Prediction and Decoding of Retinal Ganglion Cell Responses with a Probabilistic Spiking Model

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Sensory encoding in spiking neurons depends on both the integration of sensory inputs and the intrinsic dynamics and variability of spike generation. We show that the stimulus selectivity, reliability, and timing precision of primate retinal ganglion cell (RGC) light responses can be reproduced accurately with a simple model consisting of a leaky integrate-and-fire spike generator driven by a linearly filtered stimulus, a postspike current, and a Gaussian noise current. We fit model parameters for individual RGCs by maximizing the likelihood of observed spike responses to a stochastic visual stimulus. Though compact, the fitted model predicts the detailed time structure of responses to novel stimuli, accurately capturing the interaction between the spiking history and sensory stimulus selectivity. The model also accounts for the variability in responses to repeated stimuli, even when fit to data from a single (non-repeating) stimulus sequence. Finally, the model can be used to derive an explicit, maximumlikelihood decoding rule for neural spike trains, thus providing a tool for assessing the limitations that spiking variability imposes on sensory performance.

Sensory experience depends on the encoding of external events in the spiking activity of neurons. Understanding this encoding process and its effects on sensory experience requires quantitative models that can capture the relationship between stimuli and neural spike responses. To provide a clear insight into neural function, such a model should be simple, approximating or leaving out many biophysical details. But to be useful, the model must accurately describe the mapping from stimuli to spikes. Specifically, the elements of the model should capture the spatiotemporal integration of sensory inputs, and should provide insight into the statistical features governing the fidelity of neural signals. Finally, the model should be "invertible", providing an optimal rule for decoding the information contained in spike trains. The most common approach to describing neural stimulus selectivity is the "receptive field", which captures the stimulus features that best drive spiking responses (Kuffler, 1953; Hubel and Wiesel, 1968). The receptive field is often interpreted as a quasi-linear model that maps stimuli to firing rates, with an implicit assumption that spikes are then generated from a Poisson process. However, recent studies have shown that this model fails to account for key statistical features of neural spike responses (Troy and Lee, 1994; Reich et al., 1997; Berry et al., 1997; Fellous et al., 2004). In particular, neurons in the early visual pathway are capable of spiking with high temporal precision, which can be important for conveying visual information (Mainen and Sejnowski, 1995; Berry et al., 1997; Liu et al., 2001). It is therefore important to develop models that can account for the both stimulus selectivity and the statistical features of neural spike trains.

Integrate-and-fire (IF) models provide a well-known and more realistic alternative to Poisson models of spike generation. Recent studies have suggested that IF models are capable of exhibiting some of the important statistical behaviors of real neurons (Troyer and Miller, 1997; Reich et al., 1998; Shadlen and Newsome, 1998; Jolivet et al., 2003; Keat et al., 2001). However, spike dynamics can cause significant distortions in the receptive field properties estimated using standard methods like reverse correlation (Aguera y Arcas and Fairhall, 2003; Pillow and Simoncelli, 2003), and the estimation of such models is an important open problem.

In this paper, we show that a generalized IF model provides a detailed functional description of the light responses of retinal ganglion cells (RGCs) in the macaque monkey. We use a recently developed technique (Pillow et al., 2004a; Paninski et al., 2004) to fit the model to spike responses elicited by a white noise stimulus. The fitting procedure simultaneously estimates parameters that govern stimulus selectivity, history-dependence, and stochasticity (noisiness) of the neuron. We show that the fitted model accurately predicts spike responses to novel stimuli, accounting for both the stimulus-dependence and detailed statistical structure of spike trains. Surprisingly, the model also predicts the variability and precision of responses to repeated stimuli, even when fit with a single (non-repeating) stimulus. The model also bears out a simple explanation for the origins of spike timing precision, in terms of voltage slope at threshold.

Finally, the model provides a novel tool for assessing the information content of RGC spike responses. The computed probability of spike responses can be used to derive an optimal (maximum likelihood) decoding rule for extracting stimulus information from spike trains. Our results demonstrate that this method is capable of extracting information from spike trains more faithfully than a generic linear-nonlinear encoding model.

Preliminary versions of this work have been reported at (Pillow et al., 2004b; Pillow et al., 2004c).



Figure 1: Schematic diagrams of the generalized integrate-and-fire model (left), and the standard linear-nonlinear-Poisson model (right).

Materials and Methods

Experimental Measurements & Stimuli

The data presented in this paper are a subset of the data in (Uzzell and Chichilnisky, 2004); experimental methods are described in detail there. Briefly, multi-electrode extracellular recordings were obtained *in vitro* from small pieces of retina from four macaque monkeys, with retinal pigment epithelium attached, maintained at 32-36 degrees C, pH 7.4. Analysis was restricted to two physiologically-defined classes of cells that very likely correspond to ON and OFF parasol cells based on several lines of evidence (Chichilnisky and Kalmar, 2002).

The retina was stimulated with a photopic, achromatic, spatially uniform, optically reduced image of a cathode ray tube display refreshing at 120 Hz. The stimulus was a temporal sequence consisting of two intensity values, pseudo-randomly selected on every refresh. In the first three experiments (24 cells), the contrast (standard deviation divided by mean) of this stimulus was 96%, and model validation was performed on a novel stimulus sequence (7-30 sec) repeated 37-176 times. In the fourth experiment (9 cells), binary white noise stimuli of 12% and 48% contrast were used for fitting the model parameters. Validation was then performed using responses to brief (25 ms) positive and negative flashes with contrasts of 12%, 24%, 48% and 96% (flash amplitude divided by background luminance).

Generalized Integrate-and-Fire (IF) Model

We fit neural responses using a generalized integrate-and-fire (IF) model. This model, which is closely related to the "Spike Response Model" (Gerstner, 2001; Jolivet et al., 2003), consists a standard leaky integrate-and-fire compartment driven by three time-varying input currents: (1) a stimulus-dependent current I_{stim} ; (2) a spike-history dependent current I_{sp} ; and (3) a noise current I_{nse} . A schematic diagram of the model is shown in Figure 1a. The model dynamics are specified by a differential equation governing voltage of the integrator:

$$\frac{dV}{dt} = -\frac{1}{\tau}(V(t) - V_l) + I_{stim}(t) + I_{sp}(t) + I_{nse}(t),$$
(1)

and whenever V(t) = 1, a spike occurs and V(t) is reset instantaneously to zero. The input current I_{stim} is the linear convolution of the stimulus with input filter \vec{k} , which represents the neuron's spatiotemporal receptive field. We can therefore write the stimulus-dependent current as

$$I_{stim}(t) = \vec{k} \cdot \mathbf{stim}(t), \tag{2}$$

where $\mathbf{stim}(t)$ is a vector consisting of the (discretized) space-time stimulus vector just preceding time t. For a full-field stimulus (as considered here), $\mathbf{stim}(t)$ is an *n*-vector of the monitor intensities for the *n* frames preceding time t, and \vec{k} is simply the neuron's temporal receptive field.

The spike-history dependent current I_{sp} results from the injection of a fixed current waveform \hat{h} following each spike, and accounts for spike history effects such as refractoriness, burstiness and adaptation (see (Paninski et al., 2004)). Mathematically, this is equivalent to considering \hat{h} as a linear filter that is convolved with the recent spike train history:

$$I_{sp}(t) = \vec{h} \cdot \mathbf{spike_hist}(t), \tag{3}$$

where $\mathbf{spike_hist}(t)$ is a vector of (discretized) recent spiking activity at time t, matched to the size of \vec{h} .

Finally, the noise current I_{nse} consists of Gaussian white noise with standard deviation σ and represents the net contribution of all noise sources to the membrane potential:

$$I_{nse}(t) = \sigma \sqrt{\Delta t} \mathcal{N}_i, \tag{4}$$

where Δt is the time step of the simulation, and \mathcal{N}_i is a standard Gaussian random variable. The last two parameters of the model are τ , the membrane time constant, and V_l , the reversal potential of the leak current.

IF Model Fitting

The generalized IF model is specified by the parameters $\{\vec{k}, \vec{h}, \sigma, \tau, V_l\}$, where \vec{k} and \vec{h} are both multidimensional vectors. Characterizing a neuron's response requires the estimation of these parameters from a sequence of stimuli and the (extracellularly recorded) times of elicited spikes. In previous work (Pillow et al., 2004a; Paninski et al., 2004), we developed an efficient and computationally tractable algorithm for computing the IF model likelihood function, $P(\mathbf{spikes} | \mathbf{stim}, \{\vec{k}, \vec{h}, \sigma, \tau, V_l\})$, which is the probability of observing a set of spike times given a set of stimuli and a fixed setting of the model parameters. The likelihood function, described in detail in (Paninski et al., 2004), can be computed by numerically solving the Fokker-Plank equation for subthreshold voltage probability during each interspike interval. This amounts to finding the probability (under the Gaussian noise model) that voltage crossed threshold at precisely the observed spike times. We fit model parameters $\{\vec{k}, \vec{h}, \sigma, \tau, V_l\}$ by maximizing the likelihood of the observed responses. This involved searching for the model parameters (in the high-dimensional parameter space) that maximized the probability of the observed spike times for the given stimulus. In performing this search, we relied on a result from (Paninski et al., 2004): that the likelihood function for the IF model is log-concave, meaning that the likelihood is mathematically guaranteed to be free from local maxima, for any stimulus and spike train data. This result ensures that gradient ascent of the likelihood function will converge to its global maximum, and guarantees that the IF model (unlike more complicated nonlinear models like Hodgkin-Huxley), can be fit reliably and tractably using simple gradient ascent techniques. We performed parameter optimization for the IF model using 50 seconds of data: a non-repeating binary white-noise stimulus and the associated spike times for each neuron. The stimulus-selectivity (receptive field) parameters \vec{k} and spike-history parameters \vec{h} were taken to be 15 and 10 dimensional vectors, respectively. These were fit in a vector space spanned by a basis of raised cosines with log scaling of the time axis. Basis vectors were given by

$$B_i(t) = \frac{\cos(\log[t+\psi] - \phi_i) + 1}{2},$$
(5)

for t such that $\log(t + \psi) \in [\phi_i - \pi, \phi_i + \pi]$ and 0 elsewhere (so each basis vector looks like a single raised "bump"). The bump positions ϕ_i were selected so that the basis vectors "tile", or sum to 1, allowing for phase-invariant (on a log-time axis) representation of \vec{k} and \vec{h} . The resulting basis (similar to that used in (Keat et al., 2001)) has the ability to represent fine temporal structure near the time of a spike, but is constrained to be smooth at longer timescales. This allows \vec{k} and \vec{h} to be represented with relatively few parameters.

The basis for \vec{k} had a temporal extent of 40 stimulus frames (333 ms), meaning that \vec{k} effectively operates on a 40-dimensional stimulus space. The 15-dimensional space used for fitting \vec{k} (i.e. the space spanned by the basis vectors $\{B_i\}$) allowed for faster fitting of \vec{k} , but did not qualitatively affect the results; sample comparisons with \vec{k} fit in the full 40-dimensional space achieved qualitatively identical performance. The basis was also observed to accurately represent shape the spike-triggered average. The basis for \vec{h} had a temporal extent of 200 ms, though (as shown in fig. 2a and b), the actual fits of \vec{h} were essentially zero outside of the first 25-40 ms following a spike).

Finally, a robust version of maximum likelihood was used to obtain parameter estimates that were less sensitive to statistical outliers. The lowest-likelihood 5% of the interspike intervals were treated as statistical outliers and were ignored when optimizing the parameters. Empirically, this resulted in reduced estimates of the noise parameter σ , in keeping with the intuition that the model compensates for exceedingly low-probability spikes by artificially increasing the noise parameter (see Discussion). The robust estimate of σ provided an improved match to the first-spike time precision of spike trains.

Linear-Nonlinear-Poisson model

To provide a baseline for comparison with the IF model performance, we also fit data with the simplest and most widely-used model of visual responses: a Linear-Nonlinear-Poisson (LNP) model, which is illustrated schematically in Figure 1b. This model consists of a single linear filter (i.e. a linear receptive field), followed by an instantaneous nonlinear function, which accounts for response nonlinearities like rectification and saturation, followed by Poisson spike generation. This model is widely used because of its computational simplicity and the ease with which its parameters can be estimated (Chichilnisky, 2001). LNP is the implicit model underlying all reverse-correlation style methods for estimating receptive fields.

The LNP model is specified by the parameters \vec{k} and \vec{n} , where \vec{k} denotes the linear stimulus filter (as in the IF model), and \vec{n} is a parametric description of the nonlinear function converting filter output to instantaneous spike rate. In this case, \vec{k} was taken to be a 15-dimensional vector (as used above for the IF model), and \vec{n} was taken to be a linear spline with 15 knots (hence 15 fitted parameters). We initialized our estimate of these parameters using reverse correlation to compute the spiketriggered average, as described in (Chichilnisky, 2001). However, reverse correlation can provide biased estimates of \vec{k} if the stimulus distribution is non-spherical (Simoncelli et al., 2004), as it is here (i.e. binary). We therefore performed gradient ascent of the LNP model likelihood function, to find the maximum likelihood estimate of \vec{k} and \vec{n} . Although not necessarily guaranteed to converge to the global maximum, this procedure can correct biases in the estimates of the \vec{k} and \vec{h} arising from the use of non-spherical stimuli, and performs quite well in simulations.

A robustifying procedure similar to that used for the IF model (i.e. ignoring the likelihood of the lowest 5% of interspike intervals) was employed to put likelihood comparisons with the IF model on equal footing. Nearly identical performance was obtained when the LNP model was fit using reverse correlation (i.e. without further ascent of the likelihood function), and was robust to the number of parameters used in the representation of the nonlinearity \vec{n} over a wide range (8 to 25).

Quantifying Model Performance

In order to compare IF and LNP model performance at predicting responses to novel stimuli, we calculated the percent error in the peristimulus time history (PSTH) and peristimulus time variance (PSTV) predicted for responses to repeated stimuli, as well as a single-trial measure of distance between spike-trains. Figure 5b shows the percent of the variance in the PSTH accounted for by each model, or $100 * (1 - \langle (PSTH_{RGC} - PSTH_{model})^2/(PSTH_{RGC} - \langle PSTH_{RGC} \rangle)^2 \rangle$, where $\langle \cdot \rangle$ indicates an average over time. PSTHs were computed by binning each response, summing, and filtering with a Gaussian with a standard deviation 1 ms. Figure 5c shows the percent error in the PSTV for each model, which is given by $100 * \langle PSTV_{RGC} - PSTV_{model} \rangle / \langle PSTV_{RGC} \rangle$. PSTV was computed by sliding a 10-ms window across the response raster and calculating the variance (across trials) of the number of spikes in that window.

Figure 5d-f examines the similarity between pairs of spike trains to repeated presentations of a single stimulus, using a specific spike train distance measure (Victor and Purpura, 1997). The measure relies on a timescale parameter λ and is defined as the minimum cost for bringing one spike train into alignment with another by shifting and adding/deleting spikes, where adding or deleting has a cost of 1, and shifting a spike by t ms entails a cost of t/λ . For any two spike trains, this distance measure is bounded above by the sum of the number spikes in both spike trains ($\lambda = 0$) and bounded below by their spike count difference ($\lambda = \infty$). Software for computing the spike time distance measure was obtained from the author's website (Victor and Purpura, 1997).

Calculation of Likelihoods for Discrimination Task

We examined the decoding performance of the two models by using them to perform discrimination in a simple ideal observer task. We suppose that an observer is given spike trains $\{s_A, s_B\}$ in response to presentation of stimulus sequences $\{A, B\}$, and must decide which of the two stimuli is associated with which spike train. The optimal decision rule comes from comparing the likelihood of each stimulus under each observed spike train (Green and Swets, 1966):

$$R(A,B) = \frac{P(s_A|A) \cdot P(s_B|B)}{P(s_A|B) \cdot P(s_B|A)}$$

If R(A, B) is greater than unity, the correct choice is made; otherwise the stimuli are paired with the wrong spike trains. The conditional probabilities in this expression may be determined directly from any stochastic model of neural response, such as the IF or LNP models.

Results

Figure 1a illustrates the components of the generalized IF model, which consists of a leaky integrateand-fire model driven by a stimulus-dependent current, a spike-history dependent current and a Gaussian noise current. The stimulus-dependent current arises from the neuron's linear receptive field, while the spike-history current arises from the injection of a "post-spike current" \vec{h} following every spike (see Methods for details). Note that \vec{h} can assume an arbitrary shape, which allows the model to generate a diverse array of behaviors observed in real neurons, including refractoriness, spike rate adaptation, spike rate facilitation, bursting and bistability (Jolivet et al., 2004; Paninski et al., 2004). This flexibility endows the model with the ability to mimic a wider array of biologically realistic dynamical behaviors than the classic integrate-and-fire model or Poisson spiking models.

We fit the model using responses of parasol (magnocellular-projecting) RGCs to a spatially uniform, achromatic binary temporal white noise sequences (random flicker, see Methods). For these stimuli, \vec{k} represents the temporal receptive field, although in general \vec{k} can represent the neuron's spatio-temporal-chromatic receptive field. The IF model parameters were fit to spike responses from a



Figure 2: Parameters obtained from fits to RGC data for the IF model (a-c) and the LNP model (d-e). (a) Filters \vec{k} and spike-response currents \vec{h} obtained for five ON cells in one retina. (b) Corresponding filters for four OFF cells. (c) Histograms of model parameters V_l , τ and σ for all 24 cells in 3 retinas. (d) Comparison of linear filters for IF model (gray) and LNP models (black), for one ON cell (above) and one OFF cell (below). (e) Measured LNP point nonlinearities for converting filter output to instantaneous spike rate.

single (non-repeating) stimulus train. Subsequently, RGC responses to multiple repeats of a novel stimulus were recorded, and predictions generated by the IF model were assessed using several quantitative measures.

Figure 2a-c shows model parameters fit to data from a collection of ON and OFF RGCs recorded simultaneously. Note that the filters, \vec{k} , are consistent in waveform and timescale within each cell type. The same is true of the spike current waveforms, \vec{h} , which operate on a faster timescale. The biphasic shape of the \vec{h} currents allows the model to reproduce burstiness in RGC responses the initial positive component drives voltage up close to threshold following voltage reset, and the later negative component (possibly accumulated over several spikes) exerts a hyperpolarizing effect to end a burst. The larger amplitude \vec{h} in ON cells matches the burstier responses observed in ON compared to OFF cells (see Figs. 3-4). Figure 2c also shows histograms of the three scalar parameters { σ, τ, V_l } for all cells.

To provide a baseline for comparison, the same data were fit using the simplest and most widelyused model of visual responses: a Linear-Nonlinear-Poisson (LNP) model, shown schematically in Fig. 1. The model consists of a cascade of linear and nonlinear stages, followed by Poisson spike generation (see Methods for details). Although Poisson processes cannot exhibit refractoriness, bursting, or other known statistical features of spike trains, the LNP model is widely used because



Figure 3: Responses of an ON cell to a repeated stimulus. (a) Recorded responses to repeated one-second stimulus (top), simulated LNP (middle) and IF model (bottom) spike trains. Each row corresponds to the response during a single stimulus repeat; 167 repeats are shown. (b) Peristimulus time histogram (PSTH), or mean spike rate, for the RGC, LNP model and IF model. For this cell, the IF model accounts for 91% of the variance of the PSTH, while the LNP model accounts for 75%. (c): spike count variance computed in a sliding 10-ms window. (d): Magnified sections of rasters, with rows sorted in order of first spike time within the window. The four sections shown are indicated with blue brackets in **a**.

of its computational simplicity and the ease with which its parameters can be estimated using reverse correlation (Chichilnisky, 2001).

Figure 2d shows linear filters obtained for both the LNP model and the IF model, for one ON cell and one OFF cell. Note that the linear filter recovered for the LNP model (i.e. the spike-triggered average) is noticeably different than that obtained for the IF model (Figure 2d). Given that the IF model incorporates more realistic spike generation and provides more accurate predictions of real spike trains (see below), this suggests that the LNP model provides an inaccurate description of how neurons integrate visual inputs over time (Berry and Meister, 1998; Pillow and Simoncelli, 2003; Aguera y Arcas and Fairhall, 2003). Also, note that the nonlinear function of the LNP model (Figure 2e) has no direct counterpart in the IF model, whose firing rate is determined implicitly by the dynamics of the leaky integrator and spike thresholding.



Figure 4: Responses of an OFF cell to a repeated stimulus. Details identical to Fig. 3.

Predicting Responses to Novel Stimuli

Once both models were fit using a set of training data (a 50-second white noise stimulus sequence), we sought to compare their performance at predicting responses to novel stimuli. For a first comparison, we examined RGC responses to repeated presentations of a binary white noise stimulus. This allowed us to assess model accuracy in predicting not only the average response, but response variability over repeated trials.

Note that this type of analysis, commonly known as "cross-validation", provides a much more more stringent test of a model's capabilities than a simple goodness-of-fit evaluation. Rather than examining how well the model fits the training data, we evaluate its ability to generalize and predict responses to novel, randomly-generated patterns that were not presented during training. An important implication of this procedure is that we can ignore the fact that the models differ slightly in their number of parameters; fitting a model with too many parameters leads necessarily to a *decrease* in generalization performance (a problem known as "overfitting"). For example, using too many parameters to represent the post-spike current \vec{h} in the IF model (or the nonlinearity \vec{n} in the LNP model) yields an irregular (non-smooth) function that overfits detailed random fluctuations in the training data, and therefore performs poorly at predicting novel responses.

Figures 3 and 4 show recorded responses and predictions of the IF model and the LNP model, for an ON and an OFF cell, respectively. Rasters of RGC responses and corresponding simulated responses from both models illustrate that the IF model (bottom rows) captures the structure of the RGC spike trains (top rows) more faithfully than the LNP model (middle rows).

The peristimulus time histogram (PSTH, Figs. 3b & 4b) summarizes the time-varying firing rate exhibited by the data and both models. The IF model (black trace) matches the sharp peaks in the PSTH more accurately than the LNP model (red trace). Trial-to-trial variability of the responses is reflected by the peristimulus time variance (PSTV, Figs. 3c & 4c), computed by sliding a 10-ms window along the response raster and computing the variance across trials of the number of spikes in that window. Because RGC spike trains have history-dependence which makes them much less variable than a Poisson process (Uzzell and Chichilnisky, 2004), it is unsurprising that the LNP model fails to match the PSTV of the data. The IF model provides a more accurate prediction. Although integrate-and-fire models have been shown previously to be capable of reproducing realistic spike count statistics in simulation (Reich et al., 1998), it is notable that the generalized IF model does so despite the fact that the objective function (likelihood of the spike train) does not include a measure of variability and does not require (or use) repeated stimuli.

A more detailed view of spike train structure and model performance was obtained by sorting the rows of a response raster in order of the first spike time in a given window (Figs. 3d and 4d). Sorting reveals considerable structure in RGC interspike intervals, which is largely captured in the sorted responses of the IF model (bottom) but is completely absent in the sorted responses of the LNP model (middle).



Figure 5: Performance comparison across cells. Empty and filled circles represent ON and OFF cells, respectively. (a) likelihood per spike of novel RGC responses under the fitted IF model and LNP model. Value plotted is the geometric mean of the likelihood of over all spikes under each model. Gray dashed lines represent a factor of 2 above and below identity. Data from 24 cells (3 retinas) are shown. (b) percent of the variance in the PSTH accounted for by both models, for each cell. Points above the diagonal represent superior performance by IF model. (c): percent error in the peristimulus time variance for the IF and LNP models, across cells. (d) Average pairwise distance between spike trains, as a function of timescale of analysis (see Results). Black trace shows the median distance between IF model response and data. Gray trace shows median distance between LNP model response and data. (e) Same as D, for an OFF RGC. (f) Fractional increase in spike-time distance for the IF and LNP models averaged over all cells. Dashed curve: ratio of IF model distance to the average pairwise distance between RGC responses, normalized by the number of spikes from each cell and averaged across cells. Gray curve: same, for LNP model.

Summary statistics of IF model performance and comparison to the LNP model are shown in Fig. 5, for all RGCs examined. Figure 5a shows a comparison of the likelihood of responses to novel stimuli for the IF and LNP models, obtained by using the fitted parameters for each model to compute the probability of the observed responses. Using this metric, the IF model provided a significantly higher likelihood per spike for all cells, in many cases nearly two-fold. Figures 5b and c show comparisons of the similarity of PSTH and PSTV obtained from RGC spike trains and model simulations. In both cases, the IF model outperforms the LNP model for all cells.

To compare the accuracy of model predictions to the intrinsic variability in RGC spike trains, we also applied a previously-used summary measure of distances between spike trains (Victor and Purpura, 1997). This distance is the minimum cost of transforming one spike train into another using the elementary operations of adding, deleting, and shifting spikes. A timescale parameter expresses the cost of shifting per unit time relative to that of adding or deleting. Although this is an imperfect measure of model performance because it neglects any effect of stimulus or spike train history on the probabilistic cost of shifting spikes (e.g. it would not penalize a model for ignoring the refractory period), it is easy to compute and provides a direct benchmark for comparing the performance of present and future models.

The solid curves in Fig. 5d-f show the average distance between pairs of RGC responses to repeated presentations of the same stimulus. This provides a measure of intrinsic variability in spike trains. The distance falls monotonically as a function of timescale of analysis (Fig. 5d,e). Dashed and gray curves indicate distances between simulated IF and LNP model spike trains and recorded RGC spike trains. This provides a measure of the discrepancy between model predictions and data. The IF model distances were systematically higher than the intrinsic variability, indicating that IF model responses differ noticeably from RGC spike trains. However, across a wide range of timescales (1-100 ms) and in all cells, IF model distances were smaller than LNP model distances. Figure 5f shows a summary of the IF and LNP model distances expressed as a fraction of the intrinsic variability, averaged over all cells. IF model error exceeds intrinsic variability by up to $\sim 20\%$; LNP model error is roughly three-fold higher for most time scales.

We also sought to examine IF model performance at predicting responses to stimuli not drawn from the class of fixed-contrast white noise stimuli. Figure 6 shows repeated responses of an ON and and an OFF retinal ganglion cell to brief positive and negative flashes of different intensities. In this case, the IF model parameters were fit using one minute of response to binary white noise stimuli of two different contrasts (12% and 48%). Despite the fact that the flash stimuli are non-stationary and span a wide range of contrasts (12% to 96%, positive and negative), the IF model does a reasonably good job predicting the RGC responses. The rasters of the simulated IF model exhibit visual agreement with those of the ON and OFF RGCs, and provide an accurate prediction of both mean and variance of the RGC responses. Across 9 cells (5 OFF, 4 ON), the IF model accounts for between 73% and 96% (mean of 84%) of the variance in the PSTH of RGC responses. This compares with a range of 36% to 81% (mean of 68%) for the predictions of the LNP model (data not shown).



Figure 6: IF model prediction of responses to brief flashes. a-b: ON cell responses and IF model predictions. c-d: OFF cell responses and IF model predictions. The flash stimuli (shown above each raster) consisted of a grey screen followed by a brief (25 ms) flash. Flash contrasts were +12, +24, +48, and +96% (*pos. flashes*: a,c) and -12, -24, -48 and -96% (*neg. flashes*: b,d). Below each stimulus is a raster showing 25 responses of the RGC (black), below which is a raster showing 25 simulated responses of the IF model (red). Below each pair of rasters is a plot of the PSTH of the RGC and IF model on the same axes. For these two cells, the IF model accounted for 76% and 90% of the PSTH variance for the ON and OFF cells, respectively.

Spike Timing Precision

The accurate descriptions of RGC spike trains provided by the IF model present an opportunity to examine the origins of spike timing precision, which has been widely discussed in recent studies. RGCs are capable of firing spikes precisely time-locked to the onset of a stimulus (Reich et al., 1998; Berry and Meister, 1998; Keat et al., 2001; van Rossum et al., 2003; Uzzell and Chichilnisky, 2004). In some cases, the variation in the onset time of spiking across repeated stimulus presentations is as low as ≈ 1 ms. Although precise timing during periods of rapid firing may be explained by action potential refractoriness (Berry and Meister, 1998; Liu et al., 2001; Uzzell and Chichilnisky, 2004), the origin and significance of the precision in firing remains difficult to assess, particularly in the case extracellularly recorded spike data.

A simple hypothesis about the origin of such precision (Bryant and Segundo, 1976) is that it is determined by the slope of the membrane voltage as it crosses threshold. If voltage crosses threshold with a steep slope, then the noise current exerts little influence on the time of the spike; if voltage crosses threshold with a shallow slope, noise has a greater influence. Figure 7 illustrates this intuition graphically with an example from an RGC response raster. The raster shows two adjacent periods of rapid firing. The histograms below show the distributions of the time of the first spike in each period of firing, and indicate that the first period of firing exhibited more precise timing. The trace below shows the (noiseless) voltage response of the IF model, obtained using the stimulus and the parameters fit for this cell. The slope of V at threshold crossing is indicated by thick gray lines. The period of firing that begins with a steeper voltage slope exhibits much more precise timing. Qualitatively, this example supports the simple hypothesis.

Although this idea about the origin of spike timing precision has been explored in both intracellular experiments and detailed modeling studies (Mainen and Sejnowski, 1995; Reich et al., 1997; Banerjee, 2001; van Rossum et al., 2003; Gutkin et al., 2003), it has not to our knowledge been connected with recent extracellular measurements and information-theoretic analyses focusing on the phenomenology and coding significance of precision (Berry et al., 1997; Berry and Meister, 1998; Koch et al., 2004; Uzzell and Chichilnisky, 2004). The IF model, which makes a prediction about the underlying voltage path for any extracellularly-recorded spike train, allows us to examine this connection quantitatively. We performed an analysis of the firing precision obtained for each cell, in all identified firing onsets during a 7-30 second response raster. Firing onsets were defined as periods following a silence of at least 8 ms long across all trials, followed by a spike on at least 80% of trials within a window of 40 ms (see (Uzzell and Chichilnisky, 2004)). The standard deviation of the first spike time during each onset was computed, and compared to the average current produced by the IF model at those times. Figure 7c shows a scatter plot of precision (inverse of standard deviation) as a function of model input current (which is proportional to the slope dV/dt crossing threshold) across 70 such firing onsets for the cell shown in 7a. The model accounted for 89% of the variability in precision for this cell.

Figure 7d shows a similar analysis across all cells in the dataset, plotting correlation coefficients between the predicted voltage slope and first-spike precision for all cells tested. Thus, a significant



Figure 7: Precision of firing onset times. (a) 200 ms portion of stimulus and a response raster showing two periods of firing onset. Below are histograms of the time of the first spike in each event. Standard deviations: 1.3 ms (left) and 4.7 ms (right). (b) simulated voltage response from the fitted IF model with noise set to zero. Tangent at time of firing onset is shown in red. (c) Precision of first spike times (inverse of standard deviation) as a function of the mean current at the time of the first spike, over 70 isolated firing onsets. Correlation coefficient: 0.89. (d) Correlation coefficient between precision and IF model current prediction, as a function of the inverse of the average SD of the first spike time. Open circles denote ON cells and filled circles denote OFF cells.

fraction of the variability in precision was clearly explained by the membrane voltage slope at firing onset. Note that the IF model captures more of the variability in precision for ON cells (open circles) than for OFF cells (filled circles). However, for both ON and OFF cells, spike timing precision is not an intrinsic property of the cell: it varies substantially as a function of the stimulus history, and in a manner predicted by elicited currents and the intrinsic noise in the IF model.

The measure described above provides a useful means of quantifying precision, but is limited by its reliance on a somewhat arbitrary criterion for identifying firing onsets, assumes a univariate measure of precision (standard deviation), and ignores all spikes beyond the first. One indication that this approach is incomplete is that subsequent spikes sometimes exhibit more precise timing than the first spike. Figure 8a shows a period of firing in which the second spike is more precise than the first, a behavior seen in more than one third of the firing onsets recorded from this cell. Figure 8b shows a more unusual example in which the last spike is more precise than the first. Thus, restricting the analysis to the first spike fails to capture important aspects of spike train precision. Furthermore, measures of spike precision are bound up inextricably with the notion of spiking reliability. Attempts to separate precision (jitter in spike time) from reliability (probability of spike occurrence in a particular time interval) require the use of *ad hoc* criteria for defining firing events: the length of silence preceding an event and the fraction of repeats containing a spike necessary to constitute an event. Changes in these criteria will result in the identification of quite different numbers of firing onsets.

Given these difficulties, it is natural to ask whether the IF model can provide a more complete explanation for the precision and reliability of all recorded spikes. Such an account emerges naturally from the likelihood function used in fitting the IF model. The machinery for computing the likelihood of spike trains can be used to compute the probability density for a particular spike time, conditional on the stimulus and spike train history, as shown in Fig. 8c-d. Using the linear kernel \vec{k} and after-current waveform \vec{h} , we can compute the predicted intracellular current $(I_{stim} + I_{sp})$ during the interval. This input, combined with the noise current I_{nse} , determines a probability density over subthreshold voltage P(V) as a function of time, and can be used to compute the probability density of the next spike P(next spike) as a function of time.

Of course, the probability density of the next spike time depends on the particular spike history for that trial, and therefore differs slightly on each trial. Recent studies indicate that this type of history dependence can give rise to distinct patterns of response on different trials, which could be significant for the encoding of visual information across neurons (Fellous et al., 2004; Tiesinga and Toups, 2005). Figure 8e shows an example where the distribution of the next spike time following a period of silence is bimodal, a condition where standard measures of spike time precision are problematic. The bimodality observed here is accurately reflected in the (average) density of the next spike time, computed using the IF model. This indicates that the next-spike-time density encompasses summary measures of reliability as well as precision.



Figure 8: Generalized analysis of timing precision. (a) RGC response raster sorted in order of first spike time. Below: histogram of first spike (gray) and second spike (black) in the event, illustrating higher precision in the time of the second spike than the first. (b) Sorted RGC response raster, with histograms below showing the distribution of the first (gray) and *last* (black) spike in this event. Precision of the last spike is higher than the first. **Right:** Use of the IF model formalism to analyze RGC spike timing precision and reliability. (c) 170 msec stimulus fragment and corresponding RGC spike response during one trial. (d) Probability distribution over subthreshold V for central interspike interval, on a single trial. The likelihood of the next spike time (below) is given by the probability mass crossing threshold at each moment in time. Note that the probability distribution of the next spike time is bimodal. (e) Raster of repeated RGC responses to this stimulus fragment, with rows sorted in order of first spike time. Below: probability density of the next spike, averaged across 25 trials. Black trace shows model prediction, Gray bars show actual distribution.

Decoding of Spike Responses

Perhaps the most important role for a model of RGC responses is to provide a precise description of the visual information transmitted to the brain. The generalized IF model makes it possible to assess the degree to which variability in spike trains imposes limitations on the fidelity of information transmission. Specifically, the model can be used to compute the probability that an observed spike train was elicited by any given stimulus. This provides a powerful method for decoding the information contained in neural spike responses.

One method to illustrate decoding would be to simply use the model to perform stimulus reconstruction from measured spike responses. Given a particular spike response x, we can obtain the probability that it was elicited by a stimulus y via Bayes' rule:

$$P(y|x) = P(x|y)P(y)/P(x),$$
(6)

where P(x|y) is obtained from the model likelihood calculation, P(y) is the prior over the stimulus, and P(x) is a normalizing term, the marginal probability of observing response x. We can perform stimulus reconstruction by choosing the y which is the maximum or the mean of P(y|x). The maximum provides the maximum *a posteriori* (MAP) estimate for the stimulus, while the mean provides the Bayes estimate under a squared loss function.

Rather than performing a full stimulus reconstruction, which requires choosing a prior P(y) and a search for the y which achieves the maximum or mean of eq. 6, we performed a simple illustration of decoding by using the model to discriminate stimuli in a 2-alternative forced-choice (2AFC) experiment. This experiment represents a much simpler demonstration of decoding: we use the model likelihood to distinguish which of *two* stimuli was most likely given an observed neural response, whereas reconstruction consists in selecting the most likely stimulus amongst *all* possible stimuli.

In the 2AFC experiment, illustrated schematically in figure 9a, an observer is presented with two spike trains and two different stimuli, and must decide on the correct pairing of stimuli with elicited responses. The optimal decision rule given a particular model of the response is to use the pairing with the higher likelihood under that model (Green and Swets, 1966). We used the likelihood of spike responses under the IF model to discriminate pairs of stimuli, and by applying this decision rule to each pair of responses obtained over multiple repeats of the stimulus, we obtained a percent correct for the performance of the model at discriminating the pair of stimuli. As a benchmark, we compared this procedure against one in which likelihood was computed (and discrimination performed) using the LNP model. Figure 9b shows a comparison of the performance of the two procedures. The predominance of data above the diagonal indicates that, on average, the IF model provided significantly more accurate discrimination than the LNP model. Thus, stimulus decoding based on the IF model exploits information in temporal patterns of spikes that is not captured by decoding based only on firing rate.



Figure 9: Decoding responses using model-derived likelihoods. (a) Two stimulus fragments and corresponding fragments of RGC response raster. Gray boxes highlight a 50-ms interval of the first row of each response raster. A two-alternative forced-choice (2AFC) discrimination task was performed on these response fragments, where the task was to determine which stimulus gave rise to each response. The IF and LNP models were used to compute the likelihood of these responses given the "correct" and "incorrect" pairing of stimuli and responses, and the pairing with higher likelihood was selected. This discrimination procedure was applied to each row of the response raster and used to obtain a percent correct for the discrimination performance of each model. (b) Discrimination performance of the IF and LNP models. Each point corresponds to the percent correct of a 2AFC discrimination task using two randomly-selected 50-ms windows of the response. Although both models obtain perfect performance (100%) for a majority of such randomly-selected response windows, the scatter of points above the diagonal shows that when discrimination performance is imperfect, the IF model is far better at decoding the neural spike responses.

Discussion

An integrate-and-fire model, generalized to include a linear receptive field, a spike after-current and a noise source, provides an accurate description of the detailed structure of RGC spike trains elicited by visual stimuli. The model relies only on measurements of (extracellularly recorded) spike times, but provides a full description of the hypothesized underlying currents, along with detailed, accurate predictions of spiking behavior. The model can be fit reliably to responses from arbitrary stimuli using a straightforward procedure, and does not require long measurements of responses to repeated or specialized stimuli. Most importantly, the model provides a rough mechanistic intuition about the origins of spike timing precision, as well as an optimal decoding procedure that exploits temporal patterns in spike trains.

Variability, structure, and fidelity of retinal signals

In previous work, the timing precision of firing onsets and the large gaps between periods of firing have been suggested as the basis for a novel interpretation of retinal coding (Berry et al., 1997; Keat et al., 2001). In this view, the retina encodes visual information in discrete firing events whose timing and spike count encode the timing and features of the stimulus, respectively. The IF model provides a more mechanistic explanation of spike train structure. Precisely timed spikes result primarily from stimulation that causes the membrane potential to cross threshold rapidly, leaving little opportunity for voltage noise to influence the time of the spike (Fig 7) (Bryant and Segundo, 1976; Liu et al., 2001; Banerjee, 2001). Similarly, long gaps between periods of firing result from stimuli that effectively suppress spiking by providing strong currents of opposite polarity. Periods of more gradual firing rate modulation, arise from stimuli poorly matched to the linear filter, or following periods of maintained firing. Thus, the encoding process is fundamentally linear, and the precision and structure of spike trains simply reflect the interaction of the stimulus with intrinsic filtering, noise, and spike generation.

In the model, a single noise parameter accounts for all the variability in neural response, and thus summarizes the effects of noise in transduction, synaptic transfer, and cellular integration. This summary measure of response variability may serve as a parsimonious replacement for more phenomenological measures of variability, such as reliability and firing onset precision (Keat et al., 2001; Berry et al., 1997; Uzzell and Chichilnisky, 2004; Reinagel and Reid, 2000). These latter measures require *ad hoc* criteria for defining firing events, and exclude many spikes from analysis (e.g. those which do not occur during an event). Additionally, these measures cannot easily be applied in stimulus regimes where the neural response does not cleanly parse into discrete events (see, e.g., RGC responses to low-contrast flashes, fig. 6). In contrast, the noise parameter of the IF model explains many different measures of response variability (Fig 8), has a rough physiological interpretation in terms of membrane current, and can be used to predict the variability of responses to novel stimuli, in both event and non-event response regimes.

The IF model also provides a potentially significant technique for assessing the fidelity of sensory signals (Fig 9). Because the probability of an observed spike train given any stimulus can be computed directly, the optimal stimulus discrimination procedure is provided explicitly. Such an optimal procedure is essential for a meaningful investigation of the factors that limit sensory performance. Stimulus discrimination based on the IF model was significantly more accurate than discrimination based on the LNP model, indicating that temporal patterns of spikes convey information not captured in the time-varying firing rate. Stimulus decoding based on the IF model also provides a bound on the accuracy with which the brain could decode stimulus information from RGCs. This approach to describing the fidelity of neural coding in the context of a mechanistic model could provide a valuable complement to approaches based on information theory (Bialek et al., 1991).

Limitations of the IF model

Although the IF model offers many practical and theoretical advantages, it has some shortcomings. First, although it is clearly outperforms the LNP model by a variety of measures, the IF model still fails to account for $\approx 10\text{-}25\%$ of the variance in the PSTH of RGCs (Fig. 5b), and deviations from the observed data exceed the variability of repeated responses by up to $\approx 20\%$ (Fig. 5). One likely source of error is the existence of nonlinear stimulus dependencies (Hochstein and Shapley, 1976; Victor and Purpura, 1997; Benardete et al., 1992) not captured by the linear front end. Nonlinearities may be incorporated into the model without compromising its stable fitting properties by expanding the filter \vec{k} to operate on nonlinear functions of the input. In preliminary studies, the inclusion of terms with quadratic dependence on the input led to a 5-10% improvement in the percentage of explained variance. A drawback is that the elaborated model has more parameters to describe the nonlinear stimulus dependence, and thus requires more data for estimation of those parameters.

Second, the fitting procedure tends to set the noise parameter somewhat higher than is necessary to account for the timing variability observed during periods of rapid firing. This is evident in the sorted rasters of Fig. 3, where the repeated interval structure of the RGC response is more regular than that of IF model responses. The discrepancy may result from the fact that the noise parameter accounts for both true variability in neural responses and any approximation errors of the model. Specifically, nonlinear mechanisms not captured by the model require an increase in the noise parameter to keep the observed spike train likelihood from becoming prohibitively small. This shortcoming could be addressed by separately optimizing the noise parameter using an objective function that isolates the stochastic behaviors of the neural response, such as the repeat interval structure (Fig. 3) or the spike-time distance (Fig. 5). A complete solution, however, requires the incorporation of nonlinear stimulus selectivity in the model (see above). This may be particularly important for applying the IF characterization to neurons in cortex.

Third, although the IF model is clearly more realistic than models with Poisson firing, it provides only a simplified description of known mechanisms underlying neural responses. This IF model is based on current rather than conductance inputs, and has linear subthreshold dynamics. Although conductance-based IF and nonlinear Hodgkin-Huxley style models provide much more biophysically realistic descriptions of intracellular dynamics, it is unfortunately the case that the fitting of these models to extracellular data is relatively intractable. In particular, the likelihood in such models is much more difficult to compute and generally contains many local maxima. Another oversimplification of the IF model is the use of a single Gaussian white noise source to summarize all real noise sources (e.g. photon noise, synaptic failure, and channel noise); this assumption makes fitting tractable, but does not allow us to make detailed inferences about the sources of noise.

Extensions

The present findings suggest a variety of extensions. First, the front-end linear filter can be extended to incorporate spatial dependence to model neural responses to spatially (in addition to temporally) varying stimuli. Higher stimulus dimensionality means that the model will require more data for fitting, but the problem remains tractable. In pilot studies, we have obtained reasonable fits of spatiotemporal stimulus filters using 10-15 minutes of data (compared to 50 sec in the current experiment). The spatial component of these stimulus filters does not differ remarkably from that of the spike-triggered average, suggesting that the important differences between the IF and LNP models are restricted to their temporal properties.

Second, several studies (Smirnakis et al., 1997; Chander and Chichilnisky, 2001; Kim and Rieke, 2001; Baccus and Meister, 2002) have revealed slow contrast adaptation in RGCs, and integrateand-fire models can likewise exhibit a form of spike rate adaptation to stimulus variance (Rudd and Brown, 1997). Although a full study of the adaptive properties of the IF model are beyond the scope of this paper, the model's ability to predict responses to multi-contrast flashes indicate that it can account for some forms of adaptation. The shape of the \vec{h} current also gives rise to one simple form of adaptation: the delayed negative lobe of \vec{h} (figure 2b) means that several rapidly occurring spikes will give rise to a significant buildup of negative current, which produces adaptation in the response to a DC step current.

Third, the IF model can be extended to describe neural responses in sensory areas beyond the retina. We have conducted preliminary analyses which suggest that the IF model provides a reasonably accurate description of the responses of V1 simple cells to flickering bars. Additionally, in cases where responses are driven by nonlinear stimulus transformations of a known form, the model can be augmented by fitting to a stimulus already transformed by a given nonlinearity. For example, fitting the model using a quadratic function of the stimulus as input to the linear filter \vec{k} allows the model to exhibit second-order stimulus dependence, such as that observed in V1 complex cells.

Fourth, the model can be extended to the modeling of multi-cell responses by introducing postspike currents between pairs of cells. These currents can be reliably fit to multi-cell data (again, by maximizing likelihood), and preliminary exploration indicates that the the model can reproduce observed correlations in firing between cells. (Pillow et al., 2005). Finally, the fitting method is based on gradient ascent of the likelihood function, and so can be applied to data collected with any sufficiently rich set of stimuli. Thus, unlike reverse correlation approaches to estimating LNP model parameters, IF model characterization does not require white noise stimuli. This raises the possibility of characterizing light responses using stimuli that drive cortical neurons more strongly than white noise, or that more closely approximate the environment in which the visual system normally operates (Reinagel, 2001).

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