The Fine Structure of Shape Tuning in Area V4

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SUMMARY

Previous studies have shown that neurons in area V4 are involved in the processing of shapes of intermediate complexity and are sensitive to curvature. These studies also suggest that curvature-tuned neurons are position invariant. We sought to examine the mechanisms that endow V4 neurons with these properties. Consistent with previous studies, we found that response rank order to the most- and least-preferred stimuli was preserved throughout the receptive field. However, a fine-grained analysis of shape tuning revealed a surprising result: V4 neurons tuned to highly curved shapes exhibit very limited translation invariance. At a fine spatial scale, these neurons exhibit local variation in orientation. In contrast, neurons that prefer straight contours exhibit spatially invariant orientation-tuning and homogenous fine-scale orientation maps. Both of these patterns are consistent with a simple orientation-pooling model, with tuning for straight or curved shapes resulting, respectively, from pooling of homogenous or heterogeneous orientation signals inherited from early visual areas.

INTRODUCTION

Visual shape information is processed in the ventral cortical pathway, which progresses from the primary visual cortex (V1), the secondary cortex (V2), intermediate areas (V3/V4), and finally onto the inferotemporal (IT) cortex (Felleman and Van Essen, 1991). In the earlier stages, shape is encoded primarily through local orientation in V1 (Hubel and Wiesel, 1959, 1965, 1968) and combinations of orientations in V2 (Anzai et al., 2007; Tao et al., 2012). At the final stages in IT, cells have been shown to be selective for complex objects like faces (Desimone et al., 1984; Tanaka et al., 1991; Tsao et al., 2006). How this transformation is achieved remains largely unknown. In addition, the selectivity to complex features becomes more invariant to simple stimulus transformations such as size or spatial position as one traverses the ventral cortical hierarchy (Rust and Dicarlo, 2010). To understand how contours of objects are integrated into coherent percepts in the later stages, a detailed understanding of shape processing in intermediate stages like V4 is critical.

Previous studies (Pasupathy and Connor, 1999, 2001) demonstrate that neurons in monkey visual area V4 are involved in the processing of shapes of intermediate complexity and are sensitive to curvature. These studies showed that V4 neurons responded more strongly to a preferred stimulus, as compared to a null stimulus, throughout the receptive field (RF)—a form of translation invariance. However, little is known about the mechanisms that underlie shape tuning of neurons in area V4 or about the degree to which translation invariance depends on stimulus complexity. Using a dense mapping procedure, we sought to understand the detailed structure of shape selectivity within V4 RFs.

RESULTS

We analyzed responses from 93 isolated neurons in area V4 of two awake-behaving male macaques (see Experimental Procedures). The stimuli consisted of oriented bars presented alone or linked end to end to form curves or in the most tightly curved conditions: “C” shapes (Figure 1A). Bars were presented at eight orientations. Composite shapes were composed of three bars linked together to yield five categories of shapes: straight, low curvature, medium curvature, high curvature, and C shaped. Stimuli were presented in fast reverse correlation sequences (16 ms duration, exponential distributed delay between stimuli with a mean delay of 16 ms) at various locations within the RF of peripheral V4 neurons (2°–12° eccentricity) while the monkeys maintained fixation for 3 s. The composite shapes were presented on a 5 × 5 location grid centered on the RF, while the oriented bars were presented on a finer 15 × 15 location grid. The grid of locations and the size of visual stimuli were scaled with RF eccentricity to maintain the same proportions as shown in Figure 1A. A pseudorandom sequence from the combined stimulus sets was presented in each trial.

We found that the majority of neurons in our population were significantly selective to the composite contours. Example neurons with significant tuning for composite contours are illustrated in Figure 2 (neurons I, II, and III). The middle panels (labeled B) show the mean firing rate response to each of the composite forms tested (5 × 16 array) at the most responsive spatial location. The adjacent panels to the right show the Z scores of the responses after subtracting the mean spatial response (see Experimental Procedures and Figure S1A, available online, for details of assessing significance). Example neuron I is preferentially tuned to straight shapes, neuron II to medium-curvature...
shapes, and neuron III to high-curvature/C shapes. Neuron IV had a significant spatial response but no significant shape selectivity. The distribution of spatial and shape selective tuning is shown in Figure 1B. Across the population, 80 of 93 neurons showed significant shape selectivity while a smaller subset (n = 13, labeled in blue) had spatial tuning without significant shape tuning. We did not analyze this subset further.

Furthermore, among neurons with significant shape selectivity, those preferring either straight or more curved stimuli exhibited similar degrees of selectivity (Figure 1C). There was no correlation between the degree of selectivity and shape preference.

**Straight- and Low-Curvature-Tuned Neurons Exhibit Spatial Invariance**

We find that neurons that are tuned for straight (zero-curvature) or low-curvature contours are spatially invariant in their tuning. That is, they respond preferentially to the same shape in different parts of the RF. The response characteristics of an example neuron are shown in Figure 3 (example neuron I). Earlier studies (Pasupathy and Connor, 1999) examined spatial invariance by comparing the neuronal responses to the most (black bar) and least (white bar) preferred stimulus across different spatial locations, as seen in the lower right panel of Figure 3A. Our fast mapping procedure allowed us to estimate the selectivity...
for the full set of composite shapes at different spatial locations. Examination of the location-specific response maps taken from four significant response locations (Figure 3B) reveals the neuron’s full spatial invariance. The local maps show clear tuning for straight shapes, with an orientation preference that is shared across locations. This point is further clarified by plotting the shape (or set of shapes) to which the neuron preferentially responds at different locations of the stimulus grid. This is shown in Figure 3A (bottom-left panel), in which the set of shapes to which the neuron responded (greater than 90% of local peak rate) at each location are spatially superimposed (color indicates firing rate). This spatial invariance to orientation tuning is also reflected in the homogeneity of the fine-scale orientation-tuning map obtained from the bar stimuli on the 15 × 15 grid (Figure 3C). Several other examples of straight- and low-curvature-tuned neurons exhibiting spatial invariance are shown in Figure S2.

**Higher-Curvature- and C-Shape-Tuned Neurons Are Not Spatially Invariant**

In contrast, we found that neurons tuned for curved (medium to high) and C-shaped stimuli exhibited a high degree of spatial variation in their shape preference. Two such example neurons are shown in Figure 3 (neurons II and III). In both cases, comparing the relative responses evoked by the most and least preferred stimuli across locations (Figure 3A, lower right panels) suggests a degree of spatial invariance, consistent with earlier studies (Paspas and Connor, 1999). However, the pattern of selectivity to the full set of stimuli across locations reveals that the preferred stimulus varies considerably across locations. Example neuron II exhibits selectivity for distinct clusters of medium-curvature shapes in different parts of its RF (Figure 3B). The fine-scale orientation-tuning map for this neuron (Figure 3C) shows that although there is relatively sharp tuning for orientation at each location, there is a systematic variation in tuning across locations, and this variation appears to be correlated with the neuron’s spatially varying curvature preference. Note that the average fine-scale orientation response (Figure 3C, left inset) for this neuron is not tuned and therefore does not reflect the diversity of orientation tuning at the fine scale. Such a neuron would be mischaracterized as nonorientation selective if mapped at a coarse level.

**Figure 2. Diversity of Shape Tuning**

(A) Four example neurons. Top: average time course to the composite stimuli ± SD. The average was taken across all stimuli and across all locations on the 5 × 5 grid. The dotted line is the baseline rate, which was determined from a temporal window between 0 and 20 ms after stimulus onset. The dotted gray box marks the temporal window where the average firing rate exceeded the baseline rate by 4 SDs. This temporal window was used for all subsequent analysis. Bottom: spatial RFs obtained by averaging responses across the composite shape stimuli. Significant spatial locations are marked with “x” (see Experimental Procedures). Contour lines demarcate 90%, 80%, 70%, and 60% of maximum response. These were obtained by spatial interpolation of the RF.

(B) Location-specific response map at the most responsive spatial location (marked with black “x” in the spatial RF in A). The composite stimuli are overlaid on the maps for ease of reference.

(C) Shape Z score maps (see Experimental Procedures) for the same spatial location in (B). Neuron I is significantly selective for straight shapes, neuron II for medium curvature, and neuron III for high curvature/C. Neuron IV is not significantly shape selective. Contour lines demarcating Z scores at the 0.05 (outer contour) and the Bonferroni corrected (inner contour) levels are superimposed on the response maps in (B). For ease of visualization, all color maps in (B) and (C) were smoothed with a Gaussian kernel.

See also Figures S2 and S3.
Example neuron III shows similar spatially varying preference for the C stimuli and a heterogeneous fine-scale orientation map. We see evidence for tuning along both dimensions of our stimulus space: orientation (e.g., neuron III, location 4) and shape category (e.g., neuron II, locations 2 and 4). We considered if neurons selective to highly curved shapes might be less tuned to the orientation of the shape. However, at the population level, we find that orientation tuning, as indexed by circular variance (see Supplemental Experimental Procedures), is not correlated with shape preference (Figure S1C). We also considered if these neurons might be less tuned in the shape dimension (Figure S1B). Again, we find that at the population level, an index of shape tuning (see Supplemental Experimental Procedures) is not correlated with shape preference (Figure S1D). Other examples of neurons exhibiting spatial variation in shape preference are shown in Figure S3.
Heterogeneity of Feature Selectivity across RF Locations of Neurons Tuned for Higher-Curvature or C-Shaped Stimuli

To quantify the relationship between curvature preference and spatial invariance at the population level, we examined two complementary aspects of the neuronal data. First, we computed the shape preference and the preferred orientation at each location in the stimulus presentation grid where the neuron responded significantly (see Experimental Procedures). As one measure of translation invariance, we determined the preferred shape and orientation at the maximally responsive location and measured how shape and orientation preferences changed relative to those values at other spatial locations (Figure 4). We find a clear difference in spatial invariance between the population of cells that prefer straight/low curvature (local shape preference values between 0 and 1, n = 32; Figure 4A) versus those that either prefer medium curvature (local shape preference values between 1.5 and 2.5, n = 16; Figure 4B) or high-curvature/C-shaped stimuli (local shape preference values between 3 and 4, n = 20; Figure 4C). We find that those neurons that preferred straight or low curvature at the most responsive location tend to be tuned for similar orientations at other RF locations and preserve their shape preference across locations (Figure 4A). In contrast, although neurons that prefer high curvature at their maximally responsive location continue to prefer high curvature at other locations within the RF, the preferred shapes do not generally share the same orientation (Figure 4C). Similarly, neurons with preference for medium curvature at their maximally responsive location tend to prefer medium curvature at other locations, but the preferred shapes are not as sharply aligned with the reference orientation (Figure 4B). The marginal distribution of orientation preference for the straight/low-curvature neurons (Figures 4A, right histogram) was significantly different from those of the other two subpopulations (Figures 4B and 4C, right histograms; p = 0.03 and p = 0.006, respectively; see Experimental Procedures).

Second, we compared neuronal response patterns across the entire set of curved shapes between pairs of locations within the RF. For any pair of location-specific response maps where the neuron responded significantly, we estimated the empirical distribution of correlation coefficients between the response patterns (see Experimental Procedures; Figure S4). The mean pattern correlation (ρ, expected value of the empirical distribution) provides a measure of tuning similarity or invariance between pairs of locations in the RF, with values closer to 1 corresponding to spatially invariant tuning. The average pattern correlation for each neuron (averaged across all pairwise ρ values) when plotted against the average shape preference (Figure 5A) shows a power-law decay relationship. Neurons with preference for medium curvature and higher tend to have little spatial invariance. In contrast, neurons with very low-curvature preference tend to have substantial spatial invariance, with few units exhibiting low invariance.

For each location pair in our population, we also calculated the reliability of the estimated pattern correlation from the SD of the empirical distribution (see Experimental Procedures). This controls for the possibility that noisier data gave rise both to greater response heterogeneity and lower pattern correlations. A
scatterplot of pattern correlation ($\rho$) versus pattern reliability ($r$) is shown in Figure 5B for all possible location pairs across all neurons in our population. We see no difference in the reliability of our estimates for three subpopulations of location pairs: those that come from neurons with average shape preference for straight/low curvature, those that came from neurons with average shape preference for medium curvature, and those that came from neurons with average shape preference for high-curvature/C-shaped stimuli. The reliability distributions are not significantly different from each other.

Figure 5. Neurons with Preference for Curvature Have Limited Spatial Invariance

(A) Average correlation between pairs of response patterns (averaged across all possible pairs of spatially significant response locations for each neuron; see Experimental Procedures) plotted against the average shape preference for all neurons in our population ($n = 80$) shows an inverse power law relationship ($R^2 = 0.4$). Average pattern correlation is high for neurons tuned for straight/low curvature, while the pattern correlation is low for neurons tuned for high curvature/C, indicating a trade-off between curvature and spatial invariance. The three example neurons in Figures 2 and 3 are indicated.

(B) Scatterplot of pairwise pattern correlation versus pairwise pattern reliability for all possible pairs of significant response locations in our entire neuronal population. The colors indicate the average shape preference of the neuron to which the location pair belongs. Right histograms: distribution of pattern correlation for pairs that came from three sub-population of neurons: neurons with average shape preference for straight/low curvature (shape preference values between 0 and 1), those that came from neurons with average shape preference for medium curvature (shape preference values between 1.5 and 2.5), and those that came from neurons with average shape preference for high curvature/C (shape preference values between 3 and 4). The correlation distribution of the straight/low-curvature subpopulation is significantly different from those of the other two subpopulations. Bottom histograms: distribution of pattern reliability for the same three subpopulations as above. The reliability distributions are not significantly different from each other.

See also Figure S4.

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the fine-scale orientation-tuning maps in our example cells (Figure 3C) seems to reflect the cell’s shape-selective properties. It has been proposed, both from experimental observations (Chapman et al., 1991; Jin et al., 2011) and theoretical simulations (Paik and Ringach, 2011), that simple pooling of the spatially segregated afferent connections from the lateral geniculate nucleus (LGN) to the primary visual cortex (V1), might determine both the orientation-tuning characteristics of V1 neurons as well as the pinwheel structure of orientation maps in V1. We hypothesized that this pooling architecture might carry forward to downstream retinotopic extrastriate areas like V4. This hypothesis is also consistent with earlier proposals, in which neuronal responses in V4 to combinations of line elements are weighted averages of the responses evoked by individual elements (Ghose and Maunsell, 2008; Lee and Maunsell, 2010; Reynolds et al., 1999; Reynolds and Heeger, 2009), and with related proposals in MT (Heuer and Britten, 2002; Rust et al., 2006) and IT (Zoccolan et al., 2005).

To examine the degree to which the pooling of orientation signals could account for the observed complexity of shape tuning in V4 neurons, we generated location specific-response predictions to the composite shapes derived from a simple weighted average of the component responses obtained from the fine-scale orientation-tuning maps (see Experimental Procedures). We then calculated the correlation coefficient between the observed response pattern and the predicted response pattern. Note that the fine-scale orientation maps contain both a spatial response component and an orientation-tuning component. To investigate the contribution of these components, we also considered two reduced versions of the pooling model (see Experimental Procedures; Figure S5C). A space-only version was obtained by averaging across orientation at each fine-grid location. This model did not have any local orientation tuning. An orientation-only version was obtained by subtracting the space-only response from the measured data at each fine-grid location, leaving only orientation tuning. Thus, this model did not contain any local spatial information.

The predicted response maps for two example neurons (neurons II and III in Figures 2 and 3) are shown in Figure 7A (panels labeled “prediction”). Maps are shown for three different RF locations for each neuron. For the RF location marked “1”, the left panel shows the empirical data, while the other three panels show the predicted responses from the full model and the two reduced models. Shown below the predicted response maps are the corresponding sections of the fine-scale orientation map, which were used to generate the predictions. To take the example of RF location 1 in neuron II, we can see clearly that the selectivity for medium-curvature shapes pointing upward arises from the layout of the fine-scale map; the middle location is tuned to horizontal elements, the upper-left location is tuned to elements tilted 45 degrees counterclockwise, and the upper-right location is tuned to elements tilted 45 degrees clockwise (and also vertical). There is a close correspondence between the data and the predicted patterns both for the full model and the orientation-only model. The space-only model performed less well but still explained significant parts of the response ($r = 0.43$ for the space-only model versus $r = 0.58$ for the orientation-only model). Thus, both spatial and orientation components...
The present results reveal considerable heterogeneity in feature selectivity and the translation invariance of neurons in macaque area V4 and force us to reconsider the established notion that neuronal invariance increases as one traverses the ventral visual hierarchy. Consistent with the conclusions of earlier reports (Pasupathy and Connor, 1999), we find a subpopulation of V4 neurons whose stimulus tuning is maintained throughout the RF. Also consistent with earlier studies, the majority of neurons did exhibit a higher firing rate to the most preferred stimulus tested versus the most nonpreferred stimulus, across spatial locations. However, a detailed mapping of stimulus tuning reveals many neurons exhibiting considerable variability in tuning across space and very limited spatial invariance. This diversity can be captured by two underlying organizing principles.

DISCUSSION

One innovation of the current study is the use of fast reverse correlation procedures to map V4 RFs. Such techniques are common in earlier visual areas (Ringach, 2004), but previous studies in V4 have generally used longer-duration stimuli, typically with durations ranging from 200 to 500 ms and correspondingly long interstimulus intervals. The primary advantage of the fast mapping technique was that it allowed us to perform a dense mapping of shape selectivity across several locations in the RF in addition to a fine-grained mapping of the selectivity to individual oriented components of the composite shapes. This provides a more comprehensive description of contour/shape selectivity across the RF than has been possible in previous studies.

The present results reveal considerable heterogeneity in feature selectivity and the translation invariance of neurons in macaque area V4 and force us to reconsider the established notion that neuronal invariance increases as one traverses the ventral visual hierarchy. Consistent with the conclusions of earlier reports (Pasupathy and Connor, 1999), we find a subpopulation of V4 neurons whose stimulus tuning is maintained throughout the RF. Also consistent with earlier studies, the majority of neurons did exhibit a higher firing rate to the most preferred stimulus tested versus the most nonpreferred stimulus, across spatial locations. However, a detailed mapping of stimulus tuning reveals many neurons exhibiting considerable variability in tuning across space and very limited spatial invariance. This diversity can be captured by two underlying organizing principles.
The first is the dichotomy between the heterogeneity of feature selectivity across RF locations in the case of neurons tuned to higher-curvature/C shapes and its homogeneity in the case of neurons tuned to straight/low-curvature shapes. The denser sampling of the RF afforded by our method reveals that true translation invariance is largely restricted to neurons preferring straight contours. Neurons with preference for very low curvature tend to exhibit spatial invariance, but curvature/C-selective...
neurons often exhibit a high degree of variation in shape preference across their RFs. Further, curvature-tuned neurons tend to prefer curved over straight elements at different locations in the RF while varying in the orientation of the preferred shape across locations (Figures 4B and 4C). These results are echoed by our observations from a separate study where we have observed a trade-off between curvature and invariance using naturalistic images. Thus, we expect that the conclusions of the present study will generalize across different stimulus conditions. This is also supported by the control analyses presented above in which virtually identical tuning was observed when stimuli were presented for longer durations.

There is strong evidence that object recognition is quite rapid as has been demonstrated via rapid serial visual presentation (Potter and Levy, 1969) and rapid object categorizing (Bodeleón et al., 2007; Thorpe et al., 1996) paradigms, suggesting a primary involvement of the feed-forward pathway. Our study focused on neuronal selectivity to individual contour fragments, and the rapid reverse correlation procedure may have mainly isolated feed-forward contributions to the neuronal response. When we compared the shape selectivity among a sample of neurons with fast mapping procedures and longer-duration stimuli, we found striking similarities in their selectivity to the individual elements (Figure S6). It is possible that recurrent or feedback connections, mediated at longer latencies, could refine the selectivity of the initial V4 visual responses and could contribute to spatial invariance as well as to other object-centered or attention-dependent effects (Connor et al., 1996; Pasupathy and Connor, 2001; Yau et al., 2013). Further studies with dense spatiotemporal mapping are needed to fully understand neuronal selectivity to complex combinations of shape fragments.

The second organizing principle alluded to above is that the diversity of shape tuning in V4 is well accounted for by a simple pooling of local orientation signals. Much of the complexity of V4 tuning in our data set could be explained by a linear pooling of local orientation signals. Much of the complexity of the V1 cortical surface (Motter, 2009). Our control experiments presented for longer durations.

The finding that spatial invariance falls off with preference in the RF can be explained by the hypothesis that the RFs of central V4 neurons correspond to a constant-sized sampling of the V1 cortical surface (Motter, 2009). Our control experiments show that these findings are robust against the spatial characteristics of the primitives that made up the curved stimuli.

Previous assessments of spatial invariance were made using the most and least preferred stimuli, either with local curved stimuli (Pasupathy and Connor, 1999) or with larger pattern stimuli (Pasupathy and Connor, 2001), and found consistent selectivity across shifts in position half the RF size or more. Models inspired by these earlier findings utilized linear pooling mechanisms to achieve feature selectivity followed by nonlinearities such as “soft-max” selection to gain spatial invariance (Cadieu et al., 2007). The soft-max operation can be parametrically varied to yield a simple averaging operation at one end (no spatial invariance) to taking the “max” operation on the other (full spatial invariance). Consistent with the earlier studies, we find that both straight- and curve-prefering neurons do preserve a relative preference for the stimuli that are, on average, most and least preferred (Figure 3A, bottom right panels). However, the more detailed examination in our study leads us to conclude first that shift invariance is much more limited than previously appreciated, at least for local curved elements, and, further, that much of the response across the RF is well explained by linear pooling of local orientation responses.

We note that the variation in curvature tuning that we observe is consistent with previous studies using closed form contour stimuli (Carlson et al., 2011; Pasupathy and Connor, 2001) that show selectivity to different convex and concave curves positioned relative to the center of a closed form. However, the limited spatial invariance that we observe and the success of local orientation pooling in predicting responses lead us to suggest that spatial invariance to larger pattern stimuli will be much more restricted than previously suggested, falling within one of our coarse grid locations (about one-third of the RF size). Recent studies at still higher stages of processing such as IT also call into question the spatial extent of invariance in ventral stream representations, suggesting invariance is not intrinsic but is a learned attribute of those representations (Cox and DiCarlo, 2008).

It is possible that the 13 neurons excluded from our analyses due to their lack of shape selectivity are purely color selective (see, e.g., Bushnell et al., 2011). The relationship between the present findings and the recent report of segregated orientation and color domains (Tanigawa et al., 2010) remains to be explored. Since cells selective for higher curvature are not strongly tuned for orientation (Figure 3, example neurons II and III), domain segregation might be somewhat reduced if measured using composite shapes (as in our study). We do not see evidence for the response bias toward acute contour curvature as reported in a recent study (Carlson et al., 2011). This could be due to the fact that in our study we explored the fine structure of the entire RF, whereas the stimuli used in the Carlson et al. study were presented at the center of the RF and typically spanned the extent of the RF.

The finding that spatial invariance falls off with preference for more curved contours suggests a possible segregation of
function. Spatially invariant neurons selective for orientation may play a role in representing extended regions of uniform texture, where the location of the individual texture elements need not be encoded with great spatial precision. In contrast, neurons selective for curvature are likely activated when an appropriately curved contour falls at a particular location within the RF. This form of spatially selective encoding of curved contours would be useful in localizing contours, particularly at the points of high curvature that often play a critical role in defining shape (Attneave, 1954; Feldman and Singh, 2005). We note that such a code, although parsimonious, would be ambiguous for downstream neurons, which would likely integrate multiple signals to derive an unambiguous interpretation of a complex contour.

Although the trade-off between invariance and contour complexity does suggests distinct functions, we also find that V4 responses across this spectrum can be explained using a simple model that pools fine-scale orientation signals. This suggests that differences in invariance and contour complexity depend on differences in the orientation-selective inputs that are pooled to give rise to selectivity in V4. We thus suggest that these different patterns of shape tuning and invariance can be understood as arising from differences in the wiring that these different patterns of shape tuning and invariance are pooled to give rise to selectivity in V4. We thus suggest that such a code, although parsimonious, would be ambiguous for downstream neurons, which would likely integrate multiple signals to derive an unambiguous interpretation of a complex contour.

EXPERIMENTAL PROCEDURES

Electrophysiology

Neurons were recorded in area V4 in two rhesus macaques. Experimental and surgical procedures have been described previously (Reynolds et al., 1999). All procedures were approved by the Salk Institute Institutional Animal Care and Use Committee and conform to NIH guidelines. See Supplemental Experimental Procedures for further details.

Stimulus Presentation and Eye-Movement Monitoring

Stimuli were presented on a computer monitor (Sony Trinitron Multiscan, TC, 640 x 480 pixel resolution, 120 Hz) placed 57 cm from the eye. Eye position was continuously monitored with an infrared eye tracking system (240 Hz, ETL-400; ISCAN). Experimental control was handled by NIMH Cortex software (http://www.cortex.salk.edu). Trials were aborted if eye position deviated more than 1° from fixation.

Preliminary RF Mapping

At the beginning of each recording session, neuronal RFs were mapped to determine the approximate spatial extent over which stimuli elicited a visual response. Monkeys fixated a central point during which each neuron’s RF was mapped using subspace reverse correlation in which Gabor (eight orientations, 80% luminance contrast, spatial frequency 1.2 cpd, Gaussian half-width 2°) or ring stimuli (80% luminance contrast) appeared at 60 Hz. Each stimulus appeared at a random location selected from a 19 x 15 grid with 1° spacing in the inferior right visual field. The orientation of the Gabor stimulus and the color of all stimuli (one of six colors or achromatic) were randomly selected. This resulted in an estimate of the spatial RF, orientation, and color preference of each neuron. Recordings were often made from multiple electrodes, and the preferences of units on separate channels did not always match. The stimuli for the main experiment were centered on the estimated spatial RF of the best-isolated units.

Task and Stimuli

The monkey began each trial by fixating a central point for 200 ms and then maintained fixation through the trial. Each trial lasted 3 s, during which neuronal responses to a fast-reverse correlation sequence (16 ms stimulus duration, exponential distributed delay between stimuli with mean delay of 16 ms, i.e., 0 ms delay p = 1/2, 16 ms delay p = 1/4, 32 ms delay p = 1/8, and so on) were recorded. The stimuli were composed of oriented bars (eight orientations) or bar composites (16 orientations x 5 conjunction angles, total of 72 unique stimuli, Figure 1A). These latter stimuli were constructed from the conjunction of three bars at conjunction angles of 0°, 22.5°, 45°, 67.5°, and 90° between the end elements and the center. The five conjunction levels created five categories of shapes. These were enumerated as 0 (zero curvature), 1 (low curvature), 2 (medium curvature), 3 (high curvature), and 4 (Q). A pseudorandom sequence from the combined stimulus sets was presented in each trial. The composite stimuli were presented on a uniform 5 x 5 location grid (“coarse grid”) centered on the estimated spatial RF based on the preliminary mapping. The grid locations were separated by one-fourth of the RF eccentricity (for example, for a RF centered at 6°, the grid-spacing was 1.5° and the grid covered a visual extent of 3°-9°). The oriented bar stimuli were presented on a finer 15 x 15 location grid (“fine grid”) that spanned the larger 5 x 5 grid in equally spaced increments. Stimuli were scaled by RF eccentricity, such that each single bar element spanned approximately the diagonal length of the fine grid. The RFs of all neurons reported in the study were in the parasfoveal region between 2° and 12° in the inferior right visual field.

Inclusion Criteria

Only well-isolated units were considered as potential candidates (n = 251) for the analysis. Among these, only those neurons with robust visual responses were selected. The robustness of the spatiotemporal response to the visual stimuli was determined as follows: a temporal window between 60 and 120 ms after stimulus onset was used to identify a temporal interval of significant visual response. The temporal window was divided into 8.33 ms bins for determining the peristimulus time histogram (PSTH). Typical average temporal responses to the composite stimuli are shown in Figure 2A. The temporal window for each neuron, $T_{up}$, was taken as those PSTH bins where the mean firing rate averaged across all stimulus conditions exceeded the baseline rate by 4 SDs (significant time points labeled in Figure 2A, dotted gray box). The baseline rate was determined from a temporal window between 0 and 20 ms after stimulus onset. A neuron was considered a potential candidate for further analysis if it had a clear transient response peak that was contained within the larger 60–120 ms interval. A total of 23 single units were eliminated in this process. Of the remaining 228 units, we eliminated a further 135 units since their fine-scale orientation maps (described below) were not fully contained within the stimulus presentation grid (i.e., the spatial extent of the RFs of these neurons could not be fully characterized).

With the remaining 93 units, we next determined the locations within the coarse 5 x 5 stimulus grid where the neuron had significant spatial responses. We first performed a jackknife analysis on the data (Nj = 20 jackknives, each using 95% of trials). For each jackknife, j, we determined the neuronal response, $r(x,y,s,j)$ to a particular composite stimulus, s, at grid location $(x,y)$, as the average firing rate (within $T_{up}$) across stimulus repeats. The mean, $\bar{r}(x,y,s)$, and SEM, $\eta(x,y,s)$, of the responses across the 20 jackknives were then used to calculate a spatial Z score at each grid location $(x,y)$:

$$Z_{spatial}(x,y) = \frac{\bar{r}(x,y,s) - b}{\eta(x,y,s) \sqrt{N_j - 1}}$$

where $b$ was the baseline firing rate and the " operation indicates that the responses were averaged across stimuli. The grid location was marked as significant if the spatial Z score exceeded the significance level of 0.05 (corrected for 25 multiple comparisons; see Figure S1A). Spatially significant grid locations for example neurons are marked with “x” or numerals in Figures 2 and 3.

For each spatially significant grid location, we next determined whether the neuron was significantly selective to the composite stimuli at that location. We calculated a Z score for each stimulus:

$$Z_{stimulus}(x,y,s) = \frac{\bar{r}(x,y,s) - \bar{r}(x,y,s)}{\eta(x,y,s) \sqrt{N_j - 1}}$$

where $\bar{r}(x,y,s)$ is the average firing rate across all stimulus conditions at location $(x,y)$ and $\eta(x,y,s)$ is the SEM for that location. A neuron was considered selective for the stimulus if the Z score was greater than 3.0 for that stimulus. The Z score for each stimulus was then used to calculate the discriminability of the neuron's spatial RF to that stimulus.

The results of this analysis are shown in Figure 2B, where the Z score for each stimulus is plotted as a function of the spatial frequency of the stimulus. The Z score is calculated as the difference in firing rate between the stimulus and the baseline, divided by the SEM for that location. The Z score is then used to determine the discriminability of the neuron's spatial RF to that stimulus.
We define a shape selectivity index, $SSI(x,y)$, for that spatial location as the maximum of the shape 2 scores: $SSI(x,y) = \max_z Z_{\text{shape}}(x,y,z)$.

A grid location was considered significantly shape selective if the index exceeded the significance level of 0.05 (corrected for 72 × M multiple comparisons, where M was the number of significant spatial locations; see Figure S1A). A neuron was considered significantly shape selective if it had at least one spatially significant grid location that was also significantly shape selective. A total of 13 neurons failed this significance test. These neurons had significant spatial RFs, but were not significantly shape selective (Figure 1B). An example of a nonselective neuron is shown in Figure 2 (example neuron IV). We did not analyze these neurons any further. All subsequent analyses were performed on the remaining 80 neurons.

**Data Analysis**

We used the mean responses $\bar{r}(x,y)$ to generate three basic response maps: (1) location-specific response maps for the composite stimuli at each location in the $5 \times 5$ presentation grid (Figures 2B and 3B); (2) average response map, $\bar{r}(s)$, for the composite stimuli by averaging across spatially significant grid locations; and (3) fine-scale orientation-tuning maps using the same procedure as in (1) for the bar stimuli on the 15 × 15 grid (Figure 3C).

For the population analysis, we determined several metrics from the response maps for each neuron:

**Average Shape Preference**

Average shape preference was calculated by first determining the set of composite shapes, $s$, whose firing rate in the average response map, $\bar{r}(s)$, exceeded 90% of the maximum firing rate. The shape category, $c$: (0: straight, 1: low curvature, 2: medium curvature, etc.), corresponding to these shapes was weighted and averaged by their firing rates to determine the average shape preference:

$$\frac{\sum_i r_i(s) c_i}{\sum_i r_i(s)}$$

**Local Shape Preference**

Local shape preference is same as above but derived from the location-specific response maps.

**Local Preferred Shape Orientation**

Local preferred shape orientation is the orientation (0°, 22.5°, 45°, ..., 337.5°) of the local preferred shape defined above. We computed the conditional joint distribution of local shape preference and the angular deviation of preferred shape orientation, $\Delta\phi_{\text{pref}}$ (Figure 4). The computation was conditioned on the shape preference and shape orientation at the maximally responsive location for each neuron. For all spatially significant locations other than the maximally responsive location, $\Delta\phi_{\text{pref}}$ was computed as the absolute value of the angular deviation of the local preferred shape orientation from that of the preferred shape orientation at the maximally responsive location. We divided our neuronal population into three subpopulations: those that preferred straight/low curvature (local shape preference values between 0 and 1, $n = 32$; Figure 4A), those that preferred medium curvature (local shape preference values between 1.5 and 2.5, $n = 16$; Figure 4B), and those that preferred high curvature/C (local shape preference values between 3 and 4, $n = 20$; Figure 4C) at the maximally responsive location. To test whether the marginal distributions of the orientation deviation, $\Delta\phi_{\text{pref}}$, between the straight/low-curvature/prefer- ring units and the high-curvature/C-prefering units (Figures 4A and 4C, right histograms) were significantly different, we calculated the Kullback-Leibler (KL) divergence between the distributions:

$$D_{\text{KL}}(P|Q) = \sum_i P(i) \ln \frac{P(i)}{Q(i)}$$

where $P$ is the marginal distribution in Figure 4A and $Q$ is the marginal distribution in Figure 4C. This yielded a value of 0.5685. We then computed a boot- strapped set (Efron and Tibshirani, 1993) (1,000 iterations) of divergences $D_{\text{KL}}(P\|P_{\text{null}})$ with respect to the null distribution, $P_{\text{null}}$, which was obtained from a random sample (with replacement) of the combined data that underlay the two distributions $P$ and $Q$. Comparing $D_{\text{KL}}(P|Q)$ to this distribution yielded a p value of 0.008, indicating that the two marginal distributions were significantly different. Similarly, the marginal distributions between the straight/low-curvature-prefering units and the medium-curvature-prefering units (Figures 4A and 4B, right histograms) were also significantly different (p = 0.03).

**Pair-Wise Pattern Correlation, $r$ and Pair-Wise Pattern Reliability, $r$**

For any pair of spatially significant coarse grid locations, we estimated the empirical distribution of correlation coefficients between the response patterns (location-specific response maps) at the two locations using a bootstrap procedure (resampling with replacement, 1,000 iterations) (Efron and Tibshirani, 1993). The pairwise correlation $(\rho)$ was taken as the expected value of a Gaussian fit to this empirical distribution (Figure S4). The Gaussian fits were in excellent accord with the raw distributions across our data set. The pairwise pattern reliability, $r$, was defined as $r = 1 - \sigma$, where $\sigma$ was the SD of the Gaussian fit to the empirical distribution (Figure S4). The reliability served as a measure of data quality, with values closer to 1 indicating that the estimates of pattern correlation were more reliable. A scatterplot of pattern correlation versus pattern reliability for all possible location pairs in our neuronal population is shown in Figure 5B. The marginal distributions of pattern correlation (Figure 5B, right histograms) and pattern reliability (Figure 5B, lower histograms) for three subpopulations—points that came from neurons with average shape preference for straight/low curvature (average shape preference values between 0 and 1), those that came from neurons with average shape preference for high curvature (between 1.5 and 2.5), and those that came from neurons with average shape preference for high curvature/C (between 3 and 4)—were tested for statistical difference (using the same procedure described above using the KL divergence measure). The marginal distribution of pattern correlation for the low/straight neurons was significantly different from those of the high-curvature/C-prefering ($p = 0.0001$) and the medium-curvature-prefering neurons ($p = 0.001$). The distributions of pattern reliability were not significantly different from each other, indicating that differences in data quality were not an issue.

To examine the idea that local pooling of orientation signals within subregions of the RF determines the patterns of selectivity to more complex features, we generated predictions of location-specific response maps. This was done by spatially interpolating the fine-scale orientation-tuning map in a three step process: first, the pure spatial information in the fine-scale map, obtained by averaging across orientation at each fine-grid location, was subject to a two-dimensional (2D) nearest-neighbor interpolation (20 interpolation points) followed by a 2D Gaussian smoothing operator ($\sigma = 2/3\times$ the spacing between fine-grid locations); second, the pure orientation information in the map, obtained by subtracting the average orientation response from the measured data at each fine-grid location, was subject to a 2D nearest-neighbor interpolation (20 interpolation points) followed by a 2D Gaussian smoothing operator ($\sigma = 4/3\times$ the spacing between fine-grid locations); finally, the two components were combined by addition. The composite stimuli (at each coarse grid location) were then projected onto this interpolated space. The response to each component element was read off as the value of the closest orientation match in the interpolated space at the location corresponding to the center of the component element. The predicted response to each composite stimulus was taken as the average of the three component responses. We then calculated the correlation coefficient, $r_{\text{model}}$, between the response patterns in the predicted map and the observed map. Since we were only concerned with pattern selectivity and not with rate matching, the correlation measure was sufficient for our purpose.

To test for the predictive power of the model, we also calculated a null distribution of the correlation coefficients. This was done by spatially shuffling the fine tuning curves of the fine-scale orientation map within a 3 x 3 fine grid that underlay a coarse grid location (see Figure S5A), generating the predicted responses from this shuffled map (same procedure as above for the original unshuffled map) and hence the correlation coefficient between the predicted map and the observed map. This shuffling procedure preserved the orientation tuning at the fine scale while perturbing the relative spatial structure of the map within a coarse grid location. This procedure was repeated 1,000 times to give an estimate of the null distribution ($\rho_{\text{null-model}}$; see Figure S5B). The model correlation, $r_{\text{model}}$, was tested against the null distribution for significance (p = 0.05, Bootferroni corrected for M multiple comparison, where M is the number of significant spatial locations for each neuron). The model
was considered to have significant predictive power for a neuron if there was at least one spatial location that was significant, according to the above criteria. We also investigated two reduced versions of the pooling model (Figure S5C). The space-only version was obtained by averaging across orientation at each fine-grid location (Figure S5C, right upper panel). This model did not have any local orientation tuning. The orientation-only version was obtained by subtracting the average orientation response (as in the space-only model) from the measured data at each fine-grid location (Figure S5C, right lower panel). Thus, this model did not contain any local spatial information. The model correlations and null distributions for these reduced models were calculated using the same procedure described above for the full model.

The explained variance of our model was estimated by first calculating the model correlation, \( r_{\text{model}} \), as above, but on different jackknifed fractions of the data. Specifically, we calculated \( r_{\text{model}} \) between the predicted response map and the observed response maps from (1) the full data set, (2) 95% of trials (3) 90% of trials, (4) 85% of trials and (5) 80% of trials. We then performed a linear regression on the resulting \( r_{\text{model}} \) values against the reciprocal of the corresponding jackknife fraction values (1, 1/0.95, 1/0.9, 1/0.85, and 1/0.8). This procedure is designed to correct for the bias due to finite data set size (Brenner et al., 2000; Sahani and Linden, 2003; Machens et al., 2004). The square of the y-intercept of the regression line was taken as the explainable variance for that RF location. The explained variances of the reduced space-only and orientation-only models were calculated using the same procedure.

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2013.04.016.

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