Connecting psychophysical performance to neuronal response properties II: Contrast decoding and detection

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The purpose of this article is to provide mathematical insights into the results of some Monte Carlo simulations published by Tolhurst and colleagues (Clatworthy, Chirimuuta, Lauritzen, & Tolhurst, 2003; Chirimuuta & Tolhurst, 2005a). In these simulations, the contrast of a visual stimulus was encoded by a model spiking neuron or a set of such neurons. The mean spike count of each neuron was given by a sigmoidal function of contrast, the Naka-Rushton function. The actual number of spikes generated on each trial was determined by a doubly stochastic Poisson process. The spike counts were decoded using a Bayesian decoder to give an estimate of the stimulus contrast. Tolhurst and colleagues used the estimated contrast values to assess the model’s performance in a number of ways, and they uncovered several relationships between properties of the neurons and characteristics of performance.

Although this work made a substantial contribution to our understanding of the links between physiology and perceptual performance, the Monte Carlo simulations provided little insight into why the obtained patterns of results arose or how general they are. We overcame these problems by deriving equations that predict the model’s performance. We derived an approximation of the model’s decoding precision using Fisher information. We also analyzed the model’s contrast detection performance and discovered a previously unknown theoretical connection between the Naka-Rushton contrast-response function and the Weibull psychometric function. Our equations give many insights into the theoretical relationships between physiology and perceptual performance reported by Tolhurst and colleagues, explaining how they arise and how they generalize across the neuronal parameter space.

Introduction

This work was inspired by a series of articles from Tolhurst and colleagues that described Monte Carlo simulations of contrast coding in the visual cortex (Chirimuuta, Clatworthy, & Tolhurst, 2003; Clatworthy, Chirimuuta, Lauritzen, & Tolhurst, 2003; Chirimuuta & Tolhurst, 2005a, 2005b). In these simulations, the contrast of a visual stimulus was encoded by a model spiking neuron, or a set of such neurons, and the spike counts were decoded using a Bayesian decoder to give an estimate of the stimulus contrast. The contrast estimates were then used to assess the model’s decoding precision or to generate decisions in simulations of psychophysical tasks.

Although these studies made a very useful contribution to our understanding of the relationship between physiology and perceptual performance, the Monte Carlo simulations provided little insight into why the obtained patterns of results arose or how general they are. It was not clear whether the findings applied to just the specific sets of model parameters that were used in the simulations or to any parameter values. The goal of the present study was to derive equations that predict the performance of this kind of model from its parameters in order to better understand the relationships between physiology and perceptual performance and to allow the model’s performance to be calculated quickly and easily, thus removing the need for Monte Carlo simulations, which are difficult to program and time consuming to run.

Our article focuses on two key sets of results from the work of Tolhurst and colleagues. The first focus is the relationships between decoding precision and neuronal properties discovered by Clatworthy et al. (2003); we find that these relationships can be explained
by approximating the decoding precision using Fisher information. The second focus is the model’s psychometric function for detection, studied by Chirimuuta and Tolhurst (2005a); we derive an expression that approximates this function and sheds light onto one of Chirimuuta and Tolhurst’s (2005a) results. Some of the derivations and technical details are included in appendices in the supplementary information. Each supplementary appendix is labeled with a letter. Equations and figures in an appendix are labeled with the appendix’s letter, followed by a dot, followed by the equation or figure number within that appendix. Supplementary Appendix A provides a list of the main symbols used in the text and their meanings.

### The contrast coding model

The contrast coding model used by Tolhurst and colleagues consisted of an encoding stage, in which the stimulus was encoded as neuronal spike counts, and a decoding stage, which used Bayesian inference to estimate the stimulus contrast from the spike counts. The encoding stage was based on standard physiological models of the neuronal response, while the decoding stage was simply a Bayesian decoder, with no proposed physiological implementation.

The model has three elements: (a) the form of the tuning function, which specifies the mean spike count of each neuron for a given stimulus; (b) the random process that generates spikes at the given rate; and (c) the method of decoding the population response to give an estimate of the stimulus. We consider these three elements in turn.

We use the same basic terminology as May and Solomon (2015): We represent random variables using uppercase letters and their values on particular trials using corresponding lowercase letters. $X$ and $x$ represent the stimulus level, $R$ and $r(x)$ represent the mean spike count, $N$ and $n$ represent the actual spike count, and $N$ and $n$ are vectors holding the spike counts of all the neurons in the population. These variables are explained in more detail in our companion article (see the first paragraph of the section titled “The sensory coding model” in May & Solomon, 2015).

### The tuning function

The tuning function, $r(x)$, specifies the mean spike count for stimulus $x$. For visual stimulus contrast, the tuning function is called the contrast-response function. Tolhurst and colleagues modeled this function using the Naka-Rushton function (Naka & Rushton, 1966; Albrecht & Hamilton, 1982). For contrast, $c$, in linear (e.g., Michelson) units, the Naka-Rushton function has the following form:

\[
NR(c) = \frac{r_{\text{max}}^c}{c^{1/2} + c^q} + r_0. \tag{1}
\]

If we measure contrast in log units, $x = \log_b c$, then the Naka-Rushton is given by

\[
NR(x) = \frac{r_{\text{max}}^b x}{b^{1/2} + b^{q x}} + r_0, \tag{2}
\]

where

\[
z = \log_b c^{1/2}. \tag{3}
\]

See our companion article (May & Solomon, 2015) for a description of all the parameters and for plots of the Naka-Rushton function. On the log contrast scale, the gradient of the Naka-Rushton function peaks at a log contrast of $x = z$. The term $c_{1/2}$, called the semi-saturation contrast, is the contrast for which the mean response exceeds $r_0$ by $r_{\text{max}}/2$. Many authors use $c_{50}$ to represent this contrast, but we use the less common term $c_{1/2}$ (except when quoting other authors) because this form is easier to extend to other fractions of $r_{\text{max}}$, such as $c_{1/3}$ (the contrast for which the mean response exceeds $r_0$ by $r_{\text{max}}/3$, which we show to be the contrast that is coded most accurately by a model neuron with a Naka-Rushton contrast-response function with $r_0 = 0$).

In this article, whenever we use the term contrast without specifying the units, we mean log Michelson contrast. To be compatible with Clatworthy et al. (2003) and Chirimuuta et al. (2003), we always used log to base 10 in our modeling (i.e., $x = \log_{10} c$ and $z = \log_{10} c_{1/2}$); however, our equations are derived for the general case of any base, $b$.

Note that the units of the Naka-Rushton function’s output are often taken to be spikes per second, but we find it more convenient to use units of spikes without making assumptions about the time interval over which the neuron’s output is integrated.

### The random process for spike generation

The tuning function gives the mean spike count, and we now turn to the stochastic process that generates spikes at the specified rate. The Poisson process is sometimes used as a stochastic spiking model because of its considerable mathematical convenience (e.g., see Dayan & Abbott, 2001, chapter 1). The Poisson distribution is defined as

\[
P_{\text{Poisson}}(N = n | R = r(x)) = \frac{r(x)^n}{n!} \exp(-r(x)). \tag{4}
\]

In Equation 4, we use a right subscript on $P$ to indicate the type of stochastic process being used.
For the Poisson process, the variance in the number of spikes for repeated trials is equal to the mean, \( \sigma^2 = \mu \), always giving a Fano factor (ratio of variance to mean) of 1. In the visual cortex, the Fano factor is usually greater than 1 (Dean, 1981b; Tolhurst, Movshon, & Thompson, 1981; Tolhurst, Movshon, & Dean, 1983; Bradley, Skottun, Ohzawa, Sclar, & Freeman, 1987; Skottun, Bradley, Sclar, Ohzawa, & Freeman, 1987; Scobey & Gabor, 1989; Vogels, Spileers, & Orban, 1989; Snowden, Treue, & Maunsell, 1999; Durant, Clifford, Crowder, Price, & Tolhurst, 1994; Geisler & Albrecht, 1997; Bair & O'Keefe, 1998; McAdams & Maunsell, 1999; Oram, Weiner, Lestienne, & Rich mond, 1999; Durant, Clifford, Crowder, Price, & Ibbotson, 2007). To get a Fano factor greater than 1, Tolhurst and colleagues (Chirimuuta et al., 2003; Clatworthy et al., 2003; Chirimuuta & Tolhurst, 2005a, 2005b) used a doubly stochastic Poisson process, which we refer to as the Tolhurst process. This process is a Poisson process in which the mean is itself a random variable sampled from a Poisson process with mean \( \mu(x) \):

\[
P_{\text{Tolhurst}}(N = n|R = r(x)) = \sum_{\mu=0}^{\infty} P_{\text{Poisson}}(N = n|R = \mu) \times P_{\text{Poisson}}(N = \mu|R = r(x)).
\]

For this process, the mean spike count is \( r(x) \) and the variance is \( 2r(x) \), giving a Fano factor of 2. The infinite series in Equation 5 is difficult to handle, so in Supplementary Appendix B we derive a finite series expansion of the Tolhurst process that is more useful.

Since the aim of this article is to explain the modeling results of Tolhurst and colleagues, all the Monte Carlo simulations in this article use the Tolhurst process to generate spikes. However, to make our theoretical results more general, we consider two other Poisson-based spiking processes.

The first additional spiking process is the generalized Poisson distribution, devised by Consul and Jain (1973). Sakata and Harris (2009) used this process to model neuronal spiking distributions. This process is more flexible than the Tolhurst process because the Fano factor, \( F \), is set as a parameter and can take any value greater than or equal to 1. The Consul-Jain distribution takes the following form:

\[
P_{C.J}(N = n|R = r(x)) = \frac{r(x)^n}{n!\sqrt{F}} A^{n-1} \exp(-A),
\]

where

\[
A = \frac{r(x) + n(\sqrt{F} - 1)}{\sqrt{F}}.
\]

Like the Poisson and Tolhurst processes, the Consul-Jain process generates only nonnegative numbers of spikes, and the spike count variance is proportional to the mean. When \( F = 1 \), the Consul-Jain process reduces to the ordinary Poisson process, given in Equation 4. However, the Consul-Jain process with \( F = 2 \) is not identical to the Tolhurst process, even though the Fano factor is the same.

The second additional spiking process that we consider is the doubly stochastic Poisson process recently proposed by Goris, Movshon, and Simoncelli (2014). We refer to this as the Goris process. It is a Poisson process in which the mean is modulated by a multiplicative gain mechanism. The gain value on each stimulus presentation is a gamma-distributed random variable. The Goris process is described in detail in our companion article (May & Solomon, 2015). The gain fluctuations result in a Fano factor that increases with the firing rate, which is more physiologically plausible than the fixed Fano factor produced by the Tolhurst and Consul-Jain processes. For mathematical simplicity, we assume that each neuron in the population has the same gain signal and that each neuron’s Poisson spiking process is independent (see May & Solomon, 2015, for justification of these restrictions). The shared gain signal causes the neuronal responses to be correlated, with a realistic correlation structure. However, because each neuron’s Poisson spiking process is independent, a decoder that knows the gain signal on each stimulus presentation can express the spiking distributions as independent Poisson distributions. Therefore, the neurons can be decoded as if they were statistically independent (see May & Solomon, 2015, for a more detailed explanation of this).

**Bayesian population decoding**

All the results that we report from Tolhurst and colleagues were obtained using maximum-likelihood decoding. The estimated stimulus level is the one that had the highest probability of giving rise to the obtained set of spike counts; that is, it is the value of \( x \) that maximizes the likelihood, \( P(N = n|X = x) \). In Tolhurst and colleagues’ model, the neurons were statistically independent; in addition, if we use May and Solomon’s (2015) parameterization of the Goris process, the neurons are implicitly decorrelated if the decoder knows the gain signal. For statistically independent neurons, the population likelihood is then given by the product of the likelihoods of the individual neurons:

\[
P(N = n|X = x) = \prod_{j=1}^{K} P(N_j = n_j|X = x) = \prod_{j=1}^{K} P(N_j = n_j|R_j = r_j(x)).
\]

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$K$ is the number of neurons. $N_j$ is a random variable representing the spike count of neuron $j$, and $n_j$ is its value. $R_j$ is a random variable representing the mean spike count for neuron $j$, and $r_j(x)$ is its value. The second equality (Equation 9) arises because each $r_j(x)$ is a deterministic function of the stimulus value, $x$. To evaluate the probabilities in Equation 9, we use the appropriate expression, depending on which spiking process we are using. For the Consul-Jain process, we use Equation 6. For the Goris process, assuming the decoder knows the gain value, we use the gain-modulated Poisson distribution (May & Solomon, 2015; Equation 10). All the modeling in this article used the Tolhurst process, which is defined in Equation 5.

**Measuring decoding precision**

**Definition of decoding precision**

Decoding precision is usually taken to be the reciprocal of the variance of the estimated stimulus value; this is how it is defined in our companion article (May & Solomon, 2015). However, in their Monte Carlo simulations, Tolhurst and colleagues (Chirimuuta et al., 2003; Clatworthy et al., 2003) used a slightly different measure of decoding performance. They called it accuracy, defined as the reciprocal of the mean squared difference between the individual estimate and the true stimulus value:

\[
\text{accuracy} = \frac{T}{\sum (\hat{x} - x)^2},
\]

where $T$ is the number of trials ($= 10,000$), $\hat{x}$ is the estimated log contrast estimate (note that we use an uppercase $\hat{X}$ to represent the random variable and a lowercase $\hat{x}$ to represent its value on a particular trial), and the denominator is the sum over all trials. For the models that we consider in this study, the log contrast estimate is largely unbiased (except at very low performance levels), so we have mean[$\hat{X}$] = $x$. In this case, the accuracy score in Equation 10 is essentially the same as the precision. For consistency with Tolhurst and colleagues, we used their measure of decoding accuracy (Equation 10) when analyzing our Monte Carlo simulations. However, we refer to it as precision (except when directly quoting Tolhurst and colleagues) because our analytical approximations of it are formally measures of precision and because the term `accuracy` is often used to mean the inverse of bias (e.g., Smith, 1999, chapter 2). We found that, except in degenerate conditions where the model’s performance level was very low, it made a negligible difference whether we plotted Tolhurst's accuracy score or true precision. Note that in the Monte Carlo simulations in our companion article (May & Solomon, 2015), we calculated true decoding precision (i.e., reciprocal of the variance of the estimated stimulus value), not the accuracy score defined in Equation 10.

**Approximating decoding precision using Fisher information**

For an unbiased maximum-likelihood decoder, the precision cannot exceed a quantity called the Fisher information. This limit is known as the Cramér-Rao bound (Rao, 1945; Cramér, 1946; see Dayan & Abbott, 2001, pp. 120–121). For populations that give sufficiently large numbers of informative spikes, the precision of a maximum-likelihood decoder is very close to the Fisher information; when the tuning function is a sigmoid function like the Naka-Rushton function, this applies even if the population consists of a single neuron. This means that we can use the Fisher information as an analytical approximation of the decoding precision, thereby giving us a real insight into why Tolhurst and colleagues' results occurred and how general they are.

The Fisher information depends on the tuning function and the spiking process. Unfortunately, for the Tolhurst and Consul-Jain spiking processes, the derivation of an exact formula for the Fisher information turned out to be intractable. In Supplementary Appendix E, we derive approximations of the Fisher information for each of these spiking processes: $\tau_T(x)$ and $\tau_{C-J}(x)$. As long as the mean spike count of the most informative neurons is not too low, both $\tau_T(x)$ and $\tau_{C-J}(x)$ are very close to the following general approximation of the decoding precision, $\tilde{\tau}(x)$:

\[
\tilde{\tau}(x) = \frac{1}{v} \sum_{j=1}^{K} r_j(x)^2 / r_j(x),
\]

where $r_j(x)$ is the first derivative of neuron $j$'s tuning function with respect to $x$. To parameterize $\tilde{\tau}(x)$ so that it approximates $\tau_T(x)$ or $\tau_{C-J}(x)$, $v$ should be equal to the Fano factor; the Fano factor is fixed at 2 for the Tolhurst process and can take any value greater than or equal to 1 for the Consul-Jain process. The tilde (') indicates that this estimate of decoding precision is based on an approximation of the Fisher information that is not always accurate. With a Fano factor of 1, the Consul-Jain process is the ordinary Poisson, in which case $\tilde{\tau}(x)$ with $v = 1$ is exactly equal to the Fisher information (see Dayan & Abbott, 2001, chapter 3). For the Tolhurst process, and the Consul-Jain process with $F \neq 1$, $\tilde{\tau}(x)$ with $v$ equal to the Fano factor is an approximation of the Fisher information. For the Goris process, May and Solomon (2015)
showed that an appropriate estimate of the decoding precision is given exactly by Equation 11 with $v = 1/(1 - \sigma_{\tilde{v}}^2)$, where $\sigma_{\tilde{v}}^2$ is the variance of the gain signal (see equation 26 and relation 31 of May & Solomon, 2015). In this case, $v$ is not the Fano factor—the Fano factor for Goris et al.’s (2014) spiking process is variable and depends on the mean spike count. When the gain is known by the decoder, the Fisher information of the Goris process varies from trial to trial due to the fluctuating gain, and $\tilde{\tau}(x)$ with $v = 1/(1 - \sigma_{\tilde{v}}^2)$ gives the modal value of the Fisher information exactly.

In summary, Equation 11 gives a general approximation of the decoding precision for a variety of Poisson-based spiking distributions. For two processes (Poisson and Goris) it is based on an exact expression for the Fisher information; for the other two processes (Tollhurst and Consul-Jain) it is based on an approximation of the Fisher information that is accurate as long as the mean spike count of the most informative neurons is not too low.

If we expand $r_j(x)$ and $r'_j(x)$ in Equation 11 using the Naka-Rushton function (Equation 2), then we have

$$\tilde{\tau}(x) = \frac{1}{v} \sum_{j=1}^{K} \frac{r_{\text{max}} (qlb)^2 h^{q(2x+b)}}{(b^{2x} + b^{ax})^3} \quad \text{if} \quad r_0 = 0.$$  \hspace{1cm} (12)

Tolhurst and colleagues always used $r_0 = 0$ in their modeling. Since we are focusing on their modeling results, we consider only the case of $r_0 = 0$ in this article. In Equation 12, each neuronal parameter can vary from neuron to neuron, so, strictly speaking, each parameter should be indexed by the neuron number, $j$, but we omit these indices to reduce notational clutter.

In the next section, many of the populations of neurons that we analyze consist of either just a single neuron or a set of identical neurons. In this case, Equation 12 reduces to

$$\tilde{\tau}(x) = \frac{K r_{\text{max}} (qlb)^2 h^{q(2x+b)}}{v (b^{2x} + b^{ax})^3} \quad \text{if} \quad r_0 = 0,$$ \hspace{1cm} (13)

with $K = 1$ in the case of a single neuron.

1. “Increasing $R_{\text{max}}$ increases the contrast identification accuracy of single neurons at all contrasts, most obviously the peak accuracy, without changing the contrast at which accuracy is a maximum” (Clatworthy et al., 2003, p. 1991; note that they use $R_{\text{max}}$ where we use $r_{\text{max}}$). This can be summarized by saying that there is an approximately multiplicative effect of $r_{\text{max}}$ on decoding precision.
2. “The position of the accuracy peak along the contrast axis is consistently close to but, interestingly, slightly below the neuron’s $c_{s0}$” (Clatworthy et al., 2003, p. 1989).
3. “The peak value of accuracy is independent of $c_{s0}$” (Clatworthy et al., 2003, p. 1989).
4. “The relationship between the maximum accuracy and $q$ is a steep straight line on log-log coordinates” (Clatworthy et al., 2003, p. 1989).
5. “To change the maximum accuracy . . . requires only a change in the product of $R_{\text{max}}$ and number of neurons, i.e., the total number of action potentials generated on average . . . ; for a given accuracy, there is a simple trade-off between the number of neurons and the response amplitude of individual neurons” (Clatworthy et al., 2003, p. 1990).

The numerical nature of Clatworthy et al.’s method means that it gave little insight into why these relationships arose or how generally they apply. In this section we explain Clatworthy et al.’s findings by deriving equations that explain corresponding findings for the analytical approximation of decoding precision, $\tilde{\tau}(x)$, given in Equation 13.

First, we examined whether $\tilde{\tau}(x)$ was sufficiently close to the true decoding precision to make this approach valid. This is important because the Fisher information, on which $\tilde{\tau}(x)$ is based, is only an upper bound on the decoding precision and can far exceed the true decoding precision for small population sizes (Xie, 2002). In one of their investigations (shown in their figure 5A), Clatworthy et al. (2003) examined the effect of $r_{\text{max}}$ on decoding precision for a single Tolhurst-spiking neuron with Naka-Rushton exponent ($q$) equal to 2, $c_{1/2} = 0.1$, and $r_0 = 0$, $r_{\text{max}}$ took values of 5, 20, 50, 100, or 180 spikes. We replicated their methods (see Supplementary Appendix G for details) and obtained contrast decoding precision scores (calculated using Equation 10) that were essentially identical to those that we read off from their figure 5A. The small differences were almost certainly attributable to the stochastic nature of the simulations and small inaccuracies in our transcription of the data from Clatworthy et al.’s figure. The symbols in Figure 1 show these precision scores. The thin, colored curves show the true Fisher information, calculated numerically (for method, see Supplementary Appendix H). The thick, black curves show $\tilde{\tau}(x)$ with $v = 2$. The two panels in Figure 1 are identical except that Figure 1A has a linear vertical

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**Explaining Clatworthy et al.’s contrast decoding results**

Clatworthy et al. (2003) applied their Bayesian contrast decoding algorithm to single Tolhurst-spiking neurons, or populations of them, and discovered several relationships between the neuronal parameters and the peak of decoding precision along the contrast axis.
Observation 1: Multiplicative effect of increasing $r_{\text{max}}$

From Equation 13, we see that $\tilde{r}(x)$ is proportional to $r_{\text{max}}$, so increasing $r_{\text{max}}$ increases the precision by the same multiplicative factor for each contrast level; the position of the peak across contrast is unchanged, and the largest absolute change is for the peak precision. This multiplicative change gives rise to the vertical shifts seen for the black curves on the logarithmic vertical axis in Figure 1B.

Observation 2: Precision peaks slightly below the semisaturation contrast

The exact position of the peak of $\tilde{r}(x)$ along the contrast axis can be found by differentiating $\tilde{r}(x)$ with respect to $x$, setting the derivative to zero, and solving for $x$. From Equation 13, we have

$$\frac{d \tilde{r}(x)}{dx} = \frac{Kr_{\text{max}}(q \log b)^{3} b^{q(2x+1)} b^{q} (b^{q} - 2 b^{2q})}{v(b^{q} + b^{2q})^4}.$$  (14)

Setting this to zero gives

$$\frac{d \tilde{r}(x)}{dx} = 0 \Rightarrow b^{q} = 2 b^{2q}$$

$$\Rightarrow x = z - \log_{b}(2^{1/q})$$  (15)

$$\Rightarrow c = 2^{-1/q}c_{1/2}.$$  (16)
Using Equation 15 to substitute for \(x\) in Equation 2 gives (for \(r_0 = 0\))

\[
N_R \bar{r}(x) = \frac{r_{max}}{3} \quad \text{if} \quad \tilde{r}(x) \text{ is at its peak.} \tag{17}
\]

So, regardless of the values of \(v\), \(r_{max}\), \(c\), or \(c_{1/2}\), \(\tilde{r}(x)\) for a single neuron will always peak at the contrast for which the mean response is \(r_{max}/3\). We introduce the term \(c_{1/3}\) for the Michelson contrast that gives rise to a mean response of \(r_{max}/3\), to be consistent with the term \(c_{1/2}\) for the semisaturation contrast. The value of \(\log_{10}(c_{1/3})\) is indicated by the black vertical line in each panel of Figure 1, and it passes through the peak of each thick, black curve.

**Observation 3: The peak value of precision is independent of the semisaturation contrast**

The peak value of \(\tilde{r}(x)\) occurs at a log contrast given by Equation 15. Using this equation to substitute for \(x\) in Equation 13, we find that the peak value of \(\tilde{r}(x)\) is given by

\[
\max[\tilde{r}(x)] = \frac{4Kr_{max} (q \ln b)^2}{27v}, \tag{18}
\]

which is independent of \(c_{1/2}\).

**Observation 4: The relationship between the maximum precision and \(q\) is a steep straight line on log-log coordinates**

From Equation 18, we see that \(\max[\tilde{r}(x)]\) is proportional to \(q^2\), giving a straight line on log-log coordinates. Figure 2 plots \(\max[\tilde{r}(x)]\) as a function of \(q\) for a single Tolhurst-spiking neuron with \(r_{max} = 50\) spikes and shows that \(\max[\tilde{r}(x)]\) is very close to the true peak of precision for this model neuron obtained using Clatworthy et al.’s Monte Carlo methods.

**Observation 5: Linear tradeoff between \(r_{max}\) and number of neurons**

Clatworthy et al. (2003) and Chirimuuta and Tolhurst (2005a) both noted that \(r_{max}\) and the number of neurons, \(K\), trade off so that, regardless of the individual values of \(r_{max}\) and \(K\), the decoding precision is a function of their product. In fact, unless \(r_{max} \times K\) is very low, the decoding accuracy is close to being proportional to \(r_{max} \times K\). This observation is easily explained by Equation 13, which shows that \(\tilde{r}(x)\) is proportional to \(r_{max} \times K\).

The crosses in Figure 3 replot data from Clatworthy et al.’s (2003) figure 5C. These data show the
maximum decoding precision of various populations of neurons, all with $q = 2$ and $c_{1/2} = 0.1$. The near proportionality between decoding accuracy and $r_{\text{max}} \times K$ is indicated by the fact that most of the points lie on a straight line of gradient 1 on these log-log axes. The straight line in our Figure 3 is a plot of max $\tilde{s}(x)$, calculated according to Equation 18. It clearly provides a very good match to the decoding precision data for $r_{\text{max}} \times K / C^2 \leq 50$.

Note that although $\tilde{s}(x)$ gives an exact linear trade-off between $r_{\text{max}}$ and $K$, the true Fisher information for the Tolhurst process does not. As $r_{\text{max}}$ decreases, $\tilde{s}(x)$ tends to underestimate the true Fisher information.

A more accurate estimate of the decoding precision is given by $\tau_{\text{Tolhurst}}(x)$, derived in Supplementary Appendix E:

$$\tau_{\text{Tolhurst}}(x) = \sum_{j=1}^{K} H_{\text{Tolhurst}}(x) \times \frac{r_j'(x)^2}{r_j'(x)}$$  \hspace{1cm} (19)

where

$$H_{\text{Tolhurst}}(x) = \left( \frac{1}{2} - \frac{1 + 0.06630 \times r(x)}{e} \right)$$

$$\times \exp[r(x)(1/e - 1)] + \frac{1}{2}$$  \hspace{1cm} (20)

$H_{\text{Tolhurst}}(x)$ is plotted in Figure 4. As the spike rate of each individual neuron increases, $H_{\text{Tolhurst}}(x)$ for that neuron approaches 1/2, and so $\tau_{\text{Tolhurst}}(x)$ approaches $\tilde{s}(x)$ (Equation 11) with $v = 2$. However, as the spike rate decreases, $H_{\text{Tolhurst}}(x)$ increases, and so $\tau_{\text{Tolhurst}}(x)$ exceeds $\tilde{s}(x)$ and better reflects the true decoding precision.

For the Naka-Rushton function with $r_0 = 0$, we can expand Equation 19 in a similar way to Equation 12:

$$\tau_{\text{Tolhurst}}(x) = \sum_{j=1}^{K} H_{\text{Tolhurst}}(x) \times \frac{r_{\text{max}} (q \ln b)^2 b^j (2z + x)}{b^{2z} + b^j (x)^3}.$$  \hspace{1cm} (21)

Unlike $\tilde{s}(x)$, $\tau_{\text{Tolhurst}}(x)$ is specific to the Tolhurst spiking process rather than being a general approximation of the decoding precision that applies to several different spiking processes.

In each panel of Figure 5, the solid line is the same as in Figure 3, while the dashed line plots the maximum...
value of $\hat{\tau}_{\text{Tolhurst}}(x)$. As $r_{\text{max}}$ decreases, $\hat{\tau}_{\text{Tolhurst}}(x)$ and $\hat{\tau}(x)$ start to diverge by a factor that approaches $(1 - 1/ e)/0.5 = 1.264$—that is, the ratio of the maximum to minimum values of $H_{\text{Tolhurst}}(x)$.

This inexact trade-off between $r_{\text{max}}$ and $K$ applies to the Tolhurst process but not to the Poisson or Goris processes: for these, $\hat{\tau}(x)$ is derived from the exact expression for the Fisher information, so the trade-off that we derived for $\hat{\tau}(x)$ applies to these processes exactly.

### Failure of Clatworthy et al.'s observations for single neurons at low spike rates

When $r_{\text{max}}$ (or, for populations, $r_{\text{max}} \times K$) is substantially less than 50 spikes, the peak of precision does not coincide closely with that of $\hat{\tau}(x)$ or even the true Fisher information, so the explanations of Clatworthy et al.’s findings given above are no longer valid. However, most of these findings do not apply for these low $r_{\text{max}}$ values either. The failure of observations 1 and 2 at low spike rates is clear from Figure 1, the failure of observation 5 at low spike rates is shown in Figure 3, and the failure of observation 4 at low spike rates is demonstrated in Figure 6.

The median $r_{\text{max}}$ for a 200-ms stimulus is only around 5.7 spikes for V1 neurons (Geisler & Albrecth, 1997), suggesting that, with single-neuron models, the spike count has to be implausibly high for the decoding precision to be well approximated by the Fisher information. However, as shown by Clatworthy et al. (2003), Chirimuuta and Tolhurst (2005a), and our Figure 3, $r_{\text{max}}$ can be approximately traded off against the number of neurons so that what is important is the average total spike count of the population rather than that of the individual neurons. With a population code, it is possible to achieve a high total population spike count while keeping the spike counts for the individual neurons at a plausible level. This makes the Fisher information more relevant to understanding population-coding models than coding schemes based on a single neuron. So far we have considered only populations of identical neurons. We now turn to populations of differently tuned neurons.

### Population coding of contrast

Clatworthy et al. (2003) applied maximum-likelihood decoding to three different populations of 18 model Tolhurst-spiking neurons, each with $r_{\text{max}} = 10$, $r_0 = 0$, and $q = 2$. One population had $\log_{10}(c_{1/2})$ values uniformly distributed between $-3$ and 0.1, and the other two had $c_{1/2}$ values distributed according to the recorded values in either cat or monkey populations, found by arranging the neurons in ascending order of $c_{1/2}$ and then sampling the population at equal percentile intervals. We did not have access to the exact sets of cat or monkey $c_{1/2}$ values that they used, but we estimated them by fitting functions to the cat or monkey $c_{1/2}$ distributions given in Clatworthy et al.’s figure 6 and then sampling these distributions in equal percentile steps (see Supplementary Appendix I for details). The advantage of our method is that it can easily be extended to neuronal populations of any size. Having set up the populations of neurons, we then calculated decoding precision scores using Clatworthy et al.’s Monte Carlo methods (see Supplementary Appendix G for details). Figure 7 shows that our decoding precision scores are very similar to those of Clatworthy et al., confirming that our method of generating the sets of $c_{1/2}$ values is a good approximation to that of Clatworthy et al. The colored curves in Figure 7 show $\hat{\tau}(x)$, calculated using Equation 12 with $v = 2$. Since $r_{\text{max}} = 10$ in these simulations, $H_{\text{Tolhurst}}(x)$ is 0.5 for the most informative neurons, so $\hat{\tau}_{\text{Tolhurst}}(x)$ was not advantageous in using the closer but more complex approximation. Figure 7 shows that, even for these small populations of neurons, with $r_{\text{max}}$
of only 10 spikes, the decoding precision from the Monte Carlo simulations is very close to $\bar{r}(x)$ over a wide range of contrasts.

In Figure 7, the only substantial deviation of decoding precision from Fisher information occurs at the ends of the contrast range, where the precision scores shoot up. This is an artifact, discussed by Clatworthy et al., caused by the fact that in the simulations the likelihood functions were calculated over a finite range of contrasts. This means that on trials where the inferred contrast would have fallen beyond the ends of the contrast range, the inferred contrasts instead pile up on the ends of the range, artificially boosting the number of trials on which the inferred contrast takes those values. Thus, when the stimulus contrast really is at or close to one of the ends of the range, it is likely to be close to the inferred contrast, so the precision is artificially high. The Fisher information does not show this effect because it is a purely local measure, derived from the contrast-response functions at each point along the contrast axis, so it is unaffected by any bounds on the range of contrasts.

$\bar{r}(x)$ is a good approximation of the decoding precision for the Tolhurst process as long as $r_{\text{max}}$ is not too low. For low values of $r_{\text{max}}$, $\bar{r}(x)$ tends to underestimate the true Fisher information. As noted earlier, this means that there is not an exact trade-off between $r_{\text{max}}$ and the number of neurons, $K$. For low values of $r_{\text{max}}$, it is better to use $\tau_{\text{Tolhurst}}(x)$ to accurately predict decoding precision. This is demonstrated in Figure 8.

There are at least two reasons why we sometimes need an expression for the decoding precision that remains a close approximation down to very low spike rates. First, the median spike count for a 200-ms stimulus presentation is only around 5.7 spikes for a V1 neuron tuned to the stimulus (Geisler & Albrecht, 1997), and observers can make complex judgements based on exposures even shorter than that (Thorpe, Fize, & Marlot, 1996). Second, in many situations the majority of neurons will not be well tuned to the stimulus but may still contribute to task performance due to their abundance; these neurons would be expected to have a very low average spike count over the course of a stimulus presentation.

The psychometric function for contrast detection

Chirimuuta and Tolhurst (2005a) used their contrast coding model to simulate a two-alternative forced-choice (2AFC) contrast detection task, in which the observer is presented with a zero-contrast nontarget stimulus and an above-zero-contrast target stimulus and has to pick the target. Chirimuuta and Tolhurst (2005a) measured the model’s proportion of correct responses as a function of the target contrast and fitted a Weibull psychometric function to the data.

For 2AFC tasks, the Weibull function can be defined as

$$P(\text{correct}) = 1 - 0.5\exp\left[-(c/z)^\beta\right].$$  \hspace{1cm} (22)

$z$ is the threshold—that is, the target contrast that gives $P(\text{correct}) = 1 - 0.5/e = 0.816$... and $\beta$ controls the function’s shape on linear axes or slope on log axes (see May & Solomon, 2013, for an in-depth analysis of the Weibull function). In psychophysical contrast detection experiments with human observers, $\beta$ usually takes a value of about 3 (Foley & Legge, 1981; Nachmias, 1981; Mayer & Tyler, 1986; Meese, Georgeson, & Baker, 2006; Wallis, Baker, Meese, & Georgeson, 2013).
When Chirimuuta and Tolhurst’s (2005a) model had a standard Naka-Rushton contrast-response function with \( r_0 = 0 \) and \( q = 2 \), Weibull \( \beta \) for detection varied from 1.75 to 1.99 as the number of neurons increased from 1 to 23, but the model’s \( \beta \) never reached the normal human level of around 3. Chirimuuta and Tolhurst then introduced a threshold to the Naka-Rushton function by subtracting a constant value from the output and setting negative values to zero. With a threshold on the Naka-Rushton function, \( \beta \) ranged from 2.25 to 4.20, providing a better match to psychophysically obtained values. Chirimuuta and Tolhurst assumed that their failure to obtain sufficiently high Weibull \( \beta \) values with the standard Naka-Rushton function had been caused by the lack of a threshold, and they suggested that the standard, unthresholded Naka-Rushton function “may be crucially inadequate” as a model of the neuronal contrast-response function (Chirimuuta & Tolhurst, 2005a, p. 2956). However, we now show that, when \( r_0 = 0 \), the model’s psychometric function is close to a Weibull function with \( \beta = q \). Thus, the real reason that Chirimuuta and Tolhurst always obtained a \( \beta \) of about 2 with the standard Naka-Rushton function is that they always kept \( q \) at 2 in these simulations. We show that one can obtain any Weibull \( \beta \) by setting \( q \) close to the required \( \beta \) value.

Proof that, when \( r_0 = 0 \), the model’s psychometric function for 2AFC contrast detection is close to a Weibull function with \( \beta = q \)

At the low contrasts corresponding to the model’s contrast detection threshold, the population spike rate is typically so low that the Fisher information does not provide a useful approximation of perfor-
performance. Therefore, we need to use other methods. For contrast detection, the model’s performance can be derived straightforwardly from basic probability theory.

Because \( r_0 = 0 \), there is zero response to zero contrast: The nontarget stimulus can never elicit a single spike. If the target elicits at least one spike, the model will respond correctly. If the target fails to elicit any spikes, the model has to guess and will be correct half the time on a 2AFC task. In summary, the model will be correct on all 2AFC trials except half of those on which the target failed to elicit any spikes. This statement can be formalized as follows:

\[
P(\text{correct}) = 1 - 0.5P(\text{no spikes}),
\]

where \( P(\text{no spikes}) \) is the probability of getting no spikes in response to the target. We can already see that the model’s psychometric function has a similar form to the Weibull function: \( P(\text{correct}) = 1 - 0.5 \times \) something.

From Equation E.4 of Supplementary Appendix E, we see that, for a single Tolhurst-spiking neuron,

\[
P(\text{no spikes}) = P_{\text{Tolhurst}}(N = 0|R = r(c)) = \exp[(1/e - 1)r(c)].
\]

Note that since the Weibull function is defined as a function of contrast in Michelson (linear) units, we express the mean response, \( r(c) \), as a function of Michelson contrast (rather than log contrast) in Equation 24. For a population of \( K \) statistically independent neurons,

\[
P(\text{no spikes}) = \prod_{j=1}^{K} \exp[(1/e - 1)r_j(c)] = \exp\left(1/e - 1\right)\sum_{j=1}^{K} r_j(c),
\]

where \( r_j(c) \) is the contrast-response function of neuron \( j \). Using Equation 1 to expand \( r_j(c) \), we have

\[
P(\text{no spikes}) = \exp\left(1/e - 1\right)\sum_{j=1}^{K} \frac{r_{\text{max}}}{c_{1/2}}\left(e^{q}\right) + e^{q},
\]

where \( r_{\text{max}} \) and \( c_{1/2} \) are the \( r_{\text{max}} \) and \( c_{1/2} \) parameters, respectively, of neuron \( j \). Using Equation 26 to substitute for \( P(\text{no spikes}) \) in Equation 23, we get an exactly correct expression for the model’s psychometric function for contrast detection:

\[
P(\text{correct}) = 1 - 0.5\exp\left(1/e - 1\right)\sum_{j=1}^{K} \frac{r_{\text{max}}}{c_{1/2}}\left(e^{q}\right) + e^{q},
\]

If all the neurons in the population being monitored by the observer have the same contrast-response function, then Equation 27 reduces to

\[
P(\text{correct}) = 1 - 0.5\exp\left(1/e - 1\right)r_{\text{max}}K\left(e^{q}\right)/c_{1/2} + e^{q}.\]

In this case, Equation 28 shows that there is an exact linear trade-off between \( r_{\text{max}} \) and \( K \) in the psychometric function for 2AFC detection rather than the approximate trade-off that we get with the Fisher information.

Equations 27 and 28 are precisely correct, but it is illuminating to derive an approximation. As long as there are enough neurons or \( r_{\text{max}} \) is sufficiently high, the contrast detection threshold will be somewhat lower than the lowest \( c_{1/2} \) in the population. Thus, at threshold (i.e., around the middle of the psychometric function), the denominators of Equations 26 to 28 become dominated by the \( c_{1/2}^{q} \) term, and the \( e^{q} \) term will make little difference. Dropping \( e^{q} \) from the denominator of Equation 27, we get

\[
P(\text{correct}) \approx 1 - 0.5\exp[-(c/\alpha)^q],
\]

where \( \alpha \) is a constant, given by

\[
\alpha = \left(1 - 1/e\right)\sum_{j=1}^{K} \frac{\left(r_{\text{max}}\right)}{c_{1/2}}^{q} - 1/q.
\]

Relation 29 has the form of a Weibull function with \( \beta = q \). The near equality in Relation 29 approaches equality as \( K \) or \( r_{\text{max}} \) increase. \( \square \)

The derivation of Relation 29 is similar to the derivation of the psychometric function for Quick’s (1974) vector-magnitude model of contrast detection. The performance of the vector-magnitude model is exactly equivalent to that of a model in which the observer monitors a number of detectors that each have a detection probability that can be described by a Weibull function, with the same \( \beta \) for all detectors. Nachmias (1981) called this assumption of identical \( \beta \) for each detector the “homogeneity assumption,” and it is largely equivalent to the implicit assumption in the above proof that all neurons have the same \( q \). In the Discussion we expand on the links between our analysis and that of Quick (1974).

**Another attribute of the model’s psychometric function: Lapse rate**

Note that Relation 29 is a Weibull function with zero lapse rate; that is, it predicts that the model’s performance will asymptote to perfect performance, \( P(\text{correct}) = 1 \), with increasing target contrast. For
most instantiations of the model, this is very close to the truth. However, when \( r_{\text{max}} \) and \( K \) are both very low, the model’s asymptotic performance is far below 1. This is because as \( c \) increases, the denominator of Equation 26 becomes more and more dominated by the \( c^q \) term, and the \( c_{1/2}^q \) term makes less and less difference, so

\[
\lim_{c \to \infty} P(\text{no spikes}) = \exp \left( \frac{1}{e} - 1 \right) \sum_{j=1}^{K} (r_{\text{max}})^j.
\]  

(31)

For very low \( r_{\text{max}} \) and \( K \), this value can be substantially above zero so that, even for infinite contrast, the model has to guess on a significant proportion of 2AFC trials.

This behavior can be accommodated by including a “lapse rate” parameter, \( \lambda \), in the definition of the Weibull function:

\[
P(\text{correct}) = (1 - \lambda) - (0.5 - \lambda) \exp \left(-\left(\frac{c}{\alpha}\right)^q\right).
\]  

(32)

This equation approaches an asymptote of \( P(\text{correct}) = (1 - \lambda) \) as \( c \to \infty \). When \( \lambda = 0 \), Equation 32 reduces to Equation 22. We now derive an expression for the model’s “lapse rate” parameter. (Strictly speaking, the model never lapses; a low asymptotic performance level arises from a low spike rate rather than a finger error or a failure to look at the stimuli on some 2AFC trials.)

Using Equation 31 to substitute for \( P(\text{no spikes}) \) in Equation 23, we obtain the asymptotic value of \( P(\text{correct}) \) given by

\[
\lim_{c \to \infty} P(\text{correct}) = 1 - 0.5 \exp \left( \frac{1}{e} - 1 \right) \sum_{j=1}^{K} (r_{\text{max}})^j.
\]  

(33)

Since this asymptotic value of \( P(\text{correct}) \) is \( (1 - \lambda) \), we have

\[
\lambda = 0.5 \exp \left( \frac{1}{e} - 1 \right) \sum_{j=1}^{K} (r_{\text{max}})^j.
\]  

(34)

This expression for \( \lambda \) quickly approaches zero as \( r_{\text{max}} \) or \( K \) increase.

**Verifying our equations using Monte Carlo simulations**

We simulated the 2AFC contrast detection experiments for a range of model parameterizations (see Supplementary Appendix J for details of the method). For each parameterization, each neuron in the population had an identical contrast-response function. We fitted the three-parameter Weibull function (Equation 32) to the data for each model parameterization. The fitted values of \( \alpha \), \( \beta \), and \( \lambda \) are plotted as symbols in Figures 9, 10, and 11, respectively. The solid lines plot the corresponding analytical expressions (\( \alpha \) given by Equation 30, \( \beta \) given by \( q \), and \( \lambda \) given by Equation 34).

Equation 34 is the model’s true lapse rate parameter (not an approximation), so it is not surprising that it matches the fitted \( \lambda \) values extremely well in Figure 11. Our analytical expressions for \( \alpha \) and \( \beta \) are approximations that become increasingly accurate as \( K \) or \( r_{\text{max}} \) increase.

**The Consul-Jain spiking process also generates a Weibull psychometric function for detection**

The analytical expressions for the psychometric function for detection, derived above, apply only to the Tolhurst spiking process. We can also derive analogous expressions for the Consul-Jain process (which includes the ordinary Poisson, when \( F = 1 \)). Equation E.17 of Supplementary Appendix E states that, for a single Consul-Jain-spiking neuron,

\[
P(\text{no spikes}) = \exp \left( - \frac{r(c)}{\sqrt{F}} \right),
\]  

(35)

where \( F \) is the Fano factor. If we follow a series of mathematical steps analogous to Equations 23 to 29 above but use Equation 35 instead of Equation 24 to express the probability of a single neuron not spiking, we obtain an approximation of the psychometric function with the same form as Relation 29 but with

\[
\alpha = \left( \sum_{j=1}^{K} \frac{r_{\text{max}}}{(c_{1/2})^j \sqrt{F}_j} \right)^{-1/q},
\]  

(36)

where \( F_j \) is the Fano factor or neuron \( j \). Similarly, if we follow a series of steps analogous to those in Equations 31 to 34 but use Equation 35 instead of Equation 24 to express the probability of a single neuron not spiking, we obtain the following expression for the “lapse rate” parameter:

\[
\lambda = 0.5 \exp \left( - \sum_{j=1}^{K} \frac{r_{\text{max}}}{\sqrt{F}_j} \right).
\]  

(37)

**Discussion**

The purpose of this study was to explain a number of empirical modeling results reported by Tolhurst and colleagues. Their results were obtained from Monte Carlo simulations using models of spiking
neurons with Naka-Rushton contrast-response functions and the doubly stochastic Poisson spiking process defined in Equation 5. The numerical nature of the simulations meant that it was not clear why the results occurred or how they generalized across parameter space.

We addressed these problems by deriving equations to explain the model’s performance. This kind of analysis is essential if we are to understand the brain: If we use realistic models to simulate brain processes but do not understand why the models behave in the way that they do, then we have not really explained anything; we have just shifted the problem from understanding the brain to understanding the model.

We began by deriving a closed-form expression for the Tolhurst likelihood function that was more tractable than the infinite series that originally defined this function. This expression played a role in understanding two facets of the model’s performance: (a) the relationships between decoding precision and the neuronal parameters, and (b) the form of the psychometric function for contrast detection.

**Decoding precision**

To explain how the neuronal parameters map onto decoding precision, we derived an analytical approximation of the decoding precision, $\tilde{\tau}(x)$, which can be adjusted to apply to a variety of different spiking processes by setting the value of a single scalar parameter, $v$. For the Tolhurst and Consul-Jain processes, $v$ should be set to the Fano factor. For the Goris process, $v$ should be set to $1/(1 - \sigma_G^2)$, where $\sigma_G^2$...
is the variance of the gain signal. For the Tolhurst and Consul-Jain processes, $\tilde{s}(x)$ is an estimate of the Fisher information. For the Goris process, the Fisher information varies across trials, and $\tilde{s}(x)$ is its modal value.

The expression for $\tilde{s}(x)$ revealed some surprisingly simple relationships between decoding precision and the neuronal parameters, and explained the five observations of Clatworthy et al. (2003) that we investigated. For example, Equation 18 shows that,
A population of identical, statistically independent neurons, the height of the peak of decoding precision for a population of neurons, is approximately proportional to $r_{\text{max}}$. Sclar et al.’s (1990) study included one further V1 study.

The overall mean was the average of the mean $q$ values, weighted by the number of neurons in each study (this is equivalent to pooling the neurons across all three studies and finding the mean of the pooled sample).

### Table 1. Mean and median Naka-Rushton exponent, $q$, from three physiological studies of V1.

<table>
<thead>
<tr>
<th>Study</th>
<th>No. of neurons</th>
<th>Mean $q$</th>
<th>Median $q$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albrecht &amp; Hamilton (1982); cat</td>
<td>127</td>
<td>2.5</td>
<td>N/A</td>
</tr>
<tr>
<td>Albrecht &amp; Hamilton (1982); monkey</td>
<td>98</td>
<td>3.4</td>
<td>N/A</td>
</tr>
<tr>
<td>Geisler &amp; Albrecht (1997); cat</td>
<td>247</td>
<td>3.0</td>
<td>2.8</td>
</tr>
<tr>
<td>Geisler &amp; Albrecht (1997); monkey</td>
<td>71</td>
<td>2.5</td>
<td>2.0</td>
</tr>
<tr>
<td>Sclar et al. (1990); monkey</td>
<td>85</td>
<td>2.65</td>
<td>2.4</td>
</tr>
<tr>
<td>Overall mean</td>
<td></td>
<td>2.9</td>
<td></td>
</tr>
</tbody>
</table>

The psychometric function for contrast detection

To explain Chirimuuta and Tolhurst’s (2005a) finding regarding the slope of the psychometric function for contrast detection, we derived an analytical approximation of the model’s psychometric function, which we showed approaches a Weibull function with $\beta = q$ as $r_{\text{max}}$ or $K$ increase. We thus refuted Chirimuuta and Tolhurst’s conclusion that the standard Naka-Rushton function is unable to give Weibull $\beta$ values that are high enough to match those of human observers.

This example illustrates the power of the analytical approach taken here, compared with Monte Carlo simulations. Because of the considerable time that it takes to run the simulations, Chirimuuta and Tolhurst (2005a) could not feasibly explore every corner of the physiological study space.

It should be noted that our derivation of a psychometric function with the form of a Weibull function applies only to the case of $r_{0} = 0$. With nonzero $r_{0}$, the analytical form of the psychometric function is different (we have analyzed this more general case and will present it in another article). It is implausible that, in human vision, contrast detection is mediated entirely by neurons with zero spontaneous firing rate. The assumption that there is no neuronal response to zero contrast is often called the high-threshold assumption.

Under the conventional assumption of additive, stimulus-independent noise, the lack of response to zero contrast implies that there is a threshold on the output of the sensory units that lies enough standard deviations above the mean of the noise for there to be a negligible probability of a response to zero contrast. Because Chirimuuta and Tolhurst’s model has no sensory response to zero contrast, it is formally equivalent to a high-threshold model even though (with the standard Naka-Rushton function) it does not actually contain a sensory threshold. In high-threshold theory (and in Chirimuuta and Tolhurst’s model),
detection errors are always unlucky guesses on 2AFC trials that failed to elicit a response, whereas there is plenty of psychophysical evidence that incorrect responses are caused at least partly by hallucinations due to noise in the nontarget interval rather than entirely by unlucky guesses (Tanner & Swets, 1954; Swets, Tanner, & Birdsall, 1961; Nachmias, 1981; Solomon, 2007; Laming, 2013).

Despite the implausibility of Chirimuuta and Tolhurst’s model with \( r_0 = 0 \) as a model of contrast detection, we presented our analysis of it for three reasons.

1. It allows us to fully understand why Chirimuuta and Tolhurst’s (2005a) contrast detection simulations resulted in a fitted Weibull \( \beta \) that approached 2 with increasing number of neurons. This in turn allows us to refute their conclusion that the Naka-Rushton function requires a threshold to make it a plausible model of the neuronal contrast-response function.

2. Neurons with zero or negligible spontaneous firing rates do exist (e.g., see Dean, 1981a, figure 2), so it is not inconceivable that there are some organisms or experimental situations to which our analysis applies.

3. It allows us to address Tyler and Chen’s (2000) claim that high-threshold analysis of probability summation is “fundamentally flawed.” This idea is explored in the next subsection.

**High-threshold probability summation**

Our derivation of the model’s psychometric function for detection is an example of high-threshold probability summation. When \( r_0 = 0 \), each neuron acts as a detector; the observer detects the target if at least one neuron responds during the target presentation, and has to guess the correct answer otherwise. The more neurons the observer is monitoring, the greater the chance that at least one neuron will respond. The term probability summation refers to this increase in detection probability due to an increase in the number of detectors. The psychometric function in this case gives the probability that at least one neuron responds during the target presentation, or the observer guesses correctly if none respond.

Quick (1974) showed that if the detectors are statistically independent and each detector has a detection probability that is a Weibull function with the same \( \beta \), then the observer’s psychometric function will be a Weibull function with that \( \beta \)-value and with the detection threshold parameter, \( \alpha \), given by

\[
\alpha = \left( \frac{K}{\sum_{j=1}^{n} \alpha_j} \right)^{-1/\beta}.
\]

where \( \alpha_j \) is the detection threshold parameter of detector \( j \). For a very clear derivation of Equation 38, see Nachmias (1981), but note that his equation 4 has a typographical error: It is missing the minus sign on the exponent, \( \beta \). If all the detectors are identical (but statistically independent), then they all have the same \( \alpha_j \), and Equation 38 reduces to

\[
\alpha = K^{-1/\beta} \alpha_j.
\]

Thus, in Quick’s model, the Weibull \( \beta \) parameter controls how much of a reduction in detection threshold we achieve by increasing the number of detectors, \( K \). If \( \beta \) is low, then the detection threshold, \( \alpha \), decreases quickly as \( K \) increases; if \( \beta \) is high, then the detection threshold decreases more slowly as \( K \) increases.

Equation 38 gives the observer’s detection threshold according to Quick’s model, while Equation 30 gives the observer’s approximate detection threshold according to Chirimuuta and Tolhurst’s (2005a) model. Equation 38 has the exact form of Equation 30 if \( \beta = q \) and

\[
\alpha_j = \frac{(c_{1/2})_j}{\left(1 - 1/e(r_{max})\right)^{1/q}}.
\]

Thus, there is a near equivalence between Quick’s model and that of Chirimuuta and Tolhurst. If all the neurons in Chirimuuta and Tolhurst’s model have the same contrast-response function, then the model’s detection threshold approximation given by Equation 30 reduces to

\[
\alpha = K^{-1/q} \frac{c_{1/2}}{\left[1 - 1/e(r_{max})\right]^{1/q}}.
\]

Therefore, since \( \beta \approx q \) in Chirimuuta and Tolhurst’s model, their model shows approximately the same probability summation effects as Quick’s model, with detection threshold proportional to \( K^{-1/\beta} \). For the modeling in Figure 9, all the neurons were identical, so Equation 41 is equivalent to Equation 30 (which was used to generate the solid lines in Figure 9), and it is clear that this equation does accurately predict the detection threshold of Chirimuuta and Tolhurst’s model, particularly for the higher values of \( r_{max} \) or \( K \).

As noted previously, there is psychophysical evidence against Quick’s high-threshold model. Tyler and Chen (2000) went further, arguing not just that high-threshold theory has evidence against it but that the analysis of probability summation using high-threshold theory is “fundamentally flawed.” Their analysis of high-threshold theory made several basic assumptions about the high-threshold model, including the following five.

1. The observer monitors a set of channels.
2. Within each channel is a continuous signal that increases linearly with the stimulus strength.
3. Noise is added to this signal.
4. The noise might be additive (standard deviation independent of the signal level) or multiplicative (standard deviation proportional to a power function of the signal level), or the noise could be the sum of an additive and a multiplicative component.
5. A sensory threshold is applied to the noisy internal signal in each channel so that a stimulus is detected if and only if the noisy signal falls above threshold in at least one of the channels.

We should emphasize that Tyler and Chen (2000) were not arguing in support of this model; they argued that it was fundamentally flawed.

Tyler and Chen noted that if the internal noise in this model were fully multiplicative, then the internal signal-to-noise ratio would be unchanged by a change in stimulus strength. Thus, detection performance would remain the same for any stimulus level, and measurement of detection thresholds would be impossible! They concluded that even if there is a multiplicative component to the noise, detection performance must be limited by an additive component.

Tyler and Chen derived the shape of the additive noise distribution that would give rise to a Weibull psychometric function for the high-threshold model outlined previously. They showed that, for most Weibull $\beta$ values, the noise distribution deviated markedly from a Gaussian. They found this unacceptable because, according to the central limit theorem, the sum of a large number of non-Gaussian-distributed random variables is asymptotically Gaussian distributed. Therefore, if there are many different sources of noise from the stimulus to the neuronal decision mechanism, the noise is likely to be Gaussian at the decision mechanism. They then showed that high-threshold probability summation fails for additive Gaussian noise. They showed that, if the sensory threshold were low enough to be exceeded by the noisy signal in one channel 75% of the time, then the noisy signal would exceed the sensory threshold in at least one of 100 channels almost all the time, even when the stimulus intensity was reduced to zero. Thus, if the stimulus area or number of components increased so that the observer was monitoring many more channels, the observer would almost always be in a detect state, even when the stimulus was absent. The signal would have to be reduced to a physically unachievable negative contrast for the observer to be in a detect state 75% of the time. This problem does not occur with the noise distribution implied by the Weibull function because the Weibull probability density function falls to zero at the sensory threshold (see Supplementary Appendix K, especially Figure K.1); therefore, when the stimulus intensity is zero, no detector’s noisy signal will exceed its sensory threshold. However, as already noted, Tyler and Chen (2000) rejected that distribution because of its “bizarre” non-Gaussian nature when $\beta$ is not close to 4 (p. 3127). They therefore concluded that probability summation with high-threshold theory is fundamentally flawed.

Tyler and Chen’s arguments are perfectly valid, but they apply only to the set of high-threshold models defined by their assumptions (i.e., that each channel contains a continuous signal to which noise is added). With a detection model based on spiking neurons, the signal is discrete. When the spontaneous firing rate is zero, as it was in Chirimuuta and Tolhurst’s model, detection occurs when a single neuron produces a single spike. In these circumstances, it is not appropriate to approximate the neuronal signal as a continuous signal to which noise is added. It is perfectly legitimate to apply probability summation to find the probability that at least one neuron gives at least one spike. When we do this, we find that the model’s threshold is very close to being proportional to $K^{-1/\beta}$, as in standard high-threshold Weibull analysis. The contrast threshold for detection never reaches zero regardless of how many neurons the observer is monitoring, but we do not have to rely on bizarre model characteristics to achieve this. Aside from the zero spontaneous firing rate, the neurons in Chirimuuta and Tolhurst’s model have contrast-response functions and noise distributions that are physiologically plausible to a reasonable extent. One could ask what the equivalent “continuous signal plus additive noise” model is. Since Chirimuuta and Tolhurst’s model produces a psychometric function that closely approximates a Weibull function, the equivalent “continuous signal plus additive noise” model is closely approximated by the one derived by Tyler and Chen (and outlined in Supplementary Appendix K), with the “bizarre” noise distributions. However, this bizarreness comes from forcing Chirimuuta and Tolhurst’s model into the Procrustean bed of “continuous signal plus additive noise” rather than being an implausible characteristic of the model itself.

Conclusions

We derived equations that explained the performance of the contrast coding model described by Tolhurst and colleagues (Clatworthy et al., 2003; Chirimuuta & Tolhurst, 2005a). These equations gave a deep insight into the results of their Monte Carlo simulations.

As long as the stimulus contrast is high enough (and the neuronal population is large enough) to generate a sufficiently high population spike rate, the decoding precision can be closely approximated by the Fisher
information. We derived an estimate of decoding precision, $\tilde{v}(x)$, which approximates the Fisher information for a population of neurons with Tolhurst’s spiking process as long as the mean spike count of the most informative neurons is around five spikes or more. $\tilde{v}(x)$ is also an estimate of the Fisher information for the Consul-Jain spiking process. Furthermore, it gives the exact Fisher information for the Poisson process and the exact modal value of the Fisher information for the Goris process when the decoder has access to the gain signal in Goris et al.’s (2014) model of neuronal spiking. Our expression for $\tilde{v}(x)$ revealed simple relationships between the properties of the neurons and the decoding precision that hold to a good approximation when the mean count rate is sufficiently high. We used this expression to explain five relationships between decoding precision and the neuronal parameter values that Clatworthy et al. (2003) observed from their Monte Carlo simulations.

For the case of contrast detection, the spike rate is too low for the Fisher information to match decoding precision. To analyze the performance of Chirimuuta and Tolhurst’s (2005a) model in a 2AFC contrast detection task, we used basic probability theory. We derived an expression for the model’s psychometric function for contrast detection and showed that as $K$ or $r_{\text{max}}$ increase, the psychometric function asymptotically approaches a Weibull function with $\beta = q$. Our work therefore reveals a previously unknown theoretical connection between two of the most widely used functions in vision science: the Weibull psychometric function and the Naka-Rushton contrast-response function. This relationship explained why Chirimuuta and Tolhurst always obtained a Weibull $\beta$ of about 2 in their modeling (they always had $q = 2$ in their assessments of the model’s Weibull $\beta$) and allowed us to refute their conclusion that it is necessary to have a threshold on the Naka-Rushton function to achieve Weibull $\beta$ values that match those found with human observers. Their threshold on the Naka-Rushton function had a similar effect to increasing $q$, as it made the spike rate increase more abruptly with increasing contrast.

**Keywords:** Fisher information, doubly stochastic Poisson distribution, decoding, detection, probability summation

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### Footnotes

1 In this paper, we use the word *trial* in two ways. First, we use it in the way a physiologist would, to mean a stimulus presentation. Second, we use it to mean a trial on a two-alternative forced-choice (2AFC) psychophysical experiment, in which the observer is presented with two stimuli and has to make a response. To distinguish these two meanings, we always refer to the latter type of trial as a 2AFC trial.

2 In this discussion, we use the word *threshold* in two ways: (a) to refer to an internal threshold on the sensory signal, and (b) to refer to the stimulus contrast corresponding to a particular level of detection performance. We have tried to make the meaning clear in each case by using the term *sensory threshold* for the former case and the term *detection threshold* for the latter.

3 As pointed out by Mortensen (2002), Tyler and Chen’s (2000) published equation for the probability density function (PDF) of the noise (Tyler and Chen’s equation 4b) contains errors. However, Tyler and Chen’s plots of the noise PDFs (shown in their figure 2b) are correct, so we assume that the errors in Tyler and Chen’s equation 4b are typographical errors rather than fundamental problems with their analysis. To clarify matters, we present in Supplementary Appendix K a derivation of the PDF that is based on Mortensen’s derivation but is hopefully easier to follow than Mortensen’s derivation or that of Tyler and Chen.

### References


Britten, K. H., Shadlen, M. N., Newsome, W. T., &


