Visual Recognition and Inference Using Dynamic Overcomplete Sparse Learning

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We present a hierarchical architecture and learning algorithm for visual recognition and other visual inference tasks such as imagination, reconstruction of occluded images, and expectation-driven segmentation. Using properties of biological vision for guidance, we posit a stochastic generative world model and from it develop a simplified world model (SWM) based on a tractable variational approximation that is designed to enforce sparse coding. Recent developments in computational methods for learning overcomplete representations (Lewicki & Sejnowski, 2000; Teh, Welling, Osindero, & Hinton, 2003) suggest that overcompleteness can be useful for visual tasks, and we use an overcomplete dictionary learning algorithm (Kreutz-Delgado, et al., 2003) as a preprocessing stage to produce accurate, sparse codings of images.

Inference is performed by constructing a dynamic multilayer network with feedforward, feedback, and lateral connections, which is trained to approximate the SWM. Learning is done with a variant of the backpropagation-through-time algorithm, which encourages convergence to desired states within a fixed number of iterations. Vision tasks require large networks, and to make learning efficient, we take advantage of the sparsity of each layer to update only a small subset of elements in a large weight matrix at each iteration. Experiments on a set of rotated objects demonstrate various types of visual inference and show that increasing the degree of overcompleteness improves recognition performance in difficult scenes with occluded objects in clutter.

1 Introduction

Vision, whether in the brain or computer, can be characterized as the process of inferring certain unknown quantities using an input image and...
predictions or expectations based on prior exposure to the environment. Visual inference includes tasks such as recognizing objects, reconstructing missing or occluded features, imagining previously learned or entirely novel objects, and segmentation (finding which features in a cluttered image correspond to a particular object). Performing these inference tasks requires combining information about the current image (bottom-up processing) and abstract concepts of objects (top-down processing). These tasks can naturally be placed into the framework of Bayesian probabilistic models, and determining the structure and priors for such models is a great challenge for both understanding vision in the brain and application-oriented computer vision. A primary goal of this letter is to derive an effective probabilistic model of visual inference consistent with current understanding of biological vision.

A number of important properties have emerged from neuroscience:

1. Vision in the brain is a hierarchical process with information flowing from the retina to the lateral geniculate nucleus (LGN), occipital and temporal regions of the cortex (Kandel, Schwartz, & Jessel, 2000).
2. This hierarchy has extensive recurrence with reciprocal connections between most regions (Felleman & Van Essen, 1991).
3. There is extensive recurrence within cortical regions, as typified by lateral inhibition, a mechanism for how sparse coding can arise (Callaway, 2004).
4. The primary visual cortex (V1) is strikingly overcomplete, meaning there are many more cells than are needed to represent the retinal information. In humans, there are over 200 to 300 V1 neurons per each LGN neuron and a lesser degree of overcompleteness in other primates (Stevens, 2001; Ejima et al., 2003).
5. The firing patterns of cortical neurons gives evidence for sparse distributed representations, in which only a few neurons are active out of a large population, and that information is encoded in these ensembles (Vinje & Gallant, 2000; Quiroga, Reddy, Kreiman, Koch, & Fried, 2005).
6. Although there are differences among areas, the basic structure of the cortex is qualitatively similar, and the notion of cortical similarity states that the underlying cortical operation should be similar from area to area (Mountcastle, 1978; Hawkins & Blakeslee, 2004).

Since these six properties are present in animals with high visual acuity, it is reasonable to assume they are important for inference, and we will adopt them in a network model.

While many computational models of vision have been developed that incorporate some of the properties listed above (Fukushima & Miyake,
1982; Rao & Ballard, 1997; Riesenhuber & Poggio, 1999; Rolls & Milward, 2000; Lee & Mumford, 2003; Fukushima, 2005), we propose a model that takes into account all six properties. For example, the recognition models of Rolls and Milward (2000) and Riesenhuber and Poggio (1999) do not use feedback (and so are incapable of inference tasks such as reconstruction or imagination), and the dynamic system of Rao and Ballard (1997) does not use overcomplete representations. The use of learned overcomplete representations for preprocessing is a new and largely unexplored approach for visual recognition and inference algorithms. Recent developments in learning overcomplete dictionaries (Lewicki & Sejnowski, 2000; Kreutz-Delgado et al., 2003; Teh, Welling, Osindero, & Hinton, 2003) and the associated methods for sparse image coding (Murray & Kreutz-Delgado, 2006) now make possible the investigation of their utility for visual inference.

Real-world images are high-dimensional data that can be explained in terms of a much smaller number of causes, such as objects and textures. Each object in turn can appear in many different orientations but in fact is seen in only one particular orientation. For each orientation, an object can be represented with a concise set of features, such as lines, arcs, and textures. The key feature of these various types of image descriptions is that they can be represented as sparse vectors, where only a few of the many possible choices suffice as explanation. While pixel values of images have nonsparse distributions (they are unlikely to be zero), these more abstract representations are very sparse (each component is likely to be zero), and only a few nonzero components at a time succinctly describe the scene. This intuition, along with the biological evidence for sparsity, is the justification for our use of sparse prior distributions. Other advantages of sparsity include reduced metabolic cost and increased storage capacity in associative memories (Olshausen & Field, 1997).

1.1 Overview and Organization. Beginning with a hypothetical hierarchical generative world model (GWM) that is presumed to create images of objects seen in the world, we discuss in section 2 how the GWM can be used for visual inference. The GWM requires the selection of a probability distribution, and a suitable choice is required to create practical algorithms. As a first move, we consider a Boltzmann-like distribution that captures the desired top-down, bottom-up, and lateral influences between and within layers, but it is computationally intractable. Then a simplified world model (SWM) distribution is created based on a variational approximation to the Boltzmann-like distribution and specifically designed to model sparse densities (see section 2.4).

By designing a dynamic network that rapidly converges to a self-consistency condition of the SWM, we can perform inference tasks if we have the weights that parameterize the network (see section 3). The dynamic network arises as a way of estimating the fixed-point state of the SWM. Although we consider only the problem of estimating static world
models, generalization to dynamic worlds is also possible. To determine the unknown weights, we develop a learning algorithm based on the backpropagation-through-time algorithm (Williams & Peng, 1990) which operates on the preactivation state and includes a sparsity-enforcing prior (see section 4). This algorithm can be seen as a natural extension of the sparse-coding principles that are useful in modeling V1 response properties (Olshausen & Field, 1997) to the full visual inference task.

We demonstrate experimentally several types of visual inference: recognition, reconstruction, segmentation, and imagination. These simulations show that overcomplete representations can provide better recognition performance than complete codes when used in the early stages of vision (see section 6). A discussion of the biological motivations and comparison to prior work is given in section 7, and conclusions are drawn in section 8.

1.2 Notation. We use the following notation:

- $a$: Activation function parameters
- $B$: Error-related term in learning algorithm
- $c^{(m)}$: Object code for object $m$, (sparse binary code)
- $D$: Sparsity-enforcing term in learning algorithm
- $f(\cdot)$: Sigmoid activation function
- $I(\cdot)$: Indicator function: 1 if the expression is true, 0 otherwise
- $J_{PA}$: Cost function on preactivation state, minimized by learning rule
- $K$: Number of images in training set $\mathcal{Y}$
- $L_l$: Lateral weights between units in layer $l$
- $M$: Number of unique objects in training set
- $n$: Number of layers in network
- $N$: Number of elements in state vector $X$
- $r$: Number of nonzero elements $r = [r_1, \ldots, r_n]$, where $r_l$ is the number of nonzero elements in layer $l$ (diversity $\times n$)
- $s$: Size of layers, $s = [s_1, \ldots, s_n]$, where $s_l$ is the size of layer $l$
- $U_t$: Network input at time $t$
- $v$, $v$: Unit weight sum $v$ (for entire layer $v_l$), preactivation function
- $\hat{V}_l$: Preactivation state of all layers
- $\hat{W}_{lm}$: Certainty-equivalence approximation of preactivation values
- $\mathcal{W}$: Complete weight matrix for all layers (including all $\hat{W}_{lm}$ and $L_l$), $\mathcal{W} \in \mathbb{R}^{N \times N}$
- $\mathbf{x}_l$: Activation vector at layer $l$, expected values of $P(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1})$
- $X$: State vector of all layers, $X = [\mathbf{x}_1^T, \ldots, \mathbf{x}_l^T]^T$
- $\hat{\mathbf{Y}}$: Training data, $\hat{\mathbf{Y}} = [\hat{\mathbf{y}}_1^T, 0, \ldots, 0, \hat{\mathbf{y}}_n^T]^T$, where $\hat{\mathbf{y}}_1$ is a sparsely coded image and $\hat{\mathbf{y}}_n$ is an object code
- $Y_t$: Dynamic network output at time $t$
- $\mathcal{Y}, \mathcal{V}, \mathcal{U}$: Sets of multiple state vectors $Y$, $V$ (e.g., $\mathcal{Y} = \{Y^{(1)}, \ldots, Y^{(K)}\}$)
In this section, we postulate a hierarchical generative visual-world model (GWM) and discuss its properties, particularly that of independence of the nodes of a layer conditioned on its immediately neighboring layers. We then discuss how the GWM can be used for visual inference tasks such as recognition, imagination, reconstruction, and expectation-driven segmentation. Specific forms of the probability distribution in the model must be chosen, and as a starting point, we posit a Boltzmann-like distribution. Since inference with the Boltzmann-like distribution is generally intractable, a variational approximation is developed leading to a simplified world model (SWM). The key assumption of sparsely distributed activations is enforced and used extensively. In this section we consider static world models; in section 3.1, we will use dynamic networks to implement inference by settling to the fixed points of the SWM.

2.1 Hierarchical Generative Visual-World Model. Images of objects seen in the world can be thought of as being created by a hierarchical, stochastic generative model (the GWM). Although it cannot be rigorously claimed that the real world uses such a model to generate images, the idea
Figure 1: Hierarchical generative visual-world model (GWM) for objects. At each layer $z_l$, the image can be represented by a large (possibly overcomplete) sparse vector. In this generative model, each layer is a binary random vector, which, given only the layer immediately above it in the hierarchy, is independent of other higher layers.

of the GWM is a useful fiction that guides the development of learning algorithms (Hinton & Ghahramani, 1997).

For the GWM, we assume a hierarchical binary state model of the form shown in Figure 1. The number of layers is somewhat arbitrary, though there should be enough layers to capture the structure of the data to be modeled, and four to five appears to be reasonable for images of objects (Riesenhuber & Poggio, 1999; Lee & Mumford, 2003; Hinton, Osindero, & Teh, 2006). The arrows in Figure 1 indicate that each layer, given the layer directly above it, is independent of higher layers. At the highest level, the vector $z_5$ is a sparse binary coding of the object in the image, and its value is drawn from the prior distribution $P(z_5)$. The representation of the particular orientation $z_4$ of an object depends on only the object representation $z_5$. The invariant, composite, and local features, $z_3$, $z_2$, and $z_1$, depend only on the layer immediately above them, for example, $P(z_3|z_4, z_5) = P(z_3|z_4)$, and the local features $z_1$ model the image $I$. The sequence can be summarized as

$$ P(z_5|z_4) P(z_4|z_3) P(z_3|z_2) P(z_2|z_1) P(I|z_1). \quad (2.1) $$
The joint distribution of the image and generative states $z_l$ is

$$P(I, z_1, z_2, z_3, z_4, z_5) = P(I | z_1) P(z_1 | z_2) P(z_2 | z_3) P(z_3 | z_4) P(z_4 | z_5) P(z_5),$$

(2.2)

where each layer $z_l$ is a binary vector of size $s_l$.

We postulate that the $z_l$ are sparse: they have very few nonzero components (Olshausen & Field, 1997). For example, in every image, only a few of all possible objects will be present, and each object will be in only one of its possible orientations, and so forth. Sparsity is proportional to the number of zero components in a vector $z \in \mathbb{R}^n$, \textit{sparsity} $\equiv \#(z_i = 0)/n$. A related quantity, diversity, is proportional to the number of nonzero components, \textit{diversity} $\equiv \#(z_i \neq 0)/n = 1 - \text{sparsity}$. Many studies have confirmed that natural images can be represented accurately by sparse vectors, corresponding to $z_1$ (Olshausen & Field, 1996; Kreutz-Delgado et al., 2003; Murray & Kreutz-Delgado, 2006). These studies have mainly dealt with small patches of images (on the order of $8 \times 8$ to $16 \times 16$ pixels), and it is clear that features larger than such patches will be represented nonoptimally. This further redundancy in larger-scale features can be reduced at higher levels, which can also have the property of sparseness.

### 2.1.1 Neighboring Layer Conditional Probability (NLCP)

For a middle layer $z_l$ given all the other layers, we find that $z_l$ conditioned on its immediate neighbors $z_{l-1}, z_{l+1}$ is independent of all the remaining layers—for example,

$$P(z_2 | I, z_1, z_3, z_4, z_5) = \frac{P(I | z_1) P(z_1 | z_2) P(z_2 | z_3) P(z_3 | z_4) P(z_4 | z_5) P(z_5)}{P(I | z_1) P(z_1 | z_3) P(z_3 | z_4) P(z_4 | z_5) P(z_5)} $$

$$= \frac{P(z_1 | z_2) P(z_2 | z_3)}{P(z_1 | z_3)} .$$

(2.3)

For an arbitrary layer, we can find the neighboring layer conditional probability (NLCP),

$$P(z_l | z_{l-1}, z_{l+1}) = \frac{P(z_{l-1} | z_l) P(z_l | z_{l+1})}{P(z_{l-1} | z_{l+1})} \quad \text{(NLCP)} .$$

(2.4)

This important independence assumption is equivalent to saying that each layer learns about the world only through its neighboring layers (Lee &
Mumford, 2003).\(^1\) Returning to the joint distribution and substituting in the NLCPs,

\[
P(I, z_1, z_2, z_3, z_4, z_5) = P(I|z_1) \cdot P(z_1|z_2)P(z_2|z_3) \cdot P(z_3|z_4)P(z_4|z_5) \cdot P(z_5)
= P(I|z_1) \cdot P(z_2|z_1, z_3)P(z_1|z_3) \cdot P(z_4|z_3, z_5)P(z_3|z_5) \cdot P(z_5).
\]

(2.5)

So the joint can be recovered given the NLCP and additional terms. Of course, other factorizations of the joint are possible, but these are also consistent with the NLCP for their respective layers (Brook, 1964).\(^2\)

2.1.2 Properties of Generative World Model (GWM). We now summarize the four properties of our generative world model (GWM).

1. There is a hierarchy of \(n\) hidden-layer vectors \(z_1, \ldots, z_n\) that model each image \(I\).

2. Each layer is independent of all higher layers given the neighboring layer above, \(P(z_l|z_{l+1}, \ldots, z_n) = P(z_l|z_{l+1})\).

3. Each layer is independent of all lower layers given the neighboring layer below, \(P(z_l|z_{l-1}, \ldots, z_1) = P(z_l|z_{l-1})\) (as shown in Murray, 2005, section 1.2).

4. Given its immediate neighboring layers, a layer \(z_l\) is independent of all other higher and lower layers, \(P(z_l|I, z_1, \ldots, z_n) = P(z_l|z_{l-1}, z_{l+1})\).

2.2 Types of Inference: Recognition, Imagination, Reconstruction, and Expectation-Driven Segmentation. For object recognition, the goal is to infer the highest layer representation \(z_n\) given an image \(I\). However, recognition is only one type of inference that might be required. Another type is running a model generatively using a high-level object representation to imagine an image of that object. In the brain, imagining a particular instance of an object will not correspond to the level of detail in the retinal representation, but there is evidence of activity in many of the lower visual areas (such as medial temporal, V1, and V2) during imagination (Kosslyn, Thompson, & Alpert, 1997).

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\(^1\) The NLCP is closely related to the Markov blanket, which is defined for a single node in a Bayesian network as that node’s parents, children, and children’s parents. The NLCP is defined over all the units in a given layer.

\(^2\) Brook (1964) proves that any system specified by the NLCP, \(P(z_j|z_i, i \neq j) = P(z_j|z_{j-1}, z_{j+1})\), has a joint distribution that can be factored as \(P(z_1, \ldots, z_n) = \prod_{i=1}^{n} Q_i(z_i, z_{i-1})\), which is the joint factorization of a simple Markov chain. This proof is for the case of scalar \(z\), but since our \(z_l\) are binary vectors, they can be equivalently represented as scalar integer variables \(\in\{1, \ldots, 2^{s_l}\}\). Thus, any system defined by the vector NLCP is consistent with a joint distribution that can be specified as the product of neighboring-layer factors, that is, the Markov assumption in equation 2.2.
Table 1: Types of Inference That Can Be Performed with the Hierarchical Generative World Model and the Types of Information Flow Required.

<table>
<thead>
<tr>
<th>Type of Inference</th>
<th>Inputs</th>
<th>Outputs</th>
<th>Requires</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recognition</td>
<td>$(I \rightarrow z_1)$</td>
<td>$z_n$</td>
<td>Bottom Up</td>
</tr>
<tr>
<td>Imagination</td>
<td>$z_n$</td>
<td>$(z_1 \rightarrow I)$</td>
<td>N</td>
</tr>
<tr>
<td>Reconstruction</td>
<td>$(I \rightarrow z_1)$</td>
<td>$(z_1 \rightarrow I)$</td>
<td>Y</td>
</tr>
<tr>
<td>Expectation-driven segmentation</td>
<td>$(I \rightarrow z_1)$</td>
<td>$z_n$</td>
<td>Y</td>
</tr>
<tr>
<td>Expectation-driven detection</td>
<td>$(I \rightarrow z_1)$</td>
<td>$z_n$</td>
<td>Y</td>
</tr>
</tbody>
</table>

Notes: We wish to find a good approximation to the layer $z_l$ of interest. The approximation used is the conditional expected value of $z_l$ under the variational approximation, $E_Q[z_l|z_{l-1}, z_{l+1}] = x_l$, as discussed in section 2.4.

Certain types of inference involve the use of top-down influences interacting with bottom-up inputs. For example, given a partially occluded image that has been recognized by higher layers, top-down influences can be used to reconstruct the hidden parts of the object (those features that are most likely given the input). Another type of inference is expectation-driven segmentation, where a prediction is presented at a higher level that may be used to explain cluttered, incomplete, or conflicting inputs at the lowest layer and the desired output is the segmented object at the first layer (Grossberg, 1976; Rao & Ballard, 1997; Hecht-Nielsen, 1998). The expectation input (higher-layer, top-down) must come from a source external to the visual system, which in the brain could be higher cortical areas or other senses and in computer vision could be dependent on the task or provided by a user. If we wish to find which objects are in a cluttered scene (i.e., the desired output is the highest-layer object representation) based on prior knowledge of what might be there (higher-layer input), we perform expectation-driven detection. If the high-level prediction about the scene is consistent with the input, the system converges with the expectation at the highest layer, and the prediction is confirmed. If the system converges to a different pattern, this indicates that the expected object is not present (which could be considered a state of surprise). Table 1 shows types of inference and the necessary information flow (top down or bottom up) needed in the model. As discussed below, we use a sparse-coding algorithm to transform the image into the first layer representation, $z_1$, and vice versa (denoted by $\rightarrow$ in the table).

2.3 Boltzmann-Like Distributions for Layer-Conditional Probabilities. Our next task is to postulate a form for the GWM distributions $P$ that is powerful enough to generate the images seen in the world. A common choice in probabilistic modeling is the Boltzmann distribution, $P(z) = \xi^{-1} \exp(-\beta\xi(z))$, where the probabilities are related to a function $\xi$ that
assigns an energy to each state, \( \zeta \) is a normalizing function, and \( \beta \) is a constant (which is a degree-of-randomness parameter related to temperature in physical systems, \( \beta \propto T^{-1} \); Hopfield, 1982; Hinton & Sejnowski, 1983; Hertz, Palmer, & Krogh, 1991). In thermodynamics and physical systems such as magnetic materials, the energy function captures the influence of each particle on its neighbors, where lower-energy states are more probable. The energy function usually has the form \( \xi(z) = -\frac{1}{2} \sum_{ij} w_{ij} z_i z_j \), where \( w_{ij} \) is the symmetric interaction weight (\( w_{ij} = w_{ji} \)) between \( z_i \) and \( z_j \). In the context of associative memories, the weights of the energy function are adjusted so that learned patterns form low-energy basins of attraction (e.g., using the Boltzmann machine learning rule; Ackley, Hinton, & Sejnowski, 1985).

The Boltzmann distribution requires the weights \( w_{ij} \) to be symmetric and have zero self-energy \( w_{ii} = 0 \) (Kappen & Spanjers, 2000). There are three main advantages of symmetric weights. First, a dynamic network with symmetric interactions is guaranteed to be asymptotically stable and settle to a fixed point (the zero-temperature solution), which minimizes the Boltzmann energy function (Mezard, Parisi, & Virasoro, 1987). Second, there is a procedure (Gibbs sampling with simulated annealing) that generates samples from this distribution at a given nonzero temperature \( T \). Finally, given a gradual enough annealing schedule for reducing \( T \), Gibbs sampling will track the global minimum-energy state (highest probability state) of the network and guarantee convergence to the zero-temperature solution as the temperature is lowered (Geman & Geman, 1984).

While the above properties are attractive and help explain the wide interest in the Boltzmann distribution and the Boltzmann machine, they may be of limited use in practice. It often takes considerable time for a stochastic network with symmetric weights to settle to an equilibrium state, possibly longer than a brain or artificial network has to make a decision (Welling & Teh, 2003), which accounts for the interest in simplifying approximations such as mean-field annealing (Peterson & Anderson, 1987). Furthermore, it has also been argued that the use of asymmetric weights can improve performance (such as by suppressing spurious memory states) and has greater biological plausibility (Parisi, 1986; Crisanti & Sompolinsky, 1988; Sompolinsky, 1988; Gutfrued, 1990; Apolloni, Bertoni, Campadelli, & de Falco, 1991; Kappen & Spanjers, 2000; Chengxiang, Dasgupta, & Singh, 2000). An additional motivation for admitting asymmetric weights is the notion that in hierarchical networks designed for invariant recognition, the relative strengths of the feedforward and feedback pathways will need to be different. Since neurons in higher layers tend to require inputs from multiple units to activate, the relative strength of the feedback connections to those units must be stronger than the feedforward weights to enable lower layer activity (i.e., generative ability). The primary deterrent to the use of asymmetric weights is the difficulty associated with ensuring asymptotic stability of the resulting algorithms, which involves the use of significantly more complex stability arguments (Apolloni et al., 1991).
We allow for asymmetric weights and sidestep the stability issue by working within a finite-time horizon framework. The resulting simplicity of the finite horizon problem relative to the infinite horizon problem is well known in the dynamical systems literature (Bertsekas, 1995). In appendix B we design a learning rule that encourages convergence to the desired state within a small number of time steps $\tau$. Also, the use of symmetric weights is merely sufficient for fixed points to exist; it is not a necessary condition.

We use the terms Boltzmann-like and energy-like to distinguish our model (with asymmetric weights) from stricter Boltzmann distribution assumptions. The Boltzmann-like form of the NLCP is

$$P_B(z_l|z_{l-1}, z_{l+1}) = \frac{1}{\xi(z_{l-1}, z_{l+1})} \exp (-\xi(z_l, z_{l-1}, z_{l+1})) \quad \text{(NLCP-B)},$$

(2.6)

where $\xi$ is the energy-like function and $\zeta$ is a normalizing function, with

$$\xi(z_l, z_{l-1}, z_{l+1}) = -z_l^T W_{l,l-1} z_{l-1} - z_l^T L_l z_l - z_l^T W_{l,l+1} z_{l+1} - \theta_l^T z_l$$

and

$$\zeta(z_{l-1}, z_{l+1}) = \sum_{z_l} \exp (-\xi(z_l, z_{l-1}, z_{l+1})).$$

(2.7)

where $W_{l,l+1}$ are top-down weights from layer $l+1$ to $l$, $W_{l,l-1}$ are the bottom-up weights from the layer $l-1$ to $l$, $L_l$ encodes the influence of units in layer $l$ on other units in that layer (lateral weights), and $\theta_l$ is a bias vector. The summation in $\zeta$ is over all states of layer $l$. Note that if the properties of symmetric weights are desired, they can be used without changes to the variational approximation developed in section 2.4.

An important question is whether the Boltzmann-like distribution, equations 2.6 and 2.7, is adequate to model the GWM model of Figure 1. It is possible to construct densities that are not well represented by any set of weights $W, L$ in equation 2.7. However, we do not need to model an arbitrary density, only densities that are sparse and therefore have more limited forms of dependence. Algorithms related to the Boltzmann machine have shown success on real-world vision tasks (Teh & Hinton, 2001; Hinton et al., 2006) and tend to confirm that the Boltzmann-like distribution is a reasonable starting point.

### 2.4 Simplified World Model Developed with a Variational Method.

The Boltzmann-like distribution, equations 2.6 and 2.7, provides a reasonable form of the probabilities in the GWM, which allows feedforward, feedback, and lateral influences. Unfortunately, exact inference on $z_l$ given $z_{l-1}, z_{l+1}$ is intractable for reasonably sized models even when the parameters of $P_B(z_l|z_{l-1}, z_{l+1})$ are known, because of the need to sum over every possible state $z_l$ in the normalizing function $\zeta$. In this section, we use a
variational method that approximates \( P_B(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}) \) with a factorial distribution, \( P_Q(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}) \). By *variational*, we mean that there are certain parameters \( \mathbf{x}_l = \{x_{l,i}\} \) that are varied to make the distribution \( P_Q \) as close to \( P_B \) as possible. The form of \( P_Q \) is taken to be a generalized factorial Bernoulli distribution,

\[
P_Q(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}) = \prod_{i=1}^{s_l} \left( \frac{x_{l,i} - a_4}{a_1} \right) \left( \frac{x_{l,i} - a_4}{a_1} \right) \left( 1 - \frac{x_{l,i} - a_4}{a_1} \right) \left( 1 - \frac{x_{l,i} - a_4}{a_1} \right), \tag{2.8}
\]

where \( x_{l,i} \) are the variational parameters and \( a = [a_1, a_2, a_3, a_4] \) are additional constant parameters (\( a_2 \) and \( a_3 \) will be introduced later) that are used to encourage sparsity-inducing densities (see section 2.5). The dependence on \( \mathbf{z}_{l-1}, \mathbf{z}_{l+1} \) will be introduced through \( x_{l,i} \) as derived below.

A sufficient condition for equation 2.8 to be a probability distribution is that \( \frac{x_{l,i} - a_4}{a_1} + (1 - \frac{x_{l,i} - a_4}{a_1}) = 1 \) and \( \frac{x_{l,i} - a_4}{a_1} \geq 0 \), which is true for \( a_1 > 0 \) and \( x_{l,i} \geq a_4 \). The slightly generalized Bernoulli distribution, equation 2.8, is based on a shift in the logical values of \( z_{l,i} \) in the energy function from \([0, 1]\) to \([a_4, a_1 + a_4]\) (the experiments below use \([-0.05, 1.05]\), which improves computational efficiency). Our formulation encompasses the two common choices for logical levels, \([0, 1]\) and \([-1, 1]\); for example, if logical levels of \([-1, 1]\) are needed, then \( a_4 = -1, a_1 = 2 \). Collecting the \( x_{l,i} \) into vectors \( \mathbf{x}_l \) of the same size as \( \mathbf{z}_l \) for each layer, it can be shown that \( \mathbf{x}_l \) are the conditional expected values for each layer,

\[
\mathbf{x}_l = E_Q[\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}]. \tag{2.9}
\]

Note that the variational parameter vector \( \mathbf{x}_l \) is a random variable which is the minimum mean-squared error (MMSE) estimate of \( \mathbf{z}_l \) given the values of its neighboring layers (Kay, 1993).

We now find the \( x_{l,i} \) that minimize the Kullback-Leibler divergence (Cover & Thomas, 1991) between the conditional probabilities \( P_B(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}) \) and \( P_Q(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}) \),

\[
KL(P_Q || P_B) = E_Q[\log P_Q(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1})] - E_Q[\log P_B(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1})]. \tag{2.10}
\]

where \( E_Q \) is the expected-value operator with respect to the distribution \( P_Q(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1}) \). When expected value \( E_Q[\mathbf{z}_{l,i}] = x_{l,i} \) is used, the first term is

\[
E_Q[\log P_Q(\mathbf{z}_l | \mathbf{z}_{l-1}, \mathbf{z}_{l+1})] = \sum_i \left[ \frac{x_{l,i} - a_4}{a_1} \log \left( \frac{x_{l,i} - a_4}{a_1} \right) \right. \nonumber \\
+ \left. \left( \frac{a_1 - x_{l,i} + a_4}{a_1} \right) \log \left( 1 - \frac{x_{l,i} - a_4}{a_1} \right) \right]. \tag{2.11}
\]
The second term in equation 2.10 can be expanded:

\[
E_Q[\log P_B(\mathbf{z}|\mathbf{z}_{l-1}, \mathbf{z}_{l+1})] = E_Q[-\log(\xi) - \xi(\mathbf{z}_l, \mathbf{z}_{l-1}, \mathbf{z}_{l+1})]
\]
\[
= E_Q[-\log(\xi) - \mathbf{z}_i^T \mathbf{W}_{l,l-1} \mathbf{z}_{l-1} - \mathbf{z}_i^T \mathbf{L}_l \mathbf{z}_l - \mathbf{z}_i^T \mathbf{W}_{l,l+1} \mathbf{z}_{l+1} - \theta_i^T \mathbf{z}_l].
\]

Again using the expected value \( E_Q[\mathbf{z}_{l,i}] = x_{l,i} \):

\[
E_Q[\log P_B(\mathbf{z}_i|\mathbf{z}_{l-1}, \mathbf{z}_{l+1})] = -\log(\xi) - \sum_{ik} W_{ik}^- z_{l-1,k} x_{l,i} - \sum_{ik} L_{ik} x_{l,k} x_{l,i}
\]
\[
- \sum_{ik} W_{ik}^+ z_{l+1,k} x_{l,i} - \sum_i \theta_{l,i} x_{l,i} + c_l,
\]

(2.13)

where \( W_{ik}^+, W_{ik}^- \), and \( L_{ik} \) are elements of the weight matrices \( \mathbf{W}_{l,l+1}, \mathbf{W}_{l,l-1}, \) and \( \mathbf{L}_l \), respectively and, defining \( \phi_l = (\mathbf{z}_l - x_l) \), the term \( c_l = E_Q[(\mathbf{z}_l - x_l)^T \mathbf{L}_l (\mathbf{z}_l - x_l)] = E_Q[\phi_l^T \mathbf{L}_l \phi_l] \), which is zero assuming that \( L_{ii} = 0.3 \)

2.4.1 Self-Consistency Conditions of the Variational Approximation. The variational parameters \( x_{l,i} \) that minimize the distance between \( P_B \) and \( P_Q \), equation 2.10 are found by solving,

\[
\frac{\partial \text{KL}(P_Q||P_B)}{\partial x_{l,i}} = 0 = a_2 \left( \sum_k W_{ik}^- z_{l-1,k} + \sum_k L_{ik} x_{l,k} + \sum_k W_{ik}^+ z_{l+1,k} \right)
\]
\[+ \log \left( \frac{z_{l,i} + a_4}{z_{l,i} - a_4} \right) - a_3,
\]

(2.14)

using a constant term \( a_3 \) for the bias \( \theta_{l,i} \), and factoring out \( a_2 \) from \( W^+, W^- \) and \( L \) (with a slight abuse of notation, including factoring \( \frac{1}{a_1} \) into \( a_2, a_3 \); see equation 2.11). Setting equation 2.14 equal to zero and solving for \( x_{l,i} \),

\[
x_{l,i} = f(v_{l,i})
\]
\[
v_{l,i} = \sum_k W_{ik}^- z_{l-1,k} + \sum_k L_{ik} x_{l,k} + \sum_k W_{ik}^+ z_{l+1,k},
\]

(2.15)

---

3 The term \( c_l = E_Q[(\mathbf{z}_l - x_l)^T \mathbf{L}_l (\mathbf{z}_l - x_l)] = \text{Tr}[\mathbf{L}_l \Sigma_{z_l}] \), where \( \Sigma_{z_l} \) is the covariance matrix of \( \mathbf{z}_l \) under \( P_Q \). Since \( \mathbf{z}_l \) is assumed conditionally independent under \( P_Q \), the non-diagonal elements of the covariance matrix are zero. We will disallow self-feedback (i.e., \( L_{ii} = 0 \)), so that \( \text{Tr}[\mathbf{L}_l \Sigma_{z_l}] \) is zero. However, it is straightforward to handle the case when \( L_{ii} \neq 0 \) given the factorial form of \( P_Q \).

4 For simplicity, we set \( \theta_{l,i} = a_3 \) for all \( l, i \). However, this assumption can be relaxed.
where \( f(\cdot) \) is a sigmoid activation function parameterized by \( a = [a_1, a_2, a_3, a_4] \).

\[
f(v) = \frac{a_1}{1 + \exp(-a_2 v + a_3)} + a_4.
\]  

(2.16)

Defining \( \phi_{l,i} = z_{l,i} - x_{l,i} \) to be the approximation error, the state \( z \) is equal to \( x \) plus a random noise component, \( z_{l,i} = x_{l,i} + \phi_{l,i} \). This yields

\[
v_{l,i} = \sum_k W_{ik}^- (x_{l-1,k} + \phi_{l-1,k}) + \sum_k L_{ik} x_{l,k} + \sum_k W_{ik}^+ (x_{l+1,k} + \phi_{l+1,k})
\]

\[
= \sum_k W_{ik}^- x_{l-1,k} + \sum_k L_{ik} x_{l,k} + \sum_k W_{ik}^+ x_{l+1,k} + \varepsilon_{l,i},
\]  

(2.17)

where the \( \phi \) terms have been collected into \( \varepsilon_{l,i} \). By collecting all the terms for each layer into a vector, we obtain the single equation,

\[
\begin{bmatrix}
x_1 \\
x_2 \\
x_3 \\
\vdots \\
x_n
\end{bmatrix} = f
\begin{bmatrix}
L_1 & W_{12} & 0 & \cdots & 0 \\
W_{21} & L_2 & W_{23} & \vdots & \vdots \\
0 & W_{32} & L_3 & W_{34} & \vdots \\
\vdots & \vdots & \vdots & \ddots & \vdots \\
0 & \cdots & 0 & W_{n,n-1} & L_n
\end{bmatrix}
\begin{bmatrix}
x_1 \\
x_2 \\
x_3 \\
\vdots \\
x_n
\end{bmatrix} +
\begin{bmatrix}
\varepsilon_1 \\
\varepsilon_2 \\
\varepsilon_3 \\
\vdots \\
\varepsilon_n
\end{bmatrix},
\]  

(2.18)

which is the self-consistency condition for the variational approximation. The self-consistency condition, equation 2.18, is a necessary condition for the factorial NLCP (see equation 2.8) of the SWM to have been optimally fit to the GWM NLCP of equation 2.6.

2.4.2 Simplified World Model Forms. Further collecting all the estimates for each layer into a single vector \( X = [x_1^T, \ldots, x_n^T]^T \) and all the weights into a global weight matrix \( \mathbb{W} \), equation 2.18, can be written concisely:

\[
X = f(\mathbb{W} X + \varepsilon) \quad \text{(SWM-E)},
\]  

(2.19)

which is called the simplified world model on the expected values (SWM-E) and where the vector forms of the errors are

\[
\Phi = Z - X
\]

\[
\varepsilon = (\mathbb{W} - \mathbb{L})\Phi.
\]  

(2.20)
The error $\epsilon$ is generally unobservable, and later we will have to make approximations to perform inference and learn the weights $\mathbb{W}$. In particular, for inference and learning, we will neglect $\epsilon$ and use a certainty equivalence approximation (see section 4). The SWM can be written equivalently in terms of the binary state $Z$,

$$Z = f(\mathbb{W}Z - L\Phi) + \Phi \quad \text{(SWM-B),}$$

which is called the SWM on the binary state (SWM-B). The SWM can also be written in an equivalent dual form on the preactivation state. Collecting the $v_{l,i}$ into a state vector $V \equiv \mathbb{W}X + \epsilon$ (and $X = f(V)$), we have

$$V = \mathbb{W}f(V) + \epsilon \quad \text{(SWM-P),}$$

which is called the SWM on the preactivation state (SWM-P). Equations 2.19, 2.21, and 2.22 are self-consistency conditions for the SWM. We will return to these key results in section 3.1, where we discuss how to find solutions to these conditions through evolution of updates in time. Note that with a slight abuse of notation, we refer to the self-consistency conditions themselves as the SWMs.

2.4.3 Relation to Other Variational Methods. Our approach is based on using a neighboring-layer conditional probability model matched to the hierarchical NLCP GWM using a factorial variational approximation. The use of a factorial variational approximation is known as a mean-field (MF) approximation in the statistical physics community (Peterson & Anderson, 1987), where the probabilities to be approximated are either unconditional (as typically done in statistical physics) or conditioned only on visible layers (as in the Boltzmann machine). A distinguishing feature of our method is that the Boltzmann-like NLCP admits the use of an approximating distribution $P_Q$ that is factorial when conditioned on its neighboring layers, and the resulting approximation is not factorial when conditioned only on the visible layers. This modeling assumption removes less randomness (and allows more generative capability) than the MF approximation conditioned on the visible layers (as in the deterministic Boltzmann machine; Galland, 1993). This is a richer model than the deterministic Boltzmann machine, as our conditional expectations, $x_l = E_q[z_l|z_{l-1}, z_{l+1}]$, retain more randomness than the nonhidden expectations $x_l = E_q[z_l|\text{visible layers}]$ of the deterministic Boltzmann machine. Our factorial approximation is reasonable, as it is equivalent to saying that the meaningful information about the world contained in any layer is provided by its immediate neighboring layers, so if we condition on neighboring layers, then only random (“meaningless”) noise remains. The factorial Bernoulli distribution is one of the most tractable and commonly used variational approximations (Jordan,
Figure 2: (A) Activation function $f(v)$ with parameters $a = [1.1, 2.0, 4.0, -0.05]$ (see equation 2.16). The limits of the activation function are $[a_4, a_1 + a_4] = [-.05, 1.05]$, and the slope is set by $a_2$ and the bias by $a_3$. The shape of the activation function encourages sparsity by ensuring that small input activities $v < v_l$ do not produce any positive output activity. In the simulations, the values of $x = f(v)$ are thresholded so that $x = [f(v)] \in [0, 1]$; however, the values of $f'(v)$ are kept for use in the weight updates (see appendix B). (B) The probability density $P(x; a)$ of a normal random variable ($\mu = 0$, $\sigma^2 = 1$) after being transformed by the activation function, $f(v)$ in equation 2.16, is a sparsity-inducing density if the parameters $a$ are chosen properly. The parameters used are $a = [1.1, 2.0, 4.0, -0.05]$. (C) Probability $P(x; a')$ is not sparsity inducing with the standard set of parameters for sigmoid activation functions, $a' = [1, 1, 0, 0]$.

Ghahramani, Jaakkola, & Saul, 1998); however, if more accurate approximations are desired, other distributions may be used, such as the second-order methods described by Welling & Teh (2003) and Kappen and Spanjers (2000), although at higher computational cost.

2.5 Activation Functions Can Encourage Sparse Distributions. The parameterized sigmoid activation function, equation 2.16, can be used to encourage sparse activation by appropriate choice of parameters $a$. Figure 2 shows the activation function, equation 2.16, when parameterized with $a = [1.1, 2.0, 4.0, -0.05]$, which was chosen so that small levels of activation do not lead to positive values of $f(v)$. Parameters $a_2, a_3$ can be viewed as prior constraints on the network weights. Theoretically these weight scaling and bias terms could be learned, but from a practical standpoint, our networks are quite large, and the critical property of sparsity that makes learning tractable must be enforced from the early epochs or too many extra weights would be updated.

We can reasonably assume that $v = v_{il}$ as given by equation 2.15 is a normally distributed random variable due to the central limit theorem (Johnson, 2004) because $v$ is the sum of (nearly) independent and identically distributed values with bounded variance.\(^5\) The density $P(x; a)$ can

\(^5\) There will be dependence in $v$ between units that all represent the same feature or object; however, since all network layers are constrained to be sparse, these dependencies
then be found by transforming the normal density \( P(v) = \mathcal{N}(\mu, \sigma^2) \) by the activation function 2.16 (see equation 1.2.26 of Murray, 2005). For the values of \( a \) given above and for \( \mu = 0, \sigma^2 = 1 \), Figure 2B shows that \( P(x; a) \) is indeed a sharply peaked sparsity-inducing distribution. In contrast, Figure 2C shows \( P(x; a') \) after being transformed by the sigmoid activation function with parameters \( a' = [1, 1, 0, 0] \), which does not lead to a sparsity-inducing distribution. The choice of parameters \( \mu, \sigma^2 \) is also important for the transformed distributions to be sparse. For example, if \( \mu = 0 \), the variance must be less than about 2.0, or the resulting density will be bimodal. However, with the proper choice of initial conditions, we are able to ensure these conditions are met (see section 5).

3 Recurrent Dynamic Network

Recognizing that solutions to important inferencing problems correspond to solutions of the self-consistency conditions derived in section 2.4, we generalize these conditions into a dynamic network capable of converging to a solution satisfying equation 2.18 in order to estimate the states \( x_l \). We introduce a time index \( t \) for the iterations of this dynamic network, while our goal remains to estimate the state of the static SWM.

There are \( n \) layers in the network, and the vector of activations for the \( l \)th layer at time \( t \) is denoted \( x_{l,t}, l = 1, \ldots, n \), with layer sizes \( s = [s_1, \ldots, s_n] \). The network is designed to enforce rapid convergence to the self-consistency conditions (see equation 2.18) for \( x_l \), such that \( x_{l,t} \to x_l \). The state vector of all the layers at time \( t \) is denoted

\[
X_t = [x_1^T, x_2^T, \ldots, x_n^T]^T \in \mathbb{R}^N,
\]

where \( N \) is the size of the state vector (dropping the time index on \( x_l \) inside the vector for clarity). The activity in all layers \( x_l \), is enforced to be sparse, and the number of nonzero elements of the layers is denoted \( r_t = [r_1 \ldots r_n] \).

Figure 3 shows the four-layer network structure used for the experiments in this letter. Dotted lines indicate inputs and connections that are not used here but are allowed in the model.

The layers used for input and output depend on the type of inference required. In this work, inputs are usually injected at either the highest...
Figure 3: Dynamic network used in the experiments. Inputs images $I$ are first sparsely coded using the FOCUSS+ algorithm, which operates on nonoverlapping patches of the input image (see appendix A). This sparse overcomplete code $u_1$ is used as bottom-up input to the four-layer hierarchical network. Dotted lines indicate inputs ($u_3$) and connections ($L_i$) that are not used in the experiments in this letter but are allowed by the network.

or lowest layer (although in general, we may have inputs at any layer if additional types of inference are required). We define an input vector $U^X_t$ (again dropping the time index inside the vector),

$$U^X_t = [u_1^T, u_2^T, \ldots, u_n^T]^T,$$  \hspace{1cm} (3.2)

where $u_1$ is a sparsely coded input image (see appendix A) and $u_n$ is an $m$-out-of-$n$ binary code called the object code, which represents the classification of the object. The advantage of using an $m$-out-of-$n$ object code is that it allows more objects to be represented than the size $n$ of the highest layer, which is the limitation of 1-out-of-$n$ codes. The object code provides a high
representational capacity and robustness to the failure of any individual unit or neuron, both of which are desirable from a biological perspective. In addition, we can represent new objects without adjusting the size of the highest layer, \( u_n \), by creating new random object codes.

For recognition and reconstruction, the input \( u_1 \) is the coded image, and the object code input is zero, \( u_n = 0 \). When the network is used for imagination, the input is the object code presented at the highest layer \( u_n \) and random noise at \( u_2 \), and the output is the reconstructed image at the lowest layer. For expectation-driven segmentation, both \( u_1 \) and \( u_n \) inputs are used. Table 1 shows the layers used for input and output for each type of inference.

### 3.1 Dynamic Network Form.

The recurrent dynamic network (DN-E) is the time-dependent generalization of the self-consistency conditions 2.18 of the SWM-E given by

\[
\begin{bmatrix}
  x_1 \\
  x_2 \\
  x_3 \\
  \vdots \\
  x_n
\end{bmatrix}_{t+1} = \begin{bmatrix}
  L_1 & W_{12} & 0 & \cdots & 0 \\
  W_{21} & L_2 & W_{23} & \cdots & \vdots \\
  0 & W_{32} & L_3 & W_{34} & \cdots \\
  \vdots & \vdots & \ddots & \ddots & \vdots \\
  0 & \cdots & 0 & W_{n,n-1} & L_n
\end{bmatrix}
\begin{bmatrix}
  x_1 \\
  x_2 \\
  x_3 \\
  \vdots \\
  x_n
\end{bmatrix}_t + \begin{bmatrix}
  \varepsilon_1 \\
  \varepsilon_2 \\
  \varepsilon_3 \\
  \vdots \\
  \varepsilon_n
\end{bmatrix}_t + \begin{bmatrix}
  u_1 \\
  u_2 \\
  u_3 \\
  \vdots \\
  u_n
\end{bmatrix}_t
\]

(3.3)

which can be written in the compact form,

\[
X_{t+1} = f(WX_t + \varepsilon_t) + U_t^X (\text{DN-E}),
\]

(3.4)

where \( U_t^X \) is the input to the network, which can include a sparsely coded input image \( u_1 \) or a top-down \( u_n \) consisting of an object code, or both.

Our goal will be to learn a \( W \) such that network 3.4 will rapidly converge to a steady state, given transient or constant inputs \( U^X_t \). We will attempt to enforce the steady-state self-consistency behavior at finite time horizon \( t = \tau \), where the horizon \( \tau \) is a design parameter chosen large enough to ensure that information flows from top to bottom and bottom to top and small enough for rapid convergence. Because of the block structure of \( W \), information can pass only to adjacent layers during one time step \( t \). (We use the terms time step and iteration interchangeably.) For example, in a four-layer network, it takes only four time steps to propagate information from the highest to the lowest layer, while the network may require more iterations to converge to an accurate estimate. A relatively small number of iterations, on the order of 8 to 15, will be shown to work well.
Table 2: Progression of Models Developed in Sections 2 and 3.

<table>
<thead>
<tr>
<th>Model Type</th>
<th>Equation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hierarchical generative world model (GWM)</td>
<td>$P(z_l</td>
<td>z_{l-1}, z_{l+1})$</td>
</tr>
<tr>
<td>Simplified world model (SWM) (self-consistency conditions)</td>
<td>$X = f(WX + \varepsilon)$</td>
<td>Variational approximation $E_\theta[Z] = X$ leads to: (SWM-E, equation 2.19)</td>
</tr>
<tr>
<td></td>
<td>$Z = f(WZ - L\Phi) + \Phi$</td>
<td>Binary state: (SWM-B, equation 2.21)</td>
</tr>
<tr>
<td></td>
<td>$V = WF(V) + \varepsilon$</td>
<td>Equivalent preactivation state: (SWM-P, equation 2.22)</td>
</tr>
<tr>
<td>Dynamic network (DN) (discrete time)</td>
<td>$X_{t+1} = f(WX_t + \varepsilon_t) + U_{t+1}^X$</td>
<td>State update: (DN-E, equation 3.4)</td>
</tr>
<tr>
<td></td>
<td>$V_{t+1} = WF(V_t) + \varepsilon_{t+1} + U_{t+1}^V$</td>
<td>Preactivation state update: (DN-P, equation 3.5)</td>
</tr>
</tbody>
</table>

### 3.2 Preactivation State-Space Model.

In the previous section we created a dynamic network on the state vector $X_t$ based on the SWM-E. By defining an equivalent model on the preactivation vector $V_t$, we create another dynamic network, which will be used in deriving the learning algorithm (see section 4). Generalizing the preactivation model (SWM-P, equation 2.22) to a dynamic network,

$$V_{t+1} = WF(V_t) + \varepsilon_{t+1} + U_{t+1}^V$$ (DN-P),

where $V_t$ is assumed to be a gaussian vector, as discussed in section 2.5, and $U_{t+1}^V$ is the input or initial conditions for the preactivation state (compare with $U_{t+1}^X$ for the state $X$). The DN-E and DN-P are equivalent representations of a dynamic generative world model. Interpreting the layers of $V_t$ as the hidden states of the generative visual-world model, the visible world is found with the read-out map,

$$Y_t = C g(V_t) + \text{noise},$$ (3.6)

where $g(\cdot)$ is the output nonlinearity, and $C = [1, 0, \ldots, 0]$ hides the internal states. Table 2 summarizes the moves made from the generative world model of section 2 to the dynamic networks of this section.
4 Finding a Cost Function for Learning the Weights $\mathbb{W}$

The dynamic networks of the previous section can perform visual inference by being forced to approximate the self-consistency conditions of the simplified world model (SWM). This can be done assuming that the weights $\mathbb{W}$ are known. Now we turn to the problem of learning these weights given a set of training data. In this section, we proceed in a Bayesian framework assuming $\mathbb{W}$ is a random variable and derive a cost function after suitable approximations. The labeled training set is denoted $\mathcal{Y} = \{ \hat{Y}^{(1)}, \ldots, \hat{Y}^{(K)} \}$, where the $k$th element $\hat{Y}^{(k)}$ is a vector with a sparse coding of image $k$ at its first layer $\hat{y}^{(k)}_1$ and the corresponding object code $\hat{y}^{(k)}_n$ at the highest layer and zero vectors at the other layers,

$$
\hat{Y}^{(k)} = \left[ \hat{y}^T_1 \ 0 \ldots \ 0 \ \hat{y}^T_n \right]^T,
$$

where the superscript index of the pattern $k$ for each layer (i.e., $\hat{y}^{(k)}_n$) has been omitted for clarity.

The cost function for $\mathbb{W}$ is derived using the DN-P dynamics on the preactivation state $V_t$, equation 3.5. During training, for each pattern $k$, we create an input time series $U^X_t$ from the data set as follows: $U^X_t = \hat{Y}^{(k)}$ for $t = 1, 2, 3$ and $U^X_t = 0$ for $4 \leq t \leq \tau$. This choice of $U^X_t$ starts the dynamic network in the desired basin of attraction for the training pattern $\hat{Y}^{(k)} (U^X_t = \hat{Y}^{(k)}$ for $t = 1, 2, 3)$. The network is then allowed to iterate without input ($U^X_t = 0$ for $4 \leq t \leq \tau$), which with untrained weights $\mathbb{W}$ will in general not converge to the same basin of attraction. The learning process attempts to update the weights so that the training inputs are basins of attraction, and to create middle-layer states consistent with that input. The set of inputs for pattern $k$ for all the time steps is denoted $U^{(k)} = \{ U^{(k)}_1, \ldots, U^{(k)}_\tau \}$, and for the entire data set we have $U = \{ U^{(1)}, \ldots, U^{(K)} \}$. Similarly, for each pattern in the preactivation state, we have $V^{(k)} = \{ V^{(k)}_1, \ldots, V^{(k)}_\tau \}$, and for the whole data set, $V = \{ V^{(1)}, \ldots, V^{(K)} \}$.

Assuming that the weights $\mathbb{W}$ are random variables, their posterior distribution is found by Bayes’ rule,

$$
P(\mathbb{W}|V; U) = \frac{P(V|\mathbb{W}; U)P(\mathbb{W})}{P(V; U)}. \quad (4.2)
$$

Our goal is to find the weights $\mathbb{W}$ that are most likely given the data and the generative model, and we use the maximum a posteriori (MAP)
estimate,

\[ \mathcal{W} = \arg \max_{\mathcal{W}} P(\mathcal{W}|\mathcal{V}; U) \]

\[ = \arg \min_{\mathcal{W}} -\ln P(\mathcal{V}|\mathcal{W}; U) - \ln P(\mathcal{W}), \quad (4.3) \]

due to the denominator in equation 4.2 not depending on \( \mathcal{W} \). Correct assumptions about \( \mathcal{W} \) are important for successful learning, which requires some form of constraint, such as prior normalization to use all of the network’s capacity (see section 5). Assuming patterns in the training set are independent, \( P(\mathcal{V}|\mathcal{W}; U) = \prod_k P(\mathcal{V}(k)|\mathcal{W}; U^{(k)}) \),

\[ \mathcal{W} = \arg \min_{\mathcal{W}} \left[ -\sum_k \ln P(\mathcal{V}(k)|\mathcal{W}; U^{(k)}) - \ln P(\mathcal{W}) \right]. \quad (4.4) \]

Note that the dynamic system 3.5 is Markovian under our assumption that \( \varepsilon_t \) are independent (Bertsekas, 1995). Then the probability of the sequence of time steps can be factored (omitting the pattern index \( k \) on the \( V_t \) for clarity),

\[ P(\mathcal{V}(k)|\mathcal{W}; U^{(k)}) = P(V_r, V_{r-1}, V_{r-2}, \ldots, V_1|\mathcal{W}; U^{(k)}) \]

\[ = \prod_{t=1}^{r} P(V_t|V_{t-1}, \mathcal{W}; U^{(k)}), \quad (4.5) \]

from the chain rule of probabilities. The preactivation state at each time \( V_t \) can be expressed in terms of each layer \( v_{l,t} \),

\[ P(\mathcal{V}_{l}^{(k)}|\mathcal{V}_{l-1}^{(k)}, \mathcal{W}; U^{(k)}) = \prod_{l=1}^{n} P(v_{l,t}^{(k)}|V_{l-1}^{(k)}, \mathcal{W}; U^{(k)}), \quad (4.6) \]

if we assume that the layers are conditionally independent of each other at \( t \) given the state at the previous time \( V_{l-1} \). Combining equations 4.4, 4.5, and 4.6,

\[ \mathcal{W} = \arg \min_{\mathcal{W}} \left[ -\sum_k \sum_{t=1}^{r} \sum_{l=1}^{n} \ln P(v_{l,t}^{(k)}|V_{l-1}^{(k)}, \mathcal{W}; U^{(k)}) - \ln P(\mathcal{W}) \right]. \quad (4.7) \]

Since \( v_{l,t} \) is approximately normal (see section 2.5), for those layers where and when we have target values of \( y_{l,t} \) from the data set and corresponding
target states for \( v_{l,t} \), the probability of the layer is

\[
P_{\text{targ}}(v_{l,t} | V_{t-1}, \mathbb{W}, U) = \frac{1}{(2\pi \sigma_v^2)^{\frac{N}{2}}} \exp \left( -\frac{1}{2\sigma_v^2} e_{l,t}^T e_{l,t} \right),
\]

(4.8)

where \( \sigma_v^2 \) is the variance of each component (which is assumed identical).

At other layers and times, the state probabilities \( v_{l,t} \) are approximately gaussian, but we do not have a desired state, and so we enforce sparsity in these cases. We model the distributions at these layers by independent gaussians with fixed mean \( \mu \) and variance \( \sigma_s^2 \),

\[
P_{\text{spar}}(v_{l,t} | V_{t-1}, \mathbb{W}; U) = \frac{1}{(2\pi \sigma_s^2)^{\frac{N}{2}}} \exp \left( -\frac{1}{2\sigma_s^2} \|v_{l,t} - \mu\|^2 \right),
\]

(4.9)

where \( \mu = [\mu \ldots \mu]^T \) with \( \mu \) a design parameter of the appropriate size.

Introducing an indicator variable \( \beta \) that selects between \( P_{\text{spar}} \) and \( P_{\text{targ}} \), we define \( \beta_{l,t} = 1 \) if we have target values for layer \( l \) at \( t \) and \( \beta_{l,t} = 0 \) otherwise. The probability of each layer becomes

\[
P(v_{l,t} | V_{t-1}, \mathbb{W}; U) = \beta_{l,t} P_{\text{targ}}(\cdot) + (1 - \beta_{l,t}) P_{\text{spar}}(\cdot).
\]

(4.10)

Substituting equation 4.10 in equation 4.7 yields

\[
\mathbb{W} = \arg \min_{\mathbb{W}} \left\{ \sum_k \sum_{t=1}^T \left[ e_{l,t}^T (e_{l,t} \odot \beta_t) + \lambda (V_{l} - \mu)^T [(V_{l} - \mu) \odot (1 - \beta_t)] \right] \right. \\
- \ln P(\mathbb{W}) \right\},
\]

(4.11)

where \( \beta_t \in \mathbb{R}^N \) is the indicator vector for all elements of \( V_{l} \), \( \odot \) is the element-wise Hadamard vector product and the constant terms depending on \( \sigma_v^2, \sigma_s^2 \) have been combined into a new constant \( \lambda \) (again omitting the \( k \) inside the summation).

Several things should be noted about this formulation. First, the objective function is derived in relation to the preactivation vector \( V_l \) instead of the postactivation vector \( X_l \). This is done to use the gaussian form of equation 4.8 and is reminiscent of the technique in the generalized linear model literature of working with the linear structure vector of a nonlinear model (Gill, 2001). Second, the cost function, equation 4.11, is similar in form to those used in overcomplete coding algorithms, which are unsupervised, and are designed to minimize the reconstruction error using as sparse a code as possible (Olshausen & Field, 1997; Kreutz-Delgado et al., 2003).

\footnote{We assume that noise = 0 in equation 3.6 and that given \( Y_l \), we can solve for a corresponding value of \( V_l \).}
The cost function for $\mathbb{W}$, equation 4.11, is a function of the true state $V_t$ and the error $\varepsilon_t$, which we generally do not have access to. In practice, we will resolve this problem by generating estimates of the unknown $V_t$ using a current estimate of the weights from the dynamic network (DN-P) under the certainty equivalence approximation that $\varepsilon_t = 0$ (Bertsekas, 1995). Certainty equivalence is a standard technique in optimization when certain variables are random. For example, an unknown random variable can be replaced by its mean before optimizing the cost function. In our case, we estimate the unknown random $V_t$ by the dynamic network’s output $\hat{V_t}$, which is then used to find the $\mathbb{W}$ that minimizes the cost function 4.11. For each pattern in the data set, we run DN-P (see equation 3.5) using the input sequence $U^Y_t = v_h U^X_t$, where $v_h = f^{-1}(1.0)$ (see Figure 2).\(^8\) Running the network with certainty equivalence gives estimated states

$$\hat{V}_t = \mathbb{W} f(\hat{V}_{t-1}) + U^Y_t. \quad (4.12)$$

The errors $\hat{\varepsilon}_t$ used for learning are then the difference between $\hat{V}_t$ and the desired target states found from the data set,

$$\hat{\varepsilon}_t = (\hat{V}_t - v_h \hat{Y}^{(k)}) \odot \beta_t, \quad (4.13)$$

where layers with no target values are set to 0 due to the effect of $\beta_t$.

Using the cost function in equation 4.11, we find a learning algorithm for weights $\mathbb{W}$ (see appendix B) which is closely related to the backpropagation-through-time algorithm (BPTT) for training recurrent networks (Williams & Peng, 1990). The main drawback of the BPTT algorithm is that it is computationally inefficient due to the unrolling of the network for each time step. Our approach overcomes this drawback by using a small number of time steps $\tau$ and taking advantage of the sparsity of every layer to update only weights between units with some nonzero activity.

5 Algorithm Implementation

This section summarizes the implementation details of the dynamic network and learning algorithm as used in the experiments.

5.1 Preparing the Data Set. The data set consists of $K$ images representing $M$ unique objects, where in general we have many different views or transformations of each object, so $K > M$. For each object $m$, we generate a sparse object code $c^{(m)} \in \mathbb{R}^{s_n}$ (the size of the highest layer) with $r_n$ randomly selected nonzero elements, which is used as the desired value of the highest

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\(^8\) This is an approximation to $U^Y_t = f^{-1}(U^X_t)$ when $U^X_t$ is binary, assuming elements $U^Y_{j,t} = 0$ when $U^X_{j,t} = 0$. 

layer. Each image \( k \) is preprocessed and converted into a sparse code (see section 6), which is used as the first layer input, \( y_1 \). The data set of all images is \( \tilde{Y} = \{\tilde{Y}^{(1)}, \ldots, \tilde{Y}^{(K)}\} \) where each pattern is

\[
\tilde{Y}^{(k)} = \begin{bmatrix} \tilde{y}_1^T & 0 & \ldots & 0 \end{bmatrix}^T.
\]

and the highest layer is the object code, \( \tilde{y}_n^T = c^{(m)} \).

5.2 Network Initialization. The network weights are initialized with small, random values uniformly distributed within certain ranges. The initial weight ranges are feedforward and feedback weights \( W \in [-0.01, 0.01] \) and lateral weights \( L \in [-0.001, 0.000] \) (which enables only lateral inhibition, not excitation). Self-feedback is not allowed, \( L_{ii} = 0 \), and lateral weights are not used in layer 1 for computational efficiency. Feedback weights are initialized to be the transpose of the corresponding feedforward weights, \( W_{lm} = W_{ml}^T \), but are not restricted to stay symmetric during training.

5.3 Performing Inference Given Known Weights \( W \). To run the network for the experiments, we create an input time series \( U_t^X \) from the images and object codes in the data set \( \tilde{Y} \). The input can include \( \tilde{y}_1 \) or \( \tilde{y}_n \) (or both) as determined by the type of inference desired (see Table 1). For example, when the network is run for recognition, the inputs for the first few time steps are the coded image \( \tilde{y}_1 \), so that \( (U_t^X)^T = [\tilde{y}_1^T, 0, \ldots, 0]^T, t = 1, 2, 3 \), and \( U_t^X = 0, t \geq 4 \). When the network is run generatively, the object code is used as input, such that \( (U_t^X)^T = [0, \ldots, \tilde{y}_n^T]^T, t = 1, \ldots, \tau \), and the network is then run for \( \tau \) steps, after which the first layer contains a representation of an imagined image.

Given a sequence of inputs \( U_t^X \), the network is run in certainty-equivalence mode (no added noise) for a fixed number of discrete time steps, \( 0 \leq t \leq \tau \), with \( \tau \) being 8 to 15 for the experiments below. With an initial state \( \tilde{X}_0 = 0 \), the network is run using

\[
\tilde{V}_t = \mathcal{W} \tilde{X}_t
\]

\[
\tilde{X}_t = f(\tilde{V}_{t-1}) + U_t^X \quad 1 \leq t \leq \tau.
\]

The state \( \tilde{X}_t \) is further restricted to be in the unit cube, \( \tilde{X}_t \in [0, 1]^N \). To improve computational efficiency, only a limited number of nonzero elements are allowed in each layer, \( \bar{r} = [\bar{r}_1, \ldots, \bar{r}_n] \), which is enforced on \( V_t \) at each layer by allowing only the largest \( \bar{r}_i \) of them to remain nonzero.

5.4 Learning the Weights \( \mathcal{W} \). Training proceeds in an online epoch-wise fashion. In each epoch, a subset of patterns is chosen from \( \tilde{Y} \), and inputs are created with the coded image in the first layer for the first three time steps, so
that \( U_t^X = [\mathbf{y}_1^T, 0, 0, \mathbf{y}_n^T] \), \( t = 1, 2, 3 \), and \( U_t^X = 0 \), \( t \geq 4 \). Input patterns must be removed at some point during training, because otherwise, there would be no error gradient to enforce learning of reconstruction. Presenting the input for three time steps was found to give better performance than other lengths of input (see section 6.1).

The state \( \hat{X}_t \) and preactivation state \( \hat{V}_t \) from running the network (see equation 5.2) are saved for each \( t \leq \tau \). The error vector for weight updates is \( \hat{\epsilon}_t = (\hat{V}_t - v_h \hat{Y}^{(k)}) \odot \beta_t \) (see equation 4.13). Weight updates \( \Delta w_{ji} \) are given by equation B.14. In standard gradient descent, weight updates will naturally become small when errors are small. However, since we use an additional sparsity enforcing term, even if both the highest and lowest layer errors are zero, weight updates will still occur in order to sparsify middle layers. Training stops after a certain number of epochs are completed.

For computational efficiency when learning sparse patterns, only a small set of weights \( w_{ji} \) is updated for each pattern. During our simulations, \( \hat{X}_t \) is found by thresholding the activation function output \( f(\hat{V}_{t-1}) \) to \([0, 1]\), resulting in a sparse \( \hat{X}_t \) given certain conditions (see section 2.5). Weights are then updated between units only when the source unit \( \hat{X}_i, t \) is active and either the target unit \( \hat{X}_j, t \) is active or has nonzero error \( \hat{\epsilon}_j, t \). During the initial epochs of learning, there must be enough weight strength to cause activation throughout the middle layers. As learning progresses, the activity is reduced through the sparseness-enforcing term.

### 5.5 Testing for Classification

To classify an input image once the network has settled into a stable state, the last layer’s activation \( x_n \) is compared with the object codes \( c^{(m)} \) to find the class estimate,

\[
\text{Class}(x_n) = \arg \min_{m \in \{1, \ldots, M\}} \|x_n^T - c^{(m)}\|.
\]

### 5.6 Weight Normalization

In early experiments with the learning algorithm, we found that some units were much more active than others, with corresponding rows in the weight matrices much larger than average. This suggests that constraints need to be added to weight matrices to ensure that all units have reasonably equal chances of firing. These constraints can also be thought of as a way of avoiding certain units being starved of connection weights. A similar issue arose in the development of our dictionary learning algorithm (Kreutz-Delgado et al., 2003) and led us to enforce equality among the norms of each column of the weight matrix. Here, both

---

9In theory, the thresholding should not significantly affect the learning. However, due to the size of the network, it was not practical to compare thresholded versus nonthresholded performance. Even with the smallest dictionary size (64 \( \times \) 64, layer 1 input size = 4096), there are about 5,570,000 weights. Thresholding reduces the actual number of weights updated to about 45,000 per pattern, an increase in speed of over 100 times.
row and column normalization are performed on each weight matrix (feed-
forward, lateral, and feedback). Normalization values are set heuristically
for each layer, with an initial value of 1.0 and increasing layer normalization
until sufficient activity can be supported by that layer. The normalization
values remain constant during network training and are adjusted from trial
to trial.

6 Visual Recognition and Inference Experiments

In this section, we detail experiments with the learning algorithm devel-
oped above and demonstrate four types of visual inference: recognition,
reconstruction, imagination, and expectation-driven segmentation.

A set of gray-scale images was generated using the Lightwave photo-
realistic rendering software. Each of 10 objects was rotated 360 degrees
through its vertical axis in 2 degree increments, for a total of $10 \times 180 = 1800$
images, of which 1440 were used for training and the 360 remaining were
held out for testing (see Figure 4). All images were $64 \times 64$ pixels. Before im-
ages can be presented to the network, they must be sparsely coded, which
is done with a sequence of preprocessing (see Figure 5). First, each image is
edge-detected to simulate the on-center/off-center contrast enhancement
performed by the retina and LGN (see Figure 5B). Edge-detected images
are then scaled by subtracting 128 and dividing by 256, so that pixel values
are $\in [-0.5, 0.5]$. Next, each image is divided into $8 \times 8$ pixel patches and
sparsely coded with FOCUSS+ using a dictionary learned by FOCUSS-
CNDL+ as described in appendix A (see Figure 5C). Dictionaries of size $64
\times 64$, $64 \times 128$, and $64 \times 196$ were learned to compare the effect of vary-
ing degrees of overcompleteness on recognition performance. (Figures 6
to 14 in this section are from experiments with the $64 \times 196$ dictionary.)
Table 3 shows the accuracy and diversity of the image codes. As dictionary
overcompleteness increases from $64 \times 128$ to $64 \times 196$, both mean square
error (MSE) and mean diversity decrease; images are more accurately rep-
resented using a smaller number of active elements chosen from the larger
overcomplete dictionary. As seen in Figure 5C, the reconstructed images
accurately represent the edge information even though they are sparsely
coded (on average 192 of 12,288 coefficients are nonzero). Finally, the non-
negative sparse codes are thresholded to $\{0, 1\}$ binary values before being
presented to the network; any value greater than 0.02 is set to 1 (see Fig-
ure 5D). This stage, however, does introduce errors in the reconstruction
process, and the fidelity of the network’s reconstructions will be limited

10 Available online at www.newtek.com/products/lightwave/.
11 Edge detection was done with XnView software (www.xnview.com) using the edge
detect light filter, which uses the $3 \times 3$ convolution kernel $[0\ -1\ 0\ -1\ 4\ -1;\ 0\ -1\ 0]$. 
Figure 4: (A) Objects used in the experiments, showing one of the 180 views of each object. Images are 64 × 64 pixel gray scale. (B) Sample rotated object images in the data set.

6.1 Recognition with a Four-Layer Network. To test recognition performance, a four-layer network was trained using the data set described above. The training parameters of the network are given in Table 4. Note that all the lateral interactions were forced to be inhibitory or 0 and that no lateral connections were used in the first layer (as we assume the increase in sparsity produced by the FOCUSS+ iterations model the layer 1 lateral connections). Coded images were presented to the first layer of the network for the initial three time steps. Random object codes with $r_4 = 10$ nonzero elements were used on the highest layer. Training took between 11 and 22 hours depending on dictionary size using an Intel Xeon 2.8 Ghz processor. Classification performance reached 100% accuracy on the test set after 135 epochs, but training continued until 1000 epochs to increase the reconstruction accuracy at the first layer. Figure 6 shows the iterations of the network state $X_t$ during classification of a test set image. The first row
Figure 5: Several preprocessing steps are done before presentation to the network: edge detection, FOCUSS+ sparse coding, and binarization of coefficients. (A) Original images. (B) Edge-detected images. (C) Reconstructions from FOCUSS+ codes using a learned overcomplete dictionary. (D) Reconstructions from binarized FOCUSS+ codes.

Table 3: Coding Performance on $64 \times 64$ Pixel Images (Blocked into $8 \times 8$ Patches) Using Complete and Overcomplete Dictionaries.

<table>
<thead>
<tr>
<th>Dictionary Size</th>
<th>Layer 1 Size</th>
<th>MSE</th>
<th>Maximum</th>
<th>Mean</th>
<th>Minimum</th>
</tr>
</thead>
<tbody>
<tr>
<td>$64 \times 64$</td>
<td>4096</td>
<td>0.00460</td>
<td>0.0449</td>
<td>0.0266</td>
<td>0.0103</td>
</tr>
<tr>
<td>$64 \times 128$</td>
<td>8192</td>
<td>0.00398</td>
<td>0.0339</td>
<td>0.0240</td>
<td>0.0128</td>
</tr>
<tr>
<td>$61 \times 196$</td>
<td>12,288</td>
<td>0.00292</td>
<td>0.0221</td>
<td>0.0156</td>
<td>0.0085</td>
</tr>
</tbody>
</table>

Note: Mean squared-error (MSE) is calculated over all $8 \times 8$ patches in the image, and diversity $= (\# \text{non-zero coefficients})/(\text{layer 1 size})$.

shows the FOCUSS+ coded input image and the original. The next rows show the activity of each layer and the reconstructed image from the first layer. The object was presented for three time steps and then removed, so that all activity on layer 1 for $t \geq 4$ results from network feedback. As the iterations proceed, the reconstruction completes the outline of the airplane.
Figure 6: Recognition of a test set object. Each row shows the network activity $X_t$ at a time step. In layer 4, the marker ■ indicates that the unit is active and is part of the correct object code, ○ that the unit is in the object code but inactive, and × that the unit is active but should not be. When $t > 3$, there is no external input, and the reconstructed image in layer 1 is due only to network feedback. At $t = 4$ in layer 4, there are four incorrectly activated units (×), but at later times, the dynamics of the network suppress these incorrectly active units.
Table 4: Network Parameters for Training the Four-Layer Network with 64 × 196 Overcomplete Dictionary, Corresponding to Layer 1 Size of 12,288.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>s (layer size)</td>
<td>12,288, 512, 512, 256</td>
</tr>
<tr>
<td>$\bar{r}$ (maximum diversity of layer)</td>
<td>430, 100, 100, 100</td>
</tr>
<tr>
<td>$\tau$ (time iterations per pattern)</td>
<td>8</td>
</tr>
<tr>
<td>$\eta$ (learning rate)</td>
<td>0.002</td>
</tr>
<tr>
<td>$\lambda$ (regularization parameter)</td>
<td>0.005</td>
</tr>
<tr>
<td>$\mu$ (target mean for hidden layers)</td>
<td>$-4.0$</td>
</tr>
<tr>
<td>Epoch size (number of patterns)</td>
<td>100</td>
</tr>
<tr>
<td>Maximum number of epochs</td>
<td>1000</td>
</tr>
<tr>
<td>Feedforward weight range</td>
<td>$[-5.0, 5.0]$</td>
</tr>
<tr>
<td>Feedback weight range</td>
<td>$[-5.0, 5.0]$</td>
</tr>
<tr>
<td>Lateral weight range</td>
<td>$[-5.0, 0.0]$</td>
</tr>
<tr>
<td>Layer 1 norms (FB)</td>
<td>[12.0]</td>
</tr>
<tr>
<td>Layer 2 norms (FF, L, FB)</td>
<td>[12.0, 2.1, 2.1]</td>
</tr>
<tr>
<td>Layer 3 norms (FF, L, FB)</td>
<td>[5.9, 2.1, 1.5]</td>
</tr>
<tr>
<td>Layer 4 norms (FF, L)</td>
<td>[1.5, 1.5]</td>
</tr>
</tbody>
</table>

Note: For other sized dictionaries, the size of the first layer was 8192 (64 × 128 dictionary) and 4096 (64 × 64 dictionary), with all other parameters as listed in the table.

and becomes stronger in intensity. In layer 4, the marker shape indicates whether the unit is active and is part of the correct object code (■), or is part of the object code but inactive (○), or is active but should not be (×). At $t = 4$, all 10 of the highest layer units in the object code for airplane are active (■), so that the image is classified correctly; however, there are four other units active that should not be (×). At later iterations, these extra incorrect units are deactivated (or “sparsified away”) so that at $t \geq 5$, only those units in the object code are active, demonstrating the importance of lateral connections in the highest layer. Activity in layers 2 and 3 also decreases with time.

Presenting rotated test set views of the object shows that the network has learned basins of attraction for the other orientations. Figure 7 shows the state of the network at $t = 7$ after presenting various rotations of the airplane. The invariance of the representation is shown to increase from layer 1 (with nearly completely different units active) through layer 3 (with many of the same units active) to layer 4 (which has identical activity for all four orientations of the airplane).

Training on rotated objects gives the network some robustness to small translations. When tested on images translated $+/-1$ pixel in each direction, recognition accuracy is 96.9% on the test set. However, in general, we make no claim that our network has learned transformations (such as translation or scaling) that it has not seen.

The network includes many parameters (see Table 4), and learning is more sensitive to some than others. For example, the maximum activity $\bar{r}_2, \bar{r}_3$ for layers 3 and 4 can vary quite widely without noticeable effect on performance, while increasing $\bar{r}_2$ to 512 (from 100) increases the training time by more than an order of magnitude. This is because weights are
Figure 7: Each row is the network state $X_t$ at $t = 7$ after presenting various rotated images of the airplane (test set images, views unseen during training), demonstrating that multiple basins of attraction can be learned for each object. Higher layers show more invariant representations than lower layers, with layer 4 showing the fully invariant representation of the airplane.

updated only between active units, and increasing the maximum number of active units on layer 2 results in a very large number of weights to and from layer 1 that must be updated. When the diversity penalty is turned off ($\lambda = 0$), the average diversity of the second and third layers increases by about 30% and 60%, respectively, with no significant change in MSE or classification rates. This demonstrates that using the diversity penalty results in more efficient representations (more sparse), consistent with our sparse generative-world model.

We also experimented with different variations of the input time series, and these changes had more dramatic effects on performance. Two experiments were done with 12 time steps: (1) with input presented for 6 time steps and turned off for 6 steps and (2) using a linear decay $1 - t/6$ (input presented at full strength at the first time step and decaying to 0 at the sixth time step). The performance of both of these experiments was worse than the original method (input presented for three steps). For the six-step input, the recognition accuracy reached only 85% on the test set; for the decaying six-step input, the accuracy was 90% compared with 100% using the original method. It is not clear why performance drops, but there seems to be a reduction in middle layer activity. Perhaps adjusting normalization or other parameters could improve these results.
Figure 8: (A) Histogram of FOCUSS+ coefficient values before binarization. There are $2.9 \times 10^8$ elements in the zero bin, and the maximum coefficient value is 2.9. (B) Histogram and gaussian fit of a randomly chosen unit in the preactivation state $V_t$ in layer 3 before training. (C) Histogram and gaussian fit of residual $\varepsilon_t$ in layer 1 after training.

To test the gaussian assumptions made regarding $V_t$ and the errors $\varepsilon_t$, we plot histograms and normal curve fits of randomly chosen units in Figures 8B and 8C. From Figure 8B we can see that distribution of units in $V_t$ is quite normal before training. In Figure 8C, we see that after training, the errors $\varepsilon_t$ for layer 1 are less gaussian but still reasonably modeled as such. Also, there is more mass in the negative tail, indicating patterns where the target values are 1 but the network output is much lower.

6.2 Reconstruction of Occluded Images. Using the same network trained in section 6.1, reconstruction is demonstrated using occluded images from the test set. Approximately 50% of pixels are set to black by choosing a random contiguous portion of the image to hide. Figure 9 shows the network iterations during reconstruction, where an occluded image is presented for the first three time steps. By $t = 3$, the feedback connections to the first layer have reconstructed much of the outline of the copier object, showing that feedback from the second layer contains much of the orientation-dependent information for this object. Further iterations increase the completion of the outline, particularly of the bottom corner and lower-right panel. (Another example of reconstruction is shown in Figure 1.13 of Murray, 2005.)

The network also performs well when recognizing occluded objects. Accuracy is 90% on the occluded test set objects with the complete dictionary ($64 \times 64$) and 96% to 97% with the overcomplete dictionaries. Figure 9 shows that (as above) there are incorrectly activated units in layer 4 at $t = 4$, which are suppressed during later times. In contrast with Figure 6, in layer 2 here, there is more activity as time progresses, presumably due to the activation of missing features during reconstruction.

More insight into reconstruction can be gained by examining the receptive and projective fields of units in the middle layers. Considering layer 2
Figure 9: Reconstruction of an occluded input image. As early as $t = 3$, feedback from layer 2 results in reconstruction of some of the outer edges of the objects. More detail is filled in at later time steps. Layer 4 legend: ■ = unit is active and in correct object code. ○ = unit is in the object code but inactive. × = unit is active but should not be (not in object code).
Figure 10: Receptive and projective fields of four units in layer 2. For each unit, the top row shows the receptive fields (feedforward weights from layer 1 to 2), and the bottom row shows the projective field (feedback weights from layer 2 to 1). The weight vectors are converted into images by multiplying by the learned dictionary. The unit in the first column is tuned to respond to both the plane and the table, while its projective field appears to include many possible orientations of the plane.

(see Figure 10), we find that the receptive fields (top row) tend to learn a large-scale representation of a particular orientation of an object. This is mainly because the receptive fields are allowed to cover the entire first layer, and no topology is enforced on the weights. Some receptive fields (such as the first column of Figure 10) are tuned to two very different objects, suggesting that units are recruited to represent more than one object, as would be expected from an efficient distributed code. The projective fields are not as clearly specific to a particular orientation and include strong noise, which indicates there must be inhibitory feedback from other layer 2 units, contributing to the cleaner version of the layer 1 outputs when the full network is run.

6.3 Imagination: Running the Network Generatively. Imagination is the process of running the network generatively with input given as an object code at the highest layer. For this experiment, the network trained in section 6.1 is used with an object code clamped on the highest layer for all time steps. Random activity is added to the second layer at $t = 3$ so that the network has a means of choosing which view of the object to generate. It was found that increasing the feedback strength (by multiplying feedback weights by 5.0) to the first and second layers increased the activity and quality of the imagined image at the first layer. Without this increase, the layer 1 reconstruction was very likely to settle to the 0 state. Figure 11 shows the results when the object code for the knight is presented. At $t = 4$, the reconstruction is a superposition of many features from many objects but at later times, the outline of the object can be seen. The orientation of the
Figure 11: Imagination using the object code for the knight as the top-down input and the injection of random activity in layer 2 at $t = 3$. The reconstruction is a bistable (oscillating) pattern of the object from the front and side views.
generated image alternates between a front view ($t = 5, 7$) and a side view ($t = 6, 8$), which is reminiscent of the bistable percept effect. Not all trials of this experiment result in a bistable state; the majority converged to a single orientation. Interestingly, some orientations of certain objects appear to be generated much more often than other orientations. These “canonical views” represent high-probability (low-energy) states of the network. A random sample of five imagined objects is shown in Figure 12, showing that a superposition of states can also occur, which is consistent with the projective field properties shown in Figure 10.

6.4 Expectation-Driven Segmentation: Out from Clutter. In expectation-driven inference, both an input image and a top-down expectation are presented to the network, and the output can be either the highest-layer classification or the lowest-layer reconstructed image. Here, we considered the latter case, where the desired output is a segmented image reconstructed from the first layer. The same network trained in section 6.1 is used here with increased feedback strength as described in section 6.3. Cluttered input images are created by combining many objects from the data set at random translations, overlaid with a portion of the desired image (the same portion, 50%, used in the reconstruction experiment). This is a fairly difficult recognition problem, as the clutter in each image is composed of features from trained objects, so that competing features tend to confound recognition algorithms. Although the features from the clutter objects are likely to be in different locations than seen during training, it is still a more difficult task than segmentation from a randomly chosen (untrained) background.

The problem of expectation-driven segmentation is different from recognition in that we ask the network not, “What object is this?” but, “Assuming object X is here, what features in the image most likely correspond to it?” For this experiment, we present at $t = 2, 3$ the image of the occluded object in clutter and at $t = 1, \ldots, 4$ the expectation that the object is present at the
highest layer. Figure 13 shows the network states when presented with a cluttered image and top-level expectation of the knight object. The timing of the inputs was arranged so that the feedback and feedforward input first interact at \( t = 3 \) in layer 3. When \( t = 4 \), the input image is no longer presented, and the network feedback has isolated some features of the object. Later time steps show a sharper and more accurate outline of the knight, including edges that were occluded in the input image. At the highest layer, feedforward interactions from lower layers cause the correct object code to degrade. At \( t = 12 \), all the units in the object code for knight were active, as well as four incorrectly active units, which still allows correct classification. To illustrate the need for the top-down expectation input in this case, Figure 14 shows the states at \( t = 1, 4, 8 \) when no object code is presented at layer 4. The activity gradually decays, and there is no reconstruction at layer 1. Comparing Figure 11 (imagination) and Figure 13 shows that the partial information provided in the cluttered image is enough to keep the network at a stable estimate of segmentation and in this case prevent oscillations between two orientations (which occurred when only top-down input was present).

6.5 Overcompleteness Improves Recognition Performance. One of the central questions addressed in this work is how a sparse overcomplete representation in the early stages of visual processing, such as V1 in primates (Sereno et al., 1995), could be useful for visual inference. As described above, we trained networks using learned dictionaries of varying degrees of overcompleteness, \( 64 \times 64 \), \( 64 \times 128 \), and \( 64 \times 196 \), and corresponding sizes of the first layer: 4096, 8192 and 12,288. Performance was compared on the test set objects, occluded objects, and objects in clutter. The cluttered images were created by overlaying the entire object on a cluttered background, resulting in a somewhat easier problem than the occluded-object-in-clutter images used in section 6, although here no top-down expectations were used to inform the recognition. Figure 15 shows the recognition accuracy on these three image sets. For the test set (complete images), all three networks had performance at 99% to 100%, but for the occluded and cluttered images, there is a gain in accuracy when using overcomplete representations, and the effect is more pronounced for the more difficult cluttered images. For occluded objects, accuracy was 90% (324/360) for the complete dictionary and 97% (349/360) for the 3X overcomplete dictionary. The most significant improvement was with the cluttered images: accuracy was 44% (160/360) for the complete dictionary and 73% (263/360) for the 3X overcomplete dictionary. While the absolute classification rate for the cluttered images might appear low (44–73%), many of the misclassified objects were those of smaller size (e.g., the airplane and fire hydrant), which allowed more features from other larger objects to be visible and confound the recognition. In addition, neither the dictionary nor the network was trained on images
Figure 13: Expectation-driven segmentation using occluded objects over a cluttered background. The clutter input is presented at the lowest layer for $t = 2, 3$. Top-down expectations (the object code for knight) are presented at the highest layer for $t = 1, \ldots, 4$. By $t = 12$, the network converges to a segmented outline of the knight in the correct orientation at the first layer. Layer 4 legend: ■ = unit is active and in correct object code. ◦ = unit is in the object code but inactive. × = unit is active but should not be (not in object code).
Figure 14: Recognizing the occluded object in a cluttered background is difficult without top-down expectations. The same input image used in Figure 13 is presented for $t = 1, 2, 3$; however, no top-down inputs are present. A few representative time steps show that the activity gradually decays over time, and no object is reconstructed at layer 1. Layer 4 legend: ■ = unit is active and in correct object code. ◦ = unit is in the object code but inactive. × = unit is active but should not be (not in object code).

with clutter, so the network had no previous experience with this particular type of cluttered images.

7 Discussion

In this section, we discuss the motivations for our network and compare it with other recurrent and probabilistic models of vision. Additional discussion can be found in Murray (2005, sec. 1.8).

7.1 Why Sparse Overcomplete Coding and Recurrence? In the brain, early visual areas are highly overcomplete, with about 200 to 300 million neurons in V1 compared to only about 1 million neurons that represent the retina in the lateral geniculate nucleus (LGN) of the thalamus (Stevens, 2001; Ejima et al., 2003). As primate evolution has progressed, there has been an increase in the ratio of V1 to lateral geniculate nucleus (LGN) size. While even the smallest of primates shows a high degree of overcompleteness, the increase in higher primates is linked with an increase in retinal resolution
and presumably improved visual acuity (e.g., 87 times overcomplete for the tarsier monkey compared with 200 to 300 times for humans).

Mathematically, sparse coding strategies are necessary to make efficient use of overcomplete dictionaries because the dictionary elements are generically nonorthogonal. To provide a low-redundancy representation (Attnave, 1954; Barlow, 1959), a sparse set of elements must be chosen that accurately represents the input. If we have faith in the generative model postulated in Figure 1, real-world images can be accurately modeled as being caused by a small number of features and objects, supporting the choice of a sparse prior (even in the case of complete coding). Other benefits of sparse coding include making it easier to find correspondences and higher-order correlations, increasing the signal-to-noise ratio, and increasing the storage and representational capacity of associative memories (Field, 1994). Biological evidence for sparse coding ranges from the simple fact that average neural firing rates are low, 4 to 10 Hz (Kreiman, Koch, & Fried, 2000), to experiments that find sparseness in V1 increases as larger patches of natural images are presented, indicating that a concise representation can be found by deactivating redundant features, presumably through the interaction of lateral and feedback inhibition (Vinje & Gallant, 2000). One of the successes of sparse-coding theory has been the learning of receptive fields that resemble the orientation and location selectivity of V1 neurons (Olshausen & Field, 1997), and extensions have been made to model complex cells (Hoyer & Hyvärinen, 2002).
While overcompleteness and sparse coding are important features of early vision in V1, perhaps the most striking aspect of higher visual areas is the amount of lateral and feedback connections within and between areas (Felleman & Van Essen, 1991). Even in V1, lateral and feedback input from other cortical areas account for about 65% of activity, with only 35% of response directly due to feedforward connections from the LGN (Olshausen & Field, 2005). We showed in section 2 that feedback and lateral connections are required for many types of inference. In some recognition tasks, there is evidence that the brain is fast enough to complete recognition without extensive recurrent interaction (Thorpe, Fize, & Marlot, 1996). Consistent with this, our model is capable of quickly recognizing objects in tasks such as Figure 6, where the correct object code is found at $t = 5$. However, more difficult tasks such as segmentation (see Figure 13) require recurrence and would take longer for the brain (Lee, Mumford, Romero, & Lamme, 1998).

7.2 Related Work: Biologically Motivated Models of Vision. There have been many hierarchical models created to explain vision, and these fall into two main categories: feedforward only or recurrent (which include various types of feedback and lateral connections between layers). Some examples of the feedforward class are the Neocognitron model of Fukushima and Miyake (1982), VisNet of Rolls and Milward (2000), and the invariant-recognition networks of Földiáék (1991) and Riesenhuber and Poggio (1999). While many of these models use sparsity with some form of winner-take-all competition, which is usually interpreted as lateral interaction, since they do not include feedback connections, they are not capable of the range of inference described in section 2.2 and will not be discussed further here.

One of the more closely related works is the dynamic network developed by Rao and Ballard (1997). A stochastic generative model for images is presented, and a hierarchical network is developed to estimate the underlying state. Their network includes multiple layers with feedforward and feedback connections, which are interpreted as passing the residuals from predictions at higher levels back to lower levels (but with no explicit learnable lateral connections or overcomplete representations). Experiments demonstrate recognition, reconstruction of occluded images, learning of biologically plausible receptive fields, and ability to tell that an object had not been seen during training. Perhaps because of the computational requirements, only fairly limited recognition experiments were performed, using five objects (one orientation per object) and rotation-invariant recognition with two objects, each with 36 views used for training and testing (Rao, 1999).

Newer versions of the Neocognitron include feedback connections and are demonstrated for recognition and reconstruction (Fukushima, 2005). The model posits two types of cells in each region of the system, S-cells and C-cells, in analogy with the simple and complex cells categorized by Hubel and Wiesel (1959). The S-cells are feature detectors, and the C-cells
pool the output of S-cells to create invariant feature detectors. To solve the reconstruction problem, further cell types and layers are added, and many of the layers have different learning rules. In contrast, our network is able to perform various inference types without changes to the architecture or learning rule.

7.3 Related Work: Probabilistic Models in Computer Vision. Recent work in computer vision has investigated probabilistic generative models apart from direct biological motivation (Hinton et al., 2006; Hinton and Salakhutdinov, 2006; Fergus, Perona, & Zisserman, 2007; Sudderth, Torralba, Freeman, & Willsky, 2005).

Most closely related to our work is the learning algorithm of Hinton et al. (2006) for hierarchical belief networks. The network has multiple hidden layers, followed by a much larger overcomplete associative memory (whereas our overcomplete stage occurs at the second layer), and a highest layer with a 1-out-of-$n$ code for the object class. The first layer has real-valued inputs, while stochastic binary values are used at higher layers. Feedforward and feedback weights are learned, but no lateral connections are used, and during testing, only one forward-backward pass is made at each layer. When trained on a benchmark handwritten digit data set, the accuracy is competitive with the best machine learning methods, showing that generative hierarchical networks are promising for real-world vision tasks. Using a similar learning procedure with an autoencoder network architecture, Hinton and Salakhutdinov (2006) show applications to data compression and text classification.

While there are many differences between this work and our algorithm, they address the same basic question of how to train hierarchical generative models. One important difference is that Hinton et al. (2006) use stochastic units and Gibbs sampling for generative inference, while we use a nearest-layer conditional variational approximation. We believe the factorial approximation of equation 2.8 can be sufficiently accurate in the case of sparse activations and that enforcing a short enough time horizon $\tau$ makes learning computationally tractable. More experiments with known generative models will be needed to further evaluate the differences between these algorithms.

Fergus et al. (2007) develop a model for classification of object categories in unsegmented images. The first step is finding a small set of interesting features using a saliency detector. For each category, a probabilistic model is learned for these features, including their relative position and scale. Impressive detection performance is achieved on real-world data sets. In contrast to our work, which models all the features in the image, Fergus et al. (2007) use only a small number of features (fewer than 30), so that, if run generatively, their model would reconstruct only a small subset of the features in each object. Using a saliency detector improves position and scale invariance (which would benefit our network); however, using only
this small feature set reduces performance when features of a class model cannot be found.

In a related work, Sudderth et al. (2005) present a probabilistic model of object features and apply it to object categorization in real-world scenes. Similar to our model and in contrast with Fergus et al. (2007), their model is a true multiclass classifier, which allows features to be shared between models of different objects and allows for more rapid classification without the need to run multiple classifiers. As above, the small feature set limits the potential detail of generative reconstruction. However, segmentation results show that regions such as “building,” “car,” and “street” can be detected in city scenes.

8 Conclusion

We have developed a framework and learning algorithm for visual recognition and other types of inference such as imagination, reconstruction of occluded objects, and expectation-driven segmentation. Guided by properties of biological vision, particularly sparse overcomplete representations, we posit a stochastic generative world model. Visual tasks are formulated as inference problems on this model, in which inputs can be presented at the highest layer, lowest layer, or both, depending on the task. A variational approximation (the simplified world model) is developed for inference, which is generalized into a discrete-time dynamic network.

An algorithm is derived for learning the weights in the dynamic network, with sparsity-enforcing priors and error-driven learning based on the preactivated state vector. Experiments with rotated objects show that the network dynamics quickly settle into easily interpretable states. We demonstrate the importance of top-down connections for expectation-driven segmentation of cluttered and occluded images. Four types of inference were demonstrated using the same network architecture, learning algorithm, and training data. We show that an increase in overcompleteness leads directly to improved recognition and segmentation in occluded and cluttered scenes. Our intuition as to why these benefits arise is that overcomplete codes allow the formation of more basins of attraction and higher representational capacity.

Appendix A: Sparse Image Coding with Learned Overcomplete Dictionaries

The dynamic network and learning algorithm presented above require that the inputs $u_l$ at each layer be sparse vectors. To transform the input image into a suitable sparse vector, we use the focal underdetermined system solver (FOCUSS) algorithm for finding solutions to inverse problems. The FOCUSS algorithm represents data in terms of a linear combination of a small number of vectors from a dictionary, which may be overcomplete.
Other methods for sparsely coding signals include matching pursuit, basis pursuit, and sparse Bayesian learning, which were also evaluated for image coding (Murray & Kreutz-Delgado, 2006). The overcomplete dictionary is learned using the FOCUSST-CNDL (column-normalized dictionary learning) algorithm developed by Murray and Kreutz-Delgado (2001) and Kreutz-Delgado et al. (2003).

The problem that FOCUSST-CNDL addresses here is that of representing a small patch of an image $y \in \mathbb{R}^m$ using a small number of nonzero components in the source vector $x \in \mathbb{R}^n$ under the linear generative model,

$$ y = Ax, \quad \text{(A.1)} $$

where the dictionary $A$ may be overcomplete, $n \geq m$. The algorithm updates and more discussion of the FOCUSST-CNDL algorithm in this context are given in Murray (2005, section 1.A). Parameters for FOCUSST-CNDL are: data set size $= 20,000$ image patches; block size $N = 200$; dictionary size $= 64 \times 64, 64 \times 128, \text{or} 64 \times 196$; diversity measure $p = 1.0$; regularization parameter $\lambda_{\text{max}} = 2 \times 10^{-4}$; learning rate $\gamma = 0.01$; number of training epochs $= 150$; reinitialization every 50 epochs. After each dictionary update, $A$ is normalized to have unit Frobenius norm, $\|A\|_F = 1$ and equal column norms. Figure 1.18 of Murray (2005) shows the learned $64 \times 196$ dictionary after training on edge-detected patches of man-made objects (the data set described in section 6).

Once the dictionary $A$ has been learned, input images for the dynamic network (DN) are coded using the FOCUSST+ algorithm (Murray & Kreutz-Delgado, 2006). The input images are divided into consecutive nonoverlapping patches of the same $8 \times 8$ size used for dictionary learning. The FOCUSST+ algorithm consists of repeated iterations of equation 6 from Murray and Kreutz-Delgado (2006) over an image patch $y_k$ to estimate $x_k$. Each $x_k$ is updated for 15 iterations with $p = 0.5$.

### Appendix B: Derivation of Learning Algorithm for $\mathbb{W}$

The learning algorithm for the weights $\mathbb{W}$ is derived similarly to the backpropagation through time algorithm (BPTT) (Williams and Peng, 1990). Using the preactivation cost function, equation 4.11, for an individual pattern,

$$ J_{PA} = \frac{1}{2} \sum_{t=1}^{\tau} \left[ \epsilon_t^T (\epsilon_t \odot \beta_t) + \lambda (V_t - \mu)^T [(V_t - \mu) \odot (1 - \beta_t)] \right], \quad \text{(B.1)} $$

which uses states $V_t$ generated from running the network in certainty-equivalence mode. The effect of the weight prior $-\ln P(\mathbb{W})$ will not be considered in this section, as it was found that enforcing periodic weight
normalization is more computationally efficient than using prior constraints in every weight update (see section 5).

To minimize the cost $J_{PA}$, we update the weights using gradient descent,

$$
\Delta w_{ji} = -\eta \frac{\partial J_{PA}}{\partial w_{ji}} = -\eta \sum_{t=1}^{\tau} \frac{\partial J_{PA}}{\partial V_{j,t}} \cdot \frac{\partial V_{j,t}}{\partial w_{ji}},
$$

(B.2)

where $w_{ji}$ is the element from the $j$th row and $i$th column of $W$. The second term on the right is

$$
\frac{\partial V_{i,t}}{\partial w_{ji}} = \frac{\partial}{\partial w_{ji}} \mathbb{W}_j X_t = X_{i,t},
$$

(B.3)

where $\mathbb{W}_j$ is the $j$th row of $\mathbb{W}$. The first term on the right of equation B.2 is divided into two parts,

$$
\frac{\partial J_{PA}}{\partial V_{j,t}} = B_{j,t} + D_{j,t}
$$

where $B$ is related to reconstruction error and $D$ increases sparsity on those layers without desired values. Recursion expressions can now be found for $B_{j,t}$ and $D_{j,t}$. First, some notation: the $j$th row of the weight matrix $W$ is denoted $W_j$, and the element from the $j$th row and $i$th column is $w_{ji}$. Beginning with the reconstruction-enforcing term $B$ (temporarily omitting the binary indicator variable $\beta$ for notational clarity),

$$
B_{j,t} = \frac{\partial}{\partial V_{j,t}} \left[ \frac{1}{2} \sum_{\rho=1}^{\tau} e_{\rho}^T (e_{\rho} \odot \beta_{\rho}) \right],
$$

(B.5)

For $B_{j,t}$, at the last time step in equation B.2, when $t = \tau$, only the $\rho = \tau$ terms depend on $V_{j,\tau}$,

$$
B_{j,\tau} = \frac{\partial}{\partial V_{j,\tau}} \left[ \frac{1}{2} e_{\tau}^T e_{\tau} \right] = e_{\tau}^T \frac{\partial}{\partial V_{j,\tau}} e_{\tau} = e_{j,\tau},
$$

(B.6)
where $\varepsilon_{j,t}$ is the $j$th element of the error vector $\varepsilon_t$. When $t = \tau - 1,$

$$B_{j,\tau-1} = \frac{\partial}{\partial V_{j,\tau-1}} \left[ \frac{1}{2} \varepsilon_T^T \varepsilon_\tau + \frac{1}{2} \varepsilon_{\tau-1}^T \varepsilon_{\tau-1} \right]. \quad (B.7)$$

The second term on the right can be found to be $-\varepsilon_{j,\tau-1}$ as in equation B.6. For the first term,

$$\frac{\partial}{\partial V_{j,\tau-1}} \frac{1}{2} \varepsilon_T^T \varepsilon_\tau = \varepsilon_T^T \frac{\partial}{\partial V_{j,\tau-1}} \varepsilon_\tau = \varepsilon_T^T \nabla \frac{\partial}{\partial V_{j,\tau-1}} f(V_{\tau-1})$$

$$= f'(V_{j,\tau-1}) \sum_{k=1}^{N} \varepsilon_{k,\tau} w_{kj}. \quad (B.8)$$

Substituting equations, B.8 and B.6 into the expression for $B_{j,\tau-1}$ (see equation B.7),

$$B_{j,\tau-1} = \varepsilon_{j,\tau-1} + f'(V_{j,\tau-1}) \sum_{k=1}^{N} B_{k,\tau} w_{kj}. \quad (B.9)$$

The general recursion for $B_{t,j}$ is (after reintroducing the indicator variable $\beta$)

$$B_{j,t} = \begin{cases} 
\varepsilon_{j,t} + \beta_{j,t} & t = \tau \\
\varepsilon_{j,t} + f'(V_{j,t}) \sum_{k=1}^{N} B_{k,t+1} w_{kj} & 1 \leq t \leq \tau - 1
\end{cases}. \quad (B.10)$$

Turning to the sparsity-enforcing term (again omitting $\beta$),

$$D_{j,t} = \frac{\partial}{\partial V_{j,t}} \left[ \lambda \sum_{\rho=1}^{\tau} (V_{t} - \mu)^T (V_{t} - \mu) \right]. \quad (B.11)$$

Following similarly to the derivation for $B$ above, when $t = \tau,$

$$D_{j,\tau} = \frac{\partial}{\partial V_{j,\tau}} \left[ \lambda \frac{1}{2} (V_{\tau} - \mu)^T (V_{\tau} - \mu) \right] = \lambda (V_{\tau} - \mu) \frac{\partial}{\partial V_{j,\tau}} (V_{\tau} - \mu) = \lambda (V_{j,\tau} - \mu). \quad (B.12)$$
When $t < \tau$, we follow equations B.7 to B.9, and find the general recursion for $D_{j,t}$ (reintroducing $\beta$),

$$D_{j,t} = \begin{cases} 
\lambda(V_{j,t} - \mu)(1 - \beta_{j,t}) & t = \tau \\
\lambda(V_{j,t} - \mu)(1 - \beta_{j,t}) + f'(V_{j,t} - \mu) \sum_{k=1}^{N} D_{k,t+1} w_{kj} & 1 \leq t \leq \tau - 1.
\end{cases}$$

(B.13)

the recursions B.10 and B.13 are used in the final weight update,

$$\Delta w_{ji} = -\eta \sum_{t=1}^{\tau} (B_{j,t} + D_{j,t}) X_{i,t},$$

(B.14)

where the activation function derivative is

$$f'(v) = \frac{a_1 a_2 \exp(-a_2 v + a_3)^2}{(1 + \exp(-a_2 v + a_3))}.$$ 

(B.15)

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