

How close are we to understanding V1? *

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Abstract

A wide variety of papers have reviewed what is known about the function of primary visual cortex. In this article, rather than stating what is known, we attempt to estimate how much is still unknown about V1 function. In particular, we identify five problems with the current view of V1 that stem largely from experimental and theoretical biases, in addition to the contributions of non-linearities in the cortex that are not well understood. Our purpose is to open the door to new theories, a number of which we describe along with some proposals for testing them.

1 Introduction

The primary visual cortex (area V1) of mammals has been the subject of intense study for at least four decades. Hubel and Wiesel's original studies in the early 1960's created a paradigm shift by demonstrating that the responses of single neurons in the cortex could be tied to distinct image properties such as the **local orientation of contrast** (Hubel & Wiesel 1959; 1968). Since that time, the **study of V1 has become something of a miniature industry**, to the point where the annual Society for Neuroscience meeting now routinely devotes multiple sessions entirely to V1 anatomy and physiology. Without doubt, much has been learned from these efforts. However, as we shall argue here, **there remains a great deal that is still unknown about how V1 works and its role in visual system function. We believe it is quite probable that the correct theory of V1 is still far afield from the currently proposed theories.**

It may seem surprising to some that we should take such a stance. V1 does after all have a seemingly ordered appearance—a clear topographic map, and an orderly arrangement of ocular dominance and orientation columns. Many neurons are

*An earlier version of this article appeared previously as, "What is the Other 85% of V1 Doing?" In: *23 Problems in Systems Neuroscience*. T.J. Sejnowski, L. van Hemmen, eds. Oxford University Press. 2005.

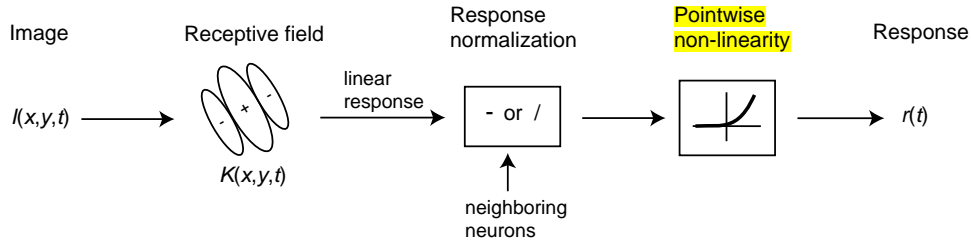


Figure 1: Standard model of V1 simple cell responses. The neuron computes a weighted sum of the image over space and time, and this result is normalized by the responses of neighboring units, and passed through a pointwise non-linearity (see e.g., Carandini et al., 1997).

demonstrably tuned for stimulus features such as **orientation**, **spatial-frequency**, **color**, **direction of motion**, and **disparity**. And there has even emerged a fairly well agreed upon “standard model” for V1 in which simple-cells compute a linearly weighted sum of the input over space and time (usually a Gabor-like function) which is then normalized by the responses of neighboring neurons and passed through a pointwise nonlinearity (Figure 1). Complex cells are similarly explained in terms of summing the outputs of a local pool of simple-cells with similar tuning properties but different positions or phases. A variety of models have been proposed for the response normalization (Heeger 1991; Geisler & Albrecht 1997; Schwartz & Simoncelli 2001; Cavanaugh et al., 2002a), but the net result is often to think of **V1 as a kind of “Gabor filter bank.”** There are numerous papers showing that this basic model fits much of the existing data well, and many scientists have come to accept this as a working model of V1 function (see e.g., Lennie, 2003a). Indeed, such models are widely used to predict psychophysical performance (Graham & Nachmias 1971; Watson et al. 1983; Anderson et al. 1991), and they have been shown to provide efficient representations of natural scenes (Olshausen & Field, 1996; Bell & Sejnowski, 1997).

But behind this picture of apparent orderliness, there lies an abundance of unexplained phenomena, a growing list of untidy findings, and an increasingly uncomfortable feeling among many about how the experiments that have led to our current view of V1 were conducted in the first place. The main problem stems from the fact that **cortical neurons are highly nonlinear**—i.e., they emit **all-or-nothing action potentials, not analog values**. They also adapt, so their response properties depend upon the history of activity. Most importantly, cortical pyramidal cells have highly elaborate dendritic trees, and realistic biophysical models that include voltage-gated channels suggest that **each thin branch could act as a non-linear subunit**, so that **any one neuron could be computing many different non-linear combinations of its inputs** (Hausser & Mel, 2003; Polsky et al., 2004), in addition to being sensitive to coincidences (Softky & Koch, 1993; Azouz & Gray, 2000, 2003).

Everyone knows that neurons are non-linear, but few have acknowledged the implications for studying cortical function. Unlike linear systems, where there exist math-

ematically tractable, textbook methods for system identification, non-linear systems can not be teased apart using some straightforward, structuralist approach. That is, there is no unique “basis set” with which one can probe the system to characterize its behavior in general.¹ Nevertheless, the structuralist approach has formed the bedrock of V1 physiology for the past four decades. Researchers have probed neurons with spots, edges, gratings and a variety of mathematically elegant functions in the hope that the true behavior of neurons can be explained in terms of some simple function of these components. However, the evidence that this approach has been successful is lacking. We simply have no reason to believe that a population of interacting neurons can be reduced in this way.

For any complex system, it seems reasonable to begin where the system acts rationally: to study the behavior under conditions where ones models are relatively effective. But for a neural system, that leaves the question as to whether such behavior represents the relevant aspect of the neurons activity—i.e., does this help us understand how neurons operate under natural conditions? Much of our understanding of V1 is derived from recording from one neuron at a time using simple stimuli (edges, gratings, spots). From this body of experiments has emerged the standard model that forms the basis for our conceptual understanding of V1. In recent years, a number of innovative studies have moved away from this basic approach, recording from multiple neurons with complex, ecologically relevant stimuli. Are these studies simply adding minor correction factors to our understanding, or will they require us to completely revamp the current theories? Are the current models close to accounting for the majority of responses in the majority of neurons in V1? How close are we to understanding V1?

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In this article we present our reasons for believing that we may have far to go in understanding V1. We identify five fundamental problems with the current view of V1 function that stem largely from experimental and theoretical biases, in addition to the contributions of non-linearities in the cortex that are not well understood. Furthermore, we attempt to quantify the level of our current understanding by considering two important factors: an estimate of the fraction of V1 neuron types that are typically characterized in experimental studies, and the fraction of variance explained in the responses of these neurons under natural viewing conditions. Together, these two factors lead us to conclude that at present we can rightfully claim to understand only 10-20% of how V1 actually operates under normal conditions.

Our aim in pointing these things out is not simply to tear down the current framework. We ourselves have attempted to account for some aspects of the standard model in terms of efficient coding principles (sparse coding), so obviously we believe that we have made a good start. Rather, our goal is to show how much room there is

¹The Volterra series expansion is often touted as a general approach for characterizing non-linear systems, but it has been of little practical value in analyzing neural systems because it requires estimating many higher-order moments. In addition, it is an overly-general, “black-box” approach that does not easily allow one to incorporate prior knowledge about the types of non-linearities known to exist in the nervous system.

for new theories, and where the weaknesses in the current theories might lie. In the second half of the article we describe a few of our favorite alternatives to the standard theories. A central conclusion that emerges from this exercise is that we need to begin seriously studying how V1 behaves using natural scenes, using multi-unit recording techniques, in addition to explicitly describing any potential biases in the gathering of data. We believe this approach can help to reveal not just how much we know about neural coding in the visual pathway, but also how much we do not know.

2 Five problems with the current view

2.1 Biased sampling of neurons

The vast majority of our knowledge about V1 function has been obtained from single unit recordings in which a single micro-electrode is brought into close proximity with a neuron in cortex. Ideally, when doing this one would like to obtain an unbiased sample from any given layer of cortex. But some biases are difficult to avoid. For instance, neurons with large cell bodies will give rise to extracellular action potentials that have larger amplitudes and propagate over larger distances than neurons with small cell bodies. Without careful spike sorting, the smaller extracellular action potentials may easily become lost in the background when in the vicinity of neurons with large extracellular action potentials. This creates a bias in sampling that is not easy to dismiss.

Even when a neuron has been successfully isolated, detailed investigation of the neuron may be bypassed if it does not respond “rationally” to standard test stimuli or fit the stereotype of what the investigator believes the neuron should do. This is especially true for higher visual areas such as V4, but it is also true for V1. Such neurons are commonly regarded as “visually unresponsive.” It is difficult to know how frequently such neurons are encountered because oftentimes they simply go unreported, or else it is simply stated that only visually responsive units were used for analysis.

While it is admittedly difficult to characterize the information processing capabilities of a neuron that seems unresponsive, it is still important to know *in what way* these neurons are unresponsive. What are the statistics of activity? Do they tend to appear bursty or tonic? Do they tend to be encountered in particular layers of cortex? And most importantly, are they merely unresponsive to bars and gratings, or are they also equally uninterpretable in their responses to a wider variety of stimuli, such as natural images? A seasoned experimentalist who has recorded from hundreds of neurons would probably have some feel for these things. But for the many readers not directly involved in collecting the data, there is no way of knowing these unreported aspects of V1 physiology. It is possible that someone may eventually come up with a theory that could account for some of these unresponsive neurons, but this can’t happen if no one knows they are there.

A related bias that arises in sampling neurons is that the process of hunting for neurons with a single micro-electrode will typically steer one towards neurons with

higher firing rates. One line of evidence suggesting that this is a significant bias comes from work estimating mean firing rates in the cortex based on energy consumption. Attwell & Laughlin (2001) and Lennie (2003b) calculate that the average activity must be relatively low—i.e., less than 1 Hz in primate cortex. However, in the single-unit literature one finds many studies in which even the spontaneous or background rates are well above 1 Hz. This suggests that the more active neurons are substantially overrepresented (Lennie 2003b). What makes matters worse is that if we assume V1 neurons exhibit an exponential firing rate distribution, as has been demonstrated for natural scenes and other stimuli (Baddeley et al., 1997), then a mean firing rate of 1 Hz would yield the distribution shown in Figure 2a. With such a distribution, only a small fraction of neurons would exhibit the sorts of firing rates normally associated with a robust response. For example, the total probability for firing rates of even 5 Hz and above is 0.007, meaning that one would have to wait 1-2 minutes on average in order to observe a one-second interval containing five or more spikes. It seems possible that such neurons could either be missed altogether, or else purposely bypassed because they do not yield enough spikes for data analysis. For example, the overall mean firing rate of V1 neurons in the Baddeley et al. study was 4.0 Hz (std. 3.6 Hz), suggesting that these neurons constitute a sub-population that were perhaps easier to find, but not necessarily representative of the population as a whole. Interestingly, the authors point out that even this rate is considered low (which they attribute to anaesthesia), as previous studies (Legendy & Salcman, 1985) report the mean firing rate to be 8.9 Hz (s.d. 7.0 Hz).

Given the variety of neurons in V1, it seems reasonable to presume there exists a heterogeneous population of neurons with different mean firing rates. If we assume some distribution over these rates, then it is possible to obtain an estimate of the fraction of the population characterized given a particular criterion response. And from that we can calculate what the observed mean rate would be for that fraction. The result of such an analysis, assuming a log-normal distribution of mean-rates with an overall mean of 1 Hz, is shown in Figure 2b. As one can see, an overall mean of 4 Hz implies that the selection criterion was somewhere between 1-2 Hz, which would capture less than 20% of the population.

Neurophysiological studies of the hippocampus provide an interesting lesson about the sorts of biases introduced by low firing rates. Prior to the use of chronic implants, in which the activity of neurons could be monitored for extended periods while a rat explored its environment, the granule cells of the dentate gyrus were thought to be mostly high rate “theta” cells (e.g., Rose et al., 1983). But it eventually became clear that the majority are actually very low rate cells (Jung & McNaughton, 1993), and that for technical reasons only high-rate interneurons were being detected in the earlier studies (W.E. Skaggs, personal communication, Jan. 2004). In fact, Thompson & Best (1989) found that nearly two-thirds of all hippocampal neurons which showed activity under anaesthesia became silent in the awake, behaving rat. This overall pattern appears to be upheld in macaque hippocampus, where the use of chronic implants now routinely yields neurons with overall firing rates below 0.1 Hz (Barnes et al., 2003), which differs by nearly two orders of magnitude from the “low baseline rates” of 8.1 Hz reported by Wirth et al. (2003) using acutely implanted electrodes.

much higher mean firing rates than brain energy consumption allows

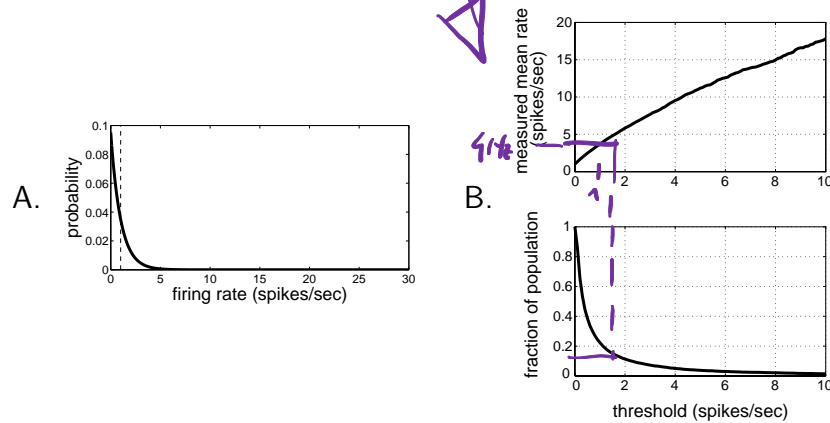


Figure 2: Sampling bias. A. Exponential firing rate distribution with a mean of 1 Hz (dashed line denotes mean). B. Resulting overall mean-rate of the population (top), and fraction of the population captured (bottom), as a result of recording from neurons only above a given mean firing-rate (threshold).

The dramatic turn of events afforded by the application of chronic implants combined with natural stimuli and behavior in the hippocampus can only make one wonder what mysteries could be unraveled when similar techniques are applied to visual cortex. What is the natural state of activity during free-viewing of natural scenes, where the animal is actively exploring its environment? What are the actual average firing rates and other statistics of activity among layer 2/3 pyramidal cells? What are the huge numbers of granule cells in macaque layer 4, which outnumber the geniculate fiber inputs by 30:1, doing? Do they provide a sparser code than their geniculate counterparts? And what about the distribution of actual receptive field sizes? Current estimates show that most parafoveal neurons in V1 have receptive field sizes on the order of 0.1 deg. But based on retinal anatomy and psychophysical performance one would expect to find a substantial number of neurons with receptive fields an order of magnitude smaller, ca. 0.01 deg. (Olshausen & Anderson, 1995). Such receptive field sizes are extremely rare, if not non-existent, in the existing data on macaque V1 neurons collected using acute recording techniques (De Valois et al., 1982; Parker & Hawken, 1988).

Overall, then, one can identify at least three different biases in the sampling of neurons:

1. preference for neurons with large cell bodies and large extra-cellular action potentials,
2. preference for “visually responsive” neurons, and
3. preference for neurons with high firing rates.

So where does this leave us? Let us be conservative. If we assume that 5-10% of neurons are missed because they have weak extracellular action potentials, another

5-10% are discarded because they are not visually unresponsive, and 50-60% are missed because of low firing rates (assuming a conservative threshold of 0.5 Hz in Figure 2), then even allowing for some overlap among these populations would yield the generous estimate that 40% of the population has actually been characterized.

2.2 Biased stimuli

Much of our current knowledge of V1 neural response properties is derived from experiments using reduced stimuli. Oftentimes these stimuli are ideal for characterizing linear systems—i.e., spots, white noise, or sinewave gratings—or else they are designed around pre-existing notions of how neurons *should* respond. The hope is that the insights gained from studying neurons using these reduced stimuli will generalize to more complex situation—e.g., natural scenes. But of course there is no guarantee that this is the case. And given the non-linearities inherent in neural responses, we have every reason to be skeptical.

Sinewave gratings are ubiquitous tools in visual system neurophysiology and psychophysics. In fact, the demand for using these stimuli is so high that some companies produce lab equipment with specialized routines designed for this purpose (e.g., Cambridge Research Systems). But sinewaves are special only because they are eigenfunctions of linear, time- or space-invariant systems. For non-linear systems, they bear no particular meaning, nor do they occupy any special status. In the auditory domain, sinewaves could be justified from the standpoint that many natural sounds are produced by oscillating membranes. However, in the visual world there are few things that naturally oscillate either spatially or temporally. The Fourier basis set is just one of many possible basis sets, and if the system is non-linear, no one basis set will necessarily provide a proper account of the system.

Bars of light, Gabor functions, Walsh patterns or any other basis set will suffer from similar problems requiring assumptions of the types of non-linearities that are present. The Gabor function has been argued to provide a good model of cortical receptive fields (Field & Tolhurst, 1986; Jones & Palmer, 1987). However, the methods used to measure the receptive field in the first place generally search for the best fitting linear model. They are not tests of how well the receptive field model actually describes the response of the neuron. Not until recent work by Gallant and colleagues (David, Vinje & Gallant, 2004) have these models been tested in ecological conditions. And as we discuss below, the results demonstrate that these models often fail to adequately capture the actual behavior of neurons.

The use of white noise and m-sequences can provide some advantage over the traditional linear systems approach, as they can provide a wider range of stimuli than a simple basis set and are thus capable of mapping out the non-linearity of a system if the non-linearities take on particular forms (e.g., Nykamp and Ringach, 2002). In addition, by analyzing the eigenvectors of the spike-triggered covariance matrix one can recover fairly complex non-linear models, such as the hypothetical subunits composing a complex cell, or suppressive dimensions in the stimulus space (Touryan et al. 2002; Rust et al. 2004).

However, there is only one way to map a non-linear system with complete confidence—

one must present the neuron with all possible stimuli. The scope of this task is truly breathtaking. Even an 8×8 pixel patch with 6 bits of grey level requires searching $2^{384} > 10^{100}$ possible combinations (a google of combinations). If we allow for temporal sensitivity and include a sequence of 10 such patches, we are exceeding 10^{1000} . With the estimated number of particles in the universe estimated to be in the range of 10^{80} , it should be clear that this is far beyond what any experimental method could explore. In theory, a non-linear neuron could behave quite rationally for all but a handful of these stimuli, so unless this handful has been measured, there is no way to be certain the neuron has been adequately characterized. The use of independent white noise can theoretically present a neuron with all possible stimuli. However, 10 hours of recording from a single neuron with a patch like that above at 30 frames per second will present just 10^6 out of the 10^{1000} possible stimuli. Using such a tiny fraction of the possible stimuli allows mapping of the non-linearities only if the non-linearities are quite smooth.

The deeper question is whether one can predict the responses of neurons from some combinatorial rule of the responses derived from a reduced set of stimuli. The response of the system to any reduced set of stimuli can not be guaranteed to provide the information needed to predict the response to an arbitrary combination of those stimuli. Of course, we will never know this until it is tested, and that is precisely the problem—the central assumption of the elementwise, reductionist approach has yet to be thoroughly tested.

We believe that the solution to these problems is to turn to natural scenes. Our intuitions for how to reduce stimuli should be guided by the sorts of structure that occur in natural scenes, not arbitrary (or even elegant) mathematical functions or stimuli that are conceptually simple or which happen to be easy to generate on a monitor. Since it is impossible to map out the response to all possible stimuli, some assumptions about the nature of the non-linearity and the stimulus space must be made. The assumption we believe is appropriate is that the non-linearities relevant to visual processing are most likely to be revealed when the system is presented with ecologically relevant stimuli.

Traditionally experimentalists have been reluctant to use natural scenes as stimuli because they seem highly variable and “uncontrolled.” But in recent years there has been significant progress in modeling the structure of natural images (Simoncelli & Olshausen, 2001), and it should soon be possible to develop parametric descriptions of natural images that could be used to generate experimental stimuli (e.g., Heeger & Bergen, 1995). In addition, there have been some recent attempts to map out the non-linearities in response to natural images (Sharpee et al, 2004). And the development of several adaptive stimulus techniques looks to be a promising avenue for determining the relevant stimulus for sensory neurons (Foldiak et al. 2004; Edin et al. 2004; O’Connor et al. 2004).

In summary, then, there are two reasons for using natural scenes as stimuli: 1) By devoting ones resources to relevant ecological stimuli, the experimentalist has a greater chance of finding and mapping the non-linearities relevant to the function of neurons, and 2) the responses to natural scenes provide an ecologically meaningful test of any neural model. Even if non-ecological stimuli are used to map a neurons

natural
stimuli

behavior, the true test that the characterization is correct is to demonstrate that one can predict the neurons behavior in ecological conditions.

2.3 Biased theories

Currently in neuroscience there is an emphasis on “telling a story.” This often encourages investigators to demonstrate when a theory explains data, not when a theory provides a poor model. In addition, editorial pressures can encourage one to make a tidy picture out of data that may actually be quite messy. This of course runs the risk of forcing a picture that does not actually exist. Theories then emerge that are centered around explaining a particular subset of published data, or which can be conveniently proven, rather than being motivated by functional considerations—i.e., how does this help the brain to solve the real problems of vision?

For instance, early work demonstrating the spatial frequency selectivity of neurons (e.g., Blakemore and Campbell, 1969) led a number of investigators toward a “Fourier view” of the cortex. Such work led to thousands of studies devoted to questions regarding frequency tuning and the relevance of this tuning to the human detection and discrimination of sinusoidal gratings. This left us with complex theories for how we detect gratings, but with little understanding of how such a system would function in the natural world.

Another example is the classification of V1 neurons into the categories of ‘simple’, ‘complex’, and ‘hypercomplex’ or ‘end-stopped’. Simple cells are noted for having oriented receptive fields organized into explicit excitatory and inhibitory subfields, whereas complex cells are tuned for orientation but are relatively insensitive to position and the sign of contrast (black-white edge vs. white-black edge). Hypercomplex cells display more complex shape selectivity and some appear most responsive to short bars or the terminations of bars of light (so-called “end-stopping”). Are these categories real, or a result of the particular way neurons were stimulated and the data analyzed?

A widely accepted theory that accounts for the distinction between simple and complex cells is that simple cells compute a (mostly linear) weighted sum of image pixels, whereas complex cells compute a sum of the squared and half-rectified outputs of simple cells of the same orientation—i.e., the so-called “energy model” (Adelson & Bergen, 1985). This theory is consistent with measurements of response modulation in response to drifting sinewave gratings, otherwise known as the “F1/F0 ratio” (Skottun et al., 1991). From this measure one finds clear evidence for a bimodal distribution of neurons, with simple-cells having ratios greater than one, and complex-cells having ratios less than one. Recently, however, it has been argued that this particular nonlinear measure tends to exaggerate or even introduce bimodality rather than reflecting an actual, intrinsic property of the data (Mechler & Ringach, 2002). When receptive fields are instead characterized by the degree of overlap between zones activated by increments or decrements in contrast, one obtains a continuous, unimodal distribution when the overlap is expressed as the normalized distance between the zones, but a bimodal distribution when expressed as an overlap index (sum of widths minus the separation divided by sum of widths plus the separation) (Mata et al.,

2005; Kagan et al., 2002). In addition, the energy model of complex cells does a poor job accounting for complex cells with a partial overlap of activating zones. Thus, the way in which response properties are characterized can have a profound effect on the resulting theoretical framework that is adopted to explain the results. The notion of two classes of neurons, “simple” and “complex,” has been firmly planted in the minds of modelers and experimentalists alike, but a closer examination of the data reveals that this classification scheme could actually be an artifact of the lens through which we view the data.

The notion of “end-stopped” neurons introduces even more questions when one considers the structure of natural images. Most natural scenes are not littered with line terminations or short bars—see for example Figure 3a. Indeed, at the scale of a V1 receptive field, the structures in this image are quite complex and they defy the simple, line drawing-like characterization of a “blocks world.” Where in such an image would one expect an end-stopped neuron to fire? By asking this question, one could possibly be led to a more ecologically relevant theory of these neurons than suggested by simple laboratory stimuli.

Another theory bias often embedded in investigations of V1 function is the notion that simple cells, complex cells, and hypercomplex cells are actually coding for the presence of edges, corners, or other 2D shape features in images. However, much of this thinking is derived from a rather ‘cartoon’ view of images. Computer vision studies provide clear evidence of the fallacy of the purely bottom up approach. One can not compute the presence even of simple edges of an object purely from the luminance discontinuities (i.e., using a filter such as a simple or complex cell model). As an example, Figure 3 demonstrates the result of processing a natural scene with the standard energy-model of a complex cell. Far from making contours explicit, this representation creates a cluttered array of orientation signals that make it difficult to discern what is actually going on in the scene. **Our perception of crisp contours, corners, and junctions in images is largely a posthoc phenomenon that is the result of massive *inferential* computations performed by the cortex, which are heavily informed by context and high-level knowledge.** It could well be that our initial introspections about scene structure are a poor guide as to the actual problems faced by the cortex.

In order to properly understand V1 function, our theories will need to be guided by functional considerations and an appreciation for the ambiguities contained in natural images, rather than being biased by simplistic notions of feature detection that are suggested by the responses of a select population of neurons recorded using simplified stimuli. One of the most challenging problems facing the cortex is that of inferring a representation of 3D surfaces from the 2D image (Nakayama et al. 1995; see also sec. 3.4 below). This is not an easy problem to solve, and it still lies beyond the abilities of modern computer vision. **It seems quite likely that V1 plays a role in solving this problem, but understanding how it does so will require going beyond bottom-up filtering models to consider how top-down information is utilized in the interpretation of images** (Olshausen, 2003; Lee & Mumford, 2003; see also sec. 3.5 below).

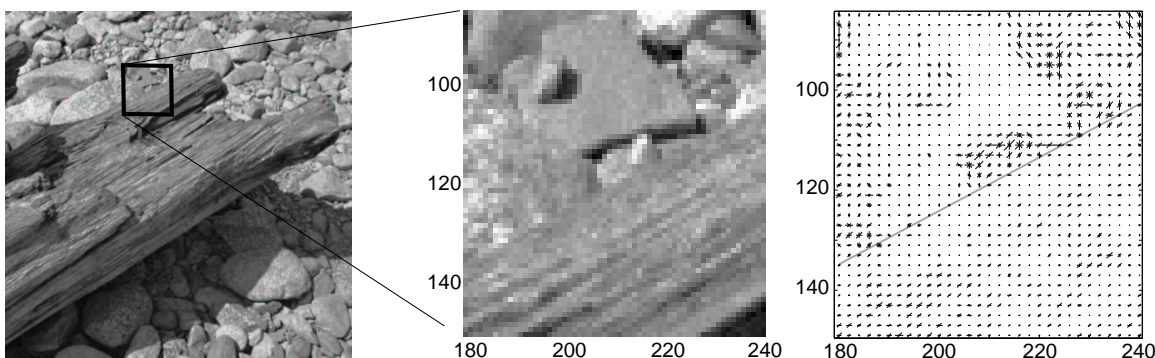


Figure 3: A natural scene (left), and an expanded section of it (middle). Far right shows the information conveyed by an array of model complex cells at four different orientations. The length of each line indicates the strength of response of a model complex cell at that location and orientation. The solid gray line shows the location of the boundary of the log in the original image. Note that very few complex cells of the appropriate orientation are responding along this contour.

2.4 Interdependence and contextual effects

It has been estimated that roughly 5% of the excitatory input in layer 4 of V1 arises from the LGN, with the majority resulting from intracortical inputs (Peters & Payne 1993; Peters et al. 1994). Thalamocortical synapses have been found to be stronger, making them more likely to be effective physiologically (Ahmed et al., 1994). Nevertheless, based on visually evoked membrane potentials, Chung and Ferster (1998) have argued that the geniculate input is responsible for just 35% of a layer 4 neurons response. This leaves 65% of the response determined by factors outside of the direct feedforward input. Using optical imaging methods, Arieli et al. (1996) showed that the population ongoing activity can account for 80% of an individual V1 neuron’s response variance, and recent work using multi-electrode arrays has shown that the ongoing activity V1 neurons is only slightly modified by visual input (Fiser et al., 2004). Thus, we are left with the real possibility that somewhere between 60-80% of the response of a V1 neuron is a function of other V1 neurons, or inputs other than those arising from LGN.

It should also be noted that recent evidence from the early blind has demonstrated that primary visual cortex has the potential for a wide range of multi-modal input. Sadato et al (1996) and Amedi et al (2003) demonstrated that both tactile Braille reading and verbal material can activate visual cortex in those that have been blind from an early age, even though no such activation occurs in those with normal sight. This implies that in the normal visual system, primary visual cortex has the potential for interactions with quite high-level sources of information.

That V1 neurons are influenced by context—i.e., the spatio-temporal structure outside of the classical receptive field (CRF)—is by now well known and has been the subject of many investigations over the past decade. Knierim & Van Essen (1992)

showed that many V1 neurons are suppressed by a field of oriented bars outside the classical receptive field of the same orientation, and Sillito et al. (1995) have shown that one can introduce quite dramatic changes in orientation tuning based on the orientation of gratings outside the CRF. Other investigators have probed the spatial specificity of the surround using grating patches and demonstrated fairly specific zones of suppression (Walker et al. 1999; Cavanagh et al. 2002b). And these studies, in addition to others (see Series et al. (2003) for a review), have likely tapped only a portion of the **interdependencies and contextual effects** that actually exist.

The problem in teasing apart contextual effects in such a piecemeal fashion is that one faces a combinatorial explosion in the possible spatial and featural configurations of surrounding stimuli such as bars or gratings. What we *really* want to know is how neurons respond within the sorts of context encountered in natural scenes. For example, given the results of Knierim & Van Essen (1993) using bar stimuli, or Sillito et al. (1995) using gratings, what should we reasonably expect to result from the sorts of context seen in the natural scene of Figure 3? Indeed, it is not even clear whether one can answer the question since the contextual structure here is so much richer and more diverse than that which has been explored experimentally. Some of the initial studies exploring the role of context in natural scenes have demonstrated pronounced nonlinear effects that tend to sparsify activity in a way that would have been hard to predict from the existing reductionist studies (Vinje & Gallant, 2000). More studies along these lines are needed, and most importantly, we need to understand how and why the context in natural scenes produces such effects.

Another striking form of interdependence exhibited by V1 neurons is in **the synchrony of activity**. Indeed, **the fact that one can even measure large-scale signals such as the local field potential or EEG implies that large numbers of neurons must be acting together**. Gray et al. (1989) demonstrated gamma-band synchronization between neurons in cat V1 when bars moved through their receptive fields in similar directions, suggesting that **synchrony is connected to a binding or segmentation process**. More recently, Wörgötter et al. (1998) have shown that receptive field sizes change significantly with the degree of synchrony exhibited in the EEG, and Maldonado et al. (2004) have shown that **periods of synchronization preferentially occur during periods of fixation as opposed to during saccades or drifts**. However, what role synchrony plays in the normal operation of V1 neurons is entirely unclear, and it is fair to say that this aspect of response variance remains a mystery.

2.5 Ecological deviance

We have argued above for experiments that measure the responses of neurons in ecological conditions even when no model is capable of predicting the results—or we should say, “especially if no model can predict the results.” Publishing findings only in conditions when a particular model works would be poor science. It is important to know not only where the current models can successfully predict neural behavior, but also **under what conditions they break down and why**. And as we have emphasized above, it is most important to know how they fare under ecological conditions. If the current models fail to predict neural responses under such conditions, then the

literature should reflect this.

In the last few years, a number of labs have begun using natural scenes as stimuli when recording from neurons in the visual pathway (Dan et al. 1996; Baddeley et al. 1996; Keyser et al 2001; Vinje & Gallant 2002; Ringach et al., 2002; Smyth et al. 2003; David et al. 2004). In particular, the Gallant lab at UC Berkeley has taken the approach of attempting to determine how well one can predict the responses of V1 neurons to natural stimuli using a variety of different models. However, assessing how well these models fare, and what it implies about our current understanding of V1, is difficult for at least three reasons.

First, one must make **several assumptions (either implicitly or explicitly) regarding what aspects of the response are relevant to the model.** **Spike counts** will show **significant variability over repeated trials** (Tohurst et al., 1983). One can take the average over a number of presentations, but this implicitly assumes that the variability can be attributed to noise. This can be questioned, especially considering that in many cases individual spikes have been shown to have relatively high reliability (Rieke et al., 1997). The trial-to-trial variability could well be due to **internally generated dynamics** that plays an important role in information processing that we simply do not as yet understand (Arieli et al., 1996; Fiser et al., 2004; see also Section 3.1). Furthermore, to take averages, one must make assumptions regarding the temporal window over which the average is computed.

Secondly, these studies are best performed with an awake, behaving animal. In such conditions, there are limitations to the spatial and temporal accuracy with which the gaze can be measured. When averaging across presentations of stimuli, an assumption must be made as to whether the same stimulus was actually presented. Again, one must make an assumption as to what spatio-temporal window to use.

The third problem is that whatever model is chosen, one is **always subject to the criticism that the model is not sufficiently elaborate.** Thus, any inability to predict the neurons response might be argued to be simply due to some missing element in the model.

For example, David, Vinje & Gallant (2004) have explored two different types of models—a linearized spatiotemporal receptive field model, in which the neuron’s response is essentially a weighted sum of the image pixels over space and time, and a “phase-separated Fourier model” which allows one to capture the phase invariance non-linearity of a complex cell. These models can typically explain between 20-40% of the response variance. Correcting for inter-trial variability improves matters somewhat (David et al., 2004), and it is possible that with more trials, and with the addition of other non-linearities such as contrast normalization, adaptation, and response saturation, the fraction of variance explained could rise even more above these levels (and this is a current direction of these studies).

We believe such reports are critically important for several reasons. First, such results create a benchmark for showing how well the standard or basic models actually predict ecologically relevant data. Secondly, these are well-established models that have been given a fair run for their money. One could imagine any number of improvements to these models, and it will be interesting to see if they fare better, but in the meantime these results provide a useful baseline for comparison. Furthermore,

these are the data that represent the ultimate goal of any computational model, and so they are crucial to presenting a complete picture of V1 function. Given the nature of the errors, as discussed below, we do not believe that the addition of simple response nonlinearities such as contrast normalization are likely to improve matters much. Given these results with both linear and Fourier power models, we conjecture the best-case scenario is that the percentage of variance explained is likely to asymptote at 30-40% with the standard model.

One of the reasons for our pessimism is due to the way in which these models fail. For example, Figure 4 shows data collected from the lab of Charles Gray at Montana State University, Bozeman, in which the activities of V1 neurons in anaesthetized cat are recorded in response to repeated presentations of a natural movie (C.M. Gray, J. Baker, S.C. Yen, personal communication, 2004). Shown (Fig. 4a) is the peristimulus time histogram (PSTH) of a typical V1 simple cell, whose receptive field as measured from an M-sequence kernel is similar to those found in the literature—i.e., a Gabor-like function that translates over time (i.e., space-time inseparable). Superimposed on this is the predicted response generated by convolving the neurons space-time receptive field (Fig. 4b) with the movie, and putting the result through a point-wise non-linearity (including a gain factor and offset term). The neuron tends to exhibit sparse, punctate responses, some of which are predicted by the receptive field model, others not. In most cases the model response undershoots the PSTH, and this can not simply be addressed by increasing the gain or narrowing the response of the model, because there are many other episodes where the model predicts responses of equal magnitude in which there is little or no response from the neuron. One could possibly obtain a better fit to the data by including additional terms modeling suppression (Rust et al., 2004) and temporal adaptation (Lesica et al. 2003), or even a spiking mechanism (Paninski et al., 2004), but we believe it is useful to see how much the linear, driving term of the model alone fares under these circumstances. Moreover, these additions are essentially single-neuron mechanisms. What seems to be suggested by our initial informal observations of multiple simultaneously recorded units is that a more complex, network non-linearity is at work here, and that describing any one neuron’s behavior will require one to include the influence of other simultaneously recorded neurons.

An important lesson of these findings is that simply mapping out receptive fields does not provide a complete understanding of V1 response properties. For example, Ringach et al. (2002) have shown that it is possible to map out receptive fields using natural scenes, and they show that it is even possible to recover some non-linear effects such as cross-orientation inhibition with this technique. However, the resulting receptive field models were not tested by comparing their predictions to the actual activity of neurons in response to natural movies. Without doing so, it is difficult assess how well such models capture the function of the neuron.

Unfortunately, journals are often unprepared to publish results when a study demonstrates the failure of a model, unless the study also presents a competing model which works well. Part of this may seem understandable since there are a variety of reasons a model might fail. However, until a benchmark is placed into the literature, it is impossible to determine how ‘good a model actually is. And given the magnitude of

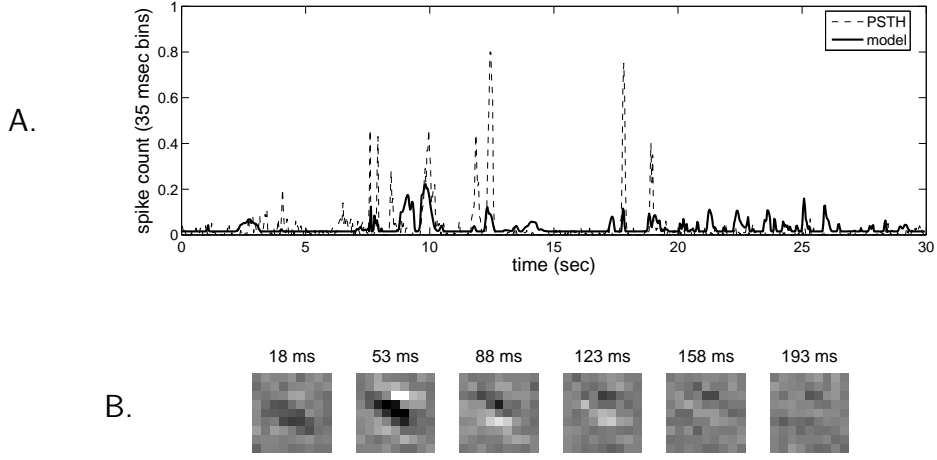


Figure 4: Activity of a V1 neuron in anaesthetized cat in response to a natural movie. A. The PSTH of the neuron’s response (dashed line), together with the predicted response (solid line) generated from the model: $\hat{r}(t) = \alpha h(\sum_{\mathbf{x}} k(\mathbf{x}, t) * I(\mathbf{x}, t) + \theta)^p + r_0$. The function $h()$ is a half-wave rectifying function, and the parameters α , p , θ , and r_0 are fit to minimize the squared error with the data. The resulting correlation coefficient in this case is 0.36. Average spike counts were obtained by averaging across 100 trials in 35 ms bins (corresponding to the frame rate). B. The kernel $k(\mathbf{x}, t)$ was measured via reverse correlation with an m-sequence, and is shown here as a series of frames in 35 ms intervals, with the center time of the interval displayed above each frame.

	Biased sampling	Biased stimuli	Biased theories	Interdepend. & context	Ecological deviance
Problem	large neurons; visually responsive neurons; neurons with high firing-rates	use of reduced stimuli such as bars, spots, & gratings	simple/complex cells; data-driven theories	influence of intra-cortical input; effect of context; synchrony	responses to natural scenes deviate from predictions of standard models
Solution	use chronically implanted electrodes, parallel recording arrays	use natural scenes, ecologically relevant stimuli	consider more functional/computational theories that solve problems of vision	examine how context affects responses in natural scenes	develop models that can account for responses to natural images

Table 1: Five problems with the current view of V1 and some possible solutions for obtaining a more complete picture.

the task before us, it could take years before a good model emerges. In the meantime, what would be most helpful is to accumulate a database of single unit or multi-unit data (stimuli and neural responses) that would allow modelers to test their best theory under ecological conditions.

Finally, it should be noted that better success has been obtained in using receptive field models to predict the responses of neurons to natural scenes in the LGN (Dan et al., 1996), or the response of cortical neurons to purely static images (Smyth et al., 2003), although they are still far from making perfect predictions. This would seem to suggest that much of the difficulty in predicting responses in cortex has to do with the effects of the massive, recurrent intra-cortical circuitry that is engaged during natural vision.

2.6 Summary

Table 1 presents a summary of the five problems we have identified with the current view of V1 that has emerged from the data collected to date, along with some of the solutions that we have suggested could possibly help in obtaining a more complete picture of V1 function.

Given the limitations described above, is it possible to quantify how well we currently understand V1 function? We attempt to estimate this as follows:

$$[\text{fraction understood}] = \left[\frac{\text{fraction of variance explained from neurons}}{\text{recorded}} \right] \times \left[\frac{\text{fraction of population}}{\text{recorded}} \right]$$

If we consider that roughly 40% of the population of neurons in V1 has actually been recorded from and characterized, together with our conjecture that 30-40% of the response variance of these neurons can be explained under natural conditions using the currently established models, then we are left to conclude that we can currently account for 12-16% of V1 function. Thus, approximately 85% of V1 function has yet to be explained (see Figure 5).²

²We have primarily drawn upon the Gallant lab’s data for obtaining the percentage

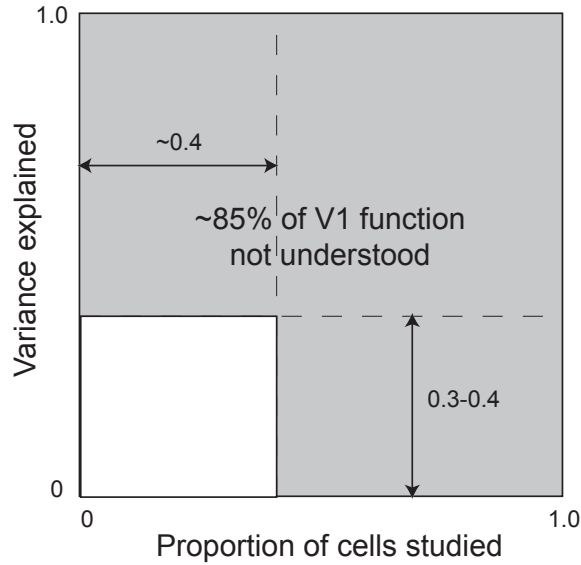


Figure 5: 85% of V1 function remains to be understood.

3 New theories

Given the above observations, it becomes clear that there is so much unexplored territory that it is very difficult to rule out theories at this point (although there are some obvious bounds dictated by neural architecture—e.g., fan-in/fan-out, the spatial extent of axonal and dendritic arbors, etc.). In the sections below, we discuss some of the theories that are plausible given our current data. However, the goal here is not to provide a detailed review of the theories currently in the literature. Rather, the goal is to provide a few examples of the range of theories that are consistent with the experimental data. It must be emphasized that considering that there may exist a large family of neurons with unknown properties, and given the low level of prediction for the neurons studied, there is still considerable room for theories dramatically different than those theories presented here.

3.1 Dynamical systems and the limits of prediction.

Imagine tracking a single molecule within a hot gas as it interacts with the surrounding molecules. The particular trajectory of one molecule will be erratic and fundamentally unpredictable without knowledge of all other molecules with potential influence. Even if we presumed the trajectory of the particular molecule was completely deterministic and following simple laws, in a gas with large numbers of interacting molecules one could never provide a prediction of the path of a single molecule except over very short distances.

of variance explained, and so we are assuming that their methods for isolating neurons are subject to the same biases in sampling discussed earlier.

In theory, the behavior of single neurons may have similar limitations. To make predictions of what a single neuron will do in the presence of a natural scene may be fundamentally impossible without knowledge of the surrounding neurons. The non-linear dynamics of interacting neurons may put bounds on how accurately the behavior of any neuron can be predicted. And at this time, we cannot say where that limit may be.

What is fascinating in many ways then is that neurons are as predictable as they are. For example, work from the Gallant lab has shown that under conditions where a particular natural scene sequence is repeated to a fixating macaque monkey, a neuron’s response from trial-to-trial is fairly reliable (e.g., Vinje & Gallant, 2000). This clearly suggests that the response is dependent in large part on the stimulus, certainly more than a molecule in the “gas model.” So how do we treat the variability that is not explained by the stimulus? We may find that the reliability of a local group of neurons is more predictable than a single neuron, which would then require multi-electrode recording to attempt to account for the remaining variance. For example, Arieli et al. (1996) have shown that much of the inter-trial variability may be explained in terms of large-scale fluctuations in ongoing activity of the surrounding population of neurons measured using optical recording, and Fiser et al. (2004) have similarly shown that ongoing population activity as measured with multi-electrode arrays is only loosely modulated by visual input. However, what role these large-scale fluctuations play in the normal processing of natural scenes has yet to be investigated.

3.2 Sparse, overcomplete representations

One effort to explain many of the non-linearities found in V1 is based on the idea that neurons are attempting to achieve some degree of gain control (Geisler & Albrecht 1992). Because any single neuron lacks the dynamic range to handle the range of contrasts in natural scenes, it is argued, the contrast response must be normalized. Here we provide a different line of reasoning to explain the observed response non-linearities of V1 neurons (further details are provided by Olshausen & Field, 1997, and Field & Wu, 2004). We argue that the spatial non-linearities primarily serve to reduce the linear dependencies that exist in an overcomplete code, and as we shall see this leads to a fundamentally different set of predictions about the population activity.

Consider the number of vectors needed to represent a particular set of data with dimensionality D (e.g., an 8×8 pixel image patch would have $D = 64$). No matter what form the data takes, such data never requires more than D linearly independent vectors to represent it. A system where data with dimensionality D is spanned by D vectors is described as “critically sampled.” Such critically sampled systems (e.g., orthonormal bases) are popular in the image coding community as they allow any input pattern to be represented uniquely, and the transform and its inverse are easily computed. The wavelet code, for example, has seen widespread use, and wavelet-like codes similar to that of the visual system have been shown to provide very high efficiency, in terms of sparsity, when coding natural scenes (e.g., Field, 1987). Some basic versions of ICA also attempt to find a critically sampled basis which minimizes

the dependencies among the vectors, and the result is a wavelet-like code with tuning much like the neurons in V1 (Bell and Sejnowski, 1997; van Hateren & van der Schaaf 1998).

However, the visual system is not using a critically sampled code. In cat V1, for example, there are 25 times as many output fibers as there are input fibers from the LGN, and in macaque V1 the ratio is on the order of 50:1. Such overcomplete codes have one potential problem: the vectors are not linearly independent. Thus, if neurons were to compute their output simply from the inner-product between their weight vector and the input, their responses will be correlated.

Figure 6a shows an example of a two-dimensional data space represented by three neurons with linearly dependent weight vectors. Even assuming the outputs of these units are half-rectified so they produce only positive values, the data are redundantly represented by such a code. The only way to remove this linear dependence is through a non-linear transform. One of the non-linear transforms that will serve this goal is shown in Figure 6b. Here, we show the iso-response curves for the same three neurons. This curvature represents an unusual non-linearity. For example, consider the responses of a unit to two different stimuli: the first stimulus aligned with the neuron’s weight vector, and a second stimulus separated by 90 degrees. The second stimulus will have no effect on the neuron on its own since its vector is orthogonal to that of the neuron. However, when added to the first vector, the combined stimulus will be on a lower iso-response curve (i.e. the neuron will have reduced its activity). In other words, the response curvature of the neuron results in a non-linearity with the characteristic “non-classical,” suppressive behavior: Stimuli which on their own have no affect on the neuron (stimuli orthogonal to the principal direction of the neuron), can modulate the behavior of an active neuron. This general non-linearity comes in several forms and includes end-stopping and cross orientation inhibition, and is what is typically meant by the term “non-classical surround.” Indeed, as Zetzsche et al (1999) note, this curvature is simply a geometric interpretation of such behaviors. With the addition of a compressive non-linearity this curvature results in the behavior described as “contrast normalization.”

In contrast to the gain control or divisive normalization theory, we argue that the non-linearities observed in V1 neurons are present primarily to allow a large (overcomplete) population of neurons to represent data using a small number of active units, a process we refer to as “sparsification.” The goal is not to develop complete independence, as the activity of any neuron partially predicts the lack of activity in neighboring neurons. However, the code allows for expanding the dimensionality of the representation without incurring the linear dependencies that would be present in a non-orthogonal code.

Importantly, this model predicts that the non-linearities are a function of the angle between the neuron’s weight vector and those surrounding it. Future multi-electrode recordings may provide the possibility to test this theory. From the computational end, we have found that our sparse coding network (Olshausen & Field, 1996; 1997) produces non-linearities much like those proposed. Our hope, then, is that many of the non-linearities that have been observed in V1 can eventually be explained within one general framework of efficient coding.

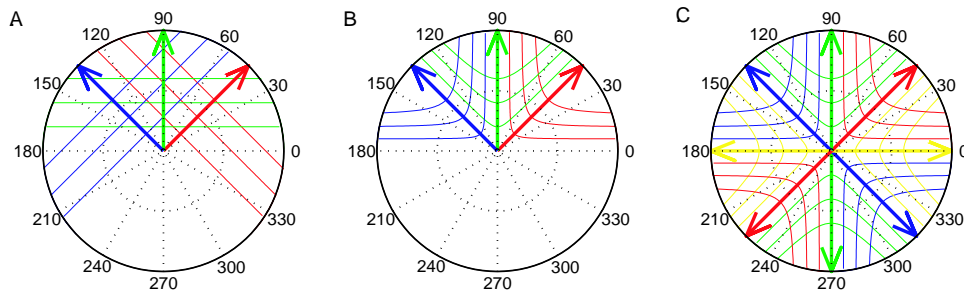


Figure 6: Overcomplete representation. A. Shown are the iso-response contours of three linear neurons (with half-wave rectification) having linearly dependent weight vectors. A stimulus falling anywhere along a given contour will result in the same response from the neuron. A stimulus falling in the upper half-plane will result in responses on all three neurons, even though only two would be required to uniquely determine its position in the space. B. Curving the response contours removes redundancy among these neurons. Now only two neurons will code for a stimulus anywhere in this space. C. A full tiling of the 2D stimulus space now requires eight neurons, which would be overcomplete as a linear code, but critically sampled given this form of non-linear response.

3.3 Contour integration

There is now considerable physiological and anatomical evidence showing that V1 neurons have a rather selective connection pattern both within layers and between layers. For example, research investigating the lateral projections of pyramidal neurons in V1 has shown that the long range lateral connections project primarily to regions of the cortex with similar orientation columns, as well as to similar ocular dominance columns and cytochrome oxidase blobs (Malach et al., 1993; Yoshioka et al., 1996). Early studies exploring the horizontal connections in V1 discovered that selective long range connections extend laterally for 2 to 5 mm parallel to the surface (Gilbert and Wiesel, 1979), and studies on the tree shrew (Rockland and Lund, 1982; Bosking et al., 1997), primate (e.g., Malach et al., 1993; Sincich & Blasdel, 2001), ferret (Ruthazer & Stryker, 1996), and cat (e.g., Gilbert & Wiesel, 1989) have all demonstrated significant specificity in the projection of these lateral connections. A number of neurophysiological studies also show that co-linearly oriented stimuli presented outside of the classical receptive field have a facilitatory effect (Kapadia et al., 1995; Kapadia et al., 2000; Polat et al. 1998). The results demonstrate that when a neuron is presented with an oriented stimulus within its receptive field, a second collinear stimulus will sometimes increase the response rate of the neuron while the same oriented stimulus presented orthogonal to the main axis of orientation (displaced laterally) will produce inhibition, or at least less facilitation.

These results suggest that V1 neurons have an orientation- and position-specific connectivity structure, beyond what is usually included in the standard model. One line of research suggests that this connectivity helps resolve the ambiguity of contours in scenes and is involved in the process of contour integration (e.g., Field et al 1993).

This follows from work showing that the amplification of locally co-aligned, oriented elements provides an effective means of identifying contours in natural scenes (Parent & Zucker, 1989; Sha’ashua & Ullman, 1988; Ben-Shahar & Zucker 2004). This type of mechanism could work in concert with the sparsification non-linearities mentioned above, since the facilitatory interactions would primarily occur among elements that are non-overlapping—i.e., receptive fields whose weight vectors are orthogonal.

An alternative theoretical perspective is that the effect of these orientation- and position-specific connections should be mainly suppressive, with the goal of removing dependencies among neurons that arise due to the structure in natural images (Schwartz & Simoncelli, 2001). In contrast to the contour integration hypothesis, which proposes that the role of horizontal connections is to *amplify* the structure of contours, this model would attempt to *attenuate* the presence of such structure in the V1 representation. Although this may be a desirable outcome in terms of redundancy reduction, we would argue that the cortex has objectives other than redundancy reduction per se (Barlow, 2001). Chief among these is to provide a meaningful representation of image structure that can be easily read out and interpreted by higher-level areas.

Finally it is important to note, with respect to the discussion in the previous section, that the type of redundancy we are talking about here is due to long-range structure in images beyond the size of a receptive field, not that which is simply due to the overlap among receptive fields. Thus, we propose that the latter should be removed via sparsification, while the former should be amplified by the long-range horizontal connections in V1.

3.4 Surface representation

We live in a three-dimensional world, and the fundamental causes of images that are of behavioral relevance are *surfaces*, not two-dimensional features such as spots, bars, edges, or gratings. Moreover, we rarely see the surface of an object in its entirety. Occlusion is the rule, not the exception, in natural scenes. It thus seems quite reasonable to think that the visual cortex has evolved effective means to parse images in terms of the three-dimensional structure of the environment—i.e., surface structure, foreground/background relationships, etc. Indeed, there is now a strong body of psychophysical evidence showing that 3D surfaces and figure-ground relationships constitute a fundamental aspect of intermediate-level representation in the visual system (Nakayama et al., 1995; see also Figure 7).

Nevertheless, it is surprising how little V1 physiology has actually been devoted to the subject of three-dimensional surface representation. Some recent studies in extrastriate cortex have begun to yield interesting findings (Nguyenkim & DeAngelis, 2003; Zhou et al., 2000; Bakin et al., 2000), but V1’s involvement in surface representation remains a mystery. Although many V1 neurons are disparity selective, this by itself does not tell us how surface structure is represented, nor how figure-ground relationships of the sort depicted in Figure 7 are resolved.

At first sight it may seem preposterous to suppose that V1 is involved in computing three-dimensional surface representations. But again, given how little we actually do

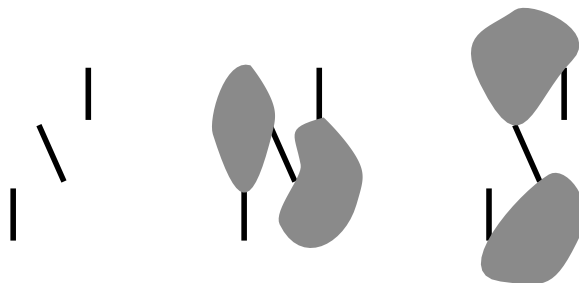


Figure 7: The three line-strokes at left are interpreted as different objects depending on the arrangement of occluders. Thus, pattern completion depends on resolving figure-ground relationships. At what level of processing is this form of completion taking place? Since it would seem to demand access to high-resolution detail in the image, it can not simply be relegated to high-level areas.

know about V1, combined with the importance of 3D surface representations for guiding behavior, it is a plausible hypothesis to consider. In addition, problems such as occlusion demand resolving figure-ground relationships in a relatively high-level representation where topography is preserved (Lee & Mumford, 2003). There is now beginning to emerge physiological evidence supporting this idea. Neurons in V1 have been shown to produce a differential response to the figure vs. background in a scene of texture elements (Lamme 1995; Zipser et al., 1996), and a substantial fraction of neurons in V1 are selective to border ownership (Zhou, 2000). In addition, Lee et al. (1998) have demonstrated evidence for a medial axis representation of surfaces in which V1 neurons become most active along the skeletal-axis of an object. It seems quite possible such effects are just the tip of the iceberg, and there could be even more effects lurking.

3.5 Top-down feedback and disambiguation

Although our perception of the visual world is usually quite clear and unambiguous, the raw image data that we start out with is not. Looking back at Figure 3, one can see that even the presence of a simple contour can be ambiguous in a natural scene. The problem is that information at the local level is insufficient to determine whether a change in luminance is due to an object boundary, simply part of a texture, or a change in reflectance. Although boundary junctions are also quite crucial to the interpretation of a scene, a number of studies have now shown that human observers are poor judges of what constitutes a boundary or junction when these features are shown in isolation (Elder et al., 1999; McDermott, 2004). Thus, the calculation of what forms a boundary is dependent on the context, which provides information about the assignment of figure and ground, surface layout, and so forth.

Arriving at the correct interpretation of an image, then, constitutes something of a chicken-egg problem between lower and higher levels of image analysis. The low-level shape features that are useful for identifying an object—edges, contours,

surface curvature and the like—are typically ambiguous in natural scenes, so they cannot be computed directly based on a local analysis of the image. Rather, they must be inferred based on global context and higher-level knowledge. However, the global context itself will not be clear until there is some degree of certainty about the presence of low-level shape features. A number of theorists have thus argued that recognition depends on information circulating through cortico-cortical feedback loops in order to disambiguate representations at both lower and higher levels in parallel (Mumford 1994; Ullman 1995; Lewicki & Sejnowski 1996; Rao & Ballard 1999; Young 2000; Lee & Mumford 2003; Hawkins & Blakeslee 2004).

An example of disambiguation at work in the visual cortex can be seen in the resolution of the aperture problem in computing the direction of motion. Because receptive fields limit the field of a view of a neuron to just a portion of an object, it is not possible for any one neuron to signal with certainty the true direction of the object in a purely bottom up fashion. Pack, Berezovskii & Born (2001) have shown that the initial phase of response of neurons in MT signals the direction of motion directly orthogonal to a contour, and that the latter phase of the response reflects the actual direction of the object that the contour is part of, presumably from the interaction with other neurons viewing other parts of the object. Interestingly, this effect does not occur under anaesthesia. A similar delayed response effect has been demonstrated in end-stopped V1 neurons as well (Pack et al., 2003).

Recent evidence from fMRI points to a disambiguation process occurring in V1 during shape perception (Murray et al., 2002). Subjects viewed a translating diamond that was partially occluded so that the vertices are invisible, resulting in a bistable percept in which the line segments forming the diamond are seen moving independently in one case, and coherently in the direction of the object motion in the other case. When subjects experience the coherent motion and shape percept, activity in the lateral occipital complex (LOC) increases while activity in V1 decreases. This is consistent with the idea that when neurons in LOC are representing the diamond, they feed back this information to V1 so as to refine the otherwise ambiguous representations of contour motion. If the refinement of activity attenuates the many incorrect responses while amplifying the few that are consistent with the global percept, the net effect could be a reduction as seen in the BOLD signal measured by fMRI. An alternative interpretation for the reduction in V1 is based on the idea of predictive coding (Rao & Ballard 1999), in which higher areas actually subtract their predictions from lower areas.

There exists a rich set of feedback connections from higher levels into V1, but little is known about the computational role of these connections. Recent experiments in which higher areas are cooled to look at the effect upon activity in lower areas seem to suggest that these connections play a role in enhancing the salience of stimuli (Hupe et al., 1998), and Shapley (2004) has concluded that top-down feedback is necessary to account for the spatial extent of surround inhibition. But we would argue that feedback has a far more important role to play in disambiguation, and as far as we know, no one has yet investigated the effect of feedback using such cooling techniques under normal conditions that would require disambiguation (e.g., natural scenes).

3.6 Dynamic routing

A challenging problem faced by any visual system is that of forming object representations that are invariant to position, scale, rotation, and other common deformations of the image data. The currently accepted, traditional view is that complex cells constitute the first stage of invariant representation by summing over the outputs of simple-cells whose outputs are half-rectified and squared—i.e., the classical “energy model” (Adelson & Bergen 1985). In this way, the neuron’s response changes only gradually as an edge is passed over its receptive field. This idea forms the basis of so-called “Pandemonium models,” in which a similar feature extraction and pooling process is essentially repeated at each stage of visual cortex (see Tarr (1999) for a review).

However, the Pandemonium model can not provide a complete account of perception because it does not preserve information about relative phase or the spatial relationships among features. Clearly, though, we have conscious access to this information. The ability to navigate, grasp, and interact with foreign objects implies that we have the ability to perceive spatial relationships among features without ever doing “object recognition.” In addition, resolving figure/ground relationships and occlusion demands that higher levels of analysis have access to information about spatial relationships as well.

One of us has proposed a model for forming invariant representations that preserves relative spatial relationships by explicitly *routing* information at each stage of processing (Olshausen et al. 1993). Rather than passively pooling, information is dynamically linked from one stage to the next by a set of control neurons that progressively remap information into an object-centered reference frame. It is thus proposed that there are two distinct classes of neurons—those conveying image/feature information, and those controlling the flow of information. The former corresponds to the invariant part, the latter to the variant part. The two are combined multiplicatively, so that mathematically it is equivalent to a bilinear model (e.g., Tenenbaum & Freeman, 2000; Grimes & Rao, 2005).

Is it possible that dynamic routing occurs in V1 and underlies the observed shift-invariant properties of complex cells? If so, there are at least two things we would expect to see: 1) that at any given moment a complex cell is effectively connected to only one or a small fraction of simple cells to which it is physically connected, and 2) that there are *control neurons* which dynamically gate these connections. Interestingly, the observed invariance properties of complex cells are just as consistent with the idea of routing as they are with pooling. What could possibly distinguish between these models is to look at the population activity: if the complex cell outputs are the result of passive pooling, then one would expect a dense, distributed representation of contours among the population of complex cells. If information is dynamically routed, though, the representation at the complex cell level would remain sparse. The control neurons, on the other hand, would look something like contrast normalized simple cells, which represent phase independent of magnitude (Zetzsche & Rohrbein, 2001).

One of the main predictions of the dynamic routing model is that the receptive fields of the invariant neurons would be expected to shift depending on the state of

the control neurons. Such effects have been seen in V4, where some neurons shift their receptive fields depending on where the animal is directing its attention (Moran & Desimone, 1985; Connor et al., 1997). And in V1, Brad Motter has shown that neurons appear to shift their receptive fields in order to compensate for the small eye movements that occur during fixation (Motter & Poggio, 1990; Motter, 1995), although Gur & Snodderly (1997) provide evidence to the contrary. Thus, there exists some evidence for dynamic routing in visual cortex, but further experiments are needed in order to characterize how and to what extent this occurs in V1 under normal viewing conditions.

4 Conclusions

Our goal in this article has been to point out that there are still substantial gaps in our knowledge of V1 function, and more importantly, that there is more room for new theories to be considered than the current conventional wisdom might allow. We have identified five specific problems with the current view of V1, emphasizing the need for using natural scenes in experiments, in addition to multi-unit recording methods, in order to obtain a more representative picture of V1 function. While the single-unit, structuralist approach has been a useful enterprise for getting a handle on basic response properties, we feel that its usefulness as a tool for investigating V1 function has been nearly exhausted. It is now time to dig deeper, using richer, ecologically relevant experimental paradigms, and developing theories that can help to elucidate how the cortex performs the computationally challenging problems of vision.

As we explore the response properties of V1 neurons using natural scenes, we are likely to uncover some interesting new phenomena that defy explanation with current models. It is at this point that we should be prepared to revisit the structuralist approach in order to tease apart what is going on. Reductionism does have its place, but it needs to be motivated by functionally and ecologically relevant questions, similar to the European tradition in ethology (Tinbergen, 1972).

At what point will we actually understand V1? This is obviously a difficult question to answer, but we believe at least three ingredients are required: 1) an unbiased sample of neurons of all types, firing rates, and layers of V1, 2) the ability to observe simultaneously the activities of hundreds of neurons in a local population, and 3) the ability to predict, or at least qualitatively model, the responses of the population under natural viewing conditions. Given the extensive feedback connections into V1, in addition to the projections from pulvinar and other sources, it seems unlikely that we will ever understand V1 in isolation. Thus, our investigations must also be guided by how V1 fits into the bigger picture of thalamo-cortical function.

5 Acknowledgments

We thank Bill Skaggs for discussions on hippocampal physiology, Charlie Gray and Jonathan Baker for sharing preliminary data, Jack Gallant for clarifying the issues involved in predicting neural responses, and Jeff Johnson and Issac Trotts for comments

on the manuscript. We also thank the two anonymous reviewers for providing many useful suggestions and pointers to relevant literature. Supported by NGIA grant HM 1582-05-C-0007 to DJF.

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