Medial Superior Temporal Area Neurons Respond to Speed Patterns in Optic Flow

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Received Nov. 12, 1996; revised Jan. 17, 1997; accepted Jan. 28, 1997.

Optic flow fields are the global patterns of visual motion generated as an observer moves through the environment (Gibson, 1950). Neurons in the dorsal region of the medial superior temporal area (MSTd) of monkey extrastriate cortex have characteristics suggesting that they might contribute to the analysis of optic flow. Their receptive fields typically cover a quadrant of the visual field, providing access to the global visual motion created by observer movement (Tanaka et al., 1986; Komatsu and Wurtz, 1988). They respond to the planar, radial, and circular patterns, which are the components of optic flow (Saito et al., 1986; Sakata et al., 1986; Tanaka and Saito, 1989; Tanaka et al., 1989; Andersen et al., 1990; Wurtz et al., 1990; Duffy and Wurtz, 1991a,b; Orban et al., 1992; Graziano et al., 1994; Lagae et al., 1994), and many change their responses when the center of motion in optic flow is shifted in the visual field to simulate different headings of observer movement (Duffy and Wurtz, 1995). Finally, some MSTd neurons compensate for the effect of pursuit eye movements that accompany observer movement through the environment (Bradley et al., 1996).

Speed of motion also varies systematically in optic flow stimuli. As an observer moves forward, the pattern of radial expansion typically contains slower motion in the center and faster motion at the edge. Although previous studies have characterized the MSTd neuronal responses to patterns of motion direction, relatively little is known about their responses to patterns of motion speed. Tanaka et al. (1989) showed that withdrawing the speed gradient reduced the response of 34 neurons responding to expanding stimuli by ~20%. Duffy and Wurtz (1991a) studied 16 neurons and found little effect of speed in all but 3. Orban et al. (1995) found that the speed tuning of 14 neurons could be regarded as being bandpass and that removing the pattern of speeds had little effect. Because speed patterns in optic flow can provide important cues about observer movement (Gibson, 1966; Rogers and Graham, 1979; Cutting et al., 1992), the insensitivity to speed patterns by MSTd neurons would represent an important exception to their suitability for optic flow field analysis.

In light of this relatively limited information on the effects of speed on MSTd neurons, and the importance of speed to optic flow, we studied preferred speeds and the effect of altering speed gradients on a larger sample of MSTd neurons that responded to either expanding or rotating optic flow stimuli. In a sample of 131 neurons, we found that different neurons were tuned to respond to different bands of stimulus speeds, and that the structure of the speed gradient was an important stimulus characteristic in many of these neurons. We believe that this sensitivity of MSTd neurons to patterns of speed, as well as patterns of direction, strengthens the view that these neurons are well suited to contribute to optic flow field analysis.

MATERIALS AND METHODS

Behavioral and neurophysiological techniques. The procedures used in these studies are identical to those reported recently (Duffy and Wurtz, 1995) and are described here only briefly. All protocols were approved by the Institute Animal Care and Use Committee and complied with Public Health Service policy on the humane care and use of laboratory animals. We recorded the activity of single cortical neurons in two adult rhesus monkeys.
The visual motion stimuli were then projected onto the screen in a pseudorandom sequence, with stimulus durations of 1 sec and interstimulus intervals of 1–1.5 sec.

The activity of single neurons was digitized using a window discriminator and stored with stimulus and behavioral event markers using the REX system (Hays et al., 1982). We recorded neuronal activity using epoxy-coated tungsten microelectrodes that were advanced with a hydraulic microdrive. Neural activity was monitored to locate the depth of physiological landmarks, and studies were initiated whenever neuronal discharges were clearly isolated.

Neuronal response amplitudes were measured as the mean neuronal discharge rate evoked by six repetitions of the 1 sec presentation of each visual motion stimulus. The responses to the visual motion stimuli were compared with activity in the same period of control trials. These control trials were pseudorandomly interleaved with the visual motion trials and consisted of visual fixation without a visual motion stimulus. Differences between stimulated and control trials were tested for statistical significance using Student’s t test ($p < 0.01$).

At the end of the experiment, electrolytic marking lesions were made along penetration tracks in three guide tubes in each hemisphere. These marks were identified in histological sections 50 μm thick, with every fourth and fifth section stained with Nissl and Gallyas methods. Drawings were made of the sections to locate the electrolytic lesions relative to anatomic landmarks, to extrapolate the position of the recording sites. These drawings indicate that at least 90% of the neurons studied were in the densely myelinated zone on the anterior bank of the superior temporal sulcus that is included in the MSTd (Komatsu and Wurtz, 1988). The remaining neurons were farther down the anterior bank closer to the lateral region of MST (MSTl), but had the same physiological characteristics.

**Visual stimuli.** The visual stimuli consisted of 360 white dots on a dark background, randomly distributed at onset, and then moving in the specified pattern. All stimuli covered the 100° × 100° tangent screen with the center of expansion–contraction for radial stimuli, and the center of rotation for circular stimuli, positioned over the fixation point at the center of the screen. Thirteen stimuli were used for preliminary classification of each neuron’s response; these included: eight directions of planar translation (uniform motion in one of eight directions at 45° intervals), two directions of radial motion (inward or outward), two directions of circular motion (clockwise or counterclockwise), and stationary dots. The radial or circular stimulus that evoked the strongest response was identified, without regard to the presence of planar translation responses, and that motion pattern was used in the subsequent speed studies.

In the radial and circular stimuli, dot speed increased with distance from the center of the pattern to create what we term normal speed gradient stimuli. The radial algorithm moved dots inward to, or outward from, the fixation point at the center of the screen, with dot speed increasing as a function of $\sin(\theta) \cdot \cos(\theta)$, where $\theta$ is the viewing angle from the center of the stimulus to a given dot (Fig. 1). This generated a simulation of translational motion away from, or toward, a frontoparallel plane. The circular algorithm moved dots clockwise or counterclockwise around the fixation point at the center of the screen, with dot speed increasing as a tangent function of distance from the center. This generated a simulation of rotational motion with reference to a frontoparallel plane.

In the normal gradient stimulus set, the radial and circular stimuli had a mean speed of 40°/sec (Fig. 1B, bold line). This mean speed refers to the speed of the dots at the halfway point between the center and edge of the stimulus. This value was chosen because it yielded subjective comparability of the overall speed in radial, circular, and planar stimuli. Because the dots were evenly distributed, and there is more area beyond the halfway point, the numerical average of dot speeds was above this value. Lower and upper limits were imposed on the dot speeds to minimize apparent stationarity of slow dots (e.g., near the center of outward radial patterns) and the streaking of fast dots (e.g., near the edges of outward radial patterns).

To test for effects of the normal speed gradient, we created nongradient stimuli lacking the dependency of dot speed on distance from the center of the stimulus, maintaining a uniform speed for all dots in a stimulus (Fig. 4A). These uniform speeds matched the mean speeds in the normal gradient speed stimuli. To explore more fully the effects of speed gradients, we applied a scaling factor to the function relating dot speed to distance from the center of the stimulus (Fig. 6). Negative scaling factors (−1.0, −1.5, and −2.0) created inverted gradients in which monumental responses were then projected onto the screen in a pseudorandom sequence, with stimulus durations of 1 sec and interstimulus intervals of 1.0–1.5 sec.

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dot speeds decreased with increasing distance from the center of the stimulus (see Fig. 6A). Positive scaling factors created normal (1.0) or exaggerated (1.5 and 2.0) gradients in which dot speeds increased with increasing distance from the center of the stimulus (see Fig. 6B). (The nongradient stimuli were implemented using a scaling factor of zero.) This approach was chosen so that all of the local dot speeds were changed while the mean speed (at the halfway point between the center and the edge) remained as a constant, pivotal value.

We also made stimuli to explore the basis of speed gradient preferences by testing local speed tuning in different parts of the stimulus area. In creating stimuli that covered only the central 50° of the screen or the area just beyond it, we imposed a software mask over the appropriate part of the nongradient speed stimuli (Fig. 8). Thus, the total number of dots differed depending on which region was presented, but dot density within the stimulated region was maintained.

To test for effects of simultaneously presented speeds, each covering the entire stimulus area, we used the transparent superimposition of a number of normal gradient speed stimuli. This created multiple speed planes with motion parallax effects generated by the motion of each plane relative to that of the other planes (Fig. 12). The number of dots in each plane was adjusted so that each of the planes had equal numbers of dots, and the total number of dots remained 360.

RESULTS

We studied 131 MSTd neurons, first determining the responses of each cell to the visual motion components of optic flow. These component stimuli consisted of 12 patterns covering the central 100° of the visual field: eight planar motion stimuli (eight directions at 45° intervals around 360°), two radial motion stimuli (inward and outward from the fixation point), and two circular motion stimuli (counterclockwise and clockwise around the fixation point). All of these neurons gave significant responses to radial and/or circular motion, with some also responding to planar motion or to radial, circular, and planar motion. Subsequent studies were conducted using the preferred radial or circular stimulus for that neuron.

Effects of mean speed in normal gradient stimuli

We varied the mean speed of radial or circular motion using the same five mean speeds between 10 and 80°/sec for all neurons. In all of these stimuli, we maintained the normal gradient of speed profiles (slower motion in the center) as shown for the example of outward radial motion in Figure 1. More than two-thirds of the neurons that showed some responses to these stimuli (68%, or 83 of 122 neurons) showed strong speed preferences; the amplitude of their response to one speed was at least twice that to another speed. The curves relating response amplitude to mean speed (Fig. 2) showed a variety of shapes that we placed into five
categories, which resemble five simple filters. If no response was significantly greater than any other, the neuron was classified as having a flat response profile (broad band; Fig. 2A). If the largest response was at one end of the curve and the smallest response was at the other end, the neuron was classified as having either an increasing or decreasing response profile (low-pass or high-pass; Fig. 2B,C). If the intermediate mean speeds evoked the smallest or largest response, the neuron was classified as having a trough or peak response profile, respectively (bandpass or band-reject; Fig. 2D,E). These response profile classifications do not bear a one-to-one correspondence with classification by optimal stimulus speed (e.g., neurons preferring the fastest speed might show a high-pass or band-reject profile).

Across the sample, speed had similar effects on the responses to both radial and circular stimuli, and Figure 3A shows the frequency of the five response classes shown in Figure 2. The most common speed response profile is that showing increasing response with increasing mean speed (42% of radial neurons and 38% of circular neurons). The frequency of optimal speed preferences is shown in Figure 3B, with the most commonly preferred speeds being the fastest and slowest (64% of radial neurons and 66% of circular neurons). Although we usually tested neurons

Figure 3. Comparison of speed preferences to radial (left column) and circular (right column) stimuli. A, Percentage of neurons tested with radial and circular stimuli that had each of the five varieties of speed response profiles. A total of 122 neurons showed some response to these stimuli; the 114 neurons that showed at least one statistically significant response were classified into one of five groups. In 13% (11 of 82) of neurons studied with radial stimuli and in 6% (2 of 32) of neurons studied with circular stimuli, no response was statistically significantly greater than any other response; those neurons were considered to have a flat response profile (first bar). In 61% (50 of 82) of neurons studied with radial stimuli and in 66% (21 of 32) of neurons studied with circular stimuli, the strongest response was at one end of the speed range, and the weakest response at the other end, with more neurons in each group preferring faster speeds. The remaining 26% (21 of 82) of neurons studied with radial stimuli and 28% (9 of 32) of neurons studied with circular stimuli either had the smallest response at one end and the peak response at an intermediate speed, or the peak response at one end and the smallest response at an intermediate speed. The classes of response profiles occurred with equal frequency for radial and circular stimuli. B, Percentage of neurons tested with radial and circular stimuli that showed the largest amplitude response at each of the stimulus speeds. The optimal speed (abscissa) was determined by averaging the responses to six stimulus presentations and selecting the speed that evoked the largest average response. A total of 122 neurons were tested with these stimuli, 93% (114 of 122) of which showed statistically significant responses to at least one stimulus. In both the radial and circular groups, the slowest and fastest speeds more commonly evoked the strongest responses, but in both groups there were substantial numbers of neurons preferring each of the speeds. C, Example of a neuron that responded to both radial and circular optic flow stimuli showing the similar preference for slower stimulus speeds for both stimuli. Same format for the graphs as in Figure 2.
only to their preferred radial or circular stimuli, those that responded to both had comparable speed profiles to both stimulus patterns. Figure 3C shows the speed profiles of such a neuron that showed preferences for slower stimulus speeds for both radial (left) and circular (right) motion.

Thus, we have made the following observations: (1) the responses of more than two-thirds of the MSTd neurons are strongly modulated by changes in the mean speed of the stimulus gradient; (2) this modulation is equally present for neurons preferring

Figure 4. The effects of removing the speed gradient in optic flow stimuli as seen in three neurons that show the range of relationships between responses to normal gradient (left) and nongradient (right) stimuli. A. Stimulus speed profiles showing the relationship between dot speed and location varying as a sine × cosine function in the normal gradient radial stimuli (left) and as a constant speed in nongradient stimuli (right). B, A neuron that showed similar responses across stimulus speeds, regardless of whether the stimuli were the normal gradient speed stimuli (left) or the nongradient speed stimuli (right). C, A neuron that showed different responses at the slowest stimulus speeds, depending on the speed gradient. The 10°/sec stimulus evoked the strongest response with normal gradient stimuli (left) and the weakest response with nongradient stimuli (right). D, A neuron that showed entirely different response profiles depending on the speed gradient. This neuron showed a preference for fast speeds with normal gradient stimuli (left), and a preference for slow speeds with nongradient stimuli (right).

Figure 5. Comparison of responses to normal gradient and nongradient stimuli in the sample of MSTd neurons. A. The percentage of neurons that showed their largest amplitude responses to the indicated speed of nongradient stimuli (open bars) and normal gradient stimuli (solid bars). The graph combines the results for radial and circular motion in Figure 3B for the gradient bars and uses the same format as Figure 3B. Removing the gradient had little effect on the overall preference for the fastest and slowest stimulus patterns. B, Percentage of neurons (ordinate) showing statistically significant differences between normal gradient and nongradient stimuli for the number of speeds indicated on the abscissa. Approximately one-third (36%, or 38 of 105) show no significant differences, almost half (46%, or 48 of 105) show one or two significant differences, and 18% (19 of 105) show three or more differences, usually at one end of the speed range. C, Percentage of neurons showing response magnitude differences expressed as a ratio between normal and nongradient stimuli. The ratio for each cell is at the speed yielding the largest ratio for that neuron. The sample is about evenly divided between those that prefer normal gradients (37%, or 39 of 105; filled bars), those without strong preferences (31%, or 32 of 105; open bar), and those that prefer nongradients (32%, or 34 of 105; shaded bars).
Effects of speed gradients in optic flow stimuli

In normal gradient stimuli, speed varies as a function of viewing angle from fixation at the center of the stimulus to a given dot: a sine \( \times \) cosine function for radial stimuli and a tangent function for circular stimuli. To determine whether these speed gradients influence MSTd responses, we created radial and circular stimuli without the normal gradient; i.e., stimuli with the same speed of dot motion throughout the stimulus (Fig. 4A). We see these stimuli with and without the speed gradient as looking alike, but with a clear sense that there are speed differences between them.

We compared the response of 105 neurons to normal gradient and nongradient stimuli at five mean speeds. Overall, the nongradient stimuli evoked only slightly less responsiveness than did the gradient stimuli; approximately two-thirds still showed a response that was at least twice the amplitude of the weakest response (62\%, or 58 of 94 for the nongradient stimuli compared with 68\% for the gradient stimuli). Figure 4B shows an example of a neuron that responded similarly to the normal gradient (left) and nongradient (right) stimuli. Nevertheless, some neurons showed substantial differences between their responses to normal gradient and nongradient stimuli. For example, the neuron in Figure 4C shows a decrease in the response to the lowest speed when the stimulus had no gradient, and the neuron in Figure 4D shows a different profile of responses, with a decrease at the fastest speed and an increase at the slowest speed for a nongradient stimulus.

Figure 5 shows the extent of these response differences between normal gradient and nongradient stimuli for our sample of neurons. The preference for faster and (to a lesser extent) slower speeds seen for gradient stimuli is preserved for nongradient stimuli (Fig. 5A). When the number of speeds at which the response differs for the normal gradient and the nongradient stimuli are compared (Fig. 5B), we find that almost two-thirds of the neurons (64\%, or 67 of 105) show significant differences in their responses at least one of the five speeds. Figure 5C shows the magnitude of the response differences between gradient and nongradient stimuli. Two-thirds of the neurons (69\%, or 73 of 105) had speeds at which the larger of the responses to either stimulus was more than one and a half times the amplitude of the smaller response. Thus, in two-thirds of the neurons, the presence of a speed gradient produced substantial changes in neuronal responses.

Because eliminating the speed gradient alters the responses of many MSTd neurons, we next determined whether maintaining the gradient but varying its shape also would alter the responses. We used the same mean speed, but for negative speed gradients, the speed decreased with increasing distance from the center of the stimulus (inverted gradients; Fig. 6A), whereas for positive gradients, the speed increased with increasing distance from the center (normal or exaggerated gradients; Fig. 6B). The speed gradients were altered by multiplying the effect of distance from the center by a value from \(-2\) to \(+2\), creating seven speed gradient stimuli covering a segment of the range of naturalistic speed gradients. Figure 6, C and D, shows the responses of two neurons that show the most common response profiles observed. The neuron in Figure 6C shows strong responses to the positive speed gradients, in contrast to the neuron in Figure 6D, which shows a clear preference for negative speed gradients.

In 122 MSTd neurons, we found a fairly uniform preference for negative and positive speed gradients (Fig. 7A). All seven gradi-
Taken together, these results show that approximately two-thirds of MSTd neurons show substantial effects of eliminating speed gradients and that almost 9 of 10 neurons prefer a positive or inverted speed gradient to stimuli without a speed gradient. This indicates that the spatial distribution of speed across these radial and circular optic flow stimuli can have substantial effects on many MSTd neurons.

Potential explanations of gradient preferences

Speed gradient preferences could result from local differences in speed tuning profiles, such as a preference for slow motion in the center of the stimulus and fast motion in the periphery. To test this hypothesis, we presented stimuli separately to the central 50° segment and the peripheral segment outside that area, dividing the stimuli approximately at the point at which local speeds are the same in all of the speed gradients (the pivotal points in Fig. 6A,B). Figure 8 illustrates the results of such experiments in two neurons having distinctly different speed gradient preferences. Both neurons showed increasing responses with increasing speeds in nongradient stimuli covering the full stimulus area (Fig. 8A) and continued to show this same increase whether the stimulus was limited to the central segment of the field (Fig. 8B) or to the peripheral segment (Fig. 8C). Thus, these neurons show no evidence of the kind of substantial differences in central and peripheral speed tuning that would seem necessary to account for the normal speed gradient preferences. This point is reinforced by noting that these neurons had very different speed gradient preferences; neuron 26KL22 showed a strong preference for negative speed gradients (Fig. 8D, left), whereas neuron 26KR5 showed a strong preference for positive speed gradients (Fig. 8D, right).

None of the 48 neurons tested had substantially different speed profiles in the central and peripheral segments. However, we saw substantial differences in overall response amplitude with six neurons showing no significant responses to stimulation of the peripheral segment. The scatter plot in Figure 9 shows the responses of the remaining 42 neurons as the slope from a least squares fit to the response profiles evoked by stimuli in the central (abscissa) and peripheral (ordinate) stimulus segments. Although this measure is insensitive to the details of a few of the response profiles included, it demonstrates the comparability of the responses evoked from the central and peripheral segments (the regression line for the sample has a slope of 0.78; r = 0.77). Thus, we see no evidence of a spatial segregation of speed preferences (e.g., preferring slow motion in the center and faster motion in the periphery) as the basis of speed gradient preferences in MSTd neurons.

One factor that might contribute to speed gradient preferences is interactions between responses that are simultaneously evoked from stimuli in different parts of the stimulus area, such as those between the central and peripheral segments. To test this hypothesis, we presented the five speeds of central motion either with no speed gradient or inverted speed gradient to stimuli without a speed gradient. Figure 10 shows the results of such studies in two neurons with nongradient full-field (Fig. 10A), central (Fig. 10B), and peripheral (Fig. 10C) responses, as well as responses to the combination of slow motion (10°/sec) in the periphery with the five speeds tested in the center (Fig. 10D). The middle column shows the responses of a neuron that preferred positive gradient stimuli and that had similar preferences for higher speeds in both central and peripheral stimuli but gave no responses when the periphery contained slow motion, even though the central segment con-
tained the otherwise preferred fast motion. In contrast, the neuron in the right column preferred negative gradient stimuli and had similar preferences for faster speeds in both central and peripheral stimuli, but this neuron’s strongest responses were recorded when the periphery contained slow motion and the central segment contained fast motion. The peripheral stimulus, therefore, clearly can alter the response to different speeds of motion in the central segment.

To examine such effects more fully, we studied 44 neurons with speed testing in the central segment at our usual five speeds, while either slow motion (as in Fig. 9D) or fast motion was presented in the periphery. The results from the sample are summarized in Figure 8.
We tested 22 neurons with stimuli containing multiple speed planes and summarized the magnitude of response variation as contrast ratios across multiple-plane stimuli (Fig. 12C). A total of 41% (9 of 22) of the neurons tested showed clear effects of the number of speed planes, with multiple-plane stimuli evoking responses with contrast ratios of more than 0.3 (Fig. 12C, right bar). In the sample, as many neurons preferred decreasing numbers of planes as preferred increasing numbers of planes. Thus, when a number of speeds are presented simultaneously in the same area, MSTd neurons do not respond merely to some preferred speed. Rather, their responses reflect, at least to some degree, the variety of different speeds presented.

The effects of the number of speed planes might rely on the particular speeds in the multiple speed plane stimuli or on the magnitude of the speed difference between those speeds. Figure 13 compares the response to the speed of a single speed plane (A) with the response to the differences between two simultaneously presented speed planes (B) for the same neuron shown in Figure 12B. Figure 13A shows the responses to normal gradient stimuli with different mean speeds, whereas Figure 13B shows the profiles of the same neuron to the difference between the speeds of two transparently overlapping planes. This neuron gave approximately the same response to all but the slowest single plane stimulus (Fig. 13A), but it showed consistently decreasing responses as speed differences increased between the two-plane stimuli (Fig. 13B). The responses to the difference in the mean speed of two-plane stimuli are not equivalent to the averaged response to those mean speeds in single-plane stimuli. However, the decline in response amplitude with increasing difference between speeds in two-plane stimuli closely resembles the profiles for multiple-plane responses in that neuron (Fig. 12B). This similarity is consistent with a response of this neuron to the relative motion within the optic flow stimulus.

We found that neurons that show sensitivity to the relative motion with increasing numbers of planes (Fig. 12B) showed similar profiles of relative motion sensitivity to the speed difference between two superimposed planes of motion (Fig. 13B). Such similarities were evident in seven of the nine neurons that had contrast ratios greater than 0.3 for multiple-plane responses (Fig. 12C). Thus, the effects of multiple speed planes can be mimicked by presenting two planes having the same overall differences in speed that are presented in the multiple speed planes stimuli. This does not appear to reflect preferences for a particular speed as much as a preference for a combination of speeds, a preference that might be related to interactions between simultaneously presented speeds.

DISCUSSION

Speed preferences of MSTd neurons

We first examined the sensitivity of MSTd neurons to the pattern of speed in optic flow stimuli by varying the mean speed within the stimuli from 10 to 80°/sec. The radial stimuli would approximate the visual experience of an observer (at our viewing distance of 50 cm from the stimulus) moving forward at speeds from 0.2 to 1.5 m/sec, a range that is included in naturalistic experience. More than two-thirds of the MSTd neurons were strongly modulated by changes in the mean speed of the stimulus, and this modulation was similar in neurons preferring radial stimuli and circular stimuli. The shape of the response profile to stimulus speeds varied across the sample of neurons, but they could be placed into five classes resembling simple filter characteristics. The most common response profile showed increasing response amplitude with increasing mean speed. These findings establish the effect of stimulus speed on the responses of many MSTd neurons, just as previous experiments established the effects of the direction of stimulus motion.

The range of MSTd speed preferences we observed is consistent with those demonstrated by Orban et al. (1995). However, they reported optimal responses mainly in the range of 15–20°/sec, whereas our results (Fig. 3B) suggest that a substantial number of MST neurons prefer faster speeds. It is worth noting that Maunsell and Van Essen (1983) found that neurons in the middle
temporal area (MT) also show a broad range of speed preferences, with some neurons preferring slower speeds (10–50°/sec) and others preferring faster speeds (>100°/sec). This conclusion is supported by the broad range of speed preferences demonstrated by Cheng et al. (1994) in MT and V4 neurons.

Our MSTd speed profiles are similar to those reported for striate cortex by Orban et al. (1981), who described speed profiles by analogy to filter characteristics, specifically low-pass, high-pass, and bandpass filters. However, MSTd neurons show an additional profile that can be termed band-reject (Figs. 2E, 3A), in an extension of the analogy to filters. The number of these neurons is small (11% of the total), but their presence might be taken as completing the spectrum of simple filters and might be viewed as supporting a filter model of these responses. The low-pass and high-pass filters provide orthogonal representations of stimulus speed that might interact to create other response characteristics.

Figure 10. Alteration of responses of two neurons to central segment stimulation by simultaneous stimulation of the peripheral segment. Schematic diagrams of the stimuli are shown on the left, and the response profiles of two neurons are shown in the middle and on the right with speed (abscissa) plotted against average response amplitude (ordinate). A, Nongradient speed stimuli containing dot motion at a uniform speed over the central 100° × 100° of the visual field. Both neurons show increasing response amplitude with faster stimuli. B, C, Stimuli containing dot motion within the central 50° × 50° of the stimulus, or motion outside the central 50° × 50°. Both neurons show increasing response amplitude with faster speeds in either the central or the peripheral stimulus segment. D, Stimuli containing slow (10°/sec) dot motion outside the central 50° × 50° of the stimulus, and five different speeds within the central 50° × 50° of the stimulus. The neuron in the middle column shows no significant responses to these stimuli, even though central stimulation alone evoked strong responses. The neuron in the right column shows its strongest responses to the combination of fast stimuli in the center and slow motion in the periphery.
such as the more selective bandpass and band-reject filters. The band-reject filters are also noteworthy because they might play a role in refining MSTd responses, since a complete set of simple filters provides a greater potential for focusing bandwidth to complex stimuli. The range of MSTd response profiles is also consistent with the observation that response profile shape varies along a continuum in MT (Maunsell and Van Essen, 1983).

The most salient point is that many MSTd neurons are sensitive to the mean speed of the optic flow stimulus, which is in contrast to the impression given by several previous studies of MSTd, including our own (Duffy and Wurtz, 1991a), based on much smaller samples of neurons.

**Effects of speed gradients**

To assess the effects of speed gradients, we created stimuli with the same radial and circular patterns of motion direction, but with no speed gradient (Fig. 4), inverted speed gradients (faster motion in the center than in the periphery; Fig. 6 A), or exaggerated gradients (much slower motion in the center; Fig. 6 B). Approximately two-thirds of the MSTd neurons showed substantial effects of eliminating speed gradients, and almost 9 of 10 neurons preferred a positive gradient or inverted speed gradient to stimuli having no speed gradient. This indicates that the spatial distribution of speed across optic flow stimuli has substantial effects on many MSTd neurons and could contribute to the role these neurons play in the analysis of optic flow.

This conclusion differs from those of previous studies in ways that may be accounted for by differences in the experiments and in the size of the sample of neurons. In the first study of speed gradient effects on MSTd neurons (Tanaka et al., 1989), one of eight radial segments in an expansion/contraction stimulus was modified to eliminate the local speed gradient, and little effect was observed. We were able to present a series of gradient stimuli that cover more of the natural range of speed gradients in optic flow, and this may have revealed effects that otherwise would not be apparent. In addition, our sample contained some MSTd neurons with relatively little sensitivity to speed, raising the possibility that studies with smaller sample sizes might have included such neurons.

We altered speed in optic flow stimuli to determine whether MSTd neurons might use this parameter as a cue for self-movement perception. However, these stimuli might also be interpreted as simulating the movement of large objects at various

Figure 12. Sensitivity to transparently superimposed optic flow patterns of different average speeds. A, Schematic diagram of three radial patterns with coextensive, transparently superimposed speed gradients, each having increasing speed with increasing distance from the center of the stimulus. B, Responses of an MSTd neuron to multiple speed planes plotted as the number of speed planes (abscissa) versus the mean response amplitude (ordinate). Speeds of each plane are shown in parentheses. Response amplitude decreased with increasing numbers of superimposed speed planes. C, Bar graph showing the percentage of neurons that showed the indicated strength of response preferences for the five multiple-plane stimuli. More than one-third (36%, or 8 of 22) of the neurons showed substantial effects of the number of stimulus planes.
Neurons. To test this hypothesis, we compared the speed profiles of peripheral stimulus segments might shape the responses of these neurons. Our experiments provide some clues about motion interactions that are in the central visual field (i.e., the house in Fig. 14 A–D). We found that 41% of the neurons tested with these stimuli, there were substantial differences between responses to stimuli presented in the central segment, depending on the speed of motion in the periphery (Figs. 10, 11). Such interactions could contribute to speed gradient preferences by altering responses to particular combinations of central and peripheral speeds.

Because these experiments suggested that speed gradient preferences might rely on interactions between simultaneously presented speed stimuli, we wanted to determine whether those interactions required some particular spatial structure of the stimulus (e.g., slow motion in the center and fast motion in the periphery). To test this possibility, we presented different speeds together as transparently overlapping speed planes each covering the entire stimulus area (Fig. 12A). We found that 41% of the neurons showed substantial changes in response amplitude as the number of speed planes increased from one to five (Fig. 12B,C), with similar effects induced by increasing the magnitude of the speed difference between two overlapping speed planes (Fig. 13). Thus, there is evidence for speed interactions even in the absence of the spatial separation of speeds in the stimulus.

The perceptual utility of neuronal sensitivity to the overlap of multiple speed planes might relate to the motion parallax in these stimuli. Figure 14C illustrates how a visual scene with substantial depth of field (i.e., the trees at various distances from the approach to the house) presents spatially overlapping features at different speeds. Motion parallax can reflect the three-dimensional structure of the environment (Gibson, 1966) and support detection of heading direction during self-movement (Rogers and Graham, 1979; Cutting et al., 1992), possibly relating the speed and direction preferences of MSTd neurons. Our experiments provide some clues about motion interactions that might underlie speed gradient and speed overlap preferences in MST neurons. However, further studies in which these prefer-
an observer moving toward the center of the scene would encounter positive (A), negative (B), and overlapping (C) speed gradients. A. When nearby features of the scene are in the peripheral visual field (the trees lining the approach to the house), and increasingly remote features are closer to the center of the visual field (the house), the observer would encounter a positive speed gradient. As indicated by the length of the arrows, the speed of feature motion would increase from the center to the periphery. B. When nearby features of the scene are in the central visual field (the house) and remote features are in the peripheral visual field (the trees), the observer would encounter a negative speed gradient. As indicated by the length of the arrows, the speed of feature motion would decrease from the center to the periphery. C. When remote features are in the central visual field (the house), and the peripheral visual field contains features at substantially varying distances (the trees), the observer sees overlapping positive speed gradients. As indicated by the length of the arrows, there would be a general increase in speed of feature motion from the center to the periphery. In addition, the overlapping gradients would create motion parallax in the scene, as indicated by the decreasing length of arrows over trees positioned at increasing distances from the observer.

Figure 14. Schematic illustrations of naturalistic circumstances in which an observer moving toward the center of the scene would encounter positive (A), negative (B), and overlapping (C) speed gradients. A. When nearby features of the scene are in the peripheral visual field (the trees lining the approach to the house), and increasingly remote features are closer to the center of the visual field (the house), the observer would encounter a positive speed gradient. As indicated by the length of the arrows, the speed of feature motion would increase from the center to the periphery. B. When nearby features of the scene are in the central visual field (the house) and remote features are in the peripheral visual field (the trees), the observer would encounter a negative speed gradient. As indicated by the length of the arrows, the speed of feature motion would decrease from the center to the periphery. C. When remote features are in the central visual field (the house), and the peripheral visual field contains features at substantially varying distances (the trees), the observer sees overlapping positive speed gradients. As indicated by the length of the arrows, there would be a general increase in speed of feature motion from the center to the periphery. In addition, the overlapping gradients would create motion parallax in the scene, as indicated by the decreasing length of arrows over trees positioned at increasing distances from the observer.

References and interaction effects are tested together in a substantial number of neurons would be required to characterize the underlying mechanisms.

REFERENCES


