Neural responses to velocity gradients in macaque cortical area MT

STEFAN TREUE1 AND RICHARD A. ANDERSEN2
1Cognitive Neuroscience Laboratory, Department of Neurology, University of Tübingen, Auf der Morgenstelle 15, 72076 Tübingen, Germany
2Division of Biology, California Institute of Technology, Pasadena
(RECEIVED December 16, 1994; ACCEPTED January 30, 1996)

Abstract
Visual motion, i.e. the pattern of changes on the retinæ caused by the motion of objects or the observer through the environment, contains important cues for the accurate perception of the three-dimensional layout of the visual scene. In this study, we investigate if neurons in the visual system, specifically in area MT of the macaque monkey, are able to differentiate between various velocity gradients. Our stimuli were random dot patterns designed to eliminate stimulus variables other than the orientation of a velocity gradient. We develop a stimulus space ("deformation space") that allows us to easily parameterize our stimuli. We demonstrate that a substantial proportion of MT cells show tuned responses to our various velocity gradients, often exceeding the response evoked by an optimized flat velocity profile. This suggests that MT cells are able to represent complex aspects of the visual environment and that their properties make them well suited as building blocks for the complex receptive-field properties encountered in higher areas, such as area MST to which many cells in area MT project.

Keywords: MT, Velocity gradients, Structure from motion, Optical flow, Surfaces

Introduction
Velocity gradients* are ubiquitous in our environment. They provide important cues about the three-dimensional (3-D) layout of the visual scene. In the large optical flow fields that result from the observer's motion through the environment they contain information about observer heading (Regan, 1985; Warren & Hannon, 1988; Crowell & Banks, 1993). Smaller velocity gradients, caused by the self-motion of objects, determine the perceived 3-D structure of moving objects in the perception of structure from motion (Rogers & Graham, 1979; Braunstein & Andersen, 1984; Siegel & Andersen, 1986; Andersen, 1989; Husain et al., 1988; Landy et al., 1991; Treue et al., 1991; Harris et al., 1992). The importance of the information contained in velocity gradients is reflected in the high sensitivity of the human visual system to detecting the presence of such motion patterns (Nakayama, 1981; Golomb et al., 1985; Nakayama et al., 1985).

The analysis of optical flow fields has received much attention from psychophysical (Gibson, 1950; Regan & Beverly, 1981, 1985; Regan, 1986; Regan et al., 1986; Royden et al., 1994; Warren et al., 1988; Warren & Hannon, 1988) and computational vision research (Koenderink, 1986; Longuet-Higgins & Prazdny, 1980; Koenderink & Van Doorn, 1981; Rieger & Lawton, 1985) demonstrating the importance of extracting the local characteristics of the flow pattern when trying to recover the 3-D layout of the visual scene from the two-dimensional (2-D) projections on the retinæ.

More recently, electrophysiological studies have attempted to localize the cortical areas involved in this analysis (Saito et al., 1986; Tanaka et al., 1986, 1989; Tanaka & Saito, 1989; Duffy & Wurtz, 1991a, b; Orban et al., 1992; Graziano et al., 1994). Area MST in the macaque monkey has been shown to contain cells with large receptive fields that are sensitive to rotating, expanding, contracting, and spiraling patterns, respectively. While naturally occurring optical flow fields can be decomposed into these more elementary motion patterns, they are still rather complex in that not only the speed but also the direction of feature motion varies as a function of spatial position. Single objects in motion and stimuli used for structure from motion studies on the other hand are often simpler in that their flow patterns are unidirectional. Such stimuli might thus serve as a bridge between the studies using very impoverished stimuli (like bars, sine-wave gratings, or simple random dot patterns) to get a detailed understanding of the physiology of motion perception and the aforementioned studies using more complex displays that are closer to natural images yet are harder to characterize. The
random dot patterns we developed for this study represent such intermediate stimuli in that they form the building blocks for complex optical flow patterns yet they can be readily parameterized.

Optical flow patterns often cover a large part of the visual field and represent many objects at different depths from the observer, while structure from motion stimuli are smaller in scale and numerosity. Despite these global differences, the two types of motion patterns are locally similar. The characteristic feature of both of them is the presence of local variations in direction and speed of motion, i.e. of velocity gradients. To perceive these motion patterns and to recover the 3-D shape of the underlying objects requires the accurate representation of the velocity gradients in the visual system. Here we introduce moving random dot patterns simulating the most elementary velocity gradient types (see Fig. 1 and Methods for details) to establish if neurons in area MT of the awake behaving monkey are able to encode the orientation of such gradients in their receptive fields. Area MT was chosen for this study since its receptive fields are smaller than those in area MST and thus seem well suited to analyze the local components of larger flow fields. At the same time the receptive-field sizes in MT match well with the size of commonly occurring structure from motion (SFM) gradients. Finally, long-lasting deficits in the perception of SFM occur with MT lesions, suggesting that this area plays a role in the perception of 3-D structure from motion (Siegel & Andersen, 1986; Andersen & Siegel, 1990).

The visual environment contains an infinite number of different velocity gradients. Since this is the first physiological study using such stimuli, we limit ourselves to stimuli containing simple linear gradients. These velocity gradients all belong to the same class and thus can be represented easily in what we call a "deformation space" (see below). The deformation space has great resemblance to the idea of a "spiral space" developed in our laboratory as a reference frame for characterizing the response properties of MST cells to rotating, expanding, and contracting stimuli (see Fig. 3; Graziano et al., 1994; for a similar approach using polar, hyperbolic, and Cartesian gratings to study V4 neurons, see Gallant et al., 1993).

Here we demonstrate that many MT neurons are tuned to velocity gradients, i.e. the response of an MT neuron to a given gradient often varies as a function of that gradients position in the deformation space.

Methods

A detailed description of our recording methods has appeared elsewhere (Snowden et al., 1991) and this section will therefore be limited to a brief overview and a detailed description of the stimuli used.

Two male rhesus monkeys were trained to fixate a small fixation point, while ignoring any other stimuli, and to signal the dimming of the fixation point by releasing a lever. The animals' eye movements and point of fixation were monitored using a scleral search-coil technique (Robinson, 1963). Visual stimulation was provided to the receptive field of individual neurons (which were about 5–10 deg eccentric) during this 4–6 s period of fixation. Electrode penetrations were made through a chamber implanted over the parietal cortex. The electrode's position within the chamber, the depth of recording, the topography of receptive-field locations, as well as the properties of the cells encountered during the penetrations were used to determine whether encountered cells were indeed in area MT.

Experimental protocol

Stimuli were presented on a large HP CRT screen or a video monitor at a viewing distance of 57 cm. They consisted of random patterns of bright, high-contrast dots upon a dark background. Each dot was approximately 1 mm in diameter, and thus subtended about 6 min arc. Dot density was 4 dots/cm².

The rate of screen refresh was 50 Hz for the CRT screen and 60 Hz for the video monitor. Each trial commenced with the onset of the fixation point. After about 1 s, a stimulus appeared if the animal had pulled the key and had not broken fixation. This stimulus lasted for 1 s, and another stimulus appeared for 1 s after a 1-s delay. The fixation point dimmed 0.2–2.0 s after the disappearance of the second stimulus; thus a complete trial lasted 4.2–6 s. The response of a neuron was determined by the average firing rate during the stimulus presentation (excluding the first 100 ms after the stimulus onset).
Stimuli

To determine the preferred direction of motion for a cell, we presented random dot patterns moving behind a circular virtual aperture. The preferred direction of the neuron determined in this way was used throughout the rest of the tests performed on this cell.

We next established the cell’s preferred speed. To determine if a cell is gradient tuned, we presented it with various linear velocity gradients all moving in its preferred direction. The average speed of each pattern was equal to the preferred speed of the cell. The velocity of every point in the pattern varied linearly as a function of its position within the gradient. Such gradients can be divided into two groups.

Shear stimuli

In these stimuli, the velocity gradient is oriented perpendicular to the direction of motion. Thus, the velocity of a particular point will be constant while it is moving across the stimulus. We call patterns in which the velocity decreases from the left of the stimulus to the right (when facing in the direction of motion) clockwise (CW) shearing stimuli\(^\dagger\) (Fig. 1a). Correspondingly, in counterclockwise (CCW) shearing stimuli (Fig. 1b), the velocity increases from the left to the right.

Compressive/stretching stimuli

In these stimuli, velocity varies along the direction of motion. Thus, a point in a compressive gradient will decrease its velocity while crossing the display (Fig. 1c). Correspondingly, points in stretching gradients will accelerate while crossing the display (Fig. 1d).

We used two different gradient slopes in our experiments. The steeper velocity gradient would start at 0 deg/s on the slower end of the display and would reach twice the preferred speed of the cell under study at the opposite end. The shallower gradient would start at half the preferred speed and reach 1.5 times the preferred speed. Note that for both stimuli the average speed as well as the speed in the center of the stimulus is equal to the preferred speed of the cell.

Points that moved across the sides of the flat velocity profiles used for determining the preferred direction and speed of a cell were simply wrapped around to the opposite side of the stimulus. This simple technique is sufficient to insure equal dot density across the stimulus for these simple patterns as well as our shear gradients.

The acceleration or deceleration of individual dots in our compression stimuli on the other hand would lead to changes in stimulus density across the display especially for those compressive gradients in which dot speeds decrease all of the way to zero at one end of the display. We therefore employed two techniques to eliminate this density cue in our displays.

Special dot wraparound. Expansive dot gradients that start with speeds of zero at one end and with even density distributions across them will remain evenly distributed. While the distribution will remain uniform, the dot density will be continuously falling because the distance between any two points will continuously increase.\(^\S\) We prevent this decrease in density in our displays by reploting any points that cross the stimulus boundaries back into the stimulus. Notice that this reploting has to be done randomly across the stimulus rather than using the wraparound method employed for the flat velocity profiles. Stimuli in which the lowest speed is larger than zero were generated by first generating a larger stimulus whose gradient starts at a speed of zero and then masking this stimulus to show only a smaller extent of the velocity gradient.

Compressive gradients are generated by first computing a stretching gradient moving in the opposite direction and then reversing the order of the individual frames making up the stimulus. Notice that in the resulting stimuli dots will disappear while approaching the zero speed stimulus edge. Thus there is no “piling-up” of dots at that edge.

Limited dot lifetimes. Replotting dots within our stretching stimuli and removing dots in the compressive stimuli generates transient events that could possibly influence the response of cells to these patterns. To insure that this transiency does not influence our findings, we introduce it into all of our stimuli. This is achieved by using dots of limited lifetimes. The dots move along a continuous path for only a short period of time—their lifetime. After its lifetime, a dot is randomly replotted within the stimulus. We used a lifetime of 300 ms for all of our random dot patterns which was long enough to not substantially affect the percept of motion while at the same time providing a significant amount of transiency, masking the transiency generated by the appearing and disappearing dots in the stretching and contractive gradient stimuli, respectively.

Our stretching, compressive, and shearing dot patterns are members of a continuous family of stimuli which only vary along one angular dimension.** This dimension is the angle between the direction of the vector describing the velocity gradient (the “gradient vector”) and the direction of dot motion in the pattern. If the gradient vector points in the direction of stimulus motion, the stimulus is stretching. If the gradient vector points in the opposite direction, the stimulus is being compressed, while gradient vectors orthogonal to the direction of pattern motion occur in shearing stimuli. These four stimulus types form the cardinal axes of a coordinate system that we term “deformation space” (Fig. 1). Stimuli that fall between these cardinal directions combine elements of stretching or compression with shear components. Note that all of the gradients contain the same velocity vectors, only spatially rearranged.

This deformation space gives us a convenient means of plotting the response of neurons to our gradient stimuli in the same way as responses of direction-selective units can be plotted in a direction of motion space or in the same way as area MST neurons’ responses to expanding, contracting, and rotating patterns can be plotted in a spiral space (see also Graziano et al., 1994).

All our stimuli were always centered on the receptive field and their size was chosen so that they would not extend beyond the boundaries of the classical receptive field, i.e. the parts of

\(\dagger\)We chose this nomenclature since in a counterclockwise rotating dot field velocity also increases from right to left when facing in the direction of motion. Note though that all of the dots in our shearing stimuli move along a straight path.

\(^\S\)This phenomenon is well known in astronomy where the finding that any two stars are moving away from each other lends support to the idea of an expanding universe.

**This is true only if all of the gradients have the same steepness of slope.
the receptive field from which we were able to elicit a response from the cell with single handheld stimuli.

Results

The aim of our study was to establish if area MT neurons are able to signal the presence and the orientation of velocity gradients in their receptive fields. In several aspects this question is similar to determining if a given cell is able to signal the direction of motion. Generally, a cell is considered direction selective if it displays a tuning curve when presented with a range of different directions. Additionally, one could require that the cell responds stronger to at least one direction of motion than to a stationary pattern. We use both of the corresponding requirements to determine if the MT cells we recorded from are able to signal the orientation of velocity gradients.

Fig. 2 shows the full set of tests we performed on every neuron considered for this study. After mapping the receptive field, we determined the cells preferred direction (Fig. 2A) and then its preferred speed. This established the most effective flat velocity profile. To determine if the cell showed gradient tuning, we ran a block of trials presenting velocity gradients of eight different orientations spaced evenly in the deformation space. Interleaved with these trials were presentations of the best flat velocity profile (as determined in the two previous blocks of trials). Fig. 2C plots the result. The dashed circle is the level of response elicited by the presentation of the flat velocity profile. The solid line connects the responses to the eight velocity gradients. The arrow is an estimate of the preferred velocity gradient of this cell. This cells shows a clear tuning to the velocity gradients with the response to several (neighboring) gradient orientations significantly ($P < 0.0001$, paired $t$-test) stronger than the flat profile.

We were able to record long enough from 25 cells (22 from one and three from the other animal) to conduct all of these tests. We classified the type of responses to the velocity gradients according to three criteria. In those cases where the responses to the two gradients slopes used were not similar, we used the one that fell more clearly into one of the classes described below.

1. Did the cell show a tuned (i.e. single-peaked) response to the velocity gradients? (tuned cells)
2. If the cell was tuned, did the response to the most effective velocity gradient clearly exceed the response to the flat velocity profile? (excitatory tuned cells)
3. If the cell was not tuned, was the response to all velocity gradients significantly less than to the flat profile? (inhibited cells)

This division of cells into classes is intended to show the existence of certain properties and not to imply that as a population MT cells fall into separable groups.

Fig. 3 shows two representative examples of each response type as well as the relative frequency of their occurrence. The response of three cells (12%) did not fit into this scheme.

About a quarter (six of 25) of the cells fell into the tuned category, while another quarter (seven of 25) were categorized as inhibited cells. More than a third (nine of 25, 36%) of the cells that could be characterized were excitatory tuned cells, our most stringent class. Thus, 60% of the cells we recorded from showed tuning to velocity gradients. The existence of excitatory tuned cells is especially important in the context of this study since it rules out an obvious possible artifact. If our stimuli are not perfectly centered on the receptive field and if the velocity tuning curve is not symmetrical around its preferred velocity, the cell would show a tuning curve when presented with our various gradient stimuli. But note that this tuning curve would never exceed the firing rate based on the preferred flat velocity gradient. By being able to demonstrate that some cells show tuning curves whose peaks exceed the response to the preferred flat velocity profile, we have ruled out this possible artifact as a general explanation for our findings.

Discussion

In the experiments presented here, we have investigated the response of area MT neurons in the awake behaving monkey to elementary velocity gradients. We have demonstrated that a substantial proportion (60%) of our sample of MT neurons respond to such gradients in a systematically tuned manner. This enables area MT to encode the shape and orientation of velocity gradients in the visual world. Furthermore, this property makes MT cells well suited to provide the building blocks for the more complex receptive-field properties encountered in area MST.

The single-lobed tuning that we found in our study suggests that the variable used in these experiments, i.e. the orientation of velocity gradients, is indeed encoded in the firing rate of MT neurons. At the same time, MT neurons are tuned to other stimulus parameters like direction of motion and stereoscopic disparity and are thus likely to be involved in the perception of those parameters too. This behavior allows MT cells to contribute to the analysis of a variety of motion stimuli but it also restricts the modulation that can be achieved by varying a single dimension. The modulation we observed, i.e. the difference in a response of a given cell for the best and worst velocity gradient, is small compared to the modulation that can be achieved with stimuli which vary in their direction of motion.

There are several possible reasons for this shallow gradient tuning:

1. The simple gradients we used did not adequately match the preferred gradients of the neurons. This would be analogous to taking a cell's direction tuning curve at an inadequate speed.
2. The response of MT neurons is dominated by the direction of motion of a pattern rather than by the shape of its velocity profile.
3. The neurons in our study were responding close to saturation, given that our stimuli were modulations around the most effective flat velocity profile.
4. The neurons encode surfaces in depth and our stimuli, lacking stereoscopic depth, were thus less than optimal.

The first argument is not likely to be correct since we could drive the cells we encountered very well, even exceeding the firing rate to the best classical stimulus. But given that MT neurons are selective for stimuli in two one-dimensional stimulus spaces, namely the direction of motion space and our deformation space, it might be more appropriate to describe the response of an MT cell in a multidimensional space that not only
Fig. 2. Example of a gradient tuned cell: (A) Response of the cell to random dot patterns moving in eight different directions spaced 45 deg apart. The preferred direction of this cell was upward to the left. (bg: background rate, i.e. the firing rate of the neuron in the absence of a stimulus). (B) Speed tuning curve of the neuron using patterns moving in the preferred direction. The preferred speed of this neuron was 8 deg/s. (C) Gradient tuning of the cell using the eight gradients presented in Fig. 1 using the preferred direction as determined in (A). The average speed of all patterns was 8 deg/s, i.e. the preferred speed. The dashed circle represents the response of the neuron to the flat velocity profile moving at the preferred speed. The arrow is an estimate of the preferred direction of the cell in the deformation space. The $P$ value denotes the significance of the difference between the cell's response to the best gradient (the one closest to the arrow) and the response to the preferred flat velocity profile.
accounts for selectivity for direction of motion and velocity gradients but also stereoscopic disparity.

The role of MT in 3-D shape perception, suggested by our results, fits well with the finding that individual MT neurons have both direction and stereoscopic tuning with complex interactions designed to disambiguate motion stimuli (Bradley et al., 1995). While we did not investigate stereoscopic tuning in this study, it is possible that our velocity gradient tuned cells show a depth tuning that enhances the response modulation caused by the various motion gradients. Such a combination of depth-from-motion and depth-from-binocular-disparity signals might underlie the perceptual similarity of these two cues to 3-D shape (Rogers & Graham, 1982).

Many neurons in area MT show an opponent center-surround organization (Allman et al., 1985) that might influence a cell's response to a particular gradient that straddles the bor-

Fig. 3. Summary of the cells: Percentages for the various cells that were tested with the velocity gradients. Each cell type (see text for details) is presented with the response profiles from two cells.
nder between the center and the surround of the receptive field. This possibility has been used in a model by Dobbins et al. (1990) to obtain estimates of the optic flow field and its spatial and temporal variation. In our recordings, we avoided such stimulus placements by limiting our stimuli to the classical receptive fields (the center) of all neurons we recorded from. Under natural viewing conditions, stimuli will often overlap the border between the center and surround of the receptive field though. Depending on the respective preferred speeds and direction in the center and the surround, this could lead to a much stronger modulation of the response of the cell to various velocity gradients. In fact, nonsymmetric surrounds such as the ones described by Xiao et al. (1994) seem particularly well suited to enhance the gradient selectivity of the classical receptive field.

Our study does not address the mechanisms responsible for the gradient tuning we observed. Such tuning could be achieved by cells whose gradient selectivity is the result of a mosaic of systematically varying preferred velocity. Such cells would not show the property of position invariance that has been observed in MST receptive fields (Lagae et al., 1994; Graziano et al., 1994) and requires more complicated neural wiring. Its existence in area MT could be tested by a study that uses small moving patches to determine the preferred speed at different locations within the receptive field of a given MT cell.

References

ALLMAN, J., MIEZIN, F. & MCGUINESS, E. (1985). Direction- and velocity-specific responses from beyond the classical receptive field in the middle temporal visual area (MT). Perception 14, 105-126.

ANDERSEN, G. J. (1989). Perception of three-dimensional structure from optical flow without locally smooth velocity. Journal of Experimental Psychology: Human Perception and Performance 15, 363-371.

ANDERSEN, R. A., & SIEGEL, R. M. (1990). Motion processing in the primate cortex. In Signal and Sense: Local and Global Order in Perceptual Maps, ed. Edelman, G. M., GALL, W. E. & COWAN, W. M. pp. 163-184, New York: John Wiley.

BRAUNSTEIN, M. L. & ANDERSEN, G. J. (1984). Shape and depth perception from parallel projections of three-dimensional motion. Journal of Experimental Psychology: Human Perception and Performance 10, 749-759.

CROWELL, J. A. & BANKS, M. S. (1993). Perceiving heading with different retinal regions and types of optic flow. Perception & Psychophysics 53, 325-337.

DOBINS, A., ZUCKER, S. W. & CYNADER, M. S. (1990). A mean field model of optic flow estimation by MT neurons. Society for Neuroscience Abstracts 16, 6.

DUFFY, C. J. & WURTZ, R. H. (1991a). Sensitivity of MST neurons to optic flow stimuli. I. A continuum of response selectivity to large-field stimuli. Journal of Neuroscience 11, 1329-1345.

DUFFY, C. J. & WURTZ, R. H. (1991b). Sensitivity of MST neurons to optic flow stimuli. II. Mechanisms of response selectivity revealed by small-field stimuli. Journal of Neuroscience 11, 1346-1359.

GALLANT, J. L., BRAUN, J. & VAN ESEN, D. C. (1993). Selectivity for polar, hyperbolic, and Cartesian gratings in macaque visual cortex. Science 259, 100-103.

GIBSON, J. J. (1950). The Perception of the Visual World. Boston, Massachusetts: Houghton Mifflin.

GOLOMB, B., ANDERSEN, R. A., NAKAYAMA, K., MACLEOD, D. I. A. & WONG, A. (1985). Visual thresholds for shearing motion in monkey and man. Vision Research 25, 813-820.

GRAZIANO, M. S. A., ANDERSEN, R. A. & SNOWDEN, R. J. (1994). Tuning of MST neurons to spiral motions. Journal of Neuroscience 14, 54-67.

HARRIS, M., FREEMAN, T. & HUGHES, J. (1992). Retinal speed gradients and the perception of surface slant. Vision Research 32, 587-590.

HUSAIN, M., TREUE, S. & ANDERSEN, R. A. (1989). Surface interpolation in 3-D structure-from-motion perception. Neural Computation 1, 324-333.

KOENDERINK, J. J. (1986). Optic flow. Vision Research 26, 161-180.

KOENDERINK, J. J. & VAN DOORN, A. J. (1981). Exospecific component of the motion parallax field. Journal of the Optical Society of America 71, 953-957.

LAGAE, L., MAES, H., RAIGUEL, S., ZIAO, D. K. & ORBAN, G. A. (1994). Responses of macaque STS neurons to optic flow components: A comparison of area MT and MST. Journal of Neurophysiology 71, 1597-1625.

LAND, M. S., DOSHER, B. A., SPERLING, G. & PERKINS, M. E. (1991). The kinetic depth effect and optic flow II. First- and second-order motion. Vision Research 31, 859-876.

LONGUET-HIGGINS, H. C. & PRADZNY, K. (1980). The interpretation of a moving retinal image. Proceedings of the Royal Society B (London) 208, 385-397.

MACNAY, D. M. (1961). Visual effects of non-redundant stimulation. Nature 192, 739-740.

NAKAYAMA, K. (1981). Differential motion hyperacuity under conditions of common image motion. Vision Research 21, 1475-1482.

NAKAYAMA, K., SILVERMAN, G., MACLEOD, D. I. A. & MULGANN, J. (1985). Sensitivity to shearing and compressive motion in random dots. Perception 14, 97-241.

ORBAN, G. A., LAGAE, L., VERRI, A., RAIGUEL, S., ZIAO, D., MAES, H. & TORRE, V. (1992). First order analysis of optical flow in monkey brain. Proceedings of the National Academy of Sciences of the U.S.A. 89, 2595-2599.

REGAN, D. (1985). Visual flow and direction of locomotion. Science 227, 1064-1065.

REGAN, D. (1986). Visual processing of four kinds of relative motion. Vision Research 26, 127-145.

REGAN, D. & BEVERLEY, K. I. (1981). How do we avoid confusing the direction we are looking and the direction we are moving? Science 215, 194-196.

REGAN, D. & BEVERLEY, K. I. (1985). Visual responses to vorticity and the neural analysis of optic flow. Journal of the Optical Society of America 2, 280-283.

REGAN, D., ERKELENS, C. J. & COLLEWIN, H. (1986). Necessary conditions for the perception of motion in depth. Investigative Ophthalmology and Visual Science 27, 584-596.

RIEGER, J. H. & LAWTON, D. T. (1985). Processing differential image motion. Journal of the Optical Society of America A 2, 354-360.

ROBINSON, D. A. (1963). A method of measuring eye movement using a sceral search coil in a magnetic field. IEEE Transactions of Biomedical Engineering 10, 137-145.

ROGERS, B. J. & GRAHAM, M. (1979). Motion parallax as an independent cue for depth perception. Perception 8, 125-134.

ROGERS, B. & GRAHAM, M. (1982). Similarities between motion parallax and stereopsis in human depth perception. Vision Research 22, 261-270.

ROYDEN, C. S., CROWELL, J. A. & BANKS, M. S. (1994). Estimating heading during eye movements. Vision Research 34, 3197-3214.

SAITO, H., YUKIE, M., TANAKA, K., HIkosaka, K., FUKADA, Y. & IwAI, E. (1986). Integration of direction signals of image motion in the superior temporal sulcus of the macaque monkey. Journal of Neuroscience 6, 145-157.

SIEGEL, R. M. & ANDERSEN, R. A. (1986). Motion perceptual deficits following ibotenic acid lesions of the middle temporal area in the behaving rhesus monkey. Society for Neuroscience Abstracts 12, 1183.

SIEGEL, R. M. & ANDERSEN, R. A. (1988). Perception of three-dimensional structure from motion in monkey and man. Nature 331, 259-261.

SNOWDEN, R. J., TREUE, S., ERICKSON, R. E. & ANDERSEN, R. A. (1991). The response of area MT and V1 neurons to transparent motion. Journal of Neuroscience 11, 2768-2785.

TANAKA, K., FUKADA, Y. & SAITO, H. (1989). Underlying mechanisms of the response specificity of expansion/contraction, and rotation cells clustered in the dorsal part of the medial superior temporal area of the macaque monkey. Journal of Neurophysiology 62, 642-656.

TANAKA, K., HIKOSAKA, K., SAITO, H., YUKIE, M., FUKADA, Y. & IwAI, E. (1986). Analysis of local and wide-field movements in the super-
terior temporal visual areas of the macaque monkey. *Journal of Neuroscience*. 6, 134–144.

Tanaka, K. & Saito, H. (1989). Analysis of motion of the visual field by direction, expansion/contraction, and rotation cells clustered in the dorsal part of the medial superior temporal area of the macaque monkey. *Journal of Neurophysiology* 62, 626–641.

Treue, S., Husain, M. & Andersen, R. (1991). Human perception of structure from motion. *Vision Research* 31, 59–75.

Warren, W. H. & Hannon, D. J. (1988). Direction of self-motion is perceived from optical flow. *Nature* 336, 162–163.

Warren, W. H., Morrie, M. W. & Kalish, M. (1988). Perception of translational heading from optical flow. *Journal of Experimental Psychology: Human Perception and Performance* 14, 646–660.

Xiao, D. K., Marcar, V. L., Raiguel, S. E. & Orban, G. A. (1994). Does the surround really surround the classical receptive field of macaque MT cells? *Society for Neuroscience Abstracts* 20, 773.