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Color-Biased Regions of the Ventral Visual Pathway Lie between Face- and Place-Selective Regions in Humans, as in Macaques

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The existence of color-processing regions in the human ventral visual pathway (VVP) has long been known from patient and imaging studies, but their location in the cortex relative to other regions, their selectivity for color compared with other properties (shape and object category), and their relationship to color-processing regions found in nonhuman primates remain unclear. We addressed these questions by scanning 13 subjects with fMRI while they viewed two versions of movie clips (colored, achromatic) of five different object classes (faces, scenes, bodies, objects, scrambled objects). We identified regions in each subject that were selective for color, faces, places, and object shape, and measured responses within these regions to the 10 conditions in independently acquired data. We report two key findings. First, the three previously reported color-biased regions (located within a band running posterior–anterior along the VVP, present in most of our subjects) were sandwiched between face-selective cortex and place-selective cortex, forming parallel bands of face, color, and place selectivity that tracked the fusiform gyrus/collateral sulcus. Second, the posterior color-biased regions showed little or no selectivity for object shape or for particular stimulus categories and showed no interaction of color preference with stimulus category, suggesting that they code color independently of shape or stimulus category; moreover, the shape-biased lateral occipital region showed no significant color bias. These observations mirror results in macaque inferior temporal cortex (Lafer-Sousa and Conway, 2013), and taken together, these results suggest a homology in which the entire tripartite face/color/place system of primates migrated onto the ventral surface in humans over the course of evolution.

Key words: color; fMRI; homology; IT; topography; VVP

Significance Statement
Here we report that color-biased cortex is sandwiched between face-selective and place-selective cortex on the bottom surface of the brain in humans. This face/color/place organization mirrors that seen on the lateral surface of the temporal lobe in macaques, suggesting that the entire tripartite system is homologous between species. This result validates the use of macaques as a model for human vision, making possible more powerful investigations into the connectivity, precise neural codes, and development of this part of the brain. In addition, we find substantial segregation of color from shape selectivity in posterior regions, as observed in macaques, indicating a considerable dissociation of the processing of shape and color in both species.

Introduction
Color is a fundamental dimension of visual experience, informing us about myriad facets of our environment. Befitting its importance for human vision, considerable evidence from patient studies (Meadows, 1974; Newcombe and Ratcliff, 1974; Heywood et al., 1987, 1991; Zeki, 1997) and human neuroimaging (McKeefry and Zeki, 1997; Hadjikhani et al., 1998; Beauchamp et al., 1999; Wade et al., 2002, 2008; Brewer et al., 2005; Simmons et al., 2007; Bendris et al., 2016) attests to its fundamental role in perception and cognition. However, the neurobiological mechanisms of color vision remain largely unexplored. While the organization of the ventral visual pathway (VVP) has been the subject of extensive research, its relationship to color processing is not well understood. Here, we report results from a functional magnetic resonance imaging (fMRI) study designed to address this question.

The VVP is a critical component of the human visual system, responsible for the processing of visual information related to faces, objects, and scenes. It is thought to consist of three main components: the fusiform face area (FFA), the ventral stream, and the dorsal stream. The FFA is thought to be specialized for face recognition, while the ventral stream is responsible for object and scene recognition. The dorsal stream, on the other hand, processes information related to motion and depth. The VVP is thought to be homologous to the visual processing system in nonhuman primates, particularly macaques.

In macaques, color processing is thought to occur in the lateral geniculate nucleus (LGN) and the superior colliculus, with additional color processing in the inferior temporal cortex (IT). In humans, color processing is thought to occur in the occipital and parietal lobes, with additional color processing in the lateral枕叶 (Trouche et al., 2007). While the VVP is thought to be homologous to the visual processing system in nonhuman primates, the relationship between color processing and VVP organization remains unclear.

In this study, we aimed to address this question by examining the organization of color processing within the VVP in humans. We used fMRI to examine the organization of color processing within the VVP in 13 healthy human subjects. We used two versions of movie clips (colored and achromatic) of five different object classes (faces, scenes, bodies, objects, and scrambled objects) to stimulate the occipital and parietal lobes. We identified regions in each subject that were selective for color, faces, places, and object shape, and measured responses within these regions to the 10 conditions in independently acquired data. We report two key findings.

First, the three previously reported color-biased regions (located within a band running posterior–anterior along the VVP, present in most of our subjects) were sandwiched between face-selective cortex and place-selective cortex, forming parallel bands of face, color, and place selectivity that tracked the fusiform gyrus/collateral sulcus. Second, the posterior color-biased regions showed little or no selectivity for object shape or for particular stimulus categories and showed no interaction of color preference with stimulus category, suggesting that they code color independently of shape or stimulus category; moreover, the shape-biased lateral occipital region showed no significant color bias.

These observations mirror results in macaque inferior temporal cortex (Lafer-Sousa and Conway, 2013), and taken together, these results suggest a homology in which the entire tripartite face/color/place system of primates migrated onto the ventral surface in humans over the course of evolution. This result validates the use of macaques as a model for human vision, making possible more powerful investigations into the connectivity, precise neural codes, and development of this part of the brain. In addition, we find substantial segregation of color from shape selectivity in posterior regions, as observed in macaques, indicating a considerable dissociation of the processing of shape and color in both species. This result also has implications for understanding the evolution of color processing in the human visual system.
implicates specific cortical regions of the ventral visual pathway (VVP) in the analysis of color. But little is known about the causal role of these regions in perception (Murphey et al., 2008; Rangarajan et al., 2014), their connectivity, the precise computations conducted within them, or the underlying neural circuits. The most powerful methods for answering these questions require a primate model. Such a model would be most informative if the processing mechanisms were homologous to those found in humans. How similar, then, are the cortical mechanisms for color vision in nonhuman primates and humans?

Macques and humans have very similar color behavior (Stoughton et al., 2012; Gagin et al., 2014), raising the possibility that the two species have similar cortical machinery for color vision. Here, we test this hypothesis by investigating whether humans show the same organization for color in the VVP as found in macaques, in which color-biased regions are sandwiched between face- and place-selective regions (Lafer-Sousa and Conway, 2013; Verhoef et al., 2015). No prior study has assessed the topographic relationship of these regions directly in humans. A strong test of the idea requires a systematic and quantitative evaluation of category, shape, and color selectivity in the same subjects. To do this, we scanned 13 human subjects using fMRI while they viewed either full-color or achromatic movie clips of faces, objects, scenes, bodies, or scrambled objects. We first determined the spatial layout of color, face, and place selectivity by projecting color, face, and place preferences onto each subject’s inflated cortical surface. We then defined color-, face-, and place-prefering regions within each subject and measured the response magnitude of each region to each of the 10 conditions in independently acquired data. These analyses enabled us to map the relative locations of shape-selective, category-selective, and color-biased regions in each subject individually and to quantify the selectivity of each region to each dimension.

Our experimental design enabled us to address a second open question: Is the processing of color and shape inextricably linked throughout the VVP? On the one hand, one might think of color as just another object property, likely processed in the same cortical regions (and neurons) as those implicated in computing object shape. On the other hand, the computations entailed in processing color, and the role of color in behavior, often diverge from those for shape (Conway, 2009, 2014), suggesting color and shape are processed by somewhat separate circuits. Some micro-electrode studies in monkeys have been interpreted as supporting the idea that color and shape are processed together (Schein and Desimone, 1990; Gallant et al., 2000; Gegenfurtner and Kiper, 2003; Shapley and Hawken, 2010; Yasuda et al., 2010), but other results in macaques (Zeki, 1980, 1983; Bartels and Zeki, 2000; Conway, 2001; Lafer-Sousa and Conway, 2013; Conway, 2014) and humans (Newcombe and Ratcliff, 1974; Milner and Heywood, 1989; Heywood et al., 1991; Bouvier and Engel, 2006; Cavina-Pratesi et al., 2010) suggest some dissociation of color and shape processing. We addressed this question by determining the extent to which regions that respond preferentially to intact shape (compared with scrambled counterparts), such as the lateral occipital region (LO), were modulated by the presence of color, and the extent to which regions responding preferentially to color were modulated by the presence of intact shape. These experiments enabled us to test the degree to which color, category, and shape preferences are segregated in the human brain and to investigate possible homologies of cortical regions in humans and macaques.
(Lafer-Sousa and Conway, 2013); these stimuli allow more stringent control of the physical parameters of the stimulus but contain no recognizable objects. The chromatic gratings were designed to be equiluminant; the achromatic gratings were designed to comprise some luminance contrast. The stimulus paradigm was similar to that used to identify color-biased regions in macaque (Lafer-Sousa and Conway, 2013). Three achromatic gratings conditions (luminance contrasts 50%, 25%, and 7%) and eight color directions defined by the cardinal and intermediate directions of the equiluminant plane in cone-opponent color space (MacLeod and Boynton, 1979; Derrington et al., 1984) were used. Block-lengths were shortened compared with those used in the macaque experiments (18 s instead of 32 s) to match the hemodynamic response function of the BOLD signal in humans (block lengths needed to be longer in the macaque experiments to account for the longer hemodynamic delay that accompanies the use of an intravascular contrast agent). Stimuli were calibrated using spectral readings taken with a PR-655 spectroradiometer (Photo Research). The spectra were multiplied with the Judd-revised Commission Internationale de l’Eclairage (CIE) 1931 color matching functions to derive CIE x,y,Y coordinates of the monitor primaries (Hansen et al., 2008) and cone excitation was calculated using the Smith and Pokorny cone fundamentals (Smith and Pokorny, 1975). Stimuli were presented as vertical trapezoid-wave gratings (Conway and Tsao, 2006). Each chromatic and achromatic grating was drifted back and forth for 18 s, switching directions every 2 s (2 cycle/s, drifting 0.75 cycles/s). Within a run, the grating blocks were interleaved with equiluminant full-field gray (100 cd/m²) (12 s) to allow the signal to return to baseline (e.g., Gray, Achromatic-25%, Gray, Color-1, Gray, Achromatic-50%, Gray, Color-8, Gray, etc). There were 9 run orders, counterbalanced across runs such that each condition happened equally often in each serial position in the run. Each achromatic grating appeared three times per run, and was shown between chromatic blocks (with neutral gray blocks interleaved). Subjects were required to fixate a central fixation cross and perform a difficult motion-detection task to maintain attention: the gratings drifted left to right, switching direction every 2 s except once per block, when the grating would drift 3 s in one direction. Subjects had to report when this occurred (via a button press in the scanner).

Retinotopic mapping. In nine subjects, we identified the borders of visual areas V1 through V4 by mapping the responses to horizontal and vertical meridians following standard procedures (Sereno et al., 1995). The stimuli were flickering (counterphase at 7.5 Hz) checkerboards restricted to wedges (subtending 55°) along either the horizontal (spanning 20° of visual angle) or vertical (spanning 15° of visual angle) meridian. Each run comprised 36 10-second blocks. Subjects were required to fixate throughout. Responses to vertical meridians were contrasted with responses to horizontal meridians and projected on each subject’s cortical surface, producing a striped map used to demarcate the boundaries of V1, V2, V3 and V4. Because the anterior border of V4 (between V4/hi/v4 and ventral occipital cortex, VO-1) is difficult to define functionally (McKeefry and Zeki, 1997; Bartels and Zeki, 2000; Tootell and Hadjikhani, 2001; Brewer et al., 2005; Winawer et al., 2010), its location was determined using an anatomical landmark, the posterior transverse collateral sulcus (ptCoS) (Winawer et al., 2010).

Data acquisition Data were acquired using a Siemens 3T MAGNETOM Tim Trio scanner (Siemens AG, Healthcare, Erlangen, Germany) with a 32-channel head coil (at the Athinoula A. Martinos Imaging Center of the McGovern Institute for Brain Research at MIT). Functional data were collected using a T2*-weighted echo planar imaging (EPI) pulse sequence sensitive to blood-oxygen-level-dependent (BOLD) contrast. Given the known susceptibility artifact of the anterior regions of the ventral temporal lobe (caused by the ear canals), we performed extensive piloting to determine the parameters that produced the highest voxel SNR. Pilot data were collected with and without iPAT (Griswold et al., 2002) and field correction; we varied the voxel-resolution, slice angle, TE, and fraction of k-space sampled. We obtained improved TSNR and BOLD sensitivity by increasing voxel resolution by a factor of 4 (from the standard 3 mm iso to 2 mm iso), forgoing image acceleration (iPAT), and collecting field maps before each run. Reducing TE (to 14 or 23 ms) enhanced the tSNR, but compromised BOLD sensitivity. TE was set to 30 ms.

The functional volumes were acquired for a restricted portion of the brain spanning the ventral surface of the temporal lobe. Each functional volume comprised 25 slices (2 mm isotropic; field of view [FOV] = 192 mm, matrix = 96 × 96 mm) oriented approximately parallel to the temporal lobe, covering the occipital cortex (V1–V4) and the full length of the temporal lobe ventral to and including some of the superior temporal sulcus (2.0 s TR, 30 ms TE, 90° flip angle, 6/8 echo fraction). The first 5 volumes of each run were discarded to allow for T1 equilibration. Field maps (2 mm isotropic, 25 slices) were collected before each functional run to measure magnetic inhomogeneities and were used to estimate spatial distortions in the functional volumes that were then removed during analysis. High-resolution T1-weighted anatomical images were collected for each subject using a multiecho MPRAGE pulse sequence (1 mm isotropic voxels; FOV = 256 mm, matrix = 256 × 256).

Data preprocessing and modeling Data were processed using Freesurfer (http://surfer.nmr.mgh.harvard.edu/) and custom MATLAB scripts. Freesurfer was used to segment white- and gray-matter structures from the anatomical volumes (Dale et al., 1999; Fischl et al., 1999, 2001). Each individual subject’s functional data were field corrected, motion corrected using rigid-body transformation to the middle image of each run, intensity normalized (nonbrain tissue was masked), spatially smoothed using an isotropic Gaussian kernel (3 mm FWHM) to improve SNR, and aligned to the individual’s anatomical volume using a rigid-body transformation determined by the standard Freesurfer’s mrinfo (Greve and Fischl, 2009). Digitally inflated (see Figs. 1, 2, 5, 6, 9) and flattened (see Figs. 2, 6, 7, 9) cortical surface reconstructions were generated using freesurfer.

Whole-volume general linear model-based analyses were performed for each subject’s even and odd runs separately. Regressors were defined as boxcar functions convolved with a gamma hemodynamic response function (Friston et al., 1994). The boxcar function for each condition included each block from that condition. Nuisance regressors for motion (three translations, three rotations) and a linear trend to capture slow drifts were included.

Analysis of color, face, place, and shape preferences across cortex (dynamic localized data) To show the large-scale spatial organization of the ventral visual pathway within each individual, statistical contrast maps were projected on the digitally inflated surface of each subject’s own cortical anatomy. Following standard practices (Julian et al., 2012), functional regions were defined as follows. Face-selective voxels were defined as those showing higher responses to video clips of faces than objects (p < 0.0001; see purple regions in Figs. 1B, 2B). Place-selective voxels were defined as those showing a higher response to places than objects (p < 0.0001; on green regions in Figs. 1B, 2B). Color-biased voxels were defined as those showing a higher response to chromatic video clips than achromatic versions of the same clips matched in luminance, averaging responses obtained in all stimulus classes (p < 0.0001; activation maps shown in Fig. 1B; blue regions in Fig. 2B). The conclusions were not affected if color-biased regions were only defined using movie clips of a single category (data not shown). Shape-selective voxels were defined as those showing a higher response to intact objects than scrambled objects (p < 0.0001; see Fig. 5).

The major conclusions presented here are derived from an analysis of individual subject data. To illustrate overall trends in the spatial layout of color-biased regions relative to face-, place-, and shape-selective regions, we generated group overlap maps. To do this, statistical contrast maps from each of the 13 individual subjects were registered to a published group-average template (Freesurfer, CVS_avg35, an average of 35 people). The group-registered contrast maps from each individual were then thresholded and each voxel was assigned a binary value (1 or 0 for above threshold or not). For a given contrast, all 13 subject maps were tallied (those showing a higher response to places than objects (p < 0.0001; on green regions in Figs. 1B, 2B) and the voxels in the resulting volume were assigned a value according to the number of subjects who had above-threshold activity at that location. Figure 2A shows the group summary map for face, color, and place.
contrasts, thresholded at 5 subjects (individual contrast maps thresholded at \( p < 0.01 \)) on the reconstructed cortical surface of the CVS_avg35 volume. Figure 5A shows the group map for the shape contrast (intact objects > scrambled objects; \( p < 0.0001 \)) as a surface histogram where voxel color (red to yellow) reflects the agreement of 2 to 5+ subjects.

The surface maps enabled us to examine the spatial segregation of the activation patterns for color, faces, and places. However, regions of interest (ROI)-based analyses make possible stronger tests of our hypotheses. Specifically, they enabled us to cross-validate the contrast by which each region was defined and to quantify the effect size and selectivity of that contrast. Effect size matters: a region may have highly significant selectivity but of small magnitude, an important distinction that is not apparent in standard activation maps showing only significance levels. By quantifying activity in ROIs, we were able to determine the profile of response of each region to each of the 10 conditions tested, providing a richer profile of the selectivity of each region. From these results, we investigated whether the selectivity for color is present to a similar degree for each stimulus category and the extent to which it depends on stimulus category (which would be manifested by an interaction of color/achromatic by category). Significant interactions between two factors (e.g., color and shape) may be evidence that the two factors depend upon different neural populations rather than being analyzed independently in two distinct neural populations within that region (Sternberg, 1969). The ROI analyses also enabled us to test whether two regions differ significantly from each other with respect to some contrast. Standard activation maps do not support such conclusions because they only show regions that reach significance in a given contrast and regions that do not—that is, they show differences in significance, but not significant differences across regions. For example, in one ROI analysis, we explicitly tested whether color-biased regions are significantly more color biased than category-selective regions and shape-selective regions, contrasting the responses in these regions to each other in an ANOVA with ROI as a factor (described below).

Functional ROIs were defined in individual subjects, in the volume, using data from even runs. The odd runs were used to quantify the responses within each ROI to each of the 10 conditions. ROIs were defined using a \( p \)-threshold criterion. For a given contrast, all contiguous voxels above the \( p \)-threshold constituted an ROI; \( p \)-threshold was \( p < 0.0001 \) for occipital face area (OFA), fusiform face area (FFA), anterior temporal lobe (ATL), parahippocampal place area (PPA), LO cortex, posterior fusiform (pFS) and posterior superior temporal sulcus (pSTS); and \( p < 0.001 \) for color-biased regions. A less stringent \( p \)-threshold was used to define color-biased voxels because color selectivity was weaker overall than shape and category selectivity. Note that the evidence for color selectivity comes not from the statistics used to define the region, but rather from the shape of the response magnitudes conducted using independently acquired data (the odd runs). In some subjects, face activations were so robust that they formed a contiguous swath extending from the OFA to the FFA; similarly, shape-selective regions sometimes formed a contiguous swath from LO to pFS and place activations sometimes extended contiguously into early visual cortex. In these cases, a published contiguity criterion (\( p < 0.001 \)) as a surface histogram where voxel color (red to yellow) reflects the agreement of 2 to 5+ subjects.

As described in the Results, the color activation along the posterior–anterior extent of the VVP was not homogeneous; three main peaks in the activation were defined: posterior color-biased region (Pc), central color-biased region (Cc), and anterior color-biased region (Ac). To quantify this inhomogeneity, in each subject, we defined two ROIs (Fig. 4B, schematic, black circles) between the peak color-biased activations (white circles): iPc (intermediate to Pc and Cc) and iAc (intermediate to Cc and Ac). These ROIs were defined in each subject individually in the volume using even runs and the nonsignificant voxels in the contrast were used to define the color-biased regions. Figure 4B bar plots show the PSCs for colored and achromatic stimuli, averaging across faces, bodies, scenes, objects, and scrambled objects.

To plot color bias as a function of shape bias (see Fig. 5C), we used an index of color selectivity (response to color − response to achromatic/p response to color + response to achromatic) and a comparable index of shape selectivity (response to intact objects − response to scrambled objects/response to intact objects + response to scrambled objects). The response to color was the average PSC to chromatic intact objects and achromatic scrambled conditions, and the response to achromatic was the average PSC to the achromatic-intact and achromatic-scrambled conditions. The response to intact objects was the average PSC to chromatic and achromatic intact object conditions; the response to scrambled objects was the average PSC to chromatic and achromatic scrambled conditions. These indices were computed at the group level to avoid complications posed by dividing by noisy negative individual responses. Error bars were calculated using bootstrapping across subjects and represent 95% confidence intervals (10,000 bootstrapped samples) (Efron, 1982).

\section*{Group analysis (random effects) of color-biased activations (dynamic localizer)}

To test the extent to which color activations were found at consistent locations in the brain across subjects (see Fig. 4), we performed a group random-effects analysis on the chromatic versus achromatic contrast, in which data from each subject were registered and normalized to a common template (Freesurfer, CVS_avg35).

\section*{Analysis of inhomogeneity in tSNR across the cortical surface}

The ability to detect significant differences in the BOLD response in each region of the cortex depends on the signal-to-noise ratio (SNR), which is known to vary across the brain due inhomogeneities in the magnetic field. To show this spatial variation in SNR, we computed temporal SNR (tSNR) for each subject as follows: the motion-corrected functional volumes were high-pass filtered with a cutoff of 0.004 Hz to remove slow drifts and, for each voxel, we computed the mean of the time course for each voxel divided by its SD. To generate a mean tSNR map from the group of subjects, the tSNR map for each subject was registered to a common template (Freesurfer, CVS_avg35) and the volumes from all subjects were averaged.

\section*{Color localization with colored gratings}

In addition to localizing color-biased cortex using a contrast of chromatic > achromatic movie clips, color-biased regions were also localized in a subset of subjects using a more controlled stimulus paradigm containing no recognizable objects (full-field drifting gratings), comparable to the one used in an earlier study of macaque monkeys (Lafer-Sousa and Conway, 2013) and similar to those used in most prior work on color localization in humans (McKeefry and Zeki, 1997; Hadikhani et al., 1998; Wade et al., 2008). This enabled us to directly compare the human data with the previously published macaque data (obtained using gratings), and establish the extent to which the natural videos recover the same regions localized with standard low-level stimuli.

In the three subjects who participated in this control experiment, color-biased activations were defined by contrasting responses to the chromatic grating conditions that elicited the weakest response in MT+ (these colors would suffer the least luminance contamination) with the achromatic grating condition that yielded the same magnitude of response in V1 (50% luminance contrast). We quantified the PSC to gratings in each of the ROIs defined using the dynamic natural stimulus, as well as regions V1, V2, and MT.
Monkey data
If the homology between humans and monkeys hypothesized in the present report is true, then color-biased regions in monkeys (Lafer-Sousa and Conway, 2013) and color-biased regions in humans (described presently) should show a similar pattern of color and shape selectivity when scanned using the same dynamic natural videos. To test this hypothesis, we scanned the two macaque subjects from Lafer-Sousa and Conway (2013) while the animals were shown the dynamic natural video stimuli using the same acquisition and preprocessing methods described in the earlier report. MION contrast agent was used in the monkey experiments. The stimulus block lengths for the dynamic localizer were extended (32 s) to account for the slower hemodynamic response function associated with MION (Vanduffel et al., 2001). The task was free viewing, as in the human experiments; animals were rewarded with juice for looking anywhere on the screen. We extracted PSC responses to the dynamic natural videos using the ROIs defined in the earlier report (independent data): the middle face patch (ML) and the set of color-biased ROIs (PLc, posterior lateral color, located in P1T; C1c, central lateral color, located in C1T; and A1c, anterior lateral color, in A1T). Bar charts (see Fig. 8A) show the mean PSC of the last 7 TRs in each stimulus block. A scatter plot (see Fig. 8B) shows the relative color and shape bias of each ROI (as computed for Fig. 5B in the humans).

Results
Topography of face, color, and place responses
We first asked to what extent the functional organization of the human VVP resembles that of macaque monkey, where parallel and adjacent processing systems have been reported for faces, places, and color, with color-biased cortex sandwiched between face-selective regions (superiorly) and place-selective regions (inferiorly) (Lafer-Sousa and Conway, 2013; Verhoef et al., 2015). To do so, 13 subjects were scanned with fMRI while viewing short video clips containing multiple stimulus categories, each presented (in different blocks) in either full color or luminance-matched grayscale (6 3 s clips/block). Color-biased activity in the ventral visual pathway was found between the more lateral face-selective regions (faces > objects contrast, p < 0.0001, purple outlines) and the more medial/ventral place-selective regions (scenes > objects, p < 0.0001, green outlines). Color-biased activity spans much of the posterior-to-anterior extent of the VVP. Three peaks in the color-biased activity were defined in most subjects: Pc (called elsewhere V4 or hV4), Cc (V4a/V8/V0 complex), and Ac. The white dashed line denotes the ptCoS, which runs transversely to the base of the CoS and defines the border between V4 and the ventral occipital complex V0 (Winawer et al., 2010; Wittfoht et al., 2013). The boundaries of early visual areas as defined by standard meridian mapping (vertical/horizontal reversals) are shown for S1. Scale bar, 1 cm.

Figure 1. Sandwiching of color-biased regions between face-selective cortex and place-selective regions. A, Frames from the video clips used to define functional ROIs. Each stimulus block comprised video clips from one stimulus category (Faces, Bodies, Scenes, Objects, and Scrambled Objects) presented in color or luminance-matched grayscale (6 3 s clips/block). B, Color-biased activation on the ventral cortical surface in four subjects (S1–S4; color scale shows exponent of the chromatic > achromatic contrast defined using video clips from all categories). Color-biased activity in the ventral visual pathway was found between the more lateral face-selective regions (faces > objects contrast, p < 0.0001, purple outlines) and the more medial/ventral place-selective regions (scenes > objects, p < 0.0001, green outlines). Color-biased activity spans much of the posterior-to-anterior extent of the VVP. Three peaks in the color-biased activity were defined in most subjects: Pc (called elsewhere V4 or hV4), Cc (V4a/V8/V0 complex), and Ac. The white dashed line denotes the ptCoS, which runs transversely to the base of the CoS and defines the border between V4 and the ventral occipital complex V0 (Winawer et al., 2010; Wittfoht et al., 2013). The boundaries of early visual areas as defined by standard meridian mapping (vertical/horizontal reversals) are shown for S1. Scale bar, 1 cm.

FmA frequently consisted of a posterior and anterior component (FFA-1/pFus-faces and FFA-2/mFus-faces; Weiner and Grill-Spector, 2010). In addition to the face-selective regions of the VVP, the dorsal face-selective region of the pSTS (Puce et al., 1998) was detected in 11/13 subjects (8/11 bilaterally). The scene-selective retrosplenial cortex (Epstein et al., 2007) and occipital place area (Julian et al., 2012) were outside of the region scanned in many subjects and were not analyzed.

Activation maps in Figure 1B show those voxels with greater activation to colored versus achromatic video clips. In each subject, the color-biased activation was sandwiched between the face-selective
regions and the place-selective regions. The volume-wide contrast (smoothed 3 mm Gaussian FWHM) of the color-activation pattern showed a mottled band of color-biased activation running posterior-to-anterior along the VVP, within which we typically observed multiple peaks. To quantify the activity, we defined three color-biased regions of interest in most subjects (the interdigitating tissue is further examined in the “Quantification of the magnitude of the color bias across the VVP” section below). In all subjects (11/13 subjects bilaterally), we observed a posterior color-biased region that we refer to as Pc (for “posterior color”); cortex, and place-selective cortex medial. This pattern was even more obvious in the inflated and flattened views (Fig. 2A, bottom) and was evident in data from most individual subjects (Fig. 2B). Three subjects also showed some color-biased activation lateral to the FFA (S5, S7, S10; Fig. 2B). In some subjects, Cc overlapped partially with the PPA (overlap indicated in cyan). Although viewing activations on the inflated surface allows us to observe their relative position on the cortical sheet, smaller activations sometimes fail to project despite being visible in the volume (Fischl et al., 1999; Operto et al., 2008). Activations that fall
in regions of high signal inhomogeneity are particularly vulnerable to imperfect registration to the anatomy, which can lead to surface projection failure (Tucholka et al., 2012). Region Ac is acutely susceptible given its small size and proximity to the ear canals and is in a region of known signal inhomogeneity (discussed in “The anterior color-biased region” section below). Despite being visible in the native volume (where ROIs were defined), it failed to visibly project to the surface in several subjects (S2, S4, S8, S9, S11, and S13). For example, Ac was not clear in the surface for subject 2 (Fig. 2B), but was obvious in the native volume for this subject (slices; see Fig. 4).

In sum, we find a parallel and adjacent structure of face, color, and place preferences on the ventral surface of the brain in humans. This pattern mirrors the systematic organization found on the lateral surface of macaque inferior temporal (IT) cortex, suggesting a broad homology between the two regions.

**Segregation of color and category in the VVP**

To what extent are color and category information processed separately in the VVP? The brain activation maps described in the previous section show some spatial segregation of preferences for color, faces, and places. As described in the Materials and Methods, ROI-based analyses make possible stronger tests of our hypotheses about the functional relationship between color and other stimulus dimensions. Therefore, category-selective (OFA, FFA, ATL, PPA) and color-biased (Pc, Cc, and Ac) ROIs were defined in each subject individually using data from even runs and the response magnitude of each region to each of the 10 conditions was quantified using the independent data from odd runs (A–F, H). For comparison, we also defined a region that responded preferentially to intact objects (defined by shape selectivity not category selectivity) that is adjacent to Pc and OFA (region LO), and a face-selective region (pSTS, F) on the lateral surface (as opposed to the ventral surface). ROIs were defined using a p-threshold criterion (for the OFA, FFA, ATL, PPA, pSTS, and LO, p < 0.0001; for the color-biased regions, p < 0.001). For each subject, an ROI’s PSC was extracted by averaging β-values across the voxels in the ROI and dividing by mean BOLD signal in the ROI. When a region appeared bilaterally, the PSC for the left hemisphere ROI was averaged with the PSC of the corresponding right hemisphere ROI for each condition. Bar plots show the average PSC across subjects for each ROI (error bars indicate within-subject SE). Participants who lacked a particular ROI were not included in the statistical analysis for that ROI. Cyan bars indicate chromatic conditions and gray bars indicate achromatic conditions (F, faces; B, bodies; S, scenes; O, objects; ScO, scrambled objects).

As expected, color-biased regions responded more strongly to colored than to achromatic stimuli; the size of this effect in each ROI can be seen in Figure 3 (significance levels are reported in Table 1). Category-selective regions showed the expected strong (Fig. 3) and significant (Table 1) selectivity for their preferred category. However, most of the ROIs were selective for both color and category (Table 1). This result reflects in part the great statistical power of the ROI method (Saxe et al., 2006), which is able to detect statistically significant but very small effect sizes such as color in the FFA (the blue bar is slightly higher than the black bar in the first pair of bars in Fig. 3E). Are the color ROIs (Fig. 3A–C) more color selective than the category-selective ROIs (Fig. 3D–F, H), and are the category-selective ROIs more category selective than the color ROIs? Because there is no uncontroversial pairing of a particular color-biased ROI with a particular category-selective ROI, we answered these questions using a stringent exhaustive analysis of all 12 pairwise ANOVAs, contrasting each color ROI with each category ROI. This analysis confirmed that each color-biased ROI is more color selective than each category-selective ROI (i.e., p < 0.01 for the interaction of ROI \times color in 12/12 pairwise ANOVAs), and each category-selective ROI is more category selective than each color ROI (i.e., p < 0.0001 for the interaction of ROI \times category in 12/12 pairwise ANOVAs, p < 0.0001). Note that the large number of statistical comparisons used here does not require correction for

**Figure 3. Segregation of color preference and category selectivity in the ventral visual pathway.** To quantify the response profiles of specific ROIs, category-selective (OFA, FFA, ATL, PPA) and color-biased (Pc, Cc, and Ac) ROIs were defined in each subject individually using data from even runs and the response magnitude of each region to each of the 10 conditions was quantified using the independent data from odd runs (A–F, H). For comparison, we also defined a region that responded preferentially to intact objects (defined by shape selectivity not category selectivity) that is adjacent to Pc and OFA (region LO), and a face-selective region (pSTS, F) on the lateral surface (as opposed to the ventral surface). ROIs were defined using a p-threshold criterion (for the OFA, FFA, ATL, PPA, pSTS, and LO, p < 0.0001; for the color-biased regions, p < 0.001). For each subject, an ROI’s PSC was extracted by averaging β-values across the voxels in the ROI and dividing by mean BOLD signal in the ROI. When a region appeared bilaterally, the PSC for the left hemisphere ROI was averaged with the PSC of the corresponding right hemisphere ROI for each condition. Bar plots show the average PSC across subjects for each ROI (error bars indicate within-subject SE). Participants who lacked a particular ROI were not included in the statistical analysis for that ROI. Cyan bars indicate chromatic conditions and gray bars indicate achromatic conditions (F, faces; B, bodies; S, scenes; O, objects; ScO, scrambled objects).
multiple comparisons because our hypothesis required that each of the tests (rather than any of them) reach significance.

The statistical tests presented thus far provide evidence of some anatomical segregation of category and color responses: color-biased regions were more color selective than category-selective regions, and category-selective regions were more category selective than color-biased regions. If color and category processing were strongly independent, then each kind of information would be carried by different populations of neurons with no interaction. This strong hypothesis predicts that responses to color and category would not interact within an ROI (instead, the effect of the two factors on the response of a voxel containing the two distinct neuronal populations would be additive). This prediction follows from an extension to fMRI (Sternberg, 2011) of the classic additive factors logic from Sternberg (1969). All seven ROIs in the ventral cortex were tested (Pc, Cc, Ac, OFA, FFA, ATL, and PPA) and only the PPA showed a significant interaction of color by category (see Table 1).

These analyses provide evidence for a double dissociation between color and category preference across the VVP, as well as statistical independence of color and category preferences within six of the seven ROIs.

### Quantification of the magnitude of the color bias across the VVP

As described above, the color-biased activations comprised a mottled band running posterior to anterior along the VVP (Figs. 1B, 2, 4A). Within this band, the three main peaks (Pc, Cc, and Ac) were identifiable in most subjects (10/13). In each subject, each peak was typically separated by tissue that did not show a significant color bias. Moreover, peaks were sufficiently stereotyped in their location within the brain such that they are apparent in a random-effects group analysis (Fig. 4B). In the "Segregation of color and category in the VVP" section, we showed that these regions were significantly more color biased than category-selective regions, but how strong is their color selectivity relative to the VVP more broadly? In particular, are the color-biased regions more color selective than the cortical tissue that immediately surrounds them?

To answer this question, we used an ROI-based analysis that enabled us to leverage the power obtained by measuring responses in many subjects (see Materials and Methods). This method is powerful because it enables one to uncover activations that may not be significant in a given ROI within a single subject, but may be significant within that ROI when pooling results from many subjects. Using this method, we asked, are the color-biased regions more color biased than the immediately intervening patches of cortex?

In each individual subject, we used data obtained in even runs to define two ROIs located between the color-biased activations with respect to the posterior–anterior axis (Fig. 4C, schematic, black circles). These included region iPCC (between Pc and Cc) and iCAC (between Cc and Ac). We used results obtained during odd runs to quantify responses. Figure 4D shows each region’s percent signal change to all chromatic stimuli (cyan bars) and all achromatic stimuli (gray bars) (“all” = faces, bodies, scenes, objects, and scrambled objects). Although both intermediate regions were not color biased in the individual subject contrast maps, they showed main effects of higher responses to chromatic than achromatic stimuli in the ROI analysis [both p < 0.01; 2-factor ANOVAs on stimulus class (faces, bodies, scenes, objects, and scrambled objects) × stimulus color]. Nonetheless, the color-biased regions were more color biased than iPCC and iCAC: Pc and Cc were each more color biased than iPCC (p < 0.001) and Cc and Ac were each more color biased than iCAC (p < 0.003; pairwise 3-way ANOVAs on ROI × color × stimulus class, each contrasting an intermediate region with one of its neighboring color regions). These results show that, although a weak color preference extends into surrounding cortex, the color-biased regions represent reproducible peaks in that spatial pattern of color bias.

### Anatomical segregation of color and shape preferences

Prior work has documented regions of cortex that respond more strongly to intact than scrambled objects, including a posterior region called LO and an anterior region often referred to as posterior fusiform (pFS) (Grill-Spector et al., 1999). How segregated is shape processing from the analysis of color? As with the case of color and category described above, this question can be tested both in terms of anatomical segregation of preferences for shape and color, and functional independence within a region. Do responses within ROIs show an interaction of sensitivity to color and shape?

Following standard practices, we defined shape selectivity as a higher response to intact objects than scrambled objects (p < 0.0001) (Malach et al., 1995). The location of cortical regions showing a shape preference was anatomically segregated from the location of color-biased regions for most of the VVP. Figure 5A shows a surface histogram of the voxelwise group overlap map for shape selectivity. The large swath of shape-selective cortex (consistent with published group-derived parcels for LO and pFS, dark blue outlines; Julian et al., 2012) showed little overlap with color-biased cortex (outlined in light blue, Fig. 2A), except for the most anterior region where some overlap of shape and color preferences was evident. This pattern suggests that color and shape processing is segregated in posterior VVP but converges at more anterior stages of processing. To test this hypothesis, we analyzed data from individual subjects. We quantified the response profiles of shape-selective ROIs (LO and pFS) and the color ROIs in each subject. Figure 5B shows the average PSC across subjects to intact (O) and scrambled objects (SO), presented with color and without color. Two-factor ANOVAs on stimulus shape (Intact/Scrambled) × color (Chromatic/Achromatic) were run separately on Pc, Cc, and LO. Because Ac overlapped slightly with pFS in 5 of 10 subjects (i.e., the ROIs comprised common voxels), quantification of the responses of these regions was not independent of each other and will be discussed in the next section. As expected, Pc and Cc showed main effects of higher responses to chromatic than achromatic stimuli (p < 0.0001). Neither region showed a preference for intact over scrambled objects; to the contrary, Pc had a significant effect in

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**Table 1. Two-factor ANOVAs on stimulus category (faces, scenes, objects, and bodies) × color (chromatic/achromatic)**

| Region | Main effect | Interaction |
|--------|-------------|-------------|
| PC     | 0.0001      | 0.0002      |
| CC     | 0.0003      | 0.003       |
| AC     | 0.003       | 0.0001      |
| OFA    | 0.02        | 0.004       |
| FFA    | 0.03        | 0.007       |
| ATL    | 0.07        | 0.002       |
| PPA    | 0.15        | 0.003       |
| PSTS   | 0.27        | 0.008       |

Significant effects are indicated in bold; p-values < 0.0005 are indicated as 0 (2-way ANOVAs, color × category).
than LO (i.e., the posterior and central color-biased were more color selective posterior color ROIs with LO. This analysis confirmed that both ran two three-factor ANOVAs, each contrasting one of the two LO, and if LO was more shape selective than the color ROIs, we plot). To test whether the color ROIs were more color biased than pairwise ANOVA) and LO was more shape selective than Pc and (yellow lines in horizontal sections show the location of the sagittal section). B, Random-effects analysis (n = 13); other conventions as for A. C, Quantification of inhomogeneity along the posterior–anterior band of color activations. Two ROIs (schematic, black circles) spatially intermediate to the color-biased regions (white circles) were defined in each subject: iPCc (intermediate to Pc and Cc) and iCAc (intermediate to Cc and Ac). Bar plots show each region’s mean percent signal change for chromatic stimuli (cyan bar) and achromatic stimuli (gray bar) (pooled across faces, bodies, scenes, objects, and scrambled objects) (error bars show within-subject SE). Color-biased regions (top row) were more color-biased than intermediate regions (bottom row) (p-values < 0.003, pairwise 3-way ANOVAs on ROI × color × stimulus class, each contrasting an intermediate region with one of its neighboring color-biased regions).

The opposite direction. Conversely, LO showed a significant preference for intact shape (p < 0.0001), but only a small preference for chromatic over achromatic stimuli (p = 0.003), which was driven mostly by the scrambled object condition (see Fig. 5B, bar plot). To test whether the color ROIs were more color biased than LO, and if LO was more shape selective than the color ROIs, we ran two three-factor ANOVAs, each contrasting one of the two posterior color ROIs with LO. This analysis confirmed that both the posterior and central color-biased were more color selective than LO (i.e., p < 0.001 for the interaction of ROI × color in each pairwise ANOVA) and LO was more shape selective than Pc and Cc (i.e., p < 0.001 for the interaction of ROI × shape in each pairwise ANOVA). These analyses confirm the dissociation of color and shape preferences among the posterior color-biased regions (Pc, Cc) and LO. This dissociation can also be appreciated in Figure 5C, which plots the color bias as a function of the shape bias for all the ROIs. In this plot, the three color-biased regions are located in the upper half of the plot and are clearly separated by a horizontal line from the other regions; the shape-biased regions (notably LO) are located on the right of the plot and could be separated by a vertical line from the color-biased regions.

We also tested whether color preferences depended on the presence of shape information or vice versa. This hypothesis predicts a superadditive interaction in which the presence of both color and shape information in a stimulus has a greater effect on the BOLD response than the sum of the two main effects. Contrary to this prediction, although all three regions (Pc, Cc, and LO) showed significant interactions of color and shape (p < 0.03 for the interaction of color × shape), these interactions went in the opposite direction, reflecting a greater sensitivity to color in scrambled rather than intact shapes (presumably because of the larger number of color edges present in the scrambled images).

Together, these results show that the processing of color and shape occupy largely segregated portions of posterior VVP. Next we turn to the more anterior portions of the VVP, including color region Ac, which showed evidence for modulation by both color and shape.

Anterior color-biased region
We defined Ac in each of the 10 subjects in whom it could be found in at least one hemisphere, and quantified the magnitude of response to each of the 10 conditions in left-out data from the same subjects (Fig. 3). As described above (see “Segmentation of color and category in the VVP”), we found in a two-factor ANOVA of color × stimulus category that this region showed a significant main effect of color, but no main effect of category and no interaction of the two. Next, we investigated whether this region was sensitive to object shape (Fig. 5) by conducting a two-factor ANOVA (intact/scrambled objects × chromatic/achromatic). In addition to the expected main effect of color (p = 0.01), this analysis found a significant main effect of shape (p = 0.01), suggesting a functional difference between this color-biased region and the two posterior color-biased regions (Pc and Cc), where we did not see significant sensitivity to intact shape. A three-factor ANOVA of ROI (Pc, Cc, and Ac) × Shape (intact/scrambled objects) × Color (chromatic/achromatic) confirmed that the anterior region was significantly more shape selective
than the posterior color regions (interaction of ROI × shape, p < 0.0001; driven by the preference for intact objects in Ac but not Pc, pairwise ANOVA, p < 0.0001). We found no interaction between color and shape information in the anterior color region (p = 0.6), suggesting that, although both kinds of information are present, each dimension is processed independently within the region (see Materials and Methods).

In Figure 5C, the color ROIs (cyan triangles) fall along the shape bias dimension in a pattern that mirrors their arrangement on the ventral surface: Pc falls to the left (prefers scrambled objects to intact objects), Ac to the right (some preference for shape), and Cc in between (no shape bias). Whereas the shape-biased regions (LO and pFS, dark blue symbols), face-selective regions (OFA and FFA, gray), and iCAc (gray triangles) all cluster on the anterior region in most subjects whereas many previous studies have found no interaction between color and shape information in the anterior color region (p = 0.6), suggesting that, although both kinds of information are present, each dimension is processed independently within the region (see Materials and Methods).

Despite being reported less often in the literature than the posterior and central color regions, the anterior color region was evident both in the random-effects group analysis (Fig. 4B) and in individual subject data (in 10/13 subjects). Why do we see the anterior region in most subjects whereas many previous studies have not? First, we went to lengths to optimize tSNR in this region (see Materials and Methods), which lies in a part of the brain notorious for signal dropout (due to the presence of the ear canals; Fig. 6). Although tSNR was still low in this part of the brain, we were also able to reliably detect (in 9/13 subjects) the anterior face region ATL, an area that has only been reliably detected (in >50% of subjects) through innovative signal enhancing efforts (Axelrod and Yovel, 2013). Also note that the inhomogeneity observed within the mottled band of color-biased activity cannot be attributed to variations in tSNR because the tSNR is strong between the color-biased regions (iPCc and iCAc). Second, more complex or engaging stimuli may be necessary to elicit activity in

they comprise a rich mixture of low-level visual features, including both luminance and colored components.

Is it possible that color activations extend beyond Ac but are obscured by low tSNR? The flattened surface view (Fig. 6B, bottom) shows the extent of signal coverage obtained presently for an example subject; regions anterior and medial to Ac are particularly impoverished (white asterisks), making it difficult to discern whether color activations span the full length of the temporal lobe (as they appear to in monkeys).

Relationship to monkey organization

The topographic relationship of the face, color, and place activation patterns, and the relative functional dissociation of color-biased and category-biased cortex in humans, mirror results we reported in macaque IT cortex (Lafer-Sousa and Conway, 2013; Verhoef et al., 2015). To further assess the extent to which color and shape information are dissociated in the monkey, we scanned the same animals from the earlier report using the dynamic natural video stimuli used here in humans. We quantified the responses to video stimuli in the set of color-biased ROIs defined in the original report: PLc, CLc, and ALc. The macaque color-biased regions responded similarly to the human color-biased regions: posterior regions (PLc and CLc) showed a color bias but no shape bias, whereas the anterior region ALc showed both a color bias and a shape bias (Fig. 8). For comparison, we quantified responses in the middle face patch ML. ML lacked a striking color bias but showed a shape bias, consistent with the results obtained in human FFA.

In addition to the face-selective regions in the human VVP (OFA, FFA, and ATL), a more dorsal face-selective region has been reported in the pSTS (Collins and Olson, 2014). We detected this region in 11/13 subjects and quantified its responses in independent data (see Fig. 3). A two-factor ANOVA on these
responses showed a main effect of category, a (weak) interaction of color × category, but no main effect of color (Table 1). The putative homology of human ventral regions and macaque IT raises the question, is there a comparable region in macaque monkeys of the human pSTS region? Freiwald and colleagues have proposed a division of the monkey face system into two streams (Voyer and Freiwald, 2013; Fisher and Freiwald, 2015). On this scheme, the ventral face stream in macaques, which follows the ventral lip of the STS and includes patches PL, ML, and AL, corresponds to regions OFA, FFA, and ATL in humans, whereas the dorsal face stream in macaques, residing along the dorsal lip of STS and comprising MD and AF (Fisher and Freiwald, 2015), corresponds to human pSTS (and perhaps aSTS). This scheme would account for the functional dissociation found in both species (Haxby et al., 2000; Pitcher et al., 2011a, 2011b, 2014; Fisher and Freiwald, 2015) between face regions identifiable by responses to static images (PL/ML/AF/AL in macaques; OFA and FFA in humans) versus those that require dynamic stimuli (the dorsal face stream, MD/AF in macaques; pSTS in humans). Figure 9 provides a direct comparison of functional data obtained in a representative monkey and human subject.

Together, these results suggest that the entire face/color/place tripartite cortical system is broadly homologous in macaque monkey and human and, moreover, that this system migrated onto the ventral surface in humans as the cortex expanded over the course of evolution (see Discussion). Establishing this homology greatly empowers studies of high-level vision in humans by making directly relevant to humans the results from invasive studies in macaques.

**Discussion**

Here, we address the functional organization of color processing in the human brain and its relationship to regions implicated in processing faces, objects, and places, using dynamic naturalistic stimuli that robustly drive the VVP. We confirm the existence of a mottled band of color-biased cortex extending along the length of the VVP, within which we identified three main peaks of activation that correspond to previously described color-preferring regions. We found that the series of color-biased regions was sandwiched between bands of face-selective and place-selective regions, mirroring prior results from macaques. In addition, we found substantial anatomical segregation of color-biased regions from those preferring shape or category. Below, we discuss two implications of this work: (1) the homology that it suggests between lateral regions of monkey cortex and ventral regions of human cortex; and (2) the segregation that it reveals of the cortical processing of color, shape, and category.

**Color regions are sandwiched between face- and place-selective regions, as in monkeys**

We identified three color-biased regions: two regions within posterior VVP, referred to here as Pc and Gc (McKeefry and Zeki, 1997; Hadjikhani et al., 1998; Wade et al., 2002; Brewer et al., 2005; Simmons et al., 2007), and an anterior region, Ac (likely corresponding to an unnamed region reported by Simmons et al., 2007). These regions were neatly sandwiched in each subject between face-selective regions on the lateral side and place-selective regions on the medial side. This systematic arrangement of three parallel sets of regions running along the posterior-to-anterior axis of ventral occipitotemporal cortex mirrors the organization of lateral IT in macaque monkey (Fig. 9; Lafer-Sousa and Conway, 2013) and suggests that the entire tripartite cortical structure responsible for face/color/place processing constitutes a “regional homology” (Orban et al., 2004) between the two species.

If this face/color/place system is homologous between humans and macaques, then why is it in such a different location within the brain in the two species? We speculate that this system was pushed onto the ventral surface of the brain over the course of human evolution by the expansion of regions engaged in language and social cognition (white arrow, Fig. 9). The human cortex expanded 10-fold since humans and macaques diverged from a common ancestor, but this expansion was not uniform. Regions involved in communicative and social behaviors, including the temporal parietal junction and the ventrolateral prefrontal cortex, expanded as much as 30-fold, whereas visual areas expanded relatively little (2- to 6-fold; Orban et al., 2004; Hill et al., 2010; Chaplin et al., 2013). Therefore, the identification of neighborhood relationships, like the parallel organization of face/color/place selectivity, may be a better cue to homology than gross anatomical correspondences. Our hypothesized homology of the face/color/place tripartite system between humans and macaques is consistent with prior hypotheses about the face system (Tsao et al., 2008; Voyer and Freiwald, 2013) and could be investigated further using tests of connectivity, cytoarchitecture, and gene expression.

**Dissociation of color from shape and category**

How segregated is the processing of color and shape in the brain? The spatial configuration of colors within a scene has a profound influence on color perception, leading some to argue that color and form are linked insetrically in visual cortical processing, especially in V-1, early in the visual-processing hierarchy (Gegen-
Conversely, color presents computational challenges somewhat distinct from those associated with shape and scene processing. The information provided by color and shape is often independent, for example, in signaling a person’s emotional state independently of face features or the edibility of a piece of fruit (e.g., bananas ripen green to yellow without changing shape); moreover, color and luminance edges are often independent in natural scenes (Hansen and Gegenfurtner, 2009). These observations suggest that color might be processed separately from shape.

We found substantial segregation of color and shape processing in the VVP, consistent with prior evidence for separate pathways in the VVP for processing surface properties (texture and color) versus geometric/shape properties of objects in humans (Cant et al., 2008, 2009; Cant and Goodale, 2009; Cavina-Pratesi et al., 2010). Specifically, the two posterior color-biased regions did not show a preference for intact objects (responding equally, if not more strongly, to their scrambled counterparts; Fig. 5B). Conversely, shape-biased cortex (LO) showed only a weak preference for color (and this was found only for scrambled, not intact, objects). This dissociation of color and form processing accords with prior fMRI studies and with double dissociations reported in neurological patients, in which color perception can be impaired while form perception is spared (e.g., patient MS, Bouvier and Engel, 2006) or vice versa (e.g., patient DF, Humphrey et al., 1994).

Objects can be defined not only by their shape but also by the more abstract category (e.g., faces or places) to which they belong. We found that the VVP showed segregation, not only for shape and color, but also for category and color: each color-biased region showed a higher response for colored than achromatic stimuli for all stimulus categories, and this effect did not differ in magnitude across categories. Further, each color-biased region was more color biased than each category-selective region. This anatomical segregation of regions preferring color versus those preferring object category suggests that category-selective regions rely primarily on luminance information; the lack of color selectivity within shape- or category-selective regions is consistent with the observation that one can recognize most objects from achromatic images (Gregory, 1977). The hypothesized dissociation of color and category processing supported by the present results could be further evaluated using adaptation (Cavina-Pratesi et al., 2010) and multivariate pattern analysis (Brouwer and Heeger, 2013).

Figure 7. Comparison of responses to movie clips versus gratings. Color activity was measured using drifting equiluminant chromatic versus achromatic gratings. A, Comparison of color activations elicited by the movie clips (left) and the gratings (right) on the digitally flattened cortical surface (S5, right hemisphere; white outlines show the location of the color-biased activations localized using the movie clips). The gratings stimulus uncovered regions Pc and Cc, but failed to uncover Ac (black arrow). B, Quantification of the responses to movie clips (top) and gratings (bottom) (n = 3 subjects); cyan bars (chromatic conditions), gray (achromatic conditions). Functional ROIs (Pc, Cc, Ac, OFA, FFA, PPA) were defined with the movie clips; V1, V2, and MT + were defined using an automatic parcellation (Freesurfer). Bars in the top graph were computed by averaging responses to all five categories.

Figure 8. Responses to color and shape in monkeys measured using movie clips. We measured responses to the movie clips (Fig. 1A) in the two animals from an earlier report (Lafer-Sousa and Conway, 2013). Bar plots (top) show the percent signal change in the functional regions defined in that study: PLc, CLc, ALc, and ML. Cyan bars show responses to colored movie clips, and gray bars show responses to achromatic movie clips, either for intact objects (O) or scrambled objects (Sc) (compare with data obtained in humans, Fig. 5B). Scatter plot (bottom) shows color as a function of shape preference; conventions as in Figure SC. The macaque color-biased regions responded similarly to the human color-biased regions; PLc and CLc showed a color bias but no shape preference, whereas ALc showed both a color bias and a shape bias. The macaque middle face patch ML showed a weak color bias and a strong shape bias. Horizontal dashed line separates shape-biased and color-biased regions (as in Figure 5).
Our findings agree with the available data from causal methods. First, the majority (72%) of cortical lesion patients who acquire achromatopsia also acquire prosopagnosia, but rarely object agnosia (8%) (Bouvier and Engel, 2006). This pattern is expected given that cortical lesions are typically larger than the scale of any single functional domain, and color and face regions tend to be adjacent. Second, the few cases of electrical stimulation of the fusiform gyrus in humans produce perceptual effects (Rangarajan et al., 2014) consistent with our findings. For example, stimulation of an electrode overlying FFA of one subject elicited the response “You turned into someone else. Your face metamorphosed. Your nose got saggy and went to the left.” Stimulation of an electrode overlying non-face-selective tissue (posterior to the FFA, likely near Pc) prompted the response, “Light moved across my eyes. It was green, purple, and yellow light together” (Rangarajan et al., 2014).

Overlap of shape and color at more anterior locations of the VVP
The perception of complex scene structure modulates multiple aspects of color perception: color constancy depends on knowledge of 3D scene structure (Bloj et al., 1999; Lafer-Sousa et al., 2015), and pan-field color filling-in fails when scene structure is scrambled (Balas and Sinha, 2007). Moreover, the memory of object colors can modulate object appearance (Hansen et al., 2006; Olkkonen et al., 2010; Witzel and Gegenfurtner, 2013) and facilitate object and scene recognition (Tanaka and Presnell, 1999; Gegenfurtner and Rieger, 2000; Oliva and Schyns, 2000; Nijboer et al., 2007) when color is specifically associated with a given object or scene. These observations suggest that color and shape interact at a high level of visual processing (Tanaka et al., 2001). In support of an interaction of color and shape processing, we found anatomical convergence of color and shape bias in the

Figure 9. Comparison of the functional topography in monkey IT and human VVP suggests a broad homology. Reconstructed cortical surfaces showing the location of face (purple), color (blue), and place (green) ROIs for a macaque monkey (A) and a human (B). Top panels show pial surface views (lateral view in the monkey; ventral view in the human); bottom panels show digitally flattened surfaces exposing functional activation buried in the superior temporal sulcus (sts; prefrontal cortex is cut off). In both species, color-biased regions (in macaque, PLc, CLc, ALc; in human, Pc, Cc and Ac) are sandwiched between face-selective regions (in macaque, PL, ML, AL; in human, OFA, FFA1, FFA2) and place-selective regions (in macaque, mPPA, APP; in human PPA). Both species show evidence for an additional face-processing stream in the sts running parallel to the ventral pathway (in macaque, MF, AF; in human, pSTS, sSTS) (Yovel and Freiwald, 2013; Fisher and Freiwald, 2015). The region of cortex separating the two face-processing streams (white arrow) is relatively larger in humans; this tissue is implicated in communicative and social behaviors and expanded considerably during evolution (Orban et al., 2004; Hill et al., 2010; Chaplin et al., 2013), accounting for the ventral (and medial) displacement of the ventral visual pathway in humans. Macaque data are from Lafer-Sousa and Conway (2013) and Verhoef et al. (2015).
most anterior color-biased region, a region implicated in color knowledge (Simmons et al., 2007). We observed a similar result in macaques, in which the anterior color-biased region (ALC) showed a shape bias whereas the more posterior color-biased regions did not (Fig. 9). This result is consistent with physiological findings of intermingled cell types in the neighborhood of ALC (cells selective for color, shape, or both) (Komatsu and Ideura, 1993; Edwards et al., 2003). We speculate that the color-biased regions of the VVP form a multistage system for the analysis of color, analogous to that observed in the adjacent face system (Lafer-Sousa and Conway, 2013; Tsao, 2014). This hypothesized architecture is consistent with functional differences between posterior and anterior color regions reported here and elsewhere (Simmons et al., 2007) and with observed differences in the color-processing deficits that can result from lesions in different anatomical locations (Miceli et al., 2001). The specific computations that these regions support is largely unknown, but the functional architecture uncovered here provides a roadmap for their investigation.

In sum, our data support two major conclusions. First, humans and macaques show striking similarity in the topographic relationship of color-biased regions, face-selective regions, and place-selective regions, suggestive of a broad interspecies regional homology. Second, color, shape, and category selectivity appear to be substantially segregated throughout much of the VVP. These results invite future work into the computations, connectivity, and developmental origins of the tripartite face/color/place system, a research program made more tractable by the system's apparent homology to macaques.

Notes

Supplemental material for this article is available at http://web.mit.edu/bcs/nklab/rosaStims.shtml. Provided is a high-resolution download of sample stimuli used to localize face, color, place, and shape biased cortex (Movie 1). This material has not been peer reviewed.

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