Representation Learning: A Review and New Perspectives

Yoshua Bengio, Aaron Courville, and Pascal Vincent

Abstract—The success of machine learning algorithms generally depends on data representation, and we hypothesize that this is because different representations can entangle and hide more or less the different explanatory factors of variation behind the data. Although specific domain knowledge can be used to help design representations, learning with generic priors can also be used, and the quest for AI is motivating the design of more powerful representation-learning algorithms implementing such priors. This paper reviews recent work in the area of unsupervised feature learning and deep learning, covering advances in probabilistic models, autoencoders, manifold learning, and deep networks. This motivates longer term unanswered questions about the appropriate objectives for learning good representations, for computing representations (i.e., inference), and the geometrical connections between representation learning, density estimation, and manifold learning.

Index Terms—Deep learning, representation learning, feature learning, unsupervised learning, Boltzmann machine, autoencoder, neural nets

INTRODUCTION

THE performance of machine learning methods is heavily **L** dependent on the choice of data representation (or features) on which they are applied. For that reason, much of the actual effort in deploying machine learning algorithms goes into the design of preprocessing pipelines and data transformations that result in a representation of the data that can support effective machine learning. Such feature engineering is important but labor intensive and highlights the weakness of current learning algorithms: Their inability to extract and organize the discriminative information from the data. Feature engineering is a way to take advantage of human ingenuity and prior knowledge to compensate for that weakness. To expand the scope and ease of applicability of machine learning, it would be highly desirable to make learning algorithms less dependent on feature engineering so that novel applications could be constructed faster, and more importantly, to make progress toward artificial intelligence (AI). An AI must fundamentally understand the world around us, and we argue that this can only be achieved if it can learn to identify and disentangle the underlying explanatory factors hidden in the observed milieu of low-level sensory data.

This paper is about representation learning, i.e., learning representations of the data that make it easier to extract useful information when building classifiers or other predictors. In the case of probabilistic models, a good representation is often one that captures the posterior

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observed input. A good representation is also one that is useful as input to a supervised predictor. Among the various ways of learning representations, this paper focuses on deep learning methods: those that are formed by the composition of multiple nonlinear transformations with the goal of yielding more abstract—and ultimately more useful—representations. Here, we survey this rapidly developing area with special emphasis on recent progress. We consider some of the fundamental questions that have been driving research in this area. Specifically, what makes one representation better than another? Given an example, how should we compute its representation, i.e., perform feature extraction? Also, what are appropriate objectives for learning good representations?

distribution of the underlying explanatory factors for the

WHY SHOULD WE CARE ABOUT LEARNING REPRESENTATIONS?

Representation learning has become a field in itself in the machine learning community, with regular workshops at the leading conferences such as NIPS and ICML, and a new conference dedicated to it, ICLR, sometimes under the header of Deep Learning or Feature Learning. Although depth is an important part of the story, many other priors are interesting and can be conveniently captured when the problem is cast as one of learning a representation, as discussed in the next section. The rapid increase in scientific activity on representation learning has been accompanied and nourished by a remarkable string of empirical successes both in academia and in industry. Below, we briefly highlight some of these high points.

2.1 Speech Recognition and Signal Processing

Speech was one of the early applications of neural networks, in particular convolutional (or time-delay) neural

networks.² The recent revival of interest in neural networks, deep learning, and representation learning has had a strong impact in the area of speech recognition, with breakthrough results [54], [56], [183], [148], [55], [86] obtained by several academics as well as researchers at industrial labs bringing these algorithms to a larger scale and into products. For example, Microsoft released in 2012 a new version of their Microsoft Audio Video Indexing Service speech system based on deep learning [183]. These authors managed to reduce the word error rate on four major benchmarks by about 30 percent (e.g., from 27.4 to 18.5 percent on RT03S) compared to state-of-the-art models based on Gaussian mixtures for the acoustic modeling and trained on the same amount of data (309 hours of speech). The relative improvement in error rate obtained by Dahl et al. [55] on a smaller large-vocabulary speech recognition benchmark (Bing mobile business search dataset, with 40 hours of speech) is between 16 and 23 percent.

Representation-learning algorithms have also been applied to music, substantially beating the state of the art in polyphonic transcription [34], with relative error improvement between 5 and 30 percent on a standard benchmark of four datasets. Deep learning also helped to win MIREX (music information retrieval) competitions, for example, in 2011 on audio tagging [81].

2.2 Object Recognition

The beginnings of deep learning in 2006 focused on the MNIST digit image classification problem [94], [23], breaking the supremacy of SVMs (1.4 percent error) on this dataset.³ The latest records are still held by deep networks: Ciresan et al. [46] currently claim the title of state of the art for the unconstrained version of the task (e.g., using a convolutional architecture), with 0.27 percent error, and Rifai et al. [169] is state of the art for the knowledge-free version of MNIST, with 0.81 percent error.

In the last few years, deep learning has moved from digits to object recognition in natural images, and the latest breakthrough has been achieved on the ImageNet dataset,⁴ bringing down the state-of-the-art error rate from 26.1 to 15.3 percent [117].

2.3 Natural Language Processing (NLP)

Besides speech recognition, there are many other NLP applications of representation learning. *Distributed representations* for symbolic data were introduced by Hinton [87], and first developed in the context of statistical language modeling by Bengio et al. [19] in the so-called *neural net language models* [10]. They are all based on learning a distributed representation for each word, called a *word embedding*. Adding a convolutional architecture, Collobert et al. [51] developed the SENNA system⁵ that shares representations across the tasks of language modeling, part-of-speech tagging, chunking, named entity recognition, semantic role labeling, and syntactic parsing. SENNA

5. Downloadable from http://ml.nec-labs.com/senna/.

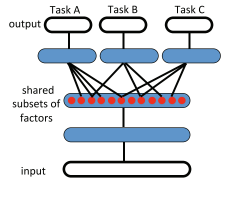


Fig. 1. Illustration of representation-learning discovering explanatory factors (middle hidden layer, in red), some explaining the input (semi-supervised setting), and some explaining target for each task. Because these subsets overlap, sharing of statistical strength helps generalization.

approaches or surpasses the state of the art on these tasks, but is simpler and much faster than traditional predictors. Learning word embeddings can be combined with learning image representations in a way that allows associating text and images. This approach has been used successfully to build Google's image search, exploiting huge quantities of data to map images and queries in the same space [218], and it has recently been extended to deeper multimodal representations [194].

The neural net language model was also improved by adding recurrence to the hidden layers [146], allowing it to beat the state of the art (smoothed n-gram models) not only in terms of perplexity (exponential of the average negative log likelihood of predicting the right next word, going down from 140 to 102), but also in terms of word error rate in speech recognition (since the language model is an important component of a speech recognition system), decreasing it from 17.2 percent (KN5 baseline) or 16.9 percent (discriminative language model) to 14.4 percent on the Wall Street Journal benchmark task. Similar models have been applied in statistical machine translation [182], [121], improving perplexity and BLEU scores. Recursive autoencoders (which generalize recurrent networks) have also been used to beat the state of the art in full sentence paraphrase detection [192], almost doubling the F1 score for paraphrase detection. Representation learning can also be used to perform word sense disambiguation [33], bringing up the accuracy from 67.8 to 70.2 percent on the subset of Senseval-3 where the system could be applied (with subject-verb-object sentences). Finally, it has also been successfully used to surpass the state of the art in sentiment analysis [70], [193].

2.4 Multitask and Transfer Learning, Domain Adaptation

Transfer learning is the ability of a learning algorithm to exploit commonalities between different learning tasks to share statistical strength and *transfer knowledge* across tasks. As discussed below, we hypothesize that representation learning algorithms have an advantage for such tasks because they learn representations that capture underlying factors, a subset of which may be relevant for each particular task, as illustrated in Fig. 1. This hypothesis seems confirmed by a number of empirical results showing

^{2.} See [9] for a review of early work in this area.

^{3.} For the knowledge-free version of the task where no image-specific prior is used, such as image deformations or convolutions.

^{4.} The 1,000-class ImageNet benchmark, whose results are detailed here: http://www.image-net.org/challenges/LSVRC/2012/results.html.

the strengths of representation learning algorithms in transfer learning scenarios.

Most impressive are the two transfer learning challenges held in 2011 and won by representation learning algorithms. First, the transfer learning challenge, presented at an ICML 2011 workshop of the same name, was won using unsupervised layerwise pretraining [12], [145]. A second transfer learning challenge was held the same year and won by Goodfellow et al. [72]. Results were presented at NIPS 2011's Challenges in Learning Hierarchical Models Workshop. In the related *domain adaptation* setup, the target remains the same but the input distribution changes [70], [43]. In the *multitask learning* setup, representation learning has also been found to be advantageous [117], [51] because of shared factors across tasks.

3 WHAT MAKES A REPRESENTATION GOOD?

3.1 Priors for Representation Learning in Al

In [16], one of us introduced the notion of AI tasks, which are challenging for current machine learning algorithms and involve complex but highly structured dependencies. One reason why explicitly dealing with representations is interesting is because they can be convenient to express many general priors about the world around us, i.e., priors that are not task specific but would be likely to be useful for a learning machine to solve AI tasks. Examples of such general-purpose priors are the following:

- *Smoothness*: Assumes that the function to be learned f is s.t. $x \approx y$ generally implies $f(x) \approx f(y)$. This most basic prior is present in most machine learning, but is insufficient to get around the curse of dimensionality; see Section 3.2.
- Multiple explanatory factors: The data generating distribution is generated by different underlying factors, and for the most part, what one learns about one factor generalizes in many configurations of the other factors. The objective to recover or at least disentangle these underlying factors of variation is discussed in Section 3.5. This assumption is behind the idea of distributed representations, discussed in Section 3.3.
- A hierarchical organization of explanatory factors: The concepts that are useful for describing the world around us can be defined in terms of other concepts, in a hierarchy, with more abstract concepts higher in the hierarchy, defined in terms of less abstract ones. This assumption is exploited with deep representations, elaborated in Section 3.4.
- Semi-supervised learning: With inputs X and target Y to predict, a subset of the factors explaining X's distribution explain much of Y, given X. Hence, representations that are useful for P(X) tend to be useful when learning $P(Y \mid X)$, allowing sharing of statistical strength between the unsupervised and supervised learning tasks, see Section 4.
- Shared factors across tasks: With many Ys of interest or many learning tasks in general, tasks (e.g., the corresponding $P(Y \mid X, \text{task})$) are explained by factors that are shared with other tasks, allowing sharing of statistical strengths across tasks, as

- discussed in the previous section (multitask and transfer learning, domain adaptation).
- Manifolds: Probability mass concentrates near regions that have a much smaller dimensionality than
 the original space where the data live. This is
 explicitly exploited in some of the autoencoder
 algorithms and other manifold-inspired algorithms
 described, respectively, in Sections 7.2 and 8.
- Natural clustering: Different values of categorical variables such as object classes are associated with separate manifolds. More precisely, the local variations on the manifold tend to preserve the value of a category, and a linear interpolation between examples of different classes in general involves going through a low-density region, i.e., $P(X \mid Y = i)$ for different *i* tend to be well separated and not overlap much. For example, this is exploited in the manifold tangent classifier (MTC) discussed in Section 8.3. This hypothesis is consistent with the idea that humans have *named* categories and classes because of such statistical structure (discovered by their brain and propagated by their culture), and machine learning tasks often involve predicting such categorical variables.
- Temporal and spatial coherence: Consecutive (from a sequence) or spatially nearby observations tend to be associated with the same value of relevant categorical concepts or result in a small move on the surface of the high-density manifold. More generally, different factors change at different temporal and spatial scales, and many categorical concepts of interest change slowly. When attempting to capture such categorical variables, this prior can be enforced by making the associated representations slowly changing, i.e., penalizing changes in values over time or space. This prior was introduced in [6] and is discussed in Section 11.3.
- Sparsity: For any given observation x, only a small fraction of the possible factors are relevant. In terms of representation, this could be represented by features that are often zero (as initially proposed by Olshausen and Field [155]), or by the fact that most of the extracted features are *insensitive* to small variations of x. This can be achieved with certain forms of priors on latent variables (peaked at 0), or by using a nonlinearity whose value is often flat at 0 (i.e., 0 and with a 0 derivative), or simply by penalizing the magnitude of the Jacobian matrix (of derivatives) of the function mapping input to representation. This is discussed in Sections 6.1.1 and 7.2.
- Simplicity of factor dependencies: In good high-level representations, the factors are related to each other through simple, typically linear dependencies. This can be seen in many laws of physics and is assumed when plugging a linear predictor on top of a learned representation.

We can view many of the above priors as ways to help the learner discover and *disentangle* some of the underlying (and a priori unknown) factors of variation that the data may reveal. This idea is pursued further in Sections 3.5 and 11.4.

3.2 Smoothness and the Curse of Dimensionality

For AI tasks such as vision and NLP, it seems hopeless to rely only on simple parametric models (such as linear models) because they cannot capture enough of the complexity of interest unless provided with the appropriate feature space. Conversely, machine learning researchers have sought flexibility in *local*⁶nonparametric learners such as kernel machines with a fixed generic local-response kernel (such as the Gaussian kernel). Unfortunately, as argued at length by Bengio and Monperrus [17], Bengio et al. [21], Bengio and LeCun [16], Bengio [11], and Bengio et al. [25], most of these algorithms only exploit the principle of local generalization, i.e., the assumption that the target function (to be learned) is smooth enough, so they rely on examples to explicitly map out the wrinkles of the target function. Generalization is mostly achieved by a form of local interpolation between neighboring training examples. Although smoothness can be a useful assumption, it is insufficient to deal with the curse of dimensionality because the number of such wrinkles (ups and downs of the target function) may grow exponentially with the number of relevant interacting factors when the data are represented in raw input space. We advocate learning algorithms that are flexible and nonparametric, but do not rely exclusively on the smoothness assumption. Instead, we propose to incorporate generic priors such as those enumerated above into representationlearning algorithms. Smoothness-based learners (such as kernel machines) and linear models can still be useful on top of such learned representations. In fact, the combination of learning a representation and kernel machine is equivalent to learning the kernel, i.e., the feature space. Kernel machines are useful, but they depend on a prior definition of a suitable similarity metric or a feature space in which naive similarity metrics suffice. We would like to use the data, along with very generic priors, to discover those features or, equivalently, a similarity function.

3.3 Distributed Representations

Good representations are expressive, meaning that a reasonably sized learned representation can capture a huge number of possible input configurations. A simple counting argument helps us to assess the expressiveness of a model producing a representation: How many parameters does it require compared to the number of input regions (or configurations) it can distinguish? Learners of one-hot representations, such as traditional clustering algorithms, Gaussian mixtures, nearest-neighbor algorithms, decision trees, or Gaussian SVMs, all require O(N) parameters (and/ or O(N) examples) to distinguish O(N) input regions. One could naively believe that one cannot do better. However, restricted Boltzmann machines (RBMs), sparse coding, autoencoders, or multilayer neural networks can all represent up to $O(2^k)$ input regions using only O(N)parameters (with k the number of nonzero elements in a

6. Local in the sense that the value of the learned function at x depends mostly on training examples $x^{(\ell)}s$ close to x.

sparse representation, and k = N in nonsparse RBMs and other dense representations). These are all distributed⁸ or sparse⁹ representations. The generalization of clustering to distributed representations is multiclustering, where either several clusterings take place in parallel or the same clustering is applied on different parts of the input, such as in the very popular hierarchical feature extraction for object recognition based on a histogram of cluster categories detected in different patches of an image [120], [48]. The exponential gain from distributed or sparse representations is discussed further in [11, Section 3.2, Fig. 3.2]. It comes about because each parameter (e.g., the parameters of one of the units in a sparse code or one of the units in an RBM) can be reused in many examples that are not simply near neighbors of each other, whereas with local generalization, different regions in input space are basically associated with their own private set of parameters, for example, as in decision trees, nearest neighbors, Gaussian SVMs, and so on. In a distributed representation, an exponentially large number of possible subsets of features or hidden units can be activated in response to a given input. In a single-layer model, each feature is typically associated with a preferred input direction, corresponding to a hyperplane in input space, and the code or representation associated with that input is precisely the pattern of activation (which features respond to the input, and how much). This is in contrast with a nondistributed representation such as the one learned by most clustering algorithms, for example, k-means, in which the representation of a given input vector is a one-hot code identifying which one of a small number of cluster centroids best represents the input.¹⁰

3.4 Depth and Abstraction

Depth is a key aspect to the representation learning strategies we consider in this paper. As we will discuss, deep architectures are often challenging to train effectively, and this has been the subject of much recent research and progress. However, despite these challenges, they carry two significant advantages that motivate our long-term interest in discovering successful training strategies for deep architectures. These advantages are: 1) deep architectures promote the *reuse* of features, and 2) deep architectures can potentially lead to progressively more *abstract* features at higher layers of representations (more removed from the data).

Feature reuse. The notion of reuse, which explains the power of distributed representations, is also at the heart of the theoretical advantages behind deep learning, i.e.,

9. Sparse representations: distributed representations where only a few of the elements can be varied at a time, i.e., k < N.

^{7.} We understand *nonparametric* as including all learning algorithms whose capacity can be increased appropriately as the amount of data and its complexity demands it, for example, including mixture models and neural networks where the number of parameters is a data-selected hyperparameter.

^{8.} Distributed representations, where k out of N representation elements or feature values can be independently varied; for example, they are not mutually exclusive. Each concept is represented by having k features being turned on or active, while each feature is involved in representing many concepts.

^{10.} As discussed in [11], things are only slightly better when allowing continuous-valued membership values, for example, in ordinary mixture models (with separate parameters for each mixture component), but the difference in representational power is still exponential [149]. The situation may also seem better with a decision tree, where each given input is associated with a one-hot code over the tree leaves, which deterministically selects associated ancestors (the path from root to node). Unfortunately, the number of different regions represented (equal to the number of leaves of the tree) still only grows linearly with the number of parameters used to specify it [15].

constructing multiple levels of representation or learning a hierarchy of features. The depth of a circuit is the length of the longest path from an input node of the circuit to an output node of the circuit. The crucial property of a deep circuit is that its number of paths, i.e., ways to reuse different parts, can grow exponentially with its depth. Formally, one can change the depth of a given circuit by changing the definition of what each node can compute, but only by a constant factor. The typical computations we allow in each node include: weighted sum, product, artificial neuron model (such as a monotone nonlinearity on top of an affine transformation), computation of a kernel, or logic gates. Theoretical results clearly show families of functions where a deep representation can be exponentially more efficient than one that is insufficiently deep [82], [83], [21], [16], [15]. If the same family of functions can be represented with fewer parameters (or more precisely with a smaller VC dimension), learning theory would suggest that it can be learned with fewer examples, yielding improvements in both computational efficiency (less nodes to visit) and statistical efficiency (fewer parameters to learn, and reuse of these parameters over many different kinds of inputs).

Abstraction and invariance. Deep architectures can lead to abstract representations because more abstract concepts can often be constructed in terms of less abstract ones. In some cases, such as in the convolutional neural network [133], we explicitly build this abstraction in via a pooling mechanism (see Section 11.2). More abstract concepts are generally invariant to most local changes of the input. That makes the representations that capture these concepts generally highly nonlinear functions of the raw input. This is obviously true of categorical concepts, where more abstract representations detect categories that cover more varied phenomena (e.g., larger manifolds with more wrinkles), and thus they potentially have greater predictive power. Abstraction can also appear in high-level continuous-valued attributes that are only sensitive to some very specific types of changes in the input. Learning these sorts of invariant features has been a long-standing goal in pattern recognition.

3.5 Disentangling Factors of Variation

Beyond being *distributed* and *invariant*, we would like our representations to *disentangle the factors of variation*. Different explanatory factors of the data tend to change independently of each other in the input distribution, and only a few at a time tend to change when one considers a sequence of consecutive real-world inputs.

Complex data arise from the rich interaction of many sources. These factors interact in a complex web that can complicate AI-related tasks such as object classification. For example, an image is composed of the interaction between one or more light sources, the object shapes, and the material properties of the various surfaces present in the image. Shadows from objects in the scene can fall on each other in complex patterns, creating the illusion of object boundaries where there are none, and can dramatically effect the perceived object shape. How can we cope with these complex interactions? How can we disentangle the objects and their shadows? Ultimately, we believe the approach we adopt for overcoming these challenges must leverage the data itself, using vast quantities of unlabeled

examples, to learn representations that separate the various explanatory sources. Doing so should give rise to a representation significantly more robust to the complex and richly structured variations extant in natural data sources for AI-related tasks.

It is important to distinguish between the related but distinct goals of learning invariant features and learning to disentangle explanatory factors. The central difference is the preservation of information. Invariant features, by definition, have reduced sensitivity in the direction of invariance. This is the goal of building features that are insensitive to variation in the data that are uninformative to the task at hand. Unfortunately, it is often difficult to determine a priori which set of features and variations will ultimately be relevant to the task at hand. Further, as is often the case in the context of deep learning methods, the feature set being trained may be destined to be used in multiple tasks that may have distinct subsets of relevant features. Considerations such as these lead us to the conclusion that the most robust approach to feature learning is to disentangle as many factors as possible, discarding as little information about the data as is practical. If some form of dimensionality reduction is desirable, then we hypothesize that the local directions of variation least represented in the training data should be first to be pruned out (as in principal components analysis (PCA), for example, which does it globally instead of around each example).

3.6 Good Criteria for Learning Representations?

One of the challenges of representation learning that distinguishes it from other machine learning tasks such as classification is the difficulty in establishing a clear objective or target for training. In the case of classification, the objective is (at least conceptually) obvious; we want to minimize the number of misclassifications on the training dataset. In the case of representation learning, our objective is far removed from the ultimate objective, which typically is learning a classifier or some other predictor. Our problem is reminiscent of the credit assignment problem encountered in reinforcement learning. We have proposed that a good representation is one that disentangles the underlying factors of variation, but how do we translate that into appropriate training criteria? Is it even necessary to do anything but maximize likelihood under a good model or can we introduce priors such as those enumerated above (possibly data-dependent ones) that help the representation better do this disentangling? This question remains clearly open but is discussed in more detail in Sections 3.5 and 11.4.

4 Building Deep Representations

In 2006, a breakthrough in feature learning and deep learning was initiated by Geoff Hinton and quickly followed up in the same year [94], [23], [161], and soon after by Lee et al. [134] and many more later. It has been extensively reviewed and discussed in [11]. A central idea, referred to as *greedy layerwise unsupervised pretraining*, was to learn a hierarchy of features one level at a time, using unsupervised feature learning to learn a new transformation at each level to be composed with the previously learned transformations; essentially, each iteration of

unsupervised feature learning adds one layer of weights to a deep neural network. Finally, the set of layers could be combined to initialize a deep supervised predictor, such as a neural network classifier, or a deep generative model, such as a deep Boltzmann machine (DBM) [176].

This paper is mostly about feature learning algorithms that can be used to form deep architectures. In particular, it was empirically observed that layerwise *stacking* of feature extraction often yielded better representations, for example, in terms of classification error [119], [65], quality of the samples generated by a probabilistic model [176], or in terms of the *invariance* properties of the learned features [71]. Whereas this section focuses on the idea of stacking single-layer models, Section 10 follows up with a discussion on joint training of all the layers.

After greedy layerwise unsupervised pretraining, the resulting deep features can be used either as input to a standard supervised machine learning predictor (such as an SVM) or as initialization for a deep supervised neural network (e.g., by appending a logistic regression layer or purely supervised layers of a multilayer neural network). The layerwise procedure can also be applied in a purely supervised setting, called the greedy layerwise supervised pretraining [23]. For example, after the first one-hiddenlayer multilayer perceptrons (MLP) is trained, its output layer is discarded and another one-hidden-layer MLP can be stacked on top of it, and so on. Although results reported in [23] were not as good as for unsupervised pretraining, they were nonetheless better than without pretraining at all. Alternatively, the *outputs* of the previous layer can be fed as extra inputs for the next layer (in addition to the raw input), as successfully done in [221]. Another variant [184] pretrains in a supervised way all the previously added layers at each step of the iteration, and in their experiments, this discriminant variant yielded better results than unsupervised pretraining.

Whereas combining single layers into a supervised model is straightforward, it is less clear how layers pretrained by unsupervised learning should be combined to form a better *unsupervised* model. We cover here some of the approaches to do so, but no clear winner emerges, and much work has to be done to validate existing proposals or improve them.

The first proposal was to stack pretrained RBMs into a deep belief network [94] or DBN, where the top layer is interpreted as an RBM and the lower layers as a directed sigmoid belief network. However, it is not clear how to approximate maximum likelihood training to further optimize this generative model. One option is the wake-sleep algorithm [94], but more work should be done to assess the efficiency of this procedure in terms of improving the generative model.

The second approach that has been put forward is to combine the RBM parameters into a DBM by basically halving the RBM weights to obtain the DBM weights [176]. The DBM can then be trained by approximate maximum likelihood as discussed in more detail later (Section 10.2). This joint training has brought substantial improvements, both in terms of likelihood and in terms of classification performance of the resulting deep feature learner [176].

Another early approach was to stack RBMs or autoencoders into a *deep autoencoder* [92]. If we have a series of encoder-decoder pairs $(f^{(i)}(\cdot), g^{(i)}(\cdot))$, then the overall encoder is the composition of the encoders, $f^{(N)}(\dots f^{(2)}(f^{(1)}(\cdot)))$, and the overall decoder is its "transpose" (often with transposed weight matrices as well), $g^{(1)}(g^{(2)}(\dots f^{(N)}(\cdot)))$. The deep autoencoder (or its regularized version, as discussed in Section 7.2) can then be jointly trained, with all the parameters optimized with respect to a global reconstruction error criterion. More work on this avenue clearly needs to be done, and it was probably avoided by fear of the challenges in training deep feedforward networks, discussed in Section 10 along with very encouraging recent results.

Yet another recently proposed approach to training deep architectures [154] is to consider the iterative construction of a *free energy function* (i.e., with no explicit latent variables, except possibly for a top-level layer of hidden units) for a deep architecture as the composition of transformations associated with lower layers, followed by top-level hidden units. The question is then how to train a model defined by an arbitrary parameterized (free) energy function. Ngiam et al. [154] have used hybrid Monte Carlo [153], but other options include contrastive divergence (CD) [88], [94], score matching [97], [99], denoising score matching [112], [210], ratio matching [98], and noise-contrastive estimation [80].

5 SINGLE-LAYER LEARNING MODULES

Within the community of researchers interested in representation learning, there has developed two broad parallel lines of inquiry: one rooted in probabilistic graphical models and one rooted in neural networks. Fundamentally, the difference between these two paradigms is whether the layered architecture of a deep learning model is to be interpreted as describing a probabilistic graphical model or as describing a computation graph. In short, are hidden units considered latent random variables or as computational nodes?

To date, the dichotomy between these two paradigms has remained in the background, perhaps because they appear to have more characteristics in common than separating them. We suggest that this is likely a function of the fact that much recent progress in both of these areas has focused on single-layer greedy learning modules and the similarities between the types of single-layer models that have been explored: mainly, the RBM on the probabilistic side and the autoencoder variants on the neural network side. Indeed, as shown by one of us [210] and others [200], in the case of the RBM, training the model via an inductive principle known as score matching [97] (to be discussed in Section 6.4.3) is essentially identical to applying a regularized reconstruction objective to an autoencoder. Another strong link between pairs of models on both sides of this divide is when the computational graph for computing representation in the neural network model corresponds exactly to the computational graph that corresponds to inference in the probabilistic model, and this happens to also correspond to the structure of graphical model itself (e.g., as in the RBM).

th in terms of likelihood and in terms of classification

The connection between these two paradigms becomes formance of the resulting deep feature learner [176].

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the case of a probabilistic model, exact inference typically becomes intractable. In the case of deep models, the computational graph diverges from the structure of the model. For example, in the case of a DBM, unrolling variational (approximate) inference into a computational graph results in a recurrent graph structure. We have performed preliminary exploration [179] of deterministic variants of deep autoencoders whose computational graph is similar to that of a DBM (in fact very close to the meanfield variational approximations associated with the Boltzmann machine), and that is one interesting intermediate point to explore (between the deterministic approaches and the graphical model approaches).

In the next few sections, we will review the major developments in single-layer training modules used to support feature learning and particularly deep learning. We divide these sections between (Section 6) the probabilistic models, with inference and training schemes that directly parameterize the generative—or decoding—pathway and (Section 7) the typically neural network-based models that directly parametrize the encoding pathway. Interestingly, some models like predictive sparse decomposition (PSD) [109] inherit both properties and will also be discussed (Section 7.2.4). We then present a different view of representation learning, based on the associated geometry and the manifold assumption, in Section 8.

First, let us consider an unsupervised single-layer representation learning algorithm spanning all three views: probabilistic, autoencoder, and manifold learning.

5.1 PCA

We will use probably the oldest feature extraction algorithm, PCA, to illustrate the probabilistic, autoencoder, and manifold views of representation learning. PCA learns a linear transformation $h = f(x) = W^T x + b$ of input $x \in \mathbb{R}^{d_x}$, where the columns of $d_x \times d_h$ matrix W form an orthogonal basis for the d_h orthogonal directions of greatest variance in the training data. The result is d_h features (the components of representation h) that are decorrelated. The three interpretations of PCA are the following: 1) It is related to probabilistic models (Section 6) such as probabilistic PCA, factor analysis, and the traditional multivariate Gaussian distribution (the leading eigenvectors of the covariance matrix are the principal components); 2) the representation it learns is essentially the same as that learned by a basic linear autoencoder (Section 7.2); and 3) it can be viewed as a simple linear form of linear manifold learning (Section 8), i.e., characterizing a lower dimensional region in input space near which the data density is peaked. Thus, PCA may be in the back of the reader's mind as a common thread relating these various viewpoints. Unfortunately, the expressive power of linear features is very limited: They cannot be stacked to form deeper, more abstract representations since the composition of linear operations yields another linear operation. Here, we focus on recent algorithms that have been developed to extract nonlinear features, which can be stacked in the construction of deep networks, although some authors simply insert a nonlinearity between learned single-layer linear projections [125], [43].

Another rich family of feature extraction techniques that this review does not cover in any detail due to space Authorized licensed use limited to: National Library Of Medicine (NLM). Downloaded on June 28,2023 at 18:56:01 UTC from IEEE Xplore. Restrictions apply.

constraints is independent component analysis or ICA [108], [8]. Instead, we refer the reader to [101], [103]. Note that while in the simplest case (complete, noise free) ICA yields linear features, in the more general case it can be equated with a linear generative model with non-Gaussian independent latent variables, similar to sparse coding (Section 6.1.1), which result in *nonlinear features*. Therefore, ICA and its variants like independent and topographic ICA [102] can and have been used to build deep networks [122], [125] (see Section 11.2). The notion of obtaining independent components also appears similar to our stated goal of disentangling underlying explanatory factors through deep networks. However, for complex real-world distributions, it is doubtful that the relationship between truly independent underlying factors and the observed high-dimensional data can be adequately characterized by a linear transformation.

PROBABILISTIC MODELS

From the probabilistic modeling perspective, the question of feature learning can be interpreted as an attempt to recover a parsimonious set of latent random variables that describe a distribution over the observed data. We can express as p(x,h) a probabilistic model over the joint space of the latent variables, h, and observed data or visible variables x. Feature values are conceived as the result of an inference process to determine the probability distribution of the latent variables given the data, i.e., $p(h \mid x)$, often referred to as the *posterior* probability. Learning is conceived in terms of estimating a set of model parameters that (locally) maximize the regularized likelihood of the training data. The probabilistic graphical model formalism gives us two possible modeling paradigms in which we can consider the question of inferring latent variables, directed and undirected graphical models, which differ in their parameterization of the joint distribution p(x, h), yielding major impact on the nature and computational costs of both inference and learning.

Directed Graphical Models

Directed latent factor models separately parameterize the conditional likelihood $p(x \mid h)$ and the *prior* p(h) to construct the joint distribution, $p(x,h) = p(x \mid h)p(h)$. Examples of this decomposition include: PCA [171], [206], sparse coding [155], sigmoid belief networks [152], and the newly introduced spike-and-slab sparse coding (S3C) model [72].

6.1.1 Explaining Away

Directed models often lead to one important property: explaining away, i.e., a priori independent causes of an event can become nonindependent given the observation of the event. Latent factor models can generally be interpreted as latent cause models, where the h activations cause the observed x. This renders the a priori independent h to be nonindependent. As a consequence, recovering the posterior distribution of h, $p(h \mid x)$ (which we use as a basis for feature representation), is often computationally challenging and can be entirely intractable, especially when h is discrete.

A classic example that illustrates the phenomenon is to imagine you are on vacation away from home and you receive a phone call from the security system company

telling you that the alarm has been activated. You begin worrying your home has been burglarized, but then you hear on the radio that a minor earthquake has been reported in the area of your home. If you happen to know from prior experience that earthquakes sometimes cause your home alarm system to activate, then suddenly you relax, confident that your home has very likely not been burglarized.

The example illustrates how the *alarm activation* rendered two otherwise entirely independent causes, burglarized and earthquake, to become dependent—in this case, the dependency is one of mutual exclusivity. Since both burglarized and earthquake are very rare events and both can cause alarm activation, the observation of one explains away the other. Despite the computational obstacles we face when attempting to recover the posterior over h, explaining away promises to provide a parsimonious $p(h \mid x)$, which can be an extremely useful characteristic of a feature encoding scheme. If one thinks of a representation as being composed of various feature detectors and estimated attributes of the observed input, it is useful to allow the different features to compete and collaborate with each other to explain the input. This is naturally achieved with directed graphical models, but can also be achieved with undirected models (see Section 6.2) such as Boltzmann machines if there are lateral connections between the corresponding units or corresponding interaction terms in the energy function that defines the probability model.

Probabilistic interpretation of PCA. PCA can be given a natural probabilistic interpretation [171], [206] as *factor analysis*:

$$p(h) = \mathcal{N}(h; 0, \sigma_h^2 \mathbf{I}),$$

$$p(x \mid h) = \mathcal{N}(x; Wh + \mu_x, \sigma_x^2 \mathbf{I}),$$
(1)

where $x \in \mathbb{R}^{d_x}$, $h \in \mathbb{R}^{d_h}$, $\mathcal{N}(v; \mu, \Sigma)$ is the multivariate normal density of v with mean μ and covariance Σ , and columns of W span the same space as leading d_h principal components, but are not constrained to be orthonormal.

Sparse coding. Like PCA, sparse coding has both a probabilistic and nonprobabilistic interpretation. Sparse coding also relates a latent representation h (either a vector of random variables or a feature vector, depending on the interpretation) to the data x through a linear mapping W, which we refer to as the *dictionary*. The difference between sparse coding and PCA is that sparse coding includes a penalty to ensure a *sparse* activation of h is used to encode each input x. From a nonprobabilistic perspective, sparse coding can be seen as recovering the code or feature vector associated with a new input x via

$$h^* = f(x) = \underset{h}{\operatorname{argmin}} \|x - Wh\|_2^2 + \lambda \|h\|_1.$$
 (2)

Learning the dictionary W can be accomplished by optimizing the following training criterion with respect to W:

$$\mathcal{J}_{SC} = \sum_{t} \|x^{(t)} - Wh^{*(t)}\|_{2}^{2}, \tag{3}$$

where $x^{(t)}$ is the tth example and $h^{*(t)}$ is the corresponding sparse code determined by (2). W is usually constrained to

exchange scaling of column i with scaling of $h_i^{(t)}$, such a constraint is necessary for the L1 penalty to have any effect).

The probabilistic interpretation of sparse coding differs from that of PCA in that instead of a Gaussian prior on the latent random variable h, we use a sparsity inducing Laplace prior (corresponding to an L1 penalty):

$$p(h) = \prod_{i}^{d_h} \frac{\lambda}{2} \exp(-\lambda |h_i|),$$

$$p(x \mid h) = \mathcal{N}(x; Wh + \mu_x, \sigma_x^2 \mathbf{I}).$$
(4)

In the case of sparse coding, because we will seek a sparse representation (i.e., one with many features set to exactly zero), we will be interested in recovering the maximum a posteriori value (MAP value of h, i.e., $h^* = \operatorname{argmax}_h p(h \mid$ x) rather than its expected value $\mathbb{E}[h \mid x]$). Under this interpretation, dictionary learning proceeds as maximizing the likelihood of the data given these MAP values of h^* : $\operatorname{argmax}_W \prod_t p(x^{(t)} \mid h^{*(t)})$ subject to the norm constraint on W. Note that this parameter learning scheme, subject to the MAP values of the latent h, is not standard practice in the probabilistic graphical model literature. Typically, the likelihood of the data $p(x) = \sum_h p(x \mid h) p(h)$ is maximized directly. In the presence of latent variables, expectation maximization is employed, where the parameters are optimized with respect to the marginal likelihood, i.e., summing or integrating the joint log likelihood over all values of the latent variables under their posterior $P(h \mid x)$, rather than considering only the single MAP value of h. The theoretical properties of this form of parameter learning are not yet well understood but seem to work well in practice (e.g., k-means versus Gaussian mixture models and Viterbi training for HMMs). Note also that the interpretation of sparse coding as a MAP estimation can be questioned [77] because, even though the interpretation of the L1 penalty as a log prior is a possible interpretation, there can be other Bayesian interpretations compatible with the training criterion.

Sparse coding is an excellent example of the power of explaining away. Even with a very overcomplete dictionary, 11 the MAP inference process used in sparse coding to find h^* can pick out the most appropriate bases and zero the others, despite them having a high degree of correlation with the input. This property arises naturally in directed graphical models such as sparse coding and is entirely due to the explaining away effect. It is not seen in commonly used undirected probabilistic models such as the RBM, nor is it seen in parametric feature encoding methods such as autoencoders. The tradeoff is that compared to methods such as RBMs and autoencoders, inference in sparse coding involves an extra inner loop of optimization to find h^* with a corresponding increase in the computational cost of feature extraction. Compared to autoencoders and RBMs, the code in sparse coding is a free variable for each example, and in that sense, the implicit encoder is nonparametric.

One might expect that the parsimony of the sparse coding representation and its explaining away effect would be advantageous and indeed it seems to be the case. Coates

have unit-norm columns (because one can arbitrarily 11. Overcomplete: with more dimensions of h than dimensions of x. Authorized licensed use limited to: National Library Of Medicine (NLM). Downloaded on June 28,2023 at 18:56:01 UTC from IEEE Xplore. Restrictions apply.

and Ng [48] demonstrated on the CIFAR-10 object classification task [116] with a patch-base feature extraction pipeline that in the regime with few (<1,000) labeled training examples per class, the sparse coding representation significantly outperformed other highly competitive encoding schemes. Possibly because of these properties and because of the very computationally efficient algorithms that have been proposed for it (in comparison with the general case of inference in the presence of explaining away), sparse coding enjoys considerable popularity as a feature learning and encoding paradigm. There are numerous examples of its successful application as a feature representation scheme, including natural image modeling [159], [109], [48], [224], audio classification [78], NLP [4], as well as being a very successful model of the early visual cortex [155]. Sparsity criteria can also be generalized successfully to yield groups of features that prefer to all be zero, but if one or a few of them are active, then the penalty for activating others in the group is small. Different group sparsity patterns can incorporate different forms of prior knowledge [110], [107], [3], [76].

S3C. S3C is one example of a promising variation on sparse coding for feature learning [73]. The S3C model possesses a set of latent binary *spike* variables together with a set of latent real-valued *slab* variables. The activation of the spike variables dictates the sparsity pattern. S3C has been applied to the CIFAR-10 and CIFAR-100 object classification tasks [116], and shows the same pattern as sparse coding of superior performance in the regime of relatively few (< 1,000) labeled examples per class [73]. In fact, in both the CIFAR-100 dataset (with 500 examples per class) and the CIFAR-10 dataset (when the number of examples is reduced to a similar range), the S3C representation actually outperforms sparse coding representations. This advantage was revealed clearly with S3C winning the NIPS '11 Transfer Learning Challenge [72].

6.2 Undirected Graphical Models

Undirected graphical models, also called Markov random fields (MRFs), parameterize the joint p(x,h) through a product of unnormalized nonnegative *clique potentials*:

$$p(x,h) = \frac{1}{Z_{\theta}} \prod_{i} \psi_{i}(x) \prod_{j} \eta_{j}(h) \prod_{k} \nu_{k}(x,h), \tag{5}$$

where $\psi_i(x)$, $\eta_j(h)$, and $\nu_k(x,h)$ are the clique potentials describing the interactions between the visible elements, between the hidden variables, and those interaction between the visible and hidden variables, respectively. The partition function Z_θ ensures that the distribution is normalized. Within the context of unsupervised feature learning, we generally see a particular form of MRF called a Boltzmann distribution with clique potentials constrained to be positive:

$$p(x,h) = \frac{1}{Z_{\theta}} \exp(-\mathcal{E}_{\theta}(x,h)), \tag{6}$$

where $\mathcal{E}_{\theta}(x,h)$ is the energy function and contains the interactions described by the MRF clique potentials and θ are the model parameters that characterize these interactions.

The Boltzmann machine was originally defined as a network of symmetrically coupled binary random variables or units. These stochastic units can be divided into two groups: 1) the *visible* units $x \in \{0,1\}^{d_x}$ that represent the data, and 2) the *hidden* or latent units $h \in \{0,1\}^{d_h}$ that mediate dependencies between the visible units through their mutual interactions. The pattern of interaction is specified through the energy function:

$$\mathcal{E}_{\theta}^{\text{BM}}(x,h) = -\frac{1}{2}x^{T}Ux - \frac{1}{2}h^{T}Vh - x^{T}Wh - b^{T}x - d^{T}h, \quad (7)$$

where $\theta = \{U, V, W, b, d\}$ are the model parameters that, respectively, encode the visible-to-visible interactions, the hidden-to-hidden interactions, the visible-to-hidden interactions, the visible self-connections, and the hidden self-connections (called biases). To avoid overparameterization, the diagonals of U and V are set to zero.

The Boltzmann machine energy function specifies the probability distribution over [x,h], via the Boltzmann distribution, (6), with the partition function Z_{θ} given by

$$Z_{\theta} = \sum_{x_1=0}^{x_1=1} \cdots \sum_{x_{d_x}=0}^{x_{d_x}=1} \sum_{h_1=0}^{h_1=1} \cdots \sum_{h_{d_h}=0}^{h_{d_h}=1} \exp\left(-\mathcal{E}_{\theta}^{BM}(x, h; \theta)\right).$$
 (8)

This joint probability distribution gives rise to the set of conditional distributions of the form:

$$P(h_i \mid x, h_{\setminus i}) = \operatorname{sigmoid}\left(\sum_j W_{ji} x_j + \sum_{i' \neq i} V_{ii'} h_{i'} + d_i\right), \quad (9)$$

$$P(x_j \mid h, x_{\setminus j}) = \operatorname{sigmoid}\left(\sum_i W_{ji} x_j + \sum_{i' \neq j} U_{jj'} x_{j'} + b_j\right). (10)$$

In general, inference in the Boltzmann machine is intractable. For example, computing the conditional probability of h_i given the visibles, $P(h_i \mid x)$, requires marginalizing over the rest of the hiddens, which implies evaluating a sum with 2^{d_h-1} terms:

$$P(h_i \mid x) = \sum_{h_1=0}^{h_1=1} \cdots \sum_{h_{i-1}=0}^{h_{i-1}=1} \sum_{h_{i+1}=0}^{h_{i+1}=1} \cdots \sum_{h_{d_k}=0}^{h_{d_h}=1} P(h \mid x).$$
 (11)

However, with some judicious choices in the pattern of interactions between the visible and hidden units, more tractable subsets of the model family are possible, as we discuss next.

RBMs. The RBM is likely the most popular subclass of Boltzmann machine [190]. It is defined by restricting the interactions in the Boltzmann energy function, in (7), to only those between h and x, i.e., $\mathcal{E}_{\theta}^{\text{RBM}}$ is $\mathcal{E}_{\theta}^{\text{BM}}$ with $U=\mathbf{0}$ and $V=\mathbf{0}$. As such, the RBM can be said to form a *bipartite* graph with the visibles and the hiddens forming two layers of vertices in the graph (and no connection between units of the same layer). With this restriction, the RBM possesses the useful property that the conditional distribution over the hidden units factorizes given the visibles:

$$P(h \mid x) = \prod_{i} P(h_i \mid x),$$

$$P(h_i = 1 \mid x) = \operatorname{sigmoid}\left(\sum_{j} W_{ji} x_j + d_i\right).$$
(12)

Likewise, the conditional distribution over the visible units given the hiddens also factorizes:

$$P(x \mid h) = \prod_{j} P(x_j \mid h),$$

$$P(x_j = 1 \mid h) = \operatorname{sigmoid}\left(\sum_{i} W_{ji} h_i + b_j\right).$$
(13)

This makes inferences readily tractable in RBMs. For example, the RBM feature representation is taken to be the set of posterior marginals $P(h_i \mid x)$ which, given the conditional independence described in (12), are immediately available. Note that this is in stark contrast to the situation with popular directed graphical models for unsupervised feature extraction, where computing the posterior probability is intractable.

Importantly, the tractability of the RBM does not extend to its partition function, which still involves summing an exponential number of terms. It does imply, however, that we can limit the number of terms to $\min\{2^{d_x}, 2^{d_h}\}$. Usually, this is still an unmanageable number of terms, and therefore we must resort to approximate methods to deal with its estimation.

It is difficult to overstate the impact the RBM has had to the fields of unsupervised feature learning and deep learning. It has been used in a truly impressive variety of applications, including fMRI image classification [180], motion and spatial transformations [201], [144], collaborative filtering [178], and natural image modeling [160], [53].

6.3 Generalizations of the RBM to Real-Valued Data

Important progress has been made in the last few years in defining generalizations of the RBM that better capture realvalued data, in particular real-valued image data, by better modeling the conditional covariance of the input pixels. The standard RBM, as discussed above, is defined with both binary visible variables $v \in \{0,1\}$ and binary latent variables $h \in \{0,1\}$. The tractability of inference and learning in the RBM has inspired many authors to extend it, via modifications of its energy function, to model other kinds of data distributions. In particular, there have been multiple attempts to develop RBM-type models of real-valued data, where $x \in \mathbb{R}^{d_x}$. The most straightforward approach to modeling real-valued observations within the RBM framework is the so-called *Gaussian RBM* (GRBM), where the only change in the RBM energy function is to the visible units biases, by adding a bias term that is quadratic in the visible units x. While it probably remains the most popular way to model real-valued data within the RBM framework, Ranzato and Hinton [160] suggest that the GRBM has proven to be a somewhat unsatisfactory model of natural images. The trained features typically do not represent sharp edges that occur at object boundaries and lead to latent representations that are not particularly useful features for classification tasks. Ranzato and Hinton [160] argue that the failure of the GRBM to adequately capture the statistical structure of natural images stems from the exclusive use of the model capacity to capture the conditional mean at the expense of the conditional covariance. Natural images, they argue, are chiefly characterized by the covariance of the pixel values, not by their

absolute values. This point is supported by the common use of preprocessing methods that standardize the global scaling of the pixel values across images in a dataset or across the pixel values within each image.

These kinds of concerns about the ability of the GRBM to model natural image data has led to the development of alternative RBM-based models that each attempt to take on this objective of better modeling nondiagonal conditional covariances. Ranzato and Hinton [160] introduced the mean and covariance RBM (mcRBM). Like the GRBM, the mcRBM is a two-layer Boltzmann machine that explicitly models the visible units as Gaussian distributed quantities. However, unlike the GRBM, the mcRBM uses its hidden layer to independently parametrize both the mean and covariance of the data through two sets of hidden units. The mcRBM is a combination of the covariance RBM (cRBM) [163], which models the conditional covariance, and the GRBM, which captures the conditional mean. While the GRBM has shown considerable potential as the basis of a highly successful phoneme recognition system [54], it seems that due to difficulties in training the mcRBM, the model has been largely superseded by the mPoT model. The mPoT model (mean-product of Student's T-distributions model) [164] is a combination of the GRBM and the product of Student's T-distributions model [216]. It is an energy-based model where the conditional distribution over the visible units conditioned on the hidden variables is a multivariate Gaussian (nondiagonal covariance) and the complementary conditional distribution over the hidden variables given the visibles are a set of independent Gamma distributions.

The PoT model has recently been generalized to the mPoT model [164] to include nonzero Gaussian means by the addition of GRBM-like hidden units, similarly to how the mcRBM generalizes the cRBM. The mPoT model has been used to synthesize large-scale natural images [164] that show large-scale features and shadowing structure. It has been used to model natural textures [113] in a *tiled-convolution* configuration (see Section 11.2).

Another recently introduced RBM-based model with the objective of having the hidden units encode both the mean and covariance information is the spike-and-slab restricted Boltzmann machine (ssRBM) [52], [53]. The ssRBM is defined as having both a real-valued "slab" variable and a binary "spike" variable associated with each unit in the hidden layer. The ssRBM has been demonstrated as a feature learning and extraction scheme in the context of CIFAR-10 object classification [116] from natural images and has performed well in the role [52], [53]. When trained convolutionally (see Section 11.2) on full CIFAR-10 natural images, the model demonstrated the ability to generate natural image samples that seem to capture the broad statistical structure of natural images better than previous parametric generative models, as illustrated with the samples of Fig. 2.

The mcRBM, mPoT, and ssRBM each set out to model real-valued data such that the hidden units encode not only the conditional mean of the data but also its conditional covariance. Other than differences in the training schemes, the most significant difference between these models is how they encode their conditional covariance. While the mcRBM and the mPoT use the activation of the hidden mcRBM and the mPoT use the activation of the hidden mcRBM.



Fig. 2. (Top) Samples from convolutionally trained μ -ssRBM from [53]. (Bottom) Images in the CIFAR-10 training set closest (L2 distance with contrast normalized training images) to *corresponding* model samples on top. The model does not appear to be overfitting particular training examples.

enforce constraints on the covariance of x, the ssRBM uses the hidden unit to pinch the precision matrix along the direction specified by the corresponding weight vector. These two ways of modeling conditional covariance diverge when the dimensionality of the hidden layer is significantly different from that of the input. In the overcomplete setting, sparse activation with the ssRBM parametrization permits variance only in the select directions of the sparsely activated hidden units. This is a property the ssRBM shares with sparse coding models [155], [78]. On the other hand, in the case of the mPoT or mcRBM, an overcomplete set of constraints on the covariance implies that capturing arbitrary covariance along a particular direction of the input requires decreasing potentially all constraints with positive projection in that direction. This perspective would suggest that the mPoT and mcRBM do not appear to be well suited to provide a sparse representation in the overcomplete setting.

6.4 RBM Parameter Estimation

Many of the RBM training methods we discuss here are applicable to more general undirected graphical models, but are particularly practical in the RBM setting. Freund and Haussler [66] proposed a learning algorithm for harmoniums (RBMs) based on projection pursuit. CD [88], [94] has been used most often to train RBMs, and many recent papers use stochastic maximum likelihood (SML) [220], [204].

As discussed in Section 6.1, in training probabilistic models, parameters are typically adapted to maximize the *likelihood of the training data* (or equivalently the log likelihood, or its penalized version, which adds a regularization term). With T training examples, the log likelihood is given by

$$\sum_{t=1}^{T} \log P(x^{(t)}; \theta) = \sum_{t=1}^{T} \log \sum_{h \in \{0,1\}^{d_h}} P(x^{(t)}, h; \theta).$$
 (14)

Gradient-based optimization requires its gradient, which for Boltzmann machines is given by

$$\frac{\partial}{\partial \theta_{i}} \sum_{t=1}^{T} \log p(x^{(t)}) = -\sum_{t=1}^{T} \mathbb{E}_{p(h|x^{(t)})} \left[\frac{\partial}{\partial \theta_{i}} \mathcal{E}_{\theta}^{\text{BM}}(x^{(t)}, h) \right] + \sum_{t=1}^{T} \mathbb{E}_{p(x,h)} \left[\frac{\partial}{\partial \theta_{i}} \mathcal{E}_{\theta}^{\text{BM}}(x, h) \right], \tag{15}$$

where we have the expectations with respect to $p(h^{(t)} \mid x^{(t)})$ under the "clamped" condition (also called the positive phase) and over the full joint p(x,h) under the "unclamped" condition (also called the negative phase). Intuitively, the gradient acts to locally move the model distribution (the negative phase distribution) toward the data distribution (positive phase distribution) by pushing down the energy of $(h,x^{(t)})$ pairs (for $h \sim P(h \mid x^{(t)})$) while pushing up the energy of (h,x) pairs (for $(h,x) \sim P(h,x)$) until the two forces are in equilibrium, at which point the sufficient statistics (gradient of the energy function) have equal expectations with x sampled from the training distribution or with x sampled from the model.

The RBM conditional independence properties imply that the expectation in the positive phase of (15) is tractable. The negative phase term—arising from the partition function's contribution to the log-likelihood gradient—is more problematic because the computation of the expectation over the joint is not tractable. The various ways of dealing with the partition function's contribution to the gradient have brought about a number of different training algorithms, many trying to approximate the log-likelihood gradient.

To approximate the expectation of the joint distribution in the negative phase contribution to the gradient, it is natural to again consider exploiting the conditional independence of the RBM to specify a Monte Carlo approximation of the expectation over the joint:

$$\mathbb{E}_{p(x,h)} \left[\frac{\partial}{\partial \theta_i} \mathcal{E}_{\theta}^{\text{RBM}}(x,h) \right] \approx \frac{1}{L} \sum_{l=1}^{L} \frac{\partial}{\partial \theta_i} \mathcal{E}_{\theta}^{\text{RBM}} \left(\tilde{x}^{(l)}, \tilde{h}^{(l)} \right), \quad (16)$$

with the samples $(\tilde{x}^{(l)}, \tilde{h}^{(l)})$ drawn by a block Gibbs Markov chain Monte Carlo (MCMC) sampling procedure:

$$\tilde{x}^{(l)} \sim P(x \mid \tilde{h}^{(l-1)}),$$

$$\tilde{h}^{(l)} \sim P(h \mid \tilde{x}^{(l)}).$$

Naively, for each gradient update step one would start a Gibbs sampling chain, wait until the chain converges to the equilibrium distribution, and then draw a sufficient number of samples to approximate the expected gradient with respect to the model (joint) distribution in (16). Then, restart the process for the next step of approximate gradient ascent on the log-likelihood. This procedure has the obvious flaw that waiting for the Gibbs chain to "burn-in" and reach equilibrium anew for each gradient update cannot form the basis of a practical training algorithm. CD [88], [94], SML [220], [204], and fast-weights persistent CD or fast-weights

persistent contrastive divergence (FPCD) [205] are all ways to avoid or reduce the need for burn in.

6.4.1 CD

CD estimation [88], [94] estimates the negative phase expectation (15) with a very short Gibbs chain (often just one step) initialized at the training data used in the positive phase. This reduces the variance of the gradient estimator and still moves in a direction that pulls the negative chain samples toward the associated positive chain samples. Much has been written about the properties and alternative interpretations of CD and its similarity to autoencoder training, for example, [42], [225], [14], [197].

6.4.2 SML

The SML algorithm (also known as persistent contrastive divergence or PCD) [220], [204] is an alternative way to sidestep an extended burn in of the negative phase Gibbs sampler. At each gradient update, rather than initializing the Gibbs chain at the positive phase sample as in CD, SML initializes the chain at the last state of the chain used for the previous update. In other words, SML uses a continually running Gibbs chain (or often a number of Gibbs chains run in parallel) from which samples are drawn to estimate the negative phase expectation. Despite the model parameters changing between updates, these changes should be small enough that only a few steps of Gibbs (in practice, often one step is used) are required to maintain samples from the equilibrium distribution of the Gibbs chain, i.e., the model distribution.

A troublesome aspect of SML is that it relies on the Gibbs chain to mix well (especially between modes) for learning to succeed. Typically, as learning progresses and the weights of the RBM grow, the ergodicity of the Gibbs sample begins to break down. If the learning rate ϵ associated with gradient ascent $\theta \leftarrow \theta + \epsilon \hat{g}$ (with $E[\hat{g}] \approx \frac{\partial \log p_{\theta}(x)}{\partial \theta}$) is not reduced to compensate, then the Gibbs sampler will diverge from the model distribution and learning will fail. Desjardins et al. [58], Cho et al. [44], and Salakhutdinov [173], [174] have all considered various forms of tempered transitions to address the failure of Gibbs chain mixing, and convincing solutions have not yet been clearly demonstrated. A recently introduced promising avenue relies on depth itself, relying on the observation that mixing between modes is much easier on deeper layers [27] (Section 9.4).

Tieleman and Hinton [205] have proposed quite a different approach to addressing potential mixing problems of SML with their FPCD, and it has also been exploited to train DBMs [173] and construct a pure sampling algorithm for RBMs [39]. FPCD builds on the surprising but robust tendency of Gibbs chains to mix better during SML learning than when the model parameters are fixed. The phenomenon is rooted in the form of the likelihood gradient itself (15). The samples drawn from the SML Gibbs chain are used in the negative phase of the gradient,

which implies that the learning update will slightly increase the energy (decrease the probability) of those samples, making the region in the neighborhood of those samples less likely to be resampled and therefore making it more likely that the samples will move somewhere else (typically going near another mode). Rather than drawing samples from the distribution of the current model (with parameters θ), FPCD exaggerates this effect by drawing samples from a local perturbation of the model with parameters θ^* and an update:

$$\theta_{t+1}^* = (1 - \eta)\theta_{t+1} + \eta\theta_t^* + \epsilon^* \frac{\partial}{\partial \theta_i} \left(\sum_{t=1}^T \log p(x^{(t)}) \right), \quad (17)$$

where ϵ^* is the relatively large fast-weight learning rate $(\epsilon^* > \epsilon)$ and $0 < \eta < 1$ (but near 1) is a forgetting factor that keeps the perturbed model close to the current model. Unlike tempering, FPCD does not converge to the model distribution as ϵ and ϵ^* go to 0, and further work is necessary to characterize the nature of its approximation to the model distribution. Nevertheless, FPCD is a popular and apparently effective means of drawing approximate samples from the model distribution that faithfully represent its diversity at the price of sometimes generating spurious samples in between two modes (because the fast weights roughly correspond to a smoothed view of the current model's energy function). It has been applied in a variety of applications [205], [165], [113], and it has been transformed into a sampling algorithm [39] that also shares this fast mixing property with herding [215] for the same reason, i.e., introducing negative correlations between consecutive samples of the chain to promote faster mixing.

6.4.3 Pseudolikelihood, Ratio Matching and More

While CD, SML, and FPCD are by far the most popular methods for training RBMs and RBM-based models, all of these methods are perhaps most naturally described as offering different approximations to maximum likelihood training. There exist other inductive principles that are alternatives to maximum likelihood that can also be used to train RBMs. In particular, these include pseudolikelihood [32] and ratio matching [98]. Both of these inductive principles attempt to avoid explicitly dealing with the partition function, and their asymptotic efficiency has been analyzed [140]. Pseudolikelihood seeks to maximize the product of all 1D conditional distributions of the form $P(x_d \mid x_{\setminus d})$, while ratio matching can be interpreted as an extension of score matching [97] to discrete data types. Both methods amount to weighted differences of the gradient of the RBM free energy¹³ evaluated at a data point and at neighboring points. One potential drawback of these methods is that depending on the parameterization of the energy function, their computational requirements may scale up to $O(n_d)$ worse than CD, SML, FPCD, or denoising score matching [112], [210], discussed below. Marlin et al. [141] empirically compared all of these methods (except denoising score matching) on a range of classification,

^{12.} When weights become large, the estimated distribution is more peaky, and the chain takes a very long time to mix, to move from mode to mode, so that practically the gradient estimator can be very poor. This is a serious chicken-and-egg problem because if sampling is not effective, then neither is the training procedure, which may seem to stall and yield even larger weights.

^{13.} The free energy $\mathcal{F}(x;\theta)$ is the energy associated with the data marginal probability, $\mathcal{F}(x;\theta) = -\log P(x) - \log Z_{\theta}$ and is tractable for the RBM.

reconstruction, and density modeling tasks and found that, in general, SML provided the best combination of overall performance and computational tractability. However, in a later study, the same authors [200] found denoising score matching to be a competitive inductive principle both in terms of classification performance (with respect to SML) and in terms of computational efficiency (with respect to analytically obtained score matching). Denoising score matching is a special case of the denoising autoencoder (DAE) training criterion (Section 7.2.2) when the reconstruction error residual equals a gradient, i.e., the score function associated with an energy function, as shown in [210].

In the spirit of the Boltzmann machine gradient (15), several approaches have been proposed to train energybased models. One is noise-contrastive estimation [80], in which the training criterion is transformed into a probabilistic classification problem: distinguish between (positive) training examples and (negative) noise samples generated by a broad distribution (such as the Gaussian). Another family of approaches, more in the spirit of CD, relies on distinguishing positive examples (of the training distribution) and negative examples obtained by perturbations of the positive examples [50], [33], [218].

DIRECTLY LEARNING A PARAMETRIC MAP FROM INPUT TO REPRESENTATION

Within the framework of probabilistic models adopted in Section 6, the learned representation is always associated with latent variables, specifically with their posterior distribution given an observed input x. Unfortunately, this posterior distribution tends to become very complicated and intractable if the model has more than a couple of interconnected layers, whether in the directed or undirected graphical model frameworks. It then becomes necessary to resort to sampling or approximate inference techniques, and to pay the associated computational and approximation error price. If the true posterior has a large number of modes that matter, then current inference techniques may face an unsurmountable challenge or endure a potentially serious approximation. This is in addition to the difficulties raised by the intractable partition function in undirected graphical models. Moreover, a posterior distribution over latent variables is not yet a simple usable feature vector that can, for example, be fed to a classifier. So, actual feature values are typically derived from that distribution, taking the latent variable's expectation (as is typically done with RBMs), their marginal probability, or finding their most likely value (as in sparse coding). If we are to extract stable deterministic numerical feature values in the end anyway, an alternative (apparently) nonprobabilistic feature learning paradigm that focuses on carrying out this part of the computation very efficiently is that of autoencoders and other directly parameterized feature or representation functions. The commonality between these methods is that they learn a direct encoding, i.e., a parametric map from inputs to their representation.

Regularized autoencoders, discussed next, also involve learning a decoding function that maps back from representation to input space. Sections 8.1 and 11.3 discuss direct encoding methods that do not require a decoder, such as semi-supervised embedding [217] and slow feature analysis [219].

7.1 Autoencoders

In the autoencoder framework [129], [37], [93], one starts by explicitly defining a feature-extracting function in a specific parameterized closed form. This function, which we will denote f_{θ} , is called the *encoder* and will allow the straightforward and efficient computation of a feature vector $h = f_{\theta}(x)$ from an input x. For each example $x^{(t)}$ from a dataset $\{x^{(1)}, \dots, x^{(T)}\}\$, we define

$$h^{(t)} = f_{\theta}(x^{(t)}),$$
 (18)

where $h^{(t)}$ is the feature vector or representation or code computed from $x^{(t)}$. Another closed-form parameterized function q_{θ} , called the *decoder*, maps from feature space back into input space, producing a reconstruction $r = g_{\theta}(h)$. Whereas probabilistic models are defined from an explicit probability function and are trained to maximize (often approximately) the data likelihood (or a proxy), autoencoders are parameterized through their encoder and decoder and are trained using a different training principle. The set of parameters θ of the encoder and decoder are learned simultaneously on the task of reconstructing as well as possibly the original input, i.e., attempting to incur the lowest possible reconstruction error L(x,r)—a measure of the discrepancy between x and its reconstruction r—over training examples. Good generalization means low reconstruction error at test examples, while having high reconstruction error for most other x configurations. To capture the structure of the data-generating distribution, it is therefore important that something in the training criterion or the parameterization prevents the autoencoder from learning the identity function, which has zero reconstruction error everywhere. This is achieved through various means in the different forms of autoencoders, as described below in more detail, and we call these regularized autoencoders. A particular form of regularization consists of constraining the code to have a low dimension, and this is what the classical autoencoder or PCA do.

In summary, basic autoencoder training consists in finding a value of parameter vector θ minimizing reconstruction error:

$$\mathcal{J}_{AE}(\theta) = \sum_{t} L(x^{(t)}, g_{\theta}(f_{\theta}(x^{(t)}))), \tag{19}$$

where $x^{(t)}$ is a training example. This minimization is usually carried out by stochastic gradient descent as in the training of MLPs. Since autoencoders were primarily developed as MLPs, predicting their input, the most commonly used forms for the encoder and decoder are affine mappings, optionally followed by a nonlinearity:

$$f_{\theta}(x) = s_f(b + Wx), \tag{20}$$

$$g_{\theta}(h) = s_g(d + W'h), \tag{21}$$

where s_f and s_q are the encoder and decoder activation functions (typically, the elementwise sigmoid or hyperbolic tangent nonlinearity, or the identity function if staying Authorized licensed use limited to: National Library Of Medicine (NLM). Downloaded on June 28,2023 at 18:56:01 UTC from IEEE Xplore. Restrictions apply.

linear). The set of parameters of such a model is $\theta = \{W, b, W', d\}$, where b and d are called encoder and decoder bias vectors and W and W' are the encoder and decoder weight matrices.

The choice of s_q and L depends largely on the input domain range and nature and is usually chosen so that L returns a negative log likelihood for the observed value of x. A natural choice for an unbounded domain is a linear decoder with a squared reconstruction error, i.e., $s_a(a) = a$ and $L(x,r) = ||x-r||^2$. If inputs are bounded between 0 and 1, however, ensuring a similarly bounded reconstruction can be achieved by using $s_g = \text{sigmoid}$. In addition, if the inputs are of a binary nature, a binary cross-entropy loss 14 is sometimes used.

If both encoder and decoder use a sigmoid nonlinearity, then $f_{\theta}(x)$ and $g_{\theta}(h)$ have the exact same form as the conditionals $P(h \mid v)$ and $P(v \mid h)$ of binary RBMs (see Section 6.2). This similarity motivated an initial study [23] of the possibility of replacing RBMs with autoencoders as the basic pretraining strategy for building deep networks, as well as the comparative analysis of autoencoder reconstruction error gradient and CD updates [14].

One notable difference in the parameterization is that RBMs use a single weight matrix, which follows naturally from their energy function, whereas the autoencoder framework allows for a different matrix in the encoder and decoder. In practice, however, weight tying in which one defines $W' = \overline{W}^T$ may be (and is most often) used, rendering the parameterizations identical. The usual training procedures, however, differ greatly between the two approaches. A practical advantage of training autoencoder variants is that they define a simple tractable optimization objective that can be used to monitor progress.

In the case of a linear autoencoder (linear encoder and decoder) with squared reconstruction error, minimizing (19) learns the same *subspace*¹⁵ as PCA. This is also true when using a sigmoid nonlinearity in the encoder [37], but not if the weights W and W' are tied $(W' = W^T)$ because W cannot be forced into being small and W' large to achieve a linear encoder.

Similarly, Le et al. [124] recently showed that adding a regularization term of the form $\sum_t \sum_i s_3(W_j x^{(t)})$ to a linear autoencoder with tied weights, where s_3 is a nonlinear convex function, yields an efficient algorithm for learning linear ICA.

7.2 Regularized Autoencoders

Like PCA, autoencoders were originally seen as a dimensionality reduction technique and thus used a bottleneck, i.e., $d_h < d_x$. On the other hand, successful uses of sparse coding and RBM approaches tend to favor overcomplete representations, i.e., $d_h > d_x$. This can allow the autoencoder to simply duplicate the input in the features with perfect reconstruction without having extracted more meaningful features. Recent research has demonstrated very successful alternative ways, called regularized autoencoders, to "constrain"

the representation, even when it is overcomplete. The effect of a bottleneck or of this regularization is that the autoencoder cannot reconstruct everything well that it is trained to reconstruct well; the training examples and generalization mean that reconstruction error is also small on test examples. An interesting justification [162] for the sparsity penalty (or any penalty that restricts in a soft way the volume of hidden configurations easily accessible by the learner) is that it acts in spirit like the partition function of RBMs by making sure that only few input configurations can have a low reconstruction error.

Alternatively, one can view the objective of the regularization applied to an autoencoder as making the representation as "constant" (insensitive) as possible with respect to changes in input. This view immediately justifies two variants of regularized autoencoders described below: Contractive autoencoders (CAEs) reduce the number of effective degrees of freedom of the representation (around each point) by making the encoder contractive, i.e., making the derivative of the encoder small (thus making the hidden units saturate), while the DAE makes the whole mapping "robust," i.e., insensitive to small random perturbations, or contractive, making sure that the reconstruction cannot stay good when moving in most directions around a training example.

7.2.1 Sparse Autoencoders

The earliest uses of single-layer autoencoders for building deep architectures by stacking them [23] considered the idea of tying the encoder weights and decoder weights to restrict capacity as well as the idea of introducing a form of sparsity regularization [161]. Sparsity in the representation can be achieved by penalizing the hidden unit biases (making these additive offset parameters more negative) [161], [134], [71], [118] or by directly penalizing the output of the hidden unit activations (making them closer to their saturating value at 0) [162], [123], [227]. Penalizing the bias runs the danger that the weights could compensate for the bias, which could hurt numerical optimization. When directly penalizing the hidden unit outputs, several variants can be found in the literature, but a clear comparative analysis is still lacking. Although the L1 penalty (i.e., simply the sum of output elements h_j in the case of sigmoid nonlinearity) would seem the most natural (because of its use in sparse coding), it is used in few papers involving sparse autoencoders. A close cousin of the L1 penalty is the Student-t penalty ($\log(1+h_i^2)$), originally proposed for sparse coding [155]. Several papers penalize the average output h_i (e.g., over a minibatch), and instead of pushing it to 0 encourage it to approach a fixed target, either through a mean-square error penalty or maybe, more sensibly (because h_i behaves like a probability), a Kullback-Liebler divergence with respect to the binomial distribution with probability ρ : $-\rho \log h_j - (1-\rho) \log(1-h_j) + \text{ constant, for }$ example, with $\rho = 0.05$.

7.2.2 DAEs

Vincent et al. [212], [213] proposed altering the training objective in (19) from mere reconstruction to that of denoising an artificially corrupted input, i.e., learning to reconstruct the clean input from a corrupted version. Authorized licensed use limited to: National Library Of Medicine (NLM). Downloaded on June 28,2023 at 18:56:01 UTC from IEEE Xplore. Restrictions apply.

^{14.} $L(x,r) = -\sum_{i=1}^{d_s} x_i \log(r_i) + (1-r_i) \log(1-r_i)$. 15. Contrary to traditional PCA loading factors, but similarly to the parameters learned by probabilistic PCA, the weight vectors learned by a linear autoencoder are not constrained to form an orthonormal basis nor to have a meaningful ordering. They will, however, span the same subspace.

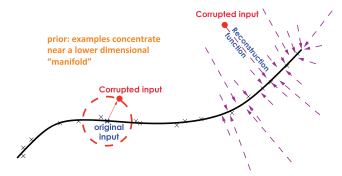


Fig. 3. When data concentrate near a lower dimensional manifold, the corruption vector is typically almost orthogonal to the manifold, and the reconstruction function learns to denoise, map from low-probability configurations (corrupted inputs) to high-probability ones (original inputs), creating a vector field aligned with the score (derivative of the estimated density).

Learning the identity is no longer enough: The learner must capture the structure of the input distribution to optimally undo the effect of the corruption process, with the reconstruction essentially being a nearby but higher density point than the corrupted input. Fig. 3 illustrates that the DAE is learning a reconstruction function that corresponds to a vector field pointing toward high-density regions (the manifold where examples concentrate).

Formally, the objective optimized by a DAE is

$$\mathcal{J}_{\text{DAE}} = \sum_{t} \mathbb{E}_{q(\tilde{x}|x^{(t)})} \left[L(x^{(t)}, g_{\theta}(f_{\theta}(\tilde{x}))) \right], \tag{22}$$

where $\mathbb{E}_{q(\tilde{x}|x^{(t)})}[\cdot]$ averages over corrupted examples \tilde{x} drawn from corruption process $q(\tilde{x} \mid x^{(t)})$. In practice, this is optimized by stochastic gradient descent, where the stochastic gradient is estimated by drawing one or a few corrupted versions of $x^{(t)}$ each time $x^{(t)}$ is considered. Corruptions considered in Vincent et al. [213] include additive isotropic Gaussian noise, salt and pepper noise for gray-scale images, and masking noise (salt or pepper only), for example, setting some randomly chosen inputs to 0 (independently per example). Masking noise has been used in most of the simulations. Qualitatively better features are reported with denoising, resulting in improved classification, and DAE features performed similarly or better than RBM features. Chen et al. [43] show that a simpler alternative with a closed-form solution can be obtained when restricting to a linear autoencoder and have successfully applied it to domain adaptation.

Vincent [210] relates DAEs to energy-based probabilistic models: DAEs basically learn in $r(\tilde{x}) - \tilde{x}$ a vector pointing in the direction of the estimated $score \frac{\partial \log p(\tilde{x})}{\partial \tilde{x}}$ (Fig. 3). In the special case of linear reconstruction and squared error, Vincent [210] shows that training an affine-sigmoid-affine DAE amounts to learning an energy-based model whose energy function is very close to that of a GRBM. Training uses a regularized variant of the *score matching* parameter estimation technique [97], [99], [112] termed *denoising score matching* [210]. Swersky [199] had shown that training GRBMs with *score matching* is equivalent to training a regular autoencoder with an additional regularization term, while follow up on the theoretical results in [210], [200] showed the practical advantage of denoising to implement

score matching efficiently. Finally, Alain and Bengio [1] generalize [210] and prove that DAEs of arbitrary parameterization with small Gaussian corruption noise are general estimators of the score.

7.2.3 CAEs

CAEs, proposed by Rifai et al. [167], follow up on DAEs and share a similar motivation of learning robust representations. CAEs achieve this by adding an analytic *contractive* penalty to (19), the Frobenius norm of the encoder's Jacobian, and results in penalizing the sensitivity of learned features to infinitesimal input variations. Let $J(x) = \frac{\partial f_{\theta}}{\partial x}(x)$ be the Jacobian matrix of the encoder at x. The CAE's training objective is

$$\mathcal{J}_{\text{CAE}} = \sum_{t} L(x^{(t)}, g_{\theta}(f_{\theta}(x^{(t)}))) + \lambda ||J(x^{(t)})||_{F}^{2}, \qquad (23)$$

where λ is a hyperparameter controlling the strength of the regularization. For an affine sigmoid encoder, the contractive penalty term is easy to compute:

$$J_{j}(x) = f_{\theta}(x)_{j} (1 - f_{\theta}(x)_{j}) W_{j},$$

$$||J(x)||_{F}^{2} = \sum_{j} (f_{\theta}(x)_{j} (1 - f_{\theta}(x)_{j}))^{2} ||W_{j}||^{2}.$$
(24)

There are at least three notable differences with DAEs, which may be partly responsible for the better performance that CAE features seem to empirically demonstrate: 1) The sensitivity of the features is penalized16 rather than that of the reconstruction. 2) The penalty is analytic rather than stochastic: An efficiently computable expression replaces what might otherwise require d_x corrupted samples to size up (i.e., the sensitivity in d_x directions). 3) A hyperparameter λ allows fine control of the tradeoff between reconstruction and robustness (while the two are mingled in a DAE). Note, however, that there is a tight connection between the DAE and the CAE: As shown in [1], a DAE with small corruption noise can be seen (through a Taylor expansion) as a type of contractive autoencoder where the contractive penalty is on the whole reconstruction function rather than just on the encoder.17

A potential disadvantage of the CAE's analytic penalty is that it amounts to only encouraging robustness to *infinitesimal* input variations. This is remedied in [168] with the CAE+H, which penalizes all higher order derivatives in an efficient stochastic manner by adding a term that encourages J(x) and $J(x+\epsilon)$ to be close:

$$\mathcal{J}_{\text{CAE+H}} = \sum_{t} L(x^{(t)}, g_{\theta}(x^{(t)})) + \lambda \|J(x^{(t)})\|_{F}^{2}
+ \gamma \mathbb{E}_{\epsilon} [\|J(x) - J(x + \epsilon)\|_{F}^{2}],$$
(25)

where $\epsilon \sim \mathcal{N}(0, \sigma^2 I)$, and γ is the associated regularization strength hyperparameter.

The DAE and CAE have been successfully used to win the final phase of the Unsupervised and Transfer Learning Challenge [145]. The representation learned by the CAE

^{16.} That is, the robustness of the representation is encouraged.

^{17.} But note that in the CAE, the decoder weights are tied to the encoder weights to avoid degenerate solutions, and this should also make the decoder contractive.

tends to be saturated rather than sparse, i.e., most hidden units are near the extremes of their range (e.g., 0 or 1), and their derivative $\frac{\partial h_i(x)}{\partial x}$ is near 0. The nonsaturated units are few and sensitive to the inputs, with their associated filters (weight vectors) together forming a basis explaining the local changes around x, as discussed in Section 8.2. Another way to get saturated (nearly binary) units is semantic hashing [175].

7.2.4 PSD

Sparse coding [155] may be viewed as a kind of autoencoder that uses a linear decoder with a squared reconstruction error, but whose nonparametric *encoder* f_{θ} performs the comparatively nontrivial and relatively costly iterative minimization of (2). A practically successful variant of sparse coding and autoencoders, named PSD [109], replaces that costly and highly nonlinear encoding step by a fast noniterative approximation during recognition (computing the learned features). PSD has been applied to object recognition in images and video [110], [111], [106], but also to audio [84], mostly within the framework of multistage convolutional deep architectures (Section 11.2). The main idea can be summarized by the following equation for the training criterion, which is simultaneously optimized with respect to hidden codes (representation) $h^{(t)}$ and with respect to parameters (W, α) :

$$\mathcal{J}_{PSD} = \sum_{t} \lambda \|h^{(t)}\|_{1} + \|x^{(t)} - Wh^{(t)}\|_{2}^{2} + \|h^{(t)} - f_{\alpha}(x^{(t)})\|_{2}^{2},$$

(26)

where $x^{(t)}$ is the input vector for example t, $h^{(t)}$ is the optimized hidden code for that example, and $f_{\alpha}(\cdot)$ is the encoding function, the simplest variant being

$$f_{\alpha}(x^{(t)}) = \tanh(b + W^T x^{(t)}),$$
 (27)

where encoding weights are the transpose of decoding weights. Many variants have been proposed, including the use of a shrinkage operation instead of the hyperbolic tangent [111]. Note that the L1 penalty on h tends to make them sparse, and how this is the same criterion as sparse coding with dictionary learning (3) except for the additional constraint that one should be able to approximate the sparse codes h with a parameterized encoder $f_{\alpha}(x)$. One can thus view PSD as an approximation to sparse coding, where we obtain a fast approximate encoder. Once PSD is trained, object representations $f_{\alpha}(x)$ are used to feed a classifier. They are computed quickly and can be further fine tuned: The encoder can be viewed as one stage or one layer of a trainable multistage system such as a feedforward neural network.

PSD can also be seen as a kind of autoencoder, where the codes h are given some freedom that can help to further improve reconstruction. One can also view the encoding penalty added on top of sparse coding as a kind of regularizer that forces the sparse codes to be nearly computable by a smooth and efficient encoder. This is in contrast with the codes obtained by complete optimization of the sparse coding criterion, which are highly nonsmooth or even nondifferentiable, a problem that motivated other approaches to smooth the inferred codes of sparse coding

[4], so a sparse coding stage could be jointly optimized along with following stages of a deep architecture.

8 REPRESENTATION LEARNING AS MANIFOLD LEARNING

Another important perspective on representation learning is based on the geometric notion of manifold. Its premise is the manifold hypothesis, according to which real-world data presented in high-dimensional spaces are expected to concentrate in the vicinity of a manifold \mathcal{M} of much lower dimensionality $d_{\mathcal{M}}$, embedded in high-dimensional input space \mathbb{R}^{d_x} . This prior seems particularly well suited for AI tasks such as those involving images, sounds, or text, for which most uniformly sampled input configurations are unlike natural stimuli. As soon as there is a notion of "representation," one can think of a manifold by considering the variations in input space which are captured by or reflected (by corresponding changes) in the learned representation. To first approximation, some directions are well preserved (the tangent directions of the manifold), while others are not (directions orthogonal to the manifolds). With this perspective, the primary unsupervised learning task is then seen as modeling the structure of the data-supporting manifold.¹⁸ The associated representation being learned can be associated with an intrinsic coordinate system on the embedded manifold. The archetypal manifold modeling algorithm is, not surprisingly, also the archetypal lowdimensional representation learning algorithm: PCA, which models a linear manifold. It was initially devised with the objective of finding the closest linear manifold to a cloud of data points. The principal components, i.e., the representation $f_{\theta}(x)$ that PCA yields for an input point x, uniquely locates its projection on that manifold: It corresponds to intrinsic coordinates on the manifold. Data manifolds for complex real-world domains are, however, expected to be strongly nonlinear. Their modeling is sometimes approached as patchworks of locally linear tangent spaces [211], [38]. The large majority of algorithms built on this geometric perspective adopt a nonparametric approach, based on a training set nearest neighbor graph [181], [172], [203], [38], [7], [62], [214], [91], [209]. In these nonparametric approaches, each high-dimensional training point has its own set of free low-dimensional embedding coordinates, which are optimized so that certain properties of the neighborhood graph computed in original high-dimensional input space are best preserved. These methods, however, do not directly learn a parameterized feature extraction function $f_{\theta}(x)$ applicable to new test points, ¹⁹ which seriously limits their use as feature extractors, except in a transductive setting. Comparatively, few nonlinear manifold learning methods have been proposed that learn a parametric map that can directly compute a representation for new points; we will focus on these.

^{18.} Actually, data points need not strictly lie on the "manifold," but the probability density is expected to fall off sharply as one moves away from it, and it may actually be constituted of several possibly disconnected manifolds with different intrinsic dimensionality.

^{19.} For several of these techniques, representations for new points can be computed using the Nyström approximation as has been proposed as an extension in [20], but this remains cumbersome and computationally expensive.

8.1 Learning a Parametric Mapping Based on a Neighborhood Graph

Some of the above nonparametric manifold learning algorithms can be modified to learn a *parametric mapping* f_{θ} , i.e., applicable to new points: Instead of having *free* low-dimensional embedding coordinate "parameters" for each training point, these coordinates are obtained through an explicitly parameterized function, as with the parametric variant [208] of t-SNE [209].

Instead, semi-supervised embedding [217] learns a direct encoding while taking into account the manifold hypothesis through a neighborhood graph. A parameterized neural network architecture simultaneously learns a manifold embedding and a classifier. The training criterion encourages training set neighbors to have similar representations.

The reduced and tightly controlled number of free parameters in such parametric methods, compared to their pure nonparametric counterparts, forces models to generalize the manifold shape nonlocally [22], which can translate into better features and final performance [209]. However, basing the modeling of manifolds on training set neighborhood relationships might be risky statistically in high-dimensional spaces (sparsely populated due to the curse of dimensionality) as, for example, most euclidean nearest neighbors risk having too little in common semantically. The nearest neighbor graph is simply not sufficiently densely populated to map out satisfyingly the wrinkles of the target manifold [17], [22], [16]. It can also become problematic computationally to consider all pairs of data points, ²⁰ which scales quadratically with training set size.

8.2 Learning to Represent Nonlinear Manifolds

Can we learn a manifold without requiring nearest neighbor searches? Yes, for example, with regularized autoencoders or PCA. In PCA, the sensitivity of the extracted components (the code) to input changes is the same regardless of position x. The tangent space is the same everywhere along the linear manifold. By contrast, for a nonlinear manifold, the tangent of the manifold changes as we move on the manifold, as illustrated in Fig. 6. In nonlinear representation-learning algorithms, it is convenient to think about the local variations in the representation as the input x is varied on the manifold, i.e., as we move among high-probability configurations. As we discuss below, the first derivative of the encoder therefore specifies the shape of the manifold (its tangent plane) around an example x lying on it. If the density was really concentrated on the manifold and the encoder had captured that, we would find the encoder derivatives to be nonzero only in the directions spanned by the tangent plane.

Let us consider sparse coding in this light: Parameter matrix W may be interpreted as a dictionary of input directions from which a *different subset* will be picked to model the local tangent space at an x on the manifold. That subset corresponds to the active, i.e., nonzero, features for input x. Nonzero component h_i will be sensitive to small changes of the input in the direction of the associated weight vector $W_{::i}$, whereas inactive features are more likely

to be stuck at 0 until a significant displacement has taken place in input space.

The *local coordinate coding* (LCC) algorithm [223] is very similar to sparse coding, but is explicitly derived from a manifold perspective. Using the same notation as that of sparse coding in (2), LCC replaces regularization term $\|h^{(t)}\|_1 = \sum_j |h_j^{(t)}|$ yielding objective

$$\mathcal{J}_{LCC} = \sum_{t} \left(\|x^{(t)} - Wh^{(t)}\|_{2}^{2} + \lambda \sum_{j} |h_{j}^{(t)}| \|W_{:,j} - x^{(t)}\|^{1+p} \right).$$
(28)

This is identical to sparse coding when p=-1, but with larger p it encourages the active anchor points for $x^{(t)}$ (i.e., the codebook vectors $W_{:,j}$ with nonnegligible $|h_j^{(t)}|$ that are combined to reconstruct $x^{(t)}$) to be not too far from $x^{(t)}$; hence the local aspect of the algorithm. An important theoretical contribution of Yu et al. [223] is to show that that any Lipschitz-smooth function $\phi:\mathcal{M}\to\mathbb{R}$ defined on a smooth nonlinear manifold \mathcal{M} embedded in \mathbb{R}^{d_x} can be well approximated by a globally linear function with respect to the resulting coding scheme (i.e., linear in h), where the accuracy of the approximation and required number d_h of anchor points depend on $d_{\mathcal{M}}$ rather than d_x . This result has been further extended with the use of local tangent directions [222], as well as to multiple layers [137].

Let us now consider the efficient noniterative "feedforward" encoders f_{θ} , used by PSD and the autoencoders reviewed in Section 7.2, which are in the form of (20) or (27). The computed representation for x will only be significantly sensitive to input space directions associated with non-saturated hidden units (see, e.g., (24) for the Jacobian of a sigmoid layer). These directions to which the representation is significantly sensitive, like in the case of PCA or sparse coding, may be viewed as spanning the tangent space of the manifold at point x.

Rifai et al. [167] empirically analyze in this light the singular value spectrum of the Jacobian (derivatives of representation vector with respect to input vector) of a trained CAE. Here, the SVD provides an ordered orthonormal basis of most sensitive directions. The spectrum is sharply decreasing, indicating a relatively small number of significantly sensitive directions. This is taken as empirical evidence that the CAE indeed modeled the tangent space of a low-dimensional manifold. The leading singular vectors form a basis for the tangent plane of the estimated manifold, as illustrated in Fig. 4. The CAE criterion is believed to achieve this thanks to its two opposing terms: The isotropic contractive penalty that encourages the representation to be equally insensitive to changes in any input directions, and the reconstruction term that pushes different training points (in particular neighbors) to have a different representation (so they may be reconstructed accurately), thus counteracting the isotropic contractive pressure only in directions tangent to the manifold.

Analyzing learned representations through the lens of the spectrum of the Jacobian and relating it to the notion of tangent space of a manifold is feasible whenever the mapping is differentiable and regardless of how it was learned, whether as direct encoding (as in autoencoder variants) or derived from latent variable inference (as in

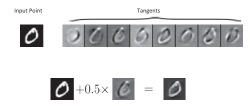


Fig. 4. The tangent vectors to the high-density manifold as estimated by a contractive autoencoder [167]. The original input is shown on the top left. Each tangent vector (images on right side of first row) corresponds to a plausible additive deformation of the original input, as illustrated on the second row, where a bit of the third singular vector is added to the original, to form a translated and deformed image. Unlike in PCA, the tangent vectors are different for different inputs because the estimated manifold is highly nonlinear.

sparse coding or RBMs). Exact low-dimensional manifold models (like PCA) would yield nonzero singular values associated to directions along the manifold, and exact zeros for directions orthogonal to the manifold. But in smooth models like the CAE or the RBM, we will instead have large versus relatively small singular values (as opposed to nonzero versus exactly zero).

8.3 Leveraging the Modeled Tangent Spaces

The local tangent space, at a point along the manifold, can be thought of as capturing locally valid transformations that were prominent in the training data. For example, Rifai et al. [169] examine the tangent directions extracted with an SVD of the Jacobian of CAEs trained on digits, images, or text-document data: They appear to correspond to small translations or rotations for images or digits and to substitutions of words within the same theme for documents. Such very local transformations along a data manifold are not expected to change class identity. To build their MTC, Rifai et al. [169] then apply techniques such as tangent distance [189] and tangent propagation [188], which were initially developed to build classifiers that are insensitive to input deformations provided as prior domain knowledge. Now, these techniques are applied using the local leading tangent directions extracted by a CAE, i.e., not using any prior domain knowledge (except the broad prior about the existence of a manifold). This approach set a new record for MNIST digit classification among prior-knowledge free approaches.2

9 CONNECTIONS BETWEEN PROBABILISTIC AND DIRECT ENCODING MODELS

The standard likelihood framework for probabilistic models decomposes the training criterion for models with parameters θ in two parts: the log-likelihood $\log P(x \mid \theta)$ (or $\log P(x \mid h, \theta)$ with latent variables h) and the prior $\log P(\theta)$ (or $\log P(h \mid \theta) + \log P(\theta)$ with latent variables).

9.1 PSD: A Probabilistic Interpretation

In the case of the PSD algorithm, a connection can be made between the above standard probabilistic view and the direct encoding computation graph. The probabilistic model of PSD is the same directed generative model $P(x \mid h)$ of sparse coding (Section 6.1.1), which only accounts for the decoder. The encoder is viewed as an approximate inference mechanism to guess $P(h \mid x)$ and initialize a MAP iterative inference (where the sparse prior P(h) is taken into account). However, in PSD, the encoder is trained jointly with the decoder, rather than simply taking the end result of iterative inference as a target to approximate. An interesting view²² to reconcile these facts is that the encoder is a parametric approximation for the MAP solution of a variational lower bound on the joint log likelihood. When MAP learning is viewed as a special case of variational learning (where the approximation of the joint log likelihood is with a dirac distribution located at the MAP solution), the variational recipe tells us to simultaneously improve the likelihood (reduce reconstruction error) and improve the variational approximation (reduce the discrepancy between the encoder output and the latent variable value). Hence, PSD sits at the intersection of probabilistic models (with latent variables) and direct encoding methods (which directly parameterize the mapping from input to representation). RBMs also sit at the intersection because their particular parameterization includes an explicit mapping from input to representation, thanks to the restricted connectivity between hidden units. However, this nice property does not extend to their natural deep generalizations, i.e., DBMs, discussed in Section 10.2.

9.2 Regularized Autoencoders Capture Local Structure of the Density

Can we also say something about the probabilistic interpretation of regularized autoencoders? Their training criterion does not fit the standard likelihood framework because this would involve a data-dependent "prior." An interesting hypothesis emerges to answer that question, out of recent theoretical results [210], [1]: The training criterion of regularized autoencoders, instead of being a form of maximum likelihood, corresponds to a different inductive principle, such as score matching. The score matching connection is discussed in Section 7.2.2 and has been shown for a particular parametrization of DAE and equivalent GRBM [210]. The work in [1] generalizes this idea to a broader class of parameterizations (arbitrary encoders and decoders) and shows that by regularizing the autoencoder so that it be contractive, one obtains that the reconstruction function and its derivative estimate first and second derivatives of the underlying data-generative density. This view can be exploited to successfully sample from autoencoders, as shown in [170], [26]. The proposed sampling algorithms are MCMCs similar to Langevin MCMC, using not just the estimated first derivative of the density, but also the estimated manifold tangents so as to stay close to manifolds of high density.

This interpretation connects well with the geometric perspective introduced in Section 8. The regularization effects (e.g., due to a sparsity regularizer, a contractive regularizer, or the denoising criterion) ask the learned representation to be as insensitive as possible to the input, while minimizing reconstruction error on the training examples forces the representation to contain just enough

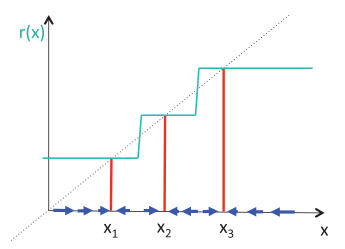


Fig. 5. Reconstruction function r(x) (green) learned by a high-capacity autoencoder on 1D input, minimizing reconstruction error at training examples $x^{(t)}$ ($r(x^{(t)})$ in red) while trying to be as constant as possible otherwise. The dotted line is the identity reconstruction (which might be obtained without the regularizer). The blue arrows show the vector field of r(x)-x pointing toward high-density peaks estimated by the model and estimating the score (log-density derivative).

information to distinguish them. The solution is that variations along the high-density manifolds are preserved, while other variations are compressed: The reconstruction function should be as constant as possible while reproducing training examples, i.e., points near a training example should be mapped to that training example (Fig. 5). The reconstruction function should map an input toward the nearest point manifold, i.e., the difference between reconstruction and input is a vector aligned with the estimated score (the derivative of the log density with respect to the input). The score can be zero on the manifold (where reconstruction error is also zero) at local maxima of the log density, but it can also be zero at local minima. It means that we cannot equate low reconstruction error with high estimated probability. The second derivatives of the log density correspond to the first derivatives of the reconstruction function, and on the manifold (where the first derivative is 0) they indicate the tangent directions of the manifold (where the first derivative remains near 0).

As illustrated in Fig. 6, the basic idea of the autoencoder sampling algorithms in [170], [26] is to make MCMC moves where one 1) moves toward the manifold by following the density gradient (i.e., applying a reconstruction) and 2) adds noise in the directions of the

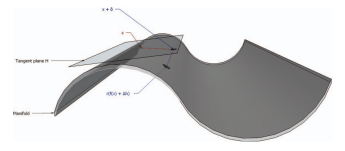


Fig. 6. Sampling from regularized autoencoders [170], [26]: Each MCMC step adds to current state x the noise δ , mostly in the direction of the estimated manifold tangent plane H and projects back toward the manifold (high-density regions) by performing a reconstruction step.

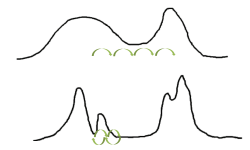


Fig. 7. Top: Early, during training, MCMC mixes easily between modes because the estimated distribution has high entropy and puts enough mass everywhere for small-steps movements (MCMC) to go from mode to mode. Bottom: Later on, training relying on good mixing can stall because estimated modes are separated by wide low-density deserts.

leading singular vectors of the reconstruction (or encoder) Jacobian, corresponding to those associated with smallest second derivative of the log density.

9.3 Learning Approximate Inference

Let us now consider from closer how a representation is computed in probabilistic models with latent variables, when iterative inference is required. There is a computation graph (possibly with random number generation in some of the nodes, in the case of MCMC) that maps inputs to representation and, in the case of deterministic inference (e.g., MAP inference or variational inference), that function could be optimized directly. This is a way to generalize PSD that has been explored in recent work on probabilistic models at the intersection of inference and learning [4], [75], [79], [177], [195], [63], where a central idea is that instead of using a *generic* inference mechanism, one can use one that is *learned* and is more efficient, taking advantage of the specifics of the type of data on which it is applied.

9.4 Sampling Challenges

A troubling challenge with many probabilistic models with latent variables like most Boltzmann machine variants is that good MCMC sampling is required as part of the learning procedure, but that sampling becomes extremely inefficient (or unreliable) as training progresses because the modes of the learned distribution become sharper, making mixing between modes very slow. Whereas initially during training a learner assigns mass almost uniformly, as training progresse, its entropy decreases, approaching the entropy of the target distribution as more examples and more computation are provided. According to our manifold and natural clustering priors of Section 3.1, the target distribution has sharp modes (manifolds) separated by extremely low density areas. Mixing then becomes more difficult because MCMC methods, by their very nature, tend to make small steps to nearby high-probability configurations. This is illustrated in Fig. 7.

Bengio et al. [27] suggest that deep representations could help mixing between such well-separated modes, based on both theoretical arguments and on empirical evidence. The idea is that if higher level representations disentangle the underlying abstract factors better, then small steps in this abstract space (e.g., swapping from one category to another) can easily be done by MCMC. The high-level representations can then be mapped back to

the input space to obtain input-level samples, as in the DBNs sampling algorithm [94]. This has been demonstrated both with DBNs and with the newly proposed algorithm for sampling from contracting and DAEs [170], [26]. This observation alone does not suffice to solve the problem of training a DBN or a DBM, but it may provide a crucial ingredient, and it makes it possible to consider successfully sampling from deep models trained by procedures that do not require an MCMC, like the stacked regularized autoencoders used in [170].

9.5 Evaluating and Monitoring Performance

It is always possible to evaluate a feature learning algorithm in terms of its usefulness with respect to a particular task (e.g., object classification), with a predictor that is fed or initialized with the learned features. In practice, we do this by saving the features learned (e.g., at regular intervals during training, to perform early stopping) and training a cheap classifier on top (such as a linear classifier). However, training the final classifier can be a substantial computational overhead (e.g., supervised fine tuning a deep neural network usually takes more training iterations than the feature learning itself), so we may want to avoid having to train a classifier for every training iteration of the unsupervised learner and every hyperparameter setting. More importantly, this may give an incomplete evaluation of the features (what would happen for other tasks?). All these issues motivate the use of methods to monitor and evaluate purely unsupervised performance. This is rather easy with all the autoencoder variants (with some caution outlined below) and rather difficult with the undirected graphical models such as the RBM and Boltzmann machines.

For autoencoder and sparse coding variants, test set reconstruction error can be readily computed, but by itself may be misleading because *larger capacity* (e.g., more features, more training time) tends to systematically lead to lower reconstruction error, even on the test set. Hence, it cannot be used reliably for selecting most hyperparameters. On the other hand, *denoising* reconstruction error is clearly immune to this problem, so that solves the problem for DAEs. Based on the connection between DAEs and CAEs uncovered in [26], [1], this immunity can be extended to DAEs, but not to the hyperparameter controlling the amount of noise or of contraction.

For RBMs and some (not too deep) Boltzmann machines, one option is the use of annealed importance sampling [150] to estimate the partition function (and thus the test log likelihood). Note that this estimator can have high variance and that it becomes less reliable (variance becomes too large) as the model becomes more interesting, with larger weights, more nonlinearity, sharper modes, and a sharper probability density function (see our previous discussion in Section 9.4). Another interesting and recently proposed option for RBMs is to track the partition function during training [59], which could be useful for early stopping and reducing the cost of ordinary AIS. For toy RBMs (e.g., 25 hidden units or less, or 25 inputs or less), the exact log likelihood can also be computed analytically, and this can be a good way to debug and verify some properties of interest.

10 GLOBAL TRAINING OF DEEP MODELS

One of the most interesting challenges raised by deep architectures is: *How should we jointly train all the levels*? In the previous section and in Section 4, we have only discussed how single-layer models could be combined to form a deep model. Here, we consider joint training of all the levels and the difficulties that may arise.

10.1 The Challenge of Training Deep Architectures

Higher level abstraction means more nonlinearity. It means that two nearby input configurations may be interpreted very differently because a few surface details change the underlying semantics, whereas most other changes in the surface details would not change the underlying semantics. The representations associated with input manifolds may be complex because the mapping from input to representation may have to unfold and distort input manifolds that generally have complicated shapes into spaces where distributions are much simpler, where relations between factors are simpler, maybe even linear or involving many (conditional) independencies. Our expectation is that modeling the joint distribution between high-level abstractions and concepts should be much easier in the sense of requiring much less data to learn. The hard part is learning a good representation that does this unfolding and disentangling. This may be at the price of a more difficult training problem, possibly involving ill conditioning and local minima.

It is only since 2006 that researchers have seriously investigated ways to train deep architectures, to the exception of the convolutional networks [133]. The first realization (Section 4) was that unsupervised or supervised layerwise training was easier and that this could be taken advantage of by stacking single-layer models into deeper ones.

It is interesting to ask why does the layerwise unsupervised pretraining procedure sometimes help a supervised learner [65]. There seems to be a more general principle at play²³ of guiding the training of intermediate representations, which may be easier than trying to learn it all in one go. This is nicely related to the curriculum learning idea [24] that it may be much easier to learn simpler concepts first and then build higher level ones on top of simpler ones. This is also coherent with the success of several deep learning algorithms that provide some such guidance for intermediate representations, like semi-supervised embedding [217].

The question of why unsupervised pretraining could be helpful was extensively studied [65], trying to dissect the answer into a regularization effect and an optimization effect. The regularization effect is clear from the experiments, where the stacked RBMs or DAEs are used to initialize a supervised classification neural network [65]. It may simply come from the use of unsupervised learning to bias the learning dynamics and initialize it in the basin of attraction of a "good" local minimum (of the training criterion), where "good" is in terms of generalization error. The underlying hypothesis exploited by this procedure is that some of the features or latent factors that are good at capturing the leading variations in the input distribution are also good at

capturing the variations in the target output random variables of interest (e.g., classes). The optimization effect is more difficult to tease out because the top two layers of a deep neural net can just overfit the training set whether the lower layers compute useful features or not, but there are several indications that optimizing the lower levels with respect to a supervised training criterion can be challenging.

One such indication is that changing the numerical conditions of the optimization procedure can have a profound impact on the joint training of a deep architecture, for example, by changing the initialization range and changing the type of nonlinearity used [68], much more so than with shallow architectures. One hypothesis to explain some of the difficulty in the optimization of deep architectures is centered on the singular values of the Jacobian matrix associated with the transformation from the features at one level into the features at the next level [68]. If these singular values are all small (less than 1), then the mapping is contractive in every direction and gradients would vanish when propagated backward through many layers. This is a problem already discussed for recurrent neural networks [18], which can be seen as very deep networks with shared parameters at each layer, when unfolded in time. This optimization difficulty has motivated the exploration of second-order methods for deep architectures and recurrent networks, in particular Hessianfree second-order methods [142], [143]. Unsupervised pretraining has also been proposed to help training recurrent networks and temporal RBMs [198], i.e., at each time step, there is a local signal to guide the discovery of good features to capture in the state variables: model with the current state (as hidden units) the joint distribution of the previous state and the current input. Natural gradient [2] methods that can be applied to networks with millions of parameters (i.e., with good scaling properties) have also been proposed [127], [157]. Cho et al. [45] propose using adaptive learning rates for RBM training, along with a novel and interesting idea for a gradient estimator that takes into account the invariance of the model to flipping hidden unit bits and inverting signs of corresponding weight vectors. At least one study indicates that the choice of initialization (to make the Jacobian of each layer closer to 1 across all its singular values) could substantially reduce the training difficulty of deep networks [68] and this is coherent with the success of the initialization procedure of echo state networks [104], as recently studied by Sutskever [196]. There are also several experimental results [68], [69], [151] showing that the choice of hidden units nonlinearity could influence both training and generalization performance, with particularly interesting results obtained with sparse rectifying units [106], [151], [69], [117]. An old idea regarding the ill-conditioning issue with neural networks is that of symmetry breaking: Part of the slowness of convergence may be due to many units moving together (like sheep) and all trying to reduce the output error for the same examples. By initializing with sparse weights [142] or by using often saturated nonlinearities (such as rectifiers as max-pooling units), gradients only flow along a few paths, which may help hidden units to specialize more quickly. Another promising idea to improve the conditioning of neural network training is to nullify the average value and slope of each hidden unit output [158], and possibly locally normalize magnitude as well [106]. The debate still rages between using online methods such as stochastic gradient descent and using second-order methods on large minibatches (of several thousand examples) [142], [123], with a variant of stochastic gradient descent recently winning an optimization challenge.²⁴

Finally, several recent results exploiting large quantities of labeled data suggest that with proper initialization and choice of nonlinearity, very deep purely supervised networks can be trained successfully without any layerwise pretraining [47], [69], [183], [117]. Researchers report that in such conditions, layerwise unsupervised pretraining brought little or no improvement over pure supervised learning from scratch when training for long enough. This reinforces the hypothesis that unsupervised pretraining acts as a prior, which may be less necessary when very large quantities of labeled data are available, but which begs the question of why this had not been discovered earlier. The latest results reported in this respect [117] are particularly interesting because they allowed drastically reducing the error rate of object recognition on a benchmark (the 1,000-class ImageNet task), where many more traditional computer vision approaches had been evaluated (http://www. image-net.org/challenges/LSVRC/2012/ results.html). The main techniques that allowed this success include the following: efficient GPU training allowing one to train longer (more than 100 million visits of examples), an aspect first reported by Lee et al. [135] and Ciresan et al. [47], a large number of labeled examples, artificially transformed examples (see Section 11.1), a large number of tasks (1,000 or 10,000 classes for ImageNet), convolutional architecture with max-pooling (see Section 11 for these latter two techniques), rectifying nonlinearities (discussed above), careful initialization (discussed above), careful parameter update and adaptive learning rate heuristics, layerwise feature normalization (across features), and a new *dropout* trick based on injecting strong binary multiplicative noise on hidden units. This trick is similar to the binary noise injection used at each layer of a stack of DAEs. Future work is hopefully going to help identify which of these elements matter most, how to generalize them across a large variety of tasks and architectures, and, in particular, contexts where most examples are unlabeled, i.e., including an unsupervised component in the training criterion.

10.2 Joint Training of DBMs

We now consider the problem of joint training of all layers of a specific unsupervised model, the DBM. Whereas much progress (albeit with many unanswered questions) has been made on jointly training all the layers of deep architectures using back-propagated gradients (i.e., mostly in the supervised setting), much less work has been done on their purely unsupervised counterpart, for example, with DBMs.²⁵ Note, however, that one could hope that the

 $^{24.\} https://sites.google.com/site/nips2011workshop/optimization-challenges.$

other promising idea to improve the conditioning of challenging because of the much harder inference problem involved.

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successful techniques described in the previous section could be applied to unsupervised learning algorithms.

Like the RBM, the DBM is another particular subset of the Boltzmann machine family of models where the units are again arranged in layers. However, unlike the RBM, the DBM possesses multiple layers of hidden units, with units in odd-numbered layers being conditionally independent given even-numbered layers, and vice versa. With respect to the Boltzmann energy function of (7), the DBM corresponds to setting U=0 and a sparse connectivity structure in both V and W. We can make the structure of the DBM more explicit by specifying its energy function. For the model with two hidden layers, it is given as

$$\mathcal{E}_{\theta}^{\text{DBM}}(v, h^{(1)}, h^{(2)}; \theta) = -v^{T}Wh^{(1)} - h^{(1)}{}^{T}Vh^{(2)} - d^{(1)}{}^{T}h^{(1)} - d^{(2)}{}^{T}h^{(2)} - b^{T}v,$$
(29)

with $\theta = \{W, V, d^{(1)}, d^{(2)}, b\}$. The DBM can also be characterized as a bipartite graph between two sets of vertices, formed by odd- and even-numbered layers (with $v := h^{(0)}$).

10.2.1 Mean-Field Approximate Inference

A key point of departure from the RBM is that the posterior distribution over the hidden units (given the visibles) is no longer tractable due to the interactions between the hidden units. Salakhutdinov and Hinton [176] resort to a meanfield approximation to the posterior. Specifically, in the case of a model with two hidden layers, we wish to approximate $P(h^{(1)}, h^{(2)} \mid v)$ with the factored distribution $Q_v(h^{(1)}, h^{(2)}) = \prod_{j=1}^{N_1} Q_v(h_j^{(1)}) \prod_{i=1}^{N_2} Q_v(h_i^{(2)})$ such that the KL divergence $\mathrm{KL}(P(h^{(1)}, h^{(2)} \mid v) \| Q_v(h^1, h^2))$ is minimized or, equivalently, that a lower bound to the log likelihood is maximized:

$$\log P(v) > \mathcal{L}(Q_v) \equiv \sum_{h^{(1)}} \sum_{h^{(2)}} Q_v(h^{(1)}, h^{(2)})$$

$$\log \left(\frac{P(v, h^{(1)}, h^{(2)})}{Q_v(h^{(1)}, h^{(2)})} \right).$$
(30)

Maximizing this lower bound with respect to the meanfield distribution $Q_v(h^1, h^2)$ (by setting derivatives to zero) yields the following mean field update equations:

$$\hat{h}_i^{(1)} \leftarrow \operatorname{sigmoid}\left(\sum_j W_{ji} v_j + \sum_k V_{ik} \hat{h}_k^{(2)} + d_i^{(1)}\right)$$
(31)

$$\hat{h}_k^{(2)} \leftarrow \operatorname{sigmoid}\left(\sum_i V_{ik} \hat{h}_i^{(1)} + d_k^{(2)}\right). \tag{32}$$

Note how the above equations ostensibly look like a *fixed* point recurrent neural network, i.e., with constant input. In the same way that an RBM can be associated with a simple autoencoder, the above mean-field update equations for the DBM can be associated with a recurrent autoencoder. In that case, the training criterion involves the reconstruction error at the last or at consecutive time steps. This type of model has been explored by Savard [179] and Seung [186] and shown to do a better job at denoising than ordinary autoencoders.

Iterating (31-32) until convergence yields the Q parameters of the "variational positive phase" of

$$\mathcal{L}(Q_{v}) = \mathbb{E}_{Q_{v}} \left[\log P(v, h^{(1)}, h^{(2)}) - \log Q_{v}(h^{(1)}, h^{(2)}) \right]$$

$$= \mathbb{E}_{Q_{v}} \left[-\mathcal{E}_{\theta}^{\text{DBM}}(v, h^{(1)}, h^{(2)}) - \log Q_{v}(h^{(1)}, h^{(2)}) \right]$$

$$- \log Z_{\theta},$$

$$\frac{\partial \mathcal{L}(Q_{v})}{\partial \theta} = -\mathbb{E}_{Q_{v}} \left[\frac{\partial \mathcal{E}_{\theta}^{\text{DBM}}(v, h^{(1)}, h^{(2)})}{\partial \theta} \right]$$

$$+ \mathbb{E}_{P} \left[\frac{\partial \mathcal{E}_{\theta}^{\text{DBM}}(v, h^{(1)}, h^{(2)})}{\partial \theta} \right].$$
(33)

This variational learning procedure leaves the "negative phase" untouched, which can thus be estimated through SML or CD [89] as in the RBM case.

10.2.2 Training DBMs

The major difference between training a DBM and an RBM is that instead of maximizing the likelihood directly, we instead choose parameters to maximize the lower bound on the likelihood given in (30). The SML-based algorithm for maximizing this lower bound is as follows:

- 1. Clamp the visible units to a training example.
- 2. Iterate over (31)-(32) until convergence.
- 3. Generate negative phase samples v^- , $h^{(1)-}$, and $h^{(2)-}$ through SML.
- 4. Compute $\partial \mathcal{L}(Q_v)/\partial \theta$ using the values obtained in steps 2 and 3.
- Finally, update the model parameters with a step of approximate stochastic gradient ascent.

While the above procedure appears to be a simple extension of the highly effective SML scheme for training RBMs, as we demonstrate in Desjardins et al. [60], this procedure seems vulnerable to falling in poor local minima that leave many hidden units effectively dead (not significantly different from its random initialization with small norm).

The failure of the SML joint training strategy was noted by Salakhutdinov and Hinton [176]. As an alternative, they proposed a greedy layerwise training strategy. This procedure consists of pretraining the layers of the DBM, in much the same way as the DBN, i.e., by stacking RBMs and training each layer to independently model the output of the previous layer. A final joint "fine tuning" is done following the above SML-based procedure.

11 BUILDING-IN INVARIANCE

It is well understood that incorporating prior domain knowledge helps machine learning. Exploring good strategies for doing so is a very important research avenue. However, if we are to advance our understanding of core machine learning principles, it is important that we keep comparisons between predictors fair and maintain a clear awareness of the prior domain knowledge used by different learning algorithms, especially when comparing their performance on benchmark problems. We have so far only presented algorithms that exploited only generic inductive biases for high-dimensional problems, thus making them potentially applicable to any high-dimensional problem. The most prevalent approach to incorporating prior knowledge is to hand design better features to feed a generic classifier and this has been used extensively in computer

vision (e.g., [138]). Here, we rather focus on how *basic* domain knowledge of the input, in particular its topological structure (e.g., bitmap images having a 2D structure), may be used to *learn* better features.

11.1 Generating Transformed Examples

Generalization performance is usually improved by providing a larger quantity of representative data. This can be achieved by generating new examples by applying small random deformations to the original training examples, using deformations that are known not to change the target variables of interest, for example, an object class is invariant to small transformations of images such as translations, rotations, scaling, or shearing. This old approach [5] has recently been applied with great success in the work of Ciresan et al. [47], who used an efficient GPU implementation ($40 \times$ speedup) to train a standard but large deep MLP on deformed MNIST digits. Using both affine and elastic deformations [187], with plain old stochastic gradient descent, they reach a record 0.32 percent classification error rate.

11.2 Convolution and Pooling

Another powerful approach is based on even more basic knowledge of merely the topological structure of the input dimensions. By this we mean, for example, the 2D layout of pixels in images or audio spectrograms, the 3D structure of videos, the 1D sequential structure of text, or of temporal sequences in general. Based on such structure, one can define local receptive fields [95] so that each low-level feature will be computed from only a subset of the input: a neighborhood in the topology (e.g., a subimage at a given position). This topological locality constraint corresponds to a layer having a very sparse weight matrix with nonzeros only allowed for topologically local connections. Computing the associated matrix products can, of course, be made much more efficient than having to handle a dense matrix, in addition to the statistical gain from a much smaller number of free parameters. In domains with such topological structure, similar input patterns are likely to appear at different positions, and nearby values (e.g., consecutive frames or nearby pixels) are likely to have stronger dependencies that are also important to model the data. In fact, these dependencies can be exploited to discover the topology [126], i.e., recover a regular grid of pixels out of a set of vectors without any order information, for example, after the elements have been arbitrarily shuffled in the same way for all examples. Thus, the same local feature computation is likely to be relevant at all translated positions of the receptive field. Hence, the idea of sweeping such a local feature extractor over the topology: This corresponds to a convolution and transforms an input into a similarly shaped feature map. Equivalently to sweeping, this may be seen as static but differently positioned replicated feature extractors that all share the same parameters. This is at the heart of convolutional networks [131], [133] that have been applied both to object recognition and to image segmentation [207]. Another hallmark of the convolutional architecture is that values computed by the same feature detector applied at several neighboring input locations are then summarized through a pooling operation,

typically taking their max or their sum. This confers upon the resulting pooled feature layer some degree of invariance to input translations, and this style of architecture (alternating selective feature extraction and invariance-creating pooling) has been the basis of convolutional networks, the Neocognitron [67] and HMAX [166] models, and argued to be the architecture used by mammalian brains for object recognition [166], [185], [61]. The output of a pooling unit will be the same irrespective of where a specific feature is located inside its pooling region. Empirically, the use of pooling seems to contribute significantly to improved classification accuracy in object classification tasks [133], [35], [36]. A successful variant of pooling connected to sparse coding is L2 pooling [103], [110], [122], for which the pool output is the square root of the possibly weighted sum of squares of filter outputs. Ideally, we would like to generalize feature pooling so as to learn what features should be pooled together, for example, as successfully done in several papers [100], [110], [122], [160], [53], [49], [76]. In this way, the pool output learns to be invariant to the variations captured by the span of the features pooled.

11.2.1 Patch-Based Training

The simplest approach for learning a convolutional layer in an unsupervised fashion is patch-based training: simply feeding a generic unsupervised feature learning algorithm with local patches extracted at random positions of the inputs. The resulting feature extractor can then be swiped over the input to produce the convolutional feature maps. That map may be used as a new input for the next layer, and the operation repeated to thus learn and stack several layers. Such an approach was recently used with independent subspace analysis [125] on 3D video blocks, reaching the state of the art on the Hollywood2, UCF, KTH, and YouTube action recognition datasets. Similarly, Coates and Ng [48] compared several feature learners with patch-based training and reached state-of-the-art results on several classification benchmarks. Interestingly, in this work, performance was almost as good with very simple k-means clustering as with more sophisticated feature learners. We, however, conjecture that this is the case only because patches are rather low-dimensional (compared to the dimension of a whole image). A large dataset might provide sufficient coverage of the space of, for example, edges prevalent in 6×6 patches so that a distributed representation is not absolutely necessary. Another plausible explanation for this success is that the clusters identified in each image patch are then pooled into a histogram of cluster counts associated with a larger subimage. Whereas the output of a regular clustering is a one-hot nondistributed code, this histogram is itself a distributed representation, and the "soft" k-means [48] representation allows not only the nearest filter, but also its neighbors to be active.

11.2.2 Convolutional and Tiled-Convolutional Training

It is possible to directly train large convolutional layers using an unsupervised criterion. An early approach [105] trained a standard but deep convolutional MLP on the task of denoising images, i.e., as a deep, convolutional, DAE. Convolutional versions of the RBM or its extensions have traded as here 28,2023 at 19:5504 HTC from 1555 Valera. Postrictions apply.

also been developed [57], [135], [202] as well as a probabilistic max-pooling operation built into convolutional deep networks [135], [136], [115]. Other unsupervised feature learning approaches that were adapted to the convolutional setting include PSD [110], [111], [106], [84], a convolutional version of sparse coding called deconvolutional networks [226], topographic ICA [122], and mPoT, which Kivinen and Williams [113] applied to modeling natural textures. Gregor and LeCun [74] and Le et al. [122] also demonstrated the technique of tiled convolution, where parameters are shared only between feature extractors whose receptive fields are k steps away (so the ones looking at immediate neighbor locations are not shared). This allows pooling units to be invariant to more than just translations and is a hybrid between convolutional networks and earlier neural networks with local connections but no weight sharing [128], [130].

11.2.3 Alternatives to Pooling

Alternatively, one can also use explicit knowledge of the expected invariants expressed mathematically to define transformations that are *robust* to a known family of input deformations, using the so-called scattering operators [139], [40], which can be computed in a way interestingly analogous to deep convolutional networks and wavelets. Like convolutional networks, the scattering operators alternate two types of operations: convolution and pooling (as a norm). Unlike convolutional networks, the proposed approach keeps at each level all of the information about the input (in a way that can be inverted) and automatically yields a very sparse (but very high dimensional) representation. Another difference is that the filters are not learned but instead are set so as to guarantee that a priori specified invariances are robustly achieved. Just a few levels were sufficient to achieve impressive results on several benchmark datasets.

11.3 Temporal Coherence and Slow Features

The principle of identifying slowly moving/changing factors in temporal/spatial data has been investigated by many [6], [219], [96], [114], [41] as a principle for finding useful representations. In particular, this idea has been applied to image sequences and as an explanation for why V1 simple and complex cells behave the way they do. A good overview can be found in [96], [31].

More recently, temporal coherence has been successfully exploited in deep architectures to model video [147]. It was also found that temporal coherence discovered visual features similar to those obtained by ordinary unsupervised feature learning [28], and a temporal coherence penalty has been *combined* with a training criterion for unsupervised feature learning [227], sparse autoencoders with L1 regularization, in this case, yielding improved classification performance.

The temporal coherence prior can be expressed in several ways, the simplest being the squared difference between feature values at times t and t+1. Other plausible temporal coherence priors include the following: First, instead of penalizing the squared change, penalizing the absolute value (or a similar sparsity penalty) would state that most of the time the change should be exactly 0, which would intuitively make sense for the real-life factors that

surround us. Second, one would expect that instead of just being slowly changing, different factors could be associated with their own different time scale. The specificity of their time scale could thus become a hint to disentangle explanatory factors. Third, one would expect that some factors should really be represented by a *group of numbers* (such as the x, y, and z position of some object in space and the pose parameters of Hinton et al. [85]) rather than by a single scalar and that these groups tend to move together. Structured sparsity penalties [110], [107], [3], [76] could be used for this purpose.

11.4 Algorithms to Disentangle Factors of Variation

The goal of building invariant features is to remove sensitivity of the representation to directions of variance in the data that are uninformative to the task at hand. However, it is often the case that the goal of feature extraction is the *disentangling* or separation of many distinct but informative factors in the data, for example, in a video of people: subject identity, action performed, subject pose relative to the camera, and so on. In this situation, the methods of generating invariant features, such as feature pooling, may be inadequate.

The process of building invariant features can be seen as consisting of two steps. First, low-level features are recovered that account for the data. Second, subsets of these low-level features are pooled together to form higher level invariant features, exemplified by the pooling and subsampling layers of convolutional neural networks. The invariant representation formed by the pooling features offers an incomplete window on the data as the detailed representation of the lower level features is abstracted away in the pooling procedure. While we would like higher level features to be more abstract and exhibit greater invariance, we have little control over what information is lost through pooling. What we really would like is for a particular feature set to be invariant to the irrelevant features and disentangle the relevant features. Unfortunately, it is often difficult to determine a priori which set of features will ultimately be relevant to the task at hand.

An interesting approach to taking advantage of some of the factors of variation known to exist in the data is the transforming autoencoder [85]: Instead of a scalar pattern detector (e.g., corresponding to the probability of the presence of a particular form in the input), one can think of the features as organized in groups that include both a pattern detector and pose parameters that specify attributes of the detected pattern. In [85], what is assumed a priori is that pairs of examples (or consecutive ones) are observed with an associated value for the corresponding change in the pose parameters. For example, an animal that controls its eyes knows what changes to its ocular motor system were applied when going from one image on its retina to the next. In that work, it is also assumed that the pose changes are the same for all the pattern detectors, and this makes sense for global changes such as image translation and camera geometry changes. Instead, we would like to discover the pose parameters and attributes that should be associated with each feature detector without having to specify ahead of time what they should be, force them to be the same for all features, and having to necessarily observe the changes in all of the pose parameters or attributes.

The approach taken recently in the MTC, discussed in Section 8.3, is interesting in this respect. Without any supervision or prior knowledge, it finds prominent local factors of variation (tangent vectors to the manifold, extracted from a CAE, interpreted as locally valid input "deformations"). Higher level features are subsequently encouraged to be *invariant* to these factors of variation so that they must depend on other characteristics. In a sense, this approach is disentangling valid local deformations along the data manifold from other, more drastic changes associated to other factors of variation such as those that affect class identity.²⁶

One solution to the problem of information loss that would fit within the feature-pooling paradigm is to consider many overlapping pools of features based on the same low-level feature set. Such a structure would have the potential to learn a redundant set of invariant features that may not cause significant loss of information. However, it is not obvious what learning principle could be applied that can ensure that the features are invariant while maintaining as much information as possible. While a DBN or a DBM (as discussed in Sections 4 and 10.2, respectively) with two hidden layers would, in principle, be able to preserve information into the "pooling" second hidden layer, there is no guarantee that the second layer features are more invariant than the "low-level" first layer features. However, there is some empirical evidence that the second layer of the DBN tends to display more invariance than the first layer [64].

A more principled approach, from the perspective of ensuring a more robust compact feature representation, can be conceived by reconsidering the disentangling of features through the lens of its generative equivalent—feature composition. Since many unsupervised learning algorithms have a generative interpretation (or a way to reconstruct inputs from their high-level representation), the generative perspective can provide insight into how to think about disentangling factors. The majority of the models currently used to construct invariant features have the interpretation that their low-level features linearly combine to construct the data.²⁷ This is a fairly rudimentary form of feature composition with significant limitations. For example, it is not possible to linearly combine a feature with a generic transformation (such as translation) to generate a transformed version of the feature. Nor can we even consider a generic color feature being linearly combined with a grayscale stimulus pattern to generate a colored pattern. It would seem that if we are to take the notion of disentangling seriously, we require a richer interaction of features than that offered by simple linear combinations.

12 CONCLUSION

This review of representation learning and deep learning has covered three major and apparently disconnected

26. The changes that affect class identity might, in input space, actually be of similar magnitude to local deformations, but not follow along the manifold, i.e., cross zones of low density.

27. As an aside, if we are given only the values of the higher level pooling features, we cannot accurately recover the data because we do not know how to apportion credit for the pooling feature values to the lower level features. This is simply the generative version of the consequences of the loss of information caused by pooling.

approaches: the probabilistic models (both the directed kind such as sparse coding and the undirected kind such as Boltzmann machines), the reconstruction-based algorithms related to autoencoders, and the geometrically motivated manifold-learning approaches. Drawing connections between these approaches is currently a very active area of research and is likely to continue to produce models and methods that take advantage of the relative strengths of each paradigm.

Practical concerns and guidelines. One of the criticisms addressed to artificial neural networks and deep learning algorithms is that they have many hyperparameters and variants and that exploring their configurations and architectures is an art. This has motivated an earlier book on the "Tricks of the Trade" [156] of which [132] is still relevant for training deep architectures, in particular what concerns initialization, ill conditioning, and stochastic gradient descent. A good and more modern compendium of good training practice, particularly adapted to training RBMs, is provided in [90], while a similar guide oriented more toward deep neural networks can be found in [13], both of which are part of a novel version of the above book. Recent work on automating hyperparameter search [29], [30], [191] is also making it more convenient, efficient, and reproducible.

Incorporating generic AI-level priors. We have covered many high-level generic priors that we believe could bring machine learning closer to AI by improving representation learning. Many of these priors relate to the assumed existence of multiple underlying factors of variation, whose variations are in some sense orthogonal to each other. They are expected to be organized at multiple levels of abstraction; hence the need for deep architectures, which also have statistical advantages because they allow to reuse parameters in a combinatorially efficient way. Only a few of these factors would typically be relevant for any particular example, justifying sparsity of representation. These factors are expected to be related to simple (e.g., linear) dependencies, with subsets of these explaining different random variables of interest (inputs, tasks) and varying in structured ways in time and space (temporal and spatial coherence). We expect future successful applications of representation learning to refine and increase that list of priors, and to incorporate most of them instead of focusing on only one. Research in training criteria that better take these priors into account is likely to move us closer to the long-term objective of discovering learning algorithms that can disentangle the underlying explanatory factors.

Inference. We anticipate that methods based on directly parameterizing a representation function will incorporate more and more of the iterative type of computation one finds in the inference procedures of probabilistic latent-variable models. There is already movement in the other direction, with probabilistic latent-variable models exploiting approximate inference mechanisms that are themselves learned (i.e., producing a parametric description of the representation function). A major appeal of probabilistic models is that the semantics of the latent variables are clear, and this allows a clean separation of the problems of modeling (choose the energy function), inference (estimating $P(h \mid x)$), and learning (optimizing the parameters), using generic tools in each case. On the other hand, doing approximate inference and not taking

loss of information caused by pooling. Other hand, doing approximate inference and not taking Authorized licensed use limited to: National Library Of Medicine (NLM). Downloaded on June 28,2023 at 18:56:01 UTC from IEEE Xplore. Restrictions apply.

that approximation into account explicitly in the approximate optimization for learning could have detrimental effects; hence the appeal of learning approximate inference. More fundamentally, there is the question of the multimodality of the posterior $P(h \mid x)$. If there are exponentially many probable configurations of values of the factors h_i that can explain x, then we seem to be stuck with very poor inference, either focusing on a single mode (MAP inference), assuming some kind of strong factorization (as in variational inference), or using an MCMC that cannot visit enough modes of $P(h \mid x)$. What we propose as food for thought is the idea of dropping the requirement of an explicit representation of the posterior and settling for an implicit representation that exploits potential structure in $P(h \mid x)$ to represent it compactly: Even though $P(h \mid x)$ may have an exponential number of modes, it may be possible to represent it with a small set of numbers. For example, consider computing a deterministic feature representation f(x) that implicitly captures the information about a highly multimodal $P(h \mid x)$, in the sense that all the questions (e.g., making some prediction about some target concept) that can be asked from $P(h \mid x)$ can also be answered from f(x).

Optimization. Much remains to be done to better understand the successes and failures of training deep architectures, both in the supervised case (with many recent successes) and the unsupervised case (where much more work needs to be done). Although regularization effects can be important on small datasets, the effects that persist on very large datasets suggest some optimization issues are involved. Are they more due to local minima (we now know there are huge numbers of them) and the dynamics of the training procedure? Or are they due mostly to ill conditioning and may be handled by approximate secondorder methods? These basic questions remain unanswered and deserve much more study.

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