Behavioral/Cognitive

A Texture Statistics Encoding Model Reveals Hierarchical Feature Selectivity across Human Visual Cortex

¹⁰Margaret M. Henderson,^{1,2,3} Michael J. Tarr,^{1,2,3} and Leila Wehbe^{1,2,3}

¹Neuroscience Institute, Carnegie Mellon University, Pittsburgh, Pennsylvania 15213, ²Department of Psychology, and ³Machine Learning Department, Carnegie Mellon University, Pittsburgh, Pennsylvania 15213

Midlevel features, such as contour and texture, provide a computational link between low- and high-level visual representations. Although the nature of midlevel representations in the brain is not fully understood, past work has suggested a texture statistics model, called the P–S model (Portilla and Simoncelli, 2000), is a candidate for predicting neural responses in areas V1–V4 as well as human behavioral data. However, it is not currently known how well this model accounts for the responses of higher visual cortex to natural scene images. To examine this, we constructed single-voxel encoding models based on P–S statistics and fit the models to fMRI data from human subjects (both sexes) from the Natural Scenes Dataset (Allen et al., 2022). We demonstrate that the texture statistics encoding model can predict the held-out responses of individual voxels in early retinotopic areas and higher-level category-selective areas. The ability of the model to reliably predict signal in higher visual cortex suggests that the representation of texture statistics features is widespread throughout the brain. Furthermore, using variance partitioning analyses, we identify which features are most uniquely predictive of brain responses and show that the contributions of higher-order texture features increase from early areas to higher areas on the ventral and lateral surfaces. We also demonstrate that patterns of sensitivity to texture statistics can be used to recover broad organizational axes within visual cortex, including dimensions that capture semantic image content. These results provide a key step forward in characterizing how midlevel feature representations emerge hierarchically across the visual system.

Key words: encoding model; fMRI; midlevel features; spectral features; texture statistics; vision

Significance Statement

Intermediate visual features, like texture, play an important role in cortical computations and may contribute to tasks like object and scene recognition. Here, we used a texture model proposed in past work to construct encoding models that predict the responses of neural populations in human visual cortex (measured with fMRI) to natural scene stimuli. We show that responses of neural populations at multiple levels of the visual system can be predicted by this model, and that the model is able to reveal an increase in the complexity of feature representations from early retinotopic cortex to higher areas of ventral and lateral visual cortex. These results support the idea that texture-like representations may play a broad underlying role in visual processing.

Introduction

Information in visual cortex is processed by a series of hierarchically organized brain regions, with the complexity of representations increasing at each level. Although there are intuitive explanations for response properties at the ends of this hierarchy (e.g., oriented spatial frequency filters in primary visual cortex; Hubel and Wiesel, 1962; Carandini et al., 2005) or object and category representations in inferotemporal cortex (Desimone et al., 1984; Grill-Spector and Weiner, 2014), the computations performed at intermediate levels have proven more challenging to describe. These intermediate or midlevel visual areas are thought to represent features like contour and texture, which play an important role in figure-ground segmentation, shape processing, and object and scene classification (Bergen and Landy, 1991; Ullman et al., 2002; Connor et al., 2007; Walther and Shen, 2014; Peirce, 2015). Thus, developing a robust model of midlevel representation is fundamental for understanding how the visual system extracts meaningful information from the environment.

Computational texture models, particularly the influential model proposed by Portilla and Simoncelli (2000), hereafter referred to as the P -S model, have proven useful in understanding the midlevel features that drive visual responses. The P–S

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Correspondence should be addressed to Margaret M. Henderson at mmhender@cmu.edu. https://doi.org/10.1523/JNEUROSCI.1822-22.2023

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Figure 1. *A*, Overview of the texture statistics model (Portilla and Simoncelli, 2000) and voxelwise encoding model fitting procedure. For each image, we first use a steerable pyramid (Simoncelli and Freeman, 1995) to decompose the image into four orientation and four frequency bands (note that only 2 orientations and frequencies are shown here for illustration). We then use the steerable pyramid representation to extract lower-level texture features, which include marginal statistics of the pyramid coefficients, and higher-level texture features, which consist of higher-order correlations of the pyramid coefficients (see below, Materials and Methods). Each of these features is computed for each pRF in a grid of candidate pRFs, using the pRF as a spatial weighting matrix. Before fitting the model, we reduce the dimensionality of the higher-level texture features using PCA. We then use ridge regression to fit a set of weights for each voxel that predicts its response to each image as a weighted sum of all the texture statistics features extracted from that image. We perform this fitting separately for each candidate pRF and use the loss on a held-out (nested) data subset to choose the best pRF. Finally, we compute the accuracy of the model (R^2) on a held-out validation set (see below, Materials and Methods). **B**–**E**, Illustrations of how the higher-level model features are computed based on coefficients of the steerable pyramid for one example pRF. The image is cropped (**B**) to yield an image patch spanning $\pm 2 \sigma$ from the pRF center; the pRF is cropped in the same way and used as a weighting matrix. The energy-auto features (**C**) are computed by correlating each magnitude feature maps corresponding to the same scale but different orientation (**D**). The energy-cross-scale features are computed by cross-correlating magnitude feature maps of steerable pyramid coefficients. Extended Data Table 1-1 shows the total number of features included in each subset.

model is constructed using a steerable pyramid decomposition (Simoncelli and Freeman, 1995) to extract features at different spatial scales and orientations and includes features at multiple levels of complexity, including simple luminance and spectral statistics, as well as higher-order correlation statistics (Fig. 1). The P–S model can account for aspects of texture sensitivity in primate visual areas such as V1, V2, and V4 (Freeman et al., 2013; Okazawa et al., 2015, 2017; Hatanaka et al., 2022), with sensitivity to the higher-order model features increasing from V1 to V2 to V4 (Freeman et al., 2013; Okazawa et al., 2013; Okaza

Hatanaka et al., 2022). In humans, functional magnetic resonance imaging (fMRI) studies suggest that P–S statistics can account for some response properties in early visual and ventral visual cortex (Baumgartner and Gegenfurtner, 2016; Long et al., 2016). The P–S model also captures key aspects of human behavior, including perceptual discrimination of texture patches (Balas, 2006; Freeman and Simoncelli, 2011), crowding (Balas et al., 2009), search performance (Rosenholtz et al., 2012), and discrimination of high-level object properties such as real-world size and animacy (Long et al., 2016, 2017).

Despite these promising results, it is not yet known how well the P-S model captures neural responses to natural scene images, particularly in higher visual cortex. The majority of past studies have used synthetic stimuli or textures that are relatively homogeneous across space (Freeman and Simoncelli, 2011; Okazawa et al., 2015, 2017). In contrast, natural scenes, which include objects and other complex spatially localized elements, may prove more challenging to characterize (Portilla and Simoncelli, 2000). One recent study did examine the ability of the P-S model to predict neural responses in macaque V1 and V4 for naturalistic stimuli (Hatanaka et al., 2022); however, this study did not consider higher visual cortex. Under the hypothesis that midlevel features also play a key role in the neural processing of object and scene categories (Connor et al., 2007; Nasr et al., 2014; Bracci et al., 2017; Groen et al., 2017; Long et al., 2018), modeling responses to natural images within higher visual cortex is a critical test of the ability of the P-S model to generalize to real-world vision.

To address this question, we developed a forward encoding model (Serences and Saproo, 2012; Naselaris and Kay, 2015) that uses P-S model texture statistics to predict voxelwise responses from the Natural Scenes Dataset (NSD; Allen et al., 2022) and used variance partitioning to probe the contributions made by each subset of texture model features. Critically, our model (1) achieves accurate predictions of held-out voxel responses throughout the visual hierarchy, with highest performance in early and midlevel areas and (2) recovers the hierarchical organization of visual cortex, with retinotopic areas best described by lower-level features and anterior category-selective areas best described by higher-level features. These and other results we present here facilitate a better understanding of how hierarchical computations give rise to midlevel representations in the brain and how midlevel feature selectivity may be used to predict the emergence of a higher-level semantic representational space.

Materials and Methods

Acquisition and preprocessing of fMRI data

We used the NSD, a large-scale, publicly available fMRI dataset. Full details on the acquisition of this data can be found in Allen et al. (2022). The NSD includes whole-brain BOLD fMRI measurements from eight human subjects (both male and female) who each viewed between 9000 and 10,000 natural scene images over the course of a year (between 30 and 40 scan sessions). Functional scanning was conducted at 7T using whole-brain gradient-echo EPI at 1.8 mm resolution and 1.6 s repetition time (TR). Images were sampled from the Microsoft Common Objects in Context (COCO) database (Lin et al., 2014). Over the course of the experiment, each image was viewed $\sim 3 \times$, for a total of $\sim 30,000$ trials per subject (fewer for some subjects who did not complete the entire experiment). Of the \sim 10,000 images viewed by each subject, \sim 9000 images were seen only by that subject, and 907 were viewed at least once by each subject. Each image was presented in color, at a size of 8.4° \times 8.4° (of visual angle), and was viewed for a duration of 3 s, with 1 s between trials. Throughout each scan, subjects performed a task in which they reported whether each image was new or old (i.e., whether it had been presented before in any session), while fixating centrally on a small fixation dot superimposed on each image.

As described in Allen et al. (2022), the functional data from all sessions were preprocessed using temporal interpolation to correct for slice timing differences and spatial interpolation to correct for head motion (Allen et al., 2022). All analyses were performed in each subject's native volumetric space (1.8 mm resolution voxels). A general linear model was used to estimate beta weights for each voxel and each individual trial (Prince et al., 2022). We obtained the beta weights from Allen et al. (2022) after this stage, and we then performed a few additional steps to prepare the beta weights for our analyses. First, beta weights for each voxel were z-scored across all trials of each scan session. To improve the signal-to-noise ratio of the data, we then averaged the beta weights for each voxel across trials where the same image was shown (approximately three trials/image), resulting in a single value for each voxel in response to each of the unique (\sim 10,000) images. Note that for subjects who did not complete the entire experiment, there were fewer than 10,000 images.

We masked out a broad region of interest (ROI) in visual cortex to include in all analyses. This region included voxels that were part of the nsdgeneral ROI described in Allen et al. (2022), which is meant to capture the general spatial extent of voxels that were responsive to the NSD image stimuli. To broaden the scope of brain areas included in our analyses, we additionally included voxels belonging to any ROI within several sets of ROI definitions, that is, any ROI in the probabilistic atlas provided in Wang et al. (2015), which includes regions of the intraparietal sulcus; any ROI belonging to an early retinotopic area based on population receptive field (pRF) mapping; and any face-selective, body-selective, or place-selective ROI identified through a functional category localizer task (Allen et al., 2022, pRF and category localizer tasks). We further thresholded voxels for inclusion according to their noise ceiling, which is a measure that captures the proportion of the response variance of voxels that can theoretically be explained by properties of the stimulus (Wu et al., 2006; Allen et al., 2022), using a threshold of 0.01. This large mask defines the extent of voxels that are included in whole-brain surface maps (see Fig. 3A). We also computed summary statistics at the ROI level using more fine-grained definitions of individual ROIs (see Fig. 3B; see below, Defining ROIs).

Defining ROIs

We defined retinotopic and category-selective ROIs based on functional localizers that were collected as part of the NSD experiment. A category localizer task (Stigliani et al., 2015) was used to define place-selective regions [parahippocampal place area (PPA), occipital place area (OPA), and retrosplenial cortex (RSC)], a face-selective region [fusiform face area (FFA); we combined FFA-1 and FFA-2 into a single FFA region], and a body-selective region [extrastriate body area (EBA)]. A pRF mapping task (sweeping bar stimuli; Benson et al., 2018) was used to define early retinotopic visual ROIs V1, V2, V3, and hV4 (Allen et al., 2022). When retinotopic and category-selective ROIs were overlapping with one another, we excluded any voxels that were overlapping from the retinotopic definition and added them to the corresponding category-selective ROI only. To ensure that the final set of category-selective ROIs were nonoverlapping, we always prioritized face-selective ROIs over place- and body-selective ROIs, and prioritized place-selective ROIs over body-selective ROIs.

Texture statistics encoding model

Overview. Our encoding model incorporated parameters for the feature selectivity of each voxel as well as its spatial selectivity (St-Yves and Naselaris, 2018; Fig. 1). The inclusion of an explicit pRF for each voxel is an aspect in which our model improves on past work (Okazawa et al., 2015; Hatanaka et al., 2022), allowing us to adaptively fit voxel responses with a wide range of receptive field positions and sizes. During model fitting, both the pRF parameters and the texture feature weights are optimized simultaneously (see below, Model fitting procedure). To fit the model, we first created a grid of candidate pRFs and extracted texture statistics features within each candidate pRF. For each pRF, we then used regularized regression to fit a linear model that predicts the activation of the voxel as a function of the texture statistics features corresponding to that pRF. The final encoding model for each voxel was computed by identifying the best-fitting model over all candidate pRFs. More detail on each step is provided in the following sections.

pRFs. We modeled each candidate pRF as a two-dimensional Gaussian (Dumoulin and Wandell, 2008; St-Yves and Naselaris, 2018). Each pRF has three parameters, x_0 , y_0 , and σ , where x_0 , y_0 and σ , respectively, indicate the center and SD of the two-dimensional Gaussian response profile as follows:

$$g_{x_0,y_0,\sigma}(x,y) = \exp\left(-\left(\frac{(x-x_0)^2}{2\sigma^2} + \frac{(y-y_0)^2}{2\sigma^2}\right)\right).$$
 (1)

Our grid of candidate pRF parameters was designed to approximate a log-polar grid, where the spacing between adjacent pRF centers is linear in terms of polar angle position (θ) and nonlinear in terms of eccentricity (r), such that candidate centers were more closely spaced closer to the center of the visual field. This nonlinear eccentricity spacing was intended to account for the cortical magnification factor in human visual cortex, where the neuronal sampling of visual space is denser close to the fovea (Duncan and Boynton, 2003). Specifically, we used 16 candidate polar angle positions that were linearly spaced, ranging from 0° to 337.5° in steps of 22.5° and 10 candidate eccentricities that were logarithmically spaced, ranging from 0° to 7°. For each of these candidate centers, we generated 10 candidate σ values that were also spaced logarithmically, ranging from 0.17° to 8.4°. The complete grid of pRF parameters over all combinations of r, θ , and σ resulted in 1600 pRFs.

Because our eccentricity range extended slightly beyond the physical image extent (an 8.4° square), some of the smaller pRFs at the largest r values were entirely nonoverlapping with the image region (in contrast, larger pRFs at these large eccentricities were still partially overlapping with the image region). To address this issue, we removed from the grid any pRFs whose rough spatial extent (center $\pm \sigma$) was nonoverlapping with the image region. This resulted in 1456 pRFs in the final grid.

Texture statistics features. As the first step of our voxelwise encoding model framework, we extracted a set of image-computable texture features that are meant to capture various aspects of local image structure (Figs. 1, 2). We used the aforementioned P–S model (Portilla and Simoncelli, 2000), which has previously been shown to predict both human behavioral judgments of textures and neural responses in midlevel visual regions such as V2 and V4 (Portilla and Simoncelli, 2000; Freeman and Simoncelli, 2011; Freeman et al., 2013; Okazawa et al., 2015). Full details of the model construction and its motivation are given in Portilla and Simoncelli (2000) and Freeman and Simoncelli (2011). Here, we provide a brief description of the model components and describe how we have incorporated it into our pRF modeling framework.

The first stage of the model consists of using a steerable pyramid (Simoncelli and Freeman, 1995) to decompose each image into a set of orientation and frequency sub-bands (We used four orientations and four frequency bands.). This step was implemented using the Python package pyrtools to construct a steerable pyramid in the frequency domain. Before processing images, we converted images to grayscale using the International Telecommunication Union Radio Communication Sector BT.709-2 standard, which consists of multiplying the RGB (red, green, blue) channels by [0.2126, 0.7152, 0.0722] and taking their sum, and resampled images to a resolution of 240 \times 240 pixels using bilinear resampling. The steerable pyramid results in spatial maps of complexvalued coefficients at each scale and orientation from which we can compute the real part and the magnitude, which correspond approximately to the responses of V1 simple and complex cells, respectively (Freeman and Simoncelli, 2011). The four frequency bands output by our pyramid were centered at \sim 0.9, 1.8, 3.6, and 7.1 cycles per degree (cpd), and the four orientation bands were centered at 0°, 45°, 90°, and 135°. The steerable pyramid additionally computes high-pass and low-pass residual images, as well as a partially reconstructed low-pass image representation at each scale.

Our complete texture statistics model included 10 total subsets of features, resulting in 641 total features (Extended Data Table 1-1), each of which we computed at each pRF grid position. For simplicity, we have divided the features into two subgroups for several of our analyses (see Fig. 5). The first subgroup, which we have termed "lower-level texture features," consists of marginal statistics (such as mean and variance) computed from either the raw image luminance values or from the outputs of the steerable pyramid. The second set of features, which we have termed "higher-level texture features," consists of higher-order correlations computed from the steerable pyramid, generated by either correlating different orientation/scale channels of the pyramid or spatially shifted versions of the same channel of the pyramid. By virtue of these cross-correlations and autocorrelations, the higher-order features are able to capture a higher degree of complexity than the lower-level model features, exhibiting sensitivity to properties like periodic, spatially repeating structure, and junctions made by contours of different orientations (Fig. 2, compare A-D and E-J). When computing each of these features, we used the Gaussian profile for the pRF of interest (Eq. 1) as a weighting matrix (similar to the pooling region used in Freeman and Simoncelli, 2011). All feature extraction steps after the initial steerable pyramid computation were done using custom code in the PyTorch package in Python.

The first set of lower-level features, termed pixel features (six features), consists of the pRF-weighted minimum, maximum, mean, variance, skew, and kurtosis of the raw pixel luminance values. The second and third sets of features, energy-mean (16 features) and linear-mean (16 features) features, are the pRF-weighted mean of the magnitude and real part, respectively, of each steerable pyramid feature channel (four orientations times four scales). The fourth set of lower-level features, marginal features (11 features), includes the pRF-weighted skew and kurtosis of the low-pass pyramid reconstruction at each scale and the pRF-weighted variance of the high-pass residual image.

The first set of higher-level texture features consists of autocorrelations of the steerable pyramid features, which are computed by correlating each feature map with spatially shifted versions of itself, allowing these features to capture the repetition of similar elements across spatial positions. Autocorrelations were computed from the magnitude of each main pyramid feature channel (energy-auto; 272 features), the low-pass reconstruction at each scale, and the high-pass residual image. The autocorrelations of the low-pass reconstructions and the high-pass residual image were treated as a single combined group of features for the variance partitioning analysis (linear-auto; 98 features), but were treated separately when performing principal component analysis (PCA; see below) as this yielded higher overall model accuracy. To compute each autocorrelation matrix, we cropped out a square region of the image that spanned approximately $\pm 2 \sigma$ from the pRF center. We then computed the weighted two-dimensional autocorrelation over this cropped image region, using the pRF profile (cropped in the same way) as a weighting matrix. From this matrix, we retained a fixed number of pixels from the center. The number of pixels (i.e., spatial shifts) retained was adjusted based on the scale of the feature map under consideration; the total number of pixels retained ranged from three pixels (shifts of ± 1 pixel) for the lowest frequency maps, to seven pixels (shifts of up to ± 3 pixels) for the highest frequency maps. As the autocorrelation matrix is diagonally symmetric, we retained only the unique values. The total number of autocorrelation features returned was independent of pRF size.

The remaining higher-level texture features consist of cross-correlations, computed by correlating different feature maps output by the steerable pyramid. All cross-correlations were computed using the entire image, weighted by the pRF profile. The first two subsets of cross-correlation features, energy-cross-orient (24 features) and linear-cross-orient (34 features) are cross-correlations of feature maps (either magnitudes or real parts) at the same scale but with different orientations; these features can thus capture image elements that include multiple orientations, such as crosses and curved lines. The next subsets of features (energycross-scale; 48 features) were computed by correlating the magnitude of feature maps having the same orientation but different scales, after upsampling the resolution of the map at the coarser scale and doubling its phase. These cross-scale comparisons are able to capture distinctions between different types of oriented elements in the image, such as object edges versus lines versus gradients. An additional subset of features (linear-cross-scale; 116 features) was computed similarly but using the real or imaginary component of the feature maps only. An additional two subsets of features were computed by correlating the low-pass residual image with spatially shifted versions of itself (within-scale) or with the lowest frequency pyramid feature map (cross-scale). The within-scale group of these features were included in the linear-cross-orient group for the variance partition analysis, whereas the cross-scale group was included in the linear-cross-scale group for the variance partition analysis, but these groups were treated separately when performing PCA (see next paragraph), as this grouping tended to result in better model performance.

To reduce the dimensionality of the texture model feature space and prevent overfitting, we performed PCA on the higher-level texture features before using them to construct encoding models. PCA was performed within each pRF separately, within each subset of higher-level



Figure 2. Visualization of the kinds of natural image features that are captured by each subset of P–S statistics, shown for one example pRF (pRF size, 1.48°). Each outlined box (*A*–*J*) corresponds to one feature subset. Names shown in blue (*A*–*D*) indicate lower-level features, names in red (*E*–*J*) indicate higher-level features. To generate this visualization, we performed PCA on the model features corresponding to each feature subset and identified the two most activating (left of arrows) and least activating (right of arrows) images for each of the first two principal components. To aid visualization of the image area that most strongly contributed to computing the features, we weighted each image according to the Gaussian profile of the pRF and cropped the image to a square centered on the pRF. The white circle indicates a radius of $\pm 2 \sigma$ from the pRF center. These visualizations are generated using only the images shown to S1, but similar results are obtained when using other sets of images. Note that some of the PCs appear to capture an empty white image patch at one end; these homogeneous images likely indicate that the first PC captures the mean value across all features of a given P–S model subset, which may roughly covary with image contrast.

texture features individually (Note that some feature groups were further subdivided when performing PCA; see above). No dimensionality reduction was performed on the lower-level texture features. Where PCA was used, the principal components (PCs) were always computed using the training data only for one subject at a time, and all data for that subject including the validation data were projected into the same subspace. We retained the minimum number of components necessary to explain 95% of the variance. Because PCA was performed on the features from one pRF at a time, this meant that the dimensionality of the features was not required to be the same across all pRFs.

Model fitting procedure. To construct the texture statistics encoding model, we modeled each voxel response as a weighted sum of the texture statistics features corresponding to each image and each pRF (plus an intercept). As described in the previous section, texture statistics features were computed in a spatially specific manner such that the feature activations for a given image depend on the pRF parameters x_0 , y_0 and σ . We solved for the weights of the encoding model for each voxel using

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Figure 3. The texture statistics encoding model accurately predicts voxel responses across visual cortex, with highest accuracy in early and midlevel areas. *A*, Model performance is shown on an inflated cortical surface (left) or flattened surface (right) for one example NSD subject, 51. For the full set of 8 subjects, see: http://www.cs.cmu.edu/~mmhender/viewers/texturemodel/fullmodel_R2/. R^2 was computed using responses to an independent set of images that were not used to fit the model (see above, Materials and Methods). For visualization purposes, values of R^2 have been transformed to $\sqrt{R^2}$, as this makes it easier to visually observe differences between color map values. White outlines and labels indicate the location of functionally defined retinotopic and category-selective ROIs in each subject. *B*, Summary of R^2 for our texture statistics model alongside several alternative low-level visual models (see above, Materials and Methods). Bar heights and error bars indicate mean ± 1 SEM across subjects, light gray dots represent individual subject. *C*, R^2 for the texture statistics model is plotted versus R^2 for AlexNet Conv1 for four selected ROIs. Each gray dot represents a single voxel, dots in shades of green indicate the mean R^2 for the texture model was significantly higher than chance, evaluated using a permutation test. Dots in shades of green indicate single subjects. Bar heights and error bars represent the mean ± 1 SEM across subjects.

ridge regression (L2-regularization), as used in previous work (Güçlü and van Gerven, 2014; Wehbe et al., 2014; Huth et al., 2016). To improve the regression fits, we used a banded ridge regression method (Nunez-Elizalde et al., 2019), which incorporated two different ridge regularization (λ) parameters, one corresponding to the lower-level texture features and one corresponding to the higher-level texture features. Both λ parameters were selected on a per-voxel basis from a set of 10 candidate λ values logarithmically spaced between 0 and 10⁵. To sample all combinations of two λ values, we created a grid of all 100 possible λ combinations. Cross-validation was used to determine the best λ parameters for each voxel; the full cross-validation procedure is as follows (Fig. 1).

First, we held out a validation set of \sim 1000 images from the \sim 10,000 total images for each subject. This set of images always consisted of the shared images in which every subject saw the same images. (Note that for subjects who did not complete the entire NSD experiment, there

were fewer images in this validation set; the minimum was 907 images; see above, Acquisition and preprocessing of fMRI data.) The remaining \sim 9000 images made up the training data. Following this, we held out a random 10% of the training data as a nested validation set. This nested set was used to select the ridge parameter as well as the best pRF parameters for each voxel. Using the remaining 90% of the training data, we then estimated regression weights for each of our candidate λ values, as well as for each of our candidate pRF models. Based on the estimated weights, we computed the loss for each λ and each pRF by generating a prediction of the nested validation data and computing the sum of the squared error for this prediction. We then selected the best pRF parameters and λ values for each voxel based on which values resulted in the lowest loss. The resulting pRF parameters and regression weights made up the final encoding model for the voxel. Finally, to estimate overall model accuracy, we generated predicted responses of each voxel on the held-out validation set and computed the coefficient of determination (R^2) between the actual and predicted response. To make plots of R^2 in surface space for each subject (Fig. 3*A*), we used the PyCortex package (Gao et al., 2015).

Permutation testing. For each voxel, we used a permutation test to evaluate whether the texture statistics encoding model resulted in higher than chance accuracy at predicting the validation set data. To perform the permutation test, we randomly shuffled (1000 times) the image labels corresponding to each voxel activation pattern (voxel activations were already averaged over presentations of each repeated image before this shuffling was performed; see above, Acquisition and preprocessing of fMRI data). Shuffling was always performed within the training set, validation set, and nested validation set (see above, Model fitting procedure) separately. For each shuffling iteration, we performed the regression procedure from scratch, including fitting the model weights and computing R^2 . However we did not refit the pRF of each voxel on the shuffled data; instead, we used the best pRF for each voxel as determined from the intact data and only refit the regression weights. To compute a one-tailed *p* value for each individual voxel, we calculated the number of iterations on which the shuffled R^2 was greater than or equal to the real R^2 , divided by the number of iterations. We then performed false discovery rate (FDR) correction on the p values for all voxels from each subject (q =0.01; Benjamini and Hochberg, 1995). We used the result of this significance test as a mask to determine which voxels to include in the subsequent variance partitioning analyses.

Variance partitioning. To determine the unique contribution of different subsets of texture features to encoding model predictive accuracy, we used a variance partitioning analysis (Groen et al., 2012; Lescroart and Gallant, 2019, for similar approaches). When performing this analysis, we always restricted the set of voxels to those that had above-chance accuracy for the full texture statistics encoding model (see previous section). The overall approach was as follows. First, we fit the full texture statistics encoding model with all feature subsets concatenated, using the method described in the previous section. We computed the prediction accuracy of the concatenated model, R_{full}^2 . Then we progressively removed one set of features at a time from the full model and fit the weights of each partial model. When constructing the partial models, we always used the features corresponding to the best pRF of the voxel (as determined based on fitting the full concatenated feature space). This ensured that the differences between models were solely because of differences in features and not to changes in the estimated pRF parameters. Then, for each partial model, we generated a prediction of the validation set data and computed R^2 for the partial model, $R^2_{partial}$. Finally the unique variance attributable to each feature subset was computed as follows:

$$R_{unique}^2 = R_{full}^2 - R_{partial}^2, \tag{2}$$

where $R_{partial}^2$ denotes the partial model where the feature set of interest was removed. We performed two versions of the variance partitioning analysis. In the first version, we grouped all the lower-level or all the higher-level texture features together (see Fig. 5); for the second version, we analyzed each of the 10 feature subsets individually (see Figs. 7, 8). For the first analysis, we additionally report the variance that is shared between the lower- and higher-level feature subsets. The shared variance between two feature subsets *A* and *B*, was computed as follows:

$$R_{sharedAB}^2 = R_{partialA}^2 + R_{partialB}^2 - R_{full}^2.$$
 (3)

To evaluate the significance of the unique variance explained by each feature subset, we used a bootstrapping analysis consisting of resampling with replacement. To make this analysis computationally feasible, we performed bootstrapping only on the validation set data (i.e., when computing R^2 but not when fitting model weights). We performed 1000 iterations of the bootstrapping test; on each iteration we resampled with replacement *n* images from the total *n* validation set images (where *n* is typically 1000, but could be fewer for the subjects who did not complete all sessions; see above, Acquisition and preprocessing of fMRI data). We

used the same resampling order for the image labels and for the voxel data so that the correspondence between images and voxel responses was intact, but the exact set of images included differed on each bootstrap iteration. Using the same resampling order, we computed R^2 for the full model and R^2 for each partial model, then used these to compute R^2_{unique} for each feature set, as described above. This resulted in a distribution of 1000 values for R^2_{unique} . We then used this distribution to compute a *p* value for whether R^2_{unique} was significantly higher than zero, by computing the number of iterations on which $R^2_{unique} \leq 0$ and dividing by the number of iterations. The *p* values for each subject were FDR corrected across all voxels (q = 0.01; Benjamini and Hochberg, 1995).

We additionally computed whether the unique variance for each feature set was significant at the ROI-averaged level for each subject. This was done by averaging the bootstrap distributions for R_{unique}^2 across all voxels in each ROI (always averaging values for the same bootstrapping iteration together) and then computing a one-tailed p value using the same method described above. This resulted in a single value for each ROI, each subject, and each feature set. We then performed multiple comparisons correction across all values using the Holm-Bonferroni method (q = 0.01; Holm, 1979). We used the Holm-Bonferroni method because the FDR procedure is not appropriate for controlling for a relatively small number of comparisons. The number of subjects with significant unique variance explained for each feature set is reported in Extended Data Tables 5-1 and 8-1. To test whether the average unique variance across all subjects was significant, we used the same averaging method just described but additionally averaged the bootstrap distributions over subjects before computing *p* values and corrected for multiple comparisons as described above.

To compute whether the unique variance explained by the lowerand higher-level texture features was significantly different for each ROI, we used the ROI-averaged bootstrap distributions to compute the distribution of differences between the lower-level unique variance and the higher-level unique variance. We then computed a two-tailed *p* value by computing the proportion of iterations for which the difference was positive and the proportion for which the difference was negative, taking the minimum and multiplying by two. We again performed multiple comparisons correction using the Holm–Bonferroni method (*q* = 0.01).

Fitting with fixed pRF size parameter. To dissociate the effects of pRF size and feature selectivity, we performed an additional analysis in which the pRF size parameter (σ) was fixed at a single value for all voxels. To achieve this, we performed our entire model fitting pipeline from scratch (see above, Model fitting procedure) with a restricted grid of candidate pRFs. This restricted pRF grid consisted of all the pRFs in our main grid having the σ value of interest. Thus, the voxels could be fit with different pRF centers but had to have the same pRF size. We performed this entire procedure for each of the 10 σ values in our original pRF grid.

Alternative low-level visual models

To assess how the accuracy of the texture statistics model compared to that of other models, we implemented several commonly used models of low-level visual features, a GIST model (Oliva and Torralba, 2001), a complex Gabor model (St-Yves and Naselaris, 2018; Henderson et al., 2023), and the first two layers of a pretrained AlexNet convolutional neural network model (Krizhevsky, 2014). The GIST model consists of spectral features (orientation and spatial frequency) that are coarsely localized in space and was implemented using MATLAB code provided by Oliva and Torralba (2001). We evaluated the GIST model with two levels of spatial resolution, a 2 \times 2 grid and a 4 \times 4 grid. To make the GIST model as comparable as possible to our texture statistics model, we used four orientation channels and four frequency channels, which match the number of channels included in the steerable pyramid. Similarly, when implementing the Gabor model, we also used four orientations and four spatial frequencies (0.36, 1.03, 2.97, and 8.57 cpd). Each Gabor model feature was computed by filtering the image with two sinusoids that were 90° out of phase, squaring the output of these two filters, summing the two outputs, and taking the square root. Henderson et al. (2023) has details on construction of a similar model. To extract features from the AlexNet model, we used the pretrained model weights for

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AlexNet available from the PyTorch model zoo, and we extracted activations from layers Conv1 and Conv2 (following the rectifying nonlinear activation function). More details on AlexNet can be found in Krizhevsky (2014). But briefly, it is a convolutional neural network model trained to perform 1000-way object classification. Thus in contrast to the Gabor, GIST, and texture statistics models, which each consist of hand-designed feature weights, the AlexNet model is task optimized and may include object-specific features.

For both the Gabor and AlexNet models, we incorporated spatial pRF parameters into the construction of the model, similar to how the pRF was incorporated into the texture statistics model. Essentially, this procedure consists of extracting features in each pRF of the grid by taking a dot product of the relevant feature maps with each pRF (Henderson et al., 2023; St-Yves and Naselaris, 2018, for similar approaches). To simplify the fitting procedure, as well as making the models more comparable to the texture statistics encoding model, we used the same pRF parameters that had already been estimated using the texture statistics encoding model (see above, Model fitting procedure). This meant that when fitting the Gabor and AlexNet models, we only had to fit the feature weights (using the set of features that had been extracted from the best pRF of each voxel). For the GIST model, the pRF parameters were not incorporated into the fitting procedure; instead, the GIST features included information from the entire image.

Data availability

All code required to reproduce our analyses is available at https://github.com/mmhenderson/texturemodel.

Results

To model neural selectivity for midlevel image structure, we constructed a forward encoding model based on image-computable texture statistics features (Figs. 1, 2; see above, Materials and Methods). The texture statistics encoding model includes parameters that capture the spatial selectivity of voxels (pRFs; Dumoulin and Wandell, 2008; St-Yves and Naselaris, 2018) as well as their selectivity for a range of features related to local image structure (computed using P-S statistics; Portilla and Simoncelli, 2000, described in more detail below). All parameters of the model were fit on a voxelwise basis, and overall accuracy for each voxel was quantified by generating predicted voxel responses to a set of held-out images that were not used during model training. We then computed the coefficient of determination (R^2) for each voxel. To facilitate comparisons across visual regions with different functional roles in midlevel visual processing, we fit and evaluated the model across a large portion of occipotemporal cortex as well as computed summary statistics for several functional ROIsearly retinotopic areas V1-hV4; scene-selective areas OPA, PPA, RSC; face-selective area FFA; and body-selective area EBA (see above, Materials and Methods).

Our model includes a range of features related to both low and midlevel properties of images. Lower-level properties include the activation in different orientation and frequency channels as extracted by a steerable pyramid (Simoncelli and Freeman, 1995) and the marginal statistics of the pixel luminance; we refer to these collectively as "lower-level" texture features (see above, Materials and Methods). The model also includes features that capture higher-order (i.e., midlevel) structure in the image, including the cross-correlations between different orientation and frequency maps output by the steerable pyramid, as well as autocorrelations computed from individual steerable pyramid maps. We refer to these collectively as "higher-level" texture features. Figure 2 provides examples of image patches that result in high and low values for the first two principal components of each subset of model features, giving some intuition for the properties that are captured by each set of features. Figure 2A-D illustrate the four subsets of lower-level model features, which capture relatively simple aspects of image structure such as mean luminance (pixel features; PC2) and the strength of horizontal and vertical orientations (energy-mean features; PC2). In contrast, Figure 2E-J illustrate the types of properties captured by the higher-level model features, which are relatively more complex. For example, the linear-auto features appear to differentiate image patches according to the presence of high-frequency spatially repeating structure in the images (items like window blinds and zebra stripes), whereas PC1 of the linear-cross-scale features differentiates patches including white lines on a black background from patches including black lines on a white background. Other subsets of the higher-level model features, such as the energycross-orient features, appear to be related to conjunctions between differently oriented elements in the image (bent knees and elbows on a person, angles created by parts of an airplane), whereas PC1 of the energy-cross-scale features appears to be related to spatial frequency but maintains some invariance to orientation. Some of the higher-level model features also appear to capture distinctions between image patches that are more angular or rectilinear versus those that include more organic and curvy shapes (see PC2 of the energy-cross-orient (G) and energy-crossscale (I) features). Although not all the principal components are easy to describe in words, this illustration provides some intuition on how our model can differentiate complex natural scene images according to multiple aspects of their midlevel image structure.

Overall, the texture statistics encoding model achieved good predictive accuracy (R^2) across multiple visual areas, with particularly strong performance in early visual cortex (Fig. 3A,B, blue bars). Although performance of the model progressively declined from V1 through more anterior category-selective visual regions, performance remained moderately high in voxels throughout the visual hierarchy, with the majority of voxels in all visual ROIs having above-chance validation set accuracy (Fig. 3D; one-tailed p values obtained using permutation test, corrected for multiple comparisons; q = 0.01). In addition, examining the spatial fit parameters of the model (Fig. 4) demonstrates that the model recovers single-voxel pRFs with properties that are consistent with past work (Dumoulin and Wandell, 2008), namely, the tendency of pRF size (σ) to increase from early to higher-level visual ROIs and the tendency of pRF size to scale with eccentricity. These results provide validation of our modeling framework and indicate that the model is able to capture a substantial portion of the response variance in voxels across multiple stages of the visual hierarchy.

We compared the performance of our model to several commonly used models for early vision, including the GIST model (implemented with a coarse 2×2 and a finer 4×4 spatial grid), a Gabor model, and the first two layers of a convolutional neural network, AlexNet (see above, Materials and Methods). The GIST and Gabor models were selected for comparison because they both capture spectral information but not higher-order correlations, meaning they should have some feature overlap with the lower-level texture model features but not the higher-level features. In contrast, AlexNet is a larger model trained on object classification and thus may encode features not captured within the texture statistics model. In particular, the increase in feature complexity from Conv1 to Conv2 of AlexNet may allow it to capture higher-order



Figure 4. pRF parameters estimated using the texture statistics encoding model (see above, Materials and Methods for details on pRF grid and fitting procedure). *A*, Distribution of pRF size (σ) across voxels in each ROI. Distributions include voxels pooled across all subjects, open circle indicates the median. *B*, Relationship between pRF σ and eccentricity, plotted for voxels pooled across early visual areas (V1, V2, V3, hV4; dark blue) or higher visual areas (OPA, PPA, RSC, FFA, EBA; light blue), across all subjects. Values of σ are averaged over all voxels having the same preferred eccentricity; error bars indicate the mean \pm SD across σ values within an eccentricity. Note that the slight boomerang shape of these plots may be related to the fact that there are few voxels within the most foveal eccentricity bins, as well as that our images are relatively small (8.4° squares), which may bias the center estimates of large pRFs toward the fovea.

correlation structure from images, similar to the higherlevel texture features in our model. As shown in Figure 3B, our texture statistics model had comparable prediction accuracy to these other models, with texture model accuracy slightly exceeding that of the 2 \times 2 GIST model, the Gabor model, and the first convolutional layer of AlexNet (Fig. 3C). The accuracy of the texture model was exceeded slightly by the second convolutional layer of AlexNet and (in some higher visual areas) the 4×4 GIST model. In the case of AlexNet Conv2, the likely reason for this advantage is that AlexNet is able to capture a wider range of features than the texture statistics model because it has many more parameters and is task optimized as opposed to being based on hand-designed features. In the case of GIST, the difference was unexpected because the GIST features are based on Gabor filter outputs and are thus relatively low level. On reflection, one possible explanation is that GIST includes spectral features extracted from multiple positions in the visual field, so it is capable of capturing higher-order aspects of response selectivity. For example, GIST can capture the responses of a voxel that is sensitive to vertical orientations in the upper left visual field and horizontal orientations in the lower right visual field. Thus, GIST is able to approximate some of the same features captured by the higher-level texture features in our model, which may explain its slight advantage in higher visual areas with larger RF sizes (e.g., PPA). It is important to note that the main goal of our modeling was interpretability rather than maximizing model accuracy or outperforming pre-existing models. That is, we developed a model that can be used to isolate the contributions of different midlevel texture features, a property that is not provided by any of our comparison models. The fact that the texture model yields competitive performance with similar models provides a good indication that its features are reliably predictive of neural responses.

Given the high overall accuracy of the texture statistics encoding model, we next asked which features were most critical to its predictive performance. To test this, we performed a variance partitioning analysis (see above, Materials and Methods), where we subdivided the model features into lower-level and higherlevel texture features (Fig. 1) and determined both the percentage of variance that was uniquely attributable to each set of features and the percentage that was shared among the two sets (Fig. 5). This analysis revealed key differences among visual areas. First, visualizing the unique variance values on a flattened cortical surface (Fig. 5A) revealed a gradient from lower to higher visual areas, where voxels in the most posterior portion of occipital cortex tended to have more unique variance explained by the lower-level features (shades of blue), whereas more anterior voxels, particularly on the lateral surface of the brain, had progressively more variance uniquely explained by the higher-level texture features (shades of red). Averaging the unique variance values across voxels within each ROI further underscored this dissociation between areas. In early retinotopic areas V1-hV4, the lower-level texture features explained more unique variance than did the higher-level texture features; but in higher category-selective visual areas FFA and EBA, this trend reversed, with the higher-level features accounting for more unique variance on average (Fig. 5B). Place-selective ROIs OPA, PPA, and RSC showed a more intermediate pattern, with the lowerand higher-level features explaining similar amounts of unique variance. Consistent with these patterns, a bootstrap significance test revealed that the average unique variance explained was significantly higher for the lower-level features than the higher-level features in V1, V2, V3, and hV4, but it was significantly higher for the higher-level features than the lower-level features in EBA (Fig. 5B; two-tailed p values computed with a bootstrap test; corrected for multiple comparisons; q = 0.01; see above, Materials and Methods). In addition to significance at the ROI-averaged level, these differences were significant in all individual subjects in V1-V3, in seven individual subjects in hV4, and in four individual subjects in EBA (Extended Data Table 5-1). Furthermore, more individual voxels in early visual areas had significant unique variance for the lower-level features than for the higher-level features, whereas more voxels in FFA and EBA had significant unique variance for the higher-level features (Fig. 5C; one-tailed p values for single voxels computed using bootstrap test; corrected for multiple comparisons; q = 0.01). Another trend evident in this analysis was that across all visual areas, a substantial portion of the variance was shared between the lower-level and higher-level texture features (Fig. 5B, gray bars), suggesting some degree of feature overlap between the lower- and higher-level feature spaces.

The distinction between low- and high-level visual areas in the previous analysis is consistent with the interpretation that these areas represent features at different levels of complexity, with early areas representing more low-level aspects of image structure and higher visual areas representing higher-order statistics. However, another factor that could potentially contribute to this distinction is receptive field size. In our procedure for computing texture statistics features, the pRF size parameter (σ) determines the size of the spatial weighting function used



Figure 5. The unique variance explained by the higher-level texture features increases from lower to higher visual areas. Unique variance explained by the lower-level and higher-level features was measured using a variance partitioning analysis (see above, Materials and Methods). *A*, Percentage variance (units of $\sqrt{R^2}$) uniquely explained by the lower-level (shades of blue) and higher-level (shades of red) features, plotted on an inflated cortical surface from three viewpoints (left) or a flattened cortical surface (right) for one example subject, S1. For the full set of 8 subjects, see: http://www.cs.cmu.edu/~mmhender/viewers/texturemodel/varpart_low_vs_high/. Transparent voxels indicate that little variance was uniquely attributable to either the lower- or higher-level features, whereas white voxels indicate a large proportion of unique variance for both feature types. *B*, Summary of the variance partitioning results averaged across ROIs and subjects. Gray bars indicate variance shared between the lower-level and higher-level features, whereas blue and red bars indicate variance unique to each set of features. Bar heights and error bars indicate median and confidence intervals (99%) for the average unique and shared variance, obtained by bootstrapping the images when computing R^2 (see above, Materials and Methods). All confidence intervals are significantly greater than zero (one-tailed *p* values computed using a bootstrap test; corrected for multiple comparisons; *q* = 0.01). Light gray dots indicate the mean variance explained for each individual subject. Extended Data Table 5-1 shows the significance of individual subjects. *C*, Proportion of individual voxels in each ROI that had a significant amount of variance uniquely explained by the lower-level (blue bars) and higher-level (red bars) texture features (one-tailed *p* values computed using a bootstrap test; bar heights and error bars indicate mean variance explained for each individual subject. Extended Data Table 5-1 shows the significance

when computing each texture feature (see above, Materials and Methods). This implies that for larger σ values, the higherorder texture statistics features may be more informative because they incorporate information about a larger portion of the visual field. As voxels in higher visual cortex tend to have larger σ than voxels in early visual cortex (Fig. 4A), it is possible that the ability of higher-level texture features to explain more variance than lower-level features within higher visual areas is driven only by larger receptive field sizes. To evaluate this possibility, we constructed new versions of our encoding models in which σ was fixed at a single value, and then refit the entire encoding model with all voxels having the same σ . Consistent with the difference in estimated receptive field size between areas, this procedure resulted in the highest performance in early areas when the fixed σ value was relatively low, but the highest performance in higher areas when σ was relatively high (Fig. 6A). To allow for a balanced comparison across early and higher

visual areas, we selected a σ close to the middle of our range of candidate values, 1.48°, for further inspection.

Importantly, when σ was fixed at 1.48° we replicated the key findings of our original model (Fig. 6*B*,*C*). We again found that in posterior early visual areas, the lower-level texture features explained a greater proportion of the model variance than the higher-level texture features, but in more anterior areas this pattern began to reverse, with the higher-level texture features explaining a progressively larger percentage of the variance. This pattern was observed across a range of σ values (Fig. 6*D*), although not for the lowest σ values (Note that as described previously, small σ values resulted in low overall R^2 for higher visual areas.). The observation that a distinction between the coding properties of low- and high-level visual areas can be recovered even when σ is fixed at a single value for all voxels indicates that the difference between these areas involves a true difference in feature selectivity, not only a difference in the scope of spatial



Figure 6. The difference in feature sensitivity between early and higher visual areas is not dependent on differences in receptive field size. We refit the entire texture statistics encoding model with a single fixed pRF size ($\sigma = 1.48^\circ$) for all voxels (see above, Materials and Methods). *A*, Overall accuracy (R^2) of models in which pRF size (σ) was fixed at a single value for all voxels, averaged across voxels in each ROI. Each colored bar represents a different σ value; gray bar represents the model in which σ was allowed to vary across voxels (similar to Fig. 3*B*). Bar height and error bars indicate the mean \pm SEM across subjects. *B*, Percentage variance (units of $\sqrt{R^2}$) unique to the lower-level (shades of blue) and higher-level (shades of red) texture features, shown on a flattened cortical surface for two example subjects. *C*, The percentage of variance that was shared among feature types (gray bars), unique to the lower-level texture features (blue bars), or unique to the higher-level texture features (red bars), averaged over voxels within each ROI. Bar heights and error bars indicate mean ± 1 SEM across subjects, light gray dots indicate individual subjects. *D*, Variance partitioning analysis performed for other values of σ , shown for selected ROIs V1, hV4, and EBA. Bar heights and error bars are as in *C*.

selectivity. This result is also consistent with (Freeman et al., 2013), who found no correlation between the receptive field size of V1 and V2 neurons and their relative sensitivity to higher-level texture statistics.

The previous analyses suggest a dissociation between lowand high-level visual areas in terms of which texture statistics features best explain their responses. However, it is not yet clear whether all subsets of the lower- and higher-level texture features contribute equally to model performance in a given brain region or whether there is redundancy among the feature subsets. Thus, to more precisely determine which texture features were most uniquely predictive of neural activation, we next performed a second variance partitioning analysis where we considered each subset of both the lower-level and higher-level feature groups separately (a total of 10 feature types; see above, Materials and Methods).

As shown in Figures 7, 8, and 9, this analysis revealed that overall model accuracy reflected unique contributions from



Figure 7. The organization of neural populations sensitive to each subset of texture model features, shown for one representative NSD subject (S1). For the full set of 8 subjects, see: http://www.cs. cmu.edu/~mmhender/viewers/texturemodel/varpart_all_subsets/. The percentage of encoding model variance (units of $\sqrt{R^2}$) uniquely explained by each individual feature type is shown on a flattened cortical surface (see above, Materials and Methods for details on feature types). The maps in shades of blue (top row) correspond to subsets of the lower-level texture features, whereas maps in shades of red (bottom two rows) correspond to subsets of higher-level texture features. Note that the linear-mean features are not plotted here, as they explained little unique variance in any ROI (Fig. 8).

multiple feature subsets. Across all areas, the feature subset that uniquely explained the largest proportion of the texture model's R^2 was the energy-mean features (Fig. 7, top middle), which are analogous to a model of complex cell responses in V1. In early areas, a small amount of variance was also explained by the pixel features, which include marginal statistics computed from the image luminance histogram, and thus may capture low-level properties like overall image brightness or contrast (Fig. 7, top left). A portion of the variance in some early areas was also explained by the linear-cross-orient features (Fig. 7, bottom left), a higher-level feature subset which consists of cross-correlations between the real parts of steerable pyramid bands with different orientations. These features may capture higher-order structure generated from the combination of different orientations, such as corners, angles, and other contour junctions (Fig. 2). Notably, the highest values of unique variance for the linear-cross-orient features tended to be observed for voxels sensitive to the vertical meridians in early visual cortex (i.e., boundaries between V1 and V2 and between V3 and hV4). Given the previously reported relationship between pRF polar angle and preferred orientation (Freeman et al., 2011; Roth et al., 2022), this retinotopic relationship may suggest these features are related to a comparison



Figure 8. Results of variance partitioning analysis across all feature types, summarized at the ROI level. The proportion of model variance uniquely explained by each feature type is shown for each ROI, averaged across all subjects. Bar heights and error bars indicate median and confidence intervals (99%) for the average unique variance, obtained by bootstrapping the images when computing R^2 . Asterisks (*) above each bar indicate that the unique variance was significantly greater than zero (1-tailed *p* values computed using a bootstrap test; corrected for multiple comparisons; *q* = 0.01; see above, Materials and Methods). Bars in shades of blue indicate feature subsets belonging to the lower-level group, whereas bars in shades of red indicate feature subsets belonging to the higher-level group. Extended Table 8-1 shows the number of individual subjects in which unique variance was significant for each feature subset.

between vertical and horizontal orientations, as well as having some association with visual field position. Moving into more anterior regions, other higherlevel features began to explain a larger proportion of the variance. The linearauto features (Fig 7, center), which are computed based on spatial autocorrelations, explained a moderate proportion of variance across several high-level visual areas, including voxels in and near the category-selective areas; this feature subset may capture periodic, spatially repeating textural elements in the images (Fig. 2). Other feature subsets that explained a notable amount of



Figure 9. Proportion of the total model variance in each ROI explained by each feature type. Variance partitioning analysis is performed in the same way as in Figure 8, but each unique variance value is expressed as a proportion of the total R^2 of the model. Results are averaged across all subjects. Blue shades indicate lower-level model features, whereas red shades indicate higher-level model features (see above, Materials and Methods).



Figure 10. PCA performed on the unique variance values for each subset of texture statistics features. We concatenated the unique variance values for each feature subset (Fig. 7) across all subjects and then performed PCA to learn a set of weights that projects the unique variance values onto a lower-dimensional subspace. *A*, The weights (loadings) for the first four PCs are plotted on a flattened cortical mesh for two example subjects (top row, S1; bottom row, S2). Similar results were obtained for the other six subjects. *B*, The percentage of variance explained by each principal component. *C*, The score for each of the feature subsets with respect to the first four principal components (each colored line indicates a different principal component). *D*, The most activating (left) and least activating (right) images are shown for each of the first four principal components. To obtain these images, we identified the images that were overlapping across all subjects (907 images) and concatenated the voxel responses to these images across all subjects. PCA weights were then used to project these voxel response patterns into principal component space, and we identified the thee images yielding the largest and smallest responses for each PC of interest.

unique variance in higher-level visual cortex included those computed from higher-order statistics of the steerable pyramid magnitude features, including the energy-cross-scale features as well as the energy-cross-orient features.

Additionally, the spatial pattern of unique variance explained by each feature type suggested some heterogeneity within individual ROIs. For example, different portions of EBA appeared to be differentially predicted by the linear-auto, energy-cross-scale, and energy-cross-orient features, and higher-level features like the linear-cross-orient features explained a larger amount of variance in the posterior portion of PPA than the anterior portion. Some of this variability may be attributable to differences in the overall accuracy of the model across voxels within an ROI (Fig. 3*A*); however, some of the variability does not appear to be related to patterns of overall accuracy. This suggests that these ROIs may contain subregions that can be delineated based on midlevel response properties.

To quantify these effects, we averaged the unique variance across voxels in each ROI (Figs. 8, 9). Again, this analysis

demonstrated a primary role for the energy-mean features in all ROIs, with this feature subset dominating the most strongly in early visual areas. The unique variance explained by the energymean features was significant at the subject-averaged level in all ROIs except EBA, and was significant in all individual subjects in V1, V2, V3, and hV4 (Extended Data Table 8-1; one-tailed p values computed using bootstrap test; corrected for multiple comparisons; q = 0.01; see above, Materials and Methods). Although the absolute amount of unique variance contributed by other feature subsets was modest in comparison to the energy-mean features, other feature subsets nonetheless contributed a significant amount of unique variance in each ROI. The pixel features made significant contributions in several areas, with the highest unique variance contributed in V1 and V2 (all eight subjects individually significant in V1 and V2). As suggested in the previous paragraphs, the linear-crossorient features also contributed to the variance explained in many areas, with unique variance explained significant at the subject-averaged level in all ROIs (significant in all eight

subjects in V1 and V2). Another higher-level feature subset that stood out was the linear-auto features, which explained a significant amount of unique variance in V3 as well as OPA, PPA, RSC, FFA, and EBA. In general, the relative proportion of unique variance explained by these higher-level texture statistics features increased from early visual areas to higher visual areas (Fig. 9). These results suggest that the performance of the texture statistics encoding model reflects contributions from multiple feature subsets within the model, with the relative importance of these feature subsets varying among visual areas. Areas across the visual cortex may encode complementary aspects of image texture, resulting in a population representation that spans a large portion of the full texture statistics feature space.

To further explore the representational space defined by the texture statistics model, we performed PCA on the unique variance values for each of the 10 feature subsets (Fig. 10). PCA identified a set of weights in voxel space that project the unique variance values onto a lower-dimensional subspace. Visualizing these weights for the first four principal components reveals several large-scale organizational motifs across visual cortex. PC1 had its highest weights for voxels in early visual cortex, with a high score for the energy-mean features; this result is not surprising given the high magnitude of unique variance values for the energy-mean features in early visual cortex (Fig. 7). PC2 had high positive weights for voxels in higher visual cortex and negative weights in early visual areas, especially for voxels sensitive to the central visual field. PC2 had a high positive score for the linear-auto features and negative score for the pixel features (Fig. 10C). Plotting the top and bottom images for PC2 indicates a potential relationship with image scale (near/far), as well as the presence of human figures and/or actions. PC3 also seemed to covary with image scale, as well as appearing to create a rough division within early visual cortex based on retinotopy, having positive weights for voxels sensitive to more central and horizontal visual field positions and negative weights for those sensitive to vertical visual field positions. This is consistent with the observation that PC3 had a strong negative score for the linear-cross-orient features, which also had high unique variance values for the vertical-meridian-preferring populations in early visual cortex (Fig. 7). As mentioned previously, these populations are also likely to be selective for vertical orientations. Consistent with this, PC3 tended to be most activated by scenes including a strong representation of the horizon (meaning voxels with a negative weight on PC3 were negatively associated with horizon-dominated scenes), possibly reflecting sensitivity for the higher-order statistics associated with naturalistic outdoor scenes. Finally, PC4 appeared to be most positively weighted for voxels in face- and body-selective regions (FFA, EBA), and more negatively weighted in PPA and RSC, especially for subject 1 (S1) as well as in periphery-preferring regions of early visual cortex. PC4 had its highest positive score for the energy-cross-scale features and a negative score for the linearcross-orient features. Based on the top and bottom images for this component (Figure 10D), PC4 appears to be sensitive to the difference between images including animals or humans versus scene images that were strongly geometric in appearance. This is consistent with past work suggesting that sensitivity to midlevel features associated with object animacy is a dominant axis of visual cortex organization (Konkle and Caramazza, 2013). Together, these results suggest that several large-scale organizational properties of visual cortex can be recovered based only on the patterns of unique variance across feature subsets in our model. Therefore, texture statistics features may provide an interpretable form of scaffolding for the emergence of higher-level semantic representational axes in the brain.

Discussion

Intermediate-level feature representations are an essential component of the human visual hierarchy, providing a link between low-level image properties and high-level conceptual information. We investigated this link by constructing voxelwise encoding models based on a set of image-computable texture statistics features (P-S statistics; Portilla and Simoncelli, 2000). In both early retinotopic and higher category-selective regions, our model generated accurate predictions of voxel responses to natural scene images that were held out during model fitting. Features contributing the most to the predictive accuracy of the model differed depending on position in the visual hierarchy; lower-level texture features explained more unique variance in early visual cortex, whereas higher-level texture features progressively explained a larger amount of unique variance in more anterior regions. At a finer-grained level, patterns of texture feature sensitivity were able to identify meaningful components of the overall representational space within visual cortex. These results increase our understanding of midlevel visual representations and highlight that such representations participate in processing even at later stages of the visual hierarchy.

Results broadly consistent with ours have been found in primate visual cortex. Multiple studies have shown that spectral features (i.e., the energy-mean features in our model), tend to have the largest regression model weights of any P-S feature type in V1, V2, and V4 (Okazawa et al., 2015, 2017; Hatanaka et al., 2022); this is consistent with our results showing the highest unique variance explained for the energy-mean features in every early visual ROI (Fig. 8). Also consistent with Hatanaka et al., (2022), we found that the pixel features had the second-largest amount of unique variance explained in V1. More generally, our finding that higher-level feature sensitivity increased from V1 to hV4, as well as increasing further in higher areas, is consistent with past studies reporting that the contributions of higher-order texture statistics are larger on average for V4 than either V1 (Hatanaka et al., 2022) or V2 (Okazawa et al., 2017). Importantly, our results expand on these past findings by demonstrating the continuous increase in feature complexity beyond early visual cortex and across a wide range of areas in the visual system.

Less aligned with past work is the relative importance of different higher-level texture feature subsets. In V1-hV4, we found that the higher-level texture feature subset that explained the most unique variance was the linear-cross-orient features. Although one study found relatively high average weights for this set of features (along with the linear-cross-scale features; Hatanaka et al., 2022), other studies did not include this feature subset in their models (Okazawa et al., 2015, 2017), and their results instead indicated a larger role for the energy-cross-position and energy-cross-scale features. These discrepancies may be the result of modeling differences across studies, such as the inclusion of certain feature subsets, particularly if some subsets carry redundant information with one another. Differences in stimulus type may also contribute to these discrepancies as two studies (Okazawa et al., 2015, 2017) used synthetic texture stimuli as opposed to natural scene images, and sensitivity to higher-order texture features may be affected by the additional

structure present in natural scene images. In particular, phase information may be important for modeling natural scene images, which are nonstationary across space, compared with spatially homogeneous texture images. This is consistent with our observation that the linear-cross-orient and linear-auto features showed an advantage over the energy-cross-orient and energy-auto features (which are both phase invariant). For the linear-cross-orient features, which contributed significant unique variance in early visual areas, higher-order selectivity may be driven by simple cells, whose responses are phase dependent. Further work will be needed to evaluate these possibilities.

Although the lower-level features explained more unique variance in V1 than the higher-level features (Fig. 5), a small but significant amount of unique variance in V1 in all eight subjects was explained by the higher-level texture features (Extended Data Table 5-1). In particular, the linear-cross-orient features, which include information about the relationships between different orientations in the image, explained a significant amount of variance in V1 (as well as V2, V3, and hV4). This result is somewhat surprising given that past work has found V1 neurons have little sensitivity to the higher-order correlation statistics of the P-S model (Freeman et al., 2013). However, other work suggests that V1 neurons are sensitive to higher-order statistical correlations (Purpura et al., 1994) and show context-dependent responses (e.g., figure-ground separation, sensitivity to illusory contours or sensitivity for perceived size; Albright and Stoner, 2002; Carandini et al., 2005; Murray et al., 2006). Such contextual sensitivity may account for the observed selectivity to higherorder image structure in our results. Given the limited temporal resolution of fMRI, we cannot distinguish whether the sensitivity to higher-level texture statistics in V1 is because of feedforward processing, lateral interactions, or feedback from higher visual regions.

Sensitivity to higher-level texture features was also observed in more anterior areas, including voxels in and around categoryselective visual regions such as EBA. When the contributions of fine-grained feature subsets were analyzed, the linear-auto texture features were among the most predictive subsets for higher visual areas, explaining a significant amount of unique variance at the ROI-averaged level in OPA, PPA, RSC, FFA, and EBA. This feature subset likely captures image periodicity, consistent with past work suggesting ventral visual areas VO1 and LOC are sensitive to the degree of spatial regularity in synthetic texture stimuli (Kohler et al., 2016). Other higher-level feature subsets also contributed significant amounts of unique variance in higher visual areas, with the energy-cross-scale features having the highest average unique variance in EBA. The cross-scale correlations included in these features may allow them to differentiate between different kinds of oriented elements, such as lines versus object edges, as well as capturing information about image scale (Fig. 2, bottom left). The sensitivity to these diverse sets of texture statistics features across higher visual cortex also appeared to be related to differences in higher-level tuning properties. Our PCA analysis across the unique variance maps revealed components that captured high-level image properties like viewing distance (scale), the presence of animate figures, and scene elements like horizons and geometric layout. Thus, midlevel representations across higher visual cortex may support the broader computational functions of these regions.

Within higher visual cortex, higher-level texture model features explained relatively more unique variance in face- and body-selective areas (FFA, EBA), whereas lower-level model features contributed more in scene-selective areas (OPA, PPA, RSC). The sensitivity to lower-level features in scene-selective areas is consistent with work showing sensitivity to simple oriented features in scene-selective cortex (Nasr and Tootell, 2012; Lescroart et al., 2015), whereas the high sensitivity to higher-level features in face- and body-selective areas is consistent with the finding that the fusiform gyrus and the middle occipital gyrus exhibit larger responses to texture stimuli that include higher-order correlations (Beason-Held et al., 1998). This difference could also indicate that the higher-level features of our model align better with the midlevel features represented within face- and body-selective areas (curvy, organic features; Ponce et al., 2017) compared with within scene selective areas (geometric, rectilinear features; Nasr et al., 2014). Processing in scene-selective cortical areas may also be more spatially global, reflecting a role in computing large-scale properties such as scene layout (Epstein and Baker, 2019). Our model features are computed over local regions of the image and, thus, may fail to capture the large-scale structure of scenes. This may also be a reason why the 4×4 GIST model, which incorporates information from multiple spatial regions of the image, yielded slightly more accurate predictions than the texture model in scene-selective areas (Fig. 3). Importantly, the use of only a single pRF for each voxel in our model serves the purpose of isolating feature selectivity from spatial selectivity, which facilitates a straightforward interpretation of our variance partitioning results based on feature selectivity only. Future work may incorporate global features with the goal of improving accuracy in scene-selective areas; however, this is beyond the scope of the present work.

Note that the overall performance of the model in more anterior regions was relatively poor compared with early areas. Thus, perhaps unsurprisingly, the texture statistics model does not fully capture the range of image features to which voxels in higher visual cortex are sensitive. This conclusion is consistent with past work suggesting that higher visual areas in macaques and humans, although exhibiting some sensitivity to P–S statistics, are also sensitive to certain high-level aspects of natural image structure that are not captured by these features (Rust and DiCarlo, 2010; Long et al., 2018). However, our results do indicate that intermediate visual features are reliably encoded within high-level visual areas and that modeling these midlevel features can contribute to a more detailed understanding of the representations in these regions.

Relatedly, the ability of the texture statistics model to predict visual responses does not imply that the model features capture the explicit computations performed within these brain areas. As with any computational model, it is possible that the true features represented by the brain are merely correlated with the model features. Indeed, the similarity in performance between the texture statistics model and other low-level models suggests that there are multiple reasonable candidates. Nevertheless, the texture statistics model provides a simple and physiologically plausible hypothesis of how the brain computes midlevel feature representations, and we demonstrated that its features can predict voxel responses in visual cortex to natural images. Moreover, the contributions of different feature subsets to the performance of the model vary meaningfully across brain regions that serve different roles in visual processing. These findings pave the way for future work examining how midlevel features contribute to higher cognitive processes such as the recognition of complex objects and scenes.

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