Response of Monkey MST Neurons to Optic Flow Stimuli with Shifted Centers of Motion

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Neurons in the dorsal region of the medial superior temporal area (MSTd) have previously been shown to respond to the expanding radial motion that occurs as an observer moves through the environment. In previous experiments, MSTd neurons were tested with radial and circular motion centered in the visual field. However, different directions of observer motion, relative to the direction of gaze, are accompanied by visual motion centered at different locations in the visual field. The present experiments investigated whether neurons that respond to radial and circular motion might respond differently when the center of motion was shifted to different regions of the visual field. About 90% of the 245 neurons studied responded differently when the center of motion was shifted away from the center of the field. The centers of motion preferred by each neuron were represented in the sample, with greater numbers of neurons preferring centers of motion closer to the center of the field. We hypothesize that each of the MSTd neurons has a center of motion field with a gradient of preferred centers of motion, and that there is an orderly arrangement of MSTd neurons with each region of the visual field being represented by a set of neurons. This arrangement creates the potential for graded responses from individual neurons for different directions of heading as an observer moves through the environment.

[Key words: visual system, motion, optic flow, extrastriate cortex, MST, monkey, FOE]

While moving through the environment, the visual world streams around observers in a pattern which reflects their motion. These optic flow fields combine the effects of all observer movements in three-dimensional space to provide visual information that can guide self-motion, stabilize posture, and reveal the structure of the environment (Gibson, 1950, 1986). All observer movements consist of translation along, and/or rotation around, three axes in space, and similarly all optic flow fields consist of visual patterns created by these motion components.

Attempts to investigate the neuronal processing underlying the analysis of optic flow fields have concentrated on the medial superior temporal area (MST) of the macaque superior temporal sulcus. Neurons in this region show direction selective visual motion responses (Desimone and Ungerleider, 1986; Tanaka et al., 1986; Ungerleider and Desimone, 1986). Those in dorsal MST (area MSTd) have very large receptive fields, often encompassing more than a quadrant of the visual field, and are most responsive to large pattern motion (Tanaka et al., 1986; Komatsu and Wurtz, 1988). Studies in anesthetized monkeys showed that these neurons respond selectively to expanding or rotating patterns (Saito et al., 1986; Tanaka et al., 1986, 1989; Tanaka and Saito, 1989; Saito, 1993) which possess some of the characteristics of optic flow fields. Studies in awake monkeys (Andersen et al., 1990, 1993; Wurtz et al., 1990; Duffy and Wurtz, 1991a,b; Orban et al., 1992; Graziano et al., 1994; Lagae et al., 1994) have demonstrated that many of these neurons respond selectively to many of the elements of optic flow, including radially expanding stimuli.

The response to radially expanding stimuli is of particular interest because it is seen by observers as they move forward. In a previous study of MSTd neurons (Duffy and Wurtz, 1991), we used computer-generated patterns of optic flow in which the radial or circular center of motion was usually in the center of the monkey’s visual field. One possibility is that these neurons might respond better if that center of motion were shifted to different regions of the visual field. To test this possibility, we studied MSTd neurons with large field optic flow stimuli using centers of motion shifted to different regions of the visual field. We find that the response of many MSTd neurons is altered when the center of motion is shifted, and that there are neurons in MSTd that respond best to such shifted centers of motion. We hypothesize that these neurons form a neuronal map of centers of motion, and that such a map might contribute to the determination of heading based on optic flow.

Brief reports of this work have appeared previously (Duffy and Wurtz, 1991c, 1993; Wurtz and Duffy, 1992; Wurtz et al., 1993).

Materials and Methods

Behavioral and physiological procedures. Two adult Rhesus monkeys (Macaca mulatta, subjects 79N and 26K) were trained to sit in a primate chair and perform a visual fixation task for liquid reward. The monkeys sat in the chair for several hours per day and then were returned to their home cages. All experimental 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.

The monkeys were prepared for recording in a single surgical session using procedures described previously (Duffy and Wurtz, 1991a). After anesthetic induction (ketamine 10 mg/kg and atropine 0.04 mg/kg, i.m.) and endotracheal intubation, inhaled Isoflurane was used to maintain a surgical plane of general anesthesia. Scleral search coils were implanted bilaterally (Judge et al., 1980), recording cylinders were placed over parietal cortex bilaterally (AP −2, ML 15), and a stainless steel head

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holder was embedded in a dental acrylic cap that covered the top of the skull. Postoperative analgesia (Flunixin, 2 mg/kg, i.m.) was administered as judged appropriate by the attending veterinarian.

After recovering from surgery, the monkeys were trained to fixate a spot of light on the tangent screen in front of them; eye position was monitored by the magnetic search coil technique (Robinson, 1963). The monkey sat 50 cm away from the center of a 100° × 100° tangent screen. Each trial began with the appearance of the projected image of a red light-emitting diode (LED): 0.25° in diameter, 2.7 cd/m² at the center of the screen. The monkey’s task was to fixate the LED image within 200 msec and maintain binocular fixation (±2.5°) for 0.5–2.5 sec. At the end of successful trials, a reinforcing tone was sounded and the monkey received a liquid reward on a variable ratio reinforcement schedule. During each fixation period, three stimuli were projected onto the screen in succession. Each stimulus had a duration of 1 sec, and interstimulus intervals were between 1 and 1.5 sec. Data points were adjusted to maintain a high success rate during training, with the monkeys successfully completing 90% of the experimental trials after a few weeks.

The behavioral task, stimulus timing, and digitizing of single neuron recordings were controlled by the real-time experimental system (REX) (Hays et al., 1982) running on a dedicated PDP 11/73 computer. Single neuron activity was recorded using a tungsten microelectrode inserted at 1 kHz for storage with markers of stimulus and behavioral events. An on-line raster display showed the occurrence of single neuron discharges with markers added to give the temporal relation to stimulus and behavioral events.

Displays used for later data analysis were spike density histograms created by replacing the pulses representing spikes with Gaussian pulses with a width corresponding to a standard deviation of 20 msec. The average dot speed was 40°/sec in all stimuli, a speed at which MST neurons consistently respond (Tanaka and Saito, 1989; Duffy and Wurtz, 1991a). In planar motion, the stationary dot moved in a rigid world without independently moving objects. Naturalistic cues, such as changes in the blur, brightness, disparity, size, and texture of moving elements in the display, were not included in these stimuli.

One difference between the stimuli in the present study and in our previous study (Duffy and Wurtz, 1991a) is the positioning of our large field stimuli. In our previous study we positioned the stimuli on the anterior bank of the sulcus that is included in the dorsal-medial region (center of gaze) regardless of the location of the receptive field. Our previous experience revealed that the receptive fields of these MSTd neurons are large and come close to the fovea; with our 100° × 100° stimuli, we found that the precise position in the receptive field made little difference in the results (Duffy and Wurtz, 1991b). In these experiments the fixation point was at the center of the screen because our previous experience had revealed that it was the best predictor of the stimulus. We always placed the stimulus in the center of the screen (center of gaze) regardless of the location of the receptive field. Our previous experience revealed that the receptive fields of these MSTd neurons are large and come close to the fovea; with our 100° × 100° stimuli, we found that the precise position in the receptive field made little difference in the results (Duffy and Wurtz, 1991b). In these experiments the fixation point was at the center of the screen because our previous experience had revealed that it was the best predictor of the stimulus. We always placed the stimulus in the center of the screen (center of gaze) regardless of the location of the receptive field. Our previous experience revealed that the receptive fields of these MSTd neurons are large and come close to the fovea; with our 100° × 100° stimuli, we found that the precise position in the receptive field made little difference in the results (Duffy and Wurtz, 1991b). In these experiments the fixation point was at the center of the screen because our previous experience had revealed that it was the best predictor of the stimulus. We always placed the stimulus in the center of the screen (center of gaze) regardless of the location of the receptive field.
We also measured the strength of the relationship between the location of the center of motion and the neuronal response. We plotted the strength of the response to each shifted center of motion stimulus on a polar plot, with the direction of the center of motion as the angle and the response amplitude as the radius. We used circular statistics (Batschelet, 1981) to measure the strength of the relationship between the direction in which the center of motion is shifted and the amplitude of the neuronal response. We tested whether the distribution of response amplitudes around 360° deviated significantly from randomness by deriving the Rayleigh z statistic for each distribution using a significance level of \( p < 0.01 \). The significance of the z values can be interpreted as indicating a clear location preference for shifted centers of motion.

Results

Responses to planar, radial, and circular motion

We studied the responses of 268 neurons on the anterior bank of the superior temporal sulcus. All neurons had large receptive fields that frequently included the fovea and a full quadrant of the visual field, and these neurons responded better to the motion of large patterned stimuli than to the motion of single spots. These characteristics are typical of neurons in the dorsal region of this MST area, MSTd (Komatsu and Wurtz, 1988).

For each cell, we first determined the responses to a standard set of 12 components of optic flow stimuli (shown at the top of Fig. 2) that were centered on the monkey's fixation point. The motion was planar (eight directions), radial (inward and outward), or circular (counterclockwise and clockwise). As we reported previously (Duffy and Wurtz, 1991a), some neurons responded best only to a single type of visual motion stimulus, either planar, circular, or radial motion, and Figure 2A shows an example of such a single component neuron which responded to outward radial motion (darkened spike density plot). Other neurons responded to two of these motion stimuli such as the double component neuron shown in Figure 2B. Others responded to planar, circular, and radial visual motion stimuli, and Figure 2C shows the responses of such a triple component neuron.

Instead of discrete subpopulations of single, double, or triple-component neurons, we again found a continuum of response selectivity for the 12 motion stimuli (Duffy and Wurtz, 1991). However, in order to describe the distribution of cells, we classified each neuron according to the types of stimuli that evoked significant responses (Fig. 3). Triple component neurons formed the largest group, most being planocirculoradial (PCR) neurons with selective responses to planar, circular, and radial stimuli. Also included in this group were nonselective excitatory (NSE) and nonselective inhibitory (NSI) neurons which responded to all three types of stimuli (frequently to all 12 stimuli used). Double-component neurons were mainly planoradial (PR) neurons with responses to a planar or radial stimulus and planocircular (PC) neurons with responses to a planar or circular stimulus. Also included in this group were nonselective inhibitory (NSI) neurons which responded to all three types of stimuli (frequently to all 12 stimuli used). Double-component neurons were mainly planoradial (PR) neurons with responses to a planar or radial stimulus and planocircular (PC) neurons with responses to a planar or circular stimulus. We also found a few circuloradial (CR) neurons with responses to a circular or radial stimulus, a category first reported by Graziano et al. (1990), but they are rare in our sample. Single component neurons are of three varieties: planar (P) neurons responded only to planar stimuli, radial (R) neurons responded only to radial stimuli, and circular (C) neurons responded only to circular stimuli.

Responses to shifted centers of motion

We used stimulus sets with shifted centers of motion which were produced by combining radial motion and circular motion with eight directions of planar motion as illustrated in Figure 1. In our sample of 268 neurons, 245 had significant responses to radial or circular stimuli; they were triple, double, and single

\[ z = \frac{N - 1}{\sqrt{N - 3}} \]
Figure 2. MST neurons respond to one, two, or three types of centered motion stimuli (planar, radial, and circular) used in initial testing of each neuron. The top drawings show the 12 large-field visual motion stimuli used to initially test all MST neurons: 8 planar (directions at 45° intervals), 2 radial (inward and outward), and 2 circular (clockwise and counterclockwise) stimuli. As in Figure 1, each box represents the 100° × 100° stimulus, and the arrows represent the direction of dot motion. A-C, Responses to each stimulus are shown for three neurons. In this and subsequent figures, the neuronal responses are represented by spike density plots averaging responses to six stimulus presentations. The vertical line on each plot marks stimulus onset and indicates a 75 spikes/sec/trial discharge rate; the 1 sec stimulus period is marked by the bold horizontal line. The numbers to the right of each sample are the cell identification numbers; R and L in the numbers indicate right and left hemisphere. A, Example of a single component radial neuron which responded best to outward radial motion (response shown in bold outline) but not to planar or circular motion. This neuron's receptive field filled the left lower quadrant. B, A double component neuron which responded best to leftward planar and outward radial motion (response shown in bold outline) but not to planar or circular motion. This neuron's receptive field extended from 20° above the horizontal meridian to 40° below the horizontal meridian, and from the vertical meridian 35° into the left hemifield. C, A triple component neuron which responded to rightward planar, outward radial, and clockwise circular motion. This neuron had a 25° × 25° receptive field extending from the fixation point into the right upper quadrant.
component neurons. We tested each of these neurons with either the radial or circular stimuli with shifted centers of motion depending upon whether the radial or circular stimulus evoked the strongest response in the initial study of the neuron. In all cases the monkey continued to fixate at the center of the screen. For the largest group of neurons (43%), the response remained largest to stimuli with the centers of motion in the center of the screen, and Figure 4A illustrates the responses of one such neuron. The second most common group of neurons (26%) had responses that were significantly larger to stimuli with the shifted centers of motion (Fig. 4B). An additional 22% of the neurons tested had mixed responses to the shift of the centers of motion, some smaller, others larger. The remaining 9% did not respond differently to the stimuli with shifted centers of motion.

Figure 4C shows how frequently the 245 neurons showed significant changes in response amplitude with the shifted center of motion stimuli. About 90% of the neurons (dark bars) had responses to one or more shifted center of motion stimuli that were significantly different from the responses to the centered stimulus. Thus, a large fraction of the neurons gave a different response to stimuli with shifted centers of motion than they did to the centered stimulus. This suggests substantial response selectivity for the location of the center of motion in these visual motion patterns.

Many neurons showed a systematic preference for centers of motion in only one region of the visual field. This can be illustrated using a polar plot to show the strength of the response for centers of motion shifted in different directions. Figure 5B shows the responses of a neuron with centers of motion primarily in the upper right quadrant. In contrast, the neuron in Figure 5A shows no such preference. The strength of this preference for shifts in given directions can be described using the Rayleigh z statistic (see Materials and Methods) for each of the neurons studied with shifted centers of motion. Figure 5C shows the range of z values for our sample of cells, and shows that 65% of the neurons yielded significant z values (dark bars). These significant values indicate a clear location preference for the direction of shifted centers of motion.

We have so far only considered the effect of shifting the center of motion of the radial or circular stimulus (the preferred stimulus) that gave the best response for each neuron, and we cannot reject the possibility that any stimulus with a shifted center of motion would lead to a different response in these cells. We therefore compared responses to stimuli with shifted centers of motion derived from the preferred radial or circular stimuli to those derived from nonpreferred radial or circular stimuli. For example, the neuron in Figure 6A responded strongly to outward radial motion but not to clockwise circular motion, and it responded more strongly to the radial stimuli with shifted centers of motion. The neuron in Figure 6B responded to outward radial motion and only somewhat less to inward radial motion, but still responded most vigorously to the shifted center of motion stimuli derived from outward radial motion. We made such comparisons for 59 neurons, and Figure 6C shows that most of the neurons (56%) showed significantly stronger responses to the preferred stimulus rather than to a nonpreferred stimulus with shifted centers of motion (solid bar). Only 10% showed stronger responses to the nonpreferred stimulus (open bar) although 34% of the neurons showed comparable responses to the preferred and nonpreferred stimuli (shaded bar). Five times as many neurons showed an increase in response when the stimuli with shifted centers of motion were derived from the preferred radial or circular stimulus as when they were not. Thus, shifting the center of motion of any radial or circular stimulus did not necessarily improve the response.

One possibility was that the center of motion selectivity revealed by these stimuli was somehow a simple product of some other response characteristic previously described. Two of the most obvious characteristics are the receptive field sizes of the neurons and their responses to the components of optic flow. We had only a sufficient number of experiments on the radial stimuli with shifted centers of motion to address these issues. We first determined center of motion preference and then compared that preference to that neuron's receptive field type, dividing receptive fields according to their overall extent as determined with a hand-held moving pattern stimuli. Three types of receptive field were recognized: those with sides of about 25° (15–40°) and restricted to one quadrant of the visual field, those with sides of about 50° (40–70°) and filling one quadrant, and those with sides of about 75° (>70°) and filling substantially more than a quadrant. Figure 7A shows the distribution of center of motion preferences for neurons having these different types of receptive fields. All receptive field types showed less preference for planar stimuli, and comparable preference for shifted and radial stimuli, except the largest fields showing fewer preferring radial stimuli. None of these receptive field types was uniquely related to a stimulus preference.
Figure 4. Neuronal responses to stimuli with shifted centers of motion. At the top, drawings show the centered outward radial stimulus (left) and the eight outward radial and planar stimuli with shifted centers of motion (right). A and B, Spike density histograms showing the responses of two neurons to the outward radial stimuli and stimuli with shifted centers of motion. Vertical line indicates 75 spikes/sec discharge rate; horizontal bar indicate 1 sec stimulus duration. Responses to stimuli with shifted centers of motion that are significantly different from the centered response (t test, p < 0.01) are drawn in bold outline. A, A neuron that responded best when the center of motion was in the center of the stimulus. This neuron is part of a subgroup (14% of the neurons) that showed significantly smaller responses to all eight of the stimuli with shifted centers. This planocircular radial neuron's receptive field was 40° × 40°, extending from the fixation point into the right lower quadrant. B, A neuron that was activated by any stimulus containing outward motion although the strongest responses were evoked by stimuli with centers of motion near the right edge. This planocircular radial neuron's receptive field included all of the 50° × 50° area of the left lower quadrant, except the 5° closest to the vertical meridian. C, Frequency with which neurons responded to shifted center of motion stimuli. Each bin in the histogram shows the percentage of neurons with significant differences between the responses to the indicated number of shifted center of motion stimuli and the centered motion stimulus (t test, p < 0.01). Over 90% of the neurons showed a change (dark bars).

We next examined whether we could predict the response of a neuron to the stimuli with shifted centers of motion if we knew the neuron's response to each of the planar, radial, and circular stimuli (the single, double, or triple component response categories shown in Fig. 2). We suspected that neurons responding to several optic flow components (nonselective excitatory, planocircular radial, and planoradial) would be the ones most responsive to the stimuli with shifted centers of motion; we thought that they might be tuned to shifted centers and that our categorization of them was simply the result of inadequate stimuli
Figure 5. Strength of directional preference for shifted centers of motion. A and B, Polar plots showing response amplitude (radius in spikes/sec) versus direction in which the center of motion was shifted (angle in degrees). A, Polar plot of a neuron showing no clear directional preference, with similar response amplitudes for all eight directions in which the center of motion was shifted. B, The neuron shows strong directional preference and responded most vigorously when the center of motion was shifted upwards (90°). This radial neuron's receptive field included all of the right hemifield. C, The distribution of Rayleigh z statistics describing the strength of the directional preferences of the neurons studied with shifted centers of motion. Those with z values three or less (open bar) were not significantly different from a random distribution around a circle (z value for the neuron in B was 0.2). Those with z values greater than 6 (solid bars) were significantly different from random (with a few exceptions due to small sample size), suggesting a unimodal relationship between the center of motion and response amplitude (z value for the neuron in A was 8). Those with z values greater than 3 but 6 or less (shaded bar) were, or were not, significant as a function of sample size. A total of 65% of the neurons yielded significant z values, indicating most had unimodal responses.

Spatial distribution of preferred centers of motion

Because many of the MSTd neurons gave their strongest response to the stimuli in which the center of motion was shifted away from the center of the stimulus, we investigated whether there was any preference among the neurons for one part of the visual field such as above or below the horizon or near or far from the center. Figure 8A shows the spatial distribution of the centers of motion of the stimuli we used, with each of the 17 boxes representing the 100° × 100° stimulus, the arrows representing the direction of dot motion, and the circles representing the center of motion. The stimuli are ordered in Figure 8A so that the one with centered motion is in the middle, those with a shifted centers of motion are placed eccentrically in the direction of the shift, and those with planar motion are at the edge of the figure as if their centers of motion were in the far periphery.

We observed all three possible response patterns among the neurons studied with these stimuli: those with their strongest responses to centered, eccentric, or peripheral centers of motion. Figure 8B–D show the responses of three neurons all of which responded to outward radial motion. The spike density histograms of their responses are located in the positions occupied by the stimuli in Figure 8A. The best responses are darkened, as are the responses that were not significantly different from the best responses. Figure 8B shows the best response to the centered motion with all other responses being significantly smaller. In Figure 8C the best responses are to an eccentric center of motion stimulus, and in Figure 8D the best response is to a planar stimulus, one that can be regarded as having a peripheral center of motion.

Figure 9A shows the distribution of preferred centers of motion for our sample of neurons that responded to outward or inward radial motion and were tested with radial stimuli with shifted centers of motion. We did similar experiments using clockwise and counterclockwise circular motion, and Figure 9C shows the comparable distribution for neurons which preferred these stimuli. The distribution of preferred centers of motion included all of the 17 shifted center sites tested, suggesting that...
Figure 6. Comparison of responses to stimuli with shifted centers of motion derived from preferred and nonpreferred radial or circular stimuli. A. Comparison between the responses of a planoradial neuron to the preferred outward radial stimuli (top row), and the nonpreferred clockwise circular stimuli (bottom row). The preferred radial stimulus with a shifted center of motion evoked stronger responses (Rayleigh z for this distribution equals 10.55, significant p < 0.001) than the nonpreferred circular shifted center of motion stimuli (z = 6.12, not significant). This illustrates the effect of stimulus preference for one type of motion (radial vs circular). This neuron’s receptive field covered the left lower quadrant, and extended 15° into the left upper quadrant. B. Comparison between the responses of a neuron that responded only somewhat better to outward radial stimuli (top row) than to inward radial stimuli (bottom row). The outward radial shifted center of motion stimuli evoked stronger responses (z = 37.42, significant p < 0.001) than the nonpreferred inward radial shifted center of motion stimuli (z = 1.90, not significant). This illustrates the effect of stimulus preference for one direction of motion (outward vs inward) within a type of motion. This neuron’s receptive field covered the left upper quadrant, and extended 10° into the left lower quadrant. Vertical line on the spike density plots indicates 75 spikes/sec; horizontal bar indicates 1 sec. C. Comparison of responses to shifted centers of motion stimuli derived from preferred and nonpreferred stimuli for all 59 cells so studied. If the neuron showed a significantly stronger response (t test, p < 0.01) to one of the preferred shifted centers of motion stimuli than to any of the nonpreferred shifted center of motion stimuli it was placed in the left bin of the histogram. Most neurons (56%) showed stronger responses to the preferred stimuli with shifted centers of motion.
to a type of center of motion preference, and no clear trends are evident.

The population might be capable of responding to a wide variety of different centers of motion.

The bar graphs in Figure 9, B and D, show the proportion of neurons preferring central, eccentric, and peripheral centers of motion. The most frequently preferred stimuli with shifted centers of motion among both the radial and circular neurons were those with eccentric centers of motion. This preference included two neurons that showed no statistically significant response to any of the 12 basic motion stimuli (see Fig. 2) but showed strong responses to stimuli with eccentric centers of motion. Although the eccentric centers of motion were the most frequently preferred stimuli, this in part reflects the fact that eight center of motion stimuli are summed together. The most frequently preferred individual stimulus is the centered one for both the radial and circular stimuli. Thus, taking the mean number of neurons along a line from periphery to center, there was an increasing preference for centers of motion closer to the center of the field.

The higher frequency with which neurons responded best to a motion stimulus centered in the visual field could reflect the fact that the initial screening study presented only radial and circular stimuli with the center of motion in the center of the field. However, we studied all of the neurons we could successfully isolate, and we usually did not know into what component class the response to our standard 12 stimulus test set would place the neuron, a procedure that should have reduced the impact of any selection bias.

Specificity for preferred centers of motion

Because many neurons had significant responses to stimuli with centers of motion in more than one location, we measured how widely spread these preferred centers were by determining the number of centers of motion stimuli evoking responses that were not significantly different from the best response. In Figure 10 we divided the responses into highly selective (responses to one center of motion), moderately selective (two or three centers of motion), and less selective (>three centers of motion). A substantial number of neurons (19%—45/236, solid bar in Fig. 10) were highly selective, and an additional 36% (85/236—shaded bars) were moderately selective so that over half of the neurons responded to centers of motion in no more than three locations. Over 95% of the neurons had significant responses to eight or fewer stimuli. We consistently found the centers of motion contiguous with one another rather than spread separately around the visual field.

One might expect that center of motion preference should possess greater spatial resolution towards the center of the visual field, providing observers with more refined information of subtle path deviations in the most common circumstances (looking where they are going). To test this prediction we determined whether the number of centers of motion stimuli to which a cell responded was related to where the preferred centers of motion were located. Figure 11 makes this comparison for all neurons (Fig. 11A) and for neurons having preferred centers of motion centered, eccentric, or peripheral in the field (Fig. 11B–D). The neurons preferring the center showed a greater percentage with a high degree of center of motion selectivity. The neurons preferring eccentric and peripheral centers of motion showed lower percentages with a high degree of center of motion selectivity.

This indication of greater spatial selectivity for centers of motion in the center of the field, prompted us to test neurons with a larger number of stimuli having centers of motion concentrated closer to the center of the field. To do so, we added a set of stimuli having centers of motion which were midway between those of the eccentric and centered stimuli (22° from the center of the stimulus, Fig. 12A). Figure 12, B and C, shows the results of such a study in a planoradial neuron that responded to inward radial motion. In this neuron, stimuli having centered or peripheral centers of motion evoked weak responses, eccentric stimuli evoked stronger responses, and pericentric stimuli evoked the strongest responses. Of the 44 neurons studied with this expanded set of 25 stimuli, half preferred pericentric stimuli, with the remainder distributed between peripheral, eccentric, and centered preferences. Thus, the more extended stimulus set revealed a more localized center of motion preference than could be seen with the set of only 17 stimuli and revealed a greater spatial
Figure 8. Location of the best responses to the center of motion stimuli. 

A. The 17 stimuli used with neurons that responded to outward radial motion. Each box represents the 100° × 100° stimulus, the arrows within each box represent the directions of visual motion, and the circles indicate the center of motion. The outer ring of boxes represent planar stimuli; their outward centers of motion can be regarded as being 90° from the center, off the screen in these stimuli. The inner ring of boxes represent stimuli with shifted centers of motion, their outward centers of motion are 45° from the center, near the edge of the screen in these stimuli. The center box represents the outward radial stimulus with its center of motion at the center of the screen. 

B–D. Spike density histograms showing the responses of three neurons to the stimuli occupying the same positions as the stimuli illustrated in A. The darkened histograms are those that were not significantly different from the largest response (t test, p < 0.01). B, A neuron that responded best to the stimulus with its center of motion at the center of the screen. This radial neuron’s receptive field was in the right hemifield. C, A neuron that responded best to stimuli with eccentric centers of motion. This radial neuron’s receptive field covered a 50° × 35° area in the left upper quadrant. D, A neuron that responded best to a planar stimulus with a peripheral center of motion. This planoradial neuron’s receptive field covered a 20° × 20° area in the right upper quadrant.

resolution near the fixation point. Thus, the central area of the visual field appears to be represented by a relatively greater proportion of the neurons (Fig. 9) and by neurons that yield their best responses to centers of motion limited to the smallest area of the visual field (Fig. 11).

With the expanded set of stimuli (25) we could also measure how close together the preferred stimuli were in the visual field. For example, Figure 12C shows that the stimuli that evoked responses comparable to the best response (solid) or significantly above control activity level (shaded) were clustered together in the visual field. This illustrates a clustering of the centers of motion that evoke strong responses from a neuron that
Figure 9. Distribution of centers of motion that evoked the strongest response. A, Distribution for neurons that preferred radial (inward or outward) stimuli. The 17 segments of the circular plot represent the 17 stimuli, with each segment representing the center of motion stimulus depicted at the corresponding position in Figure 8A. Each dot represents a neuron, and the dots are placed in the segment which corresponds to the stimulus that evoked the largest response from that neuron. All tested centers of motion were preferred by at least one neuron. B, The relative distribution of the preferred centers of motion shown in A. The graph shows the percentage of neurons that had the strongest response to center, eccentric, or peripheral centers of motion stimuli. Stimuli having eccentric centers of motion were most commonly preferred, but the stimulus with the center of motion in the center of the screen was the single stimulus preferred by the largest number of neurons. C, Distribution for neurons that preferred circular (clockwise or counterclockwise) stimuli. Again, all centers of motion were preferred by at least one neuron. D, The relative distribution of the preferred centers of motion shown in C. As with the neurons tested with radial combinations, eccentric centers of motion are most commonly preferred, but the centered center of motion stimulus was the single stimulus preferred by the greatest number of neurons.
Figure 10. Number of the 17 stimuli evoking a significant response in neurons studied with radial and circular stimuli with shifted centers of motion. Neurons are divided into three groups: highly selective neurons (solid bar) in which only one of the 17 stimuli evoked a response significantly different from the background response, moderately selective neurons (shaded bars) in which two or three stimuli evoked significant responses, less selective neurons (open bars) in which more than three stimuli evoked significant responses. Over half the neurons responded to centers of motion in no more than three locations.

was evident in all of our experiments. In the 44 neurons studied with the 25 stimuli, 97% of the 233 significant responses were adjacent to other significant responses. Thus, the preferred stimuli lie in one part of the visual field.

We examined the spatial distribution of the preferred centers of motion in more detail in 12 neurons in which we presented 33 stimuli having centers of motion within the central 40° of the visual field, arranged at 10° intervals along four axes: horizontal, vertical, and both 45° obliques (Fig. 13A). Figure 13B–D shows the results of studies on three neurons. The neuron in Figure 13B had its strongest responses to stimuli having centers of motion in the lower left corner of the visual field. The neuron in Figure 13C had its strongest responses to stimuli having centers of motion clustered around or at the center of the visual field or symmetrically around it. The neuron in Figure 13D had its strongest responses to stimuli having centers of motion asymmetrically clustered around the center of the field, but with a notch of decreased responsiveness to stimuli with the center of motion at the center of the visual field. The three examples illustrated in Figure 13 represent the three major types of spatial structure observed in these 12 neurons: a peripheral corner (5 neurons), a central peak (4 neurons), and a central notch (3 neurons). These studies suggest that the preferred center of motion, rather than being a simple segment of the visual field, may have a more complex structure which our relatively limited stimuli have not fully defined.

Figure 11. Interactions between the location of the preferred center of motion (as in Fig. 9) and the degree of center of motion selectivity (as in Fig. 10). A, The percentage of all neurons tested which showed highly (solid bar) moderately (shaded bar), or less (open bar) selective responses for center of motion stimuli. B–D, The percentages of center preferring (B), eccentric preferring (C), and peripheral preferring (D) neurons which showed highly, moderately, or less selective responses for center of motion stimuli. Center preferring neurons most commonly showed highly selective responses, whereas eccentric and peripheral preferring neurons most commonly showed moderately or less selective responses.
Discussion

Neuronal responses to shifted centers of motion

We studied MSTd neurons that responded to radial or circular motion in the middle of the visual field by presenting the preferred pattern with its center shifted to eight points around the field. We found that MSTd neurons were sensitive to such shifts: at least one of the eight shifted center stimuli produced a response significantly different from the centered stimulus in 91% of the neurons. In our sample of 245 neurons, over half gave significant responses to three or fewer stimuli; the preferred centers of motion were clustered in one part of the field and not scattered about. The neurons studied preferred centers of motion throughout all parts of the visual field; a larger number of neurons preferred the center of motion in the middle of the field and these neurons responded to centers of motion over a more limited area of the visual field. These observations suggest that the shifted center of motion, and the accompanying changes in the pattern of motion throughout the visual field, is a stimulus characteristic which influences the responses of MSTd neurons.

Hypothesis on functional organization

Based on these observations, we suggest that each of the MSTd neurons can be regarded as having a center of motion field. The responses of individual neurons would be graded according to the proximity of the center of motion of a stimulus to the preferred center of motion for the neuron, just as other visual cortical neurons have graded preferences for the direction of planar motion or for the location of a spot of light. The role of MSTd neurons in interpreting optic flow fields would not be one of qualitative feature matching, but rather one of responding to visual motion according to the degree of match between the visual input and the preferred optic flow field of the neuron. Thus, each neuron would contribute to the population response to a number of stimuli, just as many neurons would contribute to the population response evoked by any stimulus. This population encoding would have the advantage of remaining robust in spite of idiosyncracies in any individual neuron's response profile.

We suggest further that there is an orderly arrangement of MSTd neurons with each region of the visual field being represented by a set of neurons whose center of motion falls in that region of the field. The population of neurons would have a gradient in the density of the centers of motion with those near the pericentric centers of motion are shifted 22° from the center of the screen, halfway between the eccentric centers of motion (45°) and the screen center. B. Graph of response amplitudes evoked by stimuli shown in A with peripheral (diamonds), eccentric (circles), pericentric (squares), and centered (far right diamond) centers of motion. Shaded symbols indicate responses that were significantly above control activity level, solid symbols represent responses that were not significantly different from the best response. The horizontal lines indicate unstimulated control activity in the trials whose responses are represented by the same type of line. Arrows on abscissa show the direction of the shift of the center of motion. C. Plot of these responses with 25 segments representing the 25 center of motion stimuli arranged as shown in A. The shaded and solid segments show the same significant responses shown in B. Stimuli having centers of motion immediately to the right of the center of the screen evoked the strongest response which indicates a finer grain in the preferred center of motion location than can be determined by stimuli with 17 centers of motion in Figure 8. This planar-radial neuron's receptive field included all of the lower hemifield, except the 10° at the right most edge of the screen.
Figure 13. Types of neuronal responses to stimuli having centers of motion concentrated within the central 40° of the visual field. A, Diagrammatic representations of a set of 33 outward radial stimuli. The centers of motion are distributed at 10° intervals along four axes. B–D, The responses of three individual neurons. Responses are represented by ovals at positions which correspond to the relative position of the centers of motion. Response amplitude at each position is proportionate to the length of the oval and is derived from the spike density histograms as described in Materials and Methods. Activity during unstimulated control trials is represented below each plot. B, The responses of a neuron that preferred stimuli having centers of motion in the lower left corner of the screen, with decreasing response amplitude graded towards the center. This planoradial neuron’s receptive field covered the right hemifield to 30° above and below the horizontal meridian. C, The responses of a neuron that preferred stimuli having centers of motion nearer the center of the screen, with the greatest responses to stimuli in which the center of motion was at the center of the screen. This radial neuron’s receptive field extended for 40-45° from the fixation point into the right lower quadrant.
motion, with those neurons responding to more peripheral centers of the key observations that all areas of the visual field are represented by a neuron is sensitive. The schematic drawing is based on the response to neuronal activity in MT whereas the elements related to heading are more comparable to the neurons studied in MSTd. However, we believe that our data do not allow us to definitively favor any of these hypotheses. Our hypothesis of a map of headings among the MSTd neurons is particularly relevant to the recent models of Perrone (1992) and of Perrone and Stone (1994), who used a template to determine heading: a map of the visual field has different headings represented by different neural elements, and the heading is identified by the most active element. The organization of heading directions that we have hypothesized (Fig. 14) is almost identical to the heading map shown in this model. In addition, the series of constraints that they place on this model to make it biologically realistic (Perrone and Stone, 1994) produces an organization that would seem to have the same grain of heading information which could be provided by the neurons.

Relation to heading determinations
We think that MSTd neurons that respond to radial motion with centers of motion (or a focus of expansion (FOE) in the case of the radial stimulus) arrayed across the visual field could contribute to the determination of heading of observers moving through the environment. While heading may be determined by a combination of visual cues (Cutting, 1986), our findings support the conclusions of a number of recent psychophysical experiments suggesting the existence of brain mechanisms which might derive heading from optic flow. Recent psychophysical experiments using simulations of the optic flow produced by observer movement have demonstrated that humans can make heading judgements within a few degrees of the simulated heading (Warren and Hannon, 1988, 1990; Warren et al., 1988; Stone and Perrone, 1991, 1993; van den Berg 1992). Such judgements are frequently better when the eyes move (Royden et al., 1992) and fail only when only a single plane (like a wall) is present (Regan and Beverley, 1982; Warren and Hannon, 1990).

In our physiological experiments, we have tried to eliminate all cues but optic flow, simulating motion toward a single plane during visual fixation of a stationary target. The information relevant to heading judgements in our display therefore is limited. Our physiological observations would be entirely relevant, however, to determining heading in the circumstance in which the point of fixation is at a distance, without the interaction with multiple depth planes or pursuit eye movements. The forward view from an airplane during landing, used by Gibson (1986) to illustrate flow patterns, would seem to closely approximate this condition since there is radial flow with little rotation. For example, the MSTd neurons studied with the dense array of center of motion stimuli (Fig. 13) revealed patterns of center of motion sensitivity which suggests that some cells might differentiate between going where the observer is looking versus going in another direction. This is illustrated by the comparison of Figure 15, A and B, in which the flight path leading towards the landing strip (Fig. 15A) would be detected by a neuron with a preferred center of motion in the center of the field whereas a flight path leading to the lower left corner (Fig. 15B) would be detected by a neuron with the preferred center of motion off to the side. While this is a limited and highly specific example of the use of the FOE structure of these MSTd cells, they indicate the type of heading information that could be provided by the neurons.

Our experiments might also be relevant to the recent models which attempt to show how directionally selective elements might produce heading judgements (Heeger and Jepson, 1990; Warren et al., 1991; Perrone, 1992; Lappe and Rauschecker, 1993; Perrone and Stone, 1994). All attempt to use two-dimensional retinal information to reconstruct three-dimensional heading information, with some requiring differential motion in a region of the field (Longuet-Higgins and Prazdny, 1980; Rieger and Lawton, 1985; Rieger and Toet, 1985). In general, determination of directionally selective elements in the models corresponds to neuronal activity in MT whereas the elements related to heading are more comparable to the neurons studied in MSTd. However, we believe that our data do not allow us to definitively favor any of these hypotheses.

Our hypothesis of a map of headings among the MSTd neurons is particularly relevant to the recent models of Perrone (1992) and of Perrone and Stone (1994), who used a template to determine heading: a map of the visual field has different headings represented by different neural elements, and the heading is identified by the most active element. The organization of heading directions that we have hypothesized (Fig. 14) is almost identical to the heading map shown in this model. In addition, the series of constraints that they place on this model to make it biologically realistic (Perrone and Stone, 1994) produces an organization that would seem to have the same grain of heading specification suggested by our experiments. Further specification of the density of the hypothesized heading map in MSTd and the effect on this map of rotation in addition to translation might allow further convergence between the neural models and the physiological hypotheses.

Position invariance
A key point in the analysis of MST neuronal responses to the components of optic flow stimuli is whether these responses can simply result from planar motion in one part of their receptive fields. In our previous experiments (Duffy and Wurtz, 1991a), we addressed this issue by testing whether radial or circular stimuli, which were effective when presented over the full 100° × 100° field, remained effective when we presented a smaller version of the same stimulus. We found that the single component radial and circular neurons were the most likely to continue
to respond as we moved these small field stimuli into different segments of the larger field, indicating that their responses could not be explained by their response to simple planar motion. In contrast, triple component neurons showed the greatest change in response when the stimuli were moved to different regions of the field suggesting that their responses could result from sensitivity to simple planar motion. Andersen et al. (1990, 1993) and Graziano et al. (1994) used smaller stimuli (10° × 10°) and shifted them 20° to study the same issue, and also found many MST neurons that continued to respond after the stimulus was shifted. Orban and his collaborators (Orban et al., 1991; Lagaë et al., 1994) have reported a series of experiments which extend these conclusions and suggest a role for adjacent area MT in developing these response properties. In net, these experiments reach the same conclusion, that some MST neurons continue to respond when the stimulus is shifted on their receptive fields.

The continued response of some MST neurons in spite of a shift of stimulus location has been referred to as position invariance, and it has recently been argued (Graziano et al., 1994) that such position invariance would preclude the use of these neurons in the determination of heading. Our present experiments using large field stimuli, rather than the small field stimuli just discussed, reveal the limitations of position invariance. We found many cells that responded best when the center of motion fell in one region of the visual field, rather than another, and that about 90% of the MSTd neurons showed an alteration in the strength of their responses when the center of motion was shifted. When we look at the results of the previous experiments using small field stimuli, we also see shifts in the magnitude of the response even though the preferred stimulus remained the same, so that many of these observations would be better regarded as indicating relative invariance rather than absolute invariance. In the previous experiments, stimuli were shifted over relatively small distances in the field, leading to response changes that could now be explained by movement across the graded preference for centers of motion revealed by our large field stimuli. Thus, MSTd neurons could provide information related to heading.

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