Kinematic Coordinates In Which Motor Cortical Cells Encode Movement Direction

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AJEMIAN, ROBERT, DANIEL BULLOCK, AND STEPHEN GROSSBERG. Kinematic coordinates in which motor cortical cells encode movement direction. J Neurophysiol 84: 2191–2203, 2000. During goal-directed reaching in primates, a sensorimotor transformation generates a dynamical pattern of muscle activation. Within the context of this sensorimotor transformation, a fundamental question concerns the coordinate systems in which individual cells in the primary motor cortex (MI) encode movement direction. This article develops a mathematical framework that computes, as a function of the coordinate system in which an individual cell is hypothesized to operate, the spatial preferred direction (pd) of that cell as the arm configuration and hand location vary. Three coordinate systems are explicitly modeled: Cartesian spatial, shoulder-centered, and joint angle. The computed patterns of spatial pds are distinct for each of these three coordinate systems, and experimental approaches are described that can capitalize on these differences to compare the empirical adequacy of each coordinate hypothesis. One particular experiment involving curved motion was analyzed from this perspective. Out of the three coordinate systems tested, the assumption of joint angle coordinates best explained the observed cellular response properties. The mathematical framework developed in this paper can also be used to design new experiments that are capable of disambiguating between a given set of specified coordinate hypotheses.

INTRODUCTION

Activity in primary motor cortex (MI) has been implicated in a variety of aspects of movement behavior from control of movement execution to participation in movement planning. Specific examples of MI involvement in the control of kinematic or kinetic attributes of multi-joint movements include established correlations between cell firing rates and the following movement variables: movement direction (Georgopoulos et al. 1982; Schwartz et al. 1988), hand position (Georgopoulos and Massey 1985; Georgopoulos et al. 1984; Kettner et al. 1988), force (Georgopoulos et al. 1992; Kalaska et al. 1989), hand speed (Ashe and Georgopoulos 1994; Moran and Schwartz 1999a; Schwartz 1992), movement amplitude (Fu et al. 1993, 1995), and target direction (Alexander and Crutcher 1990b; Shen and Alexander 1997). Further studies have shown that cell firing rates correlate with aspects of movement planning such as movement preparation (Alexander and Crutcher 1990a; Kettner et al. 1996), target sequence information (Carpenter et al. 1999), and rapid motor adaptation (Wise et al. 1998). Cell activity therefore shows relations to a multitude of movement variables that span the sensorimotor spectrum.

Since not all MI cells are equally responsive to each of these variables, it makes sense to separately investigate distinct components of firing rate modulation. Although force or other movement variables could be analyzed with the methods employed herein, the present analysis focuses on cell response components related to a kinematic variable—movement direction—because studies have demonstrated the prevalence and strength of directional coding in MI (Ashe and Georgopoulos 1994) and because a large literature exists on center-out tasks in which movement direction is the explicitly controlled variable. Still, knowing that cell activity strongly reflects a kinematic movement variable like direction does not specify the nature of the cellular representation: Cartesian spatial coordinates, joint angle coordinates, or muscle length coordinates all might be used to represent movement direction at one neural stage or another.

For the entirety of MI, the supposition of a unique coordinate system in which movement direction is encoded may be inappropriate since a heterogeneity of coordinate systems may exist within a single brain region (Crutcher and Alexander 1990). Indeed it is well-documented that the representations that mediate motor behavior are distributed, often in a graded manner, across extensive, overlapping cortical regions (Fetz 1992; Kalaska and Crammond 1992; Mushiake et al. 1991). Therefore we restrict our analysis to the single-cell level and ask: how can one analyze the coordinate system in which an individual cell encodes movement direction? Beyond outlining a general framework for testing alternative coordinate hypotheses, we test three specific coordinate systems, Cartesian spatial, shoulder-centered, and joint angle, with regard to the data of Hocherman and Wise (1991).

Model and approach

PREFERRED DIRECTIONS IN AN INTERNAL SPACE. Georgopoulos et al. (1982) showed that the movement-related activity of many MI cells in the standard center-out task can be represented as

$$v(t) = b_0 + b_1 \cos (\omega - \omega_0)$$

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where $v$ is the cell’s average firing rate, $b_0$ is the mean movement-related activity across all directions, $b_1$ is the amplitude of the direction-dependent modulation of movement-related activity, $\omega$ is the movement direction of the hand, and $\omega_{pd}$ is the spatial preferred direction or spatial pd, i.e., the movement direction in space that elicits the maximal cellular response.

The empirical success of Eq. 1 warrants investigating, as one possibility, whether movement direction is represented in a spatial coordinate system. This hypothesis contrasts with some earlier studies where cell activity correlated strongly with muscle force (Cheney and Fetz 1980; Evarts 1968). More recently, Mussa-Ivaldi (1988) demonstrated theoretically that the observed spatial tuning can arise even if a motor cortical cell explicitly controls the time rate of change of multiple muscle lengths. From a diversity of empirical and theoretical studies, no consensus has emerged, and a variety of coordinate interpretations spanning the sensorimotor spectrum have been proposed for understanding directionally tuned cell activity in MI (Bullock and Grossberg 1988; Caminiti et al. 1990; Mussa-Ivaldi 1988; Sanger 1994; Schwartz 1992–1994; Scott and Kalaska 1997; Tanaka 1994; Zhang and Sejnowski 1999).

A key step to investigating alternative coordinate hypotheses is to distinguish between two types of representation of pds: a spatial pd and an internal pd.

SPATIAL PD. A spatial pd is that hand motion direction, as represented in extrapersonal space, to which a cell will respond maximally during small movements made from a common starting posture. What is meant here by the term “space” is the coordinate system utilized by the experimentalist in making measurements, typically a Cartesian coordinate system whose axes are aligned with the task space: e.g., the planar surface on which the monkey performs a center-out task. This coordinate system will henceforth be referred to as Cartesian spatial coordinates.

INTERNAL PD. An internal pd is that movement direction that elicits maximal cell response when represented in whatever coordinates best characterize the cellular-level encoding of movement direction. This “internal” coordinate system of a cell may be Cartesian spatial coordinates, or it could be some other coordinate system, such as a joint angle or muscle length coordinate system, which is more closely coupled to the biomechanical variables directly affected by the cell through its output connections. Thus, although the spatial pd reflects the internal pd, it is the internal pd that describes a cell’s distinctive role in the sensorimotor transformation.

For a well-defined internal coordinate system, mathematical transformations can be used to convert back and forth between a representation of direction in external space and its corresponding representation in the internal space. These transformations are in general posture dependent: the relationship between directions in the internal space and directions in external space changes as a function of posture. By using the distinctions between a spatial pd and an internal pd as well as the posture-dependent properties of the directional transformations between the spaces, a vector field method is developed that generates, for a given cell, spatial pd predictions that differ across the workspace as a function of coordinate hypothesis.

METHODS

Model arm

The analysis in this paper assumes a 2-joint or 2-degree-of-freedom (2-DOF) arm moving on a 2-dimensional (2-D) planar workspace situated within the horizontal plane passing through the shoulder. This model arm, illustrated in Fig. 1A, will be referred to as the 2-DOF planar arm. The kinematic equations describing this arm are detailed in the APPENDIX. A critical feature of the 2-DOF planar arm that simplifies our analysis is that positions map one-to-one to postures.

Modeling internal pds

One complication in adopting Eq. 1 as a general model for cell firing rates in center-out type tasks is that spatial pds have been observed to vary with hand position (Caminiti et al. 1990) and, more generally, arm posture (Scott and Kalaska 1997). To account for more of the variance in cell discharge as the center-out task base expands, additional predictor variables (such as hand position) might be added to the regression equation (Ashe and Georgopoulos 1994). Alternatively, a change in the coordinate representation of the variable of interest (Lacquaniti et al. 1995), which in this case is the preferred movement direction, might allow an equation as compact as Eq. 1 to account for a larger proportion of the variance. As part of the search for a more generally applicable tuning equation, the 2-DOF planar arm model can be used to construct alternative coordinate systems for the purpose of testing whether Eq. 1 (in which a cell’s pd is specified once and without regard to the arm’s posture) can provide a better data fit if the spatial pd is interpreted as a specific instantiation of an underlying and invariant internal pd. A constant internal pd, together with the relevant coordinate transformation, can in principle fully explain the observation of a posture-dependent spatial pd by generating a systematic prediction of the manner in which the spatial pd changes with posture.

To illustrate, suppose that the spatial pd of a cell at some reference posture is direction $\hat{A}$ and that the internal space of a cell is coordinate system $Z$. Movement direction $\hat{A}$ in space maps to movement direction $\hat{B}$ in coordinate system $Z$. Now suppose that movements are initiated from a new arm posture. How can we predict the new spatial pd, $\hat{A}'$? Assuming that $\hat{B}$ remains the cell’s pd in internal coordinate system $Z$ at the new arm posture, $\hat{A}'$ can be calculated using the reverse mapping: between directions in coordinate system $Z$ to directions in external space. In general, for all cases where the internal coordinate system is not identical to external space, $\hat{A}'$ will not be the same as $\hat{A}$, because the transformation between directions in coordinate system $Z$ and directions in external space depends on posture (i.e., as the posture changes, so does the local relationship between movement directions in the 2 coordinate systems). This type of coordinate analysis belongs to the branch of mathematics known as differential geometry.

Vector fields of spatial pds

Given a 2-DOF planar arm, hand position maps uniquely to arm posture (which is not the case when the arm possesses redundant degrees of freedom). Thus determining the spatial pd at every posture is equivalent to uniquely determining the spatial pd at every hand position in the workspace. Specifying a spatial direction and a corresponding magnitude over a field of points in space defines a vector field [in this case, a vector field of spatial pds as in Zhang and Sejnowski (1999)]. Thus an internal pd in a particular coordinate system implies a vector field of spatial pds. To illustrate, plots of vector fields of spatial pds were constructed under the assumption of each of three internal coordinate systems for a sample cell whose spatial pd is $60^\circ$ at a reference posture, as indicated in Fig. 1A.
The simplest vector field arises when the internal coordinate system in which a cell encodes movement direction is the same Cartesian spatial coordinate system in which spatial pds are measured. Psychophysical evidence (Morasso 1981) suggests that movement planning may occur in this coordinate system. Spatial pds for this case will not vary with posture because the spatial pd at the reference posture is also the cell’s internal pd; in other words, the identity transformation converts between the two representations of direction. Figure 1C shows this constant-direction vector field of spatial pds. The magnitude of each vector is unity; the APPENDIX describes how magnitudes are determined. For this and subsequent vector field plots, information regarding the direction but not magnitude of the vectors is provided.

A vector at a given point in these vector field plots represents the cell’s expected spatial pd if the center-out task were performed with that point as the movement origin. Because it is impractical to map out a cell’s vector field of spatial pds by performing the center-out task as many times as there are arrows on the simulation plots, alternative testing methods are described later.

One axis important for many mammals is the line between the proximal and distal end of a limb; e.g., between the shoulder and the hand (Maioli and Lacquaniti 1988). Psychophysical studies (Flanders et al. 1992; Soechting and Flanders 1989) have suggested the existence of a shoulder-referenced spatial coordinate system, and cell data have been interpreted in terms of a shoulder-referenced intrinsic coordinate system (Caminiti et al. 1990, 1991; Tanaka 1994). In consideration of these observations, suppose as shown in Fig. 1B that a cell’s spatial pd is computed in a mobile Cartesian spatial reference frame, one axis of which is aligned with the axis connecting the shoulder to the hand. As the shoulder-hand axis rotates (due to rotations at the shoulder and/or elbow joints), the cell’s spatial pd rotates by an equivalent amount. Thus the rotational transformation converts between representations of direction in the two spaces. Figure 1D plots the variable-direction vector field of spatial pds generated for the sample cell with a constant pd in shoulder-centered coordinates. Specifications for generating this vector field are contained in the APPENDIX.

An MI cell may encode movement in a joint angle coordinate system that represents a later stage in the sensorimotor transformation from spatial coordinates to muscle activations. Psychophysical studies on motor adaptation (Gandolfo et al. 1996; Shadmehr and Mussa-Ivaldi 1994) have implicated joint-based representations. Mussa-Ivaldi (1988) suggested that MI cell activity could be a linear function of the rate of multiple muscle length changes. More recently, Scott and Kalaska (1997) introduced a joint angle interpretation of MI cell activity, and our interpretation is similar to theirs.

Suppose that at the reference posture, \((\theta_0, \phi_0)\), a cell possesses a spatial pd, \(\mathbf{p}_{\text{sp}}\). Using the inverse of the Jacobian of the kinematic transformation from joint angle coordinates to spatial coordinates, this spatial direction can be converted to a direction in joint angle space. On assumption of a new arm posture, the Jacobian can be used to

**Cartesian spatial coordinates**

**Shoulder-centered coordinates**

**Joint angle coordinates**

convert the joint angle direction back to a spatial pd. Since the Jacobian is posture dependent, application of the inverse Jacobian followed by application of the forward Jacobian evaluated at a new posture is not equivalent to operating with the identity transformation; the composite transformation will result in a new spatial pd. The mathematical details of constructing this vector field are contained in the Appendix.

An intuitive explanation of what it means for a cell to possess an internal pd in a joint angle coordinate system is as follows. Suppose the internal pd for a cell is

\[
\begin{bmatrix}
\theta_{pd} \\
\phi_{pd}
\end{bmatrix} = \begin{bmatrix} 1 \\ 3 \end{bmatrix}
\]

where \(\theta_{pd}\) and \(\phi_{pd}\) correspond to the relative shoulder and elbow components of the preferred velocity vector in joint angle space. Such a cell responds maximally to directions of coordinated two-joint motions produced when the elbow rotation rate is three times the shoulder rotation rate. Depending on the posture, the spatial movement direction that corresponds to this movement direction in joint angle space (i.e., this joint synergy) will vary. Figure 1E depicts the vector field of spatial pds generated for the sample cell with a constant pd in joint angle coordinates.

**Global description of vector fields**

These three vector fields simulated for a sample cell clearly differ from one another. Is there any simple way to classify the differences in their structure without comparing vectors in the alternative vector fields one by one for each cell? The *curl* of a vector field is a local measure of the rotational tendency of vector field flow; that is, a measure at a point of how much the vectors rotate in the neighborhood of that point. Observing how the curl changes across the workspace helps to explain the global structure of a vector field. Below we present the distinct curls for each of the three classes of vector fields described above. The mathematical details of the derivations are reported in the Appendix.

**Cartesian spatial coordinates.** Cartesian spatial internal pds imply that the spatial pds do not change. Hence, there is no oriented flow to the vector fields, and their curls are everywhere zero.

**Shoulder-centered coordinates.** Vector fields generated under the assumption of this coordinate system yield

\[
\text{curl}(x, y) = -\cos \theta_{pd}(x_R, y_R) \frac{1}{r}
\]

where \(\theta_{pd}(x_R, y_R)\) is the spatial pd of the cell at the reference posture, and \(r\) is the distance of the hand from the shoulder. The inverse dependence on \(r\) indicates that the rotational tendency of vectors diminishes at more distal portions of the workspace.

**Joint angle coordinates.** For a cell tuned to an invariant direction in joint angle space, this internal pd can be written as a normalized joint angle velocity vector (* denotes normalization)

\[
\begin{bmatrix}
\theta_{pd}^* \\
\phi_{pd}^*
\end{bmatrix}
\]

where \(\theta_{pd}^*\) denotes the shoulder component of the preferred joint synergy and \(\phi_{pd}^*\) denotes the elbow component. The curl value for the vector field of such a cell is

\[
\text{curl}(x, y) = 2\theta_{pd}^* + \phi_{pd}^*
\]

This curl is a nonzero constant (no dependence on hand position or arm posture). Thus vectors in this vector field rotate (in sharp contrast to Cartesian spatial coordinates), and their rotational tendency is uniform throughout the workspace (in sharp contrast to shoulder-centered coordinates). The constant value depends only on the joint synergy to which the cell is tuned.

**Utility of vector fields**

Measuring the curl experimentally is problematic since it is a local measure whose accurate estimation at multiple points would require a high resolution sampling of the workspace that may be difficult to accomplish in practice. Nonetheless, for any pair of candidate coordinate systems, computation of the curl indicates whether the two coordinate systems give rise to vector fields of similar or disparate structure, and thus whether they are empirically distinguishable. Based on this fact, two distinct methods for experimentally disambiguating between distinguishable vector field structures are now described: “direct field sampling” and “indirect field sampling.”

**Direct field sampling**

This method determines spatial pds at several different workspace locations and then, using a least mean square analysis, compares the results with those predicted by the different coordinate hypotheses. For example, spatial pd predictions at the locations indicated by the thin-lined boxes in Fig. 1, C–E, can be compared with the measured spatial pds at those locations to determine which coordinate system provides the best fit. Knowledge of the vector field structures can optimize the discriminatory efficacy of the direct field sampling paradigm by enabling workspace sampling that focuses on those locations that engender very different predictions for the coordinate systems being evaluated. For 2-D planar arm movements, no experiment has been performed that directly sampled the workspace in the manner suggested above, although Caminiti et al. (1990) and Scott and Kalaska (1997) have performed experiments based on this concept (see discussion).

**Indirect field sampling**

Another method relies on investigating cortical activity during long, curved movements that sweep broadly across the workspace, thereby visiting many postures and implicitly sampling a cell’s vector field of spatial pds over a single trajectory. The pattern of movement-related activity registered by a cell along multiple such paths determines the cell’s trajectory-selectivity or its tendency to respond preferentially to certain types of trajectories. A cell’s trajectory-selectivity, if any, can serve as the signature for a specific coordinate system.

\[\text{Eq. 1} \]

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direction-dependent component of cell activity over the course of a movement path.

To complete the determination of the temporal response profile, we note that MI cell response for trained movements with unimodal speed profiles often takes the form of a phasic pulse or burstlike response. Many generative hypotheses are consistent with this shape. As the focus of this article is not on explicating the specific shape of the response, but on understanding how variations in cell response arise as a function of the directional characteristics of the movement path taken by the hand, we simply assume a generic burstlike shape for cell response in our simulations. Therefore to determine the temporal response profile, the directional component of cell activity (as determined in steps 1–3 above) is modulated by a generic Gaussian that embodies the phasic response properties of many MI cells. The Gaussian modulation is a fixed component of cell response used identically for all paths and coordinate assumptions, and it introduces no bias. Simulations showed that the precise form of the response envelope (which included different pulse shapes as well as the constant function) does not alter the results on trajectory-selectivity. A determination of trajectory-selectivity indicates that a cell responds preferentially to certain movement paths, and this path-dependent response depends on the variable directional component of cell response and not on the fixed modulatory component.

Averaging the activity over all the bins of a movement path determines the mean firing frequency over the course of the entire movement. Thus the average firing rate, \( \bar{v} \), of a cell over the course of an arbitrary trajectory can be expressed as

\[
\bar{v} = \left( \frac{1}{T} \right) \sum_{i=1}^{n} G_i(b_o + b_1 \cos(\theta(i) - \theta_o(i))) \Delta t,
\]

where \( i \) denotes the bin number, \( \Delta t \) denotes the duration of bin \( i \), \( G \) denotes the modulation of the burstlike activity by a Gaussian, and \( T = \sum_{i=1}^{n} \Delta t \) denotes the total movement time. Note that since the movement direction, \( \theta \), and spatial \( pd \), \( \theta_o \), are written as functions of the bin number (i.e., the position along the movement path), both are interpreted as varying as a function of hand position or arm posture. Therefore not only does the movement direction in general change as the hand traverses a curved path, but so too may the cell’s spatial \( pd \).

### Hocherman and Wise (1991)

The data of Hocherman and Wise (1991) are now analyzed within the framework of indirect field sampling for the purpose of evaluating the adequacy of the three internal coordinate hypotheses. That study investigated the correlation between individual motor cortical cell activity and the curvature type of end-effector motion. Briefly, a monkey was trained (by use of intermediate via points between the movement origin and target locations) to make movements of different curvature types from an origin point to each of three equidistant targets spaced at intervals of 30°. Both the arm and the targets were constrained to lie on a 2-D planar surface. The three movement types consisted of clockwise arcs, straight lines, and counterclockwise arcs; a movement of each curvature type was made to each of the three targets for a total of nine distinct trajectories, which are numerically labeled in Fig. 2. Unconstrained return movements were also part of the protocol, so even though the targets were concentrated in a 60° wedge, movement directions did span the entire 360° of the angular continuum.

Cell activities were recorded in the arm area of MI both before and during the movements. (In the actual experiment, cell recordings occurred in several different epochs, but we only simulate activity for a single movement-related epoch, which corresponds closely to their “late movement epoch.”) In the study, a neuronal modulation index, \( M_j \), was used as a normalized measure of a cell’s average movement-related activity for path \( j \) and was computed with the equation

\[
M_j = \frac{A_j - R}{A_{\text{MAX}} - R}
\]

where \( A_j \) is the cell’s average activity over movement path \( j \), \( R \) is the cell’s resting discharge rate, and \( A_{\text{MAX}} \) is the cell’s average discharge rate over that movement path (of the 9) for which the cell is maximally active. An \( M \) value close to 1 means a cell is highly active for that path, while an \( M \) value close to 0 means the cell is largely inactive. Cells were classified as trajectory selective for a certain curvature type if they were preferentially active for movements of that curvature type (see the APPENDIX). For example, a cell that was clockwise trajectory selective exhibited higher levels of activity for the clockwise trajectories (labeled in Fig. 2 as 1, 4, and 7) than for straight or counterclockwise movement counterparts. Similar definitions held for classifying cells as straight trajectory selective or counterclockwise trajectory selective.

Using the method of indirect field sampling, we simulated the experiment of Hocherman and Wise (1991) by 1) computing each model cell’s modulation index for all of the nine movement paths using their normalization procedures, 2) classifying model cells using their classification criteria, and 3) generating cellular temporal response profiles. A model cell was identified by its spatial \( pd \) at the reference posture; the population of model cells consisted of 360 cells, 1 for each degree of the angular continuum. Simulation details are found in the APPENDIX.

### RESULTS

#### Simulations of trajectory-selectivity

A key discovery of Hocherman and Wise (1991) was a strong tendency for cells to respond preferentially to movements of the curved trajectory types. Illustrations of the results of the original experiment are given in Fig. 3, \( A \) and \( C \), which show the percentages of trajectory-selective cells for each trajectory type using the strict (A) and relaxed (C) criteria to classify cells.

For the simulations run under the assumptions of Cartesian spatial, shoulder-centered, and joint angle coordinates, there were, respectively, 181, 156, and 135 task-related model cells of a total of 360 model cells. The spatial \( pd \) of these cells at the reference posture were almost entirely contained in the 0–180° range since the movement directions required to reach the targets also exist in that range. Plots in Fig. 3, \( B \) and \( D \), depict the percentages of cells that were trajectory selective for each trajectory type using each classification criterion. Under
the assumptions of both Cartesian spatial coordinates and
shoulder-centered coordinates, the vast majority of the trajec-
tory-selective model cells, 100 and 68%, respectively, were
trajectory selective for the straight trajectory type when the
strict classification criterion was used; using the relaxed crite-
ror, the percentages were 98 and 69%. These simulation
results are not consistent with the data where the vast majority
of trajectory-selective cells are of the two curved movement
types. Under the assumption of joint angle coordinates, how-
ever, the majority of model cells were (like MI cells) trajectory
selective for the curved trajectories. Furthermore, the percent-
ages of all three types of trajectory-selective cells using the
joint angle model correspond well with the data for both
classification schemes, as can be seen by comparing the graphs.

To understand the simulation results, recall Eq. 5. It implies
that, over the course of a trajectory, a cell registers significant
activity while the movement direction is parallel to the spatial
pd; the greater the deviation from colinearity, the less the
activity generated. The average firing rate of a cell for an entire
movement, then, depends on the interaction between the vector
field structure of spatial pds and the sequence of movement
directions taken by the hand. Previously, it was shown that the
hypothesis of a particular coordinate system imparts a signa-
ture structure to the vector field of spatial pds. Similarly, each
type of movement curvature (clockwise, straight, counterclock-
wise) engenders its own characteristic pattern of movement
directions. The movement direction for clockwise movements
rotates continuously and in a clockwise manner from the be-
inning of the movement to its end for a net rotation of about
90°. The reverse is true for the counterclockwise movements.
During straight movements, the movement direction never
changes. The observed ratios of trajectory-selectivity for a
given coordinate system can be understood by considering,
within the context of the task, how these characteristic move-
ment patterns interact with each vector field structure.

For example, the spatial pds of vector fields generated by the
assumption of joint angle coordinates tend to rotate in a uni-
form direction over the entire course of each trajectory. Figure
4 shows plots of a model cell’s spatial pd values over the
course of a clockwise trajectory and a straight trajectory to a
particular target under the assumption of each coordinate sys-
tem. It can be seen in the plot of the joint angle coordinate
simulation that the spatial pd is not initially aligned with the
movement direction at the beginning of the clockwise move-
ment, but the two gradually fall into alignment over the course
of the trajectory. The reverse is true for the straight trajectory.

A rotating movement direction can engender considerable
activity when paired with a rotating spatial pd if they rotate in
the same direction and if the movement direction rotates more
sharply, over the same spatial extent, than the spatial pd (a
situation that does arise in the case of joint angle coordinates
for the curved movements in this experiment). The dual rota-
tion facilitates the occurrence of an interval of overlap during
which the two directions are nearly aligned. At some point, the
movement direction “overtakes” the spatial pd, although these
directions may not be initially aligned, and this tendency
toward alignment occurs for multiple movements of the same
curvature type even when the final targets of these movements
are different. Thus the assumption of joint angle coordinates
gives rise to relatively large proportions of cells that are tra-
jectory selective for the curved trajectory types. In contrast,
under the assumption of either Cartesian spatial or shoulder-centered coordinates, there is either no tendency or a much weaker tendency for the spatial pds to rotate over the course of the trajectories, and what rotation does occur is often not unidirectional over an entire trajectory. This produces model cells that respond preferentially to straight trajectories.

Simulations of cell response profiles

In addition to simulating the average activity over the course of an entire trajectory, the model can also simulate, as shown in Fig. 5, the temporal response profiles of a cell for each of the nine different movement trajectories under the assumption of joint angle coordinates. Figure 5 shows that cell response properties vary as a function of movement curvature. For example, the relative timing of peak activity depends critically on the time-evolving relationship between the hand’s movement direction and the cell’s spatial pd for the movement path under consideration. The peak activity for the model cell occurs 150–200 ms after the onset of movement-related activity for the counterclockwise movement paths and 275–325 ms after onset for the clockwise movement paths. This predicted time lag between the peak activities can be tested experimentally. Such temporal differences in activity profiles exist for all the response envelopes we tried since these differences stem from the variable directional component of cellular response, which is highly differentiated in this paradigm as a function of curvature type. For other model cells (depending on the spatial pd at the reference posture), the relative timing of peak activity as a function of movement curvature will be reversed: the peak activity will occur sooner for the clockwise paths than for the counterclockwise paths.

This cell is typical of all model cells in two important respects: 1) its response characteristics, such as its peak firing rate, mean firing rate, and the timing of its peak firing rate, change relatively gradually from one trajectory type to the next; and 2) the mean activity levels across trajectory types are ordered in a characteristic manner, i.e., a clockwise trajectory-selective cell will be most active for the clockwise paths, least active for the counterclockwise paths, and intermittently active for the straight movement paths (the inequality is reversed for counterclockwise trajectory-selective cells). In what follows, we analyze Hocherman and Wise (1991) data with respect to the above two model cell response properties.

Comparison of model cell response properties with data

Do real MI cells exhibit graded responses such as those illustrated in Fig. 5? Instead, an MI cell might be highly modulated for clockwise trajectories but relatively silent for straight and counterclockwise movements. If curvature were explicitly encoded as a movement primitive by MI cells, then one might expect such a discretization of response characteristics. Some of the plotted response profiles in Hocherman and Wise (1991) seem to support the all-or-none view, although this type of analysis was not performed in that study. To assess
whether MI cell activity more closely conforms to the graded or categorical response characteristics, we obtained the original data files for 59 of the 76 task-related MI neurons (which included 19 of the 24 trajectory-selective cells using the strict criterion) from Hocherman and Wise (1991) and analyzed the spread in activity for movements of different curvature types. Specifically, for each trajectory-selective cell, let $\bar{A}_{cw}$, $\bar{A}_{ccw}$, and $\bar{A}_{str}$ denote cell activity averaged over each set, respectively, of clockwise movements, counterclockwise movements, and straight movements. For example, $\bar{A}_{cw}$ denotes cell activity averaged over the clockwise movement paths 1, 4, and 7 in Fig. 2. Consequently, a separation index, analogous to the modulation index, was defined for each trajectory-selective cell as

$$\frac{A_{\text{max}} - A_{\text{min}}}{A_{\text{MAX}} - R}$$

where $A_{\text{max}}$ is the largest of $\bar{A}_{cw}$, $\bar{A}_{ccw}$, and $\bar{A}_{str}$; $A_{\text{min}}$ is the least of these three averages; $A_{\text{MAX}}$ is the cell’s average discharge rate over that movement path (of the 9) for which the cell is maximally active; and $R$ is the cell’s resting discharge rate. The numerator represents the absolute spread in activity as a function of curvature type, while the denominator represents the maximum amount of movement-related activity exhibited by the cell. The ratio can range from 0 to 1 with a fraction close to 1, suggesting that the curvature-dependent activity possesses close to an all-or-none character, while a fraction close to 0 suggests that activity varies rather gradually as a function of movement curvature.

Figure 6A plots the distribution of separation indices for the population of trajectory-selective cells in Hocherman and Wise (1991). The mean and median separation indices are 0.48 and 0.43, suggesting that (outside of the small percentage of outliers present in the plot) cell response varies relatively gradually as a function of movement curvature. Figure 6B plots the corresponding distribution of simulated separation indices for the population of model trajectory-selective cells under the assumption of joint angle coordinates. The mean and median separation indices are 0.35 and 0.34. Note that for both distributions the vast majority of separation indices lie in the interval between 0.3 and 0.5. Therefore the gradual variation exemplified by the Fig. 5 model cell is a characteristic feature of both the model and the data.

A second distinctive feature of model cell response properties is the very specific ordering of mean activity as a function of curvature type. In particular, for every clockwise trajectory-selective model cell, the following condition holds: $\bar{A}_{cw} > \bar{A}_{str} > \bar{A}_{ccw}$. This condition (with the inequality accordingly reversed) also holds for every counterclockwise trajectory-selective model cell. For the population of curved trajectory-selective cells in Hocherman and Wise (1991), 89% of the cells (17/19) showed the same ordering in their activity. Thus the model reproduces not only the observed graded responses, but also the observed ordering of those responses.

**Varying simulation parameters**

There are no free parameters in the model, since the only model variable is a cell’s vector field of spatial pds, which is completely determined as a function of the working coordinate hypothesis. However, the simulations did require values for the location of the movement origin, the speed profiles of the hand, the lengths of the arm segments, and $b_0$ and $b_1$ of a cell’s tuning curve. Regarding the kinematic movement parameters, Hocherman and Wise (1991) did not have precise measurements for these quantities. Therefore while the values used in the simulations were in accord with the specifications communicated to us by Dr. Wise, we systematically varied these parameters to probe the robustness of the results regarding trajectory-selectivity. The **appendix** provides the details of these sensitivity analyses, the findings of which demonstrate the robustness of the simulation results for all three coordinate systems. Varying the cellular parameters $b_0$ and $b_1$ [which must be assumed since no center-out task is performed to determine them in Hocherman and Wise (1991)] did not alter a cell’s trajectory-selectivity as shown in the **appendix**. The use of response envelopes with different pulse functions or with the constant function made no significant difference in the simulation results. Finally, although the simulations employed a uniform distribution of spatial pds at the reference posture (as revealed in Lurito et al. 1991), distributional skewing away from nonuniformity, such as that reported in the literature (Georgopoulos et al. 1982; Scott and Kalaska 1997) did not change the character of the results under the assumption of any of the three internal coordinate systems.
Amplitude effects

In the experiment of Hocherman and Wise (1991), the curved paths were longer than the straight paths (23 cm as opposed to 20 cm), so the prevalence of curved trajectory-selective cells could conceivably result from an amplitude dependence of the cell firing function (Fu et al. 1993, 1995). However, this hypothesis conflicts with the observed ordering of cell activity by trajectory type, which indicates that, for the actual clockwise trajectory cells, $A_{cw} > A_{str} > A_{xw}$ (the order of inequalities is reversed for counterclockwise trajectory selectivity). Even if modulation indices are scaled outright by path length, the simulation results for Cartesian spatial and shoulder-centered coordinates grossly contradict this observed ordering.

Effect of other movement variables

We conducted analyses (see Appendix) to assess whether a simple dependence of cell firing rates on either hand speed or hand position could alter the relative goodness-of-fit of the three coordinate hypotheses. The inclusion of these correlations did not change the nature of the results. The joint angle coordinate hypothesis continued to fit the data well while the other two coordinate systems failed. Although we believe that amplitude and hand speed were the most pertinent task variables (aside from direction) to consider in explaining the data, these variables comprise only a subset of the known correlates of MI cell activity (see Introduction). Additional studies would be needed to assess whether correlations with other movement-related variables could provide an alternative or supplementary explanation of the Hocherman and Wise (1991) results.

Prediction: internal pd controls spatial pd and trajectory-selectivity

The simulations of the Hocherman and Wise (1991) experiment not only determine the percentages of cells selective for the different trajectory types but also imply a relationship between a cell’s internal pd and its trajectory selectivity. Specifically, a cell’s spatial pd at a reference posture maps to a cell’s internal pd; from the cell’s internal pd, a vector field of spatial pds is generated; from the cell’s vector field of spatial pds, the cell’s trajectory selectivity is determined. Thus a mapping is constructed from the spatial pd of a cell at a reference posture to the type of trajectory selectivity which that cell is predicted to possess. For example, a model cell with a spatial pd of 45° at the reference posture is clockwise trajectory selective under the assumption of joint angle coordinates. Table 1 depicts the complete predicted mapping from a cell’s spatial pd at the reference posture to its trajectory selectivity using the example of joint angle coordinates. This prediction can be tested in an experiment that determines both spatial pds through the center-out task and cellular trajectory selectivity through the curved motion task of Hocherman and Wise (1991). The end result of this composite protocol would be an empirical determination of the mapping between spatial pds at a reference posture and type of trajectory selectivity. Model mappings constructed for each internal coordinate system could then be compared with the actual mapping to assess the goodness-of-fit of alternative coordinate hypotheses.

<table>
<thead>
<tr>
<th>Spatial pd of a Model Cell at the Reference Posture, deg</th>
<th>Predicted Trajectory-Selectivity</th>
</tr>
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<tbody>
<tr>
<td>27–91</td>
<td>Clockwise</td>
</tr>
<tr>
<td>92–95</td>
<td>Indeterminate</td>
</tr>
<tr>
<td>96–105</td>
<td>Straight</td>
</tr>
<tr>
<td>106</td>
<td>Indeterminate</td>
</tr>
<tr>
<td>107–161</td>
<td>Counterclockwise</td>
</tr>
</tbody>
</table>

Predicted results for a composite protocol that conjoins the standard center-out task (Georgopoulos et al. 1982) and the curved motion task of Hocherman and Wise (1991) under the assumption of joint angle coordinates. The center-out task will determine a cell’s spatial preferred direction (pd) at a reference posture. The curved motion task will result in a cell’s being classified as either 1) trajectory selective for the clockwise, straight, or counterclockwise trajectory type (using the strict criterion of cell classification), or 2) indeterminate trajectory selective, which means that the cell is modulated by the task but cannot be classified as responding preferentially to 1 of the 3 movement types using the strict criterion (for example, the cell may respond preferentially to the clockwise movement for target A but responds preferentially to the straight movement for target B). Those cells absent from the list are not found to be task related. The table maps the dual experimental outcomes to each other on a cell-by-cell basis implicitly utilizing a cell’s assumed internal pd as the common underlying factor (and sole cellular response characteristic) in generating cell behavior for each paradigm. As this experiment has not been performed, these simulation results serve as an untested prediction, the confirmation of which would provide support for the contention that observed results on cellular trajectory-selectivity (Hocherman and Wise 1991) are derived from joint angle directional control and not from the explicit encoding of curvature as a movement primitive or from the encoding of other movement variables.

Target selectivity

In addition to classifying cells as trajectory selective, Hocherman and Wise (1991) classified cells as target selective if the cells responded preferentially to movements to a specific target as compared with the responses to movements to the other targets. For the purpose of representing the findings of Hocherman and Wise (1991) on target selectivity and of showing corresponding simulation results (using the strict criterion of classification), let $x/y/z$ indicate that $x\%$ of task-related cells are target selective for target 1, $y\%$ of cells are target selective for target 2, and $z\%$ are target selective for target 3. On the basis of Table 3 and Fig. 12b in Hocherman and Wise (1991), the percentages of excitable target-selective cells found in that study were 42/29/29. For the Cartesian spatial simulations, the percentages were 33/31/36; for the shoulder-centered simulations, the percentages were 38/16/46; for the joint angle simulations, the percentages were 46/16/38. Thus all three coordinate hypotheses roughly reproduce the results on target selectivity, and these target selectivity data cannot distinguish between the coordinate systems.

Compatibility of joint angle coordinates with prior population vector analyses

The population vector algorithm (PVA) has been used to predict movement direction over the course of a trajectory on a bin-by-bin basis with good results (for review, see Georgopoulos 1995). In standard use of the algorithm, the assumed spatial pds do not change as the hand location changes from one bin to the next. If spatial pds in actuality do vary across the workspace, the population vector should rotate away from the movement direction as the movement progresses away from
the point at which the cell’s pd was assessed. Such a mismatch between the population vector direction and the movement direction would arise because the algorithm’s invariant representation of a cell’s vectorial contribution comes to lie in a direction slightly askew from the cell’s actual preferred direction. Nonetheless, the very robustness of the PVA, as an aggregate estimator of movement direction, renders it insensitive to alternative coordinate assumptions (Georgopoulos 1996; Mussa-Ivaldi 1988; Sanger 1994). To assess sensitivity in the current case, we performed a bin-by-bin population vector simulation for all eight movements in the standard center-out task. In this PVA, computed cell activity was based on the bin(posture)-dependent spatial pds determined by the joint angle coordinate model. The trajectories were divided into 25 bins, each of length 20 ms. Under these conditions, the population vector did rotate away from the actual movement direction (due to the structure of the joint angle coordinate system), but the rotation was modest and the resultant prediction error was within the range of prior reports that used the PVA. The average amount of rotation from the beginning of a trajectory to the end of the trajectory was <10°. Further, the mean absolute differences between the population vector direction and the movement vector direction over all of the bins was <5°. That the mean signed difference between the two vector directions over all of the bins was 0° indicates the difficulty of using a population vector analysis to distinguish between coordinate systems. Thus the joint angle coordinate hypothesis is consistent with prior PVA results.

DISCUSSION

This paper presents a framework for analyzing the coordinate system in which an individual cell encodes movement direction. A cell’s preferred direction can be predicted to vary across the workspace in a distinct manner depending on the assumed internal coordinate system, and direct sampling experiments can be designed to probe these variations. Indirect sampling experiments examining cell activity over long, curved movements implicitly sample vector fields of spatial pds and can be used to choose between alternative coordinate hypotheses from the pattern of path-dependent activity. We simulated one such experiment (Hocherman and Wise 1991) under the assumption of three kinematic coordinate systems (Cartesian spatial, shoulder-centered, and joint angle) and found that joint angle coordinates robustly fit the MI data better than either of the other two coordinate systems.

These results do not imply that all MI cells encode movement direction in joint angle coordinates. First, only three coordinate systems were tested, and there may exist another coordinate system that fits the data better than joint angle coordinates. Second, even if a majority of cells within a given brain region represent movement direction in one particular coordinate system, evidence (Crutcher and Alexander 1990) suggests that there will often exist other cells in the same brain region that utilize different coordinate representations. Third, a recent investigation (Wise et al. 1998) demonstrates the capacity of motor cortex to rapidly reorganize its response properties during adaptation to a series of differentiated visuomotor tasks, perhaps implying that the CNS solves motor control problems by implementing task-specific solutions that utilize task-dependent coordinate decompositions of the sensorimotor transformation. Finally, the current analysis focuses on the representation of movement direction but, as reiterated above, cell activity likely reflects information about other movement variables as well. A more detailed exploration of the functional dependence of cell activity on multiple movement variables is warranted for clarifying these and other data.

By looking at the coordinate system in which an individual cell encodes movement direction, it becomes possible to assess how populations of cells with similar coordinate representations are distributed across a cortical area. Hocherman and Wise (1991) recorded in the supplementary motor area (SMA), dorsal premotor cortex (PMd), and ventral premotor cortex (PMv) as well as MI. In these other cortical regions, a smaller percentage of cells responded preferentially to curved movements. From the present analysis we infer that MI represents movement commands in a coordinate system possessing a stronger joint angle character than do the SMA, PMd, or PMv. This conclusion is consistent with the findings of Scott et al. (1997).

Our study uses curved movements as a means to probe the structure of a cell’s vector field of spatial pds by indirectly sampling the workspace. Another way to investigate vector field structure is by directly sampling the workspace, and two prior studies involving proximal arm movements fall into that category: Caminiti et al. (1990) and Scott and Kalaska (1997). In Caminiti et al. (1990), a 3-D center-out task was performed from three distinct movement origins that were colinear (normal to the sagittal plane), spaced 10 cm apart, and situated in a transverse plane cutting through the shoulders. Spatial pds were found to change across the workspace by a statistically significant amount. These changes were fit reasonably well by assuming that the change in a cell’s spatial pd matched the rotation of the shoulder joint from one workspace location to the next. Since the rotation of the shoulder joint from one workspace location to the next, proceeding from left to right, is virtually equivalent in this task to the rotation of the shoulder-hand axis (18 and 20° for the former as opposed to 21.8 and 21.8° for the latter), the shoulder-centered coordinates defined in this paper would fit the data about as well. Lacking information regarding the movement trajectories in joint angle space (which is here necessary since an unconstrained arm operating in 3-D space is motor redundant), we were unable to simulate this paradigm under the assumption of joint angle coordinates.

In Scott and Kalaska (1997), a monkey performed the center-out task in two different postures (natural and abducted) that corresponded to the same end-effector location in space. They noted a significant posture by direction interaction effect present in the response properties of a majority of cells and demonstrated statistically that changes in a cell’s directional preference were a major contributing factor. The difference between the mean spatial pds across the two arm orientations was significant for 48% of the 422 cells examined. Scott and Kalaska (1997) modeled these data using Cartesian spatial, joint angle, and joint torque coordinate systems. They found that joint angle coordinates best fit the data. The results were incompatible with the assumption of either Cartesian spatial or shoulder-centered coordinates.

On the basis of our analyses as well as the analyses in Caminiti et al. (1990) and Scott and Kalaska (1997), Table 2 provides an evaluation of the adequacy of the three different
coordinate systems modeled in this paper with regard to three experiments, each of which investigated proximal arm cell activity during unloaded reaching movements: Caminiti et al. (1990), Hocherman and Wise (1991), and Scott and Kalaska (1997).

The observations of Caminiti et al. (1990) and Scott and Kalaska (1997) may appear to contradict the findings of Schwartz (1992), which investigated the temporal discharge patterns of individual cells in motor cortex during the tracing of sinusoids. It was found that cell discharge patterns correspond closely (high correlation coefficient) to what would be predicted under the assumption of a fixed spatial pd (i.e., Cartesian spatial coordinates), once the time lag between the cortical signal and its controlling effect at the periphery is taken into account. One could conceivably interpret these findings as support for Cartesian spatial coordinates, although Schwartz (1992) does not address the issue of coordinate systems and makes no claims in this regard. The framework for coordinate analysis established in this paper suggests that the results of Schwartz (1992) do not support or refute any coordinate system hypothesis. Differentiating between coordinate systems requires 1) probing the organization of spatial pds across a broad postural range that includes both the central and peripheral portions of the workspace, and 2) comparing the results with those predicted by alternative coordinate systems. The height of the sinusoids in Schwartz (1992) ranged from 3 to 12 cm, and their horizontal extent was roughly 15.5 cm. Although detailed postural information was not given, the dimensions, location, and orientation of the 2-D workspace indicate that the monkeys were able to trace the sinusoids without moving the contributing joints through more than a relatively small fraction of their full range of motion. Such was not the case in Caminiti et al. (1990) or Hocherman and Wise (1991), where the larger dimensions of the workspace (30 x 10 x 10 cm and 20 x 20 cm, respectively) required a broader range of joint angles that would make changes in a cell’s spatial pd more easily discernible.

Spatial pds will not vary significantly over small postural changes under any of the three coordinate systems considered, so it is not surprising that Cartesian spatial coordinates engendered good correlations in Schwartz (1992). Further, a definitive analysis must compare correlations under the assumption of Cartesian spatial coordinates versus correlations under the assumption of alternative coordinate systems. Such comparisons are as important as broad workspace sampling, and without them, one cannot make strong inferences about coordinate systems.

Although our analysis has focused on proximal arm movements, the approach can also be applied to the investigation of distal movements. Kakei et al. (1999) performed a direct sampling experiment on movements restricted to the wrist and hand. Preferred directions of MI cells during a latency interval (final 100 ms before movement onset) were determined in three different wrist postures: pronated, supinated, and midway between pronated and supinated. On the basis of the relative size of posture-dependent shifts in cellular pds, Kakei et al. (1999) divided the population into a class of “muscle-like” (sizeable pd shift) cells (32%) and a larger class of “extrinsic-like” (limited pd shift) cells (50%). At least two considerations argue for being cautious in treating these extrinsic-like cells as truly extrinsic. First, roughly 60% of extrinsic-like cells exhibited large posture-dependent gain changes, a response feature analogously found in muscle activations but not expected of a true extrinsic coding scheme. Second, as shown by Scott and Kalaska (1997) and by our simulations, not all cells that encode direction in a purely intrinsic coordinate system will exhibit significant shifts in their pds; it depends on the specific internal pd. More research is needed to clarify the implications of the important results in Kakei et al. (1999).

The generality of the vector field framework makes it applicable to all well-defined coordinate frames, including kinematic, kinetic, and hybrid kinematic-kinetic frames. For example, a plausible hypothesis is that motor cortical cell activity reflects combinations of muscle shortening rates (Mussa-Ivaldi 1988). Support for the idea of bound muscle synergies comes from post-spike facilitation studies (Fetz and Cheney 1978, 1980; Fetz et al. 1976), which suggest that motor cortical cells typically project to motor neurons associated with more than one muscle. Generating predictions for muscle-length coordinates requires a detailed biomechanical model of the arm-muscle system and knowledge of the recruitment patterns by which multiple muscles are synergistically innervated by individual cortical cells. Extending the framework to consider a kinetic, muscle-force based coordinate system remains desirable, particularly for MI, because muscle forces ultimately drive movements and because anatomical and physiological considerations have long shown MI to provide prominent cortical input to the spinal cord and motoneurons. Further, studies have established correlations between MI cell activity and force for multijoint movements (Bullock et al. 1998; Kalaska et al. 1989; Sergio and Kalaska 1998). Unfortunately, an analysis of a muscle force coordinate system would require, in addition to a detailed biomechanical model and knowledge of cortical recruitment patterns, an understanding of all relevant elastic, inertial, and viscous forces involved in center-out hand movements. Given the difficulty in gauging these forces (which are intricately composed, highly complex, and posture-dependent), reliably constructing an explicit muscle-force based coordinate system is an exceedingly difficult task. While we did not attempt to model such a coordinate system, skeletomuscular considerations suggest that a vector field of spatial pds based on muscle forces would possess a highly curved structure. A more efficacious analysis of kinetic coordinate systems, however, can be achieved by applying the vector field framework to an analysis of postural variations of a cell’s preferred direction of force in isometric tasks (Sergio and Kalaska 1997). Finally, the framework can be extended to the analysis of a system of noncanonical coordinates defined by a set of motor primitives like those proposed by Bizzi et al. (1991) to explain the results of stimulating intermediate gray matter in the spinal

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<tbody>
<tr>
<td>Cartesian spatial</td>
<td>Inconsistent</td>
<td>Inconsistent</td>
<td>Inconsistent</td>
</tr>
<tr>
<td>Shoulder-centered Joint</td>
<td>Consistent</td>
<td>Inconsistent</td>
<td>Inconsistent</td>
</tr>
<tr>
<td>Joint angle</td>
<td>Untested</td>
<td>Consistent</td>
<td>Consistent</td>
</tr>
</tbody>
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A summary evaluation of each coordinate system with regard to each of the 3 experiments involving unloaded planar arm movements.
spatial pds is constructed as

\[ \text{curl} \mathbf{v} = \hat{\theta}^\mu_{\nu}[2] + \phi^\mu_{\nu}\left[2 + k \left\{ \sin \left( \theta + \varphi \right) - \cos \left( \theta + \varphi \right) \right\} \right] \]  

This partial derivatives, rather than being explicitly computed, can be taken directly from the first row of the inverse Jacobian to produce

\[ \text{curl} \mathbf{v} = \hat{\theta}^\mu_{\nu}[2] \]

\[ + \phi^\mu_{\nu}\left[2 + k \right\{ \sin \left( \theta + \varphi \right) - \cos \left( \theta + \varphi \right) \right\} \]  

The resulting expression can be simplified using the cosine angle addition formula to yield

\[ \text{curl} \mathbf{v} = 2\hat{\theta}^\mu_{\nu} + \phi^\mu_{\nu} \]

Remarkably, all intermediate dependencies of the curl on hand position and arm posture cancel, leaving a final expression for the curl that depends only on the joint synergy to which the cell is tuned. Thus the rotational tendency of vectors in such a vector field remains uniform across the workspace.

The rest of the APPENDIX, including simulation details and additional derivations, can be found at http://www.cns.bu.edu/pub/ajemian.

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