

A Theory of Geometric Constraints on Neural Activity for Natural Three-Dimensional Movement

Kechen Zhang¹ and Terrence J. Sejnowski^{1,2}

¹Howard Hughes Medical Institute, Computational Neurobiology Laboratory, The Salk Institute for Biological Studies, La Jolla, California 92037, and ²Department of Biology, University of California, San Diego, La Jolla, California 92093

Although the orientation of an arm in space or the static view of an object may be represented by a population of neurons in complex ways, how these variables change with movement often follows simple linear rules, reflecting the underlying geometric constraints in the physical world. A theoretical analysis is presented for how such constraints affect the average firing rates of sensory and motor neurons during natural movements with low degrees of freedom, such as a limb movement and rigid object motion. When applied to nonrigid reaching arm movements, the linear theory accounts for cosine directional tuning with linear speed modulation, predicts a curl-free spatial distribution of preferred directions, and also explains why the instantaneous motion of the hand can be recovered from the neural population activity. For three-dimensional motion of a

rigid object, the theory predicts that, to a first approximation, the response of a sensory neuron should have a preferred translational direction and a preferred rotation axis in space, both with cosine tuning functions modulated multiplicatively by speed and angular speed, respectively. Some known tuning properties of motion-sensitive neurons follow as special cases. Acceleration tuning and nonlinear speed modulation are considered in an extension of the linear theory. This general approach provides a principled method to derive mechanism-insensitive neuronal properties by exploiting the inherently low dimensionality of natural movements.

Key words: 3-D object; cortical representation; visual cortex; tuning curve; motor system; reaching movement; speed modulation; potential function; gradient field; zero curl

For natural movements, such as the motion of a rigid object or an active limb movement, many sensory receptors or muscles are involved, but the actual degrees of freedom are low because of geometric constraints in the physical world. For example, as illustrated in Figure 1, the rotation of an object alters many visual cues. How these cues vary in time is not arbitrary but is fully determined by the rigid motion, which has only 6 degrees of freedom. As a consequence, neuronal activity reflecting such natural movements also is likely to be highly constrained and to have only a few degrees of freedom.

This paper presents a theoretical analysis of how neuronal activity correlated with natural movements might be constrained by geometry. The basic theory, although essentially linear, can account for several key features of diverse neurophysiological results and generates strong predictions that are testable with current experimental techniques.

An emerging principle from this analysis is that neuronal activity tuned to movement often obeys simple generic rules as a first approximation, insensitive to the exact sensory or motor variables that are encoded and the exact computational interpretation. Such generic tuning properties are *mechanism insensitive* because they are better described as reflecting the underlying geometric constraints on movements rather than the actual computational mechanisms. This simplicity arises when sensory or motor variables represent changes in time rather than static

values. In the example shown in Figure 1, the viewpoint was fixed and the object was rotated systematically around different axes. The focus is on how neuronal responses depend on the rotation axis in three-dimensional space, given approximately the same view of the object. It is possible to derive a simple cosine tuning rule for the rotation axis, although various visual cues may depend on the static geometrical orientation of the object in complex ways. Three-dimensional object motion is a specific example; the same principles also apply to several other biological systems, including nonrigid arm movement.

DIRECTIONAL TUNING FOR ARM MOVEMENT

Although the visual and the motor examples share similar mechanism-insensitive properties, the reaching arm movement has a simpler mathematical description and more supporting experimental results and will be considered first.

Ubiquity of cosine tuning

A directional tuning curve describes how the mean firing rate of a neuron depends on the reaching direction of the hand. As illustrated in Figure 2, broad cosine-like tuning curves are very typical in many areas of the motor system of monkeys, including the primary motor cortex (Georgopoulos et al., 1986), premotor cortex (Caminiti et al., 1991), parietal cortex (Kalaska et al., 1990), cerebellum (Fortier et al., 1989), basal ganglia (Turner and Anderson, 1997), and somatosensory cortex (Cohen et al., 1994; Prud'homme and Kalaska, 1994). Although the examples shown in Figure 2 are two-dimensional, cosine tuning holds as well for three-dimensional reaching movement (Georgopoulos et al., 1986; Schwartz et al., 1988).

Received Oct. 16, 1998; revised Dec. 14, 1998; accepted Jan. 27, 1999.

We are grateful to T. D. Albright, G. T. Buračas, G. E. Hinton, R. J. Krauzlis, K. D. Miller, A. B. Schwartz, M. I. Sereno, M. P. Stryker, R. S. Turner, D. Zipser, and two anonymous reviewers for helpful comments on the analysis presented here.

Correspondence should be addressed to Dr. Terrence Sejnowski, Computational Neurobiology Lab, The Salk Institute, La Jolla, CA 92037.

Copyright © 1999 Society for Neuroscience 0270-6474/99/193122-24\$05.00/0

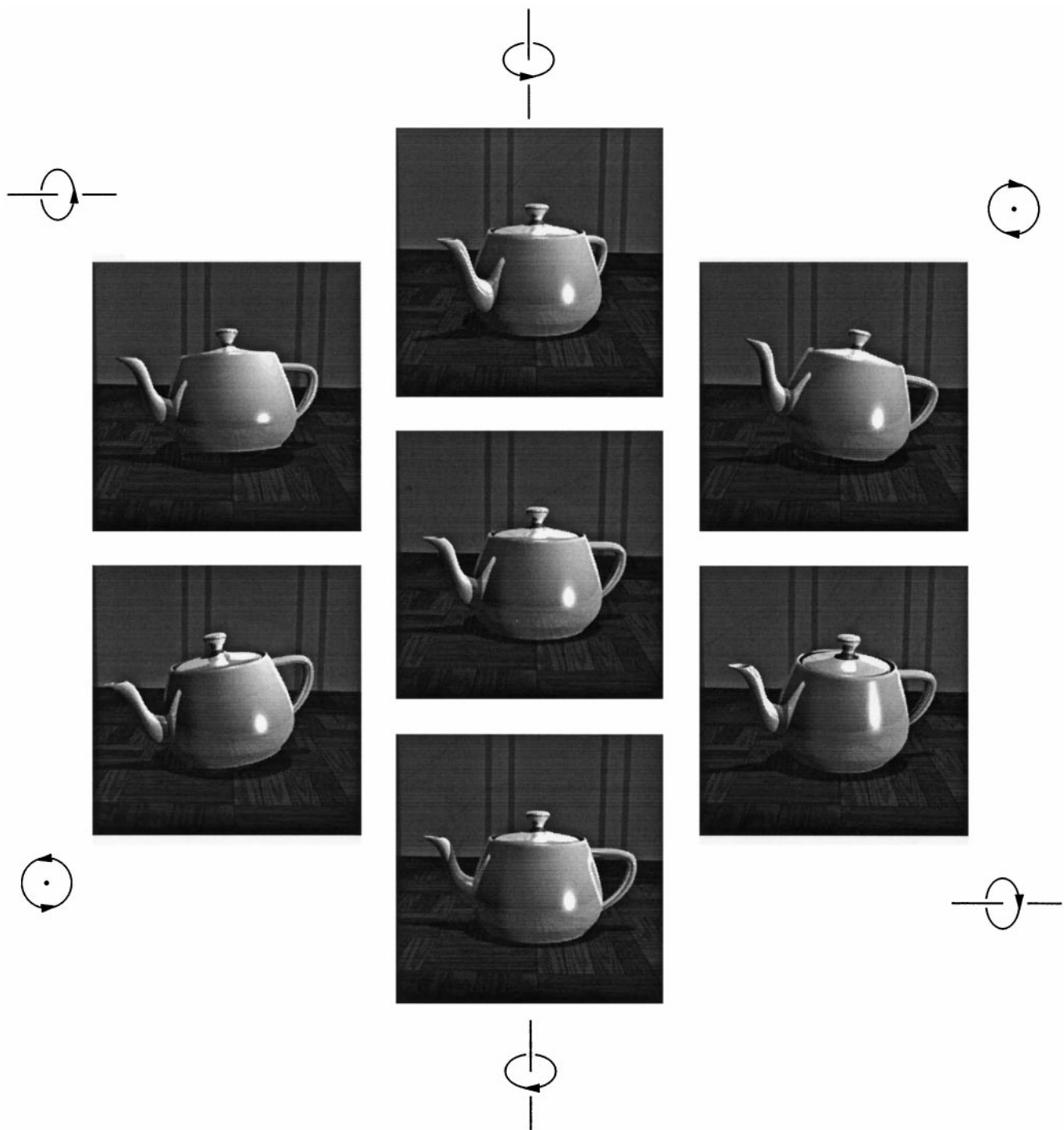


Figure 1. Axis of rotation determines how the view of an object changes instantaneously, along with various visual cues, such as shading, shadow, mirror reflection, glare, and occlusion. Rigid geometry predicts that the response of a motion-sensitive neuron, to a first approximation, should have a preferred rotation axis in three-dimensional space with cosine tuning function and linear angular speed modulation, regardless of the exact cues used and the exact computational mechanisms involved.

The ubiquity of cosine tuning is a hint that this property is generic and insensitive to the exact computational function of these neurons. For example, coding of muscle shortening rate is one theoretical mechanism that can generate cosine tuning (Mussa-Ivaldi, 1988). As another example, many somatosensory cortical cells related to reaching had cosine directional tuning, probably because of the geometry of mechanical deformation of

the skin during arm movement (Cohen et al., 1994; Prud'homme and Kalaska, 1994). Because a cosine tuning function implies a dot product between a fixed preferred direction and the actual reaching direction (Georgopoulos et al., 1986), cosine tuning by itself suggests a linear relation with reaching direction (Sanger, 1994), which could arise as an approximation to the activity in a nonlinear recurrent network (Moody and Zipser, 1998). There-

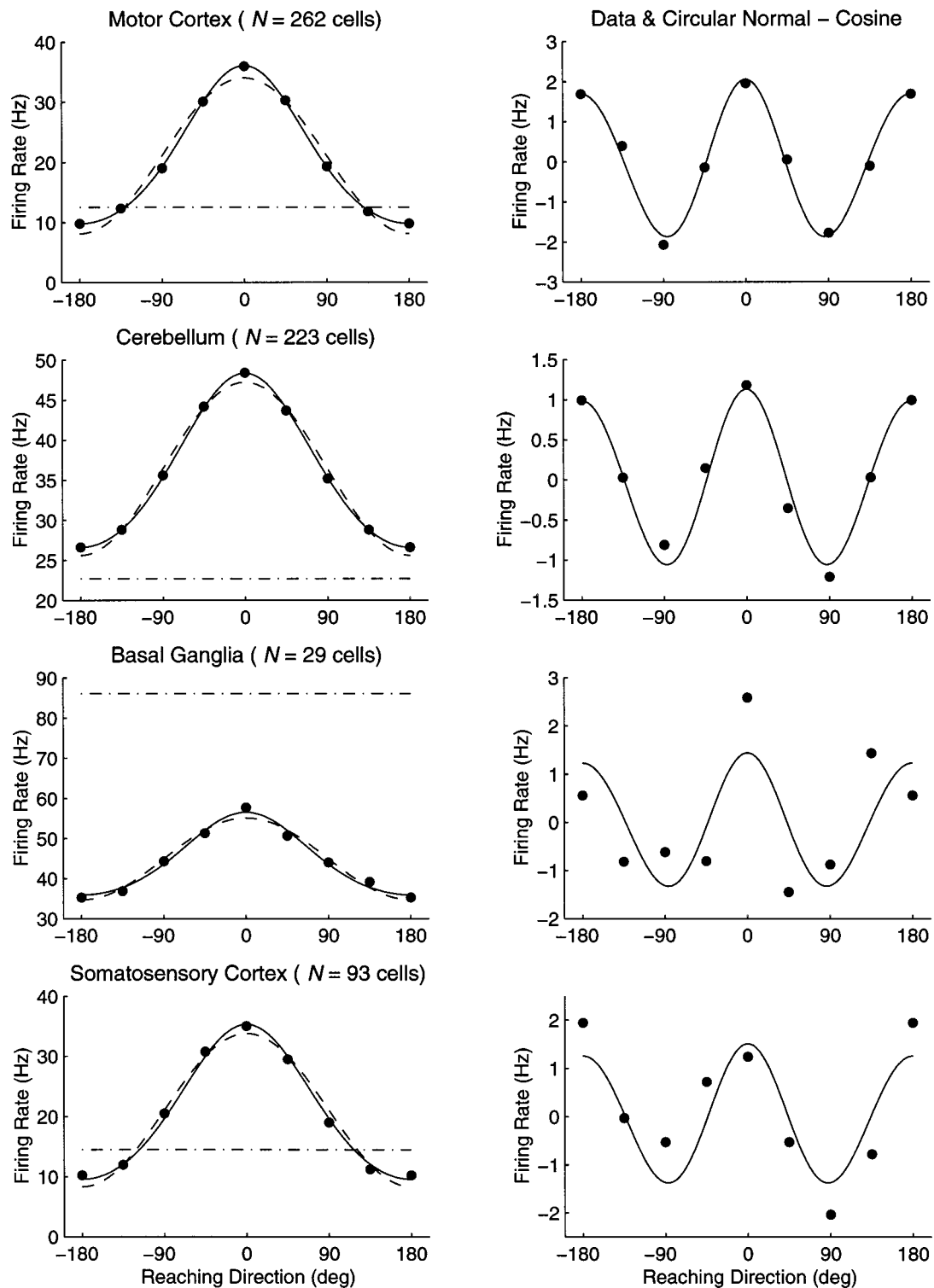


Figure 2. Cosine tuning to hand movement direction is very common in monkey motor system, here showing examples of average tuning curves in two-dimensional reaching tasks, with preferred direction taken as 0° . *Left column*, Circular normal functions (*solid curves*) fit the data (\bullet) slightly better and are slightly narrower than cosine functions (*dashed curves*). *Horizontal lines* indicate background firing rates without movement. *Right column*, Data and the circular normal functions after subtracting the cosine functions. Data from motor cortex (M1) and cerebellum (Purkinje cells plus deep nuclei) are from Figure 2 in Fortier et al. (1993), basal ganglia data (GPe) are from Figure 8B (decrease type) in Turner and Anderson (1997), and somatosensory cortex data (S1) are from Figure 11A (no load case) in Prud'homme and Kalaska (1994), with permission.

fore, cosine tuning curves should be common in a theoretical model that is approximately linear.

Basic theory

In this section we derive a general tuning rule for motor neurons and then discuss its basic properties. This example illustrates what is meant by mechanism-insensitive properties and the general theoretical argument based on geometric constraints.

Consider stereotyped reaching movement in which the configuration of the whole arm is determined completely by the hand position (x, y, z) in space. In other words, such movements have only 3 degrees of freedom. Assume that *the mean firing rate of a neuron relative to baseline is proportional to the time derivative of an unknown smooth function of hand position in space*. In other words:

$$f = f_0 + \frac{d}{dt} \Phi(x, y, z), \quad (1)$$

where f is the firing rate, f_0 is the baseline rate, and Φ is an arbitrary function of the hand position (x, y, z) . A possible small time difference between the neural activity and the arm movement may also be included, as appropriate.

The function $\Phi(x, y, z)$ could have any form and could include any function of arm configuration, such as muscle length, joint angles, or any combination of those. Mussa-Ivaldi (1988) first used muscle length to demonstrate the appearance of cosine tuning in a two-dimensional situation and pointed out that the argument could be generalized to include other muscle variables. This interesting example illustrates how cosine tuning property might emerge from some simple assumptions. The assumption in Equation 1 is more general and the formalism is simpler than that of Mussa-Ivaldi (1988) because joint angles are no longer used as intermediate variables in the derivation. This makes interpretation easier and more flexible and the curl-free condition more apparent (see below). The precise interpretation of Φ is not the focus of this paper; the only requirement is that it be a function fully determined by the hand position in the three-dimensional space.

We emphasize that although Equation 1 uses hand position as the only free variables, this does *not* require that the neuron must directly encode the hand position or end-point in particular or kinematic variables in general. Stereotypical reaching movements have only 3 degrees of freedom and can be conveniently parameterized by the hand position (x, y, z) , although other parameters can also be used without affecting the final conclusion (see below and Appendix A). A neuron related to reaching arm movement should be sensitive to changes of arm posture, which can always be expressed equivalently as changes in some functions of the hand position (x, y, z) . The simplest estimate of such changes is the first temporal derivative given in Equation 1. In other words, the above assumption only postulates a general dependence of the firing rate of a neuron on changing arm posture as a first approximation, regardless of which parameters are encoded and how they are encoded.

The assumption in Equation 1 implies that the mean firing rate of a neuron should follow the tuning rule:

$$f = f_0 + \mathbf{p} \cdot \mathbf{v}, \quad (2)$$

where $\mathbf{v} = (\dot{x}, \dot{y}, \dot{z})$ is the instantaneous reaching velocity of the hand, and the vector \mathbf{p} is the *preferred reaching direction*, given by:

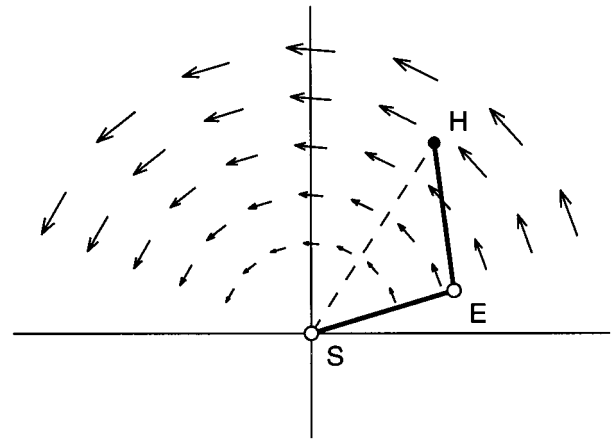


Figure 3. The preferred direction field of a hypothetical neuron that violates the curl-free condition in a planar reaching task. For each hand position, the preferred direction of this neuron is always perpendicular to the straight line from the hand (H) to the shoulder (S), and the length of the vector is proportional to the distance of HS . This vector field has constant non-zero curl everywhere in the work space. The gradient theory does not allow the existence of such a neuron.

$$\mathbf{p} = (p_x, p_y, p_z) = \nabla \Phi = \left(\frac{\partial \Phi}{\partial x}, \frac{\partial \Phi}{\partial y}, \frac{\partial \Phi}{\partial z} \right). \quad (3)$$

The derivation of this result follows immediately from the chain rule:

$$\frac{d\Phi}{dt} = \frac{\partial \Phi}{\partial x} \dot{x} + \frac{\partial \Phi}{\partial y} \dot{y} + \frac{\partial \Phi}{\partial z} \dot{z} = \mathbf{p} \cdot \mathbf{v}. \quad (4)$$

For hand movements starting from the same position (x, y, z) in space, the tuning rule in Equation 2 implies cosine directional tuning and linear speed modulation (see Eq. 12). The preferred direction vector $\mathbf{p} = \mathbf{p}(x, y, z)$ of the neuron may depend on the starting hand position. It can be regarded as a constant vector when the hand is close to its starting position.

For hand movements starting from different positions, the preferred direction vector may vary with the starting hand position (x, y, z) and thus can be visualized as a vector field (Caminiti et al., 1990; Moody and Zipser, 1998). It follows from the gradient formula in Equation 3 that this vector field of preferred direction must have zero curl:

$$\begin{aligned} \text{curl } \mathbf{p} &= \left(\frac{\partial p_z}{\partial y} - \frac{\partial p_y}{\partial z}, \frac{\partial p_x}{\partial z} - \frac{\partial p_z}{\partial x}, \frac{\partial p_y}{\partial x} - \frac{\partial p_x}{\partial y} \right) \\ &= \text{curl } \nabla \Phi = 0, \end{aligned} \quad (5)$$

because of the equality of mixed second partial derivatives of Φ . This means that the components of the preferred direction cannot vary arbitrarily with the starting hand position. An equivalent integral formulation of the curl-free condition is that the path integral of \mathbf{p} vanishes along any closed curve in three-dimensional space:

$$\oint \mathbf{p} \cdot d\mathbf{l} = 0, \quad (6)$$

with $d\mathbf{l} = (dx, dy, dz)$, assuming that there are no singularities in the vector field. This constrains how the preferred direction of a

neuron should vary with the starting hand position. Any distribution with non-zero curl can be ruled out (Fig. 3).

Human eyes are not reliable at judging whether a vector field is curl-free (see Fig. 10), so numerical computation is needed (Mussa-Ivaldi et al., 1985; Giszter et al., 1993). See Appendix B for more discussion. A vector field is curl-free if and only if it can be generated as the gradient of a potential function. A more intuitive interpretation of the curl-free condition is that when a vector field is regarded as the velocity field of a fluid, there is no net circulation along any closed path in space.

Under the curl-free condition, the net spike count (integration of the firing rate with respect to baseline over time) can be used to recover the value of the unknown potential function:

$$\Phi(x, y, z) - \Phi(x_0, y_0, z_0) = \int_0^T (f - f_0) dt, \quad (7)$$

where the integral depends only on the initial hand position (x_0, y_0, z_0) at time 0 and the final position (x, y, z) at time T , not on the exact trajectory of hand movement. For each hand position, the firing rate is the largest when the hand moves along the local gradient of the potential function, which defines \mathbf{p} .

Baseline firing rate

The theory in the preceding section does not constrain the baseline firing rate f_0 , which needs to be considered separately. By definition, the baseline firing rate is independent of the reaching direction, but it may be modulated by several other factors. For example, in the motor cortex, Kettner et al. (1988) have reported that the linear formula:

$$f_0 = a_0 + a_1x + a_2y + a_3z, \quad (8)$$

approximately described the baseline firing rate while the hand was held fixed at position (x, y, z) in the three-dimensional work space, where a_0, a_1, a_2, a_3 are constant coefficients. For reaching at speed v , a more general linear formula for the baseline firing rate is:

$$f_0 = a_0 + a_1x + a_2y + a_3z + av, \quad (9)$$

where the coefficients a_0, a_1, a_2, a_3, a are independent of the hand position (x, y, z) and the speed v , but may vary with task conditions. For instance, the baseline firing rate when the hand is held still (Fig. 2, *horizontal lines*) differs from the baseline rate defined as the average of the cosine curve during reaching. Moran and Schwartz (1999) showed that a linear speed term for baseline rate should be included in the fitting formula, although their analysis used the square root of firing rate instead of the raw firing rate. Indirect evidence for a linear speed term in baseline rate is provided by the linear effect of reaching distance (see below).

Note that in Equation 9, the baseline firing rate contains information about both the static hand position (x, y, z) and its speed v . As shown by Kettner et al. (1988), the spatial gradient of the spontaneous firing rate for static hand position tends to be consistent with the preferred direction of the same neuron. In the current theory, this means that the preferred direction $\mathbf{p} = \nabla\Phi$ tends to point in the same direction as the vector (a_1, a_2, a_3) in Equation 9. Therefore, if the potential function $\Phi = \Phi(x, y, z)$ can

be approximated as a linear function in x, y, z , we can replace Equation 9 by:

$$f_0 = a_0 + k\Phi(x, y, z) + av, \quad (10)$$

where k is a constant coefficient. In this case, the overall firing rate of a neuron that obeys the basic tuning rule in Equation 2 would convey two pieces of information: the baseline firing rate f_0 would represent the static value of the potential function Φ , and the directionally tuned part $\mathbf{p} \cdot \mathbf{v}$ would represent the spatial gradient of the same potential function.

Linear theory without gradient

If we simply postulate that the firing rate of a neuron is linearly related to the components of reaching velocity $\mathbf{v} = (v_x, v_y, v_z)$, we would have the same tuning rule:

$$f = f_0 + p_x\dot{x} + p_y\dot{y} + p_z\dot{z} = f_0 + \mathbf{p} \cdot \mathbf{v}, \quad (11)$$

where the components of the preferred direction, $(p_x, p_y, p_z) \equiv \mathbf{p}$, are three arbitrary functions of the hand position (x, y, z) . For a single starting hand position, this tuning rule is locally indistinguishable from the prediction of the gradient theory. The difference is that now the preferred direction field is not required to be the gradient of any potential function so that its global distribution in hand position space is not constrained at all. In other words, this vector field need not be curl-free. The nongradient theory is more general, allowing a circular distribution of the preferred directions as in Figure 3. The necessary and sufficient condition for the gradient theory to be true is that the preferred direction field is curl-free. The existing data cannot distinguish the two theories (see discussion below and Appendix B).

Comparison with experimental results

Data from a wide range of motor-related brain areas largely confirm the tuning rule in Equation 2 as a reasonable approximation, together with its various ramifications as follows. Theoretical predictions such as the curl-free distribution remain to be tested.

Cosine directional tuning and multiplicatively linear speed modulation

The tuning rule in Equation 2 captures two main effects: cosine directional tuning and multiplicatively linear speed modulation, as clearly seen in its equivalent form:

$$f = f_0 + pv \cos \alpha, \quad (12)$$

where $v = |\mathbf{v}|$ is the reaching speed, the proportional factor $p = |\mathbf{p}|$ is length of the preferred direction vector \mathbf{p} , and α is the angle between the instantaneous reaching direction and the preferred direction. Because the hand trajectory is approximately straight in normal reaching, the instantaneous velocity \mathbf{v} is a vector that points in the same direction as the reaching direction. If a tuning function is cosine in three-dimensional space, it must also be cosine in any two-dimensional subspace, as in the examples in Figure 2.

A cosine function is a good approximation to the directional tuning data, although a circular normal function (Eq. A17), with one more free parameter, tends to fit the data slightly better (Fig. 2). The residual can be roughly accounted for by an additional Fourier term, $\cos 2\alpha$, with an amplitude less than $\sim 10\%$ of that of the original term, $\cos \alpha$ (see further discussion in Appendix A).

The speed modulation effect predicted by Equation 12 is multiplicative; that is, the firing rate should be higher for faster

reaching speed without affecting the shape of the cosine tuning function. This is approximately true as shown by Moran and Schwartz (1999), who, however, used the square root of firing rate in analysis so that the linearity of speed modulation on raw firing rate was not directly quantified. Indirect evidence for linear speed modulation includes trajectory reconstruction and the curvature power law (see below).

Neuronal population vector

Suppose the firing rate of each neuron i in a population ($i = 1, 2, \dots, N$) follows the same tuning rule as considered above:

$$f_i = f_{i0} + \mathbf{p}_i \cdot \mathbf{v}. \quad (13)$$

The population vector \mathbf{u} is defined as the vector sum of the preferred directions \mathbf{p}_i weighted by firing rates relative to baselines (Georgopoulos et al., 1986):

$$\mathbf{u} = \sum_{i=1}^N (f_i - f_{i0}) \mathbf{p}_i = \sum_{i=1}^N \mathbf{p}_i (\mathbf{p}_i \cdot \mathbf{v}), \quad (14)$$

where in the second step, Equation 13 is used. For the population vector \mathbf{u} to be proportional to the true velocity \mathbf{v} , namely:

$$\mathbf{u} = \lambda \mathbf{v}, \quad (15)$$

the necessary and sufficient condition is that the preferred directions satisfy:

$$\sum_{i=1}^N \mathbf{p}_i \mathbf{p}_i^T = \lambda \mathbf{I}, \quad (16)$$

where λ is an arbitrary constant, \mathbf{I} is 3×3 identity matrix, each \mathbf{p}_i is a column vector, and \mathbf{p}_i^T is a row vector (Mussa-Ivaldi, 1988; Gaál, 1993; Salinas and Abbott, 1994; Sanger, 1994). In particular, when \mathbf{p}_i are distributed uniformly, as is roughly true for cells in motor cortex (Georgopoulos et al., 1988), the condition in Equation 16 is satisfied so that Equation 15 follows as a consequence. Then the population vector approximates the reaching direction and reaching velocity (Moran and Schwartz, 1999).

Trajectory reconstruction

One implication of Equation 15 is that integration over the population over time can reconstruct the hand trajectory, up to a scaling constant:

$$\mathbf{r}(t) = \mathbf{r}(0) + \frac{1}{\lambda} \int_0^t \mathbf{u}(\tau) d\tau, \quad (17)$$

where $\mathbf{r}(t)$ is hand position at time t . This is consistent with the finding that adding up the population vector head-to-tail approximately reproduced the shape of the hand trajectory (Schwartz, 1993, 1994), because head-to-tail addition is a discrete approximation to the continuous vector integration.

Curvature power law

While drawing, the hand moves more slowly when the trajectory is more highly curved, and obeys a power law:

$$\omega = B\kappa^{2/3}, \quad (18)$$

where ω is instantaneous angular velocity with respect to an instantaneous center determined by the local curvature κ of the trajectory, and B is a constant (Lacquaniti et al., 1983). Schwartz (1994) showed that the changing direction of the population vector of cells in motor cortex of monkeys followed the same power law during drawing. This is consistent with Equation 15, which requires that the population vector \mathbf{u} be proportional to the instantaneous hand velocity \mathbf{v} , up to a possible time difference. This form of the power law involves only the direction of population vector \mathbf{u} . To test its length $u = |\mathbf{u}|$ or the linearity of firing rate modulation by reaching speed, one may use the equivalent form of the power law:

$$v = Br^{1/3}, \quad (19)$$

where $v = r\omega$ is the hand speed and $r = 1/\kappa$ is the local radius for the curvature of the trajectory. The length of the population vector u is proportional to the hand speed v if and only if the population vector follows the same power law in Equation 19.

Reaching distance

Fu et al. (1993) reported a nearly linear correlation between firing rates of cells in motor cortex and reaching distance. Although this result was somewhat confounded by faster reaching for longer distances, it raises the question of the general effect of reaching distance. A linear distance effect would be consistent with the basic model in Equation 2, which implies that:

$$\int_0^T (f - f_0) dt = \int_0^T \mathbf{p} \cdot \mathbf{v} dt = \mathbf{p} \cdot \mathbf{d}, \quad (20)$$

where vector $\mathbf{d} = \mathbf{r}(T) - \mathbf{r}(0)$ is the final displacement from the starting position $\mathbf{r}(0)$, assuming the preferred direction \mathbf{p} is approximately constant along movement trajectory. The dot product $\mathbf{p} \cdot \mathbf{d}$ implies a linear relation between the reaching distance and the total spike count above baseline, together with a cosine directional tuning, regardless of the exact time course of hand velocity.

Note that in Equation 20 the baseline rate f_0 has been subtracted. Because the baseline rate itself may contain a linear speed component as in Equation 9 (Moran and Schwartz, 1999), its contribution to total spike count should be:

$$\int_0^T f_0 dt = \int_0^T (a_0 + av) dt = a_0 T + a|\mathbf{d}|, \quad (21)$$

where, for simplicity, $a_1 = a_2 = a_3 = 0$ has been assumed to ignore the effect of static hand position. Because the last term is proportional to the reaching distance $|\mathbf{d}|$ but independent of the reaching direction, it might account for the observation that the modulation of overall firing rates by reaching distance was often linear but insensitive to the reaching direction (Fu et al., 1993; Turner and Anderson, 1997).

Curl-free distribution of preferred direction

Caminiti et al. (1990, 1991) reported that the preferred direction of a motor cortical neuron often varied with the starting point of hand movement. This is allowed by the gradient theory, provided that this vector field is curl-free, according to Equation 5 or 6. A constant preferred direction field is always allowed because it has zero curl. The curl-free condition constrains how the preferred

direction of a neuron may vary in different parts of space. For example, it rules out the possibility of any circular arrangement of the preferred directions, such as that in the two-joint planar arm example shown in Figure 3. The existing data do not include enough points to compute the curl (see Appendix B). Further experiments would be needed to test whether the prediction of the gradient theory is correct.

Elbow position

Scott and Kalaska (1997) found that the preferred directions of some motor cortical cells were altered when the monkey had to reach unnaturally with the elbow raised to shoulder level. In the current theoretical framework, adding elbow position as a free parameter is equivalent to adding one rotation variable φ , for example, the angle between the horizontal plane and the plane determined by the hand, elbow, and shoulder. The same theoretical argument yields the tuning rule:

$$f = f_0 + \mathbf{p} \cdot \mathbf{v} + K \frac{d\varphi}{dt}, \quad (22)$$

where K is a coefficient that may depend on both hand position and elbow position. This formula implies two new effects. The first is that now the preferred direction vector, both its direction and length, may depend on the elbow position φ as well as the hand position (x, y, z) :

$$\mathbf{p} = \mathbf{p}(x, y, z, \varphi), \quad (23)$$

as reported by Scott and Kalaska (1997). The second effect, a new prediction, is that the firing rate may contain a component proportional to the angular speed $d\varphi/dt$ of elbow rotation.

How does this case relate to our earlier results with hand position as the only free parameter? In the preceding sections, reaching was assumed to be “stereotypical” in the sense that the elbow position can be determined completely by the hand position, ignoring forearm rotation. This assumption may not be true if the final posture sometimes depends also on the initial hand position (Soechting et al., 1995). However, when comparing reaching movements starting from the same initial hand position, it is reasonable to assume that for stereotypical reaching, the elbow angle φ can be completely determined by the hand position (x, y, z) , or $\varphi = \varphi(x, y, z)$. Then the time derivative of φ , after expanding by the chain rule, can be absorbed into the term $\mathbf{p} \cdot \mathbf{v}$, yielding the original basic tuning rule in Equation 2. In other words, the assumption of stereotypical movement reduces the total degrees of freedom to 3, eliminating the elbow position as an independent variable. Although the elbow angle can still be used as a free parameter, it is no longer independent of the hand position. Only three parameters are independent in this case, and their exact choice does not affect the general form of the tuning rule (see Appendix A for more discussion on coordinate-system independence).

Summary and discussion of more complex cases

As shown above, the basic tuning theory can naturally account for several important experimental results without making any specific assumptions about the exact variables encoded or details of the encoding. These results are generic properties independent of the exact functional interpretations. This generality makes sense

because during stereotypical movement, redundant variables are inevitably constrained by the geometry and become highly correlated, so that they are likely to show similar tuning properties of the same general type. The theory presented here has formalized this intuition.

The relationship between cosine tuning properties and geometric constraints is also apparent in the studies of muscle activities and actions during reaching and isometric tasks. Basic properties resembling those for motor cortical cells have been reported, including approximately cosine directional tuning curves (but often with a small secondary peak opposite the preferred direction), speed sensitivity, and posture dependence (Flanders and Soechting, 1990; Flanders and Herrmann, 1992; Buneo et al., 1997).

The basic theory needs to be generalized in situations where the hand position is not the only free parameter. For example, force is one variable that is often correlated with the activity of motor cortex; recent examples related to directional tuning include tasks with static load (Kalaska et al., 1989) and varying isometric forces (Georgopoulos et al., 1992; Sergio and Kalaska, 1997).

As another example, preparatory activity in motor cortex before onset of movement can reflect the upcoming reaching direction, as is especially evident during instructed delay (Georgopoulos et al., 1989a), and can change rapidly in tasks requiring mental rotation (Georgopoulos et al., 1989b) or target switching (Pellizzer et al., 1995).

Moreover, when sensory and motor components were decoupled, some neurons even from primary motor cortex were more closely related to the visual movement of a cursor on the computer screen than to the joystick position or hand movements, in both one-dimensional (Alexander and Crutcher, 1990) and two-dimensional tasks (Shen and Alexander, 1997a). By contrast, in virtual reality experiments with visual distortion, motor cortical activity mainly followed the actual limb trajectory rather than the animal's visual perception (Moran et al., 1995).

In addition, some differences exist among the neural activity from different brain areas, although they all show approximate cosine directional tuning (compare Fig. 2). For instance, compared with neurons in the motor cortex in a reaching task, the preferred directions in the cerebellum are more variable in repeated trials (Fortier et al., 1989), neurons in the parietal cortex are less sensitive to static load (Kalaska et al., 1990), and neurons in the premotor cortex are activated earlier, more transiently (Caminiti et al., 1991; Crammond and Kalaska, 1996), and affected more frequently by visual cues (Wise et al., 1992; Shen and Alexander, 1997b). In the motor cortex and elsewhere, there also exist neurons with complex properties that are either not task-related or hard to describe but still could have useful functions in a distributed network (Fetz, 1992; Zipser, 1992; Moody et al., 1998).

In most of these cases, there are additional free variables besides hand position. The linear theory may still yield useful results in these more complex cases after including these additional variables. For example, the planned movement direction is an independent variable, which could be used to describe some preparatory activity before overt hand movement. These new variables should be included when deriving the tuning rule, as demonstrated in the preceding section by adding the elbow position as a free variable in abducted reaching.

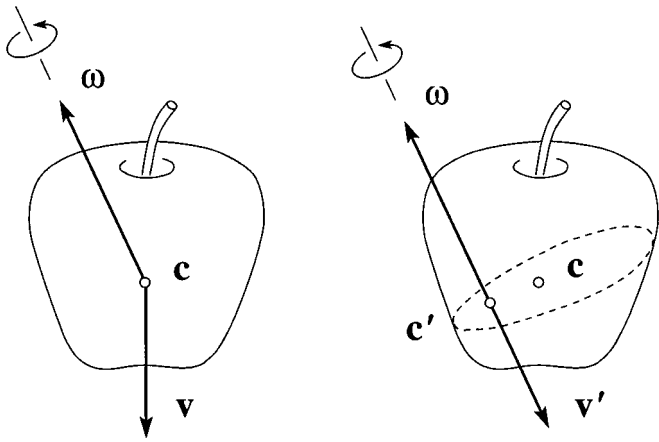


Figure 4. Arbitrary motion of a rigid object can always be decomposed instantaneously into a translation and a rotation, allowing arbitrary parallel shift of the rotation axis. The two examples shown here describe identical physical motion. Parallel shift of rotation axis affects the translation velocity but not the angular velocity ω .

REPRESENTING RIGID OBJECT MOTION

The same geometric argument for arm movement can be applied to moving rigid objects, which have additional rotational degrees of freedom around an axis in space (Fig. 1). In the following, we derive a general tuning rule for rigid motion, discuss its basic properties, and then contrast the results with concrete models of visual receptive fields.

Description of rigid object motion

Arbitrary instantaneous motion of a rigid object can always be described by a rotation plus a translation (Fig. 4), but given the same physical motion, this description is ambiguous up to an arbitrary parallel shift of the rotation axis. For example, translational velocity can always be aligned instantaneously with the angular velocity to obtain a screw motion by passing the rotation axis through the point of zero velocity in a perpendicular plane (Fig. 4).

This ambiguity disappears when the rotation axis is always required to pass through the same reference center in the object, say, the center of mass. We assume that the reference center has been chosen so that a rigid motion can be described uniquely by a translational velocity and an angular velocity. We return to this topic later.

The static position and orientation of a rigid object can be specified by six independent parameters:

$$(x, y, z, \theta_1, \theta_2, \theta_3), \tag{24}$$

where x, y, z describe the position of the reference center of the object with respect to a coordinate system fixed to the world, and $\theta_1, \theta_2, \theta_3$ are three angular variables that represent the object's orientation. The translational velocity of the object is:

$$\mathbf{v} = (\dot{x}, \dot{y}, \dot{z}). \tag{25}$$

The angular velocity $\omega = (\omega_x, \omega_y, \omega_z)^T$ in world coordinates is always linearly related to the time derivatives of the orientation variables $\dot{\theta} = (\dot{\theta}_1, \dot{\theta}_2, \dot{\theta}_3)^T$:

$$\omega = \mathbf{M}\dot{\theta}, \tag{26}$$

where \mathbf{M} is an invertible 3×3 matrix that depends only on the orientation $(\theta_1, \theta_2, \theta_3)$. For example, when Euler angles are used to describe orientation (Fig. 5), we have:

$$(\theta_1, \theta_2, \theta_3) = (\theta, \phi, \psi), \tag{27}$$

and

$$\mathbf{M} = \begin{pmatrix} \cos \phi & 0 & \sin \theta \sin \phi \\ \sin \phi & 0 & -\sin \theta \cos \phi \\ 0 & 1 & \cos \theta \end{pmatrix}, \tag{28}$$

which is invertible as long as $\det \mathbf{M} = \sin \theta \neq 0$ (Goldstein, 1980).

Only the abstract linear relation in Equation 26 is needed in the next section. The actual choice of $(\theta_1, \theta_2, \theta_3)$ is unimportant here. Because the time derivatives of different sets of variables are linearly related by a Jacobian matrix, Equation 26 always holds regardless of the exact choice of the parameterization of orientation (see also Appendix A on independence of the coordinate system).

Tuning rule for rigid motion

Consider neuronal activity associated with motion of a rigid three-dimensional object. Assume that the mean firing rate of a neuron relative to baseline, with a possible time delay, is proportional to the time derivative of a smooth function of the position and orientation of the object in three-dimensional space. In other words:

$$f = f_0 + \frac{d}{dt} \Phi(x, y, z, \theta_1, \theta_2, \theta_3), \tag{29}$$

where f is the firing rate, f_0 is the baseline rate, and Φ is an arbitrary function of object position (x, y, z) and orientation $(\theta_1, \theta_2, \theta_3)$, as described in the preceding section. This equation is analogous to Equation 1.

The exact form of function Φ need not be specified here. It may depend on both the receptive field properties of the cell and the visual appearance of the object and its surroundings. This formulation is quite general. For example, all the visual cues of the object illustrated in Figure 1 are functions of the position and orientation of the object that completely determine how light is reflected from various surfaces, whether diffuse (uniform scattering in all directions) or specular (energy concentrated around the

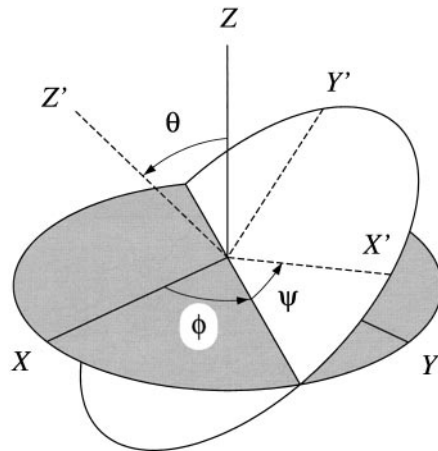


Figure 5. Euler angles (θ, ϕ, ψ) describe an arbitrary orientation of a rigid object with axes (X', Y', Z') with respect to a standard orientation with axes (X, Y, Z) .

mirror reflection direction), giving rise to various visual effects such as shading, shadows, specular reflections, and highlights (Watt and Watt, 1992). Given that all sensory cues are determined completely by the position and orientation of the object, we expect a motion-sensitive neuron to respond to changes of these variables. The simplest way to estimate these changes is to compute the first temporal derivative.

The assumption in Equation 29 allows us to derive a general tuning rule for neurons sensitive to three-dimensional object motion. Given a three-dimensional object moving at instantaneous translational velocity \mathbf{v} and angular velocity $\boldsymbol{\omega}$, the mean firing rate of a generic neuron should depend on these variables in a highly stereotyped way:

$$f = f_0 + \mathbf{p} \cdot \mathbf{v} + \mathbf{q} \cdot \boldsymbol{\omega}, \quad (30)$$

where f_0 is the background firing rate, \mathbf{p} is the *preferred translational direction*, given by:

$$\mathbf{p} = (p_x, p_y, p_z) = \left(\frac{\partial \Phi}{\partial x}, \frac{\partial \Phi}{\partial y}, \frac{\partial \Phi}{\partial z} \right), \quad (31)$$

and vector \mathbf{q} is the *preferred rotation axis*, given by:

$$\mathbf{q} = (q_1, q_2, q_3) = \mathbf{q}^* \mathbf{M}^{-1}, \quad (32)$$

with matrix \mathbf{M} as in Equation 26, and:

$$\mathbf{q}^* = (q_1^*, q_2^*, q_3^*) = \left(\frac{\partial \Phi}{\partial \theta_1}, \frac{\partial \Phi}{\partial \theta_2}, \frac{\partial \Phi}{\partial \theta_3} \right) \quad (33)$$

is an intermediate vector variable, the transformed preferred rotation axis in the orientation angle space. Both the preferred translational direction \mathbf{p} and the preferred rotation axis \mathbf{q} are vectors in the physical space. They may depend on the object and its position and orientation but not on the translational velocity \mathbf{v} and angular velocity $\boldsymbol{\omega}$. The derivation of Equation 30 follows from the chain rule:

$$\begin{aligned} \frac{d\Phi}{dt} &= \frac{\partial \Phi}{\partial x} \dot{x} + \frac{\partial \Phi}{\partial y} \dot{y} + \frac{\partial \Phi}{\partial z} \dot{z} + \frac{\partial \Phi}{\partial \theta_1} \dot{\theta}_1 + \frac{\partial \Phi}{\partial \theta_2} \dot{\theta}_2 + \frac{\partial \Phi}{\partial \theta_3} \dot{\theta}_3 \\ &= \mathbf{p} \cdot \mathbf{v} + \mathbf{q}^* \cdot \dot{\boldsymbol{\theta}} = \mathbf{p} \cdot \mathbf{v} + \mathbf{q} \cdot \boldsymbol{\omega}, \end{aligned} \quad (34)$$

where Equations 25 and 26 and the definitions in Equations 31–33 have been used. The derivation of the tuning rule does not depend on which coordinate system is used (Appendix A).

Before explaining the meaning of the tuning rule in the next section, first consider the baseline firing rate, which is not constrained by the present theory and thus requires separate consideration. The baseline firing rate may itself be modulated by several factors, and the simplest linear model is:

$$f_0 = a_0 + a_1 x + a_2 y + a_3 z + b_1 \theta_1 + b_2 \theta_2 + b_3 \theta_3 + a v + b \omega, \quad (35)$$

where a_i , b_i , a , b are constants, and the position (x, y, z) and the orientation $(\theta_1, \theta_2, \theta_3)$ of the object are included as possibly relevant factors related to the static view, together with the translational speed v and the angular speed ω for object motion, which may also be relevant. This linear equation generalizes Equation 9 for motor neurons. Similarly, Equation 10 can also be generalized by including angular position and speed. This assumes that the baseline firing rate in general may contain information about both the static configuration of an object and its instantaneous motion.

Cosine tuning and multiplicative speed modulation

The basic tuning rule in Equation 30 can be rewritten in its equivalent form:

$$f = f_0 + p v \cos \alpha + q \omega \cos \beta, \quad (36)$$

where $v = |\mathbf{v}|$ is the speed of translation, $\omega = |\boldsymbol{\omega}|$ is the angular speed of rotation, $p = |\mathbf{p}|$ is the length of the preferred direction vector, $q = |\mathbf{q}|$ is the length of the preferred rotation vector, α is the angle between vectors \mathbf{p} and \mathbf{v} , and β is the angle between vectors \mathbf{q} and $\boldsymbol{\omega}$.

In other words, given the particular view of a particular object, the response above baseline should be the sum of two components, one translational and one rotational. The translational component is proportional to the cosine of the angle between a fixed preferred translational direction and the actual translational direction. In addition, it is also modulated linearly by the speed of translation, which does not alter the shape of the tuning curve. Similarly, the rotational component is proportional to the cosine of the angle between a fixed preferred rotation axis and the actual rotation axis. In addition, the rotational component is also modulated linearly by the angular speed of rotation.

Distribution of preferred direction and preferred axis

Thus far, the view of the given object is assumed to be fixed. That is, the cosine tunings for both translation and rotation are defined with respect to a particular view of the object. When the view of the object changes, the preferred translational direction \mathbf{p} and preferred rotation axis \mathbf{q} of a motion-sensitive neuron may also change.

The theory constrains this change because the preferred translational direction \mathbf{p} and the transformed preferred rotation axis \mathbf{q}^* are derived as gradient fields in Equations 31 and 33. Here the intermediate vector \mathbf{q}^* is related to the preferred rotation axis \mathbf{q} in physical space by:

$$\mathbf{q}^* = \mathbf{q} \mathbf{M}, \quad (37)$$

according to Equation 32. In three-dimensional space, where curl is defined, the gradient field implies that any three variables taken from the six variables $(x, y, z, \theta_1, \theta_2, \theta_3)$ must be curl-free. For example, when the position (x, y, z) of the object is fixed, the distribution of the transformed preferred rotation axis in the orientation space $(\theta_1, \theta_2, \theta_3)$ must be curl-free:

$$\text{curl } \mathbf{q}^* = \left(\frac{\partial q_3^*}{\partial \theta_2} - \frac{\partial q_2^*}{\partial \theta_3}, \frac{\partial q_1^*}{\partial \theta_3} - \frac{\partial q_3^*}{\partial \theta_1}, \frac{\partial q_2^*}{\partial \theta_1} - \frac{\partial q_1^*}{\partial \theta_2} \right) = \mathbf{0}. \quad (38)$$

Any hypothetical neurons with non-zero curl can be ruled out by this condition (see below). For a gradient field, the zero curl is simply attributable to the equality of mixed second partial derivatives of the potential function, which holds also in higher dimensions. The equivalent path integral formulation is valid also in all dimensions:

$$\oint (\mathbf{p} \cdot d\mathbf{l} + \mathbf{q}^* \cdot d\boldsymbol{\theta}) = 0, \quad (39)$$

along any closed curve in the six-dimensional space, where $d\mathbf{l} = (dx, dy, dz)$, and $d\boldsymbol{\theta} = (d\theta_1, d\theta_2, d\theta_3)$. Another equivalent formu-

lation is that the potential function Φ can be constructed by the path integral:

$$\Phi(\xi) = \Phi(\xi_0) + \int_{\xi_0}^{\xi} (\mathbf{p} \cdot d\mathbf{l} + \mathbf{q}^* \cdot d\boldsymbol{\theta}), \quad (40)$$

which depends only on the end points, not on the exact path. Here $\xi = (x, y, z, \theta_1, \theta_2, \theta_3)$ is an arbitrary point in the parameter space, and ξ_0 is the value at a given initial point. Therefore, in the gradient theory, how the preferred translational direction and the preferred rotation axis of a neuron change with the view of a given object cannot be arbitrary but is highly constrained. This can provide testable predictions (see below).

Linear nongradient theory

A more general theory can be obtained by directly assuming a linear relationship between the firing rate and the components of the translational velocity $\mathbf{v} = (\dot{x}, \dot{y}, \dot{z})$ and the time derivatives of the angular variables $\boldsymbol{\theta} = (\dot{\theta}_1, \dot{\theta}_2, \dot{\theta}_3)$. This yields the same tuning rule:

$$f = f_0 + p_x \dot{x} + p_y \dot{y} + p_z \dot{z} + q_1^* \dot{\theta}_1 + q_2^* \dot{\theta}_2 + q_3^* \dot{\theta}_3 \\ = \mathbf{p} \cdot \mathbf{v} + \mathbf{q} \cdot \boldsymbol{\omega}, \quad (41)$$

where $\mathbf{p} = (p_x, p_y, p_z)$ and $\mathbf{q}^* = (q_1^*, q_2^*, q_3^*)$ are arbitrary vector fields, not necessarily gradient fields, and Equations 26 and 32 are used in the last step. This tuning rule gives the same response properties predicted by the gradient theory for a single view of the object. The difference shows up when the view changes. The nongradient theory imposes no constraint on how preferred translational direction and preferred rotation axis should vary with the view of the object. The gradient theory is more restrictive, and therefore makes stronger predictions.

Change of reference center

Because the description of the same physical motion of a rigid object is ambiguous up to a parallel shift of the rotation axis (Fig. 4), we have assumed in the above that the rotation axis always passes through the same reference point $\mathbf{c} = (x, y, z)$ in the object to ensure uniqueness of description. When a different reference center \mathbf{c}' is chosen, the form of the basic tuning rule in Equation 30 remains valid, but the preferred rotation axis is affected in a predictable way:

$$f = f_0 + \mathbf{p} \cdot \mathbf{v} + \mathbf{q} \cdot \boldsymbol{\omega} = f_0 + \mathbf{p}' \cdot \mathbf{v}' + \mathbf{q}' \cdot \boldsymbol{\omega}', \quad (42)$$

where \mathbf{v}' and $\boldsymbol{\omega}'$ are the translational velocity and angular velocity for the new reference center \mathbf{c}' , and:

$$\mathbf{p}' = \mathbf{p}, \quad (43)$$

$$\mathbf{q}' = \mathbf{q} + \mathbf{p} \times (\mathbf{c}' - \mathbf{c}) \quad (44)$$

are the new preferred translational direction and rotation axis. One can readily verify that Equation 42 is valid under Equations 43 and 44, using the relations:

$$\mathbf{v}' = \mathbf{v} + \boldsymbol{\omega} \times (\mathbf{c}' - \mathbf{c}), \quad (45)$$

$$\boldsymbol{\omega}' = \boldsymbol{\omega}. \quad (46)$$

Therefore, changing the reference center of an object has no effect on the preferred translational direction of a neuron (Eq. 43), whereas the preferred rotation axis is altered systematically

in a completely predictable manner (Eq. 44). These relations arise purely from the ambiguity of the description of rigid motion, and thus apply to both the gradient and the nongradient theories.

Summary

Simple assumptions have led to a general tuning rule for how the mean firing rate of a neuron should depend on the instantaneous motion of an arbitrary rigid object. For each given view of the object, the firing rate is predicted to be the sum of two terms, one for the translational motion component and one for the rotational motion component, both with cosine directional tuning and linear speed or angular speed modulation. In general, the preferred translational direction and the preferred rotation axis may depend on the identity of the object as well as its view. This tuning rule is a linear approximation to the geometry of rigid motion and therefore should obtain regardless of the exact computational mechanisms involved. In other words, this rule is expected to be a robust property for motion-sensitive neurons responding to realistic moving objects. When the view of the object changes, both the preferred translational direction and the preferred rotation axis of a neuron may change as well. The gradient theory provides additional constraints on such changes, whereas the nongradient theory imposes no further constraints. As a consequence, these two theories can be distinguished by further experiments. Finally, although the description of rigid motion is ambiguous up to a parallel shift of the rotation axis, the effects on the tuning rule are completely predictable and therefore convey no additional information about the response properties of a neuron.

Examples of motion-sensitive receptive field models

Many neurons in visual cortex, particularly in the dorsal stream leading to parietal cortex, respond selectively to visual motion. Here we consider three-dimensional rigid motion and examine several simple computational mechanisms that yield explicit analytical formulas for the preferred translational direction and the preferred rotation axis. For each fixed view of the object, the results are consistent with the basic tuning rule in Equation 30. For different views, however, the global gradient-field condition for the preferred axes can be violated by the idealized velocity component detectors. This shows that the neuronal behavior predicted by the gradient theory is not always identical to that of an optic-flow detector.

Velocity component detectors

As illustrated in Figure 6A, suppose the firing rate of an idealized neuron detects local motion on the image plane according to:

$$f = f_0 + \mathbf{p} \cdot \mathbf{u}, \quad (47)$$

where f_0 is the baseline firing rate, \mathbf{p} is the preferred direction of visual motion, and \mathbf{u} is the local velocity on the image plane inside a small receptive field of the detector. This local velocity detector resembles some neurons in the middle temporal area (MT) of monkey, as discussed in the section after Equation 72.

This idealized neuron obeys the basic tuning rule in Equation 30, namely:

$$f = f_0 + \mathbf{p} \cdot \mathbf{v} + \mathbf{q} \cdot \boldsymbol{\omega}, \quad (48)$$

in response to a textured rigid object moving at translational velocity \mathbf{v} and angular velocity $\boldsymbol{\omega}$. Here the preferred direction \mathbf{p}

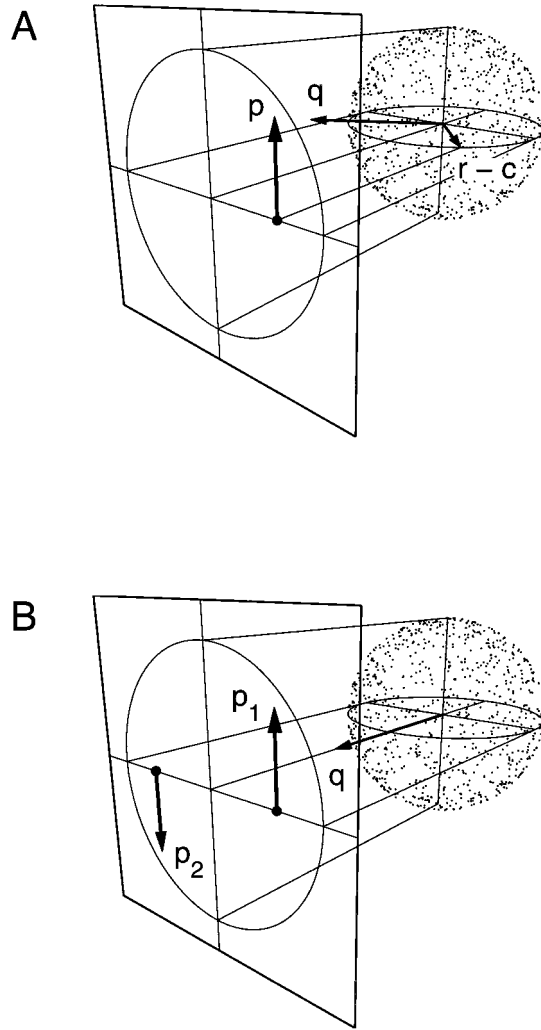


Figure 6. Examples of preferred rotation axis for velocity component detectors. Visual motion of a finely textured rotating object, here a sphere, is projected orthogonally onto the image plane. *A*, For a single local velocity detector with preferred direction \mathbf{p} , the largest response is elicited by rotation around the preferred rotation axis \mathbf{q} , which is on the horizontal plane and perpendicular to the vector $\mathbf{r} - \mathbf{c}$, with \mathbf{c} representing the center of the sphere. *B*, When the response is the sum of two local velocity detectors with preferred directions \mathbf{p}_1 and \mathbf{p}_2 , the preferred rotation axis \mathbf{q} is perpendicular to the image plane.

is the same constant vector as in Equation 47, and the preferred rotation axis is given explicitly by:

$$\mathbf{q} = (\mathbf{r} - \mathbf{c}) \times \mathbf{p}, \quad (49)$$

where \mathbf{c} is a fixed reference center in the object (center of the sphere in Fig. 6*B*), and \mathbf{r} is the coordinate of the point in the object that happens to fall into the vanishingly small receptive field of the detector. As shown in Figure 6*A*, the geometry of the situation is quite simple, with the two orthogonal vectors \mathbf{q} and $\mathbf{r} - \mathbf{c}$ both lying on the horizontal plane. For fixed angular speed, using \mathbf{q} as the rotation axis maximizes the response of the velocity detector.

To derive these formulas, note that a point in the object with coordinate \mathbf{r} has the velocity:

$$\mathbf{u} = \mathbf{v} + \boldsymbol{\omega} \times (\mathbf{r} - \mathbf{c}) \quad (50)$$

on the image plane (Fig. 6*A*). The desired equations can be obtained by inserting this into Equation 47 and using the vector identity:

$$\mathbf{p} \cdot [\boldsymbol{\omega} \times (\mathbf{r} - \mathbf{c})] = [(\mathbf{r} - \mathbf{c}) \times \mathbf{p}] \cdot \boldsymbol{\omega}. \quad (51)$$

Next, consider a higher-order neuron whose response is the sum of the outputs of several local velocity component detectors:

$$f = f_0 + \sum_{i=1}^N \mathbf{p}_i \cdot \mathbf{u}_i, \quad (52)$$

where \mathbf{p}_i is the preferred direction of detector i , and \mathbf{u}_i is the image velocity in its receptive field. This neuron also obeys the same basic tuning rule in Equation 48, with the preferred translational direction and preferred rotation axis given by:

$$\mathbf{p} = \sum_{i=1}^N \mathbf{p}_i, \quad (53)$$

$$\mathbf{q} = \sum_{i=1}^N (\mathbf{r}_i - \mathbf{c}) \times \mathbf{p}_i. \quad (54)$$

For example, when two detectors are arranged as shown in Figure 6*B*, the preferred rotation axis \mathbf{q} is perpendicular to the image plane, whereas the preferred translational axis vanishes ($\mathbf{p} = \mathbf{0}$) because the image size is constant under orthographic projection.

Finally, the basic tuning rule in Equation 48 still holds for image motion of a rigid object under a perspective projection, which projects each point (x, y, z) in the real world toward the observer at the origin $(0, 0, 0)$, leaving an image at (X, Y) in the image plane at $z = \eta$:

$$X = \eta x/z, \quad Y = \eta y/z \quad (55)$$

(Longuet-Higgins and Prazdny, 1980; Heeger and Jepson, 1990). On the image plane, suppose the velocity component detector i is located at (X_i, Y_i) with preferred direction $\mathbf{p}_i = (p_{i1}, p_{i2})$, then the preferred direction and preferred rotation axis become:

$$\mathbf{p} = \sum_{i=1}^N \frac{1}{z_i} \begin{pmatrix} p_{i1}\eta \\ p_{i2}\eta \\ -p_{i1}X_i - p_{i2}Y_i \end{pmatrix}, \quad (56)$$

$$\mathbf{q} = \sum_{i=1}^N \begin{pmatrix} -p_{i1}X_i\hat{Y}_i - p_{i2}(Y_i\hat{Y}_i + \eta\hat{Z}_i) \\ p_{i1}(X_i\hat{X}_i + \eta\hat{Z}_i) + p_{i2}Y_i\hat{X}_i \\ (-p_{i1}\hat{Y}_i + p_{i2}\hat{X}_i)\eta \end{pmatrix}, \quad (57)$$

where $(\hat{X}_i, \hat{Y}_i, \hat{Z}_i) = (x_i - x, y_i - y, z_i - z)/z_i$ with (x, y, z) the reference center of the object, and z_i is the z -coordinate in real world for the physical point that happens to activate detector i . Unlike the orthographic projection in Figure 6, this mechanism allows a neuron to respond to looming or shrinking images, because it does not confine the preferred translational direction \mathbf{p} to the image plane.

Spatiotemporal receptive field

Now consider motion-sensitive linear spatiotemporal receptive fields that obey the basic tuning rule in Equation 30 with a known potential function. Let $I(X, Y, t)$ describe the intensity of an image at location (X, Y) on the image plane at time t , ignoring color and stereo. Suppose the firing rate of a neuron with linear receptive

field $F(X, Y)$ is linearly related to how fast the overlap between the image and receptive field is changing:

$$f = f_0 + \frac{d}{dt} \langle F, I \rangle, \quad (58)$$

where the inner product is defined by:

$$\langle F, I \rangle = \iint F(X, Y) I(X, Y, t) dX dY. \quad (59)$$

This inner product can serve as the potential function postulated in Equation 29:

$$\Phi = \langle F, I \rangle. \quad (60)$$

Therefore, the basic tuning rule in Equation 30 must hold true, and the preferred translational direction and rotation axis are:

$$\mathbf{p} = (\langle F, \partial_x I \rangle, \langle F, \partial_y I \rangle, \langle F, \partial_z I \rangle), \quad (61)$$

$$\mathbf{q} = (\langle F, \partial_1 I \rangle, \langle F, \partial_2 I \rangle, \langle F, \partial_3 I \rangle) \mathbf{M}, \quad (62)$$

following Equations 31–33. Here the partial derivatives are with respect to the implicit variables $(x, y, z, \theta_1, \theta_2, \theta_3)$ for the position and orientation of a moving object that generates the image.

More generally, consider a neuron with an arbitrary linear spatiotemporal receptive field G so that its firing rate is:

$$f = f_0 + \iiint G(X, Y, \tau) I(X, Y, t - \tau) dX dY d\tau. \quad (63)$$

This becomes identical to Equation 58 for the kernel function:

$$G(X, Y, \tau) = F(X, Y) \frac{d\delta(\tau)}{d\tau}, \quad (64)$$

where δ is Dirac delta function, which can be approximated by any narrow and normalized smooth function peaked at the origin. If Equation 64 is a reasonable approximation to the spatiotemporal receptive field, then the basic tuning rule in Equation 30 as well as \mathbf{p} and \mathbf{q} given by Equations 61 and 62 becomes valid. For nonlinear mechanisms, such as squaring (Adelson and Bergen, 1985) and normalization (Heeger, 1993), the above consideration may apply only after local linearization.

Existence of a global potential function

In all the concrete examples considered above, the basic tuning rule in Equation 30 holds true for each given view of an object. However, for a single view, the gradient and nongradient theories are indistinguishable. By assumption, the nongradient theory allows arbitrary preferred translational direction \mathbf{p} and preferred rotation axis \mathbf{q} . For a given view, the gradient theory can also generate any desired constant vectors \mathbf{p} and \mathbf{q} from the gradients of the following potential function:

$$\Phi(x, y, z, \theta_1, \theta_2, \theta_3) = \mathbf{p} \cdot \mathbf{c} + \mathbf{q}^* \cdot \boldsymbol{\theta}, \quad (65)$$

where $\mathbf{q}^* = \mathbf{qM}$ is taken as a constant vector, $\mathbf{c} = (x, y, z)$ is the reference center of the object, and $\boldsymbol{\theta} = (\theta_1, \theta_2, \theta_3)$ describes the orientation of the object.

The gradient theory is globally correct only when a potential function exists for all views of the object. This is the case for the linear spatiotemporal model, where the potential function can be

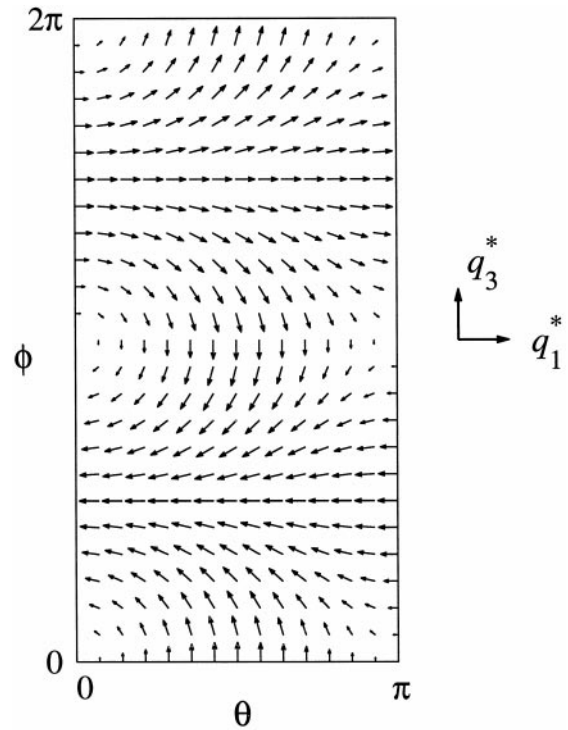


Figure 7. The gradient theory prohibits a neuron from having a constant preferred rotation axis in the physical space regardless of the view of the object, because this would allow the transformed preferred rotation axis (q_1^*, q_2^*, q_3^*) to have non-zero curl in the Euler angle space (θ, ϕ, ψ) . Here the constant preferred rotation axis $\mathbf{q} = (0, -1, 0)$ is taken as the negative Y -axis. Only two-dimensional visualization is needed here because $q_2^* = 0$, and the vector field is independent of ψ .

given explicitly (Equation 60). By contrast, for the idealized velocity component detector, a global potential function in general does not exist, as shown in Example 1 below.

Because the existence of a potential function does not depend on the choice of the coordinate system (see Appendix A), we only need to show that a potential function does not exist in the Euler angle space: $(\theta_1, \theta_2, \theta_3) = (\theta, \phi, \psi)$, assuming that the center of the object is fixed. In this three-dimensional space, a potential function exists if and only if the distribution of the transformed preferred rotation axis \mathbf{q}^* is curl free. Now consider two special examples that do not admit a global potential function:

Example 1: Constant preferred rotation axis fixed to the world. An explicit example is the model in Figure 6B, where the preferred rotation axis \mathbf{q} is the same regardless of the orientation of the spherical object. Here it is assumed that the velocity component detector has two vanishingly small receptive fields that can nevertheless detect the true local velocity components regardless of the orientation of the object. Without loss of generality, take the preferred rotation axis as a unit vector in the negative Y -axis:

$$\mathbf{q} = (0, -1, 0), \quad (66)$$

and then compute its counterpart vector in the Euler angle space:

$$\mathbf{q}^* = \mathbf{qM} = (-\sin \phi, 0, \sin \theta \cos \phi). \quad (67)$$

It is verified that $\text{curl } \mathbf{q}^*$ as defined in Equation 38 does not vanish. This proves that the desired potential function cannot exist for this hypothetical neuron (Fig. 7).

Example 2: Constant preferred rotation axis fixed to the object. A possible example is a vestibular neuron receiving input from only a single semicircular canal without other influences such as that from the otolith. Then the firing rate has a cosine tuning with respect to a preferred rotation axis fixed on the head of the animal, regardless of the orientation of the head in the world (Baker et al., 1984; Graf et al., 1993). Without loss of generality, let the preferred rotation axis be a unit vector in the positive Z' axis of the object (head), then in world coordinates this axis is:

$$\mathbf{q} = (\sin \theta \sin \phi, -\sin \theta \cos \phi, \cos \theta), \quad (68)$$

so that its counterpart vector in the Euler angle space is:

$$\mathbf{q}^* = \mathbf{qM} = (0, \cos \theta, 1). \quad (69)$$

Here curl \mathbf{q}^* is not zero, proving the nonexistence of the desired potential function.

Therefore, there are simple computational mechanisms that can violate the gradient theory. As shown above, the gradient theory prohibits a neuron from having a truly invariant preferred rotation axis fixed either to the world or to the object. These neurons, however, are allowed by the nongradient theory. In particular, the prediction of the gradient theory can differ from that of an optic-flow sensor. As another example, the preferred translational direction field for a small moving dot may also have non-zero curl when measured at different regions inside a large MST-like receptive field that has circular arrangement of local preferred directions (Saito et al., 1986). These idealized examples demonstrate that the global property of the gradient theory is quite restrictive, which, however, makes its prediction strong and refutable. Experiments could be performed to test whether the gradient theory accounts for the neuronal responses to three-dimensional object motion.

COMPARISON WITH EXPERIMENTAL RESULTS OF SINGLE NEURONS

The tuning rule for reaching arm movement in Equation 2 is a special case of the general tuning rule in Equation 30 without the rotational terms. Biological evidence from the motor system in support of the tuning rule has already been considered in the preceding sections. In this section we examine several additional biological examples that are consistent with some special cases of the general tuning rule and then discuss more comprehensive tests for moving rigid objects.

One-dimensional example: hippocampal place fields on a linear track

For one-dimensional movement, the linear tuning theory predicts only linear speed modulation, without further constraint on the tuning function. The firing rate is given by:

$$f = f_0 + \frac{d\Phi(x)}{dx} \dot{x} = f_0 + \phi(x)v, \quad (70)$$

where x is the variable of interest, $v = dx/dt$ is the speed, and $\phi(x)$ is the gradient of function Φ . This is a special case of the general tuning rule in Equation 36, keeping only the translational term, with $\cos \alpha = 1$ and $p = \phi(x)$. Function $\phi(x)$ is allowed to be completely arbitrary because a potential function:

$$\Phi(x) = \Phi(0) + \int_0^x \phi(\zeta) d\zeta \quad (71)$$

can always be constructed. For example, the firing rates of hippocampal place cells are modulated by running speed when a rat moves on a narrow track (McNaughton et al., 1983). In this one-dimensional problem, Equation 70 does not constrain the tuning function $\phi(x)$, here interpreted as a place field, describing the mean firing rate at spatial position x . When averaged over a population of simultaneously recorded place cells to get enough spikes, the average firing rate was indeed remarkably linearly related to the running speed, with a correlation coefficient >0.95 over the full range of speeds (Zhang et al., 1998), in agreement with Equation 70. A similar linear relationship was also reported recently for rats on a running wheel (Hirase et al., 1998). The theory gives a correct tuning rule without reference to the underlying biological mechanisms.

Two-dimensional example: local translational visual motion

Neurons in middle temporal area (MT or V5) of monkeys respond selectively to the direction of local visual motion (Zeki, 1974; Maunsell and Van Essen, 1983; Albright, 1984), although they are also affected by other factors, such as surround motion (Allman et al., 1985; Tanaka et al., 1986; Raiguel et al., 1995), pattern motion (Movshon et al., 1985), transparency (Stoner and Albright, 1992; Qian and Andersen, 1994), and form cues (Albright, 1992). Consider the following formula obtained by keeping only the translational term in Equation 30:

$$f = f_0 + \mathbf{p} \cdot \mathbf{v}, \quad (72)$$

which has been used for reaching arm movement (Eq. 2). This tuning rule also resembles the directional sensitivity of MT neurons (Zhang et al., 1993; Buračas and Albright, 1996), where \mathbf{p} is the preferred direction of the neuron, and \mathbf{v} is the velocity of the stimulus inside the receptive field.

This simple formula can capture two primary features of many MT neurons: a broad directional tuning curve, and speed modulation without changing the shape of the tuning curves (Rodman and Albright, 1987), while setting aside various other properties accounted for by more detailed models (Serenio, 1993; Nowlan and Sejnowski, 1995; Buračas and Albright, 1996; Simoncelli and Heeger, 1998). For many MT neurons, the tuning curves are often sharper than cosine, in which case a circular normal curve in Equation A17 might provide better fit because of its closeness to a Gaussian (Albright, 1984). Linear speed modulation is probably a reasonable approximation for some neurons when the velocity is slow, but typically firing rates often decrease after reaching a peak at an optimal speed (Maunsell and Van Essen, 1983). It would be interesting to test whether speed modulation is linear when averaged over raw firing rates for a large population of neurons, especially under ecologically plausible stimulus conditions. The above consideration may also apply to many V4 neurons, which responded to visual motion response in an MT-like manner (Cheng et al., 1994). Cosine tuning curves for translational motion have also been described in the cerebellum (Krauzlis and Lisberger, 1996) and the parietal area 7a (Siegel and Read, 1997).

Three-dimensional object motion

Spiral motion

No direct experimental data are available on how a neuron responds systematically to a realistic moving three-dimensional object with arbitrary translation and rotation. One closely related example is the broad tuning of some neurons to spiral visual

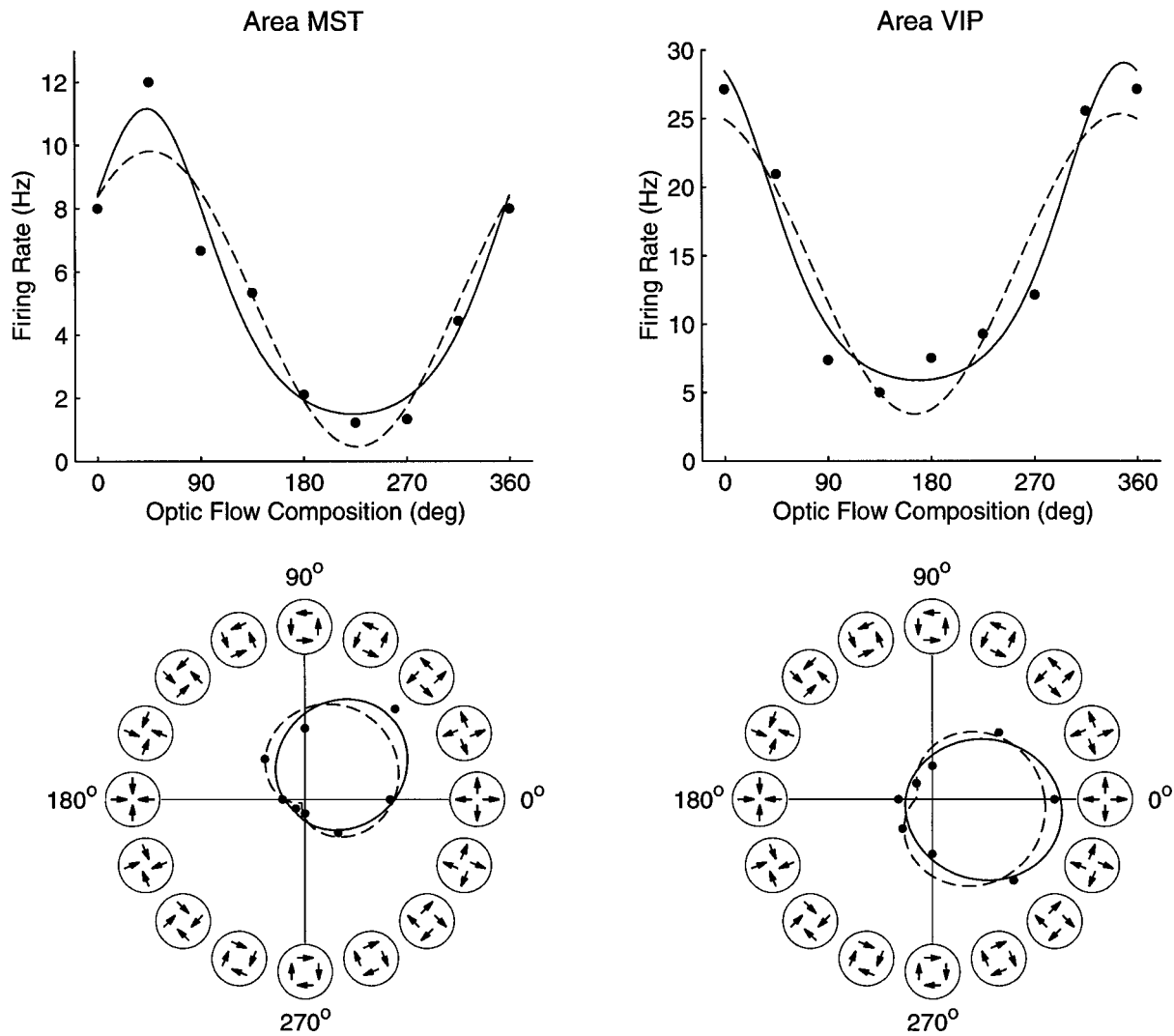


Figure 8. Visual cortical neurons sensitive to spiral optic flow may provide evidence for the existence of preferred translation direction and preferred rotation axis for three-dimensional object motion. Here the average response of a neuron depended on the combination of radial and circular motion components, which varied systematically by changing local velocity directions in the moving random dot stimulus. The solid curves are circular normal fit, and the dashed curves are cosine fit. Data of the MST neuron are from Figures 8C in Graziano et al. (1994), and the VIP neuron data are from Figure 10 in Schaafsma and Duysens (1996), with permission.

motion, which may be generated plausibly by a large moving planar object facing the observer.

As shown in Figure 8, neurons in monkey visual medial superior temporal area (MST), which receive a major input from area MT, typically respond well to wide-field random-dot spiral motion patterns (Graziano et al., 1994). Most neurons in the ventral intraparietal area (VIP) are also sensitive to visual motion (Colby et al., 1993), and some have tuning properties to spiral motion similar to those in area MST (Schaafsma and Duysens, 1996), probably due to input directly from MST and/or integration of inputs from area MT. Area 7a is at a higher level than MST and might have more complex response properties for optic flows (Siegel and Read, 1997). In theory, it is possible to build an MST-like neuron from MT-like local motion inputs, even with position-invariance properties (Saito et al., 1986; Poggio et al., 1991; Sereno and Sereno, 1991; Zhang et al., 1993). Tuning to spiral motion was predicted based on Hebbian learning of optic flow patterns (Zhang et al., 1993) and by other unsupervised learning algorithms (Wang, 1995; Zemel and Sejnowski, 1998).

To explain spiral tuning in terms of rigid motion, regard the environment itself as a large rigid object, moving relative to the observer. For the experiments mentioned above, the environment may be considered as a finely textured screen, oriented vertically, facing the observer. Translating this screen toward or away from the observer induces expansion or contraction, whereas rotating the screen around a perpendicular axis induces circular motion. According to the basic tuning rule in Equation 36, a neuron should respond to arbitrary motion of this screen with the firing rate:

$$f = f_0 + P v + Q \omega, \tag{73}$$

where v is the translational speed, ω is the angular speed, and:

$$P = p \cos \alpha, \quad Q = q \cos \beta \tag{74}$$

are constants, where α is the angle between the preferred translational direction and the actual translational direction, and β is

the angle between the preferred rotation axis and the actual rotation axis.

To see why this accounts for spiral tuning, write Equation 73 in the equivalent form:

$$f = f_0 + \mathbf{w} \cdot \mathbf{s} = f_0 + |\mathbf{w}||\mathbf{s}|\cos \gamma, \quad (75)$$

where $\mathbf{s} = (v, r\omega)$ is the *spiral composition vector* describing the stimulus, $\mathbf{w} = (P, Q/r)$ is the *preferred spiral composition vector* describing response properties of the neuron, r is a constant length, introduced to make v and $r\omega$ of the same units, and γ is the angle between \mathbf{s} and \mathbf{w} . In the polar diagrams in Figure 8, the horizontal and vertical axes correspond to the two components of the stimulus spiral composition \mathbf{s} , whose length:

$$|\mathbf{s}| = \sqrt{v^2 + (r\omega)^2} \quad (76)$$

was fixed during the experiments. It follows from Equation 75 that the response should fall off smoothly when the stimulus spiral composition becomes different from the preferred composition, in proportion to the cosine of the angle γ between them. This broad tuning to spiral motion is generally consistent with the data (Graziano et al., 1994; Schaafsma and Duysens, 1996), although a circular normal function in Equation A17, with one more free parameter, may provide better fits than a cosine function (Fig. 8).

The above interpretation implies that firing rate should scale linearly with translational speed and angular speed independently of the spiral tuning curve. The responses of most MST neurons do indeed depend on speed (Tanaka and Saito, 1989; Orban et al., 1995), and many are monotonically increasing (Duffy and Wurtz, 1997a). It would be interesting to test quantitatively how well the linearity holds when averaged over a population of cells, especially for an ecologically relevant range of motion.

The tuning rule in Equation 73 also implies that the response should depend on the focus of expansion or the translational component in the optic flow, which also occurs for many MST cells (Duffy and Wurtz, 1995, 1997b). Adding a translational velocity vector to the stimulus corresponds to translating the stimulus screen sideways, which affects the angle α in Equation 74 and thus the response in Equation 73. Changing the translational direction and the rotation axis of the stimulus screen can alter both angles α and β in Equation 74 and thus the predicted response in Equation 73.

Motion-sensitive neurons in area MST may be used for purposes such as estimating heading or self-motion (Perrone and Stone, 1994; Lappe et al., 1996) or segmenting multiple moving objects (Zemel and Sejnowski, 1998). MST responses can be affected by various factors, including, for example, surround motion (Tanaka et al., 1986; Eifuku and Wurtz, 1998), disparity (Roy et al., 1992), eye position and movement (Newsome et al., 1988; Bradley et al., 1996; Squatrito and Maioli, 1997), vestibular input (Thier and Erickson, 1992), form cues (Geesaman and Andersen, 1996), the presence of multiple objects (Recanzone et al., 1997), and attention (Treue and Maunsell, 1996). Most experiments used simplified stimuli, although more realistic stimuli were tested recently (Sakata et al., 1994; Pikel et al., 1996). Because most of these examples contain parameters other than the object's position and orientation, additional variables are needed to account for all of these effects in a model.

Further experimental test

Given all the contributing factors mentioned above, it is natural to ask how a neuron would respond to a more natural-moving

three-dimensional object. A simple geometric stimulus is easier to specify and present but may lack important sensory cues needed to predict the response of a neuron to a natural stimulus. Our analysis relies on varying the translational direction and rotation axis and might provide a convenient basic description for response properties in terms of a preferred translational direction and a preferred rotation axis.

To test directly the basic tuning rule in Equation 30 or 36, one should present realistic images of a moving three-dimensional object to motion-sensitive neurons. The simplest way to test the theory is to oscillate slightly an object around a fixed axis. The oscillation should be sufficiently small so that salient visual cues are not occluded. For sinusoidal oscillations with frequency Ω and amplitude ρ :

$$\varphi(t) = \rho \sin(\Omega t), \quad (77)$$

and the angular speed is $\omega = d\varphi/dt$ in Equation 36. Thus the theory predicts that the firing rate as a function of time should be:

$$f(t + \tau) = f_0 + q\rho\Omega \cos(\Omega t) \cos \beta, \quad (78)$$

where β is the angle between the actual rotation axis and the preferred rotation axis, q is a constant coefficient, and τ is the latency for visual response. The translational term in Equation 36 vanishes because there is no translational motion ($v = 0$). Systematically changing the orientation of the rotation axis (angle β) while keeping the view fixed allows the tuning function and the preferred rotation axis to be measured for the neuron.

Similarly, the response to translation in three-dimensional space could be tested by oscillating the whole object along a straight line:

$$x(t) = \rho \sin(\Omega t). \quad (79)$$

The translational speed is $v = dx/dt$ in Equation 36, and the firing rate is:

$$f(t + \tau) = f_0 + p\rho\Omega \cos(\Omega t) \cos \alpha, \quad (80)$$

where α is the angle between the actual translational direction and the preferred translational direction, and p is a constant coefficient. The preferred translational direction can be measured by systematically changing the translational direction (angle α) while keeping the view fixed.

For more efficient tests, the object could be rotated continuously with varying angular speed, covering all relevant views, first with respect to a fixed axis, and then systematically changing the axis. If the basic tuning rule is correct and the system is essentially linear, the tuning function and the preferred rotation axis could be computed for each view of the object. An even more efficient test is possible with a continuously time-varying rotation axis that generates tumbling movements of the object (Stone, 1998).

Eye position is one implicit factor that may affect the preferred translational direction and preferred rotation axis. The present theory allows an eye position effect but provides no additional constraints.

The linear response properties of a neuron for a given object are specified completely by its preferred translational direction, preferred rotation axis, and baseline firing rate for each given view of the object, as well as how these parameters depend on the view. All of these properties are experimentally testable and can be compared with the theoretical predictions in the preceding

sections. For example, with the center of the object fixed, the curl-free condition for a given neuron should be tested by measuring its preferred rotation axis for four or more different orientations of the object. For a full test in six-dimensional space, both the preferred translational direction and the preferred rotation axis of the neuron should be measured for seven or more different positions and orientations of the object. See Appendix B for further discussion.

If motion-sensitive neurons with similar tuning properties are clustered in the brain, then it might be possible to use functional magnetic resonance imaging techniques to test the predicted properties of the tuning rule in animal and human subjects using realistic images of moving 3-D objects as visual stimuli.

DISCUSSION

An explanation for cosine tuning

The remarkable ubiquity of approximately cosine tuning curves for a wide range of neural responses in the visual and motor systems suggests that there may be a common explanation that transcends the specific mechanisms that generate these response properties. We have shown that the low dimensionality of the geometric variables that underlie object motion and body movements could account for these observations. The gradient formulation of this general principle provides a rigorous framework for unifying the dependence of tuning curves on the axes and speeds of rotation and translation.

This theoretical framework makes a number of specific predictions. The primary prediction is the existence of preferred axes of rotation and translation for moving objects, which can be determined by systematically rotating and translating objects in the receptive field of cortical neurons. The firing rate of a neuron should fall off in proportion to the cosine of the angle between the preferred rotation axis and translational direction and the true rotation axis and translational direction. In addition, the speed and angular speed should modulate the firing rate multiplicatively without changing the shape of the directional tuning functions, somewhat related to the multiplicative gain fields in the parietal cortex for eye position (Andersen et al., 1997; Salinas and Abbott, 1997) and recent evidence for distance modulation of responses in visual cortex (Trotter et al., 1992; Dobbins et al., 1998). A secondary prediction is that the fields of preferred directions of rotation and translation for each individual neuron are curl free; these are global conditions on the overall pattern of vectors. The curl-free condition is relaxed in nongradient theory, which provides a theoretical alternative that can be experimentally tested.

Cosine tuning for the direction of arm movement characterizes many neurons in the motor cortex as well as in other parts of the motor system. If this cosine tuning with direction mainly reflects the geometrical constraint of moving in a three-dimensional space, as we propose, then the specific functions of these neurons in guiding and planning limb movements must be sought in other properties. One way to obtain this information is to measure how the preferred direction varies in space for different hand positions, because this vector field completely specifies the properties of a neuron in a linear theory. When the preferred direction field is curl free, the underlying potential function that generates the gradient field can be constructed empirically.

Cosine tuning function is an approximation to biological data. For example, the averaged directional tuning curves in Figure 2 are all slightly sharper than a cosine function. Such systematic

deviation can be accounted for only by nonlinear theories (see Appendix A). Ultimately, the underlying neural mechanisms that generate the tuning properties need to be considered in more detailed theories. These tuning properties might be the outcome of learning processes based on correlated neuronal activities induced by movement and motion. In this paper, we have focused on several analytically tractable situations to emphasize the existence of general neuronal tuning properties that are insensitive to the actual mechanisms.

How preferred rotation axes may be used to update a static representation

If the cosine tuning of motion-sensitive neurons is determined essentially by geometric constraints regardless of the actual computational functions, then what is the value of these simple response properties? For three-dimensional object motion, a population of neurons tuned to translational direction and rotation axis should carry sufficient information to determine the instantaneous motion of any given object and therefore could be used to update the static view represented elsewhere. This allows future sensory and motor states to be predicted from the current static state.

Information about the *static view* of an object is represented in the ventral visual stream in the monkey cortex, leading to the inferotemporal (IT) area (Ungerleider and Mishkin, 1982). The response of a view-sensitive neuron in IT area typically drops off smoothly as the object is rotated away from its preferred view, around either the vertical axis (Perrett et al., 1991) or other axes (Logothetis and Pauls, 1995). View-dependent representations for three-dimensional objects have been studied theoretically (Poggio and Edelman, 1990; Ullman and Basri, 1991) and have motivated several recent psychophysical experiments (Edelman and Bülhoff, 1992; Bülhoff et al., 1995; Liu et al., 1995; Sinha and Poggio, 1996). The general idea of a view-dependent representation in the IT area is consistent with recent neurophysiological results, including single-unit recordings (Perrett et al., 1991; Logothetis and Pauls, 1995; Logothetis et al., 1995) and optical imaging data (Wang et al., 1996).

Information about the *instantaneous motion* of an object is represented in the dorsal visual stream, including areas MT, MST, superior temporal polysensory area, and the parietal cortex, such as area 7a. Given a population of motion-sensitive neurons tuned to translation and rotation, it should be possible to extract complete information about the instantaneous motion of any object. For example, a six-dimensional population vector can be used to reconstruct rigid motion. More efficient reconstruction methods may also be used and implemented by a biologically plausible feedforward network (Zhang et al., 1998). The same set of neurons can extract the motion of different objects by combining the activities of input neurons differently. Instantaneous translation and rotation determine how the current view of this object is changing at the moment and could be used to update the static view representation in the IT area.

Broad tuning to static views logically implies that each static view of an object elicits a certain activity pattern in the temporal cortex, and that as the view changes, the pattern of activity also changes smoothly, depending on the axis of rotation and direction of translation. A complete representation of the dynamic state of an object would require representing information about

both the current view and how the view is changing, so that the system can effectively update its internal state in accordance with the movement of the object. Such motion information might help improve the speed and reliability of the responses of view-specific neurons to a three-dimensional object during natural movements.

Conclusion

We have shown that simple generic tuning properties arise when an encoded sensory or motor variable reflects changes rather than static configurations. By linearizing the system locally for movement-sensitive neurons, the analysis reveals mechanism-insensitive tuning properties that mainly reflect the geometry of the problem rather than the exact encoding mechanisms, which could be much more complicated. Although a nonlinear analysis is also considered (Appendix A), the basic linear theory already captures some essential features of the biological data, such as sensory responses to visual pattern motions and directional tuning for reaching movements. The analysis predicts the existence of a preferred translational direction and a preferred rotation axis in space with cosine tuning functions for representing arbitrary three-dimensional object motion. For natural movements that have an intrinsically low dimensionality, combinations of variables become highly constrained and cannot be changed arbitrarily. It is precisely for these constrained movements that the mechanism-insensitive properties studied here may become useful. By contrast, the analysis may not apply for artificial movements such as computer-generated visual motion stimuli that do not satisfy any simplifying geometry constraints that occur in the real world. The brain should have more efficient representations for those stimulus features that are consistent with commonly encountered configurations in the real world. The analysis presented here may help to predict tuning properties of motion-sensitive neurons in unknown situations by providing a basic description of expected properties with which more detailed characterizations as well as potential deviations can be contrasted.

APPENDIX A: EXTENDED THEORIES

We first reformulate the linear tuning theory for motion-sensitive neurons in general terms and then make nonlinear extensions. Here it is assumed that the natural movements of interest can be parameterized by a low-dimensional vector variable:

$$\mathbf{x} = (x_1, x_2, \dots, x_D), \quad (\text{A1})$$

with D degrees of freedom. For example, $D = 3$ for reaching with a stereotypical arm posture, and $D = 6$ for rigid object motion. D can be larger when more independent variables are included.

Linear gradient theory

Assume that the mean firing rate of a motion-sensitive neuron is linearly related to the time derivative of a potential function $\Phi(\mathbf{x})$ of the state variable \mathbf{x} . This leads to the tuning rule:

$$f = f_0 + \frac{d}{dt}\Phi(\mathbf{x}) = f_0 + \mathbf{g}(\mathbf{x}) \cdot \mathbf{v}, \quad (\text{A2})$$

where f_0 is the baseline rate, the gradient:

$$\begin{aligned} \mathbf{g}(\mathbf{x}) &= (g_1(\mathbf{x}), g_2(\mathbf{x}), \dots, g_D(\mathbf{x})) \\ &= \nabla\Phi = \left(\frac{\partial\Phi}{\partial x_1}, \frac{\partial\Phi}{\partial x_2}, \dots, \frac{\partial\Phi}{\partial x_D} \right) \end{aligned} \quad (\text{A3})$$

is the generalized preferred direction, and:

$$\mathbf{v} = (v_1, v_2, \dots, v_D) = \dot{\mathbf{x}} = (\dot{x}_1, \dot{x}_2, \dots, \dot{x}_D) \quad (\text{A4})$$

is the generalized velocity. A gradient is often treated as a vector, although mathematically it is more properly called a one-form (Flanders, 1989), a distinction, however, that has no practical consequence for this paper. The necessary and sufficient condition for the preferred direction field $\mathbf{g} = (g_1, g_2, \dots, g_D)$ to be a gradient field is that:

$$\frac{\partial g_i}{\partial x_j} = \frac{\partial g_j}{\partial x_i}. \quad (\text{A5})$$

This is equivalent to the condition that:

$$\oint \mathbf{g}(\mathbf{x}) \cdot d\mathbf{x} = 0 \quad (\text{A6})$$

for any closed path, or that the potential function can be reconstructed from the preferred direction by:

$$\Phi(\mathbf{x}) = \Phi(\mathbf{x}_0) + \int_{\mathbf{x}_0}^{\mathbf{x}} \mathbf{g}(\mathbf{x}) \cdot d\mathbf{x}, \quad (\text{A7})$$

where the integral depends only on the end points \mathbf{x} and \mathbf{x}_0 , not on the path. In three-dimensional space, these conditions are equivalent to zero curl.

Linear nongradient theory

Assume directly a linear relationship between the firing rate and the components of the generalized speed velocity \mathbf{v} . This leads to the tuning rule:

$$f = f_0 + \mathbf{g}(\mathbf{x}) \cdot \mathbf{v} \quad (\text{A8})$$

without further constraint on the distribution of the preferred direction $\mathbf{g}(\mathbf{x})$. In particular, it may not be the gradient of a potential function. Thus, in general:

$$\frac{\partial g_i}{\partial x_j} \neq \frac{\partial g_j}{\partial x_i}, \quad (\text{A9})$$

and the path integral may depend on the path. For a fixed \mathbf{x} , the local tuning property of this neuron is the same as that predicted by the gradient theory. The difference is that the nongradient theory allows an arbitrary global distribution of the preferred direction field, whereas the gradient theory admits only a gradient field.

Coordinate-system independence

Both the gradient and the nongradient theories are independent of which variables are chosen to parameterize the movements. Suppose the old vector variable \mathbf{x} and a new variable $\tilde{\mathbf{x}}$ are related by:

$$\mathbf{x} = \mathbf{h}(\tilde{\mathbf{x}}), \quad (\text{A10})$$

then the velocities $\mathbf{v} = d\mathbf{x}/dt$ and $\tilde{\mathbf{v}} = d\tilde{\mathbf{x}}/dt$ in the two coordinates are related linearly by:

$$\mathbf{v} = \mathbf{J}(\tilde{\mathbf{x}})\tilde{\mathbf{v}}, \quad (\text{A11})$$

where $\mathbf{J}(\tilde{\mathbf{x}}) = d\mathbf{h}(\tilde{\mathbf{x}})/d\tilde{\mathbf{x}}$ is the Jacobian matrix.

The tuning rule has the same form in both coordinate systems:

$$f = f_0 + \mathbf{g}(\mathbf{x}) \cdot \mathbf{v} = f_0 + \tilde{\mathbf{g}}(\tilde{\mathbf{x}}) \cdot \tilde{\mathbf{v}}, \quad (\text{A12})$$

where the new preferred direction is:

$$\tilde{\mathbf{g}}(\tilde{\mathbf{x}}) = \mathbf{g}(\mathbf{h}(\tilde{\mathbf{x}}))\mathbf{J}(\tilde{\mathbf{x}}). \quad (\text{A13})$$

If a potential function exists such that the old preferred direction is a gradient field:

$$\mathbf{g}(\mathbf{x}) = \nabla\Phi(\mathbf{x}), \quad (\text{A14})$$

then the preferred direction field is still a gradient field with respect to the new variable $\tilde{\mathbf{x}}$:

$$\tilde{\mathbf{g}}(\tilde{\mathbf{x}}) = \tilde{\nabla}\tilde{\Phi}(\tilde{\mathbf{x}}), \quad (\text{A15})$$

where the new potential function is:

$$\tilde{\Phi}(\tilde{\mathbf{x}}) = \Phi(\mathbf{h}(\tilde{\mathbf{x}})). \quad (\text{A16})$$

The choice of variable is arbitrary, but convenience favors coordinates that make the results easiest to interpret.

Nonlinear theory: circular normal tuning

The circular normal tuning function for firing rate has the general form:

$$f = A + B \exp(K \cos \alpha), \quad (\text{A17})$$

which has one more free parameter and tends to fit data better (compare Figs. 2 and 8) than a cosine tuning function of the general form:

$$f = A' + B' \cos \alpha, \quad (\text{A18})$$

where α is the angular variable of interest, and A, B, K, A', B' are parameters.

The circular normal function can mimic either a cosine or a Gaussian. When K is very small, $\exp(K \cos \alpha) \sim 1 + K \cos \alpha$ so that the circular normal function in Equation A17 approaches the cosine function in Equation A18 with $A' = A + B$ and $B' = BK$. When K is large, $\cos \alpha \sim 1 - \alpha^2/2$ so that the circular normal function approaches a narrow Gaussian function with the variance $1/K$.

How can we generate a circular normal tuning function? Because the time-derivative equation for firing rate:

$$f - f_0 = \frac{d\Phi}{dt} \quad (\text{A19})$$

can yield a cosine tuning function, as shown before, the modified equation of the form:

$$\log(af + b) = \frac{d\Phi}{dt} \quad (\text{A20})$$

should yield a circular normal tuning function, where a and b are parameters. The result is a tuning rule of the form:

$$f = A + B \exp(pv \cos \alpha), \quad (\text{A21})$$

where angle α specifies movement direction and v is movement speed. This modification introduces the following testable effects on speed modulation. If parameter p is very small, the tuning function is close to a cosine, and the firing rate increases approximately linearly with speed, as before. When the parameter p is larger, the tuning function is closer to a Gaussian, and the firing rate should increase faster than linearly with speed. The directional tuning also becomes narrower with higher speed. These

effects are similar to those shown in Figure 9A for quadratic terms. See related discussion in the next section.

Although Equation A20 can lead to a circular normal function, it does not specify how this occurs. One plausible biological mechanism is a recurrent network with appropriate lateral connections, which can generate a tuning curve closer to a circular normal function than to a cosine function (Pouget et al., 1998).

Nonlinear theory: acceleration tuning and quadratic speed modulation

In this section the basic tuning theory is generalized by including the second temporal derivative. This leads to acceleration tuning, nonlinear speed modulation, and departure from perfect cosine directional tuning.

Assume that the firing rate of a neuron contains not only the first temporal derivative of a potential function, but also the second temporal derivative of another potential function:

$$f = f_0 + \frac{d\Phi}{dt} + \frac{d^2\Psi}{dt^2}, \quad (\text{A22})$$

where f_0 is the baseline rate, $\Phi = \Phi(\mathbf{x})$ and $\Psi = \Psi(\mathbf{x})$ are two unknown potential functions of a vector variable $\mathbf{x} = (x_1, x_2, \dots, x_D)$. The special case $\Psi = 0$ reduces to what has been considered before.

This assumption leads to the new tuning rule:

$$f = f_0 + \mathbf{p}^1 \cdot \mathbf{v} + \mathbf{p}^2 \cdot \mathbf{a} + \sum_{i,j=1}^D h_{ij}v_i v_j, \quad (\text{A23})$$

where $\mathbf{v} = (v_1, v_2, \dots, v_D) = d\mathbf{x}/dt$ is generalized velocity, $\mathbf{a} = d\mathbf{v}/dt$ is generalized acceleration, $\mathbf{p}^1 = \nabla\Phi$ is the preferred direction for velocity, $\mathbf{p}^2 = \nabla\Psi$ is the preferred direction for acceleration, and $h_{ij} = \partial^2\Psi/\partial x_i \partial x_j$ is the Hessian matrix of second derivatives.

Example 1: Reaching movement

Here the vector variable $\mathbf{x} = (x_1, x_2, x_3) = (x, y, z)$ describes the hand position. The dot product terms in Equation A23 mean that both the velocity and the acceleration have preferred directions and cosine directional tuning functions, together with multiplicative linear modulation by the speed or the magnitude of acceleration. Ashe and Georgopoulos (1994) included acceleration terms in a different regression formula and found a small number of cells related to hand acceleration. Systematic tests are needed to determine whether the acceleration tuning predicted by Equation A23 really exists.

Example 2: Rigid object motion

Here the vector variable $\mathbf{x} = (x_1, x_2, \dots, x_6) = (x, y, z, \theta_1, \theta_2, \theta_3)$ describes the object's position and orientation in space. By transforming $(\theta_1, \theta_2, \theta_3)$ into the angular velocity in physical space, the tuning rule becomes:

$$f = f_0 + \mathbf{p}^1 \cdot \mathbf{v} + \mathbf{p}^2 \cdot \mathbf{a} + \mathbf{q}^1 \cdot \boldsymbol{\omega} + \mathbf{q}^2 \cdot \boldsymbol{\alpha} + \sum_{i,j=1}^3 (H_{ij}v_i v_j + I_{ij}v_i \omega_j + J_{ij}\omega_i \omega_j), \quad (\text{A24})$$

where $\mathbf{v} = (\dot{x}, \dot{y}, \dot{z})$ is the translational velocity, $\mathbf{a} = d\mathbf{v}/dt$ is the translational acceleration, $\boldsymbol{\omega} = (\omega_1, \omega_2, \omega_3)$ is the angular velocity, $\boldsymbol{\alpha} = d\boldsymbol{\omega}/dt$ is the angular acceleration, and H_{ij}, I_{ij}, J_{ij} are

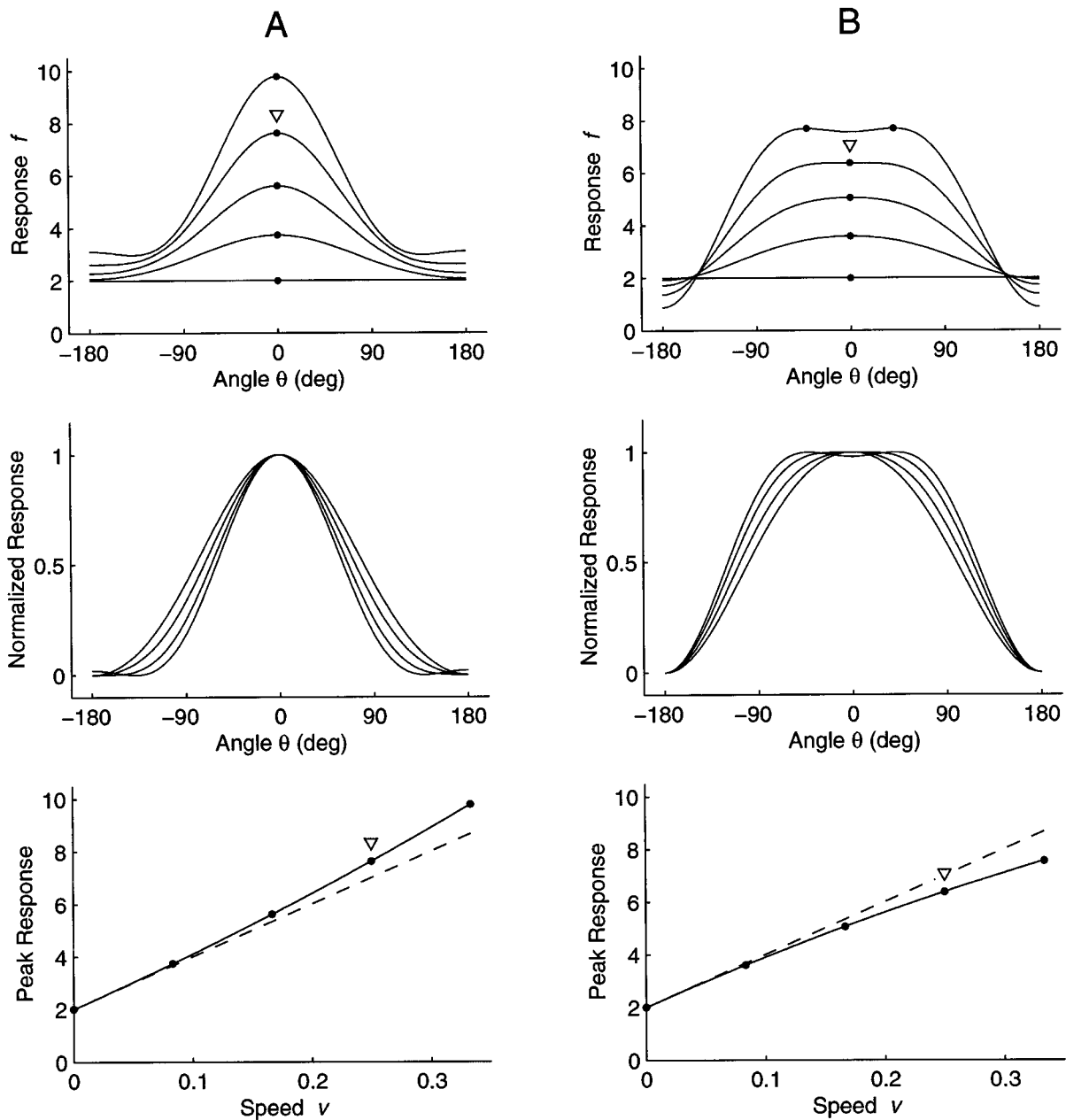


Figure 9. Effects of quadratic terms on directional tuning curve and speed modulation, here showing the tuning functions $f = 2 + 10(v + v \cos \theta \pm v^2 \cos 2\theta)$, which have a single peak when the speed $v \leq 0.25$, with the critical cases indicated by ∇ . A, Positive sign: As the speed increases, the tuning curve becomes progressively narrower, and its peak response grows more slowly than a linear function (dashed line). B, Negative sign: As the speed increases, the tuning curve becomes progressively wider, and its peak response grows more slowly than a linear function (dashed line).

determined by the Hessian matrix and matrix \mathbf{M} as in Equation 26. Therefore, angular acceleration should also have a cosine tuning with a preferred direction \mathbf{q}^2 . Although there is some evidence that some MT cells can provide information about acceleration of a visual target (Movshon et al., 1990; Lisberger et al., 1995), systematic tests for acceleration tuning with realistic objects have not been performed.

Effects of quadratic terms

The quadratic speed terms imply both nonlinear speed modulation and higher-order Fourier components for directional tuning

that are speed dependent. To see this, consider a two-dimensional reaching example with the hand velocity:

$$\mathbf{v} = (v_1, v_2) = (v \cos \phi, v \sin \phi), \tag{A25}$$

where v is the speed and ϕ is the angle of movement direction. The quadratic terms can always be written as:

$$\sum_{i,j=1}^2 h_{ij} v_i v_j = (h_{11} \cos^2 \phi + 2h_{12} \cos \phi \sin \phi + h_{22} \sin^2 \phi) v^2 = [a + b \cos(2\phi + \phi_0)] v^2, \tag{A26}$$

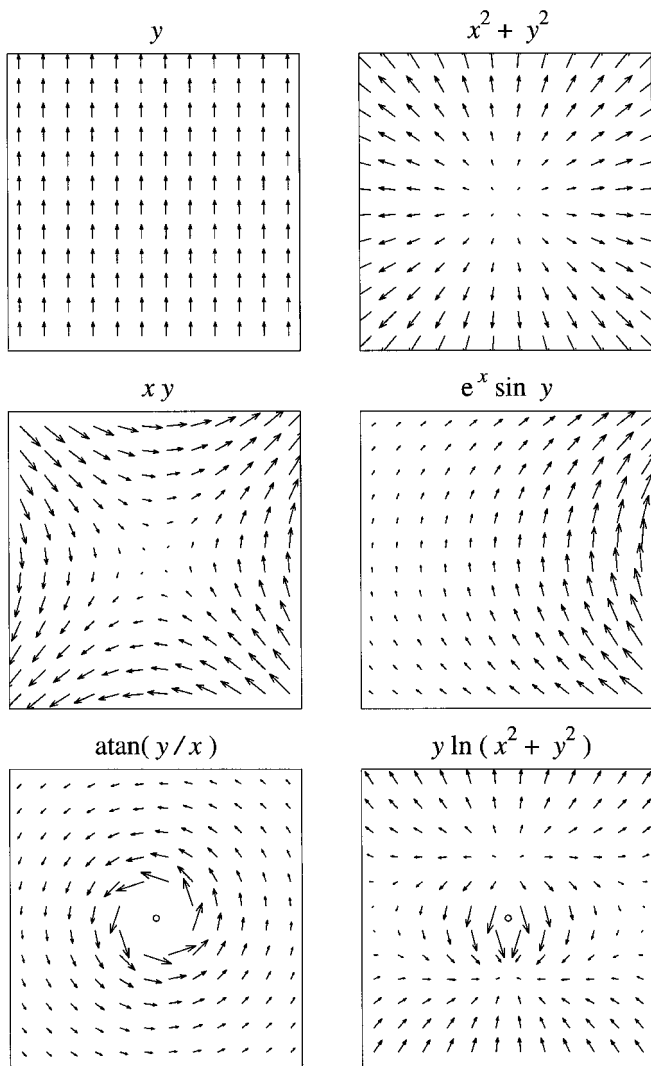


Figure 10. Curl-free vector fields may have various visual appearances. These examples, obtained as the gradients of the potential functions as indicated, have zero curl everywhere, except at a singular point in the two cases at the bottom. Plot range: always from -1 to 1 for both axes.

where $h_{12} = h_{21}$, and the coefficients a , b and the phase shift φ_0 depend only on h_{ij} . For comparison, the linear speed term can be written as $\mathbf{p}^1 \cdot \mathbf{v} = pv \cos(\phi + \varphi_1)$, with a phase shift φ_1 .

The second Fourier component can either sharpen or broaden the original cosine function, depending on its sign. As illustrated in Figure 9, if the tuning curve is sharpened by the second component, it becomes even sharper as the speed increases; if the tuning curve is broadened, then it becomes even broader as the speed increases. However, the amplitude of the second component ($\cos 2\theta$) should be no more than one-fourth of that of the first one ($\cos \theta$) to ensure that the tuning curve has only a single peak. This limits the effects from the second Fourier term. For speed modulation, the quadratic speed factor produces only a slight bend (Fig. 9) and is too weak by itself to produce U-shaped or \cap -shaped curves for some neurons in area MT (Maunsell and Van Essen, 1983; Rodman and Albright, 1987) and MST (Orban et al., 1995; Duffy and Wurtz, 1997a). Rodman and Albright (1987) also showed that the average tuning widths of MT neurons were insensitive to speed, although the typical experimental errors for individual neurons might mask the small effects shown

here. Thus, the second Fourier component with squared speed may help improve data fitting (compare Fig. 2), but probably only within a narrow range of speeds.

APPENDIX B: LINEAR VECTOR FIELD FOR DATA ANALYSIS

Linear vector field from experimental data

As shown in the main text, the preferred direction and the preferred rotation axis may be generated by gradient fields of a potential function. Here we consider how to test the gradient condition experimentally. In two and three dimensions, where the curl can be defined, a vector field generated as the gradient of any potential function is always curl free (Fig. 10). In particular, a two-dimensional vector field $(u(x, y), v(x, y))$ has both zero curl and zero divergence if and only if

$$f(z) = v(x, y) + iu(x, y) \quad (\text{B1})$$

is analytic or differentiable with respect to the complex variable $z = x + iy$, because of the Cauchy-Riemann differentiability conditions. One can use templates of curl-free vector fields generated by gradient of potential functions to interpolate arbitrary vector fields (Mussa-Ivaldi, 1992). Therefore, the curl-free condition alone is too flexible because it can locally distort fields to accommodate any sparsely sampled vector field data.

For testing the gradient-field condition or the curl-free condition with sparsely sampled data points, an additional smoothness constraint is needed. Linearity is a reasonable smoothness requirement, at least for a local region, such as in measurement of local force fields (Mussa-Ivaldi et al., 1985; Giszter et al., 1993), and in local optic flow analysis (Koenderink and van Doorn, 1976). A linear vector field has the general form:

$$\mathbf{p} = \mathbf{A}\mathbf{x} + \mathbf{b}, \quad (\text{B2})$$

where the column vector $\mathbf{x} = (x_1, x_2, \dots, x_D)^T$ contains the variables of interest, \mathbf{A} is a constant matrix:

$$\mathbf{A} = \begin{pmatrix} a_{11} & a_{12} & \cdots & a_{1D} \\ a_{21} & a_{22} & \cdots & a_{2D} \\ \vdots & \vdots & \ddots & \vdots \\ a_{D1} & a_{D2} & \cdots & a_{DD} \end{pmatrix}, \quad (\text{B3})$$

and \mathbf{b} is a constant vector. If the linear vector field is the gradient of an unknown potential function, then this function should have the general quadratic form:

$$\Phi(\mathbf{x}) = \frac{1}{2} \mathbf{x}^T \mathbf{B} \mathbf{x} + \mathbf{b}^T \mathbf{x} + C, \quad (\text{B4})$$

which has the gradient:

$$\mathbf{p} = \nabla \Phi(\mathbf{x}) = \frac{1}{2} (\mathbf{B} + \mathbf{B}^T) \mathbf{x} + \mathbf{b}, \quad (\text{B5})$$

with \mathbf{b} a constant column vector, and T for transpose. To be consistent with Equation B2, matrix \mathbf{A} must be symmetric:

$$\mathbf{A}^T = \mathbf{A}, \quad (\text{B6})$$

in which case we can set $\mathbf{B} = \mathbf{A}$. This symmetry condition in Equation B6 is the necessary and sufficient condition for a linear vector field to be a gradient field. It includes the curl-free condition for $D = 3$ as the special case, because the curl:

$$\text{curl } \mathbf{p} = (a_{32} - a_{23}, a_{13} - a_{31}, a_{21} - a_{12}) \quad (\text{B7})$$

vanishes if and only if the matrix \mathbf{A} is symmetric.

To determine \mathbf{A} and \mathbf{b} from data vectors $\mathbf{p}_1, \mathbf{p}_2, \dots, \mathbf{p}_N$ sampled at positions $\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N$, respectively, we require the total number of data points:

$$N \geq D + 1 \tag{B8}$$

in D -dimensional space, because matrix \mathbf{A} and vector \mathbf{c} contain $D^2 + D$ scalar unknowns, whereas N data points provide DN scalar values. For example, the two-dimensional case requires a minimum of three data points, and the three-dimensional case requires a minimum of four data points.

The least-square solution is:

$$\mathbf{A} = \mathbf{P}\mathbf{X}^\dagger, \tag{B9}$$

where

$$\mathbf{X}^\dagger = \mathbf{X}^T(\mathbf{X}\mathbf{X}^T)^{-1} \tag{B10}$$

is the Moore-Penrose pseudoinverse, and the matrices are defined as:

$$\mathbf{P} = [(\mathbf{p}_1 - \mathbf{p}_0), (\mathbf{p}_2 - \mathbf{p}_0), \dots, (\mathbf{p}_N - \mathbf{p}_0)], \tag{B11}$$

$$\mathbf{X} = [(\mathbf{x}_1 - \mathbf{x}_0), (\mathbf{x}_2 - \mathbf{x}_0), \dots, (\mathbf{x}_N - \mathbf{x}_0)], \tag{B12}$$

which are made of column vectors as shown, and:

$$\mathbf{p}_0 = \frac{1}{N} \sum_{k=1}^N \mathbf{p}_k, \tag{B13}$$

$$\mathbf{x}_0 = \frac{1}{N} \sum_{k=1}^N \mathbf{x}_k, \tag{B14}$$

$$\mathbf{b} = \mathbf{p}_0 - \mathbf{A}\mathbf{x}_0. \tag{B15}$$

The above solution minimizes:

$$\sum_{k=1}^N |(\mathbf{p}_k - \mathbf{p}_0) - \mathbf{A}(\mathbf{x}_k - \mathbf{x}_0)|^2, \tag{B16}$$

which vanishes only when all the data points can be fit exactly by the linear model.

Examples of the simplex method

In this section, we illustrate an alternative formulation of the curl-free condition with minimal data points in three-dimensional space, which can be extended readily to other dimensions. For local interpolation in three-dimensional space, a vector field should at a minimum be sampled at four locations 1, 2, 3, 4, not all lying in the same plane (Fig. 11). Suppose the coordinates of the four points are $\mathbf{x}_1, \mathbf{x}_2, \mathbf{x}_3, \mathbf{x}_4$, and the corresponding data vectors are $\mathbf{p}_1, \mathbf{p}_2, \mathbf{p}_3, \mathbf{p}_4$. The simplex method is based on the fact that any point \mathbf{x} in three-dimensional space can be expressed as:

$$\mathbf{x} = \lambda_1\mathbf{x}_1 + \lambda_2\mathbf{x}_2 + \lambda_3\mathbf{x}_3 + \lambda_4\mathbf{x}_4, \tag{B17}$$

where the coefficients are unique, satisfying the constraint:

$$\lambda_1 + \lambda_2 + \lambda_3 + \lambda_4 = 1. \tag{B18}$$

The linearly interpolated vector at this position is:

$$\mathbf{p} = \lambda_1\mathbf{p}_1 + \lambda_2\mathbf{p}_2 + \lambda_3\mathbf{p}_3 + \lambda_4\mathbf{p}_4. \tag{B19}$$

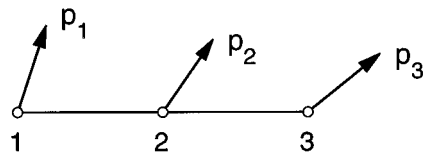
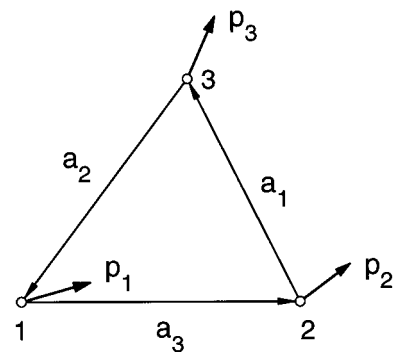
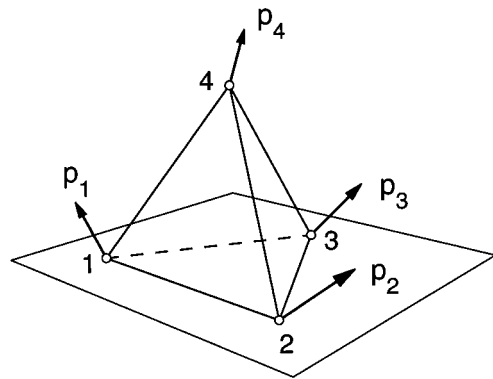


Figure 11. Interpolating a vector field linearly in three-dimensional space requires at least four data vectors to be measured at locations not all in the same plane. In two-dimensional space, at least three data vectors are required at locations not all from the same line. The case shown at the bottom is degenerate.

Similar equations with fewer variables hold for the one- and two-dimensional cases.

To derive an integral formula for the curl-free condition, first integrate along the straight line segment from \mathbf{x}_1 to \mathbf{x}_2 with a linearly interpolated vector, yielding:

$$\int_{\mathbf{x}_1}^{\mathbf{x}_2} \mathbf{p} \cdot d\mathbf{l} = \frac{1}{2} (\mathbf{p}_1 + \mathbf{p}_2) \cdot (\mathbf{x}_2 - \mathbf{x}_1). \tag{B20}$$

To evaluate the closed path integral along the triangle 123 in Figure 11, use this formula and the equality $\mathbf{a}_1 + \mathbf{a}_2 + \mathbf{a}_3 = 0$, with $\mathbf{a}_1 \equiv \mathbf{x}_3 - \mathbf{x}_2$, and so on, to obtain:

$$\oint_{123} \mathbf{p} \cdot d\mathbf{l} = -\frac{1}{2} (\mathbf{p}_1 \cdot \mathbf{a}_1 + \mathbf{p}_2 \cdot \mathbf{a}_2 + \mathbf{p}_3 \cdot \mathbf{a}_3), \tag{B21}$$

which must vanish if the field is curl free. This formula is valid in both two- and three-dimensional cases. For the tetrahedron in Figure 11, a similar closed path integral along the edges should vanish. Because there are only three independent loops, we can choose, for instance:

$$\oint_{123} \mathbf{p} \cdot d\mathbf{l} = \oint_{124} \mathbf{p} \cdot d\mathbf{l} = \oint_{234} \mathbf{p} \cdot d\mathbf{l} = 0. \quad (\text{B22})$$

Although equivalent to the matrix symmetry in Equation B6, the path integral formulas are more intuitive and express the curl-free condition in terms of directly measurable quantities, as in Equation B21.

In the experiment by Caminiti et al. (1990), the preferred direction of a motor cortical neuron was sampled at three points at equal distance, similar to the case in the bottom diagram in Figure 11. Here the linearity of the vector field only entails that:

$$\mathbf{p}_2 = \frac{1}{2} (\mathbf{p}_1 + \mathbf{p}_3), \quad (\text{B23})$$

which is a special case of Equation B19. The curl-free condition cannot be tested here. With linear interpolation, at least four starting hand positions should be sampled (Fig. 11).

REFERENCES

- Adelson EH, Bergen JR (1985) Spatiotemporal energy models for the perception of motion. *J Opt Soc Am A* 2:284–299.
- Albright TD (1984) Direction and orientation selectivity of neurons in visual area MT of the macaque. *J Neurophysiol* 52:1106–1130.
- Albright TD (1992) Form-cue invariant motion processing in primate visual cortex. *Science* 255:1141–1143.
- Alexander GE, Crutcher MD (1990) Neural representations of the target (goal) of visually guided arm movements in three motor areas of the monkey. *J Neurophysiol* 64:164–178.
- Allman JM, Miezin F, McGuinness 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 RA, Snyder LH, Bradley DC, Xing J (1997) Multimodal representation of space in the posterior parietal cortex and its use in planning movements. *Ann Rev Neurosci* 20:303–330.
- Ashe J, Georgopoulos AP (1994) Movement parameters and neural activity in motor cortex and area 5. *Cereb Cortex* 6:590–600.
- Baker J, Goldberg J, Hermann G, Peterson B (1984) Optimal response planes and canal convergence in secondary neurons in vestibular nuclei of alert cats. *Brain Res* 294:133–137.
- Bradley DC, Maxwell M, Andersen RA, Banks MS, Shenoy KV (1996) Mechanisms of heading perception in primate visual cortex. *Science* 273:1544–1547.
- Bülthoff HH, Edelman SY, Tarr MJ (1995) How are three-dimensional objects represented in the brain? *Cereb Cortex* 3:247–260.
- Buneo CA, Soechting JF, Flanders M (1997) Postural dependence of muscle actions: implications for neural control. *J Neurosci* 17:2128–2142.
- Buračas GT, Albright TD (1996) Contribution of area MT to perception of three-dimensional shape: a computational study. *Vision Res* 36:869–887.
- Caminiti R, Johnson PB, Urbano A (1990) Making arm movements within different parts of space: dynamic aspects of the primate motor cortex. *J Neurosci* 10:2039–2058.
- Caminiti R, Johnson PB, Galli C, Ferraina S, Burnod Y (1991) Making arm movements within different parts of space: premotor and motor cortical representation of a coordinate system for reaching to visual targets. *J Neurosci* 11:1182–1197.
- Cheng K, Hasegawa T, Saleem KS, Tanaka K (1994) Comparison of neuronal selectivity for stimulus speed, length, and contrast in the prestriate visual areas V4 and MT of the macaque monkey. *J Neurophysiol* 71:2269–2280.
- Cohen DA, Prud'homme MJ, Kalaska JF (1994) Tactile activity in primate primary somatosensory cortex during active arm movements: correlation with receptive field properties. *J Neurophysiol* 71:161–172.
- Colby CL, Duhamel J-R, Goldberg ME (1993) Ventral intraparietal area of the macaque: anatomic location and visual response properties. *J Neurophysiol* 69:902–914.
- Crammond DJ, Kalaska JF (1996) Differential relation of discharge in primary motor cortex and premotor cortex to movements versus actively maintained postures during a reaching task. *Exp Brain Res* 108:45–61.
- Dobbins AC, Jeo RM, Fiser J, Allman JM (1998) Distance modulation of neural activity in the visual cortex. *Science* 281:552–555.
- Duffy CJ, Wurtz RH (1995) Response of monkey MST neurons to optic flow stimuli with shifted centers of motion. *J Neurosci* 15:5192–5208.
- Duffy CJ, Wurtz RH (1997a) Medial superior temporal area neurons respond to speed patterns in optic flow. *J Neurosci* 17:2839–2851.
- Duffy CJ, Wurtz RH (1997b) Planar directional contributions to optic flow response in MST neurons. *J Neurophysiol* 77:782–796.
- Edelman S, Bülthoff HH (1992) Orientation dependence in the recognition of familiar and novel views of three-dimensional objects. *Vision Res* 32:2385–2400.
- Eifuku S, Wurtz RH (1998) Response to motion in extrastriate area MSTl: center-surround interactions. *J Neurophysiol* 80:282–296.
- Fetz EE (1992) Are movement parameters recognizably coded in the activity of single neurons? *Behav Brain Sci* 15:679–690.
- Flanders H (1989) *Differential forms with applications to the physical sciences*. New York: Dover.
- Flanders M, Herrmann U (1992) Two components of muscle activation: scaling with the speed of arm movement. *J Neurophysiol* 67:931–943.
- Flanders M, Soechting JF (1990) Arm muscle activation for static forces in three-dimensional space. *J Neurophysiol* 64:1818–1837.
- Fortier PA, Kalaska JF, Smith AM (1989) Cerebellar neuronal activity related to whole-arm reaching movements in the monkey. *J Neurophysiol* 62:198–211.
- Fortier PA, Smith AM, Kalaska JF (1993) Comparison of cerebellar and motor cortex activity during reaching: directional tuning and response variability. *J Neurophysiol* 69:1136–1149.
- Fu Q-G, Suarez JI, Ebner TJ (1993) Neuronal specification of direction and distance during reaching movements in the superior precentral premotor area and primary motor cortex of monkeys. *J Neurophysiol* 70:2097–2116.
- Gaal G (1993) Calculation of movement direction from firing activities of neurons in intrinsic co-ordinate systems defined by their preferred directions. *J Theor Biol* 162:103–130.
- Geesaman BJ, Andersen RA (1996) The analysis of complex motion patterns by form/cue invariant MSTd neurons. *J Neurosci* 16:4716–4732.
- Georgopoulos AP, Schwartz AB, Kettner RE (1986) Neuronal population coding of movement direction. *Science* 233:1416–1419.
- Georgopoulos AP, Kettner RE, Schwartz AB (1988) Primate motor cortex and free arm movements to visual targets in three-dimensional space. II. Coding of the direction of movement by a neuronal population. *J Neurosci* 8:2928–2937.
- Georgopoulos AP, Crutcher MD, Schwartz AB (1989a) Cognitive spatial-motor processes. 3. Motor cortical prediction of movement direction during an instructed delay period. *Exp Brain Res* 75:183–194.
- Georgopoulos AP, Lurito JT, Petrides M, Schwartz AB, Massey JT (1989b) Mental rotation of the neuronal population vector. *Science* 243:234–236.
- Georgopoulos AP, Ashe J, Smyrnis N, Taira M (1992) The motor cortex and the coding of force. *Science* 256:1692–1695.
- Giszter SF, Mussa-Ivaldi FA, Bizzi E (1993) Convergent force fields organized in the frog's spinal cord. *J Neurosci* 13:467–491.
- Goldstein H (1980) *Classical mechanics*, Ed 2. Reading, MA: Addison-Wesley.
- Graf W, Baker J, Peterson BW (1993) Sensorimotor transformation in the cat's vestibuloocular flex system. I. Neuronal signals coding spatial coordination of compensatory eye movements. *J Neurophysiol* 70:2425–2441.
- Graziano MS, Andersen RA, Snowden RJ (1994) Tuning of MST neurons to spiral motions. *J Neurosci* 14:54–67.
- Heeger DJ (1993) Modeling simple-cell direction selectivity with normalized, half-squared, linear operators. *J Neurophysiol* 70:1885–1898.
- Heeger DJ, Jepson A (1990) Visual perception of three-dimensional motion. *Neural Comput* 2:129–137.
- Hirase H, Czurkó A, Csicsvari J, Buzsáki G (1998) Hippocampal pyra-

- midal neurons sustain discharge frequency when the rat is “clamped” in space. *Soc Neurosci Abstr* 24:921.
- Kalaska JF, Cohen DAD, Hyde ML, Prud’homme M (1989) A comparison of movement direction-related versus load direction-related activity in primate motor cortex, using a two-dimensional reaching task. *J Neurosci* 9:2080–2102.
- Kalaska JF, Cohen DAD, Prud’homme M, Hyde ML (1990) Parietal area 5 neuronal activity encodes movement kinematics, not movement dynamics. *Exp Brain Res* 80:351–364.
- Kettner RE, Schwartz AB, Georgopoulos AP (1988) Primate motor cortex and free arm movements to visual targets in three-dimensional space. III. Positional gradients and population coding of movement direction from various movement origins. *J Neurosci* 8:2938–2947.
- Koenderink JJ, van Doorn AJ (1976) Local structure of movement parallax of the plane. *J Opt Soc Am* 66:717–723.
- Krauzlis RJ, Lisberger SG (1996) Directional organization of eye movement and visual signals in the floccular lobe of the monkey cerebellum. *Exp Brain Res* 109:289–302.
- Lacquaniti F, Terzuolo C, Viviani P (1983) The law relating the kinematic and figural aspects of drawing movements. *Acta Psychologica* 54:115–130.
- Lappe M, Bremmer F, Pekel M, Thiele A, Hoffmann K-P (1996) Optic flow processing in monkey STS: a theoretical and experimental approach. *J Neurosci* 16:6265–6285.
- Lisberger SG, O’Keefe LP, Kahlon M, Mahncke HW, Movshon JA (1995) Temporal dynamics in responses of macaque MT cells during target acceleration. *Soc Neurosci Abstr* 21:664.
- Liu, Z-L, Knill DC, Kersten D (1995) Object classification for human and ideal observers. *Vision Res* 35:549–568.
- Logothetis NK, Pauls J (1995) Psychophysical and physiological evidence for viewer-centered object representation in the primate. *Cereb Cortex* 3:270–288.
- Logothetis NK, Pauls J, Poggio T (1995) Shape representation in the inferior temporal cortex of monkeys. *Curr Biol* 5:552–563.
- Longuet-Higgins HC, Prazdny K (1980) The interpretation of a moving retinal image. *Proc R Soc Lond B Biol Sci* 208:385–397.
- Maunsell JHR, Van Essen DC (1983) Functional properties of neurons in middle temporal visual area of the macaque monkey. I. Selectivity for stimulus direction, speed, and orientation. *J Neurophysiol* 49:1127–1147.
- McNaughton BL, Barnes CA, O’Keefe J (1983) The contributions of position, direction, and velocity to single unit activity in the hippocampus of freely-moving rats. *Exp Brain Res* 52:41–49.
- Moody SL, Zipser D (1998) A model of reaching dynamics in primary motor cortex. *J Cognit Neurosci* 10:35–45.
- Moody SL, Wise SP, di Pellegrino G, Zipser D (1998) A model that accounts for activity in primate frontal cortex during a delayed matching-to-sample task. *J Neurosci* 18:399–410.
- Moran DW, Schwartz AB (1999) Motor cortical representation of speed and direction during reaching. *J Neurophysiol*, in press.
- Moran DW, Kakavand A, Schwartz AB (1995) Effect of a virtual reality illusion on motor cortical population vectors during drawing. *Soc Neurosci Abstr* 21:2076.
- Movshon JA, Adelson EH, Gizzi MS, Newsome WT (1985) The analysis of moving visual patterns. In: *Pattern recognition mechanisms* (Chagas C, Gattass R, Gross C, eds), pp 117–151. New York: Springer-Verlag.
- Movshon JA, Lisberger SG, Krauzlis RJ (1990) Visual cortical signals supporting smooth pursuit eye movements. *Cold Spring Harbor Symp Quant Biol* 55:707–716.
- Mussa-Ivaldi FA (1988) Do neurons in the motor cortex encode movement direction? An alternative hypothesis. *Neurosci Lett* 91:106–111.
- Mussa-Ivaldi FA (1992) From basis functions to basis fields: vector field approximation from sparse data. *Biol Cybern* 67:479–489.
- Mussa-Ivaldi FA, Hogan N, Bizzi E (1985) Neural, mechanical, and geometric factors subserving arm posture in humans. *J Neurosci* 5:2732–2743.
- Newsome WT, Wurtz RH, Komatsu H (1988) Relation of cortical areas MT and MST to pursuit eye movements. II. Differentiation of retinal from extraretinal inputs. *J Neurophysiol* 60:604–620.
- Nowlan S, Sejnowski TJ (1995) A selection model for motion processing in area MT of primates. *J Neurosci* 15:1195–1214.
- Orban GA, Lagae L, Raiguel S, Xiao D, Maes H (1995) The speed tuning of medial superior temporal (MST) cell responses to optic-flow components. *Perception* 24:269–285.
- Pekel M, Lappe M, Bremmer F, Thiele A, Hoffmann K-P (1996) Neuronal responses in the motion pathway of the macaque monkey to natural optic flow stimuli. *NeuroReport* 22:884–888.
- Pellizzer G, Sargent P, Georgopoulos AP (1995) Motor cortical activity in a context-recall task. *Science* 269:702–705.
- Perrett DI, Oram MW, Harries MH, Bevan R, Hietanen JK, Benson PJ, Thomas S (1991) View-centered and object-centered coding of heads in the macaque temporal cortex. *Exp Brain Res* 86:159–173.
- Perrone JA, Stone LS (1994) A model of self-motion estimation within primate extrastriate visual cortex. *Vision Res* 34:2917–2938.
- Poggio T, Edelman S (1990) A network that learns to recognize three-dimensional objects. *Nature* 343:263–266.
- Poggio T, Verri A, Torre V (1991) Green theorems and qualitative properties of the optical flow. MIT A.I. Memo 1289. Cambridge, MA: MIT.
- Pouget A, Zhang K-C, Deneve S, Latham PE (1998) Statistically efficient estimation using population code. *Neural Comput* 10:373–401.
- Prud’homme MJL, Kalaska JF (1994) Proprioceptive activity in primate primary somatosensory cortex during active arm reaching movements. *J Neurophysiol* 72:2280–2301.
- Qian N, Andersen RA (1994) Transparent motion perception as detection of unbalanced motion signals. II. Physiology. *J Neurosci* 14:7367–7380.
- Raiguel S, Van Hulle MM, Xiao D-K, Orban GA (1995) Shape and spatial distribution of receptive fields and antagonistic motion surrounds in the middle temporal area (V5) of the macaque. *Eur J Neurosci* 7:2064–2082.
- Recanzone GH, Wurtz RH, Schwarz U (1997) Responses of MT and MST neurons to one and two moving objects in the receptive field. *J Neurophysiol* 78:2904–2915.
- Rodman HR, Albright TD (1987) Coding of visual stimulus velocity in area MT of the macaque. *Vision Res* 27:2035–2048.
- Roy J-P, Komatsu H, Wurtz RH (1992) Disparity sensitivity of neurons in monkey extrastriate area MST. *J Neurosci* 12:2478–2492.
- 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. *J Neurosci* 6:145–157.
- Sakata H, Shibutani H, Ito Y, Tsurugai K, Mine S, Kusunoki M (1994) Functional properties of rotation-sensitive neurons in the posterior parietal association cortex of the monkey. *Exp Brain Res* 101:183–202.
- Salinas E, Abbott LF (1994) Vector reconstruction from firing rates. *J Comput Neurosci* 1:89–107.
- Salinas E, Abbott LF (1997) A model of multiplicative neural responses in parietal cortex. *Proc Nat Acad Sci USA* 93:11956–11961.
- Sanger TD (1994) Theoretical considerations for the analysis of population coding in motor cortex. *Neural Comput* 6:12–21.
- Schaafsma SJ, Duysens J (1996) Neurons in the ventral intraparietal area of awake macaque monkey closely resemble neurons in the dorsal part of the medial superior temporal area in their responses to optic flow patterns. *J Neurophysiol* 76:4056–4068.
- Schwartz AB (1993) Motor cortical activity during drawing movements: population representation during sinusoid tracing. *J Neurophysiol* 70:28–36.
- Schwartz AB (1994) Direct cortical representation of drawing. *Science* 265:540–542.
- Schwartz AB, Kettner RE, Georgopoulos AP (1988) Primate motor cortex and free arm movements to visual targets in three-dimensional space. I. Relations between single cell discharge and direction of movement. *J Neurosci* 8:2913–2927.
- Scott SH, Kalaska JF (1997) Reaching movements with similar hand paths but different arm orientations. I. Activity of individual cells in motor cortex. *J Neurophysiol* 77:826–852.
- Sereno ME (1993) Neural computation of pattern motion: modeling stages of motion analysis in the primate visual cortex. Cambridge, MA: MIT.
- Sereno MI, Sereno ME (1991) Learning to see rotation and dilation with a Hebb rule. In: *Advances in neural information processing systems* (Lippmann RP, Moody J, Touretzky DS, eds), pp 320–326. San Mateo, CA: Morgan Kaufmann.
- Sergio LE, Kalaska JF (1997) Systematic changes in directional tuning of motor cortex cell activity with hand location in the workspace during generation of static isometric forces in constant spatial directions. *J Neurophysiol* 78:1170–1174.
- Shen L-M, Alexander GE (1997a) Neural correlates of a spatial sensory-to-motor transformation in primary motor cortex. *J Neurophysiol* 77:1171–1194.

- Shen L-M, Alexander GE (1997b) Preferential representation of instructed target location versus limb trajectory in dorsal premotor area. *J Neurophysiol* 77:1195–1212.
- Siegel RM, Read HL (1997) Analysis of optic flow in the monkey parietal area 7a. *Cereb Cortex* 7:327–346.
- Simoncelli EP, Heeger DJ (1998) A model of neuronal responses in visual area MT. *Vision Res* 38:743–761.
- Sinha P, Poggio T (1996) Role of learning in three-dimensional form perception. *Nature* 384:460–463.
- Soechting JF, Buneo CA, Herrmann U, Flanders M (1995) Moving effortlessly in three dimensions: does Donders's law apply to arm movement? *J Neurosci* 15:6271–6280.
- Squatrito S, Maioli MG (1997) Encoding of smooth pursuit direction and eye position by neurons of area MSTd of macaque monkey. *J Neurosci* 17:3847–3860.
- Stone JV (1998) Object recognition using spatiotemporal signatures. *Vision Res* 38:947–951.
- Stoner GR, Albright TD (1992) Neural correlates of perceptual motion coherence. *Nature* 358:412–414.
- Tanaka K, Saito H-A (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. *J Neurophysiol* 62:626–641.
- Tanaka K, Hikosaka K, Saito H-A, Yukiie M, Fukada Y, Iwai E (1986) Analysis of local and wide-field movements in the superior temporal visual areas of the macaque monkey. *J Neurosci* 6:134–144.
- Thier P, Erickson RG (1992) Responses of visual-tracking neurons from cortical area MST-l to visual, eye and head motion. *Eur J Neurosci* 4:539–553.
- Treue S, Maunsell JHR (1996) Attentional modulation of visual motion processing in cortical areas MT and MST. *Nature* 382:539–541.
- Trotter Y, Celebrini S, Stricanne B, Thorpe S, Imbert M (1992) Modulation of neural stereoscopic processing in primate area V1 by the viewing distance. *Science* 257:1279–1281.
- Turner RS, Anderson ME (1997) Pallidal discharge related to the kinematics of reaching movements in two dimensions. *J Neurophysiol* 77:1051–1074.
- Ullman S, Basri R (1991) Recognition by linear combinations of models. *IEEE Trans Pattern Analysis Machine Intelligence* 13:992–1005.
- Ungerleider LG, Mishkin M (1982) Two cortical visual systems. In: *Analysis of visual behavior* (Ingle DJ, Goodale MA, Mansfield RJW, eds), pp 549–586. Cambridge, MA: MIT.
- Wang G, Tanaka K, Tanifuji M (1996) Optical imaging of functional organization in the monkey inferotemporal cortex. *Science* 272:1665–1668.
- Wang R (1995) A simple competitive account of some response properties of visual neurons in area MSTd. *Neural Comput* 7:290–306.
- Watt A, Watt M (1992) *Advanced animation and rendering techniques: theory and practice*. New York: Addison-Wesley.
- Wise SP, di Pellegrino G, Boussaoud D (1992) Primate premotor cortex: dissociation of visumotor from sensory signals. *J Neurophysiol* 68:969–972.
- Zeki SM (1974) Functional organization of a visual area in the posterior bank of the superior temporal sulcus of the rhesus monkey. *J Physiol* 236:549–573.
- Zemel RS, Sejnowski TJ (1998) A model for encoding multiple object motions and self-motion in area MST of primate visual cortex. *J Neurosci* 18:531–547.
- Zhang K-C, Sereno MI, Sereno ME (1993) Emergence of position-independent detectors of sense of rotation and dilation with Hebbian learning: an analysis. *Neural Comput* 5:597–612.
- Zhang K-C, Ginzburg I, McNaughton BL, Sejnowski TJ (1998) Interpreting neuronal population activity by reconstruction: unified framework with application to hippocampal place cells. *J Neurophysiol* 79:1017–1044.
- Zipser D (1992) Identification models of the nervous system. *Neuroscience* 47:853–862.