Partitioning neuronal variability

Robbe L T Goris¹, J Anthony Movshon¹ & Eero P Simoncelli¹,²

Responses of sensory neurons differ across repeated measurements. This variability is usually treated as stochasticity arising within neurons or neural circuits. However, some portion of the variability arises from fluctuations in excitability due to factors that are not purely sensory, such as arousal, attention and adaptation. To isolate these fluctuations, we developed a model in which spikes are generated by a Poisson process whose rate is the product of a drive that is sensory in origin and a gain summarizing stimulus-independent modulatory influences on excitability. This model provides an accurate account of response distributions of visual neurons in macaque lateral geniculate nucleus and cortical areas V1, V2 and MT, revealing that variability originates in large part from excitability fluctuations that are correlated over time and between neurons, and that increase in strength along the visual pathway. The model provides a parsimonious explanation for observed systematic dependencies of response variability and covariability on firing rate.

Neurons transmit information with sequences of action potentials. These responses are variable—repeated measurements under identical experimental conditions give different spike trains—but the origins of this variability are unknown. If spike generation were variable, it might account for response variability, but in vitro measurements indicate that it is highly reliable¹. Variability in synaptic transmission is another possible source², but its magnitude is also believed to be insufficient to account for the observed variability in spiking responses³,⁴. A more likely explanation is that the variability arises from the accumulation and amplification of small amounts of noise as signals flow through neural circuits⁵. And recent theories propose that the substantial variability in neural responses may arise from the dynamics of recurrent but largely deterministic networks⁶,⁷.

Regardless of its source, characterizing variability with simple stochastic models has proven useful in understanding the nature of neural coding. The simplest stochastic model is a Poisson process, in which spikes occur independently of one another. A hallmark of the Poisson model is that the variance of the spike count in any time interval is equal to the mean. In visual cortex, the spike-count variance typically equals or exceeds the mean but rarely falls below it⁸,⁹. This suggests that Poisson-like behavior is a ‘floor’ state of cortical variability and raises the question of the origin of the excess variance. Arousal, attention, adaptation and other contextual factors are known to modulate sensory responses¹⁰–¹². In typical electrophysiological experiments, some of these may be well controlled, but many are not. The idea that fluctuations in excitability can inflate estimates of neuronal variance has a long history⁸,⁹, and we wondered whether a more directed analysis of single-neuron responses might reveal the effect of these factors.

We formalize this hypothesis in a doubly stochastic response model in which spikes arise from a Poisson process whose rate is the product of ‘drive’ and ‘gain’ (the ‘modulated Poisson model’; Fig. 1a). The drive is a reproducible firing rate response to a sensory stimulus; the gain represents modulatory influences on excitability and can vary across repeated measurements. Under this model, trial-to-trial variability in spike counts can be partitioned into a sum of Poisson point-process variance and variance arising from fluctuations in gain. Likewise, spike-count covariation can be partitioned into point-process covariance and covariance arising from correlated gain fluctuations.

We found that this model provides an excellent account of single-neuron response distributions in macaque visual thalamus and cortex. Inferred gain fluctuations are correlated over long timescales (minutes), are larger in anesthetized than in awake animals and (in anesthetized animals) increase in strength along the visual hierarchy. They are also shared across wide areas of primary visual cortex, while point-process variance is more localized. Together, these results suggest that much of the response variability of sensory neurons arises from fluctuations in excitability that are correlated over time and between neurons and that increase in strength along the visual pathway.

Portions of these results have been presented at conferences (R.L.T. Goris, J.A. Movshon and E.P. Simoncelli, Soc. Neurosci. Abstr. 311.01, 2013; Cosyne Abstr., I-37, 2012), and two other groups have recently presented models similar to the modulated Poisson model (ref. 13 and I.-C. Lin, M. Okun, M. Carandini and K. Harris, Cosyne Abstr., III-11, 2014), though their application and conclusions are somewhat different from our own.

RESULTS

Modulated Poisson framework

Consider the most commonly used rate model, in which the spike count, N, follows a Poisson distribution:

\[
p(N | \mu, \Delta t) = \frac{(\mu \Delta t)^N}{N!} \exp(-\mu \Delta t)
\]

(1)

¹Center for Neural Science, New York University, New York, New York, USA. ²Howard Hughes Medical Institute, New York University, New York, New York, USA. Correspondence should be addressed to E.P.S. (eero.simoncelli@nyu.edu).

Received 21 November 2013; accepted 30 March 2014; published online 28 April 2014; doi:10.1038/nn.3711
Figure 1 The modulated Poisson model accounts for spike count variability. (a) Model diagram. Spikes are generated by a Poisson process whose rate is the product of two signals: a stimulus-dependent drive, \( f(S) \), and a stimulus-independent gain, \( G \), that is assumed to fluctuate slowly relative to the duration of experimental trials. (b) Variance-to-mean relation of the neural responses of a single V1 neuron stimulated with gratings drifting in different directions (gray dots), compared with predictions of the Poisson model (red line) and the modulated Poisson model (blue line). Responses were computed by counting spikes in a 1,000-ms window following response onset. Means and variances were calculated over 125 repetitions of each stimulus. The inset shows this relation measured over variable-duration windows for three drift directions (green, black and orange). Each data point is obtained from a randomly selected epoch of the corresponding raster with duration drawn uniformly from the range 1–1,000 ms (the orange data are taken from the inset raster). (c) Spike count distributions (gray histograms) measured for different stimulus drift directions compared to the best-fitting probability densities of the Poisson (red) and gamma-modulated Poisson (blue) models. (d) Log-probability of the cell responses under the Poisson model (red triangle) and the modulated Poisson model (blue triangle). Histograms illustrate the expected range of the log-probability statistic (computed with a 1,000 run parametric bootstrap) for the Poisson model (red) and the modulated Poisson model (blue). (e) Variance-to-mean relations predicted by the modulated Poisson model and an additive model for weak (light gray) to strong (black) fluctuations in gain.

where \( \mu \) is the mean spike rate and \( \Delta t \) the duration of the counting window. Assume that the rate arises from the product of two positive-valued signals:

\[
\mu = f(S)G
\]  

(2)

where \( f(S) \) is some function of the stimulus and \( G \) is a stimulus-independent gain (Fig. 1a). In this case, the mean and variance of the spike count in a time interval \( \Delta t \) are both equal to \( f(S)G\Delta t \).

If the gain signal is unobserved and varies over repeated trials, the observed spike counts over those trials will have a variance larger than predicted by the Poisson model. Assume the gain has a mean value of 1 but fluctuates randomly on a timescale that is slow relative to the interval over which spikes are counted. In this case, the net distribution of spike counts is a mixture of Poisson distributions (Online Methods) whose variance can be decomposed into two components:

\[
\text{var}[N | S, \Delta t] = f(S)\Delta t + \sigma_G^2 (f(S)\Delta t)^2
\]  

(3)

The first term is the expected variance of the Poisson distribution conditioned on a gain value of 1, which is equal to the mean count \( f(S)\Delta t \). The second term is the variance of the expected spike count conditioned on the stimulus drive and is proportional to the square of the first term, with a proportionality factor equal to the variance of the gain signal, \( \sigma_G^2 \). Thus, in the presence of gain fluctuations, the spike-count variance exceeds the mean by an amount that is proportional to the square of the mean. Note that the spike-count variance does not depend on the firing rate or duration of the counting window per se, but on their product, \( f(S)\Delta t \).

The variance-to-mean relationship expressed in equation (3) is inherent in systems with modulatory effects and depends on gain only through its variance: all gain distributions with the same variance will produce the same variance-to-mean relationship. To fit the model to data, the form of the gain distribution must be specified. If we assume that the gain follows a gamma distribution, the overall spike-count distribution will follow a negative binomial (Online Methods):

\[
p(N | S, \Delta t) = \frac{\Gamma(N + 1/\sigma_G^2)}{\Gamma(N + 1)\Gamma(1/\sigma_G^2)} \times \frac{(\sigma_G^2 f(S)\Delta t)^N}{(\sigma_G^2 f(S)\Delta t + 1)^{N + 1/\sigma_G^2}}
\]  

(4)

where \( \Gamma(.) \) represents the standard gamma function. This distribution is parameterized by the variance of the gain \( \sigma_G^2 \) and the mean spike count \( f(S)\Delta t \), and is readily fit to neural data.

Gain fluctuations increase along the visual hierarchy

We measured responses of neurons in the lateral geniculate nucleus (LGN) and areas V1, V2 and MT to drifting sinusoidal gratings of the preferred size and speed, varying either in spatial frequency (12 spatial frequencies, ranging from 0 to 10 cycles/deg) or in drift direction (16 equally spaced directions). Each grating was presented for 1,000 ms and repeated at least five times. Responses were computed by counting spikes in a 1,000-ms window after response onset. We fit both the Poisson and modulated Poisson models to the responses of each individual neuron.

The variance-to-mean relationship of an example V1 cell is well described by the modulated Poisson model (Fig. 1b). As predicted by equation (3), this relationship does not depend on firing rate or measurement interval per se (inset, Fig. 1b). Comparing histograms of the measured spike counts with the predictions of the models reveals that the modulated Poisson model captures the full response distributions much better than the standard Poisson model (Fig. 1c). When the mean spike count is low, both models produce similar distributions. But as the mean spike count grows, model predictions diverge and the...
Fluctuations in excitability modulate neuronal activity through visual thalamus and cortex. Neurons in LGN, V1, V2 and MT all exhibit super-Poisson variability (Fig. 2a). Within each area, a share comparable to the fraction due to variations in the stimulus drive (47%). The latter is dependent on the set of stimuli and the tuning properties of the neuron. To focus our analysis on the variability across repeated measurements, we consider the portion of within-condition variance that is explained by the excitability fluctuations. For the example neuron in Figure 1, this fraction is 89.6%.

In our model, stronger gain fluctuations lead to a more rapidly accelerating variance-to-mean relationship, which deviates more and more from the Poisson expectation as spike count grows (Fig. 1e). This accelerating relationship implies that the ratio of variance to mean (the Fano factor, commonly used to quantify point process variability) can vary within a single spike train and over short time scales. As such, a single Fano factor provides an incomplete and potentially biased measure of neuronal variability.

In the modulated Poisson model, the rate arises from the product of two positive-valued signals, one a function of the stimulus and the other arising elsewhere. An alternative model adds these signals as originating from three different sources: the stimulus attributes (direction of motion), a Poisson point process and gamma-distributed fluctuations in excitability. To estimate the relative contribution of each source, we used the modulated Poisson model to partition the spike-count variance (Online Methods). Surprisingly, Poisson noise accounts for only a small fraction of the total variance (5.5%). The gain fluctuations account for nearly half of the variance (47.5%), a share comparable to the fraction due to variations in the stimulus drive (47%). The latter is dependent on the set of stimuli and the tuning properties of the neuron. To focus our analysis on the variability across repeated measurements, we consider the portion of within-condition variance that is explained by the excitability fluctuations. For the example neuron in Figure 1, this fraction is 89.6%.

In our model, stronger gain fluctuations lead to a more rapidly accelerating variance-to-mean relationship, which deviates more and more from the Poisson expectation as spike count grows (Fig. 1e). This accelerating relationship implies that the ratio of variance to mean (the Fano factor, commonly used to quantify point process variability) can vary within a single spike train and over short time scales. As such, a single Fano factor provides an incomplete and potentially biased measure of neuronal variability.

In the modulated Poisson model, the rate arises from the product of two positive-valued signals, one a function of the stimulus and the other arising elsewhere. An alternative model adds these signals as originating from three different sources: the stimulus attributes (direction of motion), a Poisson point process and gamma-distributed fluctuations in excitability. To estimate the relative contribution of each source, we used the modulated Poisson model to partition the spike-count variance (Online Methods). Surprisingly, Poisson noise accounts for only a small fraction of the total variance (5.5%). The gain fluctuations account for nearly half of the variance (47.5%), a share comparable to the fraction due to variations in the stimulus drive (47%). The latter is dependent on the set of stimuli and the tuning properties of the neuron. To focus our analysis on the variability across repeated measurements, we consider the portion of within-condition variance that is explained by the excitability fluctuations. For the example neuron in Figure 1, this fraction is 89.6%.

In our model, stronger gain fluctuations lead to a more rapidly accelerating variance-to-mean relationship, which deviates more and more from the Poisson expectation as spike count grows (Fig. 1e). This accelerating relationship implies that the ratio of variance to mean (the Fano factor, commonly used to quantify point process variability) can vary within a single spike train and over short time scales. As such, a single Fano factor provides an incomplete and potentially biased measure of neuronal variability.

In the modulated Poisson model, the rate arises from the product of two positive-valued signals, one a function of the stimulus and the other arising elsewhere. An alternative model adds these signals as originating from three different sources: the stimulus attributes (direction of motion), a Poisson point process and gamma-distributed fluctuations in excitability. To estimate the relative contribution of each source, we used the modulated Poisson model to partition the spike-count variance (Online Methods). Surprisingly, Poisson noise accounts for only a small fraction of the total variance (5.5%). The gain fluctuations account for nearly half of the variance (47.5%), a share comparable to the fraction due to variations in the stimulus drive (47%). The latter is dependent on the set of stimuli and the tuning properties of the neuron. To focus our analysis on the variability across repeated measurements, we consider the portion of within-condition variance that is explained by the excitability fluctuations. For the example neuron in Figure 1, this fraction is 89.6%.

In our model, stronger gain fluctuations lead to a more rapidly accelerating variance-to-mean relationship, which deviates more and more from the Poisson expectation as spike count grows (Fig. 1e). This accelerating relationship implies that the ratio of variance to mean (the Fano factor, commonly used to quantify point process variability) can vary within a single spike train and over short time scales. As such, a single Fano factor provides an incomplete and potentially biased measure of neuronal variability.

In the modulated Poisson model, the rate arises from the product of two positive-valued signals, one a function of the stimulus and the other arising elsewhere. An alternative model adds these signals as originating from three different sources: the stimulus attributes (direction of motion), a Poisson point process and gamma-distributed fluctuations in excitability. To estimate the relative contribution of each source, we used the modulated Poisson model to partition the spike-count variance (Online Methods). Surprisingly, Poisson noise accounts for only a small fraction of the total variance (5.5%). The gain fluctuations account for nearly half of the variance (47.5%), a share comparable to the fraction due to variations in the stimulus drive (47%). The latter is dependent on the set of stimuli and the tuning properties of the neuron. To focus our analysis on the variability across repeated measurements, we consider the portion of within-condition variance that is explained by the excitability fluctuations. For the example neuron in Figure 1, this fraction is 89.6%.

In our model, stronger gain fluctuations lead to a more rapidly accelerating variance-to-mean relationship, which deviates more and more from the Poisson expectation as spike count grows (Fig. 1e). This accelerating relationship implies that the ratio of variance to mean (the Fano factor, commonly used to quantify point process variability) can vary within a single spike train and over short time scales. As such, a single Fano factor provides an incomplete and potentially biased measure of neuronal variability.
Response correlations in the modulated Poisson framework

Trial-to-trial response fluctuations are often correlated among simultaneously recorded neurons. Pairwise response correlations can arise when neurons receive shared sensory input, but also when they are subject to correlated modulatory influences that are not sensory in origin. Our model provides a vehicle for separating the effects of these two contributions. Specifically, for doubly stochastic processes, the spike-count covariance can be decomposed in a manner that is analogous to the variance decomposition introduced in equation (3) (Online Methods):

\[
\text{cov}[N_i, N_j | S, \Delta t] = \tau_{Pij} \sqrt{f_i(S)f_j(S)} \Delta \tau^2 + \tau_{Gij} \sigma_{Gi} \sigma_{Gj} f_i(S)f_j(S) \Delta \tau^2
\]

(5)

where \( \tau_{Pij} \) indicates the point-process correlation (assumed to be independent of stimulus \( S \)) and \( \tau_{Gij} \) the gain correlation. The first term is the expected covariance for the spike counts of neurons \( i \) and \( j \) conditioned on a gain value of 1. The second component is the covariance of the conditional expectations that arises from correlated gain fluctuations and can generate spike-count correlations even when the two point processes are independent. Equation (5) reduces to the expression for spike-count variance (equation (3)) when \( i = j \). The spike-count correlation is obtained by dividing this equation by the square root of the product of the spike-count variances of neurons \( i \) and \( j \), as expressed in equation (3), yielding a complicated dependence of spike-count correlation on stimulus drive and on the variance of the gain signals in the two neurons.

Studies of response correlation typically combine normalized responses across conditions to estimate a single spike-count correlation for a pair of neurons. But equation (5) implies that, even if the point-process and gain correlations are both stable properties of a given cell pair, measured spike-count correlations can vary dramatically with stimulus drive. Specifically, when the geometric mean of the stimulus-driven response of the two neurons is low, the spike-count correlation will approximate the point-process correlation \( \tau_{Pij} \). In contrast, when this geometric mean is high, the spike-count correlation will be dominated by the gain correlation \( \tau_{Gij} \). If \( \tau_{Pij} \) and \( \tau_{Gij} \) differ, the spike-count correlation will depend on the stimulus conditions, and any single estimate will provide an incomplete and potentially biased measure of neuronal covariability.

We analyzed the responses of four populations of simultaneously recorded neurons in the superficial layers of macaque primary visual cortex. The recordings were made over 2.5 h, during which gratings drifting in 72 equally spaced directions were presented for 1,280 ms each, interleaved with a 1,280 ms blank screen and repeated 50 times in random order. We analyzed 379 well-isolated units (62, 94, 87 and 136 from the four data sets). We fit the modulated Poisson model, including the values of the underlying point-process and excitability correlations \( \tau_{Pij} \) and \( \tau_{Gij} \) to the joint responses of pairs of neurons in each data set (Online Methods).

In the actual and predicted response correlations for three example pairs of V1 cells (Fig. 3), the mean spike count of each neuron depended on the direction of motion of the stimulus (Fig. 3a–c), and the overall spike-count correlation estimated for each pair from
the combined normalized responses ($r_{ac}$) was small and positive. But estimates of spike-count correlation computed separately for different response levels reveal a variety of different behaviors, including a decrease with mean response strength (Fig. 3d), an independence of response strength (Fig. 3e) or an increase with response strength (Fig. 3f). The modulated Poisson model can mimic each of these behaviors and, more generally, can predict the spike-count correlation as a function of the mean responses of the two neurons (Fig. 3g–i).

The structure of correlations in primary visual cortex

V1 spike-count correlations have been shown to depend on cortical distance and tuning similarity, but, as revealed by the examples in Figure 3, this measure mixes two underlying sources of correlation. We wondered whether these two sources might be differently structured across the neuronal population.

We examined how point-process correlation and excitability correlation depend on electrode separation and on the similarity of tuning (Fig. 4 and Online Methods). Point-process correlations were on average smaller than excitability correlations ($P < 0.001$, $F_{1,34} = 24.16$, analysis of covariance) and fell more rapidly with electrode distance ($P < 0.001$, $F_{1,34} = 18.05$). Note that neurons recorded on the same electrode (the leftmost datum in Fig. 4a,b) might have inflated the significance of the latter difference, since they are vulnerable to spike-sorting errors that can alter the measured correlation. Here, we found that excluding this datum did not alter the conclusion ($P < 0.001$, $F_{1,32} = 41.17$). Analysis of tuning similarity effects (Fig. 4c) revealed that the average point-process and excitability correlations both decrease gradually with tuning correlation ($P = 0.6$, $F_{1,14} = 0.28$) (Fig. 4d).

In summary, we find that the relationship between response correlations and mean responses in V1 depends on both interneuronal distance and tuning similarity: spike count correlations are most likely to decrease with response strength for nearby neurons with similar tuning (Fig. 3d). As interneuronal distance or tuning dissimilarity grows, response correlations initially tend to be independent of the mean responses (Fig. 3e) and eventually exhibit an increasing relation (Fig. 3f).

The temporal structure of gain fluctuations

We assume that the Poisson point-process variance is independent across trials, so all temporal structure in neural responses that is not explained by the stimulus must arise from structure in the gain signal. As normalized spiking activity evolved over 2.5 h for three simultaneously recorded neurons (Fig. 5a), response strength rose and fell on a time scale of minutes, suggesting that excitability varies slowly. This is consistent with the slow decay of the autocorrelation of the gain (Fig. 5b), computed from temporal correlations in the trial-by-trial responses (Online Methods). At a time lag of 5 s, 307 of 379 neurons had a positive autocorrelation ($t$-test, $P < 0.05$). For longer time lags, this fraction gradually decreased (at a time lag of 2 min, $r > 0$ for 185 of 379 neurons; at a time lag of 10 min, this fraction dropped to 81 of 379). Although the average profiles for the four data sets differ in detail, they share a slow falloff, indicating that changes in excitability persist for many minutes (Fig. 5c).

Trial-to-trial changes in excitability are often correlated across neurons (Fig. 4). It might therefore be expected that the slow component of excitability fluctuations is also shared across neurons. However, across neurons, excitability correlations decreased dramatically at all time lags exceeding 0 s (Fig. 5d). The slow component of excitability fluctuations thus appears to be local.

Effects of anesthesia

We have shown that neural response variability in visual cortex of anesthetized macaques originates largely from excitability fluctuations.
Poisson spiking models have been common in neuroscience since the 1950s (refs. 24–26) and are implicit in all analyses in which responses are summarized solely with mean spike counts. They provide the simplest statistical description of the data, are readily fit to data and capture the basic fact that spike-count variances grow with the mean. Poisson processes can mimic the spiking behaviors of integrate-and-fire models, as well as the responses of model neurons embedded in cortical networks with balanced configurations of excitatory and inhibitory inputs. But neuronal responses generally exhibit super-Poisson variability, and the model presented here provides a simple but effective means of explaining this behavior. Some authors have reported sub-Poisson variability, especially in brief time bins immediately after a sudden stimulus onset. This is likely a consequence of neuronal refractoriness, which could be naturally incorporated into our modeling framework through modulatory spike-feedback terms in a generalized linear model.

Our work complements recent work on the temporal dynamics of firing-rate variability. One study used the mean-matched Fano factor as a proxy for firing-rate variability, but it did not explore the dependence of this value on the mean. Another estimated firing-rate variability by subtracting an estimate of the point-process variance from the measured spike-count variance. The estimate of point-process variance is taken from the minimum of an ensemble of experimental measurements and is therefore vulnerable to the well-known problems associated with estimating extreme values of sample distributions. In contrast, we found it useful to make explicit distributional assumptions: specifically, that gain is drawn from a gamma distribution and that the spikes arise from a Poisson process. The resulting negative binomial spike-count distribution (a particular continuous mixture of Poisson distributions) is easily fit to data and validated (Fig. 1d; see also ref. 37), and its parameterization as a modulated Poisson distribution provides a natural interpretation in terms of excitability fluctuations. The spike-count distribution bears some resemblance to the discrete mixture-of-Poissons model that has been successfully used to account for super-Poisson variability, but it has the added advantages that it has far fewer parameters and provides an interpretation in terms of modulatory input. It is important to note, however, that our analyses of joint behaviors are based solely on correlations. A full model for joint neural responses would allow better fitting and validation, but it requires an explicit description of both the joint point-process distribution and the joint distribution of gain signals.

Our analyses indicate that correlated fluctuations in gain account for a substantial portion of spike-count correlation. This has been long suspected, but it has been difficult to segregate the effects of shared sensory inputs from shared modulatory influences. Our model provides an explicit method for achieving this (equation (5)). In V1, this analysis revealed that point-process correlation and excitability correlation had different structures. Point-process correlations decreased rapidly with cortical distance, while excitability correlations changed less. Both point-process and excitability correlations increased with tuning similarity. Together, these patterns suggest that point-process correlations are caused by inputs that are shared within local functional circuits while excitability correlations likely

---

**Figure 6** Analysis of spike-count variance for a population of MT neurons recorded in awake, behaving macaques. (a) Variance-to-mean relation for 307 MT cells. Each data point illustrates the mean and variance of the spike count in a 2,000-ms window of one cell for one stimulus condition. (b) Distribution of stimulus-independent fluctuations in gain, summarized by the coefficient of variation of the gain (top) and fraction of within-condition variance explained by gain fluctuations (bottom). (c) The autocorrelation function of the gain, averaged across units (trials are assumed to be separated by 5 s).
arise from fluctuations in modulatory signals that are shared across larger populations of neurons (for example, sensory adaptation, metabolic resource availability, attentional signals, reward signals).

The analysis in Figure 2 indicates that the portion of spike-count variability attributable to modulatory fluctuations increases as one ascends the visual hierarchy. This presumably depends on the context in which responses are measured, including the choice of stimuli, the presence or absence of anesthesia and the cognitive state of the animal. For example, attentional mechanisms are widely believed to act by increasing the gain of neurons involved in a task. Relative to conditions of uncontrolled attention, we might expect that this would lead to a net increase in the mean, but a decrease in the variance, of the modulatory input. Under the modulated Poisson model, this would result in an increase in spike count, accompanied by a decrease in Fano factor, as has been reported in area V4 (ref. 39).

The corresponding predictions for spike-count correlation are more complicated, since the relative contributions of the point-process and gain correlations depend on the stimulus drive to each cell, as well as the variance of the gain of each cell (equation (5)). But if the gain correlation is larger than the point-process correlation and the two cells are receiving similar stimulus drive, our model predicts that a reduced modulatory variance would decrease spike-count correlation, as has also been observed

The modulated Poisson model has broad implications for the characterization of neuronal response variability. It is common practice to summarize variability with the Fano factor (the ratio of the spike-count variance to its mean). This measure is highly dependent on the conditions under which responses are measured. For example, the Fano factor for the neuron in Figure 1b would be near 1 if estimated from trials in which the response was low (less than 1 spike per time bin), but would be significantly higher if estimated from trials with larger responses. Thus, the Fano factor, by itself, does not provide a reliable summary of neural variability. In contrast, spike-count variability of the modulated Poisson model may be decomposed into a sum of two distinct values, one corresponding to the Poisson component (which is always equal to the mean) and the second arising from unobserved modulatory influences, which grows as the square of the mean (equation (3)). The amplitude of this second term, which corresponds to the variance of the gain, provides a compact summary of variability exceeding that of the Poisson model.

An analogous issue arises with the common practice of summarizing the covariation of pairs of neurons with spike-count correlation. This measure can depend critically on the conditions under which responses are measured and thus provides a potentially biased summary of covariability. Indeed, previous studies have noted that neural correlations can increase with firing rate15 and that this behavior can be explained by models in which spikes arise when a Gaussian-distributed membrane voltage crosses a threshold15,40,41. Our model is quite different but accounts for this phenomenon, as well as the more diverse range of behaviors seen across different cell pairs, by summarizing covariation with two distinct values. One value represents the point-process correlation (which could arise from noise in common sensory inputs32) and the other the correlation of the modulatory influences. Whether response correlations primarily reflect the former or the latter depends on both the magnitude of the stimulus drive and the variance of the gain signals in the two neurons.

This insight suggests a resolution for a recent controversy regarding the nature and magnitude of cortical correlations. Despite decades of experimental evidence that cortical cells exhibit modest but significant spike-count correlations, on the order of 0.1–0.3 (ref. 15), a recent study in macaque V1 found correlations to be much closer to zero and concluded that previously reported values arose from confounding experimental effects17. However, owing to modest firing rates and short counting windows, spike counts were rather small in this study and Fano factors were closer to 1 than in most other studies13.

The interpretation arising from our model and data is that the correlation values reported in ref. 17 primarily reflect point-process correlations, which are dominant at low spike counts and fall rapidly with cortical distance, whereas most previously reported values reflect correlated modulation, which is only evident at high spike counts but is generally more substantial and falls more slowly with cortical distance. Consistent with this, ref. 17 also reported substantially larger spike count correlations for the subset of neuronal pairs with the highest responses.

Our analysis suggests that fluctuations in gain are correlated over long time scales within but not across neurons. This implies that the mechanisms underlying slow drifts in response gain differ from the mechanisms that give rise to instantaneous gain correlations. The local nature of these drifts rules out the possibility that they result simply from global state changes induced by factors such as anesthesia or arousal. That said, comparison of responses in area MT of behaving and anesthetized macaque revealed that cortex is less stable under anesthesia: fluctuations in gain were significantly stronger and slower in the anesthetized cortex.

What are the implications of our findings for understanding the representation of sensory information in the brain? We believe that the gradual increase in the strength of modulatory fluctuations along the visual pathway reflects the gradual transformation and combination of visual signals with information from other sources, such as other sensory inputs and top-down signals representing attention, arousal, metabolic state, reward expectations, emotional state and so forth. Regardless of the source of modulatory variability, its increasing weight as information ascends cortical pathways raises the question of why it does not overwhelm the sensory information encoded in higher-level visual areas5, with deleterious effects on the accuracy of behaviors that arise from decoding those neural responses43,44. It is perhaps worth noting that information encoded in the relative responses of groups of neurons will not be affected by common modulatory changes, such as those resulting from attentional focus11. Resolving this mystery is a critical step in understanding how stable perception of the visual world can arise from apparently volatile neural activity, a step that should be made easier by the principled and proven analysis presented here.

METHODS
Methods and any associated references are available in the online version of the paper.

ACKNOWLEDGMENTS
We are grateful to R. Kumbhani and N. Rabinowitz for discussions and to members of the Movshon laboratory for sharing their data. This work was supported by US National Institutes of Health grants EY04440, EY022428, the Howard Hughes Medical Institute and postdoctoral fellowships from the Fund for Scientific Research of Flanders and the Belgian American Educational Foundation awarded to R.L.T.G.

AUTHOR CONTRIBUTIONS
R.L.T.G., J.A.M. and E.P.S. designed research; R.L.T.G. analyzed data; and R.L.T.G., J.A.M. and E.P.S. wrote the paper.

COMPETING FINANCIAL INTERESTS
The authors declare no competing financial interests.

Reprints and permissions information is available online at http://www.nature.com/reprints/index.html.
1. Mainen, Z.F. & Sejnowski, T.J. Reliability of spike timing in neocortical neurons. Science 268, 1503–1506 (1995).
2. Allen, C. & Stevens, C.F. An evaluation of causes for unreliability of synaptic transmission. Proc. Natl. Acad. Sci. USA 91, 10380–10383 (1994).
3. Softky, W.R. & Koch, C. The highly irregular firing of cortical cells is inconsistent with temporal integration of small EPSPs. J. Neurosci. 13, 334–350 (1993).
4. Stevens, C.F. & Zador, A. When is an integrate-and-fire neuron like a Poisson neuron? in Advances in Neural Information Processing Systems Vol. 8 (eds. Mozer, M., Touretzky, D.S. & Hasselmo, M.) 103–109 (MIT Press, 1996).
5. Shadlen, M.N. & Newsome, W.T. The variable discharge of cortical neurons: implications for connectivity, computation, and information coding. J. Neurosci. 18, 3870–3896 (1998).
6. van Vreeswijk, C. & Sompolinsky, H. Chaos in neuronal networks with balanced excitatory and inhibitory activity. Science 274, 1724–1726 (1996).
7. Vogels, T.P. & Abbott, L.F. Signal propagation in networks of integrate-and-fire neurons. J. Neurosci. 25, 10786–10795 (2005).
8. Tomko, G.J. & Crapper, D.R. Neuronal variability: non-stationary responses to identical visual stimuli. Brain Res. 79, 409–418 (1974).
9. Tolhurst, D.J., Movshon, J.A. & Thompson, I.D. The dependence of response amplitude and variance of cat visual cortical neurones on stimulus contrast. Exp. Brain Res. 41, 414–419 (1981).
10. Kato, H.K., Chu, M.W., Isacson, J.S. & Komiyama, T. Dynamic sensory representations in the olfactory bulb: modulation by wakefulness and experience. Neuron 76, 962–975 (2012).
11. Luck, S.J., Chelazzi, L., Hillyard, S.A. & Desimone, R. Neural mechanisms of spatial selective attention in areas V1, V2, and V4 of macaque visual cortex. J. Neurophysiol. 77, 24–42 (1997).
12. Benuccio, A., Saleem, A.B. & Carandini, M. Adaptation maintains population homeostasis in primary visual cortex. Nat. Neurosci. 16, 724–729 (2013).
13. Eckner, A.S. et al. State dependence of noise correlations in macaque primary visual cortex. Neuron 82, 235–248 (2014).
14. Churchland, A.K. et al. Variance as a signature of neural computations during decision-making. Neuron 69, 818–831 (2011).
15. Cohen, M.R. & Kohn, A. Measuring and interpreting neuronal correlations. Nat. Neurosci. 14, 811–819 (2011).
16. Brody, C.D. Correlations without synchrony. Neural Comput. 11, 1537–1551 (1999).
17. Ecker, A.S. et al. Decorrelated neuronal firing in cortical microcircuits. Science 327, 584–587 (2010).
18. Zohary, E., Shadlen, M.N. & Newsome, W.T. Correlated neuronal discharge rate and its implications for psychophysical performance. Nature 370, 140–143 (1994).
19. Smith, M.A. & Kohn, A. Spatial and temporal scales of neuronal correlation in primary visual cortex. J. Neurosci. 28, 12921–12930 (2008).
20. Graf, A.B.A., Kohn, A., Jazayeri, M. & Movshon, J.A. Decoding the activity of neuronal populations in macaque primary visual cortex. Nat. Neurosci. 14, 239–245 (2011).
21. Pillow, J.W., Shlens, J., Chichilnisky, E.J. & Simoncelli, E.P. A model-based spike sorting algorithm for removing correlation artifacts in multi-neuron recordings. PLOS ONE 8, e62123 (2013).
22. Britten, K.H., Shadlen, M.N., Newsome, W.T. & Movshon, J.A. The analysis of visual motion: a comparison of neuronal and psychophysical performance. J. Neurosci. 12, 4745–4765 (1992).
23. Sherman, S.M. & Guillery, R.W. On the actions that one nerve cell can have on another: distinguishing “drivers” from “modulators”. Proc. Natl. Acad. Sci. USA 95, 7121–7126 (1998).
24. Kuffler, S.W., Fitzugh, R. & Barlow, H.B. Maintained activity in the cat’s retina in light and darkness. J. Gen. Physiol. 40, 685–702 (1957).
25. Grossman, R.G. & Wiestrern, L.J. Discharge patterns of neurons in cochlear nucleus. Science 134, 99–101 (1961).
26. Siebert, W.M. Frequency discrimination in auditory system—place or periodicity mechanisms? Proc. IEEE 58, 723–730 (1970).
27. Geisler, W.S. & Albrecht, D.G. Bayesian analysis of identification performance in monkey visual cortex: nonlinear mechanisms and stimulus certainty. Vision Res. 35, 2723–2730 (1995).
28. Chichtrand, M.M. et al. Stimulus onset quenches neural variability: a widespread cortical phenomenon. Nat. Neursc. 13, 369–378 (2010).
29. Gure, M., Beylln, A. & Snodderly, D.M. Response variability in primary visual cortex (V1) of alert monkey. J. Neurosci. 17, 2914–2920 (1997).
30. Gershon, E.D., Wiener, M.C., Latham, P.E. & Richmond, B.J. Coding strategies in monkey V1 and inferior temporal cortices. J. Neurophysiol. 79, 1135–1144 (1998).
31. Oram, M.W., Wiener, M.C., Lestienne, R. & Richmond, B.J. The stochastic nature of precisely timed spike patterns in visual system neural responses. J. Neurophysiol. 81, 3021–3033 (1999).
32. Barbieri, R., Quirk, M., Frank, L.M., Wilson, M.A. & Brown, E.N. Construction and analysis of non-Poisson stimulus-response models of neural spike train activity. J. Neurosci. Methods 105, 25–37 (2001).
33. Kara, P., Reinagel, P. & Reid, R.C. Low response variability in simultaneously recorded retinal, thalamic, and cortical neurons. Neuron 27, 635–646 (2000).
34. Amassigham, A., Chen, T.-L., Geman, S., Harrison, M. & Sheinberg, D. Spike count reliability and the Poisson hypothesis. J. Neurosci. 26, 801–809 (2006).
35. Truccolo, W., Eden, U.T., Fellows, M.R., Donoghue, J.P. & Brown, E.N. A point process framework for relating neural spiking activity to spiking history, neural ensemble, and extrinsic covariate effects. J. Neurophysiol. 93, 1074–1089 (2005).
36. Pillow, J.W. Spatio-temporal correlations and visual signaling in a complete neuronal population. Nature 454, 995–999 (2008).
37. Pillow, J.W. & Scott, J.G. Fully Bayesian inference for neural models with negative-binomial spiking, in Advances in Neural Information Processing Systems Vol. 25 (eds. Bartlett, P., Pereira, F.C.N., Burges, C.J.C., Bottou, L. & Weinberger, K.Q.) 1907–1915 (MIT Press, 2012).
38. Wiener, M.C. & Richmond, B.J. Decoding spike trains instant by instant using order statistics and the mixture-of-Poissons model. J. Neurosci. 23, 2394–2406 (2003).
39. Cohen, M.R. & Maunsell, J.H.R. Attention improves performance primarily by reducing interneuronal correlations. Nat. Neurosci. 12, 1594–1600 (2009).
40. Dorn, J.D. & Ringach, D.L. Estimating membrane voltage correlations from extracellular spike trains. J. Neurophysiol. 89, 2271–2278 (2003).
41. de la Rocha, J., Doiron, B., Shea-Brown, E., Josic, K. & Reyes, A. Correlation between neural spike trains increases with firing rate. Nature 448, 802–806 (2007).
42. Vidne, M. et al. Modeling the impact of common noise inputs on the network activity of retinal ganglion cells. J. Comput. Neurosci. 33, 97–121 (2012).
43. Goris, R.L.T., Putzeys, T., Wagemans, J. & Wichmann, F.A. A neural population model for visual pattern detection. Psychol. Rev. 120, 472–496 (2013).
44. van den Berg, R., Shin, H., Chou, W.-C., George, R. & Ma, W.J. Variability in encoding precision accounts for visual short-term memory limitations. Proc. Natl. Acad. Sci. USA 109, 8780–8785 (2012).
ONLINE METHODS

Surgical preparation. We recorded from 18 anesthetized, paralyzed, adult macaque monkeys of either sex (3 Macaca nemestrina, 1 M. mulatta and 14 M. cynomolgus). Our standard procedures for the surgical preparation of animals and single-unit recordings have been reported in detail previously43. Briefly, experiments typically lasted 5–6 d, during which we maintained anesthesia with infusion of sufentanil citrate (6–30 μg kg⁻¹ h⁻¹) and paralysis with infusion of vecuronium bromide (Norcuron; 0.1 mg kg⁻¹ h⁻¹) in isotonic dextrose-Normosol solution. We monitored vital signs (heart rate, lung pressure, end-tidal pCO₂, EEG, body temperature, urine flow and osmolality) and maintained them within appropriate physiological ranges. Pupils were dilated with topical atropine. The eyes were protected with gas-permeable contact lenses and refracted with supplementary lenses chosen through direct ophthalmoscopy. At the conclusion of data collection, the animal was killed with an overdose of sodium pentobarbital. All procedures were conducted in compliance with the National Institute of Health Guide for the Care and Use of Laboratory Animals and with the approval of the New York University Animal Welfare Committee.

Unit recording. Extracellular recordings were made with quartz-platinum-tungsten microelectrodes (Thomas Recording), advanced mechanically into the brain through a craniotomy and small durotomy. Electrode insertion angle and location varied across experiments, depending on the targeted area. We distinguished V1 from V2 on the basis of depth from the cortical surface and changes in the receptive field location of the recorded units. Area MT was identified from the brisk direction-selective responses of isolated neurons. We made recordings from every single unit with a spike waveform that rose sufficiently above noise to be isolated. Stimuli were presented in random order. Data are reported from every unit for which we completed at least five repetitions.

Visual stimulation. We presented visual stimuli on a gamma-corrected CRT monitor (Eizo T966; mean luminance, 33 cd/m²) at a resolution of 1,280 × 960 with a refresh rate of 120 Hz. Stimuli were presented using Expo software (http://corevision.cns.nyu.edu/expo/) on an Apple Macintosh computer. For each isolated unit, we first determined its ocular dominance and occluded the non-preferred eye. We presented circularly windowed sinusoidal gratings stimuli to map each cell’s receptive field, determined its preferred size and speed, and then measured selectivity for orientation or spatial frequency.

Analysis of single-electrode recordings. Responses were computed by counting spikes in a 1,000-ms window following response onset. We estimated latency for each cell by choosing the latency that maximized the variance of the tuning curve44. The modulated Poisson model describes a doubly stochastic process (also known as a Cox process47), in which spiking responses are generated by a Poisson process whose rate is the product of drive and gain (equations (1) and (2)). To make the model identifiable, we assume that the gain, G, is constant within trials and is distributed across trials according to a gamma distribution with a mean of one and variance of \( \sigma_G^2 \):

\[
p(G, r, s) = G^{-1} \exp(-G/s) 
\]

with shape parameter \( r = 1/\sigma_G^2 \) and scale parameter \( s = \sigma_G^2 \). The mean spike count of the modulated Poisson model is the product of the gain, G, the stimulus driven rate, f(S), and the bin duration, Δt, and is thus also distributed according to a gamma distribution, with parameters \( r = 1/\sigma_G^2 \) and \( s = \sigma_G^2 f(S)Δt \). The spike-count distribution is a gamma mixture of Poisson distributions, and marginalizing over the gain variable yields a negative binomial distribution48 (equation (4)) for the spike count:

\[
p(N, r, s) = \frac{G(N + 1)\Gamma(N + r)}{\Gamma(N + 1)\Gamma(r)} \left( \frac{1}{1 + s} \right)^N \left( \frac{s}{1 + s} \right)^r \]

With parameters \( r \) and \( s \) described above, this distribution has a mean of \( rs = f(S)Δt \) and a variance of \( rs + rs^2 = f(S)Δt + \sigma_G^2 f(S)^2 Δt^2 \), as expressed in equation (3). The negative binomial can also be derived as a Poisson-gamma mixture of Gaussian distributions49, a form that allows efficient inference50 but lacks the modulatory interpretation provided here.

We fit both the ordinary and modulated Poisson models to the responses of each individual neuron. The maximum likelihood estimator of the parameter \( \mu \) that characterizes a Poisson distribution is the sample mean, which we estimate separately for each stimulus condition. The modulated Poisson model has two parameters \( (r \) and \( s \)) that are related to the stimulus drive and the variance of the gain (see above). We used a simplex algorithm (the Matlab function ‘fminsearch’) to search for the value of \( (r, s) \) that minimized the negative log likelihood of the full set of observed spike counts.

To assess the models’ relative goodness-of-fit, we performed a 100-fold cross-validation. We fitted both models to a data set consisting of all trials except for one randomly chosen trial per stimulus condition and compared the average log-probability of the held-out data under each model (Fig. 2b). To evaluate the models’ absolute goodness of fit, we compared the log-probability of the data with that of a large number of simulated data sets drawn from the fitted model. Each simulated data set had the same number of repeats as the observed data. We consider a model fit acceptable if the log-probability of the real data lies within the central 95% of the distribution of log-probability of the simulated data. This test can detect both underdispersion and overdispersion.

We used the modulated Poisson model to estimate the fraction of spike-count variance that arises from the gain signal. As is standard in ANOVA, one can partition the sum of squares into components arising from variations in the stimulus (\( S_{\text{stim}} \)), the gain signal (\( S_G \)) and the point process (\( S_{\text{pp}} \)):

\[
\sum_k (N_k - \bar{N})^2 = \sum_k (N_k - \bar{N}_k)^2 + \sum_k (\bar{N}_k - \bar{N})^2
\]

where \( N_k \) is the spike count on the \( k \)th trial, \( \bar{N}_k \) is the spike count averaged over those trials in which the presented stimulus was the same as that of the \( k \)th trial, and \( \bar{N} \) is the spike count averaged over all trials. The second line follows from the first given the spike-count variance of the modulated Poisson model, as expressed in equation (3). The fraction of within-condition variance that arises from excitability fluctuations is given by \( S_G/S_{\text{pp}} \).

Analysis of multielectrode array recordings. Full details regarding these data sets may be found in ref. 20. Briefly, an array containing 96 fixed electrodes was used to record from single units in the superficial layers of macaque primary visual cortex. The animal was anesthetized and stimulated with sinusoidal gratings. We included all units that could be tracked throughout the entire experiment and whose mean response exceeded 2 spikes/s for at least one stimulus condition. We opted to leave out one set of array recordings (data set 3 in ref. 20) because the responses exhibited uncharacteristically strong correlations across time and neurons; we suspect the cortical surface was still recovering from array insertion. In our modeling framework, pairwise response correlations can result from both correlated point processes and correlated gain fluctuations. To separate these two sources, we used the covariance decomposition formula in equation (5) and searched for the point-process correlation and gain correlation that maximized the likelihood of the observed response relations. Rather than fitting response correlations directly, we performed the optimization on \( z \)-transformed spike-count correlations:

\[
z = \frac{1}{2} \left( \frac{1 + r}{1 - r} \right)
\]

where \( r \) is the spike-count correlation and \( S \) is the natural logarithm. The advantages of this Fisher transformation include a quicker convergence to normality and variance stabilization. We used a simplex algorithm to find the \( (G_{\text{pp}}, f_{\text{pp}}) \) values that maximized the likelihood of the observed response correlations under a normal residual distribution. This approach worked well for most neuronal pairs (Fig. 3), but the model parameters are not always well constrained. For instance, when the modulation of the direction tuning curve is weak for both neurons, many parameter combinations will yield a similar likelihood. For the population analysis (Fig. 4), we therefore took the uncertainty associated with the parameter estimates into account. We modeled the population distribution of the mean parameter values with a bivariate normal distribution and searched for the parameter values that maximized the likelihood across all neuronal
pairs. Poorly constrained pairs of neurons have a flatter likelihood function and are therefore down-weighted in this analysis. Although we consider this approach statistically preferable, simple averaging of the parameter estimates yielded similar results.

To study the temporal structure of the gain signal, we estimated its autocorrelation function. We first removed the stimulus-induced temporal structure by subtracting the appropriate stimulus-elicited mean response from the observed responses. We then estimated the remaining covariance between pairs of responses separated by different time lags. To obtain the autocorrelation of the gain signal at a given time lag, this covariance is normalized by the estimated variance of the gain signal, $S_g/(N - 1)$.

45. Cavanaugh, J.R., Bair, W. & Movshon, J.A. Nature and interaction of signals from the receptive field center and surround in macaque V1 neurons. J. Neurophysiol. 88, 2530–2546 (2002).
46. Smith, M.A., Majaj, N.J. & Movshon, J.A. Dynamics of motion signaling by neurons in macaque area MT. Nat. Neurosci. 8, 220–228 (2005).
47. Cox, D.R. Some statistical methods connected with series of events. J. R. Stat. Soc. Ser. A 17, 129–164 (1955).
48. Greenwood, M. & Yule, G.U. An inquiry into the nature of frequency distributions of multiple happenings, with particular reference to the occurrence of multiple attacks of disease or repeated accidents. J. R. Stat. Soc. A 83, 255–279 (1920).
49. Polson, N.G., Scott, J.G. & Windle, J. Bayesian inference for logistic models using Polya-gamma latent variables. J. Am. Stat. Assoc. 108, 1339–1349 (2013).