out that X_{LRF} can be exactly SX, as given fixed b, p, and j, SX[:,j,p,b] corresponds to entries of the input signal X[:,p,b] restrained to a LRF \mathcal{R}_j of size ω . Therefore,

$$\exists \operatorname{LRF}_j = [i_1, \dots, i_{\omega}] \text{ s.t. } SX[:, j, p, b] = X[\operatorname{LRF}_j, p, b]$$
(59)

The elements of LRF_j can be found by doing a lookup in the one-hot vectors of S, provided each kernel weight occurs exactly once in each LRF. We have:

$$R_j[k] = i_k \text{ s.t. } S[:, i_k, j][k] = 1$$
 (60)

This lookup needs not be computed each time and can be done beforehand. Finally, if we define $LRF = [LRF_1, ..., LRF_m]$, (59) gives:

$$SX = X[LRF,:,:] \tag{61}$$

The equation (61) is computed with only $\omega \times m$ assignations and can be simply implemented with automatic differentiation in commonly used deep learning libraries.

Benchmarks

To support our theoretical analysis, we benchmark three methods for computing the tensor contraction SX:

- naively using dense multiplication,
- using sparse classes of deep learning libraries,
- using the LRF based method we described above.

We run the benchmarks under the sparse priors mentioned in Section 3.2.2. For each method, we make 100 runs of computations of SX, with S and X being randomly generated according to the assumptions. In Table 4, we report the mean times. The values of the hyperparameters were each time n=m=N=M=B=256, and $\omega=4$. The computations were done on graphical processing units (GPU)¹.

Method	Mean time
Naive	950 ms
Sparse	413 ms
LRF	365 ms

Table 4: Benchmark (100 runs each)

We observe that the LRF method is faster, since the sparse implementation isn't optimized for this use case.

3.2.4 Influence of symmetries

In the case of images, or other signals over a grid, the grid structure of the domain defines the weight sharing scheme S of the convolution operation. For example, for a layer $\mathcal{L}: X \mapsto Y = h(\widehat{\Theta SX})$, and given fixed b, p, q, the classical convolution can be rewritten as

$$y[j] = h\left(\sum_{k=1}^{\omega} \theta[k] \sum_{i=1}^{n} S[k, i, j] x[i]\right)$$
(62)

$$= h\left(\sum_{k=1}^{\omega} \theta[k] x_{LRF}[k,j]\right) \tag{63}$$

Where $x_{LRF}[k,j]$ can be obtained by matching (63) with the expression given by the definition (see Definition 36). So, S[k,:,j] is a one-hot vector

¹The GPU was an Nvidia Geforce GTX1060, the CPU was an Intel Xeon E5-1630v4 at 3.70GHz with 4 cores. We used tensorflow and the function gather_nd for the assignations.

that specifies which input neuron in the LRF of y is associated with the k-th kernel weight, and the index of the 1 is determined by $x_{\tt LRF}[k,j]$. That is to say, in the ternary representation, Θ captures the information the layer has learned, whereas S captures how the layer exploits the symmetries of the input domain.

Visual construction

Visually, constructing S amounts to move a rectangular grid over the pixel domain, as depicted on Figure 13 where each point represents the center of a pixel, and the moving rectangle represents the LRF of its center j. Each of its squares represents a kernel weight $\theta[k]$ which are associated with the pixel $x_{\text{LRF}}[k,j]$ that falls in it.

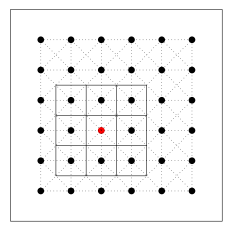


Figure 13: Weight assignement of convolutions on pixel domains

The case of images is very regular, in the sense that every pixels are regularly spaced out, so that obtained S is circulant along its last two ranks. This is a consequence of the translational symmetries of the input domain, which underlie the definition of the convolution, as seen in Chapter 2.

• What happens if we loose these symmetries?

To answer this question, we make the following experiment (Vialatte et al., 2016):

- 1. We distort the domain by moving the pixels randomly. The radial displacement is uniformly random with the angle, and its radius follows a Gaussian distribution $\mathcal{N}(0,\sigma)$.
- 2. Then we compare performances of shallow CNNs expressed under the ternary representation, for which S is constructed similarly than with the above visual construction, for different values of σ .

The visual construction of *S* on distorted domain is depicted by Figure 14.

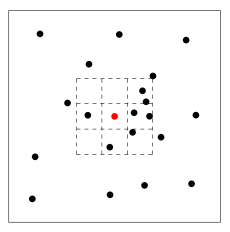


Figure 14: Weight assignement in generalized convolution on distorted domains

We run a classification task with standard hyperparameters on a toy dataset (we used MNIST, LeCun et al., 1998). The results are reported in Figure 15.

The bigger is σ , the less accurate are the symmetries of the input domain, up to a point where the ternary representation becomes almost equivalent to a dense layer. The results illustrate nicely this evolution, and stress out the importance of trying to leverage symmetries when defining new convolutions.

Moreover, they indicate that the ternary representation also allows to improve performances compared to using dense layers, providing we are able to create a relevant weight sharing scheme *S* to exploit symmetries.

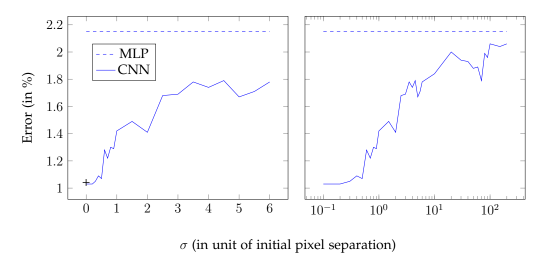


Figure 15: Error in function of the standard deviation σ , for generalized CNNs and an MLP, each with 500 weights.

For example, in the case the underlying graph structure of the input is embedded in a two-dimensional Euclidean space, S can be created just as in this experiment.

3.2.5 Experiments with general graphs

The neural contraction is not usable off-the-shelf for general graphs since the weight sharing scheme S needs to be specified. One strategy is to learn it alongside the weight kernel Θ , see Section 3.3. Another one consists in inferring it from the graph itself, see Section 3.4. But first let us try some ideas based on randomizations.

Supervised classification of graph signals

We test the idea to fill the schemes S with one-hot vectors randomly, but with respect to the sparse priors mentioned in Section 3.2.2. We apply neural contractions in a depthwise separable fashion similarly than for depthwise separable convolutions (Chollet, 2016). That is, we first propagate the neurons values on the propagation graph for each of the N input

feature maps independently. In the process, we use one different realization of S per input feature map. Then we apply a fully connected layer in the feature space (which is a convolution with trivial sliding window). Since this amounts to do M (one for each output feature map) learned weighted averages of N random realizations, we call that a Monte-Carlo (MC) module. We call $Monte-Carlo\ Networks$ (MCNet) the corresponding neural networks.

We replicated the experimental setup done by Defferrard et al., 2016 on the 20NEWS dataset. We refer the reader to Section 1.3.4 for the preliminary discussion on this experiment. We used the architecture depicted on Figure 16. It starts with 2 MC modules with 64 output feature maps, followed by a trivial convolution with 1 output feature map, then a FC layer with 500 hidden units and a final FC layer for classification. A residual connection is added between the input and the first FC layer. For the MC modules, we used the graph connecting 2 nearest neighbors. Therefore this architecture can be seen as a small random perturbation of the MLP from Table 1. Each layer except for the last is followed by 20% dropout. As usual, we use reLU activations except for the last that has softmax activation.

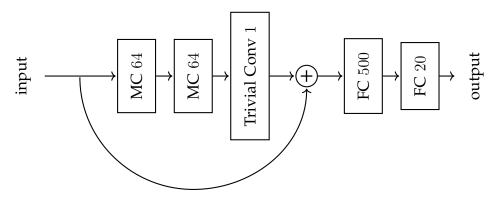


Figure 16: Diagram of the MCNet architecture used

After 100 runs of 10 epochs, MCNet obtained a mean accuracy of $70.74 \pm 2.19\%$, which is a gain that is statistically significant compared to ChebNet (65.76%,

see Table 1), but not against the former MLP ($71.46 \pm 0.08\%$). It is to be noted that the variance of MCNet is quite high and cherry picking the max instead of considering the mean improves the result to 72.62%, which is better than the MLP's max at 72.25%. Therefore, in some random realizations, MCNet is capable of leveraging the underlying graph structure (and being state-of-the-art).

This is the least to be expected. But when the quality of the graph is low, there may not be anything better to demonstrate. A dataset with a graph that suits well the underpining structure of the domain is often resembling a grid graph to some extent. Therefore we expect the model that will be presented in Section 3.4 to be more relevant in most cases.

Semi-supervised classification of nodes

A similar kind of randomization based idea that come in mind when transposing the question to the semi-supervised task is to use dropout on the edges of the graph. We call that *graph dropout*. We test this approach on citation networks, for which we set $\omega=1$ (*i.e.* one graph). The obtained architecture amounts to a GCN (Kipf and Welling, 2016) to which graph dropout is applied. We report and discuss our results in Section 3.3.6, Table 8, where we compare them with other models.

3.3 Learning the weight sharing scheme

3.3.1 Discussion

In the ternary representation $Y = h(\widehat{\Theta SX})$ of a layer \mathcal{L} , the weight kernel Θ is usually the only operand that is learned, and the role of S is to label the edges of the propagation graph P with these weights. Recall that, as noted after Definition 98, S needs not be sparse and composed of one-hot vectors. In that case, the labeling is done linearly as depicted previously in Figure 12. Therefore, the weight sharing scheme S can also be updated during the learning phase. This can be interpreted as learning a convolution-like operator on the underlying graph G (Vialatte et al., 2017).

Remark. When S does not have the sparse priors mentioned in Section 3.2.2, we must not have more parameters in S and Θ than in W, so that the weight sharing still makes sense. If we call l the number of edges in the resulting propagation graph P, then the former assumption requires $l\omega + \omega NM \leq lNM$ or equivalently $\frac{1}{\omega} \geq \frac{1}{NM} + \frac{1}{l}$. It implies that the number of weights per filter ω must be lower than the total number of filters NM and than the number of edges l, which is always the case in practice.

3.3.2 Experimental settings

In our experiments, we learn the scheme S and the kernel Θ simultaneously.

Constraints for supervised classification of graph signals

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 first rank (the rank that is contracted in the product ΘS).

$$\forall (k, i, j), \ S[k, i, j] \in [0, 1] \tag{64}$$

$$\forall (i,j), \ \sum_{k=1}^{\omega} S[k,i,j] = 1$$
 (65)

Therefore, the vectors on the first rank of S can be interpreted as performing a positive weighted average of the parameters in Θ . Also, we choose to aim our study at graph signals. Hence, we consider a graph $G = \langle V, E \rangle$ and that the layer is EC. For this reason, Proposition 96 tells us that the connectivity matrix W of the layer is masked by the adjacency matrix A. Therefore, S is also masked by A *i.e.* we impose the constraint that

$$A[i,j] = 0 \Rightarrow \forall k, S[k,i,j] = 0 \tag{66}$$

Constraints for semi-supervised classification of nodes

For the case of citation networks, we consider only one graph and thus we set $\omega=1$ so that S and Θ are matrices. Additionally, we impose rows of the scheme S to be normalized so that each node receives a normalized information from its neighborhood. In our experiments, we choose to use softmax normalization because it is of standard usage (but other normalizations could do as well).

Name

In (Vialatte et al., 2017), we used the term *Local Receptive Graph layers*, since Proposition 97 relates G with local receptive fields defined by W. However it was cumbersome so in this manuscript we will just call them *Graph Contraction Layer* in reference to the neural contraction (see Definition 98). We call *Graph Contraction Networks* (GCT) the corresponding neural networks. The main addition to the paper (Vialatte et al., 2017) from Section 3.3 is Section 3.3.6.

Initialization

In supervised settings, we introduce three types of initialization for the scheme S. The last two have the sparse priors mentioned in Section 3.2.2:

- 1. uniform random: parameters of S are simply initialized with a uniform random distribution with limits as described by Glorot and Bengio, 2010.
- 2. random one-hot: one-hot vectors are distributed randomly on the S[:,i,j], with the constraint that for each LRF, a particular one-hot vector can only be distributed at most once more than any other.
- 3. circulant one-hot: one-hot vectors are distributed in a circulant fashion on the S[:,i,j], so that on Euclidean domains, the initial state of S correspond exactly to the weight sharing scheme of a standard convolution.

In semi-supervised settings, we initialize the scheme S using the normalized adjacency matrix \widetilde{A} to which we add a Gaussian noise $\mathcal{N}(0,\sigma)$, where the standard deviation σ is determined following Glorot and Bengio, 2010.

3.3.3 Experiments with grid graphs

We experiment GCTs on the MNIST dataset (see Section 1.3.3). We use a shallow architecture made of a single ternary layer with 50 feature maps, without pooling, followed by a FC layer of 300 neurons, 50% dropout, and terminated by a FC layer of 10 neurons with softmax activation. ReLu activations are used. Input layers are regularized by a factor weight of 10^{-5} (Ng, 2004). We optimize with ADAM (Kingma and Ba, 2014) up to 100 epochs and fine-tune (while S is frozen) for up to 50 additional epochs.

For the underlying graph structure, we consider a grid graph that connects each pixel to itself and its 4 nearest neighbors (or less on the borders). We also consider 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). We test the model under two setups: either the ordering of the node is unknown, and then we use random one-hot initialization for S; either an ordering of the node is known, and then we use circulant one-hot initialization for S which we freeze in this state. We use the number of nearest neighbors as for the dimension of the first 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. On MNIST, training architectures that learn S took about twice longer. Table 5 summarizes the obtained results. The ordering is unknown for the first result given, and known for the second result between parenthesis.

Conv ₅ x ₅	Grid ¹	$Grid^2$	$Grid^3$	
(0.87)	1.24(1.21)	1.02(0.91)	0.93(0.91)	
Grid ⁴ Grid ⁵		Grid ⁶	Grid ¹⁰	
0.90(0.87)	0.93(0.80)	1.00(0.74)	0.93(0.84)	

Table 5: Error rates (in %) on powers of the grid graphs on MNIST.

We observe that even without knowledge of the underlying Euclidean structure, grid GCTs obtain comparable performances as CNNs, and when the ordering is known, they match them. We also noticed that after training, even though the one-hot vectors used for initialization had changed to floating point values, their most significant dimension was almost always the same. That suggests there is room to improve the initialization

and the optimization.

In Figure 17, we plot the test error rate for various normalizations when using the square of the grid graph, as a function of the number of epochs of training, only to find that they have little influence on the performance but sometimes improve it a bit. Thus, we will treat them as optional hyperparameters.

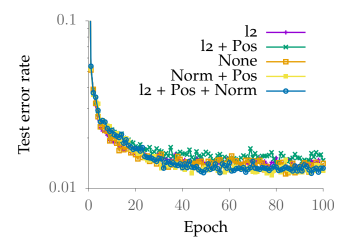


Figure 17: Evolution of the test error rate when learning MNIST using the square of a grid graph and for various normalizations, as a function of the epoch of training. The legend reads: "l2" means ℓ_2 normalization of weights is used (with weights 10^{-5}), "Pos" means parameters in S are forced to being positive, and "Norm" means that the ℓ_1 norm of each vector in the first dimension of S is forced to 1.

3.3.4 Experiments with covariance graphs

As underlying graph structure, we use a thresholded covariance matrix obtained by using all the training examples. We choose the threshold so that the number of remaining edges corresponds to a certain density p (5x5 convolutions correspond approximately to a density of p = 3%). We also infer a graph based on the k nearest neighbors of the inverse of the values of this covariance matrix (k-NN). The latter two are using no prior

about the signal underlying structure. The pixels of the input images are shuffled and the same re-ordering of the pixels is used for every image. Dimension of the first rank of S is chosen equal to k and its weights are initialized random uniformly. GCTs are also compared with models obtained when replacing the first layer by a fully connected or convolutional one. Architecture used is the same as in the previous section. Results are reported on table 6.

MLP	Conv ₅ x ₅	Thresholded ($p = 3\%$)	k-NN ($k = 25$)
1.44	1.39	1.06	0.96

Table 6: Error rates (in %) on scrambled MNIST.

We observe that GCTs outperform the CNN and the MLP on scrambled MNIST. This is remarkable because that suggests it has been able to exploit information about the underlying structure.

3.3.5 Improved convolutions on shallow architectures

On CIFAR-10 (see Section 1.3.3), we made experiments on shallow CNN architectures and replaced convolutions by receptive graphs. We report results on a variant of AlexNet (Krizhevsky et al., 2012) using little distortion on the input that we forked from a tutorial of tensorflow (Abadi et al., 2015). It is composed of two 5x5 convolutional layers of 64 feature maps, with max pooling and local response normalization (Krizhevsky et al., 2012), followed by two fully connected layers of 384 and 192 neurons. On CIFAR-10, training architectures that learn S took about 2.5 times longer. We compare two different graph 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 freezed on the 125 last ones. Circulant one-hot initialization is used. These are weak classifiers for CIFAR-10 but they are enough to analyze the usefulness of the proposed layer. Experiments are run five times each. Means and standard deviations of accuracies are reported in table 7. "Pos" means parameters in S are forced to being positive, "Norm" means that the ℓ_1 norm of each vector in the third dimension of S is forced to 1, "Both" means both constraints are applied, and "None" means none are used.

Support	Learn S	Learn S None Pos		Norm	Both
Conv ₅ x ₅	No	/	/	/	86.8 ± 0.2
Conv5x5 Yes		87.4 ± 0.1	87.1 ± 0.2	87.1 ± 0.2	87.2 ± 0.3
Grid ²	Yes	87.3 ± 0.2	87.3 ± 0.1	87.5 ± 0.1	87.4 ± 0.1

Table 7: Accuracies (in %) of shallow networks on CIFAR-10.

The GCTs are able to outperform the corresponding CNNs by a small but statistically significant amount in a shallow architecture.

Learning the scheme *S* implies a memory overhead due to the increase in the number of weights of each layer, thus why we limited this experiment to a shallow architecture. An example of strategy to extend this experiment to deeper architectures is to tie the schemes of each layer together, or to reuse a same scheme (previously learned) for all layers.

3.3.6 Benchmarks on citation networks

We benchmark multiple graph convolutional methods on citation networks. We refer the reader to Section 1.3.3 for a description of the datasets, and to Section 1.3.5 for a review of the tested models. The models we benchmark are:

- Graph Convolution Network (GCN, Kipf and Welling, 2016),
- Graph Attention Network (GAT, Velickovic et al., 2017),
- Topology Adaptive GCN (TAGCN, Du et al., 2017),
- Addition of graph dropout to GCN (GCN*, from Section 3.2.5),
- Graph Contraction Network (GCT, this work).

To proceed, we forked the official code repository of Velickovic et al., 2017, in order to replicate their environmental setup, from which we implemented the other tested models. This entailed a few differences to be noted between our implementation of GCN and TAGCN from the original ones: a bias is added, the right products XW are computed first and are followed by 50% dropout, and the best model on the validation set is saved to be reused at test time. We also used 50% graph dropout for GAT, TAGCN, and GCT, except on the Pubmed dataset for which we saw that it was detrimental. The inputs of each layer undergo 50% dropout, and the value of graph dropout of GCN* is also 50%. Every model is composed of two hidden layers. GAT models uses 8 heads with 8 hidden units (which amounts to 64 hidden units) on the first layer, and 1 head on the second one (except on Pubmed where it also uses 8) of also 8 hidden units. The number of hidden units for other models was grid searched in $\{16, 32, 64\}^{1}$. The polynomial degree of TAGCN was 2. For comparison, we also ran the experiment with an MLP composed of 500 hidden neurons. The results are reported in Table 8.

¹Most of the time the value of 16 was selected, sometimes 32, and never 64.

Dataset	MLP	GCN	GAT	TAGCN	GCN*	GCT
Cora	58.8 ± 0.9	81.8 ± 0.9	83.3 ± 0.6	82.9 ± 0.7	83.4 ± 0.7	83.3 ± 0.7
Citeseer	56.7 ± 1.1	72.2 ± 0.6	72.1 ± 0.6	71.7 ± 0.7	72.5 ± 0.8	72.7 ± 0.5
Pubmed	72.6 ± 0.9	79.0 ± 0.5	78.3 ± 0.7	78.9 ± 0.5	78.2 ± 0.7	79.2 ± 0.4

Table 8: Mean accuracy (in %) and standard deviation after 100 runs

Even though GCT models performed best overall, the differences with the second bests are not statistically significant. In particular, GCN models enjoy a bump in performances in this setup compared to the experiments in the original paper. The addition of graph dropout put them back on par with the other models on CORA and Citeseer (they were already ahead on Pubmed). We also noticed that the results we obtained for TAGCN on Pubmed were significantly worse than claimed by the authors. We contacted them to check for details and are awaiting response, so we may amend this result in a future version of this document¹. It is worth noting that the MLP performed a lot worse as expected since it did not exploit the graph structure.

¹However it is possible that this is due to the experimental setup.

3.4 Inferring the weight sharing scheme

3.4.1 Methodology

As we saw in Section 3.2.4 (Figure 13), the classical convolution on grid graphs can be obtained visually by translating a rectangular window over the pixel domain. The idea of this section is to define similarly a convolution on graph domains, using a notion of translation defined on graphs (Pasdeloup et al., 2017b). The translations we use match Euclidean translations on 2D grid graphs (Grelier et al., 2016), and extend them on general ones, by preserving three simple key properties: injectivity, edge-constraint, and neighborhood-preservation, which we will detail later.

Given a set of graph translations $\mathcal{T} = \{t_i, i \in \{1, 2, ..., \kappa - 1\}\} \cup \{t_0 = \operatorname{Id}\}$, the corresponding convolution can be expressed using the same formalism we used in Chapter 2, as:

$$w * x = \sum_{t \in \mathcal{T}} w[t] t(x) \tag{67}$$

where x is a signal over vertices of a graph $G = \langle V, E \rangle$, and w is a signal defined on $\mathcal{T} \subset \Phi_{\text{EC}}(G)$. Depending on the algebraic nature of \mathcal{T} (that we will discuss later), the theoretical results obtained in Chapter 2 hold. Under the ternary representation, finding such translations amounts to infer the weight sharing scheme S.

By denoting $w_i = w[t_i]$, we obtain the more familiar expression:

$$\forall u \in V, w * x[u] = \sum_{i=0}^{\kappa-1} w_i x[t_i^{-1}(u)]$$
 (68)

This expression extends the definition of convolutions from grid graphs to general graphs, with the use of graph translations. We use it to obtain extended CNNs that can be applied on graphs. In this manuscript, we

coin the term *Translation-convolutional neural network* (TCNN) to refer to them, but they are the same as the extended convolutions we defined in Pasdeloup et al., 2017a.

Besides defining translation-convolution on general graphs, designing a TCNN also implies extending the other building blocks of classical CNNs. As a counterpart for pooling operations, we define a graph subsampling and apply strided translation-convolution. The translations of the subsampled graph are derived from those of the base graph to define translation-convolution at the downscaled level. We will also use a weak form of data augmentation using these translations. Figure 18 depicts the proposed methodology (Lassance et al., 2018).

3.4.2 Translations

Let a graph $G = \langle V, E \rangle$. We suppose the graph is connected, as conversely the process can be applied to each connected component of G. We denote by d the max degree of the graph and n = |V| the number of vertices.

Definition 99. Candidate-translation

A *candidate-translation* is a function $\phi: U \to V$, where $U \subset V$ and such that:

• ϕ is injective:

$$\forall v, v' \in U, \phi(v) = \phi(v') \Rightarrow v = v',$$

• ϕ is edge-constrained:

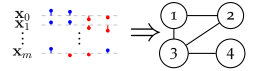
$$\forall v \in U, (v, \phi(v)) \in E,$$

• ϕ is strongly neighborhood-preserving:

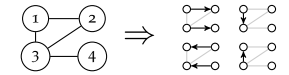
$$\forall v, v' \in U, (v, v') \in E \Leftrightarrow (\phi(v), \phi(v')) \in E.$$

The cardinal |V - U| is called the *loss* of ϕ . A translation for which V = U is called a *lossless* translation. Two candidate-translations ϕ and ϕ' are said to be *aligned* if $\exists v \in U, \phi(v) = \phi'(v)$. We define $N_r(v)$ as the set of vertices that are at most r-hop away from a vertex $v \in V$.

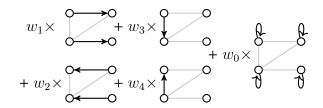
Step o (optional): infer a graph (see Pasdeloup et al., 2017a)



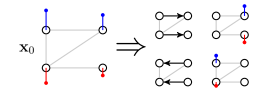
Step 1: infer translations



Step 2: infer the weight sharing scheme



Step 3: infer data augmentation



Step 4: infer subsampling and weight sharing scheme

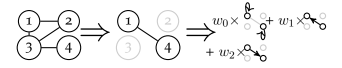


Figure 18: Outline of the proposed method

Definition 100. Translation

A *translation* in a graph G is a candidate-translation such that there is no aligned translation with a strictly smaller loss, or is the identity function.

If the graph is a 2D grid, obtained translations are exactly natural translations on images Grelier et al., 2016.

Because translations and candidate-translations need not be surjective on V, we introduce a zero vertex¹ denoted 0_V , such that any candidate translation $\phi: U \to V$ is extended as a function $\phi: V \to V \cup \{\bot\}$

$$\forall v \notin U, \phi(v) = 0_V \tag{69}$$

Finding translations is an NP-complete problem (Pasdeloup et al., 2017b). So in practice, we will first search locally. For this reason, we define local translations:

Definition 101. Local translation

A *local translation* of center $v \in V$ is a translation in the subgraph of G induced by $N_2(v)$, that has v in its definition domain.

With the help of local translations, we can construct proxies to global translations.

Definition 102. Proxy-translations

A family of *proxy-translations* $(\psi_p)_{p=0,..\kappa-1}$ initialized by $v_0 \in V$ is defined algorithmically as follows:

1. We place an indexing kernel on $N_1(v_0)$ i.e.

$$N_1(v_0) = \{v_0, v_1, ..., v_{\kappa-1}\}$$
 with $\forall p, \psi_p(v_0) = v_p$,

2. We move this kernel using each local translation ϕ of center v_0 :

$$\forall p, \psi_p(\phi(v_0)) = \phi(v_p),$$

¹Sometimes called *black hole.* (Grelier et al., 2016)

3. We repeat 2) from each new center reached until saturation. If a center is being reached again, we keep the indexing that minimizes the sum of losses of the local translations that has lead to it.

We explain this algorithm in more details in the next Section 3.4.3. A family of proxy-translations defines a translation-convolution as follows:

Definition 103. Translation-convolution layer

Let $(\psi_p)_{p=0,...,\kappa-1}$ be a family of proxy-translations identified on G s.t. $\psi_0 = \operatorname{Id}$. The translation-convolution layer $\mathcal{L}: x \mapsto y$ is defined as:

$$\forall v \in V, y[v] = h\left(\sum_{p=0}^{\kappa-1} w_p x[\psi_p(v)] + b\right)$$

where h is the activation function, b is the bias term, and with the convention that $x[0_V] = 0$.

3.4.3 Finding proxy-translations

We describe in three steps how we efficiently find proxy-translations.

First step: finding local translations

For each vertex $v \in G$, we identify all local translations using a bruteforce algorithm. This process requires finding all translations in all induced subgraphs. There are n such subgraphs, each one contains at most d local translations. Finding a translation can be performed by looking at all possible injections from 1-hop vertices around the central vertex to any vertex that is at most 2-hops away. We conclude that it requires at most $\mathcal{O}(ndd^{2(d+1)})$ elementary operations and is thus linear with the order of the graph. On the other hand, it suggests that sparsity of the graph is a key criterion in order to maintain the complexity reasonable.

Figure 19 depicts an example of a grid graph and the induced subgraph around vertex v_0 . Figure 20 depicts all obtained translations in the induced subgraph.

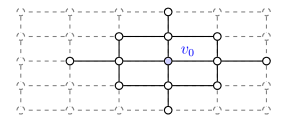


Figure 19: Grid graph (in dashed grey) and the subgraph induced by $N_2(v_0)$ (in black).

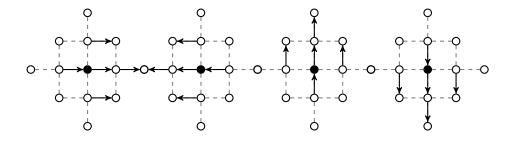


Figure 20: Translations (black arrows) in the induced subgraph (dashed grey) around v_0 (filled in black) that contains v_0 and only some of its neighbors.

Second step: move a small localized kernel

Given an arbitrary¹ vertex $v_0 \in V$, we place an indexing kernel on $N_1(v_0)$ i.e. $N_1(v_0) = \{v_0, v_1, ..., v_{\kappa-1}\}$. Then we move it using every local translations of center v_0 , repeating this process for each center that is reached for the first time. We stop when the kernel has been moved everywhere in the graph. In case of multiple paths leading to the same destination,

¹In practice we run several experiments while changing the initial vertex and keep the best obtained result.

we keep the indexing that minimizes the sum of loss of the series of local translations. We henceforth obtain an indexing of at most κ objects of $N_1(v)$ for every $v \in V$.

This process is depicted in Figure 21. Since it requires moving the kernel everywhere, its complexity is $\mathcal{O}(nd^2)$.

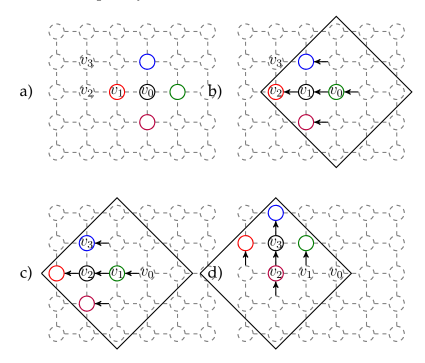


Figure 21: Illustration of the translation of a small indexing kernel using translations in each induced subgraph. Kernel is initialized around v_0 (a), then moved left around v_1 (b) using the induced subgraph around v_0 , then moved left again around v_2 (c) using the induced subgraph around v_1 then moved up around v_3 (d) using the induced subgraph around v_2 . At the end of the process, the kernel has been localized around each vertex in the graph.

Final step: identifying proxy-translations

Finally, by looking at the indexings obtained in the previous step, we obtain a family of proxy-translations defined globally on *G*. More precisely,

each index defines its own proxy-translation. Note that they are not translations because only the local properties have been propagated through the second step, so there can exist aligned candidates with smaller losses. Because of the constraint to keep the paths with the minimum sum of losses, they are good proxies to translations on G.

An illustration on a grid graph is given in Figure 22. The complexity is $\mathcal{O}(nd)$. Overall, all three steps are linear in n.

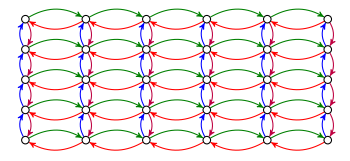


Figure 22: Proxy-translations in G obtained after moving the small kernel around each vertex. Each color corresponds to one translation.

3.4.4 Subsampling

Downscaling is a tricky part of the process because it supposes one can somehow regularly sample vectors. As a matter of fact, a non-regular sampling is likely to produce a highly irregular downscaled graph, on which looking for translations irremediably leads to poor accuracy, as we noticed in our experiments.

We rather define the translations of the strided graph using the previously found proxy-translations on G.

First step: extended convolution with stride \boldsymbol{r}

Given an arbitrary initial vertex $v_0 \in V$, the set of kept vertices $V_{\downarrow r}$ is defined inductively as follows:

- $V_{\downarrow r}^0 = \{v_0\},$
- $\forall t \in \mathbb{N}, V_{\downarrow r}^{t+1} = V_{\downarrow r}^t \cup \{v \in V, \forall v' \in V_{\downarrow}^t, v \notin N_{r-1}(v') \land \exists v' \in V_{\downarrow r}^t, v \in N_r(v')\}.$

This sequence is non-decreasing and bounded by V, so it eventually becomes stationary and we obtain $V_{\downarrow r} = \lim_t V_{\downarrow r}^t$. $V_{\downarrow r}$ is the set of output neurons of the extended convolution layer with stride r. Figure 23 illustrate the first downscaling $V_{\downarrow 2}$ on a grid graph.

Second step: convolutions for the strided graph

Using the proxy-translations on G, we move a localized r-hop indexing kernel over G. At each location, we associate the vertices of $V_{\downarrow r}$ with indices of the kernel, thus obtaining what we define as induced $_{\downarrow r}$ -translations on the set $V_{\downarrow r}$. In other words, when the kernel is centered on v_0 , if $v_1 \in V_{\downarrow r}$ is associated with the index p_0 , we obtain $\phi_{p_0}^{\downarrow r}(v_0) = v_1$. Subsequent convolutions at lower scales are defined using these induced $_{\downarrow r}$ -translations.

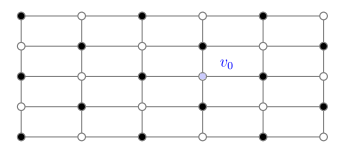


Figure 23: Downscaling of the grid graph. Disregarded vertices are filled.

3.4.5 Data augmentation

Once proxy-translations are obtained on G, we use them to move training signals, artificially creating new ones. Note that this type of data-augmentation is weaker than for images since no flipping, scaling or rotations are used.

3.4.6 Experiments

Matching CNNs on supervised image classification

On CIFAR-10 (see Section 1.3.3), our models are based on a variant of a deep residual network, namely PreActResNet18 (He et al., 2016b). We tested different combinations of graph support and data augmentation. For the graph support, we use either a regular 2D grid or either an inferred graph obtained by keeping the four neighbors that covary the most. In the second case, which correspond to scrambled CIFAR-10 (see Section 1.3.3), no structure prior have been fed to the process, so the results can be compared with those of Lin et al., 2015. We also report results obtained with ChebNet (Defferrard et al., 2016), where only convolutional layers have been replaced for comparison. Table 9 summarizes our results. In particular, it is interesting to note that results obtained without any structure prior (91.07%) are only 2.7% away from the baseline using classical CNNs on images (93.80%). This gap is even smaller (less than 1%) when using the grid prior. Also, on scrambled CIFAR-10 (case without prior) our method significantly outperforms the others.

Experiments on a fMRI dataset

We used a shallow network on the PINES dataset (see Section 1.3.3). The results reported on Table 10 show that our method was able to improve over CNNs, MLPs and other graph-based extended convolutional neural networks.

Support	MLP	CNN	Grid Graph		Covariance Graph
Support			ChebNet ^c	Proposed	Proposed
Full Data Augmentation	78.62% ^{a,b}	93.80%	85.13%	93.94%	92.57%
Data Augmentation - Flip		92.73%	84.41%	92.94%	91.29%
Graph Data Augmentation		$92.10\%^d$		92.81%	91.07%
None	69.62%	87.78%		88.83%	85.88% ^a

^a No prior about the structure

Table 9: CIFAR-10 and scrambled CIFAR-10 comparison table.

Support	None		Neighborhood Graph		
Method	MLP	CNN (1x1 kernels)	ChebNet ^c	Proposed	
Accuracy	82.62%	84.30%	82.80%	85.08%	

Table 10: PINES fMRI comparison table.

3.5 Conclusion

In this chapter, we developed a new perspective to look at neural networks. This led us to discover that neural layers can be formulated with a ternary operation, that we called *neural contraction*, between an input signal X, a weight kernel Θ , and a weight sharing scheme S. We studied this representation and proposed efficient implementations. We saw it can represent any kind of layer. In particular we showed how related works from the literature can be represented with neural contractions. Also, we used it see the influence of symmetries, and concluded on their critical role in the success of convolutions. Then, we tested models on the task of graph signals classification and on the task of semi-supervised classification of nodes. To construct the scheme S, we tested ideas based on randomizations, from which we derived what we called *Monte-Carlo Neu-ral Networks* (MCNN), and a technique we called *graph dropout*. We also tested to learn the scheme S, obtaining *Graph Contraction Networks* (GCT).

^b Lin et al., 2015

^c Defferrard et al., 2016

^d Data augmentation done with covariance graph

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Finally, we tested to infer the scheme S from a set of translations inferred from the domain, which we called Translation-Convolution Neural Networks (TCNN). On image and fMRI datasets, GCTs and TCNNs obtained performances that match those of CNNs. On scrambled image datasets, TCNNs obtained almost the same performance as in the non-scrambled case, outperforming alternatives by a large margin. On text documents, MCNNs beat the other graph convolution alternative. On citation networks, GCTs set new state-of-the-art results, but by small margins that are not statistically significant.

Conclusion

In this manuscript, after a presentation of the fields of research, we developed a theory on convolution of graph signals, and proposed new models that extend deep learning to graph domains.

In Chapter 2, we formulated two constructions of convolutions of signals defined on a vertex set V, based on a group Γ acting on V. The φ convolution can be employed when Γ and V are in one-to-one correspondence via an equivariant map φ , while the M-convolution is a more convenient formulation that can be employed when Γ is abelian. We proved that the characterization by equivariance to Γ , inherited from group convolutions holds. Then we introduced two properties that bind these convolutions with the edge set E: edge-constraint (EC) and locality-preservation (LP). In view of describing operators that are used in deep learning, we proposed formulations with kernels of smaller supports, and proved that the weight sharing property holds. We demonstrated that the existence of convolutions on a graph can be characterized by the existence of Cayley subgraphs. For some graphs, their Cayley subgraphs might not be well representative of their topology. Therefore, we suggested a few strategies and we extended the previous results with convolutions based on groupoids rather than on groups. We constructed two types of groupoids, from partial transformations and paths, and were able to extend the results but also with some limitations. With the first type of groupoid this almost amounted to partition the graph into Cayley subgraphs, whereas with the second one it included degenerated cases.

In Chapter 3 we proposed a novel layer representation for extending CNN architectures to other input domains. This representation is based on a ternary operation that we called neural contraction, from which we derived new models: Monte-Carlo Neural Networks (MCNN), Graph Contraction Networks (GCT), Translation-Convolutional Neural Networks (TCNN); and a new technique: graph dropout. We also showed how to represent related models from the literature with neural contractions. The MCNN is a first idea exploiting the neural contraction. Roughly speaking, it is based on randomizing the structure that is leveraged, then averaging. So we didn't expect much from it but it fared well on a text categorization task. The GCT is based on the idea of learning how the weights are shared while learning them. It sets new state-of-the-art performances on the task of semi-supervised classification of nodes in citation networks, but outperforms only by a small margin that is not statistically significant. In particular, we also observed that using graph dropout also significantly improved the results of alternative models on this type of tasks. The TCNN relies on constructing a convolution based on graph translations. It sets a new state-of-the-art on classification of scrambled images by a large margin, and performs well on a fMRI dataset which is structured by a graph resembling a grid graph.

We tested these models in two types of tasks: supervised classification of graph signals and semi-supervised classification of nodes. The first task is historically the one that gave visibility to CNNs. However, we do not know of a dataset with a very unusual graph structure that is well fitted for the first task, or if it exists, it might not be the best graph to describe the underlying structure. This is why is practice, experiments for the supervised task are done with graphs resembling grid graphs to some extent. This is not the case for the semi-supervised task.

In the end, both task can be abstracted to a more general one. For example, let us consider a dataset represented by a matrix X, of shape $B \times N$,

where B is the number of instances and N is the number of features. The linear part of the GCN layer formulation $Y = AX\Theta$ exploits both a graph on the rows (that we saw can be learned with the GCT) and a complete graph on the columns. Thus this layer amounts to two half layers, a sparse and a dense one. This idea can be generalized given a dataset represented by a tensor X of rank r. The formulation would then be $Y = g(X, A_1, A_2, \ldots, A_r)$ where g realizes every tensor contractions between X and each A_i along the corresponding ranks. In that case, the A_i can be seen as a learnable normalized adjacency matrix corresponding to a graph structure on the ith rank. This idea can originate a class of neural networks that can be called *multiary neural networks*.

As pointed out in the introduction, this work participates in making CNNs more generic, and thus applicable to a broader range of real world problems. In the process, we also advanced our understanding of convolutions, providing a thorough description with a set of expressions, mathematical results and theorems about how to extend them on graph domains while preserving key properties, and how to characterize them. We hope that the reader had pleasure reading this manuscript and that it gave him ideas and shed new lights!

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Résumé en français

Ce manuscrit est une thèse soumise pour candidater au grade de docteur. Il est dévolu à deux sujets. Le premier traite d'extensions de la convolution discrète aux signaux sur graphe. Le second traite d'extensions de l'ensemble de définition des réseaux de neurones à des graphes. Les deux sujets sont reliés car les réseaux de neurones peuvent tirer profit de la structure sous-jacente de leur ensemble de définition à l'aide de convolutions.

Dans un premier temps, nous présentons les notions relatives à nos sujets d'étude. L'un deux, l'apprentissage profond, est le domaine de recherche qui se concentre sur une classe particulière de fonctions: les réseaux de neurones. Puisque nous essayons de suivre une approche rigoureuse, nous commençons par définir proprement leurs ensembles de définition et d'arrivée, qui peuvent être modélisés par des espaces tensoriels. En particulier, nous donnons des définitions originales pour les tenseurs, qui sont appropriées dans le cadre de l'étude de réseaux de neurones. Nous expliquons aussi comment les données sont traitées et manipulées. Nous définissons certaines opérations binaires qui sont importantes pour notre étude : contraction tensorielle et convolution. Ensuite, nous définissons les réseaux de neurones, discutons leur interprétation biologique, montrons comment ils apprennent, et relatons quelques avancées historiques. Puis, nous introduisons des couches classiques, en particulier les couches convolutionelles pour lesquelles nous démontrons un premier résultat qui avance notre réflexion. Puis, nous présentons le domaine de recherche

relatif à l'apprentissage profond sur graphes. Nous commençons par des définitions relatives aux graphes, puis nous décrivons des cas d'usage. Enfin, nous faisons une revue des modèles à l'état de l'art en deux parties, l'une sur les méthodes spectrales, l'autre sur les méthodes dans le domaine des sommets du graphe.

Au cours du deuxième chapitre, nous formalisons mathématiquement les principes de l'extension de la convolution aux signaux sur graphes. Si le graphe n'est pas un graphe grille, il n'existe pas de manière naturelle d'étendre la convolution Euclidienne. Nous commençons par analyser les raisons pour lesquelles la convolution Euclidienne est utile en apprentissage profond. En particulier, nous rappelons une caractérisation classique : la classe des opérateurs de convolutions est exactement la même que celle des fonctions linéaires qui sont équivariantes aux translations. Nous nous demandons donc sur quels domaines une convolution avec ces propriétés peut être obtenue de manière naturelle. Cela nous amène à considérer la théorie des représentations et les convolutions définies sur des groupes. Puisque la convolution Euclidienne est juste un cas particulier de la convolution de groupes, cela est parfaitement pertinent de diriger notre construction dans cette direction. Ensuite, nous cherchons à transférer la définition de la convolution de groupes sur les sommets, par le biais de son groupe symétrique. Pour obtenir la caractérisation désirée, nous constatons que nous avons besoin de baser la convolution sur les actions du groupe, plutôt que sur ses éléments. Nous parvenons à l'obtenir, à condition de fixer une carte équivariante entre le groupe actif et les sommets. Puis, nous proposons une expression mixte de cette convolution, définie entre un signal sur le groupe actif et un signal sur les sommets, pour laquelle nous démontrons que la caractérisation reste valable sous condition de commutativité. Puis, nous introduisons le rôle de l'ensemble des arrêtes du graphe pour voir comment il devrait influencer la construction. En particulier, nous définissons la notion de contrainte par les arrêtes et la notion de préservation de la localisation. Pour chacune, nous obtenons une caractérisation de graphes admettant une construction naturelle de convolutions avec cette propriété. Nous analysons les notions de localités et de partage de poids, et proposons une formulation dans le cas de petits noyaux de convolution. Grâce aux théorèmes obtenus, nous sommes capable de décrire les convolutions sur n'importe quel graphe, en tant que convolutions sur des sous-graphes appropriés. Ensuite, nous assouplissons certains contraintes de cette construction pour mieux l'adapter à des graphes quelconques. Nous expliquons pourquoi une construction basées sur des groupes est peu intéressante pour certains graphes, et nous introduisons la notion de groupoïde. Nous étendons la construction précédente avec des groupoïdes de transformations partielles, et démontrons que la caractérisation par équivariance est préservée. Enfin, nous étendons encore notre construction avec un autre type de groupoïdes que appelons groupoïdes de chemins. Les groupoïdes de chemins permettent de traiter le cas le plus général, et pour eux nous obtenons aussi la caractérisation à condition de fixer un manière de traverser les sommets, mais au prix d'inclure des cas dégénérés.

Dans le chapitre final, nous cherchons à comprendre comment les réseaux de neurones peuvent être étendus à des ensembles de définition pour lesquels ils n'étaient pas conçus pour y être applicable. Dans ce but, nous proposons une interprétation des opérations linéaires sous-jacentes afin d'améliorer notre intuition. Dans un premier temps, nous démontrons l'évident en expliquant avec plus de détails comment un espace tensoriel peut être interprété en temps qu'espace neuronal, tout en jonglant entre les représentations tensorielles et les représentations à bases de signaux. Puis, nous proposons une représentation basée sur des graphes. Entre deux couches, un graphe de propagation décrit cette propagation. Sur la couche en entrée, les neurones peuvent avoir une structure sous-

jacente de graphe. Nous montrons une relation entre ces graphes, que nous obtenons si, et seulement si, les champs de réceptions locaux des neurones sont entremêlés. En introduisant la notion de partage de poids dans notre analyse, nous découvrons qu'une couche, quel que soit son ensemble de définition, peut être formulée par une opération ternaire et linéaire, que nous appelons contraction neurale. Ses opérandes sont le signal d'entrée X, le noyaux de poids Θ , et le schéma de partage de poids S. Nous écrivons ΘSX . Nous étudions ses propriétés, et sa généricité en comparaison avec les méthodes de la littérature. Nous proposons une méthode pour l'implémenter de manière efficace. Grâce à une expérience qui se base sur cette opération, nous observons en quoi l'exploitation de symétries est bénéfique, ce qui justifie l'utilisation de convolutions. A l'aide d'autres expériences, nous explorons des idées basées sur des tirages aléatoires pour appliquer cette représentation ternaire dans le cas de graphes quelconques. Puis, nous étudions la possibilité d'apprendre comment les poids sont partagés, ce qui revient à apprendre à la fois S et Θ . Nous explorons cette piste pour des domaines de graphes, avec des expériences sur des grilles, sur des graphes de covariance et sur des réseaux de citations. Enfin, nous testons un exemple d'architecture convolutionelle appliquées pour des signaux sur graphe. La convolution est construite à partir de translations sur graphe qui définissent le schéma de partage de poids S de la couche convolutionelle. Nous présentons des modèles de translations et les approximations, les couches de souséchantillonage, ainsi que la technique d'augmentation de données que nous utilisons. Au travers d'expériences, nous appliquons ce modèle à des graphes grilles et autres graphes y ressemblant.

Dans la conclusion, nous rappelons les différents résultats, avancées, et nouveaux modèles de réseaux de neurones que nous avons présentés tout au long de ce mémoire.