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A ratio model of perceived speed in the human visual system

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The perceived speed of moving images changes over time. Prolonged viewing of a pattern (adaptation) leads to an exponential decrease in its perceived speed. Similarly, responses of neurones tuned to motion reduce exponentially over time. It is tempting to link these phenomena. However, under certain conditions, perceived speed increases after adaptation and the time course of these perceptual effects varies widely. We propose a model that comprises two temporally tuned mechanisms whose sensitivities reduce exponentially over time. Perceived speed is taken as the ratio of these filters' outputs. The model captures increases and decreases in perceived speed following adaptation and describes our data well with just four free parameters. Whilst the model captures perceptual time courses that vary widely, parameter estimates for the time constants of the underlying filters are in good agreement with estimates of the time course of adaptation of direction selective neurones in the mammalian visual system.

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1. INTRODUCTION

Given that the early stages of visual processing are relatively well understood, it is surprising that there is no consensus as to how the human visual system encodes the 32 speed of a moving image. There are at least three classes of model that have attempted to characterize this funda-34 mental property of vision: First, motion energy models (we include here also gradient and Reichardt approaches 36 which may be differently motivated but are formally equivalent to motion energy models; Adelson & Bergen 38 1985; van Santen & Sperling 1985; Watson & Ahumada 1985), second, response frequency models (Barlow & Hill 1963; Clifford & Langley 1996; Bex et al. 1999) and most recently, Bayesian models of speed perception (Ascher & Grzywacz 2000; Hurlimann et al. 2002; Weiss et al. 2002). None of these models can capture what we know about how perceived speed may vary following adaptation to a 45 moving pattern.

46 Motion energy models assume that non-directional 47 spatiotemporal filters are added and subtracted in 48 quadrature pairs to create direction selective units. A 49 squaring of the units' outputs ensures a smooth response 50 and a differencing of left and right sensitive units gives an 51 unambiguous directional, but not speed, signal. This class 52 of model confounds velocity and contrast and, although 53 scaling the detector's output by some 'static' detector's 54 response (Adelson & Bergen 1986) would eliminate 55 contrast dependence, the motion energy model does not 56 concern itself with how an ensemble of detectors might 57 encode speed. Clearly an array of 'motion detectors' each 58 tuned to a different speed might be envisaged but such a 59 'labelled lines' approach to speed encoding is not 60 consistent with much of what we know about human 61

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speed perception. For instance, adaptation to a moving stimulus reduces the perceived speed of that stimulus and all slower speeds in the same direction and may increase the perceived speed of faster stimuli (Carlson 1962; Clymer 1973; Thompson 1981; Smith & Edgar 1994). Similarly, reducing the contrast of a slowly moving pattern reduces its perceived speed (Thompson 1982; Stone & Thompson 1992; Hawken et al. 1994). These findings do not point towards a 'place' or labelled lines model of speed in which the output of an ensemble of narrowly tuned overlapping channels determines speed (Heeger 1987; Grzywacz & Yuille 1990). Nor do they fit well with the Bayesian class of models that assume that a prior will 'distort' our estimate of speed. Whilst this class of model can account for distortions in one direction, e.g. reductions in perceived speed, they cannot simultaneously predict both increases and decreases in perceived speed that are contingent upon prevailing viewing conditions without the invocation of a somewhat arbitrary second prior.

Another approach, which we term the Response 113 Frequency model, has been to postulate that speed is 114 encoded in the frequency of response of direction-selective 115 mechanisms (Barlow & Hill 1963). This approach 116 envisages a 'frequency' code similar to that assumed to 117 underlie contrast coding. Such a proposal readily explains 118 the reduction in perceived speed of patterns following 119 adaptation in a fashion analogous to Blakemore et al.'s 120 121 (1973) model of contrast adaptation. Recently, a number 122 of studies have attempted to elucidate the nature and 123 function of speed adaptation by measuring the time course 124 of its effects (Clifford & Langley 1996; Bex et al. 1999). 125 The decay of perceived speed as a function of adaptation 126 duration has been found to be well fit by a simple 127 exponential model of the reduction of spike rates in cat 128

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129 cortical cells after motion adaptation (Giaschi et al. 1993). 130 Upon the basis of these findings it has been proposed that the exponential decay of spike rates may form the neural 131 132 substrate for motion adaptation. This suggestion has a long history and offers a potentially powerful model for 133 many perceptual phenomena (Sutherland 1961; Barlow & 134 135 Hill 1963) but in its simplest form this model struggles to 136 explain well-established phenomena. For instance, how 137 could observed increases in speed following adaptation (Carlson 1962; Rapoport 1964; Clymer 1973; Thompson 138 1981; Smith & Edgar 1994) be explained? Moreover, 139 140 whilst a large range of time courses for perceptual decay in 141 perceived speed has been found (Clifford & Ibbotson 2002), physiological estimates of reduction in spike rates 142 143 converge on a much smaller range (Maddess & Laughlin 1985; Giaschi et al. 1993; Ibbotson & Mark 1996). The 144 solution to these problems may lie in a ratio model of 145 speed encoding for which there is much psychophysical 146 evidence (Harris 1980; Thompson 1982; Harris 1986; 147 148Smith & Edgar 1994).

149 We have previously measured speed adaptation and its 150 subsequent recovery and have shown that decreases in 151 perceived speed after adaptation were consistent with a 152 ratio model of perceived speed (Hammett et al. 2000). 153 Here we report the results of a parametric study of perceived speed as a function of adaptation duration, test 154 and adaptation speed. We measured the perceived speed 155 of sinusoidal gratings that drifted over a wide range of 156 speeds after adapting to a moving grating for one of six 157 adaptation durations. Six adaptation speeds were investi-158 gated with adaptation gratings always moving in the same 159 160 direction as the test gratings. In order to ensure that contrast adaptation did not confound adaptation to speed 161 per se (Blakemore et al. 1973; Thompson 1981; Hammett 162 163 et al. 1994) we ensured that the contrast of adapting 164 patterns was much lower than that of the test patterns. We 165 show that both the increases and decreases in perceived speed observed in these psychophysical measurements are 166 well described by a model that assumes speed is encoded 167 168 as the ratio of two temporal filters whose sensitivities decay exponentially over time. 169

172 2. METHODS

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173 (a) Apparatus & stimuli

All stimuli were horizontally oriented sinusoidal gratings 174 generated on a VSG 2/3W (Cambridge Research Systems) 175 waveform generator and displayed on an EIZO 6600-M 176 Monochrome monitor. The monitor was gamma corrected 177 using the CRS OPTICAL photometric system. Mean 178 luminance was 32 cd m^{-2} and the frame rate was 100 Hz. 179 The active display subtended $30^{\circ} \times 24^{\circ}$ and the gratings were 180 presented in circular windows (diameter=6°) situated such 181 that the inner edge was 1° to the left (adapting patterns) or 1° 182 to the right (test patterns) of a small dark fixation spot. The 183 spatial frequency was always 1 c $^{\circ-1}$ and the test and adapting 184 patterns drifted upwards at one of a range of speeds $(2-20^{\circ} \text{ s}^{-1})$ 185 . The stimuli were windowed with sharp edges in both space 186 and time. The adapting contrast was 10% and the test 187 contrast was 50%. The viewing distance was 57 cm. 188

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190 (**b**) *Procedure*

At the beginning of each trial two stimuli were presentedsimultaneously for 500 ms. The subjects' task was to indicate

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which of the two patterns (left or right) appeared faster by 193 pressing a button. An adapting pattern was subsequently 194 presented to the left of the central fixation point for each of 195 five durations (4, 8, 16, 32 or 64 s). Following adaptation the 196 adapting stimulus was replaced with a blank field of mean 197 luminance for 10 ms. Subsequently two stimuli were 198 presented simultaneously for 500 ms. A standard pattern 199 was located in the original position of the adapting pattern, 200 and a test pattern, whose speed was controlled by a modified 201 PEST procedure (Taylor & Creelman 1967) was located to 202 the right of the fixation point. The subject indicated which 203 pattern appeared faster after each presentation. The speed of 204 the test patterns was altered by independent PEST pro-205 cedures that were set to converge on the 50% point using six 206 interleaved staircases. After each such run subjects rested for 207 at least 2 min in order to minimize build up of adaptation. 208 Thirty such runs were taken and the 50% point of the 209 resultant psychometric function was estimated by Probit 210 (Finney 1971). The mean of four such estimates was taken as 211 the PSE. 212

The experiments were conducted binocularly in a semidarkened room with no head restraint. One of the subjects was naïve to the purpose of the experiment, the other was aware of the general purpose of the experiment. Both subjects were experienced psychophysical observers. 213

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3. RESULTS

The results (along with the best fit of the model described below) are shown in figure 2. For the higher adaptation speeds (>12° s⁻¹), the effect of adaptation was to reduce perceived speed at all test speeds $(2-20^{\circ} \text{ s}^{-1})$. Perceived speed reduced quasi-exponentially as a function of adaptation duration. This is in good agreement with previous psychophysical studies (e.g. Bex et al. 1999; Hammett et al. 2000). However, for the lower adaptation speeds (2 and $4^{\circ} s^{-1}$), the effect of adaptation depended upon test speed. At low-test speeds, adaptation resulted in a reduction in perceived speed similar to that observed for high adaptation speeds. However, under conditions where adaptation speed was low (2 and $4^{\circ} s^{-1}$) and test speed was higher $(>8^{\circ} s^{-1})$ perceived speed increased quasiexponentially as a function of adaptation duration. Whilst no such increase as a function of time has been reported before, the basic effect of an increase in speed under such conditions is well documented (Thompson 1981). Thus the psychophysical data indicate that adaptation can lead to both increases and decreases in perceived speed depending upon adapt and test speeds and that these effects take a quasi-exponential form. In the following section we develop a simple ratio model, based upon physiologically plausible temporal filters, in order to capture these characteristics of speed perception.

(a) The model

The model assumes that perceived speed is based upon the ratio of a low pass and band-pass temporal filter. The model employs the filters recently proposed by Perrone (2005). These filters provide a good fit to typical tuning functions in macaque V1. The low-pass filter takes the form

$$p(\omega) = \sqrt{a^2 + b^2},$$
 (1) 255
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286 Figure 1. The filter sensitivities for p (solid line) and m287 (broken line) are shown in the upper panel. The decay of peak 288 filter response as a function of adaptation duration is illustrated in the lower panel for cases of near-optimal 289 adaptation (i.e. at values of ω_a close to peak filter 290 sensitivity). The decay functions have been normalized with 291 respect to peak filter response. The decay functions are shown 292 for p (solid line) at $\omega_a = 2$ Hz and for m (broken line) at $\omega_a =$ 293 12 Hz using the best fit of the model to data averaged over 294 both subjects. 295

²⁹⁶ where ²⁹⁷

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 $a = ((2\pi\omega\tau_1)^2 + 1)^{-(9/2)}$ and $b = ((2\pi\omega\tau_2)^2 + 1)^{-(10/2)}$,

300 and the band pass filter is given by

$$\begin{array}{l} 301\\ 302\\ 303 \end{array} \quad m(\omega) = \frac{\omega}{k} p(\omega), \tag{2}$$

304 where ω is temporal frequency. Following Perrone, we 305 have used values of 0.0072, 0.0043 for the time constants 306 (measured in seconds) τ_1 and τ_2 , respectively. The 307 parameter k is set to 4. We have adhered to Perrone's 308 convention of labelling the low-pass and band-pass filters p 309 and m, respectively. As noted by Perrone, a simple ratio of 310 $m(\omega)/p(\omega)$ is proportional to ω , a desirable feature of a 311 speed encoding mechanism. Note that there are no free 312 parameters at this stage of the model. Figure 1 (upper 313 panel) shows the filter sensitivities for these parameter 314 values. The filter sensitivities are determined by an 315 additive combination of the two low pass filters, a and b, 316 the features of which are in turn determined by the time 317 constants τ_1 and τ_2 . Due to the differences in time 318 constants used here, the upper limb of $p(\omega)$ is dominated 319 by the component b. For frequencies greater than $(2\pi\tau_2)^{-1}$ (i.e. 37 Hz) the expression tends to $(2\pi\omega\tau_2)^{-10}$ and thus 320

tends to a slope of -10 on a log-log plot. The component, 321 a, plays its role on the lower limb of the combined filter 322 $p(\omega)$ by controlling the corner frequency (i.e. the 323 frequency at which response is attenuated 3 dB). For the 324 filters used here the corner frequency is 8 Hz. For the 325 band-pass filter, $m(\omega)$, upper and lower limbs are similarly 326 dominated by the terms a and b. The term k scales the 327 filter's response. For the parameter values used here, peak 328 response of $m(\omega)$ is at 10.8 Hz. 329

The model comprises four free parameters that determine the time constants of exponential decay and asymptotic response attenuation for the low pass and band-pass filters. Thus the filters' sensitivities decay over time such that their responses as a function of speed and time are given by:

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$$P(\omega,t) = p(\omega_{\psi}) - K_{p} p(\omega_{a}) \left\{ 1 - \mathrm{e}^{-t/T_{p}} \right\}, \tag{3}$$

$$M(\omega, t) = m(\omega_{\psi}) - K_m m(\omega_a) \{ 1 - e^{-t/T_m} \}.$$
 (4) 338
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Where ω_a and ω_{ψ} are the adaptation and test frequencies, 340 respectively, and t is adaptation duration measured in 341 seconds. The free parameters T_p and T_m are the time 342 constants, measured in seconds, of the adaptation 343 dependent reduction in filter sensitivities. The other free 344 parameters K_p and K_m determine the magnitude of the 345 reduction in the filters' sensitivities resulting from 346 adaptation, that is the maximum (asymptotic) response 347 attenuation. Thus the model assumes that filter sensi-348 tivities will decay exponentially over time by an amount 349 determined by the product of the free parameter K_p (or 350 K_m) and a term that reflects the relative sensitivity of the 351 filter to the adaptation frequency $p(\omega_a)$. Figure 1 (lower 352 panel) illustrates how the peak responses of these filters 353 decrease for the case where adaptation frequency is near 354 the peak sensitivity of the respective filters. For the case 355 $\omega_a = \omega_{\psi}$ a value of unity for K_p or K_m could result in zeroing 356 the filter's response as t tends to infinity. Although K_p and 357 K_m were not constrained in the model fitting, it was 358 anticipated that they would lie between 0 and 1 so as to 359 limit the effect of adaptation on the filter responses to 360 plausible values. 361

Perceived speed S is given by the ratio of the output of these two filters such that:

$$S = \frac{M(\omega, t)}{P(\omega, t)}.$$

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In order to estimate the time constants and saturation levels of the filters we found the best-fitting values of these free parameters using an error minimization function (*fminsearch*) in MATLAB 7.0.1 (Mathworks Inc). The resultant best fits (solved for all adaptation speeds, test speeds and adaptation durations simultaneously) are shown in figure 2, alongside the psychophysical data.

The psychophysical estimates of perceived speed in the 374 absence of adaptation (at t_0) were near veridical (the 375 average standard error was within 4.2% of the physical 376 speed). However, in order to avoid the model fit being 377 biased by measurement error at t_0 perceived speed for each 378 condition was scaled to be veridical at t_0 and this scaling 379 factor was subsequently applied to all other points within 380 381 the condition. This is equivalent to constraining the model 382 fit such that perceived speed is veridical in the absence of 383 adaptation. The model parameters for the best fits for each 384 subject are shown in table 1. The time constants are in





Figure 2. Perceived speed (solid symbols) as a function of adaptation duration and the best fit of the model (solid lines). Each panel represents the results for one adaptation speed (indicated above the graph). Each test speed is represented by a different colour. The results for each subject (SB and DS) are shown separately. Error bars represent \pm s.e.m.

427 good agreement with those reported for the decay in 428 response of single directionally selective neurones in the 429 cat cortex (Giaschi et al. 1993; around 5-8 s) and our 430 previous estimates of the underlying filters' time constants 431 (Hammett et al. 2000; 8 and 7.25 s for T_p and T_m , 432 respectively). The model captures the essential character-433 istic of both increases and decreases in perceived speed, 434 contingent upon the adapt-test relationship. The model 435 captures both increases and decreases in perceived speed 436 because p may decrease proportionately more or less than 437 m as a function of adaptation duration. Overall, the model 438 captures around 96% of the variance of the data (table 2). 439 There are, however, a few conditions where the fits are less 440 adequate. These tend to be at moderate test speeds (8 and 441 $12^{\circ} \text{ s}^{-1}$). However, even here, the lowest r^2 value is 0.42 442 and is typically much higher than that. Encouragingly, 443 these less adequate fits are not confined to the same 444 conditions across subjects and it seems reasonable to 445 assume that the lower r^2 values are a reflection of the 446 447 transitional nature of these conditions. Note that whilst low adaptation speeds (e.g. 2 and 4° s^{-1}) yield increases in 448

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perceived speed at high test speeds, high adaptation speeds (e.g. 16 and 20° s⁻¹) yield reductions in perceived speed. Thus, the model must render a point of inflection at some intermediate speed in order to capture both effects. The model does this smoothly but tends to miss the subjects' actual point of inflections. Given this, and the fact that the model still adequately fits all conditions, it seems reasonable to assume that the lower r^2 values for these conditions is attributable to 'under-constraining' the model with respect to the point of inflection rather than to a systematic error in the model.

4. DISCUSSION

Previous researchers have suggested that the neural 505 substrate for *decreases* in perceived speed may be the 506 decrease in single unit activity of cells sensitive to motion 507 (Barlow & Hill 1963; Clifford & Langley 1996; Bex *et al.* 508 1999). Whilst this approach has intuitive appeal, it fails to account for the *increases* in perceived speed after 510 adaptation that are reported here and elsewhere 511 (Thompson 1981). Similarly, whilst motion energy 512

13	Table	1.	Model	parameters	for	the	best	fits	for	each	subjec	ct.
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parameter	DS	SB
T_{p}	13.42	6.3
T_m	10.26	6.1
K _p	0.05	0.0
K_m^P	0.18	0.1

Table 2. r^2 values for the model fit for each test speed and overall fit at all test speeds.

test speed (° s $^{-1}$)	DS	SB
2	0.9979	0.9983
4	0.9988	0.9971
8	0.8507	0.9404
12	0.4225	0.6149
16	0.9785	0.8950
20	0.9842	0.9796
overall	0.9774	0.9616

535 models have been highly successful at describing how 536 direction of motion may be computed, they require a 537 'labelled line' approach to speed encoding that is not 538 consistent with much psychophysical data. Furthermore, 539 it is not clear how they should account for increases in 540 perceived speed. A final class of model that uses Bayesian 541 priors (e.g. Hurlimann et al. 2002) to explain reductions in 542 perceived speed under certain circumstances should also 543 have difficulty in accommodating both increases and 544 decreases in speed after adaptation without assuming 545 that the prior can change. 546

The present results indicate that perceived speed is 547 exquisitely sensitive to prevailing viewing conditions and 548 can increase or decrease over time, depending upon 549 previously seen speeds. The time course of these velocity 550 after-effects is well described by an exponential function, 551 regardless of whether perceived speed increases or Q3 552 decreases over time. These findings are in agreement 553 with previous psychophysical studies that have measured a 554 much more limited range of adaptation speeds and 555 durations. Around 96% of the variance in our data can 556 be resolved by a simple ratio model of perceived speed that 557 has only four free parameters that control the time 558 constants and asymptotic response attenuation of a low-559 pass and a band-pass temporal filter. The filters are 560 consistent with the temporal tuning properties of cortical 561 cells in the macaque (Foster et al. 1985; Hawken et al. 562 1996) and bear a close resemblance to those suggested by 563 564 Q2 Anderson and Burr's (1985) psychophysical study. The time constants of the best fitting model are consistent with 565 the empirically derived time constants reported for the 566 decay in response of single cortical neurones in the cat 567 (Giaschi et al. 1993). Furthermore, psychophysical 568 569 estimates of the time constants of adaptation vary widely, 570 from 1 to 16 s (Clifford & Ibbotson 2002), depending 571 upon stimulus parameters. In the present study, our more 572 extensive psychophysical estimates of perceived speed 573 yielded time constants for best fitting exponentials over an 574 even greater range (from 2 to 46 s for conditions where Q4 575 perceived speed reduced). Despite these wide differences 576 in time constants for the perceptual phenomenon,

estimates of the model's filter time constants are very 577 similar to the estimates obtained for single cells whilst 578 their ratio simultaneously captures the wider ranging 579 psychophysically derived time constants. Thus the model 580 581 captures large changes in perceptual dynamics with 582 physiologically plausible and stable underlying filter 583 dynamics.

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We conclude that a model that employs physiologically plausible temporal filters and that assumes that speed is encoded as the ratio of the output of those filters adequately captures many of the characteristics of perceived speed with just four free parameters. The question of what may constitute the neural substrate of these filters is clearly of great interest. It has not escaped our notice that Perrone (2005) labelled his low-pass and band-pass filters p and m, respectively, and that both parvo- and magno-cellular pathways appear to be implicated in the creation of direction selective cells in V1(DeValois et al. 2000). However, the very different contrast gain (Kaplan & Shapley 1986; Sclar et al. 1990), and presumably adaptation, properties of the two pathways may mitigate against such a scheme. We are currently investigating whether the ratio of magno- and parvocellular responses may be implicated in the computation of object speed.

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REFERENCES

- Adelson, E. H. & Bergen, J. R. 1985 Spatiotemporal energy models for the perception of motion. J. Opt. Soc. Am. A 2, 284-299.
- Adelson, E. H. & Bergen, J. R. 1986 The extraction of spatiotemporal energy in human and machine vision. Proc. workshop on motion: representation and analysis, pp. 151-155. Charleston, SC.
- Ascher, D. & Grzywacz, N. M. 2000 A Bayesian model for the measurement of visual velocity. Vision Res. 40, 3427-3434. (doi:10.1016/S0042-6989(00)00176-0.)
- Barlow, H. B. & Hill, R. M. 1963 Evidence for a physiological explanation of the waterfall phenomenon and figural aftereffects. Nature 200, 1345-1347.
- Bex, P. J., Bedingham, S. & Hammett, S. T. 1999 Apparent speed and speed sensitivity during adaptation to motion. J. Opt. Soc. Am. A 16, 2817-2824.
- Blakemore, C., Muncey, J. P. & Ridley, R. M. 1973 Stimulus specificity in the human visual system. Vision Res. 13, 1915-1931. (doi:10.1016/0042-6989(73)90063-1.)
- Carlson, V. R. 1962 Adaptation in the perception of visual velocity. J. Exp. Psychol. 64, 192-197.
- Clifford, C. W. & Ibbotson, M. R. 2002 Fundamental mechanisms of visual motion detection: models, cells and functions. Prog. Neurobiol. 68, 409-437. (doi:10.1016/ S0301-0082(02)00154-5.)
- Clifford, C. W. & Langley, K. 1996 Psychophysics of motion adaptation parallels insect electrophysiology. Curr. Biol. 6, 1340-1342. (doi:10.1016/S0960-9822(02)70721-5.)
- Clymer, A. 1973 The effects of seen motion on the apparent speed of subsequent test velocities: speed tuning of movement aftereffects. New York: Columbia University.
- De Valois, R. L., Cottaris, N. P., Mahon, L. E., Elfar, S. D. & Wilson, J. A. 2000 Spatial and temporal receptive fields of

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- 6 S. T. Hammett and others A ratio model of perceived speed in the human visual system
- 641 geniculate and cortical cells and directional selectivity.
 642 *Vision Res.* 40, 3685–3702. (doi:10.1016/S0042-6989(00)
 643 00210-8.)
- Finney, D. 1971 *Probit analysis.* Cambridge: Cambridge
 University Press.
 Foster K H. Coche J. D. Markov, A. B. Mar
- Foster, K. H., Gaska, J. P., Nagler, M. & Pollen, D. A. 1985
 Spatial and temporal frequency selectivity of neurones in visual cortical areas V1 and V2 of the macaque monkey. *Physiol.* 365, 331–363.
- Giaschi, D., Douglas, R., Marlin, S. & Cynader, M. 1993
 The time course of direction-selective adaptation in simple and complex cells in cat striate cortex. *J. Neurophysiol.* 70, 2024–2034.
- 653 Grzywacz, N. M. & Yuille, A. L. 1990 A model for the
 654 estimate of local image velocity by cells in the visual cortex.
 655 *Proc. R. Soc. B Biol. Sci.* 239, 129–161.
- Hammett, S. T., Snowden, R. J. & Smith, A. T. 1994
 Perceived contrast as a function of adaptation duration. *Vision Res.* 34, 31–40. (doi:10.1016/0042-6989(94)
 90254-2.)
- Hammett, S. T., Thompson, P. G. & Bedingham, S. 2000
 The dynamics of velocity adaptation in human vision. *Curr. Biol.* 10, 1123–1126. (doi:10.1016/S0960-9822(00)
 00698-9.)
- Harris, M. G. 1980 Velocity specificity of the flicker to pattern sensitivity ratio in human vision. *Vision Res.* 20, 687–691. (doi:10.1016/0042-6989(80)90093-0.)
- Harris, M. G. 1986 The perception of moving stimuli:
 a model of spatiotemporal coding in human vision. *Vision Res.* 26, 1281–1287. (doi:10.1016/0042-6989(86)
 90109-4.)
- Hawken, M. J., Gegenfurtner, K. R. & Tang, C. 1994
 Contrast dependence of colour and luminance motion mechanisms in human vision. *Nature* 367, 268–270. (doi:10.1038/367268a0.)
- Hawken, M. J., Shapley, R. M. & Grosof, D. H. 1996
 Temporal-frequency selectivity in monkey visual cortex. *Vis. Neurosci.* 13, 477–492.
- Heeger, D. J. 1987 Model for the extraction of image flow.
 J. Opt. Soc. Am. A 4, 1455–1471.
- ⁶⁷⁸ Hurlimann, F., Kiper, D. C. & Carandini, M. 2002 Testing
 ⁶⁷⁹ the Bayesian model of perceived speed. *Vision Res.* 42,
 ⁶⁸⁰ 2253–2257. (doi:10.1016/S0042-6989(02)00119-0.)

- Ibbotson, M. R. & Mark, R. F. 1996 Impulse responses distinguish two classes of directional motion-sensitive neurons in the nucleus of the optic tract. *J. Neurophysiol.* 75, 996–1007.
- Kaplan, E. & Shapley, R. M. 1986 The primate retina contains two types of ganglion cells, with high and low contrast sensitiviy. *Proc. Natl Acad. Sci. USA* 83, 2755–2757.
- Maddess, T. & Laughlin, S. B. 1985 Adaptation of the motion sensitive neuron H-1 is generated locally and governed by contrast frequency. *Proc. R. Soc. B* **225**, 251–275.
- Perrone, J. A. 2005 Economy of scale: a motion sensor with variable speed tuning. *J. Vision* 5, 28–33. (doi:10.1167/5.1.3.)
- Rapoport, J. 1964 Adaptation in the perception of rotary motion. J. Exp. Psychol. 67, 263–267.
- Sclar, G., Maunsell, J. H. R. & Lennie, P. 1990 Coding of image contrast in the central visual pathways of the macaque monkey. *Vision Res.* **30**, 1–10. (doi:10.1016/ 0042-6989(90)90123-3.)
- Smith, A. T. & Edgar, G. K. 1994 Antagonistic comparison of temporal frequency filter outputs as a basis for speed perception. *Vision Res.* 34, 253–265.
- Stone, L. S. & Thompson, P. 1992 Human speed perception is contrast dependent. *Vision Res.* 32, 1535–1549. (doi:10. 1016/0042-6989(92)90209-2.)
- Sutherland, N. 1961 Figural aftereffects and aparent size. *Q. J. Exp. Psychol.* **13**, 222–228.
- Taylor, M. & Creelman, C. 1967 Pest: efficient estimates on probability functions. J. Acc. Soc. Am. 41, 782–787.
- Thompson, P. 1981 Velocity after-effects: the effects of adaptation to moving stimuli on the perception of subsequently seen moving stimuli. *Vision Res.* 21, 337–345. (doi:10.1016/0042-6989(81)90161-9.)
- Thompson, P. 1982 Perceived rate of movement depends on contrast. *Vision Res.* **22**, 377–380. (doi:10.1016/0042-6989(82)90153-5.)
- van Santen, J. P. & Sperling, G. 1985 Elaborated Reichardt detectors. J. Opt. Soc. Am. A 2, 300–321.
- Watson, A. B. & Ahumada Jr, A. J. 1985 Model of human visual-motion sensing. J. Opt. Soc. Am. A 2, 322–341.
- Weiss, Y., Simoncelli, E. P. & Adelson, E. H. 2002 Motion illusions as optimal percepts. *Nat. Neurosci.* 5, 598–604. (doi:10.1038/nn858.)

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