The Visual Representation of 3D Object Orientation in Parietal Cortex

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An accurate representation of three-dimensional (3D) object orientation is essential for interacting with the environment. Where and how the brain visually encodes 3D object orientation remains unknown, but prior studies suggest the caudal intraparietal area (CIP) may be involved. Here, we develop rigorous analytical methods for quantifying 3D orientation tuning curves, and use these tools to study the neural coding of surface orientation. Specifically, we show that single neurons in area CIP of the rhesus macaque jointly encode the slant and tilt of a planar surface, and that across the population, the distribution of preferred slant-tilts is not statistically different from uniform. This suggests that all slant-tilt combinations are equally represented in area CIP. Furthermore, some CIP neurons are found to also represent the third rotational degree of freedom that determines the orientation of the image pattern on the planar surface. Together, the present results suggest that CIP is a critical neural locus for the encoding of all three rotational degrees of freedom specifying an object’s 3D spatial orientation.

Introduction

Interactions with objects often require determining their 3D spatial orientation based on visual information. Following Hubel and Wiesel’s (1959) discovery of orientation selectivity in primary visual cortex (V1), a diverse range of mechanisms for encoding orientation in the two-dimensional (2D) frontoparallel plane was discovered (DeAngelis et al., 1993; Ringach et al., 1997; Rosenberg et al., 2010). However, extensions to the encoding of 3D object orientation remain surprisingly limited (Taira et al., 2000; Hinkle and Connor, 2002; Liu et al., 2004; Sanada et al., 2012). This reflects that the leap from 2D orientation (a single degree of freedom) to 3D orientation (up to 3 df) is complex for reasons beyond the gap in number of variables. For example, unlike translations, 3D rotations are noncommutative: changing the order of two successive rotations, $R_1$ and $R_2$, generally leads to different final orientations ($R_1 \times R_2 \neq R_2 \times R_1$). Rotations in 3D are also nontrivial in that singularities (orientations where angular variables become locally undefined) are inescapable (Craig, 2005). The consequences of these considerations are far reaching, imposing fundamental limits on the control of the eyes, head, and limbs (Tweed et al., 1999; Crawford et al., 2003; Angelaki and Hess, 2004). Despite its clear behavioral relevance, several factors including inherent mathematical complexity, the use of stimuli confounding shape with orientation, and a lack of suitable analysis methods have greatly limited our understanding of how 3D object orientation is visually encoded.

Here, we investigate how 3D surface orientation is encoded by neurons in area CIP of the macaque monkey. Two angular variables called slant (rotation toward/away from the observer) and tilt (rotation about the line-of-sight) are widely used to parameterize the 3D orientation of a plane (Stevens, 1983). Previous work showed that CIP neurons are tuned for planar tilt, but did not measure slant tuning (Taira et al., 2000). Thus, it is currently unknown whether CIP indeed encodes 3D orientation. In this study, we show that CIP neurons jointly encode slant and tilt, and rigorous analytical methods are developed to provide a quantitative description of how these angular variables are visually encoded. Across the population, the distribution of slant-tilt preferences is found to be statistically indistinguishable from uniform, suggesting that all slant-tilt combinations are equally represented in area CIP. Importantly, we also find that some CIP neurons are sensitive to all three rotational degrees of freedom needed to fully specify an object’s 3D orientation. This work thus suggests that CIP plays a fundamental role in the visual encoding of 3D object orientation, and develops tools necessary for properly characterizing how the brain creates an ecologically complete representation of 3D spatial orientation.

Materials and Methods

Animal preparation. All surgeries and experimental procedures were approved by the Institutional Animal Care and Use Committee, and were in accordance with NIH guidelines. Three male rhesus monkeys (Macaca mulatta) weighing between 5.0 and 7.5 kg were surgically implanted with a lightweight ddrin ring for head restraint and a removable recording grid for guiding electrode penetrations. In separate surgeries, each eye...
was implanted with a scleral search coil for monitoring eye position. After recovery, standard operant conditioning procedures were used to train the monkeys to fixate a visual target within 2° of visual angle. Tilt was varied with red–green anaglyphs. Joint slant-tilt tuning curves were measured in this library. Binocular disparity cues were generated by rendering the stimuli to be viewable only through a single eye. The screen on which visual stimuli were displayed was otherwise black for the first 300 and last 50 ms, and a planar vector may be accomplished using the following rotation matrices:

\[
R_x(\alpha) = \begin{bmatrix}
1 & 0 & 0 \\
0 & \cos(\alpha) & -\sin(\alpha) \\
0 & \sin(\alpha) & \cos(\alpha)
\end{bmatrix},
\]

\[
R_y(\beta) = \begin{bmatrix}
\cos(\beta) & 0 & \sin(\beta) \\
0 & 1 & 0 \\
-\sin(\beta) & 0 & \cos(\beta)
\end{bmatrix},
\]

\[
R_z(\gamma) = \begin{bmatrix}
\cos(\gamma) & -\sin(\gamma) & 0 \\
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\end{bmatrix}.
\]

When describing an object’s spatial orientation, it is important to distinguish between the dimensionality of the space in which the rotations occur and the number of rotational degrees of freedom. Most vision science studies to date focus on 2D orientation (i.e., in two dimensions of motion). The visual responses typically had both transient and sustained components. To calculate the visual response latency, an average spike density function across all trials and the number of rotational degrees of freedom of slant and tilt (Stevens, 1983) that describe the direction of the plane’s unit normal vector (Fig. 2A). Because slant describes rotation toward/away and tilt describes rotation about the line-of-sight, slant-tilt distinguishes rotations that change the extent of an object’s depth variation. Joint slant-tilt tuning curves were measured in this library. Binocular disparity cues were generated by rendering the stimuli to be viewable only through a single eye. The screen on which visual stimuli were displayed was otherwise black for the first 300 and last 50 ms, and a planar vector may be accomplished using the following rotation matrices:

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\]
space) where there is only one rotational degree of freedom and the axis of rotation is perpendicular to the XY plane (i.e., about the Z axis). For 3D orientation (i.e., in three dimensions of space), the axis of rotation can point along any direction in 3D space, providing up to two additional degrees of freedom. Thus, whereas 2D orientation describes rotations around a single axis (i.e., 1 df), 3D orientation describes rotations with three degrees of freedom. Thus, whereas 2D orientation describes rotations around a single axis (i.e., 1 df), 3D orientation describes rotations with three degrees of freedom. Thus, whereas 2D orientation describes rotations around a single axis (i.e., 1 df), 3D orientation describes rotations with three degrees of freedom.

Here, we parameterize the two rotational degrees of freedom describing the direction of a plane’s unit normal vector in terms of slant and tilt (Stevens, 1983). Because 3D rotations are noncommutative, parameterizing 3D orientation requires defining a rotation order. We define the frontoparallel plane (whose normal, \( \hat{n} = [0 \ 0 \ 1]^T \), is aligned with the Z axis) as the reference orientation (Fig. 2Aii). Note that tilt (rotation about the Z axis) is undefined for a frontoparallel plane, reflecting a singularity in the coordinates (i.e., rotation about the Z axis does not change the direction of a frontoparallel plane’s normal vector). Thus, slant (rotation about the Y axis) must be the first rotation (Fig. 2Aii), and tilt must be the second rotation (Fig. 2Aiii).

The configuration space of 2 df rotations (e.g., slant and tilt) is the surface of a unit sphere, denoted \( S^2 \) (Mardia and Jupp, 2000; Vershova and LaValle, 2004; Craig, 2005). For a plane, slant-tilt parameterizes this sphere according to \( R_s(t) \cdot R_t(s) \cdot \hat{n} \), resulting in spherical coordinates \( p = (x, y, z) \):

\[
\begin{align*}
    x &= \cos(t) \cdot \sin(s), \\
    y &= \sin(t) \cdot \sin(s), \\
    z &= \cos(s)
\end{align*}
\]

where \((0 \leq s < 90^\circ)\) and \((0 \leq t < 360^\circ)\) are slant and tilt, respectively. Slant is restricted to be \(<90^\circ\) to eliminate orientations where the plane self-occludes (Cowan et al., 2002). For a planar surface, spatial orientation is unique only up to the sign of the normal vector (i.e., \( \hat{n} = -\hat{n} \)). This axial symmetry imposes antipodal symmetry on the unit sphere (i.e., points \( p \) and \(-p\) are equivalent) describing the configuration space of planar slant-tilt. As a consequence of this geometry, neuronal slant-tilt tuning curves measured with a plane may be visualized on a disc via an equator projection of the sphere’s northern hemisphere in which slant and tilt are arranged using the \((x, y)\) coordinates in Equation 2 (Fig. 2B, C).

Fitting slant-tilt tuning curves. Because a unit sphere describes two rotational degrees of freedom, slant-tilt tuning curves must be fit with a spherical function. However, since two planar orientations \((s, t)\) and \((-s, t + 180^\circ)\) are physically identical (i.e., the configuration space of planar surface orientation is antipodally symmetric), that function must also be antipodally symmetric. The analysis of antipodally symmetric spherical data is commonplace in the field of geomagnetism, and a widely used descriptive model from that literature is the Bingham function (Bingham, 1974; Gubbin and Herrero-Bervera, 2007). We next parameterize the Bingham function as a model for planar slant-tilt tuning curves. Consider a function \( \Psi : S^2 \to \mathbb{R}^1 \) that maps points from the sphere to the real number line.

\[
\Psi(x) = \exp(x^T Ax)
\]
where $\mathbf{x}$ is a unit vector and $\mathbf{A} = \mathbf{A}^T \in \mathbb{R}^{3 \times 3}$ is a $3 \times 3$ symmetric matrix. Let $\mathbf{M}$ denote the matrix of eigenvectors of $\mathbf{A}$ and let $\mathbf{Z}$ be a diagonal matrix whose nonzero entries, $\xi_1, \xi_2, \xi_3$, are the eigenvalues of $\mathbf{A}$. In general, $\mathbf{M}$ can be expressed as a special orthogonal matrix, that is a rotation matrix, such that $\mathbf{M}^T \mathbf{M} = \mathbf{I}$ and $\det(\mathbf{M}) = \pm 1$. We can thus write $\mathbf{A} = \mathbf{M} \mathbf{Z} \mathbf{M}^T$, which is defined by six parameters since there are three eigenvalues plus three angular parameters for the rotation matrix $\mathbf{M}$ (elements $a_{ij} = a_{ji}$ in $\mathbf{A}$). Since columns can be reordered and negated as necessary to maintain $\det(\mathbf{M}) = \pm 1$, then without loss of generality, we order the eigenvalues as $\xi_1 \geq \xi_2 \geq \xi_3$.

$\Psi(\mathbf{x})$ can now be parameterized by a response amplitude, preferred slant and tilt, concentration (bandwidth) parameters, and an additional rotation parameter. Let $\mathbf{y} = \mathbf{M}^T \mathbf{x}$ so that

$$\Psi(\mathbf{x}) = \exp(\mathbf{y}^T \mathbf{Z} \mathbf{y}) = \exp(y_1' \xi_1 + y_2' \xi_2 + y_3' \xi_3).$$

Since we are only concerned with unit vectors (i.e., points on the spherical surface), then $y_1' = 1 - y_1' - y_2'$, and $\Psi(\mathbf{x})$ can be written

$$\Psi(\mathbf{x}) = \exp(y_1' (\xi_1 - \xi_3) + \xi_2 + y_2' (\xi_2 - \xi_3)) = \exp(\xi_1) \cdot \exp(y_1' (\xi_1 - \xi_2) + y_2' (\xi_2 - \xi_3)).$$

Denoting the columns of $\mathbf{M}$ as $\mu_1, \mu_2$, and $\mu_3$, we can express this in terms of $\mathbf{x}$:

$$\Psi(\mathbf{x}) = \exp(y_1' \xi_1) \cdot \exp\left(\frac{\xi_1 - \xi_3}{\xi_1 - \xi_2} (\mu_1^T \mathbf{x})^2 + \frac{\xi_1 - \xi_2}{\xi_1 - \xi_3} (\mu_2^T \mathbf{x})^2\right).$$

In simplified form and introducing a DC offset,

$$\Psi(\mathbf{x}) = dc + G \cdot \exp(\lambda_1 (\mu_1^T \mathbf{x})^2 + \lambda_2 (\mu_2^T \mathbf{x})^2),$$

where the three orthonormal vectors ($\mu_1, \mu_2, \mu_3$) are defined as follows:

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>dc</td>
<td>DC offset</td>
</tr>
<tr>
<td>$G \geq 0$</td>
<td>Gain</td>
</tr>
<tr>
<td>$0^\circ \leq \phi &lt; 90^\circ$</td>
<td>Preferred slant</td>
</tr>
<tr>
<td>$0^\circ \leq \phi &lt; 360^\circ$</td>
<td>Preferred tilt</td>
</tr>
<tr>
<td>$0^\circ \leq \phi &lt; 180^\circ$</td>
<td>Rotation about the preferred slant-tilt</td>
</tr>
<tr>
<td>$\lambda_1 \leq 0$</td>
<td>Tuning aspect ratio (&quot;ellipticohedral&quot;)</td>
</tr>
<tr>
<td>$\lambda_2 \geq 0$</td>
<td>Tuning bandwidth</td>
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</table>

Because the individual columns of $\mathbf{M}$ are orthonormal, any one of them may be expressed as the cross product of the other two. Consequently, $\mu_3$ drops out of Equation 7. The DC term is a constant firing rate, $G$ sets the response amplitude, and the parameters $s^*$ and $t^*$, are the preferred slant and tilt, respectively. The preferred slant-tilt is set by $\mu_3$ and the tuning bandwidth is set by $\lambda_2$. Larger positive values of $\lambda_1$ make the tuning curve narrower. The $\lambda_1$ term determines the aspect ratio of the function, with $\lambda_1 = 0$ being circularly symmetric and increasingly negative values making the function more elliptical (e.g., creating broader slant than tilt tuning). Last, $\phi$ rotates the function about the preferred slant-tilt. A geometric description of the Bingham function and summary of its seven parameters is presented in Figure 3. To avoid local minima when fitting, each tuning curve was fit with multiple initial parameter sets and perturbation techniques were applied to the converged parameter sets. The solution resulting in the highest correlation between data and model was retained.

When parameterized for the slant and tilt of a planar surface, the Bingham function may be visualized on the slant-tilt disc described above and illustrated in Figure 2, B and C, by removing the $z$ coordinate through an equatorial projection of the sphere’s northern hemisphere.
Results

Single neurons in area CIP encode the slant-tilt of a planar surface

Previous work showed that approximately half of CIP neurons are tuned for the tilt of a plane at a fixed slant, but did not examine their sensitivity to changes in slant (Taira et al., 2000; Tsutsui et al., 2001). To determine whether CIP neurons jointly encode slant and tilt, we recorded the responses of 161 visually responsive cells (Fig. 1) while varying both the slant and tilt of a planar surface rendered on an LCD with a checkerboard texture pattern and binocular disparity cues. Tilt tuning curves measured at different slants are shown for a single neuron in Figure 2C along with the joint slant-tilt tuning curve. This cell responded most strongly to planar surfaces leaning toward the monkey, but there was a wide range of diversity in the slant-tilt tuning curves across the population. To characterize these diverse responses quantitatively, it was necessary to develop analytical tools appropriate for characterizing neuronal tuning curves for 3D object orientation. This is done in the Materials and Methods, where the Bingham function (Bingham, 1974; Mardia and Jupp, 2000) is parameterized as a model for slant-tilt tuning curves. In Figure 3, the Bingham function is described geometrically and its parameters are summarized.

In Figure 4, example CIP slant-tilt tuning curves are shown along with Bingham function fits. The cell in Figure 4A preferred a frontoparallel plane (the tuning curve is nearly centered on the slant-tilt disc, $s = 2^\circ$) and the tuning was approximately circularly symmetric ($\lambda_1 = -0.21$; tuning is isotropic when $\lambda_1 = 0$). The cell in Figure 4B preferred an oblique tilt with the lower left side of the plane closest to the monkey. The fitted tilt and slant parameters were $t = 231^\circ$ and $s = 39^\circ$, respectively. Hence, the peak of the tuning curve is positioned in the lower left of the slant-tilt disc and centered at an intermediate radial distance from the origin ($s = 0^\circ$). For the cell in Figure 4C, the high firing rates on opposite sides of the slant-tilt disc (compare the example in the third row of Fig. 3B) reflect the cell’s large slant preference ($s = 87^\circ$) combined with the fact that opposite sides of the slant-tilt disc

(3) with $A = A' \in \mathbb{R}^{4 \times 4}$, $\hat{A} \in \mathbb{R}^{3 \times 3}$ and $\hat{A} \in \mathbb{R}^{2 \times 2}$ symmetric matrix, in Equation 3. This defines a 1 df Bingham function, which is equivalent to a von Mises function with $\pi$ periodicity. Planar slant-tuning curves should thus be well described by a $\pi$ periodic von Mises function. Importantly, the Bingham function is not limited to describing planar surface orientation. For example, neuronal tuning curves for the slant-tilt of a bar can be described using a similar parameterization in which the two angular variables map differently to the sphere to reflect that a bar’s tilt is defined in the frontoparallel plane. This is done in the Materials and Methods, where the Bingham function (Bingham, 1974; Mardia and Jupp, 2000) is parameterized as a model for slant-tilt tuning curves. In Figure 3, the Bingham function is described geometrically and its parameters are summarized.

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(Fig. 3). Under this projection, the singularity at $s = 0^\circ$ (the frontoparallel plane) where all tilts map to the same point (the north pole of the sphere) lies at the origin. Cross-sections through the origin “wrap around,” reflecting the fact that points on opposite edges of the slant-tilt disc correspond to identical orientations. Two examples in which this wrap around is apparent are shown: (1) in the third row of Figure 3B and (2) in Figure 4C. Cross-sections of a Bingham function which pass through the origin of the slant-tilt disc (i.e., slant tuning curves) reduce to setting the third angular variable is measured at all sampled slant-tilt combinations. Antipodal symmetry ($\slant = -\slant$) reflects the fact that points on opposite sides of the slant-tilt disc correspond to similar orientations (see Materials and Methods).
correspond to similar orientations (a consequence of the antipodal symmetry of the space describing planar surface orientation, see Materials and Methods). This cell’s tuning was relatively isotropic ($\lambda_1 = -0.34$) and narrow ($\lambda_2 = 1.21$). The cell in Fig. 4D also preferred large slants ($s = 86^\circ$), but its tuning was anisotropic ($\lambda_1 = -1.35$) and broader ($\lambda_2 = 0.40$), with broader tuning along the tilt than slant axis ($\phi = 100^\circ$). Tuning is broader along the tilt axis when $\phi$ is near $90^\circ$, and broader along the slant axis when $\phi$ is near $0^\circ$. This anisotropy results in the crescent shape of the tuning curve. As these examples illustrate, the Bingham function generally provided a strong fit to the data: the average fit correlation was $r = 0.88 \pm 0.09$ SD, $N = 161$ (Fig. 4). However, visual inspection indicated that 17 of these cells had multiple distinct modes in their tuning, and therefore lacked a unique slant-tilt preference (Fig. 5). Their tuning curves were thus poorly described by the Bingham function (average fit correlation: $r = 0.68 \pm 0.11$ SD). Interestingly, the majority of them (15/17) were encountered in left hemisphere recordings (monkeys X and P), although they were intermixed with slant-tilt selective cells (no clustering was observed). Except where otherwise noted, these cells were excluded from further analysis. The remaining 144 neurons were all selective for a unique slant-tilt and their tuning curves were well described by the Bingham function (average $r = 0.91 \pm 0.05$ SD). This result demonstrates that CIP neurons jointly encode the slant and tilt of a planar surface.

**Area CIP encodes a uniform distribution of planar slant-tilts**

The computation of planar slant-tilt (Marr’s “2.5D sketch”) is critical to constructing an accurate 3D spatial representation of the visual scene (Marr, 1982). However, whether or not the visual encoding of slant-tilt is unbiased in the number of neurons representing any particular orientation, or subset of orientations, remains untested in any neural population. In part, this reflects that analytical methods appropriate for quantifying 3D object orientation-tuning curves did not previously exist. One possible finding is that all orientations are uniformly encoded, which may be ideal for guiding interactions with objects (e.g., grasping) independent of the orientation in which they are encountered. Alternatively, the distribution of orientation preferences may be systematically biased (e.g., for ground-plane estimation), in a way that makes the population more sensitive to certain orientations (Gu et al., 2010). Using the Bingham function fits to estimate the preferred slants and tilts of the 144 slant-tilt selective neurons, we next examine the distribution of surface orientations represented by the CIP population.

As described in the Materials and Methods, slant and tilt are spherical coordinates. To examine the distribution of these angular variables, we performed a Lambert-like cylindrical projection of the slant-tilt preferences (Snyder, 1987). Under this area-preserving coordinate transformation, tilt is represented directly in the angular variable (e.g., the difference between tilts of $0^\circ$ and $30^\circ$ is the same as between $240^\circ$ and $270^\circ$) and slant is represented in the cosine of the angle (Fig. 6A). Uniformity or bias in the distribution of orientation preferences found in this transformed space implies the same over the spherical surface. The joint distribution of slant and tilt preferences following the equal area

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**Figure 5.** Tuning curves of cells not selective for planar slant-tilt. Six tuning curves illustrating the range of observed responses that were not tuned for a unique slant-tilt. Responses are baseline subtracted and firing rate is color coded. **A**, Tuning curves of three neurons for which sensitivity to the orientation of the frontoparallel checkerboard image was not tested. Like the slant-tilt tuning curves plotted in Figure 4, the response at the origin was to a frontoparallel plane with a checkerboard image with $\theta = 0^\circ$ (Fig. 5A). **B**, Tuning curves of three neurons for which sensitivity to the orientation of the frontoparallel checkerboard image was tested. The average of these responses is plotted at the origin. Tilt tuning curves measured at different slants are shown for each of these three neurons (as in Fig. 2C). Error bars are SEM.

**Figure 6.** Area CIP encodes a uniform distribution of slant-tilts. **A**, Slant–tilt is a spherical coordinate system describing the direction of a planar surface’s unit normal vector. For a frontoparallel plane, the normal vector (red arrow), $\mathbf{n}$, aligns with the Z axis of the reference frame (compare Fig. 2A). The pair of antipodal yellow points connected by a yellow line segment represents a unique 3D surface orientation. The red curve traces the path that the plane’s normal would travel from a frontoparallel orientation to the orientation specified by the pair of yellow points assuming motion along the first coordinate (slant) followed by motion along the second (tilt). The projection cylinder preserving area in the joint distribution (a Lambert-like projection) is also shown, with blue arrows illustrating the transformation. **B**, The joint distribution of slant-tilts following this transformation. **C**, Marginal distribution of slant-tuning preferences. **D**, Marginal distribution of tilt tuning preferences. In **B-D**, tilt is plotted directly in the angular variable and slant is plotted in the cosine of the angle.
projection is shown in Figure 6B. This distribution is not significantly different from uniform ($\chi^2, p = 0.26$). The marginal distribution of slant preferences (in the cosine of the angle) is shown in Figure 6C, and the marginal distribution of tilt preferences is shown in Figure 6D. Neither of these distributions is significantly different from uniform ($\chi^2, p \approx 0.27$). Unlike V1 that over-represents the cardinal orientations of a contour in the 2D frontoparallel plane (Li et al., 2003), these results suggest that CIP may represent all possible combinations of slant and tilt equally.

Population histograms showing the distributions of the other shape-determining parameters of the Bingham function are shown in Figure 7. These parameters, described geometrically in Figures 3 and 7A, rotate the intermediate and minor axes of the tuning curve about the preferred slant-tilt ($\phi$, Fig. 7B), set its aspect ratio—allowing tuning to be broader along one axis ($\lambda_1$, Fig. 7C), and set the overall tuning bandwidth ($\lambda_2$, Fig. 7D). Most typically, the intermediate and minor axes were aligned with the azimuth and elevation of the sphere, respectively (Fig. 7B). This indicates that variation in tuning bandwidth tended to occur across the tilt and slant (as opposed to oblique) axes (Fig. 8A). Because the shape of a tuning curve is at a minimum specified by a peak and a bandwidth, we further examined whether the two parameters ($\lambda_1$ and $\phi$) allowing the Bingham function to be more elongated along a particular axis were necessary to accurately describe CIP slant-tilt tuning curves. The tuning curves of the 144 slant-tilt selective neurons were fit with a model made simpler by setting $\lambda_1 = 0$, which also eliminates the $\phi$ parameter (see Eqs. 7, 8). Removing these two parameters constrains the Bingham function to be isotropic (i.e., equal slant and tilt tuning bandwidths) and is therefore a minimal model accounting for the geometry of planar slant-tilt with only five parameters (DC offset, gain, slant preference, tilt preference, and bandwidth). This function typically provided an adequate description of CIP slant-tilt tuning curves (average $r = 0.84 \pm 0.09$ SD). Using Akaike’s Information Criterion corrected for finite sample size (Burnham and Anderson, 2002) to compare model fits, the unconstrained (seven parameter) Bingham function was found to provide a better description of the majority of tuning curves (127/144; 88%; Fig. 8B). This suggests that the unconstrained Bingham function is generally the simplest model providing an accurate description of CIP planar slant-tilt tuning curves.

**Sensitivity to the third rotational degree-of-freedom**
A complete assessment of a neuron’s orientation-tuning properties requires measuring its sensitivity to all three rotational degrees of freedom describing 3D object orientation. Prior studies focused on tilt and to a lesser extent slant, but sensitivity to all 3 df was never before tested. For a planar surface, this third degree of freedom can be parameterized as a rotation about the normal vector ($0^\circ \leq \theta < 360^\circ$), and describes the orientation of the image on the face of the plane. To test for sensitivity to this third degree of freedom, we interleaved with the slant-tilt measurements a frontoparallel plane with the checkerboard image at different orientations for 92 CIP neurons (Fig. 9A). Note that this rotation is
distinct from tilt (Fig. 2A) since it never changes the direction of the plane’s normal vector. Of these 92 cells, 86 were slant-tilt selective (Fig. 4) and 6 were not (Fig. 5). An ANOVA was used to determine whether the firing rate of each cell depended significantly (p < 0.05) on the image’s orientation (Fig. 9B). The responses of approximately 16% (14/86) of the slant-tilt selective neurons were significantly tuned for the third rotational degree of freedom, and most of these preferred intermediate slants (Fig. 9C). Because the tuning curves of cells with intermediate slant preferences will be steepest near frontoparallel, they will be highly sensitive to orientation changes in this region (Purushothaman and Bradley, 2005). This suggests that if sensitivity to the third angular variable was always tested near inflection points of the slant-tilt tuning curve (i.e., where the derivative is greatest in magnitude), then significant tuning may have been observed more often. Interestingly, a higher percentage of cells lacking a unique slant-tilt preference, 67% (4/6) were tuned for the orientation of the checkerboard image. The “4-prong star” appearance of some of their tuning curves (Fig. 5) may thus be attributable to this sensitivity to the orientation of the image within the plane combined with the 90° periodicity of the cross-hatching of a checkerboard pattern (Fig. 9A). This suggests that these cells may be directly involved in encoding the orientation of surface images.

Discussion

In this study, we performed a quantitative analysis of how 3D surface orientation is visually encoded. The fundamental advance making this possible was the development of computational tools designed to address the mathematical complexities of 3D rotations. These tools were designed to be highly flexible in that they may be parameterized to quantify orientation tuning curves measured for different objects with one, two, or three rotational degrees of freedom. Here, we focused on 2 df rotations, using the tools to characterize how individual neurons in area CIP of the macaque monkey jointly encode the slant and tilt of a planar surface. Across the population, the distribution of slant-tilt preferences was statistically indistinguishable from uniform, tiling the surface of a sphere. From a Bayesian perspective, this suggests that CIP population activity encodes a uniform prior over 3D orientation (i.e., one in which all orientations are equally likely). The sensitivity of CIP, for example to discriminate two similar orientations, may therefore be independent of the objects’ absolute orientations in space (Gu et al., 2010). Such uniform coverage of 3D orientation may also be ideal for guiding interactions with objects, consistent with CIP’s projection to AIP, an area implicated in the grasping of objects (Nakamura et al., 2001). Importantly, we also showed that single CIP neurons can visually encode all three rotational degrees of freedom needed to fully specify an object’s 3D spatial orientation. These findings thus suggest that area CIP has a fundamental role in the visual encoding of 3D object orientation.

The slant-tilt parameterization of 3D orientation distinguishes rotations that change the extent of an object’s depth variation (slant) from those that do not (tilt). This is conceptually appealing since sensitivity to these two angular variables emerges sequentially within the visual system. Specifically, tilt selectivity emerges first as a robust tuning property in primary visual cortex (Hubel and Wiesel, 1959; Xu et al., 2002; Rosenberg et al., 2010) where there is limited, if any, representation of slant (Blakemore et al., 1979; Nelson et al., 1977; Bridge and Cumming, 2001; Sanada and Ohzawa, 2006). Several previous reports suggest that slant selectivity may be found in a number of extrastriate visual areas (Hinkle and Connor, 2002; Nguyenkim and DeAngelis, 2003; Liu et al., 2004), but slant tuning was only measured systematically in area V4, and no study characterized the joint representation of slant and tilt. The present results build qualitatively upon these and other earlier findings. For instance, prior work in CIP only measured tilt tuning (Taira et al., 2000), whereas we demonstrated that single neurons can visually encode all three rotational degrees of freedom. In area MT, tilt tuning was reported to emerge at large slants for stimuli rendered as linear gradients of horizontal disparity (Nguyenkim and DeAngelis, 2003). However, because of the nonlinear relationship between disparity and depth (Cormack and Fox, 1985), the stimulus became increasingly curved as the slant angle increased, confounding shape with orientation tuning. In addition, MT neurons that were tilt selective at large slants also responded strongly to small slants. It thus remains unclear whether MT neurons show band-pass tuning for slant-tilt, as we found in CIP using stimuli without a shape–orientation confound. In a study conducted in the dorsal medial superior temporal area, stimuli were rendered such that two surfaces leaning in opposite directions produced identical retinal stimulation, and therefore the results cannot be inter-
preted as surface orientation tuning (Sugihara et al., 2002). One of the clearest prior demonstrations of 3D orientation selectivity was in V4, where tuning was measured using a bar (Hinkle and Connor, 2002). However, because the data were not quantified analytically, the distribution of slant and tilt preferences represented in area V4 remains unclear. In future studies it will be interesting to determine whether other visual areas exhibiting 3D orientation selectivity encode a uniform distribution of orientations, or whether they are biased in some way that creates greater sensitivity to particular orientations. It will also be important to examine whether the 3D orientation selectivity of single neurons generalizes across objects (e.g., bars and planes). At the same time that the symmetries of achiral stimuli such as bars and planes are convenient for demonstrating 3D orientation selectivity, such objects only partially sample the full space of 3D rotations. In particular, they either constrain or eliminate the third degree of freedom (Fig. 9A) and make 1 or 2 df rotations antipodally symmetric. Although antipodally symmetric tuning (i.e., tuning that is 180° rather than 360° periodic) is an inherent property of some neurons encoding either orientation (DeAngelis et al., 1993; Rosenberg and Issa, 2011) or direction of motion (Rosenberg et al., 2008), further examination may reveal that this is not the case in CIP. Indeed, observations from this study suggest that CIP neurons may generally be tuned for all three rotational degrees of freedom describing object orientation. First, since the orientation of a frontoparallel checkerboard image was sufficient to drive a tuned response for some neurons, a more orientable image such as a face (i.e., one having a periodicity of 360°) may be more likely to reveal significant tuning. Second, for a planar surface, the third rotational degree of freedom only affects the texture pattern on the face of the plane. The presentation of an object with a more elaborate (nonplanar) 3D shape where any rotation is signaled both by disparity and texture changes would be more likely to elicit tuned responses for all three rotational degrees of freedom. Considering these observations, finding that the orientation of a frontoparallel checkerboard image can elicit tuned responses suggests that CIP neurons are exquisitely sensitive to all three rotational degrees of freedom describing an object’s 3D spatial orientation. To fully explore the space of 3D rotations, it will be necessary to measure responses to stimuli with some symmetry-breaking feature (e.g., a bar or plane with a curve at one end or edge). Tuning over this space can be quantified using a Bingham function parameterized for all three rotational degrees of freedom. Last, an intriguing question to ask is whether CIP (or any other 3D orientation selective area) dynamically changes its coverage of spatial orientation in a task-dependent manner. If so, this would imply that the representation of 3D object orientation updates in a context-dependent manner similar to the contextual remapping of the representation of space that occurs in the hippocampus (Smith and Mizumori, 2006). Consistent with this possibility, contextual features such as required grip force, viewing conditions, and intended object manipulation modify grasp representations in both macaque AIP (Baumann et al., 2009), which receives input from CIP (Nakamura et al., 2001), and the human homolog of AIP (Verhagen et al., 2008; Maranong et al., 2011).

References


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