Apparent speed and speed sensitivity during adaptation to motion

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Adaptation, a change in response to a sustained stimulus, can be demonstrated in motion perception by velocity aftereffects—changes in the apparent speed of a moving pattern following adaptation. We measured changes in the apparent speed of sinusoidal gratings drifting at 4 or 7.5 deg/s during 30 s of adaptation followed by 30 s of recovery. The apparent speed of the patterns fell to approximately half the unadapted apparent speed, and the time constants of adaptation were much faster (5 s) than for recovery (22 s). Part of the loss of apparent speed (approximately 12%) was related to a loss of apparent contrast with adaptation. Sensitivity to speed increments and speed decrements increased during adaptation and was well described by a Weber fraction based on apparent speed. The results suggest that adaptation to motion, like light adaptation, may serve to improve an observer's sensitivity to the prevailing environment. © 1999 Optical Society of America


1. INTRODUCTION

After prolonged inspection of a moving pattern, a subsequently viewed static image can appear to move in the opposite direction to that of the adapting pattern. This phenomenon is called the motion aftereffect (MAE) and has been studied in considerable detail (for a review see Refs. 1 and 2). A somewhat less-well-studied phenomenon that can occur following adaptation to motion concerns a change in the apparent speed of a subsequently viewed moving pattern. This was first reported by Wohlgemuth,3 who showed that the apparent speed of a drifting black-and-white-striped belt was reduced following 30 s of adaptation. This effect is now known as the velocity aftereffect4 (VAE) and has been confirmed with several psychophysical techniques, including magnitude estimation,5–7 speed matching,4,8–12 and estimation of the time that it would take for a line to travel a particular distance.13

Many studies of VAE’s have investigated combinations of adaptation and test speeds, directions, and contrasts. Several authors have reported that perceived velocity is reduced following adaptation to a stimulus moving faster than, and in the same direction as that of, the target.4,7–9,11,13 Adaptation to a pattern moving more slowly than, and in the same direction as that of, the test stimulus has been found to result in an increase in perceived speed for rotary patterns5 but no change or a small decrease for drifting vertical gratings.8 When adaptation is in the direction opposite to that of the target, perceived velocity has been reported to increase,14 to be variably affected,7 or to decrease,8 but only if the adaptation pattern moves as fast as or faster than the test pattern.9 Smith and Edgar10 have modeled this complex collection of results with a ratio model of motion processing.15 The model is based on the antagonistic comparison of two temporal channels, one low pass and the second bandpass16–21 (cf. alternative results implicating three temporal channels22,23). To account for both increases and decreases in apparent speed that can occur under certain conditions, the model assumes that the bandpass channel is more adaptable than the low-pass channel.

Psychophysical studies of the time course of motion adaptation and recovery from adaptation have concentrated on the conventional MAE. While there is some evidence of MAE’s lasting up to three days for spiral patterns,24 most studies of the time course of the MAE (Refs. 25–29) and the VAE (Ref. 12) have found much shorter time constants that are well fitted by exponential functions. The exponential buildup and decay of adaptation to motion compares well with physiological measures of neural response changes in area 17 of cats30–33 and in area H1 of flies.34 Giaschi et al.33 also recorded recovery rates, finding them to be somewhat slower than adaptation rates (approximately 8 s compared with 5 s). They also found an initial rapid decline in response rate, followed by a much slower steady decline that was best fitted by combined fast and slow exponential functions.

The aim of the present investigation was to measure the changes in apparent velocity of medium-speed and high-speed drifting sine-wave gratings during a 30-s adaptation period and the subsequent recovery of normal apparent speed. Recently, Clifford and Langley12 measured sensitivity to sinusoidal oscillations in speed during speed adaptation. They found that subjects were able to detect oscillations after a few seconds of adaptation that they were unable to detect at the beginning of the adap-
We sought to confirm this result and to relate any changes in speed sensitivity to changes in apparent speed.

2. METHODS

One of the authors (PB) and a paid, naive volunteer (RW) served as observers. Stimuli were generated on a Macintosh 8600/200 using software adapted from the VideoToolbox routines\(^{35}\) and were displayed on an Eizo Flexscan 6600 gray-scale monitor at a frame rate of 120 Hz and mean luminance (\(L_0\)) of 50 cd/m\(^2\). The luminance of the display was linearized with pseudo-12-bit resolution\(^{36}\) and calibrated with an OptiCal photometer. Pseudo-12-bit resolution in this case allowed the presentation of 2\(^8\) gray levels from a possible range of 2\(^{12}\) levels. The display measured 20 cm horizontally (640 pixels) and 15 cm vertically (480 pixels) and was in a dark room 57 cm from the observer.

Stimuli in all cases were horizontal sinusoidal gratings [2 cycles per degree (c/deg)] drifting within stationary Gaussian windows (\(\sigma_{x,y} = 1\) deg):

\[
L_{x,y} = L_0[1 + C \exp(- (x^2 + y^2)/2\sigma^2)] \sin(2 \pi x/\lambda),
\]

centered at 2° to the left or the right of a central, black fixation cross.

A. Procedure

Each run consisted of 16 sessions, where each session was a 30-s adaptation period followed by a 30-s recovery period. There was a match interval every 3 s throughout adaptation and recovery. The adapting pattern was presented continuously throughout the adapting period, then for 1 s every 3 s throughout the recovery period. The match pattern was presented for 1 s every 3 s throughout the adaptation and recovery periods. The temporal frequency of the adapting pattern was 8 or 15 Hz (4 or 7.5 deg/s), and its location (left or right of fixation) and direction of motion (up/down) were randomized across sessions. The starting phases of the adapting and match gratings were randomized. The match pattern was presented on the opposite side of fixation to the adapting pattern, and its direction of movement was randomized from trial to trial to minimize the buildup of adaptation to the match pattern itself. The speed of the match pattern on any trial was controlled by an independent QUEST staircase.\(^{37}\) There were 20 independent QUEST staircases, one for every test interval. Each QUEST was initialized with a random starting point and concentrated observations near the point of subjective speed equality for each match interval. The observer’s task was to maintain steady fixation throughout each session and, following each match interval, to indicate whether the pattern to the left or the right of fixation had been moving faster. The observer was allowed a rest period of not less than 1 min between sessions before proceeding to the next adaptation and recovery session in the run. At least four but typically eight runs of 16 trials each were made for each condition, and all conditions were randomly interleaved.

B. Experiment 1: Apparent Speed Changes in Adaptation and Recovery

Figure 1 shows a typical psychometric function for the naive observer (RW) for speed matches of a 15-Hz standard at 21 s (out of 30 s) into the adaptation period; error bars show the binomial standard deviations. The binomial distribution is approximately normally distributed for large samples, and its standard deviation for each data point was estimated with standard methods\(^{38}\) at

\[
\left(0.5 (pq/n)^{0.5} + (1/2n),
\right)
\]

where \(p\) = proportion correct, \(r = 1 - \) proportion correct, and \(1/2n = \) correction for continuity.

The raw data have been fitted (solid curve) by a cumulative normal function by a least-squares fit; dashed curves show the upper and lower 95% confidence fits, estimated with standard methods.\(^{39}\) The speed was taken as the speed at which observers indicated that the adapting pattern was faster on 50% of trials. For each observer 20 similar psychometric functions were collected at 3-s intervals throughout each run: ten during adaptation and ten during recovery.

Figure 2 shows speed matches (in log Hz) at 3-s intervals throughout the run for two observers [PB, Fig. 2(a); RW, Fig. 2(b)] for two standard speeds, 15 Hz (squares) and 8 Hz (circles). Error bars show 95% confidence intervals. The data have been fitted by exponential functions by a least-squares fit, weighted by the 95% confidence intervals. The exponential functions have the following form:

Adaptation:

\[
(S_a) + (S - S_a) \exp(-t/t),
\]

Recovery:

\[
S - (S - S_a) \exp(-t/t),
\]

where \(S\) is standard speed, \(a\) is proportion of speed attenuation, \(t\) is time in seconds, and \(\tau\) is time constant of the exponential function for adaptation and recovery.

The adaptation functions are shown by the black curves, and the recovery functions are shown by the gray
curves. The results show that the apparent speed of the adapting pattern slowed exponentially during adaptation with a time constant of approximately 5–6 s (mean 5 s). Recovery was much slower, taking approximately 9–29 s (mean 22 s). The magnitude of attenuation was slightly greater at 15 Hz, where adapted apparent speed fell to 44% of unadapted apparent speed (mean across observers), than at 8 Hz, where adapted apparent speed fell to 57% of unadapted levels.

C. Experiment 2: Apparent Contrast Changes in Adaptation and Recovery
Several researchers have shown that the apparent contrast of temporally modulated gratings can decrease with adaptation, the magnitude and the rate of contrast attenuation depending on the spatial frequency, the temporal frequency, and the contrast of the adapting pattern. It is also known that the apparent speed of slowly drifting gratings depends on contrast, although the results are inconclusive at higher temporal frequencies. While some studies report that apparent speed increases with contrast for temporal frequencies up to 10 Hz for 2- and 3-c/deg gratings, others report that apparent speed is invariant of contrast at 8 Hz for 1-c/deg gratings and that the relationship reverses at 16 Hz for 1–8-c/deg gratings. It is therefore possible that a change in the apparent contrast of the adapting pattern could contribute to its change in apparent speed. In experiments 2 and 3, we measured the contribution of the change in apparent contrast to the change in apparent speed during adaptation. The change in apparent contrast during adaptation and recovery was measured in experiment 2.

Stimuli and procedure were as in experiment 1, except that the adapting and match speeds were equal and the observer was required to indicate the pattern of higher contrast. The temporal frequencies of the adapting and match gratings were equal (8 or 15 Hz) throughout the 30-s adaptation and recovery periods. The contrast of the adapting grating was fixed (50%), and the contrast of the match grating for each matching interval was varied according to an independent QUEST staircase. Figure 3 shows the change in apparent contrast during a 30-s adaptation period followed by a 30-s recovery period, and the results are presented in the same format as that of the speed-matching results shown in Fig. 2. Adaptation contrast matches are again fitted by black curves, and recovery contrast matches are fitted by gray curves with functions of the form shown in expressions (3) and (4). The results show that the apparent contrast of the adapting grating decreased exponentially during adaptation, to approximately 70% of the physical contrast (mean...
counterphase flickering gratings. Previous study of apparent contrast during adaptation for are in very good agreement with similar conditions in a adaptation and especially of recovery. The adaptation results there was greater variability in the time courses of adap-
tion with a time constant of between 13 and 39 s (mean = 25.3 s). Although there was little difference between the observers in the magnitude of contrast attenuation, there was greater variability in the time courses of adap-
tion and especially of recovery. The adaptation results are in very good agreement with similar conditions in a previous study of apparent contrast during adaptation for counterphase flickering gratings.42

D. Experiment 3: Speed Matching II: Perceived
Contrast Equated
The results of experiment 2 show the changes in apparent contrast for drifting gratings during 30 s of adaptation and recovery. In experiment 3 we repeated the speed-matching task of experiment 1 but simultaneously varied the physical contrast of the match grating so that their apparent contrasts were equal. The procedure was as in experiment 1, except that the contrast and the speed of the match grating were varied from trial to trial. The physical contrast of the adapting grating was fixed throughout all runs (at 50%, although its apparent contrast varied; see experiment 2). The physical contrast of the match grating was varied according to the best-fitting function (as shown in Fig. 3) that equated its apparent contrast to that of the adapting grating, and its speed was varied according to an independent QUEST staircase for each matching interval.

Figure 4 shows the change in apparent speed of a 15- and an 8-Hz drifting grating during a 30-s adaptation period followed by a 30-s recovery period for two observers [PB, Fig. 2(a); RW, Fig. 2(b)]. The results are presented in the same format as that of the speed-matching results shown in Fig. 2. For comparison, the best-fitting functions of the basic speed-matching task (physical contrast equated) are replotted from Fig. 2 as dashed curves. Adaptation contrast matches are again fitted by black curves, and recovery contrast matches are fitted by gray curves with functions of the form shown in expressions (3) and (4). The results show that the apparent speed of the adapting grating decreases exponentially during adapta-
tion with a time constant of approximately 4–5 s (mean = 4.2 s). Recovery is once again slower than adapta-
tion, with a time constant of between 9 and 24 s (mean = 17.3 s). The time constants of adaptation and recovery are similar to those recorded with physical contrast equated.

As in experiment 1, the magnitude of attenuation was slightly greater at 15 Hz than at 8 Hz. At 15 Hz, adapted apparent speed fell to 52% of unadapted apparent speed (compared with 44% in experiment 1), and so equating apparent contrast reduced the loss of apparent speed by 8%. At 15 Hz, adapted apparent speed fell to 72% of unadapted apparent speed (compared with 57% in experiment 1), and so equating apparent contrast reduced the loss of apparent speed by 15%. These results show that part of the loss of apparent speed is evidently attributable to a loss of apparent contrast, but there is a sizable speed loss that persists when this contrast-related speed change is eliminated.

E. Experiment 4: Speed Sensitivity in Adaptation and Recovery
The results of experiments 1–3 show that apparent speed decreases exponentially during adaptation and slowly returns to unadapted levels during recovery. In experiment 4 we measured changes in sensitivity to speed increments and decrements throughout the adaptation and recovery periods and compared Weber fractions for speed increments based on physical and apparent speed. Stimuli and procedure were similar to those of experiment 1, except that two adapting patterns were present throughout the adapting period and for 1 s every 3 s throughout the recovery period. As above, the patterns moved at 8 or 15 Hz and were 2 deg to the left and the right of the central fixation cross. The direction of motion of both patterns was fixed throughout each adapta-
tion and recovery period but was randomized across the periods. Observers reported that it was easier to main-
tain steady fixation when the patterns moved in opposite directions on either side of fixation, and so we incorpo-
rated this constraint. This spatial configuration († + † or † + †) gave the appearance of rotational or shearing motion, and it has been shown that rotational configura-
tions of motion can lead to elevated adaptation effects for
conventional MAE's. However, control data for the experienced observer (PB) in the present task were unaffected by this constraint.

There was a test interval every 3 s throughout a session. During the test interval, one of the patterns (at random) increased in speed, then returned to the baseline adapting speed (8 or 15 Hz). To avoid temporal artifacts arising from abrupt speed changes, we smoothed the speed change within a Gaussian envelope with a standard deviation of 32 frames (27 ms). The observer was notified that a test interval was in progress because the fixation cross was switched to white, then back to black again at the end of the test interval. This signaled the observer to respond (with a button press) whether the pattern on the left or the right of fixation had changed speed. The size of the speed increment on any trial was varied from trial to trial according to an independent QUEST procedure that concentrated observations near a threshold of 75% correct for each testing interval.

Figure 5 shows a typical psychometric function for the naive observer (RW) for speed increment detection for a 15-Hz adapting pattern at 21 s (out of 30 s) into the adaptation period. The raw data have been fitted (solid curve) by a cumulative normal function by a least \( \chi^2 \) fit, dashed curves show the upper and lower 95% confidence fits, and error bars show the binomial standard deviations. The speed increment threshold was taken as the speed at which observers identified the faster pattern on 75% of trials. For each observer 20 similar psychometric functions were collected at 3-s intervals throughout each run: ten during adaptation and ten during recovery.

Figure 6 shows speed increment thresholds at 3-s intervals throughout the run for two observers [PB, Fig. 6(a); RW, Fig. 6(b)] for two standard speeds, 15 and 8 Hz. Error bars show 95% confidence intervals. The data have been fitted by Weber fractions based on the apparent speed estimates for each observer, as shown in Fig. 2. The basic Weber fraction [expression (5): the lower threshold of motion, \( \sigma \), plus a proportion of speed, \( k_1 \)] provided an adequate fit to the data. However, superior fits were obtained with the addition of a second quadratic term [Eq. (6) below, where \( \sigma \) and \( k_1 \) are unchanged but a proportion of speed squared, \( k_2 \), was included]:

\[
\Delta v_{\text{thresh}} = \sigma + k_1 S_a,
\]

\[
\Delta v_{\text{thresh}} = \sigma + k_1 S_a + k_2 S_a^2,
\]

where \( S_a \) = apparent speed, \( k_1 \) = standard Weber fraction, \( k_2 \) = quadratic term, and \( \sigma \) = lower threshold of motion.

For further comparisons between these Weber fractions for speed increments, see Fig. 1 of Simpson et al. It can be seen that the data are well fitted by the predicted increment threshold from Weber's law based on apparent speed. Weber's law based on physical speed is a flat function that does not describe the data very well. For observer PB's speed increments, the best-fitting parameters for \( k_1 \), \( k_2 \), and \( \sigma \) were, respectively, 0.083, 0.012, and 0.039, and for observer RW they were 0.136 and 0.008 for \( k_1 \) and \( k_2 \), and \( \sigma \) tended to 0. These parameters are higher than estimates of Weber fractions for speed in unadapted conditions, but this is perhaps not surprising given that speed change was smoothed in a Gaussian envelope in the present task. In adapted conditions, detection of speed oscillations approaches 100% correct for 16%–20% modulations and are in broad agreement with our 75% thresholds of 8%–14% speed change.

![Fig. 5. Typical psychometric function for speed increment detection. The data are for the naive observer after 21 s of adaptation to a standard grating drifting at 15 Hz. The x axis shows the speed increment (\( \Delta v \)), and the y axis shows the proportion of observations in which the match appeared faster. Error bars show binomial standard deviations. The solid curve shows the best-fitting cumulative normal function, and the dashed curves show the upper and lower 95% confidence limits to the fit.](image)

![Fig. 6. Speed increment sensitivity as a function of adaptation and recovery duration for 8-Hz (circles) and 15-Hz (squares) adapting gratings for (a) one of the authors (PB) and (b) a naive observer (RW). Observers adapted to continuous motion for the first 30 s and then recovered for 30 s; speed increment sensitivity was measured at 3-s intervals throughout. Error bars show 95% confidence intervals. The data have been fitted by Weber fractions based on the apparent speed measured in experiment 2: black curves for adaptation and gray curves for recovery.](image)
Adaptation, a change in response to a sustained stimulus, can be demonstrated in motion perception by the illusory motion of a static pattern (the motion aftereffect (MAE)) and by changes in the apparent speed of moving patterns (the velocity aftereffect (VAE)). The magnitude and the time course of these aftereffects depend on the combination of adapting and test stimuli as well the method employed to measure them (for a review see Refs. 1 and 2).

Experiment 1 confirms the classic observation that the apparent contrast of a drifting pattern is reduced during adaptation. Adaptation in other species (5 s in cats, 1–2 s in flies) suggests that changes in apparent contrast can affect the match speed. In unadapted conditions the apparent speed of 8-Hz drifting gratings has been found to be invariant of contrast or to decrease at low contrasts. The reduction in the magnitude of speed attenuation at 8 Hz is consistent with the latter results. At 16 Hz, apparent speed can increase at low contrasts near detection threshold and when there is a 3–5-fold difference in the contrast of standard and match gratings. This suggests that lowering the contrast of the match pattern (to equate the apparent contrast of adapting and match gratings) should increase its apparent speed, and therefore it should match the apparent speed of the adapting pattern at an even lower physical speed. However, experiment 3 shows that lowering the contrast of the match pattern (to equate apparent contrast) reduced the match speed. The effect of contrast at 15 Hz was much less than at 8 Hz, in the direction expected from studies of apparent speed and contrast in unadapted conditions. Also, the contrast of our stimuli was much higher (50% for the adapting grating and a minimum of 33% for the match grating), and the contrast differences were much lower, than in speed studies with no adaptation to motion. In general, experiment 3 shows that when the apparent contrasts of the adapted and unadapted drifting patterns are equated, the loss of apparent speed was reduced (by approximately 15% at 8 Hz and 8% at 15 Hz). In a similar speed-matching study, Thompson varied the physical contrast of the match grating so that the apparent contrasts of the adapting and match gratings were equal on some trials. He also found a small reduction in the magnitude of the VAE that was at most approximately 10%. Taken together, these results suggest that only approximately 12% of the loss of apparent speed is dependent on a loss of apparent contrast.

C. Enhanced Speed Sensitivity during Adaptation to Motion

In a final experiment, we examined sensitivity to speed change for patterns whose apparent speed had been attenuated by adaptation. We found that speed increment and speed decrement thresholds decreased during adaptation and steadily returned to unadapted levels during recovery. This result confirms that sensitivity to changes in speed can increase during adaptation to course of changes in spike rates during adaptation to motion in other species (5 s in cats, 1–2 s in flies). The results are also consistent with physiological studies of feline neurons showing that recovery from adaptation is slower than adaptation, but the absolute values differ considerably (8 s in cats compared with 22 s in the present study).

B. Contribution of Contrast Attenuation to Speed Attenuation

Experiment 2 confirms that the apparent contrast of temporally modulated gratings rapidly decreases with adaptation (τ = 2 s), and here we report that recovery of normal contrast perception is also exponential but with a much slower time constant (approximately 25 s). It is well-known that the apparent speed of a grating can depend on its contrast, suggesting that changes in apparent contrast could affect the match speed. In unadapted conditions the apparent speed of 8-Hz drifting gratings has been found to be invariant of contrast or to decrease at low contrasts. The reduction in the magnitude of speed attenuation at 8 Hz is consistent with the latter results. At 16 Hz, apparent speed can increase at low contrasts near detection threshold and when there is a 3–5-fold difference in the contrast of standard and match gratings. This suggests that lowering the contrast of the match pattern (to equate the apparent contrast of adapting and match gratings) should increase its apparent speed, and therefore it should match the apparent speed of the adapting pattern at an even lower physical speed. However, experiment 3 shows that lowering the contrast of the match pattern (to equate apparent contrast) reduced the match speed. The effect of contrast at 15 Hz was much less than at 8 Hz, in the direction expected from studies of apparent speed and contrast in unadapted conditions. Also, the contrast of our stimuli was much higher (50% for the adapting grating and a minimum of 33% for the match grating), and the contrast differences were much lower, than in speed studies with no adaptation to motion. In general, experiment 3 shows that when the apparent contrasts of the adapted and unadapted drifting patterns are equated, the loss of apparent speed was reduced (by approximately 15% at 8 Hz and 8% at 15 Hz). In a similar speed-matching study, Thompson varied the physical contrast of the match grating so that the apparent contrasts of the adapting and match gratings were equal on some trials. He also found a small reduction in the magnitude of the VAE that was at most approximately 10%. Taken together, these results suggest that only approximately 12% of the loss of apparent speed is dependent on a loss of apparent contrast.
motion. As the physical speed of the stimulus did not change throughout the run (except for the small increments in the test intervals), Weber fractions for speed increment sensitivity based on the physical speed of the stimulus would be constant, but the data do not have this form. Together with the suggestions of an anonymous reviewer, we can offer three possible sources for the increase in speed sensitivity:

1. Sensitivity could be proportional to the stability of apparent speed. Thus, at the start of the adaptation period, apparent speed is varying more than it is later in the adaptation period, when the exponential function asymptotes and apparent speed is relatively stable. However, this possibility is unlikely because apparent speed is also relatively stable toward the end of recovery, but thresholds are higher in those conditions.

2. The Weber fraction is not constant throughout the run but varies with the level of adaptation.

3. A similar proposal to that of proposal 2 is that the Weber fraction is constant but the stimulus level is decreasing. Both these proposals mean that sensitivity to apparent speed effectively increases. We prefer this proposal because speed sensitivity can be compared with estimates of apparent speed determined in the speed-matching task, while it is difficult to test proposal 2. Experiment 4 shows that the speed increment and the decrement sensitivity calculated from the apparent speed in any test interval with a constant Weber fraction provide a very good fit to the data.

Several other groups have examined the effects of adaptation on increment sensitivity but for luminance and contrast. Adaptation mechanisms have been extensively studied for luminance coding that has a remarkable response range of approximately 8 orders of magnitude. Several adaptation mechanisms, including optical factors, photoreceptor nonlinearities, and synaptic adaptation, extend the luminance response range and help accommodate more information within a single cell. The net effect is a computationally advantageous removal of the background signal (for a discussion see Ref. 48).

Similarly, some studies of contrast increment sensitivity have shown that, under some conditions, contrast increment sensitivity can improve following adaptation. Greenlee and Heitger compared unadapted contrast increment thresholds for a 2-c/deg grating with thresholds following adaptation to a high-contrast grating (80%). They found that the slope of the contrast discrimination function was reduced by adaptation and that thresholds for high-contrast gratings (greater than 50%) were significantly lower following adaptation. The results were described by a leftward shift in the contrast response function and were taken as evidence that adaptation serves to linearize the contrast response function of a mechanism in the region near the prevailing contrast level. Wilson and Humanski measured contrast increment thresholds for a D6 stimulus (the sixth derivative of a Gaussian in x multiplied by a Gaussian in y) centered at 3 or 9 c/deg following adaptation to a high-contrast grating (99%). Adaptation reduced the slope of the contrast discrimination function. In some cases the adapted contrast discrimination function crossed the unadapted function at high pedestal contrasts (60%), indicating an improvement in contrast increment sensitivity. However, other researchers have been unable to replicate these findings. Ross et al. measured contrast increment sensitivity for a 2-c/deg grating following adaptation to a 40% contrast grating, and both gratings counterphase flickered at 8.8 Hz. They found that while adaptation and masking raised thresholds at low contrasts, they had no effect at higher contrasts. The failure to find a reduction in thresholds at higher contrasts was attributed to the lower contrast of the adapting pattern (40%) and the lower mean luminance of the display (15 cd/m² compared with 80–100 cd/m²). For 5-c/deg gratings Maatannen and Koenderink confirmed that contrast detection thresholds and apparent contrast were reduced but found that contrast discrimination thresholds remained unchanged following adaptation to a 50% contrast grating. Their failure to replicate Greenlee and Heitger’s findings was attributable to the lower stimulus contrasts—while the physical contrasts of the stimuli were comparable (75% and 80%), the spatial Gaussian windowing employed by Maatannen and Koenderink increased detection thresholds and therefore reduced its multiple of contrast threshold.

With these reservations in mind, it is tempting to conclude that adaptation to motion serves to increase sensitivity to the prevailing speed, as has been suggested for luminance adaptation and in some studies of contrast adaptation.

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