Efficient Learning of Sparse Invariant Representations

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Abstract

We propose a simple and efficient algorithm for learning sparse invariant representations from unlabeled data with fast inference. When trained on short movies sequences, the learned features are selective to a range of orientations and spatial frequencies, but robust to a wide range of positions, similar to complex cells in the primary visual cortex. We give a hierarchical version of the algorithm, and give guarantees of fast convergence under certain conditions.

1 Introduction

Learning representations that are invariant to irrelevant transformations of the input is an important step towards building recognition systems automatically. Invariance is a key property of some cells in the mammalian visual cortex. Cells in high-level areas of the visual cortex respond to objects categories, and are invariant to a wide range of variations on the object (pose, illumination, confirmation, instance, etc). The simplest known example of invariant representations in visual cortex are the complex cells of V1 that respond to edges of a given orientation but are activated by a wide range of positions of the edge. Many artificial object recognition systems have built-in invariances, such as the translational invariance of convolutional network [1], or SIFT descriptors [2]. An important question is how can useful invariant representations of the visual world be learned from unlabeled samples.

In this paper we introduce an algorithm for learning features that are invariant (or robust) to common image transformations that typically occur between successive frames of a video or statistically within a single frame. While the method is quite simple, it is also computationally efficient, and possesses provable bounds on the speed of inference. The first component of the model is a layer of sparse coding. Sparse coding [3] constructs a dictionary matrix $W$ so that input vectors can be represented by a linear combination of a small number of columns of the dictionary matrix. Inference of the feature vector $z$ representing an input vector $x$ is performed by finding the $z$ that minimizes the following energy function

$$E_1(W, x, z) = \frac{1}{2} \|x - Wz\|^2 + \alpha |z|_{L^1}$$

where $\alpha$ is a positive constant. The dictionary matrix $W$ is learned by minimizing $\min_z E(W, x^k, z)$ averaged over a set of training samples $x^k, k = 1 \ldots K$, while constraining the columns of $W$ to have norm 1.

The first idea of the proposed method is to accumulate sparse feature vectors representing successive frames in a video, or versions of an image that are distorted by transformations that do not affect the nature of its content.

$$z^* = \sum_t \argmin_{z} \frac{1}{2} \|x_t - Wz\|^2 + \alpha |z|_{L^1}$$

1This paper was completed in June 2010, and submitted (unsuccessfully) to NIPS 2010.
where the sum runs over the distorted images \( x_t \). The second idea is to connect a second sparse coding layer on top of the first one that will capture dependencies between components of the accumulated sparse code vector. This second layer models vector \( z^* \) using an invariant code \( u \), which is the minimum of the following energy function

\[
E_2(A, z^*, u) = \alpha \sum_i z_i^* e^{(-Au)_i} + \beta |u|
\]

(3)

where \(|u|\) denotes the L1 norm of \( u \), \( A \) is a matrix, and \( \beta \) is a positive constant controlling the sparsity of \( u \). Unlike with traditional sparse coding, in this method the dictionary matrix interacts multiplicatively with the input \( z^* \). As in traditional sparse coding, the matrix \( A \) is trained by gradient descent to minimize the average energy for the optimal \( u \) over a training set of vectors \( z^* \) obtained as stated above. The columns of \( A \) are constrained to be normalized to 1. Essentially, the matrix \( A \) will connect a component of \( u \) to a set of components of \( z^* \) if these components of \( z^* \) co-occur frequently. When a component of \( u \) turns on, it has the effect of lowering the coefficients of the components of \( |z_i^*| \) to which it is strongly connected through the \( A \) matrix. To put it another way, if a set of components of \( z^* \) often turn on together, the matrix \( A \) will connect them to a component of \( u \). Turning on this component of \( u \) will lower the overall energy (if \( \beta \) is small enough) because the whole set of components of \( z^* \) will see their coefficient being lowered (the exponential terms).

Hence, each unit of \( u \) will connect units of that often turn on together within a sequence of images. These units will typically represent distorted version of a feature.

The energies (1) and (5) can be naturally combined into a single combined model of \( z \) and \( u \) as explained in section 2. There the second layer \( u \) is essentially modulating sparsity of the first layer \( z \). Single model of the image is more natural. For the invariance properties we didn’t find much qualitative difference and since the former has provable inference bounds we presented the results for separate training. However the a two layer model should capture the statistics of an image. To demonstrate this we compared the in-painting capability of one and two layer models and found that two layer model does better job. For these experiments, the combined two layer model is necessary. We also found that despite the assumptions of the fast inference are not satisfied for the two layer model, empirically the inference is fast in this case as well.

1.1 Prior work on Invariant Feature Learning

The first way to implement invariance is to take a known invariance, such as translational invariance in images, in put it directly into the architecture. This has been highly successful in convolutional neural networks (1) and SIFT descriptors (2) and its derivatives. The major drawback of this approach is that it works for known invariances, but not unknown invariances such as invariance to instance of an object. A system that would discover invariance on its own would be desired.

Second type of invariance implementation is considered in the framework of sparse coding or independent component analysis. The idea is to change a cost function on hidden units in a way that would prefer co-occurring units to be close together in some space (4)(5). This is achieved by pooling units close in space together. This groups different inputs together producing a form of invariance. The drawback of this approach is that it requires some sort of imbedding in space and that the filters have to arrange themselves.

In the third approach, rather then forcing units to arrange themselves, we let them learn whatever representations they want to learn and instead figure out which to pool together. In (6)(7), this was achieved by modulating covariance of the simple units with complex units.

The fourth approach to invariance uses the following idea: If the inputs follow one another in time they are likely a consequence of the same cause. We would like to discover that cause and therefore look for representations that are common for all frames. This was achieved in several ways. In slow feature analysis (8)(9)(10) one forces the representation to change slowly. In temporal product network (11) one breaks the input into two representations - one that is common to all frames and one that is complementary. In (12) the idea is similar but in addition the complementary representation specifies movement. In the simplest instance of hierarchical temporal memory (13) one forms groups based on transition matrix between states. The (14) is a structured model of video.
A lot of the approaches for learning invariance are inspired by the fact that the neo-cortex learns to create invariant representations. Consequently these approaches are not focused on creating efficient algorithms. In this paper, we give an efficient learning algorithm that falls into the framework of third and fourth approaches. The basic idea is to modulate the sparsity of sparse coding units using higher level units that are also sparse. The fourth approach is implemented by using the same higher level representation for several consecutive time frames. In the form our model is similar to that of [6, 7] but a little simpler. In a sense comparing our model to [6, 7] is similar to comparing sparse coding to independent component analysis. Independent component analysis is a probabilistic model, whereas sparse coding attempts to reconstruct input in terms of few active hidden units. The advantage of sparse coding is that it is simpler and easier to optimize. There exist several very efficient inference and learning algorithms [15, 16, 17, 18, 19] and sparse coding has been applied to a large number problems. It is this simplicity that allows efficient training of our model. The inference algorithm is closely derived from the fast iterative shrinkage-thresholding algorithm (FISTA) [15] and has a convergence rate of $1/k^2$ where $k$ is the number of iterations.

2 The Model

The model described above comprises two separately trained modules, whose inference is performed separately. However, one can devise a unified model with a single energy function that is conceptually simpler:

$$E(W, A, x, z, u) = \frac{1}{2} \sum_t \| x_t - W z_t \|^2 + \sum_i \alpha_t | g(u)_i | + h(u)$$

(5)

$$g(u)_i = (1 + e^{-(A u)_i})/2, \quad h(u) = \beta |u|$$

Given a set of inputs $x^\gamma$, the goal of training is to minimize $\sum_\gamma E(W, A, x^\gamma, z^\gamma, u^\gamma)$. We do this by choosing one input $x$ at a time, minimizing (5) over $z$ and $u$ with $W$ and $A$ fixed, then fixing the resulting $z_{\text{min}}, u_{\text{min}}$ and taking step in a negative gradient direction of $W, A$ (stochastic gradient descent). An algorithm for finding $z_{\text{min}}, u_{\text{min}}$ is given in section 4. It consists of taking step in $z$ and $u$ separately, each of which lowers the energy.

Note: The $g$ functions in (5) is different from that of the simple (split) model. The reason is that, in our experiments, either $u$ units lower the sparsity of $z$ too much, not resulting in a sparse $z$ code or the units $u$ do not turn on at all.

2.1 A Toy Example

We now describe a toy example that illustrates the main idea of the model [20]. The input, with $n_t = 1$, is an image patch consisting of a subset of the set of parallel lines of four different orientations and ten different positions per orientation. However for any given input, only lines with the same orientation can be present, Figure 4 (different orientations have equal probability and for a given orientation a line of this orientation is present with probability 0.2 independently of others). This is a toy example of a texture. Training sparse coding on this input results in filters similar to one in Figure 4. We see that a given simple unit responds to one particular line. The noisy filters correspond to simple units that are inactive - this happens because there are only 40 discrete inputs. In realistic data such as natural images, we have a continuum and typically all units are used.

Clearly, sparse coding cannot capture all the statistics present in the data. The simple units are not independent. We would like to learn that that units corresponding to lines of a given orientation usually turn on simultaneously. We trained (5) on this data resulting in the filters in the Figure 4c. The filters of the simple units of this full model are similar to those obtained by training just the sparse coding. The invariant units pool together simple units with filters corresponding to lines of the same orientation. This makes the invariant units invariant to the pattern of lines and dependent only on the orientation. Only four invariant units were active corresponding to the four groups. As in sparse coding, on a realistic data such as natural images, all invariant units become active and distribute themselves with overlapping filters as we will see below.

Let us now discuss the motivation behind introducing a sequence of inputs ($n_t > 1$) in (5). Inputs that follow one another in time are usually a consequence of the same cause. We would like to
Figure 1: Toy example trained using model (5). a) Randomly selected input image patches. The input patches are generated as follows. Pick one of the four orientations at random. Consider all lines of this orientation. Put any such line into the image independently with probability 0.2. b) Learned sparse coding filters. A given active unit responds to a particular line. c) Learned filters of the invariant units. Each row corresponds to an invariant unit. The sparse coding filters are ordered according to the strength of their connection to the invariant unit. There are only four active units (arrows) and each responds to a given orientation, invariant to which lines of a given orientation are present.

discover that cause. This cause is something that is present at all frames and therefore we are looking for a single representation $u$ in (5) that is common to all the frames.

Another interesting point about the model (5) is that nonzero $u$ lowers the sparsity coefficient of units of $z$ that belong to a group making them more likely to become activated. This means that the model can utilize higher level information (which group is present) to modulate the activity of the lower layer. This is a desirable property for multi-layer systems because different parts of the system should propagate their belief to other parts. In our invariance experiments the results for the unified model were very similar to the results of the simple (split) model. Below we show the results of this simple model because it is simple and because we provably know an efficient inference algorithm. However in the section 4 we will revisit the full system, generalize it to an $n$-layer system, give an algorithm for training it, and prove that under some assumptions of convexity, the algorithm again has a provably efficient inference. In the final section we use the full system for in-painting and show that it generalizes better than a single layer system.

3 Efficient Inference for the Simplified Model

Here we discuss how to find $u$ efficiently and give the numerical results of the paper. The results for the full model (5) were similar.

3.1 FISTA training algorithm

The advantage of (3) compared to (5) is that the fast iterative shrinkage-thresholding algorithm (FISTA) [15] applies to it directly. FISTA applies to problems of the form $E(u) = f(u) + \hat{g}(u)$ where:

- $f$ is continuously differentiable, convex and Lipschitz, that is $\|\nabla f(u_1) - \nabla f(u_2)\| \leq L(f)\|u_1 - u_2\|$. The $L(f) > 0$ is the Lipschitz constant of $\nabla f$.
- $\hat{g}$ is continuous, convex and possibly non-smooth

The problem is assumed to have solution $E^* = E(u^*)$. In our case $f(u) = \alpha|z_t|\rho(u)$ and $\hat{g}(u) = h(u)$ which satisfies these assumptions ($A$ is initialized with nonnegative entries which stay nonnegative during the algorithm without a need to force it). This solution converges with bound $E(u_k) - E(u^*) \leq 2\alpha L(f)\|u_0 - u^*\|^2/(k + 1)^2$ where $u_k$ is the value of $u$ at the $k$-th
iteration and $\alpha$ is a constant. The cost of each iteration is $O(mn)$ where $n$ is the input size and $m$ is the output size. More precisely the cost is one matrix multiplications by $A$ and by $A^t$ plus $O(m+n)$ cost. We used the back-tracking version of the algorithm to find $L$ which contains a fixed number of $O(mn)$ operations (independent of desired error). It is a standard knowledge and easy to see that the algorithm applies to the sparse coding (1) as well.

3.2 Results

The input to the network was prepared as follows. We converted all the images of the Berkeley data-set into gray-scale images. We locally removed the mean for each pixel by subtracting a Gaussian-weighted average of the nearby pixels. The width of the Gaussian was 9 pixels. Then, we locally normalized the contrast by dividing each pixel by Gaussian-weighted standard deviation of the nearby pixels (with a small cutoff to prevent blow-ups). The width of the Gaussian was also 9 pixels. Then, we picked a $20 \times 20$ window in the image and, for a randomly chosen direction and magnitude, we moved it for $n_t = 3$ frames and extracted them. The magnitude of the displacement was random in the range of $1 - 2$ pixels. A very large collection of such triplets of frames was extracted. We trained the sparse coding algorithm with 400 code units in $z$ on each individual frame (not on the $n_t$ concatenated frames). After training we found the sparse codes for each frame. There were 100 units in the layer of invariant units $u$. For larger a system with 1000 simple units and 400 invariant units, see the supplementary material.

![Figure 2: Connectivity between invariant units and simple units.](image)

**Left:** Each row correspond to an invariant unit. For a given invariant unit, the filters of the simple units were ordered according to the size of the weight to the invariant unit. The strongest 9 are plotted. **Middle:** Each square correspond to an invariant unit. 25 out of 100 invariant units were selected at random. The sparse coding filters were fitted with gabor functions. The center position of each line is the center of the Gabor filter. The orientation is the orientation of the sine-wave of the Gabor filter. The brightness is proportional to the strength of connection between invariant and simple units. For each invariant unit the weights were normalized for drawing so that the strongest connection is white and zero connections are black. **Right:** Each square (circle) corresponds to an invariant unit. Each dot correspond to a simple unit. The distance from the center is the frequency of the Gabor fit. The angle is twice the orientation of the Gabor fit (twice because angles related by $\pi$ are equivalent). We see that invariant units typically learn to group together units with similar orientation and frequency. There are few other types of filters as well. The units in the middle and right panel correspond to each other and correspond to the units in the left panel reading panels left to right and then down. See the supplementary material for all the filters the system: $20 \times 20$ input patches, 1000 simple units, 400 invariant units.

The results are shown in the Figure 2. We see that many invariant cells learn to group together filters of similar orientation and frequency but at several positions and thus learn invariance with respect to translations. However there are other types of filters as well. Remember that the algorithm learns statistical co-occurrence between features, whether in time or in space.
The values of the weights give us important information about the properties of the system. However ultimately we are interested in how the system responds to an input. We study the response of these units to a commonly occurring input - an edge. Specifically the inputs are given by the following function.

\[
X(x, y) = e^{-0.25v^2} \sin(v)
\]

\[
v = \vec{k} \cdot \vec{r} + kb
\]

\[
\vec{k} = k(\cos \theta, \sin \theta) \text{ and } \vec{r} = (x, y)
\]

where \((x, y)\) is the position of a pixel from the center of a patch, \(b\) a real number specifying distance of the edge from the center and \(\theta\) is the orientation of the edge from the \(x\) axis. This is not an edge function, but a function obtained on an edge after local mean subtraction.

The responses of the simple units and the invariant units are shown in the Figure. See caption for description. As expected the sparse coding units respond to edges in a narrow range of positions and relatively narrow range of orientations. Invariant cells on the other hand are able to pool different sparse coding units together and become invariant to a larger range of positions. Thus the invariant units do indeed have the desired invariance properties.

Note that for large sparsities the regions have clear boundaries and are quite sharp. This is similar to the standard implementation of convolutional net, where the pooling regions are squares (with clear boundaries). It is probably more preferable to have regions that overlap as happens at lower sparsities since one would prefer smoother responses rather then jumps across boundaries.

4 Theoretical analysis of the full model and its multi-layer generalization

In this section we return to the full model (5). We generalize it to an \(n\) layer system, give an inference algorithm and outline the proof that under certain assumptions of convexity the algorithm has the fast \(1/k^2\) convergence of FISTA, there \(k\) is the iteration number.
The basic idea of minimizing over \( z, u \) in (5) is to alternate between taking energy-lowering step in \( z \) while fixing \( u \) and taking energy-lowering step in \( u \) while fixing \( z \). Note that both of the restricted problems (problem in \( z \) fixing \( u \) and problem in \( u \) fixing \( z \)) satisfy conditions of the FISTA algorithm. This will allow us to take steps of appropriate size that are guaranteed to lower the total energy. Before that however, we generalize the problem, which will reveal its structure and which does not introduce any additional complexity.

Consider system consisting of \( n \) layers with units \( z_a \) in the \( a \)-th layer with \( a = 1, \ldots, n \). We define \( z = (z_1, \ldots, z_n) \) that is all the vectors \( z_a \) concatenated. We define two sets of functions. Let \( e_\alpha(z_a) \) be continuously differentiable, convex and Lipschitz functions. There can be several such functions per layer, which is denoted by index \( \beta \). Let \( g^\beta_\alpha(z_a) \) be continuous and convex functions, not necessarily smooth. For convenience we define \( z_0 = 0, z_{n+1} = 0, g_0 = 1 \) and \( e_{n+1} = 1 \).

We define the energy of the system to be

\[
E = \sum_{a=0}^{n} \sum_{\beta} g^\beta_\alpha(z_a) e_\alpha(z_{a+1}) = \sum_{a=0}^{n} g_\alpha(z_a) e_\alpha(z_{a+1})
\]

(9)

where in the second equality we drop the \( \beta \) from the notation for simplicity. We will omit writing the \( \beta \) for the rest of the paper. The equation (5) is a special case of (9) with \( n = 2, g_0 = 1, e_1(z_1) = (1/2)||x-Wz_1||^2, g_1(z_1) = |z_1|, e_2(z_2) = \alpha(1 + e^{-Az_2})/2, g_2(z_2) = \beta|z_2| \) with \( \bar{z} \geq 0 \) and \( e_3 = 1 \).

Now observe that given \( a \in \{1, \ldots, n\} \) the problem in \( z_a \) keeping other variables fixed satisfies the conditions of the FISTA algorithm. We can define a step in the variable \( z_a \) to be (in analogy to (5) eq. (2.6)):

\[
p_{L_a}(z_a) = \argmin_{z_a} \left\{ g(z_a') e_{a+1}(z_{a+1}) + \frac{L}{2} \left| z_a' - \left( z_a - \frac{1}{L} g_a(z_{a-1}) \nabla e_a(z_a) \right) \right|^2 \right\}
\]

\[
= \text{sh}_{\frac{\left| z_a - \frac{1}{L} g_a(z_{a-1}) \nabla e_a(z_a) \right|}{L_a}} (z_a - \frac{1}{L_a} g_a(z_{a-1}) \nabla e_a(z_a))
\]

(10)

where the later equality holds if \( g_a(z_a) = |z_a| \). Here sh is the shrinkage function \( \text{sh}_\alpha(z) = \text{sign}(z)(|z| - \alpha)_+ \). In the case where \( z_a \) is restricted to be nonnegative we need to use \( (z - \alpha)_+ \) instead of the shrinkage function.

Let us describe the algorithm for minimizing (9) with respect to \( z \) (we will write it explicitly below). In the order from \( a = 1 \) to \( a = n \), take the step \( z'_a = p_{L_a}(z_a) \) in (10). Repeat until desired accuracy.

The \( L_a \)'s have to be chosen so that

\[
g_{a-1}(z_{a-1}) e_a(z'_a) \leq g_{a-1}(z_{a-1}) e_a(z_a) + \langle z'_a - z_{a-1}, g_{a-1}(z_{a-1}) \nabla e_a(z_{a-1}) \rangle + \frac{1}{2} || z'_a - z_a ||^2 \]

(11)

This can be assured by taking \( L_a \geq L(g_{a-1} e_a) \) where the later \( L \) denotes the Lipschitz constant of its argument. Otherwise, as used in our simulations, it can be found by backtracking, see below.

This will assure that each steps lowers the total energy (9) and hence the overall procedure will keep lowering it. In fact the step \( p_{L_a} \) with such chosen \( L \) is in some sense a step with ideal step size. Let us now write the algorithm explicitly:

**Hierarchical (F)ISTA.**

Step 0. Take \( L_0 > 0 \), some \( \eta > 1 \) and \( z_0^a \in \mathbb{R} \). Set \( z_1 = z_0^a, t_1 = 1, a = 1, \ldots, n \).

Step k. \((k \geq 1)\).

Loop a=1:n { Backtracking }

\[
\text{Find smallest nonnegative integer } i^a_k \text{ such that with } L^a = \eta^i_k L_{k-1}^a
\]

\[
g_{a-1}(z_{a-1}^{k-1}) e_a(p_{L^a}(z^a_k)) \leq g_{a-1}(z_{a-1}^{k-1}) e(z^a_k) + \langle p_{L^a}(z^a_k) - z^a_k, g_{a-1}(z_{a-1}^{k-1}) \nabla e(z^a_k) \rangle + \frac{1}{2} || p_{L^a}(z^a_k) - z^a_k ||^2
\]

(12)

Set \( L_k^{a} = \eta^i_k L_{k-1}^{a} \}

Compute

\[
\bar{z}^a_k = p_{L_k^a}(z^a_k)
\]

(14)
\[ r_k = 0 \quad \text{or} \quad t_{k+1} = \frac{1 + \sqrt{1 + 4r_k^2}}{2}; \quad r_k = \frac{t_k - 1}{t_{k+1}} \]  

(15)

\[
\text{Loop a=1:n}\{ \begin{array}{l}
\hat{z}_{k+1}^a = \hat{z}_k^a + r_k(\hat{z}_k^a - z_{k-1}^a) \\
\end{array} \}
\]  

(16)

The algorithm described above is this algorithm with the choice \( r_k = 0 \) in the second last line.

Let us discuss the \( r_k \)'s. For single layer system \((n = 1)\) \( r_k = 0 \) choice is called ISTA and has convergence bound of \( E(z_k) - E(z^*) \leq \alpha L\|z_0 - z^*\|^2/2k \). The other choice of \( r_k \) is the FISTA algorithm. The convergence rate of FISTA is much better than that of ISTA, having \( k^2 \) in the denominator.

For hierarchical system, the choice \( r_k = 0 \) guarantees that each step lowers the energy. The question is whether introducing the other choice of \( r_k \) would speed up convergence to the FISTA convergence rate. The trouble is that the in general the product \( \gamma \) would certainly not be guaranteed (imagine starting at a non-minimal point with zero derivative).

The proof in [15] shows that if the algorithm consists of applying the sequence of \( P_a \)'s and these \( P_a \)'s satisfy Lemma 2.3, then the algorithm converges with rate \( E(z_k) - E(z_a) \leq 2L\max\|z_0 - z_a\|^2/(k+1)^2 \). Thus we need to prove Lemma 2.3. We start with the analog of Lemma 2.2 of [15].

Lemma 2.3: Assume that \( \forall a \in \{0, \ldots, n\}, e_a \) is continuously differentiable, Lipschitz, \( g_a \) is continuous, \( g_a e_{a+1} \) is convex and \( P_a \) is defined by the sequence of \( p_{L_a} \)'s of [10] as described above. Then for any \( z, \hat{z} \)

\[
E(\hat{z}) - E(P_L(z)) \geq \frac{L}{2}(P_L(z) - z)^2 + L(z - \hat{z}, P_L(z) - z)
\]

(17)

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Lemma 2.2: For any \( z \), one has \( z'_a = p_{L_a}^1(z_a) \) if and only if there exist \( \gamma_a(z_a) \in \partial g_a(z_a) \), the subdifferential of \( g_a(\cdot) \), such that

\[
g_a(z_a)\nabla e_a(z_a) + L(z'_a - z_a) + e(z_{a+1})\gamma_a(z_a) = 0
\]

(18)

This lemma follows trivially from the definition of \( p_{L_a} \) as in [15].

Proof of Lemma 2.3: Define \( z' = P_L(z) \). From convexity we have

\[
g_a(\hat{z}_a)e_{a+1}(\hat{z}_{a+1}) \geq g_a(z'_a)e_{a+1}(z_{a+1}) + (\langle \hat{z}_a - z'_a, e_{a+1}(z_{a+1})\rangle + \langle z'_a - z_{a+1}, g_a(z'_a)\nabla e_{a+1}(z_{a+1}) \rangle)
\]

(19)

Next we have the property (22). However the \( z_{a-1} \) should be primed (\( z'_{a-1} \)) because the \( z_{a-1} \) has already been updated. Due to space limitations we won’t write out all the calculations but specify the sequence of operations. The details are written out in the supplementary material. We take the first term on the left side of (20), \( E(\hat{z}) \) and express it in its terms (9). Then, replace the terms using the convexity equations and substitute \( \gamma(\hat{z}_a) \)'s using the Lemma 2.2. Then we take the second term of the left side of (20), \( E(P_L(z)) \), again express it using (9), and use the inequalities (22). Putting it all together, all the gradient terms cancel and the other terms combine to give Lemma 2.3. This completes the proof.
5 Conclusions

We introduced simple and efficient algorithm from learning invariant representation from unlabelled data. The method takes advantage of temporal consistency in sequential image data. In the future we plan to use the invariant features discovered by the method to hierarchical vision architectures, and apply it to recognition problems.

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A Supplementary material for Efficient Learning of Sparse Invariant Representations

1) We give details of the Proof of Lemma 2.3.
2) We show all of the invariant filters for system of: 20x20 patches input patches, 1000 simple units, 400 invariant units.

A.1 Lemma 2.3

Lemma 2.3: Assume that \( \forall a \in \{0, \ldots, n\}, \ e_a \) is continuously differentiable, Lipschitz, \( g_a \) is continuous, \( g_a e_{a+1} \) is convex and \( p_L \) is defined by the sequence of \( p_L \)'s in the paper. Then for any \( z, \hat{z} \)

\[
E(\hat{z}) - E(p_L(z)) \geq \frac{L}{2}|p_L(z) - z|^2 + L(z - \hat{z}, p_L(z) - z) \tag{20}
\]

Proof of Lemma 2.3: Define \( z' = p_L(z) \). We first collect the inequalities that we will need.
From convexity we have

\[
g_a(z_a) e_{a+1}(\hat{z}_{a+1}) \geq g_a(z_a') e_{a+1}(z_{a+1}) + \langle \hat{z}_a - z_a', e_{a+1}(z_{a+1}) \rangle + \langle z_{a+1} - z_{a+1}, g_a(z_a') \nabla e_{a+1}(z_{a+1}) \rangle \tag{21}
\]

Next we have the property for step \( p_L \) that guarantees that the energy is lowered in each step.

\[
g_{a-1}(z'_{a-1}) e_a(z_a) \leq g_{a-1}(z'_{a-1}) e_a(z_a) + \langle z_a' - z_a, g_{a-1}(z'_{a-1}) \nabla e_a(z_a) \rangle + (L/2)|z_a' - z_a|^2 \tag{22}
\]

Finally we have the Lemma 2.2

\[
g(z'_{a-1}) \nabla e(z_a) + L(z_a' - z_a) + e(z_{a+1}) \gamma(z_a) = 0 \tag{23}
\]

Now we can put these equations together. The steps are: Write out the left side of (20) in terms of the definition of \( E \). Use inequalities (21) and (22). Eliminate \( \gamma \)'s using (23). Simplify. Here are the details:

\[
E(\hat{z}) - E(z') = \sum_a g_a(\hat{z}_a) e_{a+1}(\hat{z}_{a+1}) - g_a(z'_a) e_{a+1}(z'_{a+1})
\]

\[
\geq \sum_a g_a(\hat{z}_a) e_{a+1}(z_{a+1}) + \langle \hat{z}_a - z'_a, e_{a+1}(z_{a+1}) \rangle + \langle z_{a+1} - z_{a+1}, g_a(z'_a) \nabla e_{a+1}(z_{a+1}) \rangle
\]

\[
- g_{a-1}(z'_{a-1}) e_a(z_a) - \langle z'_a - z_a, g_{a-1}(z'_{a-1}) \nabla e_a(z_a) \rangle - (L/2)|z'_a - z_a|^2
\]

\[
= \sum_a g_a(\hat{z}_a) e_{a+1}(z_{a+1}) - L(\hat{z}_a - z'_a, z'_a - z_a) - \langle \hat{z}_a - z'_a, g(z'_{a-1}) \nabla e(z_a) \rangle
\]

\[
+ \langle z_{a+1} - z_{a+1}, g_a(z'_a) \nabla e_{a+1}(z_{a+1}) \rangle
\]

\[
- g_{a-1}(z'_{a-1}) e_a(z_a) - \langle z'_a - z_a, g_{a-1}(z'_{a-1}) \nabla e_a(z_a) \rangle - (L/2)|z'_a - z_a|^2
\]

\[
= \sum_a -L(\hat{z}_a - z'_a, z' - z_a) - (L/2)|z' - z|^2
\]

which is the formula (22). Note that in the line 5 and in the first term of lines 6 we shifted \( a \) by one. This completes the proof.

A.2 Simple unit and invariant unit filters. \( \alpha = 0.5, \beta = 0.3 \)
Figure 4: Sparse coding filters. Inputs 20x20 images patches, preprocessed. code: 1000 simple units.
Figure 5: Selection of invariant units. See Figure 2a of the paper for explanation. System: 20x20 patches, 1000 simple units, 400 invariant units.
Figure 6: All 400 invariant units. See Figure 2b of the paper for explanation. System: 20x20 patches, 1000 simple units, 400 invariant units.
Figure 7: All 400 invariant units. Position of the invariant cells (centers of gabor fits). System: 20x20 patches, 1000 simple units, 400 invariant units.
Figure 8: All 400 invariant units. See Figure 2c of the paper for explanation. System: 20x20 patches, 1000 simple units, 400 invariant units.