Variability and Information in a Neural Code of the Cat Lateral Geniculate Nucleus

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Liu, Robert C., Svilen Tzonev, Sergei Rebrik, and Kenneth D. Miller. Variability and information in a neural code of the cat lateral geniculate nucleus. J Neurophysiol 86: 2789–2806, 2001. A central theme in neural coding concerns the role of response variability and noise in determining the information transmission of neurons. This issue was investigated in single cells of the lateral geniculate nucleus of barbiturate-anesthetized cats by quantifying the degree of precision in and the information transmission properties of individual spike train responses to full field, binary (bright or dark), flashing stimuli. We found that neuronal responses could be highly reproducible in their spike timing (≈1–2 ms standard deviation) and spike count (≈0.3 ratio of variance/mean, compared with 1.0 expected for a Poisson process). This degree of precision only became apparent when an adequate length of the stimulus sequence was specified to determine the neural response, emphasizing that the variables relevant to a cell’s response must be controlled to observe the cell’s intrinsic response precision. Responses could carry as much as 3.5 bits/spike of information about the stimulus, a rate that was within a factor of two of the limit the spike train could transmit. Moreover, there appeared to be little sign of redundancy in coding: on average, longer response sequences carried at least as much information about the stimulus as would be obtained by adding together the information carried by shorter response sequences considered independently. There also was no direct evidence found for synergy between response sequences. These results could largely, but not entirely, be explained by a simple model of the response in which one filters the stimulus by the cell’s impulse response kernel, thresholds the result at a fairly high level, and incorporates a postspike refractory period.

INTRODUCTION

To understand the coding of information by neurons, it is important to quantify the variability in their responses. When this variability is driven by changes in the stimulus, the neurons can use this to distinguish between stimuli. On the other hand, when this variability occurs in repeated responses to the same stimulus, it acts as noise that reduces the neurons’ potential capacity to code information.

The study of neuronal variability has recently seen a rebirth of interest in association with the renewed use of information-theoretic techniques for analyzing neural coding (Bair 1999; Borst and Theunissen 1999; Buracas and Albright 1999; de Ruyter van Steveninck et al. 1997; Meister and Berry 1999; Rieke et al. 1997; Victor 1999). In the visual system, the precision of spike times and counts has been investigated in several neural areas, although only a few have looked at the lateral geniculate nucleus (LGN) (Guido and Sherman 1998; Hartveit and Heggelund 1994; Kara et al. 2000; Keat et al. 2001; Reich et al. 1997; Reinagel and Reid 2000; Sestokas and Lehmkuhle 1988). In this paper, we further explore the degree of precision found in LGN neurons of barbiturate-anesthetized cat by examining both spike count and timing measures. We go on to quantify the amount of information transmitted by neurons about the stimulus and to determine the degree to which models of response based on linear integration of inputs can account for the observed precision.

A unique feature of the present approach is that we closely examined the dependence of neuronal variability on the degree of specification of the stimulus. To do this, we employed a pseudorandom binary stimulus known as an M-sequence (Sutter 1992). We focused only on characterizing the neurons’ response to temporally varying stimuli by showing full-field bright and dark frames, ignoring the center-surround spatial structure of LGN neurons. M-sequences provide a statistically efficient and convenient method for analyzing responses because they have the nice property that every sequence of bright and dark frames of a given length (up to some limit) is repeated the same number of times somewhere throughout the sequence (see METHODS). This allowed us to simultaneously examine the responses—both the mean response and the variability in the response—to every sequence of a given length, giving us a detailed characterization of the neural code for such sequences. By varying this length, we examined how much of the stimulus had to be specified to maximize the precision of a neuron’s response: e.g., if the neuron’s response was influenced by the last 10 frames and only 5 frames were specified, then the response would be averaged over the unspecified frames, causing the neuron’s responses to appear more variable than they would be if the stimulus were fully specified. The variability remaining when the stimulus was fully specified reflected the neuron’s intrinsic response variability.

It is common to characterize a cell’s response by its linear temporal kernel, which—as computed from an M-sequence stimulus and neglecting normalization (see METHODS)—is the difference between its mean response to a single bright frame and its mean response to a single dark frame. We found that...
average responses to a single bright or dark frame within a sequence showed Poisson-like spike count variability and temporal dispersion over tens of milliseconds, and the kernel was correspondingly temporally broad. But by specifying more of the stimulus—e.g., specifying eight consecutive frames—the response could become far more precise, with sub-Poisson spike count variability and temporal precision of 1–2 ms. The information conveyed by the neuron correspondingly increased, containing as much as 3.5 bits/spike about longer stimulus sequences. We found that this information depended on the specification of spike times down to 1-ms resolution and that the information in consecutive spikes showed little redundancy or synergy. Finally, we determined that the precision obtained when multiple frames were specified could be largely, but not entirely, explained if the spike rate arose from a filtering of the stimulus by the cell’s temporal kernel followed by thresholding, along with imposition of a postspike refractory period.

Some of this work was previously presented in abstract form (Liu et al. 2000; Tzonev et al. 1997).

METHODS

Experiments

We performed experiments on adult cats under a protocol approved by the University of California, San Francisco Committee on Animal Research. Cats were initially anesthetized with isoflurane (1–5%), and placed on a feedback-controlled heating pad to maintain body temperature at 37.5–38°C. We established an intravenous line and thereafter maintained anesthesia via thiopental sodium or pentobarbital sodium (the latter was given once anesthesia was stable). The heart rate, respiratory rate, core temperature, O2 saturation, expiratory CO2, and lung pressure were all continually monitored. After performing a tracheotomy, the animal was respirated with nitrous oxide in a 1:1 ratio with oxygen. We performed a craniotomy, and then paralyzed the animal by infusing gallamine (10 mg \( \cdot \) kg\(^{-1} \cdot \) h\(^{-1} \)) in lactated Ringers. The electroencephalogram (EEG) was subsequently monitored continuously. We reflected the optic disk onto a white background using a fiber optic light source, and inserted contact lenses to focus the eyes at a distance of 35–40 cm.

We recorded extracellularly using tetrodes (Gray et al. 1995) advanced through a guide tube inserted to within a few millimeters of the topography across repeated penetrations to published accounts.

Stimulus

For visual stimulation, sequences of full-field bright and dark frames were presented on a computer monitor at the rate of 120 Hz, yielding a frame duration of \( t_f \approx 8.3 \) ms. Each frame varied randomly between bright or dark, with a photopic mean luminance; contrast [measured as \( (L - D)/(L + D) \) where \( L \) and \( D \) were the luminances of bright and dark frames, respectively] for each full sequence was chosen from 6, 14, 20, 40, or 80%.

We generated random frames using a binary M-sequence, which is essentially a stream of pseudorandom bits having some special properties (see following text). A bit value of 1 corresponded to a bright frame, and 0 corresponded to a dark frame. An M-sequence of order \( n \) consists of \( 2^n - 1 \) bits. The full sequence can be viewed as a collage of overlapping \( k \)-bit sequences, \( k \leq n \), drawn from the list of all possible binary combinations of \( k \) bits. For example, for \( k = 2 \), the possible binary combinations are: \( (0, 0), (1, 0), (2, 10), \) and \( (3, 11) \). Thus a portion of the full sequence consisting of the bits 0110100 can be decomposed as the overlapping combination of the sequences \( (1), (3), (2), (1), (2), (0) \). The same decomposition procedure can be applied for any \( k \). The M-sequence has the convenient property that all subsequences of length \( k \leq n \) randomly appear within the full sequence the same number of times, namely \( 2^n - k \) occurrences (except that the all-zero sequence of length \( k \) appears \( 2^n - k - 1 \) times). Because of this statistical regularity of the M-sequence, it is an excellent tool for the investigation of a cell’s neural code.

Analysis

Cells were selected for analysis based on the following criteria. To ensure single cell isolation, we chose only cells with clearly isolated clusters in the various two-dimensional projections of the four-electrode amplitude space; clusters with clipped responses due to amplifier saturation were avoided. To achieve reasonable estimates of the information rates, \( \approx 1,000 \) spikes were required during the whole stimulus. Finally, only cells with on or off linear temporal kernels (see following text) were studied, since this formed the basis for the definition of response events. In total, 12 cells (4 ON, 8 OFF) in one cat were studied at five contrast levels—80% (9 cells), 40% (6 cells), 20% (3 cells), 14% (1 cell), and 6% (2 cells)—yielding a total of 21 trials.

Response events and precision analysis

To study the precision of spikes, we attempted to classify each individual spike as part of a spike event evoked in response to a specific sequence of \( k \) frames. This was done by applying the following algorithm, described here for an OFF cell. We determined the average stimulus before a spike, and defined the cell’s mean conditional latency (conditioned on a spike) as the time to the zero-crossing between peak and trough in the spike-triggered-average stimulus (illustrated in Fig. 1). Then, as shown in Fig. 2, for each spike in the train, we looked back in time from the spike by the mean conditional latency and found the closest OFF transition (bright frame followed by dark frame) within a window of \( \pm 1.5 \) frames; the spike was assigned to that transition. If there was no such transition, the spike was unclassified. We characterized sequences by their length \( k \) and the location \( t \) of the transition within the sequence (e.g., \( k = 8, t = 3 \) labeled an 8-frame sequence with a transition at the onset of the 3rd frame—that is, between the 4th and 3rd frames, where the 1st frame was the latest in time). For a given choice of \( k \) and \( t \), a given transition was uniquely associated with a surrounding sequence, and the spike was assigned to that sequence. All spikes associated with the same sequence were labeled as part of the same event. The percentage of total spikes that were unclassified served as a measure of the level of “spontaneous” activity that was not driven by transitions.

Once the events were identified for a given choice of \( k \) and \( t \), the probability that a specific sequence produced an event was computed by dividing the number of times some spike response (\( \geq 1 \) spike) was obtained for that sequence, by the total number of presentations of that sequence (i.e., \( 2 \times 2^{k-t} \) times). This quantity was called the event probability.

We assessed the timing precision of the first spike in an event for each sequence consisting of a specified number of frames, \( k \), with transition location \( t \). A distribution for the times to the first spike in an
VARIABILITY AND INFORMATION IN THE CAT LGN

FIG. 1. The spike-triggered-average stimulus and the temporal kernel for an
OFF cell. The vertical axis represents the stimulus luminance on a linear scale,
normalized and shifted so that +1 represents the bright frame luminance L, −1
represents the dark frame luminance D, and 0 represents the mean luminance
(L + D)/2. —, the spike-triggered average; - - - - , the temporal kernel, obtained
by normalizing (in the frequency domain) the spike-triggered average by the
stimulus spectrum, up to a cutoff of 90 Hz (see METHODS). The temporal kernel
represents the cell’s temporal receptive field: it is the linear filter that, when
applied to the stimulus, best predicts the cell’s response in the sense of least
mean-square error. Both functions show a strong bright-to-dark transition in
the stimulus ~32 ms before the spike occurred. This was defined as the cell’s
mean conditional latency.

The information in the spike train about the stimulus was quantified
using the “direct” method (de Ruyter van Steveninck et al. 1997;
Strong et al. 1998a,b). This method estimates the mutual information
between stimulus and response “directly” from the spike trains with-
out regard to the details of the stimulus/response relationship and with
very few assumptions about the coding strategy. This method relies on
the fact that the mutual information between the stimulus and re-
sponse can be written as the difference of two spike train entropies.
First, the maximum amount of information that a spike train response \( R \) can provide about the stimulus is just given by the entropy of the
spike train itself, \( H(R) \). This is estimated from the probability distri-
bution of spike responses over the course of the whole experiment
without specific knowledge of the stimulus. Second, the information
the spike train carries about the stimulus is reduced from this max-
imum by the degree to which there is variability or noise \( N \) in the
repeated responses to an identical stimulus, as measured by the spike
train noise entropy, \( H(N) \). This is estimated from the probability
distribution of spike responses to multiple, identical presentations of
the same stimulus, averaged over stimuli.

With the M-sequence, responses to the repeated presentations of
each k-frame stimulus sequence were easily obtained. For each oc-
currence of a specific k-frame sequence, the response beginning at
a delay \( \tau \) (ranging from 0 to 130 ms) relative to the onset of the initial
frame of the sequence was divided into bins of size \( \Delta \tau \) (usually 1 ms)
containing the number of spikes in each bin. These bins were com-
bined to form spike “words” of length \( T = M\Delta \tau \), where \( M \) was an
integer number of bins. For example, for \( M = 3 \), the joining of three
bins containing 2, 0, and 1 spikes, respectively, would yield the word
201 (note that the absence of spikes in a bin can be informative, and
its contribution was included).

We then computed the entropies for each choice of \( k, T, \) and \( \Delta \tau \)
by building the probability distribution of these words—a cross-the
whole experiment for \( H_{i=1}^{T-k+1}(R) \) and across the multiple repeats of the \( k \)
th-frame stimulus sequence \( \{ i = 1, \ldots, 2^k \} \) at time-shift \( \tau \) for
\( H_{i=1}^{T-k+1}(N) \). Note that the location of a transition, \( t \), within the
k-frame sequence was now irrelevant and not specified; instead all
k-frame sequences contributed equally to this analysis. Both \( T, \Delta \tau \)
were varied to obtain estimates of the entropy on different time scales.
For a given \( T, \Delta \tau \), the average information about the k-frame
sequence that began at time \( \tau \) before a response word was then given by
\( H_{i=1}^{T-k+1}(R) - (H_{i=1}^{T-k+1}(N)) \), where \( (H(N)) \) was the average noise
entropy across all k-frame stimulus sequences (i.e., average over \( i \)).
We assigned the information about k-frame sequences, for the given
\( T, \Delta \tau \), as the maximum information across \( \tau \) (see following text).

First though, for each combination of \( T, \Delta \tau, k, \) and \( \tau \), we corrected
for finite-data errors. This was done by computing the mutual infor-
mation for different partitions of the data: the whole data set, and
the average over each half of the set, over each third, and each fourth.
This average information was then plotted as a function of the number
of partitions \( N \), and fit to the functional form, \( I = I_0 + I/K+1/N \) (Strong et
al. 1998b). \( I_0 \), therefore represented the true information rate
extracted from the limit of infinite data for a given \( T, \Delta \tau, k, \) and \( \tau \).
Note, however, that when the amount of the data were too small, even

Information analysis

The information in the spike train about the stimulus was quantified
by the “direct” method (de Ruyter van Steveninck et al. 1997;
Strong et al. 1998a,b). This method estimates the mutual information
between stimulus and response “directly” from the spike trains with-
out regard to the details of the stimulus/response relationship and with
this correction failed. Empirically, this occurred when the ratio of $I_2$ to $I_1$ became large. We used a ratio of $2 \times 10^{-3}$ as the border between sufficient and insufficient data and show results only for cases in which data were sufficient by this criterion. In practice, the corrections for finite data were typically tiny, and the point of this procedure was primarily to screen out cases (e.g., too-large $k$ or too-large $T$) for which data were insufficient.

Given the corrected information, we assigned the information about $k$-frame sequences as follows. For the given $k$, $T$, and $\Delta T$, we determined the $\tau$ that maximized the information. The information, $I$, was then assigned to be the average information over the bins within $\pm 4$ ms around this maximum. (We chose this to correspond to a frame width, so that averaging smoothed out any frame-related artifacts.) The information rate of the spike train, in units of bits/time, was $I/(\Delta t \Delta T)$. We converted this to units of bits/spike $I_0$ by dividing by the neuron’s average spike rate, $r$, assessed over the entire two-M-sequence stimulus: $I_0 = I/(r \Delta T)$.

The method worked well only for relatively short response words. Long response words required long stimulus sequences to minimize the randomizing effect of different stimulus contexts on early or late portions of the response word. However, since each sequence repeated $2 \times 2^{14-k}$ times, as $k$ increased, our estimate of the entropies degraded due to sampling problems. Thus to consider very long response words, we employed a different strategy; we estimated a lower bound on the information carried by the spike train about the stimulus by applying the direct method to the two repeats of the full M-sequence. Assuming that the only thing in common between the two presentations of the M-sequence was the stimulus itself and that therefore the noise in the two cases were uncorrelated, the information that one response $R_1$ carried about the second response $R_2$, $I_{x,y}(R_1, R_2)$ should be a lower bound to the information between either response $R_i$ and the stimulus $f$, $I_{s,x}(f, R_i)$ (Strong et al. 1998b). We took each response to be the spike train generated by each full M-sequence, minus the first and last 200 ms. We then computed each spike train’s entropy, $H_{x,y}(R_j)$, $j = 1, 2$, for words of length $T$, and the joint entropy, $H_{x,y}(R_1, R_2)$, for the co-occurrence of words in the two spike trains. These were computed from the probability distributions for words by using overlapping intervals (incremented by $\Delta t$, to increase the effective number of samples). To correct for finite-data errors, data size scaling was applied in this case directly to the entropy estimates (rather than to the mutual information as in the data size scaling described above); an example is shown in Fig. 3A. The mutual information between the two responses was then

$$I_{x,y}(R_1, R_2) = H_{x,y}(R_1) + H_{x,y}(R_2) - H_{x,y}(R_1, R_2)$$

In general, the dependence of the information on word length $T$ for a given bin size $\Delta T$ was small. Hence, to summarize the dependence for a particular bin size, the infinite-word-length limit was taken by obtaining a linear fit to the plots of the (infinite data limit) entropies versus $1/T$, and using the $y$ intercept as the (infinite word limit) entropy rates in the calculation of the information rate. The fit was performed only over the range of $1/T$ where sufficient data were available to accurately estimate the entropy rates, as illustrated in Fig. 3B. In practice, $T$’s ranged from 8 to 48 ms. Finally, the information per second from words of spikes was converted into the information per spike by dividing by the mean spike rate across the whole experiment.

**Models**

We constructed quasi-linear threshold models of driven LGN spiking activity to investigate whether the observed precision could be explained by simple mechanisms. All models convolved the full M-sequence stimulus, binned at one-sixth the frame period, with the cell’s temporal kernel to generate a firing function, $f(t)$ (linear part). These responses were thresholded and perhaps squared (nonlinear part) to generate firing rates

![Figure 3](http://jn.physiology.org/)

**FIG. 3.** Estimate of the lower bound to the average information per spike between the stimulus and words of spikes. A: the entropy of 48 ms words within the spike train, binned in 1-ms bins, scaled with data fraction in a controllable fashion. B: as the inverse length of the spike words decreased, the amount of data (for fixed recording length) decreased, and the single spike train and joint spike train entropy estimations began to fail for words longer than ~50 ms. An infinite word length extrapolation for the entropy rates was obtained by fitting to the region where the data were sufficient. $r(t)$, as follows. We defined $r(t) = \lambda_0 \left[ (r(t) - 0)^p \right]^q$, where $[x]^k = x, x > 0$; $= 0$, otherwise; $p = 1$ for a linear function and $p = 2$ for a quadratic function; and $\lambda_0$ was chosen to make the mean of $r(t)$ equal to the observed mean firing rate. The value of the threshold $\theta$ was fit as described in the following text. Finally, spikes were generated as a Poisson process from these rates, perhaps along with a refractory period, as will be described in the following text.

The temporal kernel was determined as the spike-triggered-average stimulus, divided by the autocorrelation (or in Fourier space, the power spectrum) of the M-sequence stimulus (the power in the M-sequence at frequency $f$ is proportional to $\sin \left( |fr_s|f \right)^2$, where $r_f = 120$ Hz is the frame rate). This division yields the linear filter that, applied to the stimulus, gives the best estimate of the shape of the impulse response in the sense of least mean-square error (Rieke et al. 1997). The spike-triggered average and temporal kernel for one cell can be seen in Fig. 1. The division is done in Fourier space, where it simplifies to a frequency-by-frequency division; otherwise it would involve multiplying one matrix by the inverse of another matrix. However, one does not want to continue dividing up to arbitrarily high frequencies where the power in the stimulus approaches zero, as this will just amplify high-frequency noise. We chose to do the division up to some cutoff frequency, and to set all power above that cutoff frequency to zero. To choose a cutoff frequency, we tried cutoffs from 75 to 100 Hz in 5-Hz steps. For each cutoff, we applied the corresponding filter to the M sequence to obtain the output $f(t)$, converted this to a rate function $r(t)$ as described in the preceding text using $p = 1$, and chose the threshold $\theta$ as that which minimized the mean-square error difference between the predicted Poisson rate function and the eight-frame PSTH for the actual data. We then chose the cutoff frequency that gave the least mean-square error; this best cutoff was 90 Hz. This kernel was used subsequently in all models to draw actual spikes for PSTH comparison (see following text).

The conversion from $r(t)$ to spikes was as follows. We interpolated $r(t)$ to achieve a temporal resolution of 1/60 of a frame (the spike-
trigged average and temporal kernel had been computed in bins of 1/6 of a frame or ~1.39 ms). For the simple Poisson case, spikes were then generated in each time bin $\Delta t$ with probability $r(t)\Delta t$, using $\Delta t = 139$ $\mu s$. For the case of a Poisson process with a refractory period, a free firing rate, $q$ (Berry and Meister 1998), was generated assuming a specific refractory period, $\mu$, by taking $q(t) = r(t)[1 - r(t)\mu]$. Spikes were then drawn as in the Poisson case but using $q(t)$ rather than $r(t)$. In the case of only an absolute refractory period, the probability of a spike was set to zero for $\mu$ ms after each spike. We also tried adding an exponential recovery after the absolute refractory period, setting $\mu = \mu_{\text{abs}} + \mu_{\text{rel}}$, where $\mu_{\text{abs}}$ was the absolute refractory period and $\mu_{\text{rel}}$ was the exponential recovery of the probability from zero up to $q(t)$. This implementation for a relative refractory period is reasonable when $\mu_{\text{rel}}$ is smaller than the characteristic time over which the firing rate remains relatively constant.

For each of the models, an optimal threshold and refractory period(s) (if applicable) were selected simultaneously to minimize the mean-square error between the real data and the model of the segment of the eight-frame PSTHs defined by the 18 ~1.39 ms bins before and the 7 bins after the end of the eight-frame sequence. This was done by trying every threshold from 1 to 5 in steps of 0.2, (if applicable) absolute refractory periods from 1 to 4 ms and relative refractory periods from 0.5 to 4 ms in steps of 0.5 ms for which $q(t)$ remained positive, and then selecting the combination of threshold and refractory periods that gave the least mean-square error. These ranges seem reasonable because in no case was the optimum parameter at an extreme of the range explored for that parameter. The mean firing rate over the whole stimulus in the model was typically matched to within a few percent of the data’s mean.

**RESULTS**

Full-frame, binary, 14-bit M-sequence stimuli were presented at different contrast levels. In general, this stimulus drove cells in the LGN well. Average spike rates across all cells and stimulus conditions ranged from 4.6 to 25.3 Hz. Neural responses were usually triggered by transitions from either bright to dark frames (OFF cell), or vice versa (ON cell); we referred to two-frame sequences of bright/dark or dark/bright as an OFF or ON transition, respectively. Each cell’s polarity was determined by reverse correlating the spike train with the M-sequence stimulus. Figure 1 presents the spike-triggered-average stimulus for one of our good ON cells (cell 4, 80% contrast) that had a strongly driven response producing nearly 7,000 spikes. We use this cell to illustrate the main results of our analysis. A spike at time 0 for this cell was generally preceded by a transition from bright to dark ~32 ms earlier. This time delay was referred to as the cell’s mean conditional latency. Figure 1 also illustrates the cell’s temporal kernel (see METHODS), which represents the cell’s temporal receptive field and has the same 32 ms mean conditional latency; we will return to this later.

An initial 1,200 frames (10 s) from the beginning of the M-sequence were presented to adapt the cells to the stimulus ensemble before showing the M-sequences used in data analysis. After the conditioning, two repeats of the full M-sequence were displayed without delay. A total of $2 \times 2^{14-k}$ repetitions of each $k$-frame sequence ($k \leq 14$) occurred, e.g., 128 repeats of each eight-frame sequence. Because of this convenient property, it was natural to focus on responses to the set of $k$-frame sequences for different $k$.

**Mean response: the PSTH matrix**

The M-sequence stimulus presented frames of random stimuli in series rather than in isolation. To obtain an average response to a specific stimulus sequence, we extracted the individual spike responses to the multiple presentations of that sequence in the full M sequence. Consider first the case of one-frame stimuli. The average response to single bright or dark frames of stimuli was generated in the form of a matrix of PSTHs (Fig. 4). The shading in each 1-ms bin corresponds to the total number of spikes from all presentations of this sequence at that time relative to the frame onset. Note that there was a nonzero spike rate even at the time origin that was nearly the same for both bright and dark frames. This reflects the fact that at early times, the spikes were responses to earlier frames over which we had averaged. The response to the particular bright or dark frame was most clear at ~32 ms as expected from the cell’s mean conditional latency.

One advantage of visualizing a PSTH matrix is in the ability to display the neuron’s average responses to stimuli more complex than just a single frame, as shown in Fig. 5 for two-frame sequences. This clearly shows that spikes tended to be generated near the mean conditional latency in response to

FIG. 4. Peristimulus time histogram (PSTH) matrix of cell 4’s spike responses to 1-frame sequences (i.e., to bright frames or dark frames). Responses were histogrammed in 1-ms bins relative to time 0, defined as the time of onset of the stimulus frame, f1. The stimulus itself is illustrated to the left of time 0 (gray represents a bright frame, black represents a dark frame). For 1-frame sequences, virtually no spikes were observed in response to a bright frame (stimulus 1) at approximately a mean conditional latency (32 ms) from its onset, while a large number of spikes were seen at a similar time after a dark stimulus (stimulus 0). Responses to stimulus 0 at ~32 ms largely represent responses to the 1/2 of cases in which the frame preceding it was a bright frame, creating an OFF transition. Similarly, many spikes are seen in response to stimulus 1 ~1 frame later (~40 ms), representing responses to the 1/2 of cases in which the bright frame was followed by a dark frame. Conventions for this and future PSTH-matrix figures: time 0 is defined relative to the sequence as shown at the top of the matrix; e.g., here time 0 is the time of onset of the single frame of the stimulus, f1, for multiple-frame stimuli, the last frame in time would be f1, the preceding frame f2, etc., so that if $k$ would be the 4th frame in reverse temporal order. The stimulus frame sequence is shown to the left of time 0 in temporal sequence from left to right (left being earlier in time), with dark representing a dark frame and gray representing a bright frame.
an OFF transition (stimulus 2), whereas spiking was clearly suppressed near the mean conditional latency by an ON transition (stimulus 1). Note that the response to a dark frame (stimulus 0 in Fig. 4) was now broken down according to whether the preceding frame was dark or bright (stimuli 0 and 2, respectively, in Fig. 5).

Figure 6 displays the PSTH matrix (with 1-ms time bins) for the response to seven-frame sequences, sorted according to the rightmost two frames, f1 and f2 (we usually numbered frames in a k-frame sequence consecutively as fn, n = 1, . . . , k, with f1 the latest in time and fk the earliest). This grouped together all responses to sequences with an OFF transition in the most recent two frames. As expected, a large vertical band of spikes centered at ∼32 ms appeared in response to the OFF transition. One striking feature was the slight slant in time of the OFF response band near 32 ms. Qualitatively, for this cell, the time to the first spike was correlated with the amount of time the stimulus had been bright prior to the final transition to dark: the longer this time, the earlier the occurrence of the first spike in the response.

Moreover, the spikes in this band were noticeably isolated in time on both sides by regions of virtually no spikes, suggesting that there was a high degree of temporal precision in the response when seven frames of the stimulus were specified. To examine this, each spike should ideally be classified as part of a response to a particular sequence. In the PSTH matrix though, each spike occurred multiple times, each time associated with a different time frame and sequence. Hence, echoes of the main OFF response appeared in the other quadrants of the PSTH matrix where an OFF transition occurred earlier in the sequence.

Event classification

To classify a spike to a unique sequence, a search was performed to find the OFF transition that was most likely to be responsible for a given spike. All spikes classified to the same transition were then grouped together as the spike “event” in response to the sequence containing that transition (see Methods). In practice, this algorithm reproduced the event structure...
quite well, as can be seen from the comparison of Figs. 7 and 8. These show the PSTH matrix and the extracted unique spike events, respectively, for the 1/4 of eight-frame sequences having an off transition in their final two frames. The band of spikes near 32 ms was clearly reproduced in the spike events. Virtually all spikes in the train were accounted for by this technique; only 1.8% of the spikes were unclassified. (Note that spikes placed at random would show 5/16, or 31%, unclassified.)

In general, for the group data across all cells, 10 of 21 trials had unclassified percentages <5%, while for the remaining 11 trials this was larger than 5%. Qualitatively, the unclassified percentage was correlated with the degree to which spikes were locked to the stimulus as evidenced by visual isolation of spikes around the mean conditional latency in the PSTH matrix. When the spikes around the mean conditional latency could be visibly isolated (10 of 21 trials), the algorithm appeared to yield fairly low unclassified percentages (9 of those 10 trials). The one exception was a 40% contrast trial for an on cell in which the events in response to an on transition were fairly well isolated yet the unclassified percentage was nevertheless high (26%), probably because spikes were also produced without a transition when the stimulus had been bright for several frames. In cases when locking was evident but poor (5 of 21 trials had bands of increased spiking, but these were not well isolated) or when spiking was more indiscriminate (6 of 21 trials had poorly distinguishable bands), the unclassified percentage tended to be larger (10 of these 11 trials had unclassified percentages above 5%). The one exception was a 6% contrast trial for an off cell with a weak linear kernel–its events were not well isolated, but its unclassified percentage was nevertheless low (3.5%).

For each sequence, we defined its event probability to be the percentage of its occurrences that evoked an event of one or more spikes.

Response variability

SPIKE TIMING PRECISION. Using the binary k-frame sequences to characterize the stimulus, and the spike events to characterize the response, we turn to the next issue of this paper: a study of the reliability and precision of responses and their dependence on the stimulus. The timing precision of these events was examined by determining the jitter in the time of the first spike in the events associated with a particular sequence. This is shown in Fig. 9A for the only possible two-frame sequence with an off transition. This sequence generated a spike response 49% of the time, and the time of the first spike had a

FIG. 7. PSTH matrix of cell 4’s spike responses to 8-frame sequences containing an off transition between frames f2 and f1, histogrammed in 1-ms bins relative to the onset of stimulus frame f1 (maximum spike rate of 688 Hz).

FIG. 8. Cell 4’s extracted spike event responses to 8-frame sequences containing an off transition between frames f2 and f1, histogrammed in 1-ms bins relative to the onset of frame f1. The quality of the algorithm for associating spikes with a particular sequence can be judged by comparing this against the PSTH matrix for 8-frame sequences in Fig. 7 (or more objectively by determining the percentage of unclassified spikes, see text).
standard deviation of 3.25 ± 0.04 ms. Because the responses to all possible combinations of stimulus frames before and after the two frames of the transition were averaged together, this standard deviation represented the precision achieved by the two frames of the OFF transition alone, when the other frames were unspecified. Its value was already less than the standard deviation expected (7.2 ms) if the first spike times were distributed uniformly over the three-frame search window that defined events.

When eight frames of the stimulus were specified, with an OFF transition occurring between frames f2 and f1 (which we denote as an \("f2-f1 transition\)\), every sequence had <3 ms SD in the time to the first spike, including sequences with both high and low event probability (Fig. 9B). The median across sequences of this standard deviation was 1.28 ± 0.03 ms (median ± square root of the jackknife variance for the median sequence). These results suggested that a significant part of the timing jitter in response to two-frame stimuli was simply due to imprecise specification of the stimulus history. In particular, the average first spike times differed significantly for different eight frame sequences, decreasing with increasing event probability, as plotted in Fig. 9C. This naturally broadened the width of the distribution of first spike times when the responses to different stimulus sequences were averaged together.

Because the timing precision clearly varied with the exact stimulus sequence, we wanted a more generic measure of the overall variability of the response for a given level of stimulus specification. We selected the median, over sequences, of the standard deviation of the time to the first spike in an event as a robust index for this purpose. Figure 10 plots this median standard deviation as a function of the number of frames specified. The upper and lower interquartile ranges are also depicted, showing that in some cases, the distribution of standard deviations is clearly asymmetric. Three curves are shown, corresponding to the location within the sequence of the OFF transition. The f2–f1 curve (OFF transition between f1 and f2) indicates that the median precision improved until three to four frames before the two frames encompassing the OFF transition were specified (5–6 total frames). When four frames before the transition were specified, specifying additional frames after the transition (7 frames on the f3–f2 curve or 8 frames on the f4–f3 curve) did not substantially alter the median precision, suggesting that these frames had little effect on the overall timing precision. This plateau in the precision likely reflected the intrinsic variability of the cell because further stimulus specification did not further increase the precision. In the group

![Figure 9](http://jn.physiology.org/doi/10.1152/jn.00486.2001)

**FIG. 9.** Standard deviation of the distribution of times to the 1st spike in an event for a given sequence plotted against the probability that an event was evoked for that sequence. A: when only the 2 frames of the OFF transition were specified, the single 2-frame sequence had a standard deviation of 3.25 ± 0.04 ms and an event probability of 49%. When a total of 8 frames were specified, including the OFF transition between f2 and f1, all sequences had sub-3 ms standard deviations, with a median standard deviation of 1.28 ± 0.03 ms. The median sequence is shown as a filled circle. C: mean first-spike times plotted against the event probability for case B.

![Figure 10](http://jn.physiology.org/doi/10.1152/jn.00486.2001)

**FIG. 10.** Median SD, across all k-frame sequences, of the distribution of times to the 1st spike, plotted against the number of frames in the stimulus window, k. The 3 curves correspond to different locations of the OFF transition e.g., f2–f1 indicated the OFF transition was between frames f1 (the latest in time) and f2. The upper and lower interquartile ranges for each point indicate the width of the distribution of SDs across sequences. When no frames were specified after the transition (curve f2–f1), the median SD leveled off at ~1.3 ms, once 5–6 total frames were specified. When 1 or 2 frames were specified after the transition, the curves simply shifted to the right, indicating that the 3–4 frames before the 2 frames encompassing the transition were primarily responsible for improving the timing precision. Note that the f3–f2 curve is offset to the right by 0.1 frame for clarity.
data, the precision of all trials with <5% unclassified percentage (with the exception of a 6% contrast trial) improved with increased specification of the frames before the transition; the median standard deviation decreased on average by 41 ± 13% \((n = 9)\) from the case where only two frames were specified to the case where eight frames were specified with the ON or OFF transition between frames \(f_4\) and \(f_3\). When all trials were considered regardless of percentage unclassified, a decrease of 31 ± 18% \((n = 21)\) was found on average.

Figure 11A plots the dependence of the median standard deviation of the time to first spike on the percentage of unclassified spikes across the population of cells, for eight-frame sequences with \(f_4\)–\(f_3\) transitions. The cluster of trials having unclassified percentages <5% clearly exhibited high timing precision (with 1 exception for a 6% contrast trial)—mean of 1.56 ± 0.39 (SD) ms \((n = 9\), excluding the outlier). Two trials (1 cell at 40% contrast, another at 80% contrast) had high unclassified percentages (26 and 22%, respectively) but nevertheless had small median standard deviations \((1.97 ± 0.06\) and \(2.28 ± 0.13\) ms, respectively). The remaining nine trials that had >5% unclassified spikes clustered at ~4.01 ± 0.58 ms. Many of these trials were less well driven, as evidenced by their generally lower firing rate, as shown in Fig. 11B. Because this group of trials often responded more diffusely in time, making classification of spikes difficult, their poorer precision was not surprising. However, given that their precision was well below the 7.2 ms expected from random placement of spikes, it seems likely that this reflected a true property of the cells rather than an artifact of the classification method.

The timing precision showed only weak dependence on the event probability, that is, on the reliability with which a sequence evoked a response. Within the group data, the Spearman rank-order correlation was statistically significant \((P < 0.05)\) when both all of the data and part of the data were analyzed (see METHODS) in only 8 of 21 trials. It was not significant for both conditions in another 7 of 21 trials. In the remaining six trials, the significance level changed between the two conditions. The fact that 13 of 21 trials showed no clear correlation suggested that the dependence of first-spike-time standard deviation on event probability was not strong. That is, the temporal precision of response was not simply a result of a “strong” stimulus: even responses that were infrequently evoked could nonetheless be evoked at fairly precise times when they did occur. Hence, reliability and timing precision were not strongly coupled.

SPIKE COUNT PRECISION. The timing precision analysis focused on how the stimulus affected the jitter of a single spike (namely the 1st spike in an event). To study the precision of the remaining spikes in an event, we analyzed the precision of the number of spikes in the events evoked by a stimulus. This spike count precision was characterized by examining the variance in the number of spikes per event versus the mean number of spikes in an event. In the case of a Poisson process, the variance is equal to the mean. At the other extreme, the minimum possible variance for a discrete counting process with a given mean \(m\) is obtained if the number of spikes in every event is either \(\text{ceil}(m)\) (the smallest integer \(\geq m\)) or \(\text{floor}(m)\) (the largest integer \(\leq m\)). This minimum variance varies periodically with the mean, dropping to zero at each integer and forming a scalloped curve between integers.

Figure 12A plots cell 4’s spike count variance for the single two-frame OFF sequence against its mean spike count. Also shown are the line expected for a Poisson process and the scalloped curve representing the minimum possible variance. The variance for this sequence clearly fell close to the Poisson limit. When the stimulus history specification was expanded to seven frames, with the OFF transition between \(f_2\) and \(f_1\) (Fig. 12B), most of the sequences remained Poisson-like, but a few began to have sub-Poisson responses. However, if we consider eight-frame sequences with the OFF transition between \(f_4\) and \(f_3\) frames (Fig. 12C), meaning that we specify two frames after the transition frames as well as four frames before, the variance for almost all sequences was significantly less than Poisson, falling in many cases close to the minimum-variance limit. These results were summarized by examining the median across sequences of the Fano factor, which is the ratio of the
The marked suppression of the noise by the specification of the frames after the transition can be straightforwardly understood. Figure 9B shows that when no frames were specified after the transition, there was a cluster of sequences that had event probabilities near 50% regardless of how many frames were specified into the past. Once one frame was specified after the transition frames, however, the event probabilities diverged so that many sequences produced events with nearly unit probability, while other sequences produced events with very small probability. This reflected the fact that, for this cell, the event produced by a transition from bright to dark (10) could be suppressed by a subsequent transition back to bright (101). On the other hand, if no subsequent transition occurred (100), an event was virtually always produced. This occurred almost irrespective of what happened before the off transition. The variance in spike count for each of the two cases (101 and 100) could be small. However, by not specifying the frame after the transition, as in the f2–f1 curve of Fig. 13, the two cases were averaged together, producing a large variance and a Fano factor close to unity. Thus simply increasing the stimulus history was not always enough to obtain precise responses; enough frames both before and after the off transition had to be specified to maximize the precision (note that frames after the transition are still within the causal range where the linear kernel is sensitive to the stimulus).

In the group data, such large reductions in the spike count Fano factor were not very common. Comparing the median Fano factor for the two-frame case to the eight-frame, f4–f3 transition case, there was an average 27 ± 21% (n = 21) reduction across all trials. Considering only those trials with unclassified percentages <5% yielded a 38 ± 24% (n = 10) reduction; trials with unclassified percentages >5%, 17 ± 9% (n = 11). The Fano factor itself was generally around or below 1 in nearly all cases, as shown in Fig. 14. No strong dependence of the Fano factor on the unclassified percentage appeared in the data, except that Fano factors <0.5 occurred only in trials with <5% of spikes unclassified. Moreover, note that, among the low-unclassified-percentage trials, good timing pre-
precission did not necessarily imply good count precision (compare Fig. 11A to Fig. 14).

Information transmission

The spike timing and count variability measures discussed above gave some indication of the precision of LGN neurons. How much information did this level of precision allow the cells to transmit?

To address this, we changed our analysis method. The preceding analyses of variability depended on defining events that associated each spike with a unique sequence that evoked it. This required specifying both sequence length and the location within the sequence of the transition (because spikes were associated with transitions and these 2 facts uniquely linked transitions to sequences). For the information analysis, we instead considered all sequences of a given length, without regard for the presence of a transition, and simply examined the response at some fixed time interval after the initiation of the sequence.

We computed information using the direct method (see METHODS). We binned time into discrete units of size $\Delta t$, typically 1 ms, and defined the “letters” of the response “alphabet” as the number of spikes in a bin (0 or 1 for 1-ms bins).

A string of $M$ such letters formed a response “word”—for $M = 1$, the word was simply the number of spikes in a single bin. Ideally, the choice of bin size should reflect the degree of temporal resolution in the code, while the word size should reflect the longest time scale of temporal correlations in the code. The timing precision analysis suggested that a reasonable bin size was $\sim$1 ms. Initially ignoring correlations between bins, we calculated the information about $k$-frame stimuli by considering only single-bin words at this resolution (Fig. 15). The information grew with time from the onset of the stimulus sequence, provided that further stimulus frames continued to be specified, up to at least nine frames. At this point, the maximum information was $\sim$3.5 bits/spike and appeared to be nearing a plateau. The existence of a plateau was reasonable since a given response time bin should give little or no information about stimulus frames that occurred far in the past. For longer sequences ($k \geq 4$), the information began to drop from its peak at $\sim$24–26 ms after the onset of the last frame in the sequence, or $\sim$16–18 ms after the onset of the first unspecified frame. This suggests that 16–18 ms was the minimum delay for a frame to significantly influence the response. This was in rough agreement with our previous results that one and perhaps two frames after the transition frames can influence the spike count by vetoing or allowing spikes induced by the transition; if the response occurs 32 ms after the transition, then these frames would have onsets $\sim$15 and 24 ms before the response that they influence.

We also compared the maximal observed information rate of 3.5 bits/spike to the cell’s maximum possible information rate, as measured by the entropy of its spike train. Achieving this maximum would imply that all of the cell’s response variability (as measured in single 1-ms bins) was used to encode the stimulus. In fact, the coding efficiency, the ratio of the actual information coded to that which could possibly be encoded, was $\sim$51% (for $k = 9$), so that the cell transmitted information in individual 1-ms bins at a level that was within a factor of two of its limit.

We next examined the role of time resolution in information encoding by varying the binwidth. We considered 8-ms words
of the spike train, and binned these words using either 1-, 2-, 4-, or 8-ms resolution. If the precise timing of the spikes at these resolutions within the word were important for transmitting information, then we expected more information at smaller bins than larger bins. Finer resolution increases the possible information the spike train can code; if the actual information coded also grows, then the coding efficiency would not significantly change with increasing resolution. On the other hand, a fall-off of the coding efficiency would indicate that the increased resolution is not being used to code information. We computed maximum information rates for eight-frame sequences to ensure that there were sufficient repeats of each sequence to allow us to estimate the information for multiple-bin response words. The information rate increased from 2.4 bits per spike at 8-ms bins to 3.1 bits per spike at 2-ms bins, a 29% increase (Fig. 16A), while the spike train entropy increased by 36% over the same range. That is, \( \frac{0.29}{0.36} = 81\% \) of the increase in entropy associated with this increase in resolution was used to code information. As a result, the coding efficiency stayed relatively flat, decreasing only \(-5\%\) from a bin size of 8 to 2 ms. Thus the position of spikes at \( \leq 2\)-ms resolution was significant for coding information. Improving the resolution by a factor of 2 from 2 to 1 ms yielded an additional 3% increase in information to 3.2 bits per spike, compared with an increase in entropy of 15%, suggesting that only 20% of the entropy change encoded information. Thus while more information was encoded at this finer resolution, there was a diminishing return as the noise became a proportionately larger contributor to the cell’s increased variability.

### Redundancy or synergy in coding

Given that a temporal resolution down to 1 ms was useful, another important question to address is the manner in which patterns of spikes in these bins contributed to information transmission. Three possibilities exist: different 1-ms bins may code information independently; they may encode information redundantly, so that \( M \)-bin words code less information than \( M \) times the one-bin-word information; or they may interact synergistically so that \( M \)-bin words code more than \( M \) times the one-bin-word information. Note that the degree of redundancy or synergy may change with \( M \)—for some word sizes, the responses may be more redundant, whereas for other word sizes, they may become synergistic.

We investigated the degree of synergy and redundancy in the LGN responses in three ways. First, we compared the information in 8-ms words with 1-ms bins to that found in 1-ms words. For our example cell, the 3.2 bits per spike for 8-ms words with 1-ms bins was close to the 3.3 bits per spike for 1-ms words found for eight-frame sequences in Fig. 15, indicating only a little redundancy and no synergy between the responses of adjacent 1-ms bins. This near independence of spikes in 8-ms words was not simply due to Poisson firing since the distribution of the number of spikes within the 8-ms window showed a much larger probability for two spikes (0.21) than would be expected from the square of the one spike probability (0.008). This result suggests that the cell was bursting, although the bursts apparently did not lead to a large level of redundancy. This could happen because redundant patterns (such as bursts) might be used synergistically to code for the stimulus. It is important to point out that our measure looks at the average level of redundancy or synergy so that the combination of different groups of redundant and synergetic spikes could appear independent at this time scale.

Second, to determine whether this lack of significant redundancy or synergy survives at longer time scales, we examined the information in much longer words. Unfortunately, the direct method as applied to the repetitions of the \( k \)-frame sequences could not be used to study response word lengths longer than \(-8\) ms (for 8-frame stimuli) due to data insufficiency. Instead, we estimated a lower bound on the information in the entire response to the full \( M \)-sequence by using the two repeats of the full \( M \)-sequence (see METHODS). Assuming that the stimulus was the only common drive for the two responses, then the information between the responses to the two repeats bounded from below the information either could carry about the stimulus. We compared this lower bound extracted in the limit of infinitely long response words to the exact information rates computed from eight-frame sequences and 8-ms responses (Fig. 16). The lower bound came reasonably close to the information rates computed from 8-ms words across all bin sizes considered. This implies that there is little redundancy over times longer than 8 ms but leaves open the possibility of synergy (if the true infinite-word information were much higher than our lower bound).

As a final test of the redundancy or synergy between spikes, we compared the exact information transmitted by individual 1-ms bins to the lower bound on the information...
transmitted by infinitely long words of 1-ms bins. Figure 17 plots these two measures for all cells and trials in the data in terms of both bits per spike and per second. Nearly all of the trials fell close to the diagonal line where the information from single bins equaled the lower bound to the information from infinitely many bins. To the right of this diagonal line, coding is synergetic: more information is conveyed on average by combinations of spikes in 1-ms bins than by single 1-ms bins. To the left of this diagonal line, coding is redundant: less information is transmitted on average by the words of bins than by the single 1-ms bin. Hence, the fact that the trials aligned close to, but to the left of, the diagonal suggested that there was at most a slight amount of redundancy for real cells. Since the infinite word information is a lower bound, we can only be certain that the true information lay to the right of the plotted data—that is, there was little or no redundancy and possibly some synergy.

Comparison to quasi-linear threshold models

The results reported here suggest that, while there was variation among the population, visual thalamic cells could exhibit very precise responses that conveyed considerable amounts of information per spike on average. Responses of these cells are often modeled as resulting from the convolution of the cell’s temporal kernel (shown in Fig. 1) with the stimulus, followed by a nonlinear thresholding to generate a firing rate (see METHODS). Is the degree of precision consistent with this picture?

We first chose the threshold that gave a best match of the model PSTH to the data and assumed that spikes were generated randomly according to an inhomogenous Poisson process with the model PSTH (appropriately scaled to yield the same mean rate as the data). The best-matched model gave a broader and more symmetric PSTH than was observed in the data, suggesting that the precision of model spikes was significantly worse. Extracting spike events as described above allowed us to directly compare the precision of the model (Fig. 18B) to the data (Fig. 18A), in response to eight-frame sequences that had a bright to dark transition in the rightmost two frames. The model spike events were clearly more diffuse in time and did not capture the details of the dependence of response onset times on stimulus sequence. We next considered models in which spikes were generated from a Poisson process with an absolute refractory period. We considered this for the case in which the firing rate was a linear function (Fig. 18C) or a quadratic function (not shown) of the thresholded filter output. In each case, refractory period and threshold were chosen together to optimally match the data (least mean-square error in PSTH). The PSTH of the linear refractory model was slightly narrower than that derived without a refractory period but continued to be wider than the data and to not show the temporal irregularity of the data. The model using a quadratic function gave results similar to, but slightly poorer than, those of the linear refractory model, so we do not consider it further. A relative refractory period in addition to an absolute refractory period also yielded quantitatively similar results as the case of an absolute refractory period alone.

The model’s failure to capture the detailed structure of response onset times is specifically due to an underestimation of longer onset times, while shorter onset times were well reproduced by the model (Fig. 19A). This discrepancy can be understood from an examination of the PSTH matrices (Fig. 18): it appears that when two or more consecutive dark frames preceded the bright to dark transition, this lengthened first-spike times in the data; but this effect was not picked up by any of the models. The models also reasonably reproduced the mean spike counts observed in the data, but showed a tendency to underestimate smaller mean counts and overestimate higher ones (Fig. 19B).

The Poisson model did a poor job of reproducing the observed variability in spike timing or spike count (Fig. 20, ○). The inaccuracy in spike count precision is not surprising, because a Poisson model will always have a Fano factor of 1. However, the model incorporating a refractory period came much closer to reproducing the precision of the data (Fig. 20, △). This model tended to slightly overestimate smaller first-spike-time standard deviations and Fano factors and to under-

![FIG. 17. Group data on the information in 1-ms bins plotted against the lower bound for the information in infinitely long words of 1-ms bins. The former was computed from the multiple repeats of 9-frame sequences, while the latter was estimated from the two repeats of the full M sequence. A: data plotted as information per spike. B: data plotted as information per second. In both representations, the data fell close to the diagonal line where the 1-ms bin information equals the infinitely long word information. This suggests that there was at most a minimal amount of redundancy, and possibly synergy, in the coding by successive spikes.](http://jn.physiology.org/ijp importância dos dados científicos em uma análise adequada de dados genéticos)
estimate larger ones, showing less overall diversity of first-spike-time standard deviations and Fano factors than the data.

**Discussion**

We have found that LGN neurons can show great precision in their responses to M-sequence stimuli. For at least a subset of cells, spikes occur in discrete events triggered by an on or off transition with spike rates close to zero at other times. The time of the first spike in an event can be precise to 1–2 ms, and this precision can be maintained even for unreliable events (events that occur with low probability). The four frames before the transition frames influence the event timing, so these frames must be specified to discern the cell’s spike timing precision. The number of spikes in an event can also show great precision, with Fano factor (ratio of variance to mean) approaching 0.3 (vs. a value of 1 expected for a Poisson process). The frames after a transition can “veto” or allow an event so that two frames after the transition frames as well as the four before must be specified to discern the cell’s spike count precision. This precision of response allows cells to carry up to 3.5 bits per spike of information about the stimulus. The coding efficiency of information transmitted in 1-ms bins can be within a factor of two of the limit set by the spike train’s entropy—a limit that is achieved when all of the cell’s variability is used to code information. The coding efficiency remains relatively constant as the temporal resolution for specifying spike times increases to at least 2 ms, and still more information is gained by increasing resolution to 1 ms, indicating that the timing of spikes at these resolutions carries information about the stimulus. By comparing the information carried by 1-ms response words to that in 8-ms words and to the lower bound on the information transmitted by infinitely long response words, we find that there is at most only a modest amount of redundancy in the coding by successive spikes, and we find no evidence for synergy. Finally, this precision can be largely, but not entirely, accounted for by a model in which firing rate is generated by filtering the stimulus with the cell’s temporal kernel and applying a threshold, followed by spike generation as a Poisson process with an absolute refractory period.

**Previous work on spike timing and count precision**

Our work adds to a growing body of work finding high response precision and high information rates in the LGN in response to full-field noise stimuli. Keat et al. (2001), in work contemporary with the present work, found 1–2 ms SD for the time to the first spike in an event in response to full-field Gaussian white noise in close agreement with the present results for binary white noise. Reinagel and Reid (2000) reported a particularly low width (SD) of 0.6 ms for one PSTH peak in one cell’s response to a full-field “naturalistic” noise stimulus but did not more generally report on timing precision. Both of these papers and Kara et al. (2000) demonstrated sub-Poissonian Fano factors in LGN responses to full-field Gaussian noise in agreement with the present findings. Comparable precision of spike timing and count in response to full-field noise stimuli has been reported in the retina (Berry and Meister 1998; Berry et al. 1997; Kara et al. 2000; Keat et al. 2001).

Measures of response to other stimuli often do not show similar precision. Thus Guido and Sherman (1998) measured the jitter in the time to first spike in responses to spots flashed in the center of the LGN cell receptive field and reported standard deviations ranging from ~3 to 35 ms, depending on the mode of firing (burst vs. tonic). The greater variability seen in this case of a single flashed spot is akin to the spread of the PSTH seen when only a single frame is specified (Fig. 4) and may reflect the lack of specification of the cell’s initial state. That is, when stimulated only by a blank screen (before stimulus onset), spontaneous activities may lead a cell to wander...
through a state space of comparable diversity to that created by the set of binary stimulus sequences that could precede a single frame in our experiments. Similar reasoning might also explain why statically flashed, spatially nonuniform stimuli have produced Fano factors larger than 1 in several LGN studies (Hartveit and Heggelund 1994; Levine et al. 1996; Sestokas and Lehmkuhle 1988). Reich et al. (1997) reported a PSTH standard deviation of 5 ms for one LGN cell in response to a slowly drifting sine grating, but at least some of this jitter was due to a slow drift in response phase across many trials, which may have represented a slow change in cell state; responses over a small set of adjacent trials showed considerably greater precision.

**Specifying neuronal state**

The idea that responses to temporally modulated stimuli can show great precision, even while responses to more static stimuli may show greater variability, has already a long history (e.g., Buracas et al. 1998; de Ruyter van Steveninck et al. 1997; Mainen and Sejnowski 1995) and has stirred controversy (e.g., Egelhaaf and Warzecha 1999). Our findings add a focus on stimulus history, showing that sufficient specification of a temporally varying stimulus is key to revealing neural precision. By extension, this emphasizes the importance of control of neuronal state: noise may not be intrinsic to a neuron or a piece of neural tissue but may instead simply represent variables that are not under the experimenter’s control. While a dynamic stimulus may control neural firing and thus control a given cell’s state, lack of a stimulus (a blank screen) yields spontaneous activities that are stochastic, being triggered at least in part by spontaneous quantal events in photoreceptors (Mastronarde 1989), and these in turn may lead a cell’s state to wander in an uncontrolled way, presenting an uncontrolled initial condition at the moment of a flashed stimulus. A related argument was made by Buracas et al. (1998), who showed that whether or not a given stimulus evoked a spike in a cell of area

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**FIG. 19.** Comparison between the mean 1st-spike times and spike counts for the data and the models, plotted for all 8-frame sequences with an OFF transition between f4 and f3. A: the 1st-spike times in the data were well approximated by all of the models except that larger times were underestimated. B: the spike counts were generally well predicted by the models, although there was a small tendency to underestimate small counts and overestimate high counts.

**FIG. 20.** Comparison between the spike timing SDs (A) and spike count Fano factors (B) for the data and the models plotted for all 8-frame sequences with an OFF transition between f4 and f3. The Poisson model poorly matches the precision of the data. The model with refractoriness does considerably better but shows some tendency to overestimate smaller first-spike times/Fano factors and underestimate larger ones. The points with 0 SD and Fano factor showed only one non-0 spike response across the 128 trials.
MT was strongly correlated to the local field potential at the given time and place.

It is interesting that, at least for our binary stimuli, specification of 8 frames (67 ms) seems adequate to specify the LGN state to sufficient precision to saturate spike timing and count precision, while 9–10 frames (75–83 ms) saturate the information coded by spikes. These numbers are in rough agreement with the width of the cell’s temporal kernel (Fig. 1), which differs from zero over a span of ~65–70 ms.

Previous work on neuronal information transmission in the LGN

The information rates we have found—2–3.5 bits/spike, 20–90 bits/s—are similar to those found by others in LGN who, like us, used “direct” methods (Eckhorn and Pöpel 1975; Reinagel and Reid 2000). These methods directly estimate the information carried by the spike train about the stimulus, without a requirement for explicit decoding, by assaying certain stimulus and response probability distributions (Eckhorn and Pöpel 1974; Strong et al. 1998a,b). Indirect methods, such as the stimulus reconstruction method (Rieke et al. 1997), rely on being able to “decode” the response. These methods provide only a lower bound to the information rates: any information that is successfully decoded was present, but there is no guarantee that all information that was present was successfully decoded. Rates found using indirect methods in LGN have generally been quite low—only ~2 bits/s (Dan et al. 1998; McClurkin et al. 1991; Reinagel et al. 1999)—suggesting that much information present in the LGN spike trains was missed by those methods.

Stimuli better matched to the receptive field may yield more information. Eckhorn and Pöpel (1975) found that spatially uniform stimuli yield lower LGN transmission rates (25–40 bits/s at the best flash rate) than spots isolated at the receptive field center (60–80 bits/s) (Eckhorn and Pöpel 1975). Their spot and full-field stimuli were only briefly flashed at slow, periodic intervals (~30 Hz). Full-field stimuli modulated randomly at higher rates drive relatively high LGN information rates, as shown both by our work and that of Reinagel and Reid (2000). The latter sees a range of information rates similar to what we have found even though their naturalistic stimulus distribution contained much more entropy than our binary distribution (924 bits/s in their distribution vs. 120 bits/s in ours). That suggests that we may be seeing the limits of what an LGN cell can code, at least to full-field stimuli. On the other hand, the results of Eckhorn and Pöpel (1975) suggest that both we and Reinagel and Reid (2000) might have seen even higher information rates if we had restricted flashes to the cells’ receptive field centers.

High information rates like those reported here have also been observed in a variety of other systems, including retina, visual cortex, and insect motion-detecting neurons (e.g., Berry et al. 1997; Buracas et al. 1998; de Ruyter van Steveninck et al. 1997; Reich et al. 2000; Strong et al. 1998a), suggesting that the precision found here may not be a special property of LGN or thalamic neurons.

Minimal redundancy

The issue of redundancy or synergy in the neural code has been addressed in numerous papers, but we are aware of only a few (Brenner et al. 2000; Reinagel and Reid 2000) that have looked at the issue in terms of temporal coding in a single neuron rather than population coding across multiple neurons. Reinagel and Reid (2000) found that LGN neurons can sometimes code more information on average in patterns of spikes than if those spikes were considered independently. The synergy they reported, however, was at most only ~20%, and many neurons were slightly redundant (~10%) or only very weakly synergetic. Our results are consistent with this in the sense that we also observe at most only mild redundancies in the coding by individual neurons. We cannot rule out syner-gies, but to the degree that our lower bound closely approximates the true information, the fact that none of our neurons lay to the right of the independence line in Fig. 17 suggests that there are also no large synergies in the coding by individual neurons.

Models of response generation

We have found that a simple model of response generation, based on thresholding the output of the cell’s temporal kernel applied to the stimulus and imposing a refractory period, can match much but not all of the precision of response that we observed. To achieve this result, it was critical that the cell’s temporal kernel be used and not simply the spike-triggered average; use of the latter gave noticeably less precision in both timing and spike count (not shown).

The discrepancies between the precision of the model and the observed data most likely arise from the linear filter model rather than the specifics of the spike generation mechanism. The PSTH matrix generated by the model is somewhat wider and considerably more regular than that of the data. Most strikingly, the model fails to show the lengthening of first-spike times observed in the data when two or more consecutive dark frames preceded the bright/dark transition. This yields a less “jagged” left edge for the model PSTH compared with the data PSTH. This jagged edge is dominated by a response’s first spikes, which are unaffected by refractoriness. Accordingly, the error is unlikely to be in our model of spike generation and refractoriness but rather in the model of PSTH generation by linear filtering. This is also suggested by the fact that the temporal kernel and the spike-triggered average both give similarly smooth leading edges (data not shown), so it seems unlikely that a better filter would alter this result. It is further suggested by the results of Kara et al. (2000), who found that they could successfully model the spike count variability of LGN cells by beginning with the observed PSTH (rather than deriving the PSTH from a filter as we are doing) and adding both absolute (~1 ms) and relative (~20 ms) refractory periods extracted from the cell’s interspike interval distribution.

Accounting for the observed PSTH presumably requires a more complex nonlinearity in our model of firing-rate generation than the thresholding used here; it would be interesting to determine whether contrast-gain-control mechanisms (Shapley and Victor 1978; Victor 1987) might be sufficient to reproduce the response onsets and improve the agreement between the model and data precision measures. Nonetheless, it should be noted that the model as it stands is significantly nonlinear. The optimal threshold value (optimal in the sense of least mean-
square error in matching the data PSTH) was 80% of the root-mean-square of the output of the filtering of the stimulus by the temporal kernel (see legend to Fig. 18); that is, it was necessary to set a significant fraction of positive filter outputs to zero. The optimal absolute refractory period was 3 ms, long compared with probable biophysical absolute refractory periods of $\approx 1$ ms. (When both an absolute and a relative refractory period were used, the optimum was similar, 2.5 ms absolute plus 0.5 ms relative refractory period.)

An alternative approach to modeling the neural responses observed here is to dispense with a firing rate model altogether and instead directly model the spike generation process. Berry et al. (1997) found that responses of retinal neurons to full-field noise stimuli consisted of brief response events surrounded by substantial periods of zero spike rate, similar to the cases in our experiment in which most spikes could be accounted for by response events locked to ON or OFF stimulus transitions. This has led the same group more recently (Keat et al. 2001) to suggest that a rate description of such responses, in which spike probability is zero for extended periods interrupted by brief events, may be inadequate. Instead they proposed predicting the spikes themselves rather than a spike rate by regarding the output of a cell’s linear filter applied to the stimulus as a voltage-like variable rather than a rate and counting upward-going threshold crossings of this voltage as spike times. Parameterizing the filter, adding a spike-induced “hyperpolarization” to represent refractoriness, and adding appropriate noise yielded a 20-parameter model (15 parameters describing the filter and 5 additional parameters). They showed that such a model, fit individually to each cell by optimizing a cost function incorporating precision measures, could do a good job of replicating the cell’s spiking events and their statistics for both retinal and LGN cells in response to Gaussian noise stimuli. We have no reason to doubt that the same models would well describe the responses to binary noise stimuli studied here.

Conclusion

LGN cells can show remarkable precision in their responses and code information at high rates and with high coding efficiency. Revealing this precision requires sufficient specification of the stimulus history. This points to the possibility that measurements of neuronal precision may be limited as much by the degree to which the experimenter controls the variables relevant to a cell’s response as by the intrinsic precision of neural processing.

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