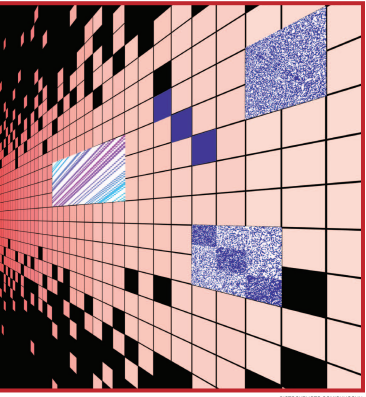
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# Theoretical Foundations of Deep Learning via Sparse Representations

A multilayer sparse model and its connection to convolutional neural networks



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odeling data is the way we-scientists-believe that information should be explained and handled. Indeed, models play a central role in practically every task in signal and image processing and machine learning. Sparse representation theory (we shall refer to it as Sparseland) puts forward an emerging, highly effective, and universal model. Its core idea is the description of data as a linear combination of few atoms taken from a dictionary of such fundamental elements.

Our prime objective in this article is to review a recently introduced [1] model-based explanation of deep learning, which relies on sparse modeling of data. We start by presenting the general story of Sparseland, describing its key achievements. We then turn to describe the convolutional-sparse-coding (CSC) model and present a multilayered (ML) extension of it, both being special cases of the general sparsity-inspired model. We show how ML-CSC leads to a solid and systematic theoretical justification of the architectures used in deep learning, along with a novel ability of analyzing the performance of the resulting machines. As such, this work offers a unique and first of its kind theoretical view for a field that has been, until recently, considered as purely heuristic.

#### Introduction

The field of sparse and redundant representations has made a major leap in the past two decades. Starting with a series of infant ideas and few mathematical observations, it grew to become a mature and highly influential discipline. Sparseland puts forward a universal mathematical model for describing the inherent low dimensionality that may exist in natural data sources. This model suggests a description of signals as linear combinations of few columns, called atoms, from a given redundant matrix, termed dictionary. In other words, these signals admit sparse representations with respect to their corresponding dictionary of prototype signals.

The Sparseland model gradually became central in signal and image processing and machine-learning applications, leading to state-of-the-art results in a wide variety of tasks and across many different domains. A partial explanation for the great appeal that

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this model has had is based on the theoretical foundations that accompany its construction, providing a solid support for many of the developed algorithms arising in this field. Indeed, in the broad scientific arena of data processing, Sparseland is quite unique due to the synergy that exists between theory, algorithms, and applications.

This article embarks from the story of Sparseland to discuss two recent special cases of it—CSC and ML-CSC. As we shall see, these descendant models pave a clear and surprising highway between sparse modeling and deep-learning architectures. This article's main goal is to present a novel theory for explaining deep (convolutional) neural networks and their origins, all through the language of sparse representations.

Clearly, our work is not the only one nor the first to theoretically explain deep learning. Indeed, various such attempts have already appeared in the literature (e.g., [2]–[13]). Broadly speaking, this knowledge stands on three pillars—the architectures used, the algorithms and optimization aspects involved, and the data these all serve. A comprehensive theory should cover all three, but this seems to be a tough challenge. The existing works typically cover one or two of these pillars, such as in the following examples:

- Architecture: The work by Giryes et. al. [9], showed that the used architectures tend to preserve distances, and explained the relevance of this property for classification. The work reported in [7] and [8] analyzed the capacity of these architectures to cover a specific family of functions.
- Algorithms: Vidal's work [5] explained why local minima are not to be feared, as they may coincide with the objective's global minimum. Chaudhari and Soatto proved [6] that the stochastic gradient descent algorithm induces an implicit regularization that is related to the information-bottleneck objective [3].
- Data: Bruna and Mallat [4] motivated the architectures by the data invariances that should be taken care of. Baraniuk's team
   [2] developed a probabilistic generative model for the data that, in turn, justifies the architectures used.

In this context we should note that our work covers the data by modeling it, and the architectures as emerging from the model. Our work is close in spirit to [2] but also markedly different.

A central idea that will accompany us in this article refers to the fact that Sparseland and its aforementioned descendants are all generative models. By this we mean that they offer a description of the signals of interest by proposing a synthesis procedure for their creation. We argue that such generative models facilitate a systematic pathway for algorithm design, while also enabling a theoretical analysis of their performance. Indeed, we will see throughout this article how these two benefits go hand in hand. By relying on the generative model, we will analyze certain feedforward convolutional neural networks (CNNs), identify key theoretical weaknesses in them, and then tackle these by proposing new architectures. Surprisingly, this journey will lead us to some of the well-known feed-forward CNNs used today.

Standing at the horizon of this work is our desire to present what is called the *ML convolutional sparse modeling* idea, as it will be the grounds on which we derive all the previously

mentioned claimed results. Thus, we take this as our running title and build the article, section by section, focusing each time on another word in it. We start by explaining better what we mean by the term *modeling*, and then move to describe *sparse modeling*, essentially conveying the story of Sparseland. Then we shift to *convolutional sparse modeling*, presenting this model along with a recent and novel analysis of it that relies on local sparsity. We conclude by presenting *ML convolutional sparse modeling*, tying this to the realm of deep learning, just as promised.

Before we start our journey, a few comments are in order.

- 1) Quoting Ron Kimmel (Computer Science Department at the Technion–Israel), this grand task of attaching a theory to deep learning behaves like a magic mirror, in which every researcher sees himself. This explains the so-diverse explanations that have been accumulated, relying on information theory [3], passing through wavelets and invariances [4], proposing a sparse modeling point of view [1], and going all the way to partial differential equations [10]. Indeed, it was David Donoho (Department of Statistics at Stanford University) who strengthened this vivid description by mentioning that this magical mirror is taken straight from Cinderella's story, as it is not just showing to each researcher his/her reflection, but also accompanies this with warm compliments, assuring that their view is truly the best.
- 2) This article focuses mostly on the theoretical sides of the models we shall discuss, but without delving into the proofs for the theorems we will state. This implies two things: 1) less emphasis will be put on applications and experiments; and 2) the content of this article is somewhat involved due to the delicate theoretical statements brought, so be patient.
- 3) Echoed in this article is a keynote talk given by Michael Elad during the 2017 IEEE International Conference on Image Processing in Beijing, as we follow closely this lecture, both in style and content. The recent part of the results presented (on CSC and ML-CSC) can be found in [1] and [14], but the description of the path from Sparseland to deep learning as posed here differs substantially.

## **Modeling**

## Our data is structured

Engineers and researchers rarely stop to wonder about our core ability to process signals—we simply take it for granted. Why is it that we can denoise signals? Why can we compress them, or recover them from various degradations, or find anomalies in them, or even recognize their content? The algorithms that tackle these and many other tasks—including separation, segmentation, identification, interpolation, extrapolation, clustering, prediction, synthesis, and many more—all rely on one fundamental property that only meaningful data sources obey: they are all structured.

Each source of information we encounter in our everyday lives exhibits an inner structure that is unique to it, and which can be characterized in various ways. We may allude to redundancy in the data, assume it satisfies a self-similarity property, suggest it is compressible due to low entropy, or even mention the possibility of embedding it into a low-dimensional manifold. No matter what the assumption is, the bottom line is the same—the data we operate on is structured. This is true for images of various kinds: video sequences; audio signals; three-dimensional (3-D) objects given as meshes or point clouds; financial time series; data on graphs, as is the case in social or traffic networks; text files such as e-mails and other documents; and more. In fact,

we could go as far as stating that if a given data is unstructured (e.g., being independent and identically distributed random noise), it would be of no interest to us, since processing it would be virtually futile.

So, coming back to our earlier question, the reason we can process data is the aforementioned structure, which facilitates this ability in all its manifestations. Indeed, the fields of signal and image processing and machine learning are mostly about identi-

fying the structure that exists in a given information source, and then exploiting it to achieve the processing goals. This brings us to discuss models and the central role they play in data processing.

# Identifying structure via models

An appealing approach for identifying structure in a given information source is imposing a (parametric) model on it, explicitly stating a series of mathematical properties that the data is believed to satisfy. Such constraints lead to a dimensionality reduction that is so characteristic of models and their modus operandi. We should note, however, that models are not the only avenue for identifying structure—the alternative being a non-parametric approach that simply describes the data distribution by accumulating many of its instances. We will not dwell on this option in this article, as our focus is on models and their role in data processing.

Consider the following example, brought to clarify our discussion. Assume that we are given a measurement vector  $\mathbf{y} \in \mathbb{R}^n$ , and all that is known to us is that it is built from an ideal signal of some sort,  $\mathbf{x} \in \mathbb{R}^n$ , contaminated by white additive Gaussian noise of zero mean and unit variance, i.e.,  $\mathbf{y} = \mathbf{x} + \mathbf{e}$ , where  $\mathbf{e} \sim \mathcal{N}(0,\mathbf{I})$ . Could we propose a method to clean the signal  $\mathbf{y}$  from the noise? The answer is negative! Characterizing the noise alone cannot suffice for handling the denoising task, as there are infinitely many possible ways to separate  $\mathbf{y}$  into a signal and a noise vector, where the estimated noise matches its desired statistical properties.

Now, suppose that we are given additional information on the unknown  $\mathbf{x}$ , believed to belong to the family of piece-wise constant (PWC) signals, with the tendency to have as few jumps as possible. In other words, we are given a model for the underlying structure. Could we leverage this extra information to denoise  $\mathbf{y}$ ? The answer is positive—we can seek the simplest PWC signal that is closest to  $\mathbf{y}$  in such a way that the error matches the expected noise energy. This can be formulated mathematically in some way or another, leading to an optimization problem whose solu-

tion is the denoised result. What have we done here? We imposed a model on our unknown, forcing the result to be likely under the believed structure, thus enabling the denoising operation. The same thought process underlies the solution of almost any task in signal and image processing and machine learning, either explicitly or implicitly.

As yet another example for a model in image processing and its impact, consider the JPEG compression algorithm [15]. We

are well aware of the impressive ability of this method to compress images by factor of ten to 20 with hardly any noticeable artifacts. What are the origins of this success? The answer is twofold: the inner structure that exists in images, and the model that JPEG harnesses to exploit it. Images are redundant, as we already claimed, allowing for the core possibility of such compression to take place. However, the structure alone cannot suffice to get the actual compression, as a model is needed to capture

this redundancy. In the case of JPEG, the model exposes this structure through the belief that small image patches (of size  $8\times 8$  pixels) taken from natural images tend to concentrate their energy in the lower-frequency part of the spectrum once operated upon by the discrete cosine transform (DCT). Thus, few transform coefficients can be kept while the rest can be discarded, leading to the desired compression result. One should nevertheless wonder, will this algorithm perform just as well on other signal sources? The answer is not necessarily positive, suggesting that every information source should be fitted with a proper model.

#### The evolution of models

This article's main goal

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(convolutional) neural

A careful survey of the literature in image processing reveals an evolution of models that have been proposed and adopted over the years. We will not provide here an exhaustive list of all of these models, but we do mention a few central ideas such as Markov random fields for describing the relation between neighboring pixels [16], Laplacian smoothness of various sorts [17], total variation [18] as an edge-preserving regularization, wavelets' sparsity [19], [20], and Gaussian mixture models (GMMs) [21], [22]. With the introduction of better models, performance improved in a wide front of applications in image processing.

Consider, for example, the classic problem of image denoising, on which thousands of papers have been written. Our ability to remove noise from images has advanced immensely over the years. Indeed, the progress made has reached the point where this problem is regarded by many in our field as nearly solved [23], [24]. Performance in denoising has improved steadily over time, and this improvement was enabled mostly by the introduction of better and more effective models for natural images. The same progress applies to image deblurring, inpainting, superresolution, compression, and many other tasks.

In our initial description of the role of models, we stated that these are expressing what the data is believed to satisfy, eluding to the fact that models cannot be proven to be correct, just as a formula in physics cannot be claimed to describe our world perfectly. Rather, models can be compared and contrasted, or simply tested empirically to see whether they fit reality sufficiently well. This is perhaps the place to disclose that models are almost always wrong, as they tend to explain reality in a simple manner at the cost of its oversimplification. Does this mean that models are necessarily useless? Not at all. While they do carry an error in them, if this deviation is small enough, then such models are priceless and extremely useful for our processing needs. How small is small enough? This is a tough question that has not been addressed in the literature. Here we will simply be satisfied with the assumption that this error should be substantially smaller compared to the estimation error the model leads to. Note that, often times,

the acceptable relative error is dictated by the application or problem that one is trying to solve by employing the model.

For a model to succeed in its mission of treating signals, it must be a good compromise between simplicity, reliability, and flexibility. Simplicity is crucial since this implies that algorithms harnessing this model are relatively easy and pleasant to work with. However, simplicity is not sufficient. We could suggest, for example, a model that simply assumes that the signal of interest is zero. This is the simplest model imaginable, and yet it is also

useless. So, next to simplicity, comes the second force of reliability—we must offer a model that does justice to the data served, capturing its true essence. The third virtue is flexibility, implying that the model can be tuned to better fit the data source, thus erring less. Every model is torn between these three forces, and we are constantly seeking simple, reliable, and flexible models that improve over their predecessors.

# Models—Summary

Models are central for enabling the processing of structured data sources. In image processing, models take a leading part in addressing many tasks, such as denoising, deblurring, and all the other inverse problems, compression, anomaly detection, sampling, recognition, separation, and more. We hope that this perspective on data sources and their inner structure clarifies the picture, putting decades of research activity in the fields of signal and image processing and machine learning in a proper perspective.

In the endless quest for a model that explains reality, one that has been of central importance in the past two decades is Sparseland. This model slowly and consistently carved its path to the lead, fueled by both great empirical success and impressive accompanying theory that added a much needed color to it. We turn now to present this model. We remind the reader that this is yet another station in our road toward providing a potential explanation of deep learning using sparsity-inspired models.

## **Sparse modeling**

# On the origin of sparsity-based modeling

Simplicity as a driving force for explaining natural phenomena has a central role in sciences. While Occam's razor is perhaps the earliest of this manifestation (though in a philosophical or religious context), a more recent and relevant quote from Wrinch and Jeffrey (1921) [25] reads: "The existence of simple laws is, then, apparently, to be regarded as a quality of nature; and accordingly we may infer that it is justifiable to prefer a simple law to a more complex one that fits our observations slightly better."

When it comes to the description of data, sparsity is an ultimate expression of simplicity, which explains the great attraction it has. While it is hard to pinpoint the exact appearance of the concept of sparsity in characterizing data priors, it is quite clear

> that this idea became widely recognized with the arrival of the wavelet revolution that took place during the late 1980s and early 1990s. The key observation was that this particular transformation, when applied to many different signals or images, produced representations that were naturally sparse. This, in turn, was leveraged in various ways, both empirically and theoretically. Almost in parallel, approximation theorists started discussing the dichotomy between linear and nonlinear approximation, emphasizing further the role of sparsity in signal analysis. These are the

prime origins of the field of Sparseland, which borrowed the core idea that signal representations should be redundant and sparse, while putting aside many other features of wavelets, such as (bi)orthogonality, multiscale analysis, frame theory interpretation, and more.

Signs of Sparseland appeared already in the early and mid-1990s, with the seminal papers on greedy- (Zhang and Mallat, [26]) and relaxation-based (Chen, Donoho, and Saunders, [27]) pursuit algorithms, and even the introduction of the concept of dictionary learning by Olshausen and Field [28]. However, it is our opinion that this field was truly born only few years later, in 2000, with the publication of the paper by Donoho and Huo [29], which was the first to show that the basis pursuit (BP) is provably exact under some conditions. This work dared and defined a new language, setting the stage for thousands of follow-up papers. Sparseland started with a massive theoretical effort, which slowly expanded and diffused to practical algorithms and applications, leading in many cases to state-of-the-art results in a wide variety of disciplines. The knowledge in this field as it stands today relies heavily on numerical optimization and linear algebra, and parts of it have a definite machine-learning flavor.

## Introduction to Sparseland

So, what is this model? How can it capture structure in a data source? Let us demonstrate this for  $8 \times 8$  image patches, in the spirit of the description of the JPEG algorithm mentioned previously. Assume that we are given a family of patches extracted from natural images. The Sparseland model starts by preparing a dictionary—a set of atom patches of the same size,  $8 \times 8$  pixels. For example, consider a dictionary containing 256 such atoms. Then, the model assumption is that every incoming patch could be described as a linear combination of only few

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atoms from the dictionary. The word *few* here is crucial, as every patch could be easily described as a linear combination of 64 linearly independent atoms, a fact that leads to no structure whatsoever.

Let's take a closer look at this model, as depicted in Figure 1(a). We started with a patch of size  $8 \times 8$  pixels, thus 64 values. The first thing to observe is that we have converted it to a vector of length 256 carrying the weights of each of the atoms in the mixture that generates this patch. Thus, our

representation is redundant. However, this vector is also very sparse, since only few of the atoms participate in this construction. Imagine, for example, that only three atoms are used. In this case, the complete information about the original patch is carried by six values: three stating which atoms are involved, and three determining their weights. Thus, this model manages to reduce the dimensionality of the patch information, a key property in exposing the structure in our data.

An interesting analogy can be drawn between this model and our world's chemistry. The model's dictionary and its atoms should remind the reader of Mendeleev's periodic table. In our world, every molecule is built of few of the fundamental elements from this table, and this parallels our model assumption that states that signals are created from few atoms as well. As such, we can regard the Sparseland model as an adoption of the core rational of chemistry to data description.

Let's make the Sparseland model somewhat more precise, and introduce notations that will serve us throughout this arti-

Sparseland has a universal ability to describe information content faithfully and effectively, due to the exponential number of possible supports that a reasonable-sized dictionary enables.

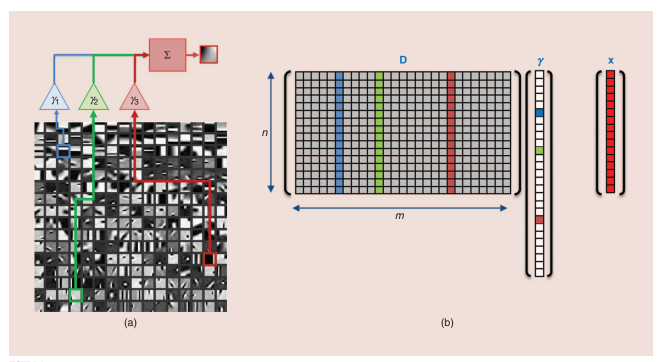
cle [30]. The signal we operate on is denoted by  $\mathbf{x} \in \mathbb{R}^n$ , and the dictionary  $\mathbf{D}$  is a matrix of size  $n \times m$ , in which each of its m columns is an atom. The sparse and redundant representation is the vector  $\mathbf{\gamma} \in \mathbb{R}^m$ , which multiplies  $\mathbf{D}$  to create  $\mathbf{x}$ , i.e.,  $\mathbf{x} = \mathbf{D}\mathbf{\gamma}$ , as shown in Figure 1(b). The vector  $\mathbf{\gamma}$  has only few (say, k) nonzeros, and thus it creates a linear combination of k atoms from  $\mathbf{D}$  to construct  $\mathbf{x}$ . We shall denote by  $\|\mathbf{\gamma}\|_0$  the number of nonzeros in  $\mathbf{\gamma}$ . This  $\ell_0$  is not a formal norm as it does not satisfy the homo-

geneity property. Nevertheless, throughout this article, we shall refer to this as a *regular norm*, with the understanding of its limitations. So, put formally, here is the definition of Sparseland signals.

## Definition

The set of Sparseland signals of cardinality k over the dictionary **D** is defined as  $\mathcal{M}\{\mathbf{D}, k\}$ . A signal **x** belongs in  $\mathcal{M}\{\mathbf{D}, k\}$  if it can be described as  $\mathbf{x} = \mathbf{D}\boldsymbol{\gamma}$ , where  $\|\boldsymbol{\gamma}\|_0 \le k$ .

Observe that this is a generative model in the sense that it describes how (according to our belief) the signals have been created from k atoms from  $\mathbf{D}$ . We mentioned previously that models must be simple to be appealing, and, in this spirit, we must ask: Can the Sparseland model be considered as simple? Well, the answer is not so easy, since this model raises major difficulties in its deployment, and this might explain its late adoption. In the following, we mention several such difficulties, given in ascending order of complexity.



**FIGURE 1.** The decomposition of an image patch into its three building blocks—atoms from the dictionary. This model can be expressed as  $\mathbf{D}\gamma = \mathbf{x}$ , where  $\|\gamma\|_0 = 3$ .

# Sparseland difficulties: Atom decomposition

The term *atom-decomposition* refers to the most fundamental problem of identifying the atoms that construct a given signal. Consider the following example: We are given a dictionary having 2,000 atoms, and a signal known to be composed of a mixture of 15 of these. Our goal now is to identify these 15 atoms. How should this be done? The natural option to consider is an exhaustive search over all the possibilities of choosing 15 atoms out of the 2,000, and checking per each whether they fit the measurements. The number of such possibilities to check stands on an order of 2.4e + 37, and even if each of this takes 1 ps, billions of years will be required to complete this task!

Put formally, atom decomposition can be described as the following constrained optimization problem:

$$\min_{\mathbf{r}} \| \boldsymbol{\gamma} \|_{0} \quad \text{s.t.} \quad \mathbf{x} = \mathbf{D} \boldsymbol{\gamma}. \tag{1}$$

This problem seeks the sparsest explanation of  $\mathbf{x}$  as a linear combination of atoms from  $\mathbf{D}$ . In a more practical version of the atom-decomposition problem, we may assume that the signal we get,  $\mathbf{y}$ , is an  $\epsilon$ -contaminated version of  $\mathbf{x}$ , and then the optimization task becomes

$$\min_{\boldsymbol{\gamma}} \| \boldsymbol{\gamma} \|_{0} \quad \text{s.t.} \quad \| \boldsymbol{y} - \mathbf{D} \boldsymbol{\gamma} \|_{2} \le \epsilon. \tag{2}$$

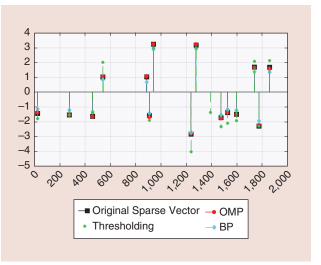
Both (1) and (2) are known to be NP-hard, implying that their complexity grows exponentially with the number of atoms in **D**. So, are we stuck?

The answer to this difficulty came in the form of approximation algorithms, originally meant to provide exactly that—an approximate solution to the previously given problems. In this context, we mention greedy methods such as the orthogonal matching pursuit (OMP) [31] and the thresholding algorithm, and relaxation formulations such as the BP [27].

While it is beyond the scope of this article to provide a detailed description of these algorithms, we will say a few words on each, as we will return to use them later when we get closer to the connection to neural networks. BP takes the problem posed in (2) and relaxes it by replacing the  $\ell_0$  by an  $\ell_1$ -norm. With this change, the problem is convex and manageable in reasonable time.

Greedy methods such as the OMP solve the problem posed in (2) by adding one nonzero at a time to the solution, trying to reduce the error  $\|\mathbf{y} - \mathbf{D}\boldsymbol{\gamma}\|_2$  as much as possible at each step, and stopping when this error goes below  $\epsilon$ . The thresholding algorithm is the simplest and crudest of all pursuit methods—it multiplies  $\mathbf{y}$  by  $\mathbf{D}^T$ , and applies a simple shrinkage on the resulting vector, nulling small entries and leaving the rest almost untouched.

Figure 2 presents an experiment in which these three algorithms were applied on the scenario we described previously, in which **D** has 2,000 atoms, and an approximate atom decomposition is performed on noisy signals that are known to be created from few of these atoms. The results shown suggest that these three algorithms tend to succeed rather well in their mission.



**FIGURE 2.** An illustration of OMP, BP, and thresholding in approximating the solution of the atom decomposition problem, for a dictionary with 2.000 atoms.

## Sparseland difficulties: Theoretical foundations

Can the success of the pursuit algorithms be explained and justified? One of the grand achievements of the field of Sparseland is the theoretical analysis that accompanies many of these pursuit algorithms, claiming their guaranteed success under some conditions on the cardinality of the unknown representation and the dictionary properties. Hundreds of papers offering such results were authored in the past two decades, providing Sparseland with the necessary theoretical foundations. This activity essentially resolved the atom-decomposition difficulty to the point where we can safely assume that this task is doable—reliably, efficiently, and accurately.

Let us illustrate the theoretical side of Sparseland by providing a small sample from these results. We bring one such representative theorem that discusses the terms of success of the BP. This is the first among a series of such theoretical results that will be stated throughout this article. Common to them all is our sincere effort to choose the simplest results, and these will rely on a simple property of the dictionary **D** called the *mutual coherence*.

#### Definition

Given the dictionary  $\mathbf{D} \in \mathbb{R}^{n \times m}$ , assume that the columns of this matrix,  $\{\mathbf{d}_i\}_{i=1}^m$  are  $\ell_2$ -normalized. The mutual coherence  $\mu(\mathbf{D})$  is defined as the maximal absolute inner product between different atoms in  $\mathbf{D}$  [32]:

$$\mu(\mathbf{D}) = \max_{1 \le i \le |\mathbf{d}|} \mathbf{d}_i^T \mathbf{d}_j \,. \tag{3}$$

Clearly,  $0 \le \mu(\mathbf{D}) \le 1$  and, as we will see, the smaller this value is, the better our theoretical guarantees become. We note that other characterizations of  $\mathbf{D}$  exist and have been used quite extensively in developing the theory of Sparseland. These include the restricted isometry property (RIP) [33], the exact recovery condition (ERC) [34], the Babel function [35], the

Spark [36], and others. As mentioned previously, we stick with  $\mu(\mathbf{D})$  in this article due to its simplicity, and this may come at the cost of getting weaker guarantees.

We are now ready to state our first theorem: The story starts with an arbitrary sparse vector  $\gamma$  that generates a Sparseland signal  $\mathbf{x} = \mathbf{D}\gamma$ . We get a noisy version of this signal,  $\mathbf{y} = \mathbf{x} + \mathbf{e}$ , and our goal is to obtain an evaluation of a sparse representation that should be as close as possible to the original  $\gamma$ . The following result claims that BP is guaranteed to provide such performance.

#### Theorem 1

Given  $\mathbf{y} = \mathbf{D}\boldsymbol{\gamma} + \mathbf{e}$ , where  $\mathbf{e}$  is an energy-bounded noise,  $\|\mathbf{e}\|_2 \le \epsilon$ , and  $\boldsymbol{\gamma}$  is sufficiently sparse,

$$\|\boldsymbol{\gamma}\|_0 < \frac{1}{4} \left(1 + \frac{1}{\mu(\mathbf{D})}\right),\tag{4}$$

then the BP solution, given by

$$\hat{\boldsymbol{\gamma}} = \arg\min_{\mathbf{y}} \|\boldsymbol{\gamma}\|_{1} \text{ s.t. } \|\mathbf{D}\boldsymbol{\gamma} - \mathbf{y}\|_{2} \le \epsilon,$$
 (5)

leads to a stable result.

$$\|\hat{\boldsymbol{\gamma}} - \boldsymbol{\gamma}\|_2 \le \frac{4\epsilon^2}{1 - \mu(\mathbf{D})(4\|\boldsymbol{\gamma}\|_0 - 1)}.$$
 (6)

A few comments are in order:

- Observe that if we assume  $\epsilon = 0$ , then Theorem 1 essentially guarantees a perfect recovery of the original  $\gamma$ .
- The result stated is a worst-case one, claiming a perfect recovery under the conditions posed and for an adversarial noise. Far stronger claims exist, in which the noise model is more realistic (for example, random Gaussian), and then the language is changed to a probabilistic statement (i.e., success with probability close to 1) under much milder conditions (see, for example, [37] and [38]).
- The literature on Sparseland offers many similar such theorems, either improving Theorem 1, or referring to other pursuit algorithms.

# Sparseland difficulties: The quest for the dictionary

Now that we are not so worried anymore about solving the problems posed in (1) and (2), we turn to discuss a far greater difficulty—how can we get the dictionary **D**? Clearly, everything that we do with this model relies on a proper choice of this matrix. Sweeping through the relevant data processing literature, we may see attempts to

- use the Sparseland model to fill-in missing parts in natural images [39], [40]
- deploy this model for audio processing (e.g., [41] and [42])
- plan to exploit it for processing seismic data [43], [44]
- process volumes of hyperspectral imaging [45], [46].

Each of these applications and many others out there call for a separate and adapted dictionary, so how can **D** be chosen or found?

The early steps in Sparseland were made using known transforms as dictionaries. The intuition behind this idea was that carefully tailored transforms that match specific signals or have

particular properties could serve as the dictionaries we are after. Note that the dictionary represents the inverse transform, as it multiplies the representation to construct the signal. In this spirit, wavelets of various kinds were used for one-dimensional (1-D) signals [19], two-dimensional (2-D) DCT for image patches, and curvelets [47], contourlets [48], and shearlets [20] were suggested for images.

While seeming reasonable and elegant, this approach was found to be quite limited in real applications. The obvious reasons for this weakness were the partial match that exists between a chosen transform and true data sources, and the lack of flexibility in these transforms that would enable them to cope with special and narrow families of signals (e.g., face images, or financial data). The breakthrough in this quest of getting appropriate dictionaries came in the form of dictionary learning. The idea is quite simple: if we are given a large set of signals believed to emerge from a Sparseland generator, we can ask what is the best dictionary that can describe these sparsely. In the past decade, we have seen a wealth of dictionary-learning algorithms, varied in their computational steps, in their objectives, and in their basic assumptions on the required D. These algorithms gave Sparseland the necessary boost to become a leading model, due to the added ability to adapt to any data source, and match to its content faithfully. In this article, we will not dwell too long on this branch of work, despite its centrality, as our focus will be the model evolution we are about to present.

# Sparseland difficulties: Model validity

We are listing difficulties that the Sparseland model encountered, and, in this framework, we mentioned the atom-decomposition task and the pursuit algorithms that came to resolve it. We also described the quest for the dictionary and the central role of dictionary-learning approaches in this field. Beyond these, perhaps the prime difficulty that Sparseland poses is encapsulated in the following questions: Why should this model be trusted to perform well on a variety of signal sources? Clearly, images are not made of atoms, and there is no dictionary behind the scene that explains our data sources. So, what is the appeal in this specific model?

The answers to those questions are still being built, and they take two possible paths. On the empirical front, Sparseland has been deployed successfully in a wide variety of fields and for various tasks, leading time and again to satisfactory and even state-of-the-art results. So, one obvious answer is the simple statement "We tried it, and it works!" Again, a model cannot be proven to be correct, but it can be tested with real data, and this is the essence of this answer.

The second branch of answers to the natural repulse from Sparseland has been more theoretically oriented, tying this model to other, well-known and better established, models, showing that it generalizes and strengthens them. Connections have been established between Sparseland and Markov random field models, GMMs, and other union-of-subspaces constructions [30]. Indeed, the general picture obtained from the answers previously given suggests that Sparseland has a universal ability to describe information content faithfully and effectively, due to the exponential

number of possible supports that a reasonable-sized dictionary enables.

# Sparseland difficulties: Summary

The bottom line to all of this discussion is the fact that, as opposed to our first impression, Sparseland is a simple yet flexible model, and one that can be trusted. The difficulties mentioned have been fully resolved and answered constructively, and today we are armed with a wide set of algorithms and supporting theory for deploying Sparseland successfully for processing signals.

The interest in this field has grown impressively over the years, and this is clearly manifested by the exponential growth of papers published in the arena. Another testimony to the interest in Sparseland is seen by the wealth of books published in the past decade in this field—see [19], [30], and [49]–[56]. This attention brought us to offer a new massive open online course (MOOC), covering the theory and practice of this field. This MOOC, given under edX, started in October 2017, and already has more than 2,000 enrolled students. It is expected to be open continuously for all those who are interested in getting to know more about this field.

# Local versus global in sparse modeling

We now discuss the practicalities of deploying Sparseland in image processing. The common practice in many of such algorithms, and especially the better performing ones, is to operate on small and fully overlapping patches. The prime reason for this mode of work is the desire to harness dictionary-learning methods, and these are possible only for low-dimensional signals such as patches. The prior used in such algorithms is therefore the assumption that every patch extracted from the image is believed to have a sparse representation with respect to a commonly built dictionary.

After years of using this approach, questions started surfacing regarding this local model assumption, and the underlying global model that may operate behind the scene. Consider the following questions, all referring to a global signal  ${\bf X}$  that is believed to obey such a local behavior—having a sparse representation with respect to a local dictionary  ${\bf D}$  for each of its extracted patches:

- How can such a signal be synthesized?
- Do such signals exist for any **D**?
- How should the pursuit be done to fully exploit the believed structure in **X**?
- How should **D** be trained for such signals?

These tough questions started being addressed in recent works [14], [57], and this brings us to our next section, in which we dive into a special case of Sparseland—the CSC model, and resolve this global-local gap.

## **Convolutional sparse modeling**

## Introducing the CSC model

The CSC model offers a unique construction of signals based on the following relationship:

$$\mathbf{X} = \sum_{i=1}^{m} \mathbf{d}_{i} * \Gamma_{i}. \tag{7}$$

Note that all of our equations and derivations will be built under the assumption that the treated signals are 1-D, but this model and all our derivations apply to 2-D and even higher dimensions just as well. In our notations,  $\mathbf{X} \in \mathbb{R}^N$  is the constructed global signal, the set  $\{\mathbf{d}_i\}_{i=1}^m \in \mathbb{R}^n$  are m local filters of small support  $(n \ll N)$ , and  $\Gamma_i \in \mathbb{R}^N$  are sparse vectors convolved with their corresponding filters. For simplicity and without loss of generality, we shall assume hereafter that the convolution operations used are all cyclic. Adopting a more intuitive view of the CSC model, we can say that  $\mathbf{X}$  is assumed to be built of many instances of the small m filters, spread all over the signal domain. To be exact, the flipped version of the filters are the ones shifted in all spatial locations, since this flip is part of the convolution definition. This spread is obtained by the convolution of  $\mathbf{d}_i$  by the sparse feature map  $\Gamma_i$ .

And here is another, more convenient, view of the CSC model. We start by constructing m circulant matrices  $\mathbf{C}_i \in \mathbb{R}^{N \times N}$ , representing the convolutions by the filters  $\mathbf{d}_i$ . Each of these matrices is banded, with only n populated diagonals. Thus, the global signal  $\mathbf{X}$  can be described as

$$\mathbf{X} = \sum_{i=1}^{m} \mathbf{C}_{i} \mathbf{\Gamma}_{i} = \begin{bmatrix} \mathbf{C}_{1} \ \mathbf{C}_{2} \ \cdots \ \mathbf{C}_{m} \end{bmatrix} \begin{bmatrix} \mathbf{\Gamma}_{1} \\ \mathbf{\Gamma}_{2} \\ \vdots \\ \mathbf{\Gamma}_{m} \end{bmatrix} = \mathbf{D} \mathbf{\Gamma}.$$
 (8)

The concatenated circulant matrices form a global dictionary  $\mathbf{D} \in \mathbb{R}^{N \times mN}$ , and the gathered sparse vectors  $\mathbf{\Gamma}_i$  lead to the global sparse vector  $\mathbf{\Gamma} \in \mathbb{R}^{mN}$ . Thus, just as said previously, CSC is a special case of Sparseland, built around a very structured dictionary being a union of banded and circulant matrices. Figure 3 presents the obtained dictionary  $\mathbf{D}$ , where we permute its columns to obtain a sliding block diagonal form. Each of the small blocks in this diagonal is the same, denoted by  $\mathbf{D}_L$ —this is a local dictionary of size  $n \times m$ , containing the m filters as its columns.

# Why should we consider the CSC?

Why are we interested in the CSC model? Because it resolves the global-local gap we mentioned previously. Suppose that we extract an n-length patch from  $\mathbf{X}$  taken in location i. This is denoted by multiplying  $\mathbf{X}$  by the patchextractor operator  $\mathbf{R}_i \in \mathbb{R}^{n \times N}, \mathbf{p}_i = \mathbf{R}_i \mathbf{X}$ . Using the relation  $\mathbf{X} = \mathbf{D}\mathbf{\Gamma}$ , we have that  $\mathbf{p}_i = \mathbf{R}_i \mathbf{X} = \mathbf{R}_i \mathbf{D}\mathbf{\Gamma}$ . Note that the multiplication  $\mathbf{R}_i \mathbf{D}$  extracts n rows from the dictionary, and most of their content is simply zero. To remove their empty columns, we introduce the stripe extraction operator  $\mathbf{S}_i \in \mathbb{R}^{(2n-1)m \times mN}$  that extracts the nonzero part in this set of rows:  $\mathbf{R}_i \mathbf{D} = \mathbf{R}_i \mathbf{D} \mathbf{S}_i^T \mathbf{S}_i$ . Armed with this definition, we observe that  $\mathbf{p}_i$  can be expressed as  $\mathbf{p}_i = \mathbf{R}_i \mathbf{D} \mathbf{S}_i^T \mathbf{S}_i \mathbf{\Gamma} = \Omega \mathbf{\gamma}_i$ . The structure of  $\Omega$ , referred to as the *stripe dictionary*, appears in Figure 3, showing that  $\Omega = \mathbf{R}_i \mathbf{D} \mathbf{S}_i^T$  is a fixed

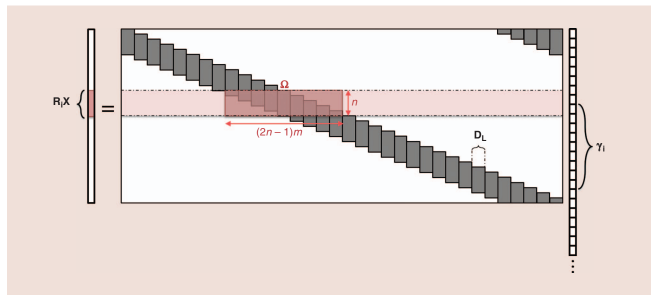


FIGURE 3. The global CSC model.

matrix regardless of *i*. The notation  $\gamma_i = \mathbf{S}_i \Gamma$  stands for a stripe of (2n-1)m elements from  $\Gamma$ .

And now we get to the main point of all of this description: As we move from location i to i+1, the patch  $\mathbf{p}_{i+1} = \mathbf{R}_{i+1} \mathbf{X}$  equals  $\Omega \gamma_{i+1}$ . The stripe vector  $\gamma_{i+1}$  is a shifted version of  $\gamma_i$  by m elements. Other than that, we observe that the extracted patches are all getting a sparse description of their content with respect to a common dictionary  $\Omega$ , just as assumed by the locally operating algorithms. Thus, CSC furnishes a way to make our local modeling assumption valid, while also posing a clear global model for  $\mathbf{X}$ . In other words, the CSC model offers a path toward answering all of the questions that we have posed in the context of the global-local gap.

# CSC: New theoretical foundations

Realizing that CSC may well be the missing link to all of the locally operating image processing algorithms, and recalling that it is a special case of Sparseland, we should wonder whether the classic existing theory of Sparseland provides sufficient foundations for it as well. Using Theorem 1 to the case in which the signal we operate upon is  $\mathbf{Y} = \mathbf{D}\Gamma + \mathbf{E}$ , where  $\|\mathbf{E}\|_2 \le \epsilon$  and  $\mathbf{D}$  is a convolutional dictionary, the condition for success of the BP is

$$\|\Gamma\|_0 < \frac{1}{4} \left(1 + \frac{1}{\mu(\mathbf{D})}\right). \tag{9}$$

Interestingly, the Welch bound offers a lower bound on the best achievable mutual coherence of our dictionary [58], being

$$\mu(\mathbf{D}) \ge \sqrt{\frac{m-1}{m(2n-1)-1}}$$
 (10)

For example, for m=2 filters of length n=200, this value is  $\approx 0.035$ , and this value necessarily grows as m is increased. This implies that the bound for success of the BP stands on  $\|\Gamma\|_0 < 7.3$ . In other words, we allow the entire vector  $\Gamma$  to have only seven nonzeros (independently of N, the size of  $\mathbf{X}$ ) for

the ability to guarantee that BP will recover a close-enough sparse representation. Clearly, such a statement is meaningless, and the unavoidable conclusion is that the classic theory of Sparseland provides no solid foundations whatsoever to the CSC model.

The recent work reported in [14] offers an elegant way to resolve this difficulty, by moving to a new, local, measure of sparsity. Rather than counting the number of nonzeros in the entire vector  $\Gamma$ , we run through all the stripe representations,  $\gamma_i = \mathbf{S}_i \Gamma$  for i = 1, 2, 3, ..., N, and define the relevant cardinality of  $\Gamma$  as the maximal number of nonzeros in these stripes. Formally, this measure can be defined as follows.

## Definition

Given the global vector  $\Gamma$ , we define its local cardinality as

$$\| \mathbf{\Gamma} \|_{0,\infty}^s = \max_{1 \le i \le N} \| \mathbf{\gamma}_i \|_0. \tag{11}$$

In this terminology, it is an  $\ell_{0,\infty}$  measure since we count number of nonzeros, but also maximize over the set of stripes. The superscript s stands for the fact that we sweep through all the stripes in  $\Gamma$ , skipping m elements from  $\gamma_i$  to  $\gamma_{i+1}$ .

Intuitively, if  $\| \Gamma \|_{0,\infty}^s$  is small, this implies that all the stripes are sparse, and thus each patch in  $\mathbf{X} = \mathbf{D}\Gamma$ , has a sparse representation w.r.t. the dictionary  $\Omega$ . Recall that this is exactly the model assumption we mentioned previously when operating locally. Armed with this local-sparsity definition, we are now ready to define CSC signals, in the same spirit of the Sparseland definition:

#### Definition

The set of CSC signals of cardinality k over the convolutional dictionary  $\mathbf{D}$  is defined as  $\mathcal{S}\{D,k\}$ . A signal  $\mathbf{X}$  belongs to  $\mathcal{S}\{D,k\}$  if it can be described as  $\mathbf{X} = \mathbf{D}\mathbf{\Gamma}$ , where  $\|\mathbf{\Gamma}\|_{0,\infty}^{s} \leq k$ .

Could we use these new notions of the local sparsity and the CSC signals in order derive novel and stronger guarantees for the

success of pursuit algorithms for the CSC model? The answer, as given in [14], is positive. We now present one such result, among several others that are given in that work, referring in this case to the BP algorithm. We note that the notation  $\|\mathbf{E}\|_{2,\infty}^p$  stands for computing the  $\ell_2$ -norm on fully overlapping sliding patches (hence the letter "p") extracted from  $\mathbf{E}$ , seeking the most energetic patch. As such, this quantifies a local measure of the noise, which serves the following theorem.

Theorem 2

Given  $Y = D\Gamma + E$ , and  $\Gamma$  that is sufficiently locally sparse,

$$\|\mathbf{\Gamma}\|_{0,\infty}^s < \frac{1}{3} \left(1 + \frac{1}{\mu(\mathbf{D})}\right),\tag{12}$$

the BP algorithm

$$\hat{\mathbf{\Gamma}} = \arg\min_{\mathbf{\Gamma}} \frac{1}{2} \|\mathbf{Y} - \mathbf{D}\mathbf{\Gamma}\|_{2}^{2} + \lambda \|\mathbf{\Gamma}\|_{1}$$
 (13)

with  $\lambda = 4 \| \mathbf{E} \|_{2,\infty}^p$  satisfies the following:

- The support of  $\hat{\Gamma}$  is contained in that of the original  $\Gamma$ .
- The result is stable:  $\|\hat{\Gamma} \Gamma\|_{\infty} \le 7.5 \|\mathbf{E}\|_{2,\infty}^p$ .
- Every entry in Γ bigger than  $7.5 \| \mathbf{E} \|_{2.\infty}^p$  is found.
- The solution  $\hat{\Gamma}$  is unique.

Note that the expression on the number of nonzeros permitted

is similar to the one in the classic Sparseland analysis, being  $O(1/\mu(\mathbf{D}))$ . However, in this theorem this quantity refers to the allowed number of nonzeros in each stripe, which means that the overall number of permitted nonzeros in  $\Gamma$  becomes proportional to O(N). As such, this is a much stronger and more practical outcome, providing the necessory property of the property and the transfer of the property of the proper

sary guarantee for various recent works that used the BP with the CSC model for various applications [59]–[64].

## CSC: Operating locally while getting global optimality

The last topic we would like to address in this section is the matter of computing the global BP solution for the CSC model while operating locally on small patches. As we are about to see, this serves as a clarifying bridge to traditional image processing algorithms that operate on patches. We discuss one such algorithm, originally presented in [65]—the slice-based pursuit.

To present this algorithm, we break  $\Gamma$  into small nonoverlapping blocks of length m each, denoted as  $\alpha_i$  and termed *needles*. A key observation this method relies on is the ability to decompose the global signal  $\mathbf{X}$  into the sum of small pieces, which we call *slices*, given by  $\mathbf{s}_i = \mathbf{D}_L \alpha_i$ . By positioning each of these in their location in the signal canvas, we can construct the full vector  $\mathbf{X}$ :

$$\mathbf{X} = \mathbf{D}\mathbf{\Gamma} = \sum_{i=1}^{m} \mathbf{R}_{i}^{T} \mathbf{D}_{L} \boldsymbol{\alpha}_{i} = \sum_{i=1}^{m} \mathbf{R}_{i}^{T} \mathbf{s}_{i}.$$
 (14)

By being the transposed of the patch-extraction operator,  $\mathbf{R}_i^T \in \mathbb{R}^{N \times n}$  places an *n*-dimensional patch into its corresponding location in the *N*-dimensional signal  $\mathbf{X}$  by padding it with

N-n zeros. Then, armed with this new way to represent **X**, the BP penalty can be restated in terms of the needles and the slices,

$$\min_{\mathbf{s}:\mathbf{j}_{i},(\boldsymbol{\alpha})\neq 2} \left\| \mathbf{Y} - \sum_{i=1}^{m} \mathbf{R}_{i}^{T} \mathbf{s}_{i} \right\|_{2}^{2} + \lambda \sum_{i=1}^{m} \|\boldsymbol{\alpha}_{i}\|_{1}$$
s.t.  $\{\mathbf{s}_{i} = \mathbf{D}_{L}\boldsymbol{\alpha}_{i}\}_{i=1}^{N}$ . (15)

This problem is entirely equivalent to the original BP penalty, being just as convex. Using the alternating direction method of multipliers (ADMM) [66] to handle the constraints (see the details in [65]), we obtain a global pursuit algorithm that iterates between local BP on each slice (that is easily managed by least-angle regression (LARS) [67] or any other efficient solver), and an update of the slices that aggregates the temporal results into one estimated global signal. Interestingly, if this algorithm initializes the slices to be patches from Y and applies only one such round of operations, the resulting process aligns with the patch-based sparsity-inspired image denoising algorithm [68]. Figure 4 shows the slices and their corresponding patches in an image, and as can be clearly seen, the slices are simpler and thus easier to represent sparsely.

The aforementioned scheme can be extended to serve for the training of the CSC filters,  $\mathbf{D}_L$ . All that is needed is to insert a "dictionary-update" stage in each iteration, in which we update

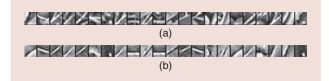
 $\mathbf{D}_L$  based on the given slices. This stage can be performed using the regular K-SVD algorithm [69] or any other dictionary-learning alternative, where the patches to train on are these slices. As such, the overall algorithm relies on local operations and can therefore leverage all the efficient tools developed for Sparseland over the last 15 years, while still

solving the global CSC model—its learning and pursuit.

## ML convolutional sparse modeling

At last, we arrive to the main serving of this article: connecting Sparseland and the CSC model to deep learning and CNNs. Preliminary signs of this connection could already be vaguely identified, as there are some similarities between these two disciplines. More specifically, both involve the presence of convolution operations, sparsifying operations such as shrinkage or rectified linear unit (ReLU), and both rely on learning from data to better fit the treatment to the given information source.

These early signs did not go unnoticed, and they motivated a series of fascinating contributions that aimed to bring an explanation to CNNs. For instance, Bruna and Mallat [4] borrowed



**FIGURE 4.** A comparison between (a) patches and (b) their respective slices, extracted from natural images. As can be seen, the slices are much simpler, and thus easier to represent sparsely.

The CSC model offers a

path toward answering all

of the questions that we

of the global-local gap.

have posed in the context

elements from wavelet theory to demonstrate how one can build a network of convolutional operators while providing invariance to shifts and deformations—properties that deep CNNs are claimed to have. Another interesting line of work comes from the observation that several pursuit algorithms can be decomposed as several iterations of linear operators and a nonlinear threshold, and therefore allow for a CNN-based implementation. The seminal learned iterative soft thresholding algorithm (LISTA) [70] showed that one can indeed train such a network by unrolling iterative shrinkage iterations, while achieving significantly faster convergence. In fact, CNN-based pursuits can even be shown to outperform traditional sparse coding methods in some challenging cases [71]. In this section our goal is to make this connection much more precise and principled, and thus we start by briefly recalling the way CNNs operate.

## A brief review of conv-nets

We shall assume that we are given an input image  $\mathbf{Y} \in \mathbb{R}^N$ , of dimensions  $\sqrt{N} \times \sqrt{N}$ . A classic feed-forward CNN operates on  $\mathbf{Y}$  by applying series of convolutions and nonlinearities [72]. Our goal in this section is to clearly formulate these steps.

In the first layer,  $\mathbf{Y}$  is passed through a set of  $m_1$  convolutions, using small-support filters of size  $\sqrt{n_0} \times \sqrt{n_0}$ . In this case as well, these convolutions are assumed cyclic. The output of these convolutions is augmented by a bias value and then passed through a scalar-wise ReLU of the form ReLU(z) = max(0,z), and the result  $\mathbf{Z}_1$  is stored in the form of a 3-D tensor of size  $\sqrt{N} \times \sqrt{N} \times m_1$ , as illustrated in Figure 5. In matrix-vector form, we could say that  $\mathbf{Y}$  has been multiplied by a convolutional matrix  $\mathbf{W}_1$  of size  $Nm_1 \times N$ , built by concatenating vertically a set of  $m_1$  circulant matrices of size  $N \times N$ . This is followed by the ReLU step:  $\mathbf{Z}_1$  = Re LU( $\mathbf{W}_1\mathbf{Y} + \mathbf{b}_1$ ). Note that  $\mathbf{b}_1$  is a vector of length  $Nm_1$ , applying a different threshold per each filter in the resulting tensor.

The second layer obtains the tensor  $\mathbf{Z}_1$  and applies the same set of operations:  $m_2$  convolutions with small spatial support  $(\sqrt{n_1} \times \sqrt{n_1})$  and across all  $m_1$  channels, and a ReLU nonlin-

earity. Each such filter weights together (i.e., this implements a 2-D convolution across all feature maps) the  $m_1$  channels of  $\mathbf{Z}_1$ , which explains the length mentioned previously. Thus, the output of the second layer is given by  $\mathbf{Z}_2 = \text{Re LU}(\mathbf{W}_2\mathbf{Z}_1 + \mathbf{b}_2)$ , where  $\mathbf{W}_2$  is a vertical amalgam of  $m_2$  convolutional matrices of size  $N \times Nm_1$ .

The story may proceed this way with additional layers sharing the same functional structure. A variation could be pooling or stride operations, both coming to reduce the spatial resolution of the resulting tensor. In this article, we focus on the stride option, which is easily absorbed within the aforementioned description by subsampling the rows in the corresponding convolution matrices. We note that recent work suggests that pooling can be replaced by stride without a performance degradation ([73], [74]).

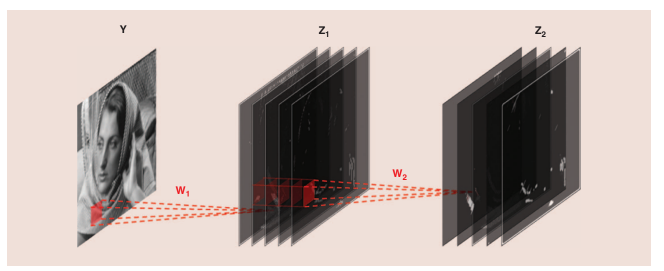
To summarize, for the two layered feed-forward CNNs, the relation between input and output is given by

$$f(\mathbf{Y}) = \operatorname{Re} \operatorname{LU}(\mathbf{W}_{2} \operatorname{Re} \operatorname{LU}(\mathbf{W}_{1} \mathbf{Y} + \mathbf{b}_{1}) + \mathbf{b}_{2}). \tag{16}$$

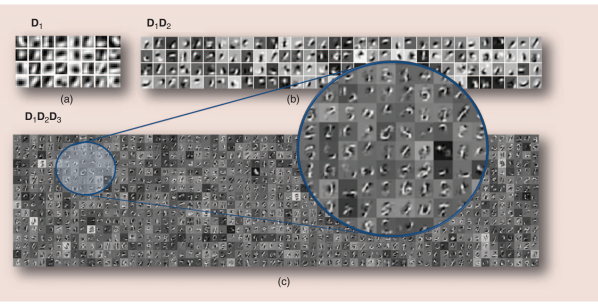
# Introducing the ML-CSC model

We return now to the CSC model with an intention to extend it by introducing its ML version. Our starting point is the belief that the ideal signal we operate on,  $\mathbf{X}$ , is emerging from the CSC model, just as described in the section "Convolutional Sparse Modeling." Thus,  $\mathbf{X} = \mathbf{D}_1 \mathbf{\Gamma}_1$ , where  $\mathbf{D}_1$  is a convolutional dictionary of size  $N \times Nm_1$ , and  $\mathbf{\Gamma}_1$  is locally sparse, i.e.,  $\|\mathbf{\Gamma}_1\|_{0,\infty}^{|\mathbf{S}|} = k_1 \ll m_1(2n_0 - 1)$ , where  $m_1$  is the number of filters in this model and  $n_0$  is their length.

We now add another layer to this signal: We assume that  $\Gamma_1$  itself is also believed to be produced from a CSC model of a similar form,  $\Gamma_1 = \mathbf{D}_2 \Gamma_2$ , where  $\mathbf{D}_2$  is another convolutional dictionary of size  $Nm_1 \times Nm_2$ , and  $\|\Gamma_2\|_{0,\infty}^s = k_2 \ll m_2(2n_1m_1 - 1)$ , where  $m_2$  is the number of filters in this model and  $n_1$  is their length. This structure can be cascaded in the same manner for K layers, leading to what we refer to as the ML-CSC model. A formal definition of this model is given next.



**FIGURE 5.** An illustration of the forward pass of a CNN. The first feature map is given by  $\mathbf{Z}_1 = \text{Re} \, \text{LU} (\mathbf{W}_1 \mathbf{Y} + \mathbf{b}_1)$ , where  $\mathbf{W}_1$  is a convolutional operator.



**FIGURE 6.** An ML-CSC model trained on the MNIST data set. (a) The local filters of the dictionary  $\mathbf{D}_1$ . (b) The local filters of the effective dictionary  $\mathbf{D}^{(2)} = \mathbf{D}_1 \mathbf{D}_2$ . (c) Some of the 1,024 local atoms of the effective dictionary  $\mathbf{D}^{(3)}$ , which are global atoms of size  $28 \times 28$ .

## Definition

The set of *K*-layered ML-CSC signals of cardinalities  $\{k_1, k_2, ..., k_K\}$  over the convolutional dictionaries  $\{\mathbf{D}_1, \mathbf{D}_2, ..., \mathbf{D}_K\}$  is defined as  $\mathcal{MS}\{\{\mathbf{D}_i\}_{i=1}^K, \{k_i\}_{i=1}^K\}$ . A signal **X** belongs to this set if it can be described as

$$\mathbf{X} = \mathbf{D}_{1} \Gamma_{1} \quad \text{s.t.} \quad \| \Gamma_{1} \|_{0,\infty}^{s} \leq k_{1}$$

$$\Gamma_{1} = \mathbf{D}_{2} \Gamma_{2} \quad \text{s.t.} \quad \| \Gamma_{2} \|_{0,\infty}^{s} \leq k_{2}$$

$$\Gamma_{2} = \mathbf{D}_{3} \Gamma_{3} \quad \text{s.t.} \quad \| \Gamma_{3} \|_{0,\infty}^{s} \leq k_{3}$$

$$\vdots$$

$$\Gamma_{K-1} = \mathbf{D}_{K} \Gamma_{K} \quad \text{s.t.} \quad \| \Gamma_{K} \|_{0,\infty}^{s} \leq k_{K}. \tag{17}$$

Observe that, given the aforementioned relations, one obviously gets  $\mathbf{X} = \mathbf{D}_1 \mathbf{D}_2 \cdots \mathbf{D}_K \Gamma_K$ . Thus, our overall model is a special case of Sparseland, with an effective dictionary being the multiplication of all the CSC matrices  $\{\mathbf{D}_i\}_{i=1}^K$ . It can be shown that this effective dictionary has an overall convolutional form, as shown in [75], and thus we could even go as far as saying that ML-CSC is a special case of the CSC model.

Indeed, ML-CSC injects more structure into the CSC model by requiring all the intermediate representations,  $\Gamma_1, \Gamma_2, ..., \Gamma_{K-1}$  to be locally sparse as well. The implication of this assumption is the belief that  $\mathbf{X}$  could be composed in various ways, all leading to the same signal. In its most elementary form,  $\mathbf{X}$  can be built from a sparse set of atoms from  $\mathbf{D}_1$  by  $\mathbf{D}_1\Gamma_1$ . However, the very same vector  $\mathbf{X}$  could be built using yet another sparse set of elements from the effective dictionary  $\mathbf{D}_1\mathbf{D}_2$  by  $\mathbf{D}_1\mathbf{D}_2\Gamma_2$ . Which are the atoms in this effective dictionary? Every column in  $\mathbf{D}_1\mathbf{D}_2$  is built as a local linear combination of atoms from  $\mathbf{D}_1$ , whose coefficients are the atoms in  $\mathbf{D}_2$ . Moreover, due to the locality of the atoms in  $\mathbf{D}_2$ , such combinations are only of atoms in  $\mathbf{D}_1$  that are spatially close. Thus, we could refer to the atoms of  $\mathbf{D}_1\mathbf{D}_2$ 

as *molecules*. The same signal can be described this way using  $\mathbf{D}_1\mathbf{D}_2\mathbf{D}_3$ , and so on, all the way to  $\mathbf{D}_1\mathbf{D}_2\cdots\mathbf{D}_K\Gamma_K$ .

Perhaps the following explanation could help in giving intuition to this wealth of descriptions of  $\mathbf{X}$ . A human-being body can be described as a sparse combination of atoms, but he/she could also be described as a sparse combination of molecules, a sparse composition of cells, tissues, and even body parts. There are many ways to describe the formation of the human body, all leading to the same final construction, and each adopting a different resolution of fundamental elements. The same spirit exists in the ML-CSC model.

To better illustrate the ML-CSC and how it operates, Figure 6 presents a three-layered CSC model trained on the MNIST database of handwritten digits. The results presented here refer to a dictionary-learning task for the ML-CSC, as described in [75]. We will not explain the details of the algorithm to obtain these dictionaries, but rather concentrate on the results obtained. This figure shows the  $m_1 = 32$  filters that construct the dictionary  $\mathbf{D}_1$ , and these look like crude atoms identifying edges or blobs. It also shows the  $m_2 = 128$  filters corresponding to the effective dictionary  $\mathbf{D}_1\mathbf{D}_2$ , and its fundamental elements are elongated and curved edges. Note that there is no point in presenting  $\mathbf{D}_2$ alone as it has almost no intuitive meaning. When multiplied by  $\mathbf{D}_1$ , it results with filters that are referred to as *molecules*. In the same spirit, the  $m_3 = 1,024$  filters of  $\mathbf{D}_1 \mathbf{D}_2 \mathbf{D}_3$  contain large portions of digits as its atoms. Any given digit image could be described as a sparse combination over  $\mathbf{D}_1$ ,  $\mathbf{D}_1\mathbf{D}_2$ , or  $\mathbf{D}_1\mathbf{D}_2\mathbf{D}_3$ , in each case using different fundamental elements set to form the construction.

# Pursuit algorithms for ML-CSC signals: First steps

Pursuit algorithms were mentioned throughout this article, making their first appearance in the context of Sparseland, and later

appearing again in the CSC. The pursuit task is essentially a projection operation, seeking the signal closest to the given data while belonging to the model, be it Sparseland, the CSC model, or the ML-CSC. When dealing with a signal  $\mathbf{X}$  believed to belong to the ML-CSC model, a noisy signal  $\mathbf{Y} = \mathbf{X} + \mathbf{E}$  ( $\|\mathbf{E}\|_2 \le \epsilon$ ) is projected to the model by solving the following pursuit problem:

Find 
$$\{\boldsymbol{\Gamma}_{i}\}_{i=1}^{K}$$
 s.t. 
$$\begin{cases} \|\mathbf{Y} - \mathbf{D}_{1}\boldsymbol{\Gamma}_{1}\|_{2} \leq \epsilon &, & \|\boldsymbol{\Gamma}_{1}\|_{0,\infty}^{s} \leq k_{1} \\ \boldsymbol{\Gamma}_{1} = \mathbf{D}_{2}\boldsymbol{\Gamma}_{2} &, & \|\boldsymbol{\Gamma}_{2}\|_{0,\infty}^{s} \leq k_{2} \\ \boldsymbol{\Gamma}_{2} = \mathbf{D}_{3}\boldsymbol{\Gamma}_{3} &, & \|\boldsymbol{\Gamma}_{3}\|_{0,\infty}^{s} \leq k_{3} \\ \vdots & & & \\ \boldsymbol{\Gamma}_{K-1} = \mathbf{D}_{K}\boldsymbol{\Gamma}_{K} &, & \|\boldsymbol{\Gamma}_{K}\|_{0,\infty}^{s} \leq k_{K} \end{cases}$$
(18)

We refer to this as the *deep coding problem (DCP)*. Figure 7 illustrates a typical result of such a problem, still in the context of the MNIST database. As before, we present these without diving into the details of how this solution was obtained. See [75] for this information. As can be seen, the very same signal  $\mathbf{X}$  is created by sparsely combining 209 atoms from  $\mathbf{D}_1$ , or very sparsely combining 47 molecules from  $\mathbf{D}_1\mathbf{D}_2$ , or extremely sparsely merging ten body parts from the dictionary  $\mathbf{D}_1\mathbf{D}_2\mathbf{D}_3$ .

The DCP problem is NP-hard. Indeed, a single layer version of it is known to be NP-hard [76], and the additional constraints can be merged into an overall problem that can be given the same

structure as the single layer one, thus exposing its NP-hardness. Therefore, just as in the general atom-decomposition problem discussed in the section "Sparse Modeling," approximation algorithms are required for its solution. We are interested in making our first steps in developing such a pursuit algorithm, and thus our starting point is an adaptation of the thresholding algorithm, due to its simplicity.

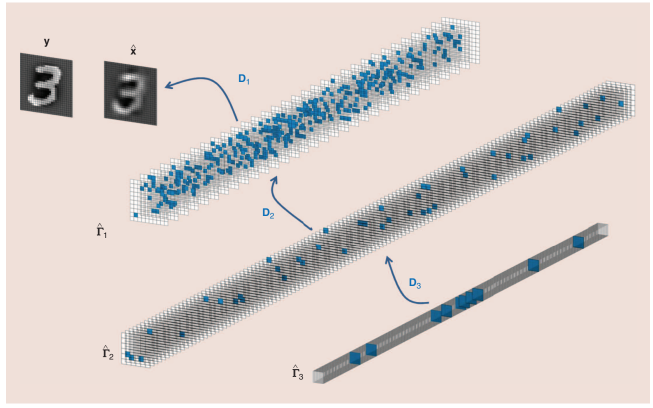
For a single-layer CSC model ( $\mathbf{Y} = \mathbf{D}_1 \mathbf{\Gamma}_1 + \mathbf{E}$  where  $\|\mathbf{\Gamma}_1\|_{0,\infty}^s \leq k_1$ ), the thresholding algorithm multiplies  $\mathbf{Y}$  by  $\mathbf{D}_1^T$ , and applies a simple shrinkage on the resulting vector, nulling small entries and leaving the rest almost untouched. This shrinkage can admit one of several forms, as shown in Figure 8: hard thresholding, soft thresholding, and one-sided soft thresholding, if the representation  $\mathbf{\Gamma}_1$  is assumed to be nonnegative. Put formally, this implies that the estimated representation is given by

$$\hat{\Gamma}_1 = \mathcal{T}_{T_1}(\mathbf{D}_1^T \mathbf{Y}), \tag{19}$$

where the operator  $\mathcal{T}_{T_1}(\mathbf{V})$  operates element-wise, thresholding the entries of the vector  $\mathbf{V}$  by the values found in the vector  $T_1$ .

As we add more layers to the CSC model, we can apply the thresholding algorithm sequentially, each layer at a time. Thus, once  $\Gamma_1$  has been estimated, we can now apply a second thresholding algorithm to evaluate  $\Gamma_2$  by

$$\hat{\boldsymbol{\Gamma}}_2 = \mathcal{T}_{T_2}(\mathbf{D}_2^T \hat{\boldsymbol{\Gamma}}_1) 
= \mathcal{T}_{T_2}(\mathbf{D}_2^T \mathcal{T}_{T_1}(\mathbf{D}_1^T \mathbf{Y})).$$
(20)



**FIGURE 1.** Decompositions of an image from MNIST in terms of its nested sparse features  $\gamma_i$  and multilayer convolutional dictionaries  $D_i$ .

This can be continued all the way to the Kth layer. Comparing (20) with the one written previously for the feed-forward CNN,

$$f(\mathbf{Y}) = \text{ReLU}(\mathbf{W}_2 \text{ReLU}(\mathbf{W}_1 \mathbf{Y} + \mathbf{b}_1) + \mathbf{b}_2), \tag{21}$$

reveals a striking resemblance. These two formulas express the same procedure: a set of convolutions applied on  $\mathbf{Y}$ , followed by a nonlinearity with a threshold, and proceeding this way as we dive into inner layers. The transposed dictionary  $\mathbf{D}_i$  plays the role of a set of convolutions, the threshold  $T_k$  parallels the bias vector  $\mathbf{b}_k$ , and the shrinkage operation stands for the ReLU. So, the inevitable conclusion is this: Assuming that our signals

emerge from the ML-CSC model, the layered-thresholding algorithm for decomposing a given measurements vector **Y** is completely equivalent to the forward pass in CNNs.

The pursuit algorithm we have just presented, or the forward pass of the CNN for that matter, estimates the sparse representations  $\{\Gamma_k\}_{k=1}^K$  that explain our signal. Why bother computing these hidden vectors? An additional assumption in our story is the fact that the labels associated with the data we are given are believed to depend on these represented.

sentations in a linearly separable way, and thus given  $\{\hat{\Gamma}_k\}_{k=1}^K$ , classification (or regression) is facilitated. In this spirit, a proper estimation of these vectors implies better classification results.

# ML-CSC: A path to theoretical analysis of CNNs

The conclusion about the relation between the CNN's forward-pass and a layered-thresholding pursuit algorithm is thrilling by itself, but this connection has far more profound consequences. Now that we consider the input signal  $\mathbf{X}$  as belonging to the ML-CSC model ( $\mathbf{X} \in \mathcal{MS}\{\{\mathbf{D}_i\}_{i=1}^K, \{k_i\}_{i=1}^K\}\}$ ), given a noisy version of it,  $\mathbf{Y} = \mathbf{X} + \mathbf{E}$ , the goal of the pursuit is quite clear—approximating the sparse representations  $\{\boldsymbol{\Gamma}_i\}_{i=1}^K$  explaining this signal's construction. Thus, we may pose intriguing new questions such as whether the sought representations are stably recoverable by the layered-thresholding algorithm.

Put more broadly, the model we have imposed on the input signals enables a new path of theoretical study of CNNs of unprecedented form, analyzing the performance of given architectures, and perhaps suggesting new ones in a systematic fashion. In the following, we give a taste from this new theoretical path, by analyzing the performance of the layered-thresholding algorithm, and considering alternatives to it.

Let us start by giving an answer to the question posed about the prospects of success of the layered-thresholding algorithm, in the form of the following theorem.

#### Theorem 3

Given  $\mathbf{Y} = \mathbf{X} + \mathbf{E}$ , where  $\mathbf{X} \in \mathcal{MS}(\{\mathbf{D}_i\}_{i=1}^K, \{k_i\}_{i=1}^K\})$  and  $\mathbf{E}$  is a bounded noise disturbance, if  $\Gamma_i$  are sufficiently locally sparse,

$$\|\mathbf{\Gamma}_{i}\|_{0,\infty}^{s} < \frac{1}{2} \left(1 + \frac{1}{\mu(\mathbf{D}_{i})} \cdot \frac{|\mathbf{\Gamma}_{i}^{\min}|}{|\mathbf{\Gamma}_{i}^{\max}|}\right) - \frac{1}{\mu(\mathbf{D}_{i})} \cdot \frac{\epsilon_{i}}{|\mathbf{\Gamma}_{i}^{\max}|}, \quad (22)$$

then the layered-hard thresholding algorithm (using the proper thresholds) finds the correct supports of all these representations, and in addition,  $\|\hat{\mathbf{\Gamma}}_i - \mathbf{\Gamma}_i\|_{2,\infty}^p \le \epsilon_i^2$ , where we define  $\epsilon_0 = \|\mathbf{E}\|_{2,\infty}^p$  and for  $1 \le i \le K$ ,

$$\epsilon_{i} = \sqrt{\|\mathbf{\Gamma}_{i}\|_{0,\infty}^{p}} \cdot \left(\epsilon_{i-1} + \mu(\mathbf{D}_{i})(\|\mathbf{\Gamma}_{i}\|_{0,\infty}^{s} - 1)\|\mathbf{\Gamma}_{i}^{\max}\|\right). \tag{23}$$

Putting many details aside, the main message we get here is quite exciting: we obtain a success guarantee for the forward-pass of CNN to find the proper locations of the nonzeros in all the representations  $\{\Gamma_i\}_{i=1}^K$ . Furthermore, the conditions for this success rely on the local sparsity of these representations and the mutual

coherence of the dictionaries involved.

On the downside, however, we do see several problems with the aforementioned result. First, observe that the conditions for success depend on the contrast factor  $|\Gamma_i^{\min}|/|\Gamma_i^{\max}|$  between the smallest and the largest nonzero in the representations. This implies that, for high-contrasted vectors, the conditions for success are much stricter. This sensitivity is not an artifact of the analysis, but rather a true and known difficulty that the thresholding algorithm carries with it.

Another troubling issue is an error growth that is seen as we proceed through the layers of the model. This growth is expected, due to the sequentiality of the layered-thresholding algorithm, propagating the errors and magnifying them from one layer to the next. Indeed, another problem is the fact that even if **E** is zero, namely, we are lucky to operate on a pure ML-CSC signal, the layered-thresholding algorithm induces an error in the inner layers, just as well.

# ML-CSC: Better pursuit and implications

In the previously given analysis, we have exposed good features of the layered thresholding, alongside some sensitivities and weaknesses. Recall that the thresholding algorithm is the

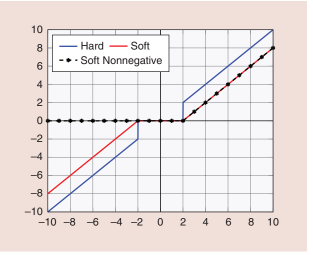


FIGURE 8. Hard-, soft-, and one-sided soft-thresholding operators.

**Assuming that our signals** 

emerge from the ML-CSC

thresholding algorithm

for decomposing a given

measurements vector Y is

completely equivalent to

the forward pass in CNNs.

model, the lavered-

simplest and crudest possible pursuit for sparsity-based signals. Why should we stick to it, if we are aware of better techniques? This takes us to the next discussion, in which we propose an alternative layered pursuit, based on the BP.

Consider the first layer of the ML-CSC model, described by the relations  $\mathbf{X} = \mathbf{D}_1 \mathbf{\Gamma}_1$  and  $\|\mathbf{\Gamma}_1\|_{0,\infty}^s \le k_1$ . This is, in fact, a regular CSC model, for which Theorem 2 provides the terms of success of the BP, when applied on a noisy version of such a signal,  $\mathbf{Y} = \mathbf{X} + \mathbf{E}$ . Thus, solving

$$\hat{\mathbf{\Gamma}}_{1} = \arg\min_{\Gamma_{1}} \frac{1}{2} \| \mathbf{Y} - \mathbf{D}_{1} \mathbf{\Gamma}_{1} \|_{2}^{2} + \lambda_{1} \| \mathbf{\Gamma}_{1} \|_{1}$$
 (24)

with a properly chosen  $\lambda_1$  is guaranteed to perform well, giving a stable estimate of  $\Gamma_1$ .

Consider now the second layer in the ML-CSC model, characterized by  $\Gamma_1 = \mathbf{D}_2 \Gamma_2$  and  $\|\Gamma_2\|_{0,\infty}^s \le k_2$ . Even though we are not given  $\Gamma_1$ , but rather a noisy version of it,  $\hat{\Gamma}_1$ , our analysis of the first stage provides an assessment of the noise power in this estimate. Thus, a second BP can be performed,

$$\hat{\mathbf{\Gamma}}_{2} = \arg\min_{\mathbf{\Gamma}_{2}} \frac{1}{2} \| \hat{\mathbf{\Gamma}}_{1} - \mathbf{D}_{2} \mathbf{\Gamma}_{2} \|_{2}^{2} + \lambda_{2} \| \mathbf{\Gamma}_{2} \|_{1}$$
 (25)

with a properly chosen  $\lambda_2$ , and this again leads to a guaranteed stable estimate of  $\Gamma_2$ .

We may proceed in this manner, proposing the layered-BP algorithm. Interestingly, such an algorithmic structure, coined *deconvolutional networks* [77], was proposed in the deep-learning literature in 2010, without a solid theoretical justification. Could we offer such a justification, now that our point of view is model based? The following theorem offers terms for the success of the layered-BP algorithm.

#### Theorem 4

Given  $\mathbf{Y} = \mathbf{X} + \mathbf{E}$ , where  $\mathbf{X} \in \mathcal{MS}\{\{\mathbf{D}_i\}_{i=1}^K, \{k_i\}_{i=1}^K\}$  and  $\mathbf{E}$  is a bounded noise disturbance, if  $\Gamma_i$  are sufficiently locally sparse,

$$\|\mathbf{\Gamma}_i\|_{0,\infty}^s < \frac{1}{3} \left(1 + \frac{1}{\mu(\mathbf{D}_i)}\right),\tag{26}$$

then the layered BP (using the proper coefficients  $\lambda_i$ ) is guaranteed to perform well:

- The support of  $\hat{\Gamma}_i$  is contained within that of the original  $\Gamma_i$  for all  $1 \le i \le K$ .
- The estimation is stable,

$$\|\hat{\Gamma}_i - \Gamma_i\|_{2,\infty}^p \le \epsilon_i, \tag{27}$$

where

$$\epsilon_i = 7.5^i \| \mathbf{E} \|_{2,\infty}^p \cdot \prod_{j=1}^i \sqrt{\| \mathbf{\Gamma}_i \|_{0,\infty}^p} \quad \forall \quad 1 \le i \le K.$$
 (28)

Every entry in  $\Gamma_i$  greater than  $\epsilon_i / \sqrt{\|\Gamma_i\|_{0,\infty}^p}$  is detected and included in the support, for all  $1 \le i \le K$ .

As we can see, the terms of success of the layered BP are far better than those of the layered thresholding. In particular, this method is no longer sensitive to the contrast of the nonzeros in  $\Gamma_i$ , and in addition, if  ${\bf E}$  is zero, then this algorithm leads to a perfect recovery of all the sparse representations. On the down side, however, this algorithm also leads to an error growth as we dive into the layers if  ${\bf E} \neq 0$ . This is an artifact of our choice to perform the pursuit layer-wise sequentially, rather than holistically solving the entire problem in parallel. Indeed, the work reported in [75] offers an alternative, free of this flaw.

The last topic we would like to discuss in this section is the matter of the actual algorithm to use when implementing the layered-BP. Our goal is the solution of this chain of problems:

$$\left\{ \hat{\mathbf{\Gamma}}_{k} = \arg\min_{\Gamma} \frac{1}{2} \| \hat{\mathbf{\Gamma}}_{k-1} - \mathbf{D}_{k} \mathbf{\Gamma} \|_{2}^{2} + \lambda_{k} \| \mathbf{\Gamma} \|_{1} \right\}_{k=1}^{K}, \quad (29)$$

where we have defined  $\hat{\boldsymbol{\Gamma}}_0 = \mathbf{Y}$ .

An appealing approximate solver for the core BP problem  $\min_{\Gamma}(1/2)\|\mathbf{Y} - \mathbf{D}\Gamma\|_2^2 + \lambda \|\Gamma\|_1$  is the iterative soft thresholding algorithm (ISTA) [78], which applies the following iterative procedure (assuming the dictionary has been normalized so that  $\|\mathbf{D}\|_2 = 1$ )

$$\hat{\mathbf{\Gamma}}^{t} = \mathcal{T}_{\lambda} (\hat{\mathbf{\Gamma}}^{t-1} + \mathbf{D}^{T} (\mathbf{Y} - \mathbf{D}\hat{\mathbf{\Gamma}}^{t-1})). \tag{30}$$

Thus, if each of the K layers of the model is managed using J iterations of the ISTA algorithm, and these steps are unfolded to form a long sequential process, we get an architecture with  $K \cdot J$  layers of a linear convolution operation followed by an elementwise nonlinearity, very much reminiscent of the recurrent neural network structure. For example, for a model with K=10 layers and using J=50 iterations of ISTA, the network obtained contains 500 layers (with a recurrent structure), and this gives a very illuminating perspective to the depth of typical networks.

Observe that our insight, as developed here, suggests that each J iterations in this scheme should use the same dictionary  $\mathbf{D}_k$ , and thus when learning the network we can force this parameter sharing, reducing the number of overall learned parameters by factor 50. Moreover, recalling the comment on LISTA networks and their advantage in terms of number of unfolding [70], if one frees the convolutional operators from being defined by the respective dictionary, such a network could be implemented with significantly fewer layers.

## **Concluding remarks**

#### Summary

We started this article by highlighting the importance of models in data processing, and then turned to describe one of these in depth: Sparseland, a systematic and highly effective such model. We then moved to CSC, with the hope to better treat images and other high-dimensional signals while operating locally, and accompanied this migration with the introduction of a new theory to substantiate this model. This brought us naturally to the ML-CSC, which gave us a bridge to the realm of deep learning.

Specifically, we have shown that handling ML-CSC signals amounts to CNNs of various forms, all emerging naturally as deployments of pursuit algorithms. Furthermore, we have seen that this new way of looking at these architectures can be accompanied by an unprecedented ability to study their performance, and identify the weaknesses and strengths of the ingredients involved. This was the broad story of this article, and there are two main take-home messages emerging from it.

The backbone of our story is three parametric models for managing signals:

Sparseland, CSC, and ML-CSC. All three are generative models, offering an explanation of the data by means of how it is synthesized. We have seen that this choice of models is extremely beneficial for both designing algorithms for serving

such signals, and enabling their theoretical analysis.

■ The ML-CSC model puts forward an appealing way to explain the motivation and origin of some common deep-learning architectures. As such, this model and the line of thinking behind it poses a new platform for understanding and further developing deep-learning solutions. We hope that this work will provide some of the foundations for the much-desired theoretical justification of deep learning.

# Future research directions and open problems

The ideas we have outlined in this article open up numerous new directions for further research. Some of the more promising ones are given next.

■ Dictionary learning: A key topic that we have deliberately paid less attention to in this article is the need to train the obtained networks to achieve their processing goals. In the context of the ML-CSC model, this parallels the need to learn the dictionaries  $\{\mathbf{D}_k\}_{i=k}^K$  and the threshold vectors. Observe that all the theoretical study we have proposed relied on an unsupervised regime, in which the labels play no role. This should remind the readers of autoencoders, in which sparsity is the driving force behind the learning mechanism. Further work is required to extend the knowledge on dictionary learning, both practical and theoretical, to bring it to the ML-CSC model, and with this offer new ways to learn neural networks. The first such attempt appears in [75], and some of its results have been given in this article.

The idea of training a series of cascaded dictionaries for getting an architecture mimicking that of deep learning has in fact appeared in earlier work [77], [79]–[83]. However, these attempts took an applicative and practical point of view and were never posed within the context of a solid mathematical ML model of the data, as in this work. In [77] and [79], the authors learned a set of convolutional dictionaries over multiple layers of abstraction in an unsupervised manner. In [80] and [81], the authors suggested using backpropagation rules for learning ML (nonconvolutional) dictionaries for CIFAR-10 classification, motivated by earlier work [82], [83]

Put more broadly, the model we have imposed on the input signals enables a new path of theoretical study of CNNs of unprecedented form, analyzing the performance of given architectures, and perhaps suggesting new ones in a systematic fashion.

- that showed how this was possible for a single layer setting. We believe that some of these ideas could become helpful in developing novel ML dictionary-learning algorithms, while relating to our formal model, thus preserving the relevance of its theoretical analysis.
- Pursuit algorithms: We introduced layered versions of the thresholding and the BP, and both are clearly suboptimal when it comes to projecting a signal to the ML-CSC model. Could we do better than these? The answer is positive, and such a progress has already appeared in [75].
- Still, there is room for further improvements, both in handling the various layers in parallel and not sequentially, and also in better serving the local sparsity we are after. Recall that better pursuit algorithms implies new architectures and, with them, new horizons to the deep-learning practice.
- Theoretical analysis: The study presented here represents the very first steps in a road that could take us to far better and more informative performance bounds. The analysis we have shown is a worst-case one, and, as such, it is too pessimistic. Also, all of it relies strongly on the mutual coherence, a worst-case study that tends to present a gloomy perspective on the prospects of the investigated algorithms. In addition, as we migrate from the layered pursuit algorithms to more sophisticated solutions, we may expect better and more encouraging bounds.

More broadly, a complete theory for deep learning cannot limit itself to the data model and the architectures emerging from it, as this work does. Missing in our picture are answers to intriguing questions on the learning, optimization, and generalization performance facets of neural networks. We believe that these topics could be addressed while still relying on data models, of the kind posed in this work.

- Improving the model: Labels play no role in the study presented in this article, and we have replaced them by the quest to recover the proper supports of the ideal representations or vectors close to them in terms of the ℓ₂ norm. While we did explain the underlying assumption behind this line of thinking, a broader model that takes the labels into account is very much needed. Indeed, rather than modeling the incoming signals, perhaps we should focus our modeling on the function that connects these to their corresponding labels.
- Deployment to practice: There is no doubt in our minds that the true and ultimate test for the theory we present will be its ability to further push the practice and performance of deep learning. The provided explanations on the used CNN architectures is a great start, but it must be followed by better understanding of more advanced ideas such as pooling, batch-normalization, dropout, and many other ideas that were found useful in practice. Beyond these, we hope to see this theory lead to new ideas that are simply impossible to develop without the systematic approach that the model-based theory unravels.

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