# Simple shape feature computation across modalities: convergence and divergence between the ventral and dorsal visual streams

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Shape processing, whether by seeing or touching, is pivotal to object recognition and manipulation. Although the low-level signals are initially processed by different modality-specific neural circuits, multimodal responses to object shapes have been reported along both ventral and dorsal visual pathways. To understand this transitional process, we conducted visual and haptic shape perception fMRI experiments to test basic shape features (i.e. curvature and rectilinear) across the visual pathways. Using a combination of region-of-interest-based support vector machine decoding analysis and voxel selection method, we found that the top visual-discriminative voxels in the left occipital cortex (OC) could also classify haptic shape features, and the top haptic-discriminative voxels in the left posterior parietal cortex (PPC) could also classify visual shape features. Furthermore, these voxels could decode shape features in a cross-modal manner, suggesting shared neural computation across visual and haptic modalities. In the univariate analysis, the top haptic-discriminative voxels in the left PPC showed haptic rectilinear feature preference, whereas the top visual-discriminative voxels in the left OC showed no significant shape feature preference in either of the two modalities. Together, these results suggest that mid-level shape features are represented in a modality-independent manner in both the ventral and dorsal streams.

Key words: fMRI; cross-modal; multimodal; haptic; visual cortex; shape.

## Introduction

How the brain computes different types of signals (e.g. optic vs. haptic), initially processed by distinct modality-specific neural systems, into more abstract, modality-independent information contents, is a fundamental question in cognitive neuroscience. One such example is shape computation. Although shape computation in human is commonly considered in the visual processing context for object recognition (e.g. Kourtzi and Kanwisher 2000; Kourtzi et al. 2003; Haushofer et al. 2008), it can also be achieved through the haptic system. Multimodal shape representation has been extensively studied at the object level (Amedi et al. 2001; Amedi et al. 2002; Amedi et al. 2007; Stilla and Sathian 2008; Lacey et al. 2009b; Amedi et al. 2010; Lacey and Sathian 2014; Lee Masson et al. 2016a). Whether the neural computation for earlier-level shape features independent of object context could occur in a multimodal fashion remains uninvestigated and is the question of the current study.

The neural representation of object shape accessed from visual and haptic inputs has been extensively investigated using univariate and multivariate analyses, revealing distributed shape representations in the brain (e.g. Stilla and Sathian 2008; see Lacey and Sathian 2014 for a review; Lee Masson et al. 2016b; Erdogan et al. 2016). For visual shape processing, the involvement of the ventral visual pathway is well established, with extensive evidence from functional neuroimaging, neuropsychological, and neurophysiological studies (Haushofer et al. 2008; Peelen and Caramazza 2012; Bracci and Op de Beeck 2016; see reviews in Grill-Spector et al. 2001; Grill-Spector and Weiner 2014). Within the ventral visual pathway, the lateral occipital cortex (LOC) also showed shape sensitivity when sighted people haptically explored object shapes (Amedi et al. 2001, 2002) and even when early/congenitally blind individuals, who primarily construct object shape knowledge through haptic experiences, process objects through touch, verbal inputs or sound-substitute devices (Amedi et al. 2007, 2010; Striem-Amit et al. 2012; Peelen et al. 2014; Striem-Amit and Amedi 2014; Xu et al. 2022; see reviews in Lacey et al. 2009b; Bi et al. 2016). Although spatially overlapping activations may house functionally independent neural populations (e.g. Wurm and Caramazza 2019), one study further showed cross-modal neural similarity between visual and haptic object representation in the LOC, providing evidence for shared neural representation of objects in this territory (Erdogan et al. 2016).

The dorsal visual pathway along the superior occipital and parietal cortex, whose functionality has been classically assumed to be the processing of spatial- and action-related visual information (Ungerleider and Mishkin 1982; Goodale and Milner 1992), also showed sensitivity to visual object shapes (see Freud et al. 2016 for a review). Neurons selective to simple visual 2D geometric shapes (e.g. triangle, square, circle) were found in the macaque lateral intraparietal area (LIP) (Sereno and Maunsell 1998); the human superior intraparietal sulcus (IPS) represents various object shape features (e.g. absence or presence of a hole, different shape outlines, different combination of simple objects) in a visual short-term memory task (Xu and Chun 2006); human IPS1/IPS2 exhibited adaptation to visual 2D and 3D objects independently of image transformations, mirroring the pattern of LOC (Konen and Kastner 2008; see Xu 2018 for a review). Haptic inputs also elicit shape-related responses in the parietal cortex, with activity strength in the bilateral posterior central sulci and superior parietal lobule (SPL) being modulated by haptic shape complexity (number of curves in a haptic curve counting task, Yang et al. 2021). The neural representational similarity of haptic objects in the SPL and anterior IPS correlated with the object shape similarity (Fabbri et al. 2016; Lee Masson et al. 2016).

In what processing stage(s) does the transition from separate primary sensory cortices to modality-independent neural convergence happen? Recent visual research has focused on mid-level visual shape features (e.g. curvature), which were visual features of intermediate complexity, considered as conjunctions of lowlevel visual features (e.g. luminance, contrast, orientation, spatial frequency) and building blocks that underlie higher-level visual cortex domain organizations (Nasr et al. 2014; Peirce 2015; Long et al. 2018; Tang et al. 2018). The object domain distribution has been shown to be (at least partly) multimodal (e.g. Wolbers et al. 2011; Wang et al. 2015b; van den Hurk et al. 2017). Is it possible that the shared neural structures across visual and haptic modalities are already present at the mid-level shape feature processing stage before the recognition of a holistic object? To test this possibility, we chose curvature/rectilinear lines and sphere/cube plastic models as visual and haptic stimuli, respectively, in consideration that they correspond to more natural basic components for object recognition within each sensory modality respectively. We conducted fMRI experiments in which subjects visually perceived and haptically explored these stimuli and performed analyses to examine the following questions: (i) Are there brain regions showing shape sensitivity (i.e. successful within-modality decoding of curvature vs. rectilinear) for both visual and haptic inputs? (ii) Do these regions represent visual shape and haptic shape with common neural structures (i.e. successful cross-modal decoding for curvature vs. rectilinear)? (iii) Do the multimodal/cross-modal regions have specific shape preferences in a multimodal fashion (i.e. univariate response magnitude differences)?

## Materials and methods Participants

Twenty-one healthy subjects (8 females, age: mean  $\pm$  SD. 23  $\pm$  2 years, range 19–26 years) were recruited among the students at Peking University and Beijing Normal University. Out of the 21 subjects, 14 participated in both the haptic and visual versions of the shape feature experiments, and all analyses were performed on their data. The other seven subjects only took part in the haptic experiment for their unwillingness to participate in the other one. All subjects were right-handed, had normal or corrected-to-normal vision and hearing, and had no history of psychiatric or neurological disease. All of them provided informed consent and received monetary compensation for their participation. All experimental protocols were approved by the Human Subject Review Committee at Peking University.

## Stimuli

The visual stimuli (see Fig. 1B) were made following a previous study looking at visual mid-level features (Nasr et al. 2014).

There were three conditions: rectilinear, curvature, and straightlines (the straight-line condition was not included in the analysis because it did not correspond to either of the haptic shape features and was designed for another study). Each condition included four patterns of arrays. Each pattern of the array contained 40 nonoverlapping shapes distributed randomly across the display. The orientations of arrays for each pattern varied in 22.5° steps, resulting in 16 pictures for each pattern and 64 pictures in total for each condition. The thickness and total length of the lines were equal across conditions.

For the haptic stimuli, basic 3D components with rectilinear and curvature features were chosen—cube and sphere. The cube and sphere models (see Fig. 1A) were made by a 3D printer (JGMAKER, Z-603S, industrial-grade precision). The bases of the models were fixed on an elongated cardboard that was used for passing the stimuli to the subjects during fMRI scanning. To control the range of hand motion, the great circle of the sphere and each surface of the cube were made with the same perimeter of 18 cm, resulting in relatively comparable total touchable surface areas (cube: 101.25 cm<sup>2</sup>, sphere: 102.73 cm<sup>2</sup>) and volumes (cube: 91.13 cm<sup>3</sup>; sphere: 97.94 cm<sup>3</sup>) between the two models.

### Procedures

The haptic experiment (Fig. 1A) consisted of two functional runs, each lasting for 270 s. In each run, after a 12-s-long fixation, the experimenter presented each stimulus to the subjects' right hand in a pseudorandom order. The subjects were instructed to explore each shape when they heard "Start" and stop exploring when they heard "Stop." There were three conditions: the haptic exploration of a cube, a sphere, and the motor control. In the motor control condition, the subjects were asked to pantomime the haptic exploration of the previous stimulus with no stimulus presented to them. Each exploration or pantomiming lasted 6 s and was followed by an 8-s interstimulus interval. Each condition was presented six times per run. During the whole haptic experiment, subjects were blindfolded and wore rubber gloves to prevent the undesired effect of visual input or tactile sensation of surface texture. The haptic experiment was always conducted first to prevent the subjects from recalling and imagining the specific visual stimuli when haptically exploring.

The visual fMRI experiment (Fig. 1B) consisted of four runs. Each run lasted for 300 s and included 12 blocks (three conditions, each presented four times). Each block lasted 16 s, and the interblock interval was 8 s. The order of the blocks was counterbalanced across conditions within each run, and the order of the four runs was counterbalanced across subjects. Within each block, there were 16 trials, consisting of an 800-ms presentation of a picture followed by a 200-ms interval. The subjects were asked to fixate on a cross in the center of the screen for 10 s at the beginning of each run and then perform a one-back task that required them to press a button with their right index finger when the current stimulus was the same as the one right before it. For each condition, there were six blocks where the one-back event appeared once, three blocks where the one-back event appeared twice, and no one-back events in the remaining three blocks. The total number of one-back events was matched across conditions.

### Functional imaging

Images were acquired using a Siemens Prisma 3-T scanner with a 20-channel phase-array head coil at the Imaging Center for MRI Research, Peking University. The participants lay supine with their heads fixed with foam pads to minimize head movement. Functional imaging data were acquired with a simultaneous

### A. Haptic Experiment



**Fig. 1.** Procedure of the haptic experiment (A) and the visual experiment (B). In the haptic experiment, subjects were blindfolded and cued by auditory signals to start and stop exploring the stimulus with their right hand. A baseline motor control condition was included, where the subjects pantomimed the haptic exploration of the former stimulus without any actual stimulus input. In the visual experiment, subjects were asked to perform a one-back task that required them to press a button with their right index finger when the current stimulus was the same as the one immediately before it.

multi-slice (SMS) echo-planar sequence: 62 axial slices, 2.0 mm thickness; 0.3 mm gap; multi-band factor=2; TR=2,000 ms; TE=30 ms; FA=90°; matrix size= $112 \times 112$ ; FoV= $224 \times 224$ ; voxel size= $2 \times 2 \times 2$  mm<sup>3</sup>.

T1-weighted anatomical images were acquired using a 3D MPRAGE sequence: 192 sagittal slices; 1 mm thickness; TR=2,530 ms; TE=2.98 ms; inversion time=1,100 ms; FA=7°; FoV=224 × 256 mm<sup>2</sup>; voxel size= $0.5 \times 0.5 \times 1$  mm<sup>3</sup>, interpolated; matrix size=448 × 512.

### Data preprocessing

Data preprocessing was performed with Statistical Parametric Mapping software (SPM12; http://www.fil.ion.ucl.ac.uk/spm/ software/spm12/) in MATLAB R2020a (MathWorks). The first few volumes (six in the haptic experiment, five in the visual experiment), which were acquired during the initial fixation period, were discarded. The remaining volumes underwent slice timing correction, head motion correction, and removal of low-frequency drift using a temporal high-pass filter (cutoff: 0.008 Hz), and normalization to the Montreal Neurological Institute (MNI) space via the unified segmentation procedure. The functional images were resampled to 2 mm isotropic voxels. For univariate analyses, the data were further spatially smoothed using a 6 mm full width at half maximum (FWHM) Gaussian kernel. The data for multivariate analyses were not smoothed during this step.

## Data analysis

Functional data were analyzed using the general linear model (GLM) in SPM12. GLMs were built for each subject in each run. For all analyses of the visual experiment and the univariate analysis of the haptic experiment, each condition was included as a predictor of interest in each GLM. Thus, for the visual experiment, there were three predictors of interest (i.e. curvature, rectilinear, straight line); for the haptic experiment, there were three predictors of interest (i.e. cube, sphere, motor control). For the multivariate analysis of the haptic experiment, due to the limited number of runs (n=2), the GLM included every single stimulus presentation as a predictor of interest, resulting in 18 predictors of interest (i.e. three conditions × six presentations). In addition, six predictors of no interest corresponding to the head motion parameters were also included in each GLM. These predictors

were convolved with a canonical hemodynamic response function (HRF). The beta images for multivariate analysis underwent a moderate level of smoothing with a 2 mm FWHM Gaussian kernel (Op de Beeck 2010; Gardumi et al. 2016; Hendriks et al. 2017).

The localization of perception regions. We constrained analyses within a whole-brain gray matter mask. The mask was defined as the voxels whose probability was higher than 1/3 in the SPM12 gray matter template and within the cerebral regions  $(1^{#}-90^{#})$  in the Automated Anatomical Labeling (AAL) template, resulting in 122,694 voxels (981,552 mm<sup>3</sup>). We contrasted haptic exploration with motor control to identify the regions related to haptic perception, and contrasted visual feature viewing with fixation to identify the regions related to visual perception. The group-level statistical parametric maps were thresholded at P < 0.05 (voxel-level FDR-corrected, cluster size > 100 voxels). The regions that survived the statistical tests (see Results) were defined as the regions of interest (ROIs) and further underwent multivariate and univariate analyses.

ROI-based SVM decoding. The estimated beta values of the conditions (i.e. curvature and rectilinear) in each perception ROI were extracted, standardized across voxels within each subject and fed into the support vector machine (SVM) classifier implemented in the LIBSVM library (Chang and Lin 2011) in MATLAB. In the within-modality decoding analysis, the classifier was trained and tested to discriminate between curvature and rectilinear features using the leave-one-run-out cross-validation method (see Fig. 2B-C for the schematics), and the accuracy of each ROI were tested against chance-level (i.e. 50%) using a one-sample t-test (one-tailed) with R (R Core Team 2019). In this step, all ROIs underwent both within-haptic and within-visual decoding analyses. Any ROIs whose decoding accuracies were significantly above chance within either of the modalities were considered to be processing the shape features of that modality. As no ROIs showed successful decoding within both haptic and visual modalities (see Results), we conducted a more sensitive top-discriminative-voxelbased decoding analysis.

Top-discriminative-voxel-based decoding. To reduce the potential noise introduced by uninformative voxels, feature selection (Norman et al. 2006) was performed before testing multimodal decoding for each ROI. We began by running a searchlight-based withinmodality decoding across the perception regions. A search sphere was centered on each voxel, with a radius of 10 mm, resulting in 515 voxels. The estimated beta weights within the sphere were extracted and standardized across voxels, and underwent the within-modality decoding analysis using the same method as in the ROI-based decoding. The predicted accuracy was assigned to the centered voxel. Following the searchlight analysis, the voxels with the top N% highest accuracies in each ROI were selected as the most discriminative voxels for individual subjects. Specifically, to determine the number of N, we plotted the within-modal decoding accuracy averaged across subjects against the percentage of top accuracy voxels which varied from 5 to 100% with a step of 5% (Supplementary Fig. 3A-B). The percentage (N%) at which the mean accuracy reached its maximum was selected for subsequent analyses. It should be noticed that selecting the top discriminative voxels to decode within the same modality could have the problem of circular reasoning (Kriegeskorte et al. 2009). Thus, we selected voxels within one modality and tested them in the other modality. Note that here we only focused on regions which already showed successful within-modality decoding in the above ROI-based analysis. Once the decoding accuracy of the selected voxels in an ROI was significantly above chance, the ROI was considered to contain multimodal neural populations, and

cross-modal decoding was then tested with these voxels. Crossvalidation for cross-modal decoding was performed using the data of all runs from one modality as the training set and all runs of the other modality as the testing set, and vice versa (see Fig. 2D for the schematic). The mean accuracy across all subjects was tested against chancel-level (i.e. 50%) using a one-sample t-test (one-tailed) with R.

Shape feature preference analysis. To test whether the multimodal/cross-modal ROIs showed any shape feature preference and whether the preference is multimodal, we also performed univariate contrast analysis for the curvature and rectilinear conditions in both the visual and haptic modalities. The whole-brain preference effects were analyzed using the group-level analysis module in SPM12. The estimated beta values for curvature and rectilinear conditions of the top discriminative voxels identified as multimodal/cross-modal were extracted and averaged across voxels for each ROI and each subject, then were tested using a  $2 \times 2$  (modality  $\times$  feature) repeated-measures ANOVA.

## Results

To test whether and how the ventral and dorsal visual pathways support the visual and haptic processes of the curvature/rectilinear features, the following analyses were performed: (i) multimodal decoding analyses, which tested whether regions showing shape decoding in one modality also show shape decoding in the other modality; (ii) cross-modal decoding analyses, which tested whether regions identified as multimodal also have common neural shape representations across visual and haptic inputs; and (iii) univariate shape feature preference analyses, which explored whether there is any region showing specific shape preference in both modalities.

### Multimodal shape feature decoding results

We first defined haptic and visual perceptual ROIs using univariate contrasts. We identified the bilateral postcentral gyri (postCG, extending to the precentral gyrus on the left side), right precentral gyrus (preCG), bilateral parietal opercula (PO), and the posterior parts of the bilateral supplementary motor areas (pos-SMA) by contrasting haptic exploration with motor control in the haptic experiment. By contrasting shape feature viewing with fixation in the visual experiment, we identified the bilateral occipital cortices (OCs) in the ventral visual pathway, the bilateral posterior parietal cortices (PPCs) within the dorsal visual pathway, as well as bilateral middle frontal gyri (MFG), the anterior parts of the bilateral supplementary motor areas (ant-SMA), right inferior frontal gyrus (IFG), and right insula (see Fig. 2A). We could only observe small overlapping clusters between visual and haptic modalities in the left OC, bilateral middle OCs, and bilateral postCG at a rather lenient threshold (P < 0.05, uncorrected, Supplementary Fig. 1A). No overlapping region between modalities was observed after multiple-comparison correction. We also contrasted haptic exploration with fixation to define the haptic regions. The result is highly similar with those defined by contrasting haptic exploration with motor control after multiple-comparison correction (Supplementary Fig. 1B).

We performed shape feature decoding (i.e. curvature vs. rectilinear) within each modality in the above-identified ROIs. As shown in Fig. 2(E) and Table 1, the bilateral OC could decode visual shape features (left: mean accuracy  $\pm$  SD=0.69 $\pm$ 0.15,  $t_{(13)}$ =4.58, P=0.003; right: mean accuracy  $\pm$  SD=0.64 $\pm$ 0.16,  $t_{(13)}$ =3.31, P=0.01, FDR-corrected), whereas the bilateral PPC



**Fig. 2.** Shape feature decoding in perceptual ROIs. (A) the regions of interest (ROIs) defined by univariate activity contrasts at the threshold of P < 0.05, voxel-wise FDR-corrected, cluster size > 100 voxels; (B–D) the analysis schemes of the within-modality and cross-modality SVM decoding. The classifier was trained and tested using a leave-one-run-out cross-validation method for the within-visual (B) and within-haptic (C) decoding. For cross-modal decoding (D), the classifier was trained with the data of all runs from one experiment and was tested with the data from the other, and vice versa; (E) results of ROI-based shape feature decoding of the perceptual ROIs. For each region, the accuracies were averaged across subjects and then tested against the chance-level accuracy (50%) using one-tailed one-sample t-tests. \*P < 0.05, \*\*P < 0.01, #\*P < 0.01, FDR-corrected.

(left: mean accuracy  $\pm$  SD=0.65 $\pm$ 0.16, t<sub>(13)</sub>=3.73, P=0.01; right: mean accuracy  $\pm$  SD=0.64 $\pm$ 0.10, t<sub>(13)</sub>=5.44, P=0.0008, FDR-corrected), bilateral MFG (left: mean accuracy  $\pm$  SD=0.59 $\pm$ 0.10, t<sub>(13)</sub>=3.51, P=0.01; right: mean accuracy  $\pm$  SD=0.59 $\pm$ 0.12, t<sub>(13)</sub>=2.84, P=0.03, FDR-corrected), ant-SMA (mean accuracy  $\pm$  SD=0.57 $\pm$ 0.09, t<sub>(13)</sub>=2.66, P=0.03, FDR-corrected), and bilateral

postCG (left: mean accuracy  $\pm$  SD=0.63 $\pm$ 0.08, t<sub>(13)</sub>=6.27, P=0.0004; right: mean accuracy  $\pm$  SD=0.62 $\pm$ 0.13, t<sub>(13)</sub>=3.33, P=0.01, FDR-corrected) could decode shape features significantly above chance within the haptic modality. None of the ROIs showed significant shape feature decoding in both modalities at the entire ROI level.

| Table 1. | ROI-based | decoding | results |
|----------|-----------|----------|---------|
|----------|-----------|----------|---------|

|                  | Within visual modality            | r      |                         | Within haptic modality |                   |                         |  |
|------------------|-----------------------------------|--------|-------------------------|------------------------|-------------------|-------------------------|--|
| ROI              | Mean Accuracy $\pm$ SD $T_{(13)}$ |        | P-value (FDR-corrected) | Mean accuracy $\pm$ SD | T <sub>(13)</sub> | P-value (FDR-corrected) |  |
| Visually defined | d ROIs                            |        |                         |                        |                   |                         |  |
| Left OC          | $0.69 \pm 0.15$                   | 4.58** | 0.003                   | $0.53 \pm 0.07$        | 1.46              | 0.24                    |  |
| Right OC         | $0.64 \pm 0.16$                   | 3.31*  | 0.01                    | $0.51 \pm 0.08$        | 0.66              | 0.43                    |  |
| Left PPC         | $0.57 \pm 0.19$                   | 1.42   | 0.24                    | $0.65 \pm 0.16$        | 3.73**            | 0.01                    |  |
| Right PPC        | $0.51 \pm 0.17$                   | 0.19   | 0.57                    | $0.64 \pm 0.10$        | 5.44***           | 0.0008                  |  |
| Left MFG         | $0.54 \pm 0.19$                   | 0.72   | 0.43                    | $0.59 \pm 0.10$        | 3.51*             | 0.01                    |  |
| Right MFG        | $0.48 \pm 0.15$                   | -0.46  | 0.81                    | $0.59 \pm 0.12$        | 2.84*             | 0.03                    |  |
| Right IFG        | $0.50 \pm 0.10$                   | 0      | 0.63                    | $0.51 \pm 0.14$        | 0.24              | 0.57                    |  |
| Right insula     | $0.46 \pm 0.14$                   | -0.94  | 0.91                    | $0.48 \pm 0.10$        | -0.79             | 0.90                    |  |
| SMA-ant          | $0.51 \pm 0.22$                   | 0.15   | 0.57                    | $0.57 \pm 0.09$        | 2.66*             | 0.03                    |  |
| Haptically defin | ned ROIs                          |        |                         |                        |                   |                         |  |
| Left postCG      | $0.54 \pm 0.19$                   | 0.72   | 0.43                    | $0.63 \pm 0.08$        | 6.27***           | 0.0004                  |  |
| Right postCG     | $0.54 \pm 0.17$                   | 0.77   | 0.43                    | $0.62 \pm 0.13$        | 3.33*             | 0.01                    |  |
| Left PO          | $0.41 \pm 0.13$                   | -2.5   | 0.99                    | $0.51 \pm 0.10$        | 0.45              | 0.50                    |  |
| Right PO         | $0.46 \pm 0.14$                   | -1.16  | 0.93                    | $0.53 \pm 0.11$        | 0.99              | 0.36                    |  |
| Right preCG      | $0.56 \pm 0.18$                   | 1.34   | 0.24                    | $0.46 \pm 0.09$        | -1.5              | 0.95                    |  |
| SMA-pos          | $0.53 \pm 0.16$                   | 0.61   | 0.44                    | $0.53 \pm 0.08$        | 1.35              | 0.24                    |  |

\*P < 0.05, \*\*P < 0.01, \*\*\*P < 0.001

Considering that uninformative voxels can induce noise and reduce classifier performance (Norman et al. 2006) and thus lead to false negatives, we carried out a further analysis to address this issue. Specifically, to increase the potential signals, we selected the top N% discriminative voxels within each ROI in its dominant modality (i.e. the modality in which ROI-based decoding was significantly above chance at the entire ROI level) and used these voxels to decode the shape features from the other modality for individual subjects (see Materials and Methods). We plotted the decoding accuracies across the full range of top voxel percentages in Supplementary Fig. 3(A-B), and identified the top N% voxels corresponding to the optimal decoding accuracy for the visual/haptic dominant ROIs (see Table 2 for the specific top N% value for each ROI), and then performed the analyses in the other modality on the identified voxels (see Fig. 3A). For the visual-dominant bilateral OC ROIs, only the activity pattern of the visual-discriminative voxels (top 20%) in the left OC could decode haptic shape features significantly above chance (mean accuracy  $\pm$  SD=0.57 $\pm$ 0.11, t<sub>(13)</sub>=2.43, P=0.03, FDR-corrected; see also Table 2). Among the haptic shape featuresensitive ROIs, the haptic-discriminative voxels in the left PPC (top 30%) could decode visual shape features (mean accuracy  $\pm$  SD=0.65  $\pm$  0.18,  $t_{(13)}$ =3.08, P=0.03, FDR-corrected; see also Table 2). Thus, the left OC and PPC top informative voxels could discriminate shape features in both visual and haptic modalities. We also presented the results of the full range of the discriminative voxels (see Fig. 3B). This full-range picture again showed that only the left OC and the left PPC exhibited successful shape feature decoding in both modalities across a relatively large top-voxel percentage range, which indicated that the findings were not specific to any particularly highlighted percentage.

### Cross-modal shape feature decoding results

As the top discriminative voxels in the left OC and left PPC showed multimodal effects, we further tested whether these voxels exhibited shared neural population coding of shape features across visual and haptic modalities, or they coded shape features in an independent way within each modality (Wurm and Caramazza 2019). We used the above-identified top discriminative

voxels to perform the cross-modal decoding analysis. As shown in Fig. 3(A) and Table 2, the top visual-discriminative voxels in left OC and top haptic-discriminative voxels in left PPC could also decode shape features cross-modally (OC: mean accuracy  $\pm$  SD=0.53  $\pm$ 0.06,  $t_{(13)}$ =2.25, P=0.05, FDR-corrected; PPC: mean accuracy  $\pm$  SD=0.52  $\pm$ 0.04,  $t_{(13)}$ =2.13, P=0.04, FDR-corrected). By separating the two ways of cross-modal decoding (i.e. training with the haptic data and testing with the visual data, and vice versa), we found that the successful cross-modal decoding was mainly driven by the visual-to-haptic direction (Fig. 3A, bottom panel; also see Supplementary Fig. 3C for confusion matrices).

We also performed whole-brain searchlight analyses for within-modal and cross-modal decoding (see Supplementary Material for the method). Significant visual shape feature decoding was observed in the bilateral LOC, and haptic shape feature decoding in the bilateral PPC (voxel-level P < 0.001, cluster-level FWE corrected P < 0.05; Supplementary Fig. 4A). However, no overlapping clusters between modalities were observed under any thresholds adjusted for multiple-comparison. No clusters survived multiple-comparison correction for cross-modal shape feature decoding, either. Under a lenient threshold (uncorrected P < 0.05), we found overlapping areas between the two modalities along both ventral and dorsal pathways, including the left OC and left PPC observed as multimodal and cross-modal in the ROI analyses, as well as in the left postCG, right preCG, and bilateral MFG (Supplementary Fig. 4B).

# Univariate results: shape preference across modalities

The previous SVM analyses examined whether neural activity patterns could discriminate between different shape features, without distinguishing the potential functional preference for each shape feature (i.e. curvature vs. rectilinear). Here, we tested whether the shape feature preference was similar or different across visual and haptic modalities in the top discriminative voxels in the left OC and left PPC (Fig. 4A). As shown in Fig. 4(B), in the left OC, there was a significant main effect of modality ( $F_{(1, 13)} = 147.3$ ;  $P = 1.82 \times 10^{-8}$ ;  $\eta^2 = 0.92$ ; stronger responses in visual modality), but neither a significant main effect of

| Table 2. | Top- | disc | rim | ina | tive- | voxel- | based | decoding  | results. |
|----------|------|------|-----|-----|-------|--------|-------|-----------|----------|
|          |      |      |     |     |       |        |       | · · · · · |          |

| ROI               | Decoding modality | Top N% (number) of voxels | Mean accuracy $\pm$ SD | T <sub>(13)</sub> | P-value (FDR-corrected) |
|-------------------|-------------------|---------------------------|------------------------|-------------------|-------------------------|
| Top visual-discri | iminative voxels  |                           |                        |                   |                         |
| Left OC           | Haptic            | 20% (475)                 | $0.57 \pm 0.11$        | 2.43*             | 0.03                    |
| Right OC          | Haptic            | 20% (622)                 | $0.51 \pm 0.07$        | 0.3               | 0.39                    |
| Top haptic-discr  | iminative voxels  |                           |                        |                   |                         |
| ant-SMA           | Visual            | 55% (136)                 | $0.51 \pm 0.21$        | 0.16              | 0.58                    |
| Left PPC          | Visual            | 30% (154)                 | $0.65 \pm 0.18$        | 3.08*             | 0.03                    |
| Left postCG       | Visual            | 30% (603)                 | $0.56 \pm 0.16$        | 1.45              | 0.30                    |
| Left MFG          | Visual            | 60% (184)                 | $0.50 \pm 0.16$        | 0                 | 0.58                    |
| Right PPC         | Visual            | 60% (277)                 | $0.50 \pm 0.10$        | 0                 | 0.58                    |
| Right MFG         | Visual            | 65% (137)                 | $0.53 \pm 0.15$        | -1.47             | 0.92                    |
| Right postCG      | Visual            | 35% (215)                 | $0.46 \pm 0.09$        | 0.67              | 0.58                    |
| Multimodal voxe   | els               |                           |                        |                   |                         |
| Left              | Cross-modal       | 20% (475)                 | $0.53 \pm 0.06$        | 2.25*             | 0.05                    |
| OC                | Visual-to-haptic  |                           | $0.55 \pm 0.09$        | 2.01*             | 0.05                    |
|                   | Haptic-to-visual  |                           | $0.52 \pm 0.05$        | 1.47              | 0.08                    |
| Left PPC          | Cross-modal       | 30% (154)                 | $0.52 \pm 0.04$        | 2.13*             | 0.04                    |
|                   | Visual-to-haptic  |                           | $0.54 \pm 0.07$        | 2.31*             | 0.04                    |
|                   | Haptic-to-visual  |                           | $0.50 \pm 0.07$        | 0                 | 0.50                    |

 $^{*}P < 0.05,\,^{**}P < 0.01,\,^{***}P < 0.001$ 

feature ( $F_{(1, 13)} = 0.03$ ; P = 0.88;  $\eta^2 = 1.88 \times e^{-3}$ ) nor an interaction ( $F_{(1, 13)} = 0.09$ ; P = 0.78;  $\eta^2 = 6.54 \times e^{-3}$ ) was observed. For the left PPC, the main effect of feature ( $F_{(1, 13)} = 23.77$ ;  $P = 3.03 \times 10^{-4}$ ;  $\eta^2 = 0.64$ ; stronger responses to rectilinear features) and modality ( $F_{(1, 13)} = 52.53$ ;  $P = 6.48 \times 10^{-6}$ ;  $\eta^2 = 0.79$ ; preferring visual modality) as well as the interaction between feature and modality ( $F_{(1, 13)} = 16.41$ ;  $P = 1.37 \times 10^{-3}$ ;  $\eta^2 = 0.60$ ) were all significant. The interaction was in the direction of stronger rectilinear preference in the haptic than in the visual experiment. Indeed, post-hoc tests showed that the rectilinear-preference effect was significant in the haptic modality ( $t_{(13)} = 8.29$ ,  $P = 3.02 \times 10^{-6}$ , FDR-corrected), but not in the visual modality ( $t_{(13)} = 1.73$ , P = 0.11, FDR-corrected).

We also ran a voxel-wise contrast analysis across whole-brain for each experiment. No voxels survived multiple-comparison correction in the visual experiment. In the haptic experiment, bilateral SPL and postCG showed significant preference to rectilinear feature (FDR-corrected; see Supplementary Fig. 5 for the visualization of the statistical maps), converging with the ROIwise result.

## Discussion

To investigate where and at what processing stage neural shape computations become multimodal and cross-modal, we conducted visual and haptic fMRI experiments using contrasting shape features (rectilinear vs. curvature). Our results indicate that voxels in the ventral visual region left OC and the dorsal visual region left PPC exhibit multimodal shape representation properties, as they were able to discriminate rectilinear vs. curvature shape features from both visual and haptic inputs, based on at least partly common neural representations as shown by cross-modal decoding success. Additionally, our specific shape feature preference test revealed different relationships between modality and shape pattern across regions, with the PPC preferring the haptic rectilinear shape, whereas OC showing no significant shape preference in either of the two modalities.

The findings that OC and PPC showed shape feature representation in both visual and haptic experiments are generally consistent with previous studies examining object shape processing in these territories across multiple modalities (see Introduction). Importantly, they indicate that sensitivity to shape across modalities is already present at the level of simple shape features without real object contexts, and the positive crossmodal decoding results demonstrate shared neural shape representation across modalities. It is worth noting that the visual and haptic stimuli in our experiments were not identical, given the intrinsic difference of the two sensory systems. However, both types of stimuli included curved vs. rectilinear geometry elements, albeit in an abstract manner. The positive cross-modal decoding effects-classifiers trained on one modality (i.e. visual rectilinear vs. curvature) successfully decoded corresponding features in the other modality (i.e. haptic cube vs. sphere)—in OC and PPC were thus particularly informative, indicating that the representations shared between modalities were rather abstract. The imbalance of cross-modal decoding (i.e. strong visual-to-haptic decoding but insignificant haptic-to-visual decoding) is also informative, indicating that the haptic perception may involve multiple processing components not shared by the visual modality (discussed further below).

More fine-grained location comparison also suggests potentially interesting differences. By plotting the distribution of the top-discriminative voxels in the left OC across subjects (Fig. 4A), we observed that the highly overlapping voxels located at the ventral occipital area, which is consistent with the peak crossmodal cluster in the left OC in the whole-brain searchlight results (peak coordinates: -37, -77, -11; see Supplementary Fig. 4B). This area is more ventral and posterior to the commonly proposed multimodal (object) shape area (LOtv), which has been previously shown to be activated by both visual and haptic perception of objects in sighted people and by haptic object perception in people without visual experience (Amedi et al. 2001, 2002, 2010; Snow et al. 2014; Erdogan et al. 2016; Lee Masson et al. 2016). For the parietal lobe, although the shape representations probed using haptic objects were observed in SPL (Erdogan et al. 2016; Lee Masson et al. 2016; Yang et al. 2021) and IPS (Amedi et al. 2002; Stilla and Sathian 2008; Lacey et al. 2009a; Snow et al. 2014; Fabbri et al. 2016), those probed using visual objects were mainly distributed in IPS1/IPS2/superior IPS (Freud et al. 2017; Konen and Kastner 2008; Xu and Chun 2006; see Xu 2018 for a review). The PPC cluster we observed as multimodal and cross-modal located mainly in the SPL (also see the only parietal cluster showing

#### A. Top-discriminative-voxel-based decoding



### B. Top-discriminative-voxel-based decoding across different numbers of voxels



**Fig. 3.** Results of top-discriminative-voxel-based multimodal and cross-modal decoding. (A) Accuracies of decoding with the optimal number of voxels in each shape feature-sensitive ROI. The top discriminative voxels were defined in individual subjects (see Materials and Methods), and the obtained accuracies were tested against the chance-level accuracy (50%) using one-tailed one-sample t-tests. The cyan boxplots indicate within-haptic decoding with the top visual-discriminative voxels; the orange boxplots indicate within-visual decoding with the top haptic-discriminative voxels; the green boxplots demonstrate cross-modal decoding using the identified multimodal voxels, with different textures indicating different cross-modal directions.  $^{\#}P < 0.1$ ,  $^{*P} < 0.05$ ,  $^{**P} < 0.01$ ,  $^{**P} < 0.001$ , FDR-corrected; (B) multimodal and cross-modal decoding accuracies varying with different numbers of top discriminative voxels in the shape feature-sensitive ROIs. Shadings indicate standard errors.  $^{*P} < 0.05$ ,  $^{**P} < 0.01$ ,  $^{**P} < 0.001$ , uncorrected.

overlap between multimodal and cross-modal decoding in the whole-brain searchlight analysis result in Supplementary Fig. 4B), largely corresponds to some of the previously identified haptic object shape representation region. Further investigation is needed to determine whether these subtle anatomical differences between the previously reported object-shape multimodal clusters and the current shape-feature multimodal/cross-modal clusters reflect meaningful shifts from lower-level shape elements to higher-level object shape representations by comparing different levels of shape processing within the same study. Additionally, we only observed significant cross-modal representation in the left hemisphere, whether this laterality was simply driven



**Fig. 4.** Shape feature preference in the left OC and left PPC. (A) The visualization of the distributions of the top 20% visual-discriminative voxels in the left OC and the top 30% haptic-discriminative voxels in the left PPC, which exhibited both multimodal and cross-modal multivariate representation of shape features. Different colors indicate the number of participants in each voxel; (B) the comparisons of activation strengths of the top discriminative voxels between the shape feature conditions. Boxplots indicate the beta values (condition vs. baseline) of each condition in each experiment. The asterisks indicate whether the effects were significant in the shape  $\times$  modality repeated-measures ANOVA and in the simple main effect analyses (for the left PPC). \*P < 0.05, \*\*P < 0.01, \*\*P < 0.001, n.s., not significant, FDR-corrected.

by the use of the right hand in the current task or reflect underlying neural mechanism of cross-modal representation remains to be investigated.

There are other clusters that have been previously reported to be sensitive to object shape in multiple modalities, such as middle occipital gyrus (MOG, James et al. 2002; Lee Masson et al. 2016). Under the lenient threshold (Supplementary Fig. 4B), MOG indeed showed a trend of sensitivity to both visual and haptic shape discrimination, but no trend of cross-modal decoding. This result pattern may be due to the fact that the shape features were still represented in the modality-specific format but were housed in a similar region, or because the representation was not abstract enough to generalize across curvature/rectilinear in 3D-shapes and 2D line arrays.

Although our target was to examine the multimodal and crossmodal representation for visual and haptic shape features, it is possible that the successful decoding of the haptic modality resulted from different patterns of motor sequences. We tested this by using the classifier trained on the haptic data to predict the motor control (pantomime) condition (see Supplementary Material for the method of haptic-to-pantomime decoding). The results showed that, except for the SMA, all ROIs could classify pantomiming for sphere vs. cube significantly above chance (Supplementary Fig. 6), suggesting that the representations of haptic exploration contain components of specific motor sequences. However, the motor sequence was not sufficient to support the cross-modal decoding, as indicated by the lack of successful visual-to-pantomime decoding result (Supplementary Fig. 7, and see Supplementary Material for the method). As the pantomime condition was intended as a baseline with limited number of trials, without perfectly matching the sphere and cube shape conditions, and always followed the haptic exploration in the current study, further testing based on well-planned experimental design is needed to determine the independent contributions of shape perception and motor sequence in the common neural representation with visual system. The key point here is that the observed cross-modal shape effects between visual and haptic processes cannot be explained by the potential shared components between haptic and motoric processes.

Another interpretation for the cross-modal result worth considering is that visual imagery was involved during haptic perception. The haptic stimuli, by virtue of being 3D, were closer to real objects and might more easily induce visual imagery. The visual shape features could possibly be seen as elements of the visual imagery of haptic 3D stimuli. Thus, the cross-modal effect in the visual cortex might be driven by visual imagery during haptic exploration. However, this possibility is challenged by the absence of visual-to-pantomime decoding result (Supplementary Fig. 7), as the pantomime of haptic exploration could also induce visual imagery. Moreover, previous studies have reported object shape representation in the lateral occipital/occipitotemporal cortices using word stimuli in congenitally blind people, i.e. without visual experience or visual imagery (Peelen et al. 2014; Xu et al. 2022). For mid-level features, Murty et al. (Murty et al. 2020) focused on the fusiform face area and did not observe significant difference between haptically processed cuboid and spheroid stimuli in congenitally blind individuals. More broad exploration of the mid-level feature representation in congenitally blind people to fully take away the contribution of visual imagery is warranted in future studies.

Although both ventral and dorsal pathways showed multimodal and cross-modal shape feature discriminative properties, the univariate activity profiles of the shape feature preference across modalities provide clues about the potential functional differences in the two pathways. The OC multi-modal voxels showed a preference for visual stimuli, with curvature and rectilinear stimuli evoking comparably stronger activity than haptic stimuli. Despite the preference for visual input, shape features could be decoded across modalities based on multivariate activity patterns. This is in line with the classical role of the ventral visual pathway supporting object shape identification (Amedi et al. 2001, 2002; Kourtzi and Kanwisher 2001; Kourtzi et al. 2003; Haushofer et al. 2008), and offers further evidence for the representation of highly abstract shape feature knowledge that is extracted from different sensory modalities and very different geometric configurations. Note that although no significant shape feature preference was observed for the left OC multimodal voxels, we observed a trend of shape feature preference pattern under a lenient threshold (uncorrected P < 0.05) within the visual cortex, with bilateral lateral fusiform gyri showing a trend of stronger responses to curvature, and the very posterior parts of the bilateral LOC preferring rectilinear. These findings were largely in line with previous studies (Nasr et al. 2014; Yue et al. 2014, 2020; Fan et al. 2021). In contrast, although the dorsal pathway also exhibits evidence for (at least partly) shared neural representation of cross-modal shape features, there is an additional functional preference difference. PPC multimodal voxels show an intriguing interaction between modality and shape, with a stronger preference for the rectilinear feature when touching. The stronger preference for rectilinear stimuli in haptic processes here is potentially aligned with one of the primary functionalities of the parietal lobe as processing sensory information for action (reaching and grasping; Binkofski and Buxbaum 2013; Culham et al. 2003; Fabbri et al. 2016; Goodale and Milner 1992; Rizzolatti and Matelli 2003), as objects that humans tend to manipulate (i.e. haptically interact with) associate with more rectilinear features (Fan et al. 2021).

In conclusion, we observed both convergent and different profiles in the ventral and dorsal visual pathways for mid-level geometric shape feature processing. Both streams contained voxels whose neural patterns could distinguish between geometric shape features within both visual and haptic modalities, with sufficient shared neural representation to enable successful crossmodal decoding. Furthermore, the discriminative voxels in the dorsal stream had a preference for rectilinear shape features in the haptic modality, whereas those in the ventral stream had no specific shape feature preference in either modality. Together, these findings suggest that both ventral and dorsal visual streams process mid-level shape features invariant to input modality, albeit potentially for different computational goals (i.e. recognition vs. action).

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# Author contributions

Shuang Tian: Data curation, Formal Analysis, Investigation, Methodology, Resources, Validation, Visualization, Writing original draft, Writing—review and editing. Yuankun Chen: Investigation, Methodology, Resources. Ze Fu: Methodology, Resources. Xiaoying Wang: Conceptualization, Data curation, Formal Analysis, Funding acquisition, Investigation, Methodology, Project administration, Resources, Supervision, Validation, Writing—original draft, Writing—review and editing. Yanchao Bi: Conceptualization, Funding acquisition, Investigation, Methodology, Project administration, Resources, Supervision, Writing—original draft, Writing—review and editing.

## Supplementary material

Supplementary material is available at Cerebral Cortex online.

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# Data availability

All data reported in this study can be accessed by contacting with the corresponding authors.

# References

- Amedi A, Malach R, Hendler T, Peled S, Zohary E. Visuo-haptic objectrelated activation in the ventral visual pathway. Nat Neurosci. 2001:4(3):324–330.
- Amedi A, Jacobson G, Hendler T, Malach R, Zohary E. Convergence of visual and tactile shape processing in the human lateral occipital complex. *Cereb Cortex*. 2002:12(11):1202–1212.
- Amedi A, Stern WM, Camprodon JA, Bermpohl F, Merabet L, Rotman S, Hemond C, Meijer P, Pascual-Leone A. Shape conveyed by visual-to-auditory sensory substitution activates the lateral occipital complex. *Nat Neurosci*. 2007:10(6):687–689.
- Amedi A, Raz N, Azulay H, Malach R, Zohary E. Cortical activity during tactile exploration of objects in blind and sighted humans. Restor Neurol Neurosci. 2010:28(2):143–156.
- Bi Y, Wang X, Caramazza A. Object domain and modality in the ventral visual pathway. *Trends Cogn Sci.* 2016:20(4):282–290.
- Binkofski F, Buxbaum LJ. Two action systems in the human brain. Brain Lang. 2013:127(2):222–229.
- Bracci S, Op de Beeck H. Dissociations and associations between shape and category representations in the two visual pathways. *J Neurosci.* 2016:36(2):432–444.
- Chang C-C, Lin C-J. LIBSVM. ACM Trans Intell Syst Technol. 2011:2(3): 1–27.
- Culham JC, Danckert SL, DeSouza JFX, Gati JS, Menon RS, Goodale MA. Visually guided grasping produces fMRI activation in dorsal but not ventral stream brain areas. *Exp Brain Res.* 2003:153(2): 180–189.
- Erdogan G, Chen Q, Garcea FE, Mahon BZ, Jacobs RA. Multisensory part-based representations of objects in human lateral occipital cortex. J Cogn Neurosci. 2016:28(6):869–881.
- Fabbri S, Stubbs KM, Cusack R, Culham JC. Disentangling representations of object and grasp properties in the human brain. *J Neurosci.* 2016:36(29):7648–7662.
- Fan S, Wang X, Wang X, Wei T, Bi Y. Topography of visual features in the human ventral visual pathway. *Neurosci Bull.* 2021:37(10): 1454–1468.
- Freud E, Plaut DC, Behrmann M. 'What' is happening in the dorsal visual pathway. Trends Cogn Sci. 2016:20(10):773–784.
- Freud E, Culham JC, Plaut DC, Behrmann M. The large-scale organization of shape processing in the ventral and dorsal pathways. *Elife.* 2017:6:e27576.

- Gardumi A, Ivanov D, Hausfeld L, Valente G, Formisano E, Uludağ K. The effect of spatial resolution on decoding accuracy in fMRI multivariate pattern analysis. *NeuroImage*. 2016:132:32–42.
- Goodale MA, Milner AD. Separate visual pathways for perception and action. Trends Neurosci. 1992:15(1):20–25.
- Grill-Spector K, Weiner KS. The functional architecture of the ventral temporal cortex and its role in categorization. Nat Rev Neurosci. 2014:15:536–548.
- Grill-Spector K, Kourtzi Z, Kanwisher N. The lateral occipital complex and its role in object recognition. Vis Res. 2001:41(10–11): 1409–1422.
- Haushofer J, Livingstone MS, Kanwisher N. Multivariate patterns in object-selective cortex dissociate perceptual and physical shape similarity. PLoS Biol. 2008;6(7):1459–1467.
- Hendriks MHA, Daniels N, Pegado F, de Beeck HPO. The effect of spatial smoothing on representational similarity in a simple motor paradigm. *Front Neurol.* 2017:8:1–11.
- van den Hurk J, Van Baelen M, Op de Beeck HP. Development of visual category selectivity in ventral visual cortex does not require visual experience. *Proc Natl Acad Sci.* 2017:114(22):E4501–E4510.
- James TW, Humphrey GK, Gati JS, Servos P, Menon RS, Goodale MA. Haptic study of three-dimensional objects activates extrastriate visual areas. Neuropsychologia. 2002:40(10):1706–1714.
- Konen CS, Kastner S. Two hierarchically organized neural systems for object information in human visual cortex. Nat Neurosci. 2008:11(2):224–231.
- Kourtzi Z, Kanwisher N. Cortical regions involved in perceiving object shape. J Neurosci. 2000:20(9):3310–3318.
- Kourtzi Z, Kanwisher N. Representation of perceived object shape by the human lateral occipital complex. *Science*. 2001:293(5534): 1506–1509.
- Kourtzi Z, Erb M, Grodd W, Bülthoff HH. Representation of the perceived 3-D object shape in the human lateral occipital complex. *Cereb Cortex*. 2003:13(9):911–920.
- Kriegeskorte N, Simmons WK, Bellgowan PS, Baker CI. Circular analysis in systems neuroscience: the dangers of double dipping. Nat Neurosci. 2009:12(5):535–540.
- Lacey S, Sathian K. Visuo-haptic multisensory object recognition, categorization, and representation. Front Psychol. 2014:5: 1–15.
- Lacey S, Flueckiger P, Stilla R, Lava M, Sathian K. Object familiarity modulates the relationship between visual object imagery and haptic shape perception. *NeuroImage*. 2009a:49(3):1977–1990.
- Lacey S, Tal N, Amedi A, Sathian K. A putative model of multisensory object representation. Brain Topogr. 2009b:21(3–4):269–274.
- Lee Masson H, Bulthé J, Op De Beeck HP, Wallraven C. Visual and haptic shape processing in the human brain: unisensory processing, multisensory convergence, and top-down influences. *Cereb Cortex.* 2016:26(8):3402–3412.
- Long B, Yu C-PP, Konkle T. Mid-level visual features underlie the high-level categorical organization of the ventral stream. Proc Natl Acad Sci. 2018:115(38):E9015–E9024.
- Murty NAR, Teng S, Beeler D, Mynick A, Oliva A, Kanwisher N. Visual experience is not necessary for the development of faceselectivity in the lateral fusiform gyrus. Proc Natl Acad Sci. 2020:117(37):23011–23020.
- Nasr S, Echavarria CE, Tootell RBH. Thinking outside the box: rectilinear shapes selectively activate scene-selective cortex. J Neurosci. 2014:34(20):6721–6735.
- Norman KA, Polyn SM, Detre GJ, Haxby JV. Beyond mind-reading: multi-voxel pattern analysis of fMRI data. Trends Cogn Sci. 2006:10(9):424–430.

- Op de Beeck HP. Against hyperacuity in brain reading: spatial smoothing does not hurt multivariate fMRI analyses? *NeuroImage*. 2010:49(3):1943–1948.
- Peelen MV, Caramazza A. Conceptual object representations in human anterior temporal cortex. J Neurosci. 2012:32(45): 15728–15736.
- Peelen MV, He C, Han Z, Caramazza A, Bi Y. Nonvisual and visual object shape representations in occipitotemporal cortex: evidence from congenitally blind and sighted adults. J Neurosci. 2014:34(1):163–170.
- Peirce JW. Understanding mid-level representations in visual processing. J Vis. 2015:15(7):5.
- R Core Team. R: a language and environment for statistical computing. Vol. 3. Vienna: R Foundation for Statistical Computing; 2019
- Rizzolatti G, Matelli M. Two different streams form the dorsal visual system: anatomy and functions. *Exp Brain Res.* 2003:153(2): 146–157.
- Sereno AB, Maunsell JHR. Shape selectivity in primate lateral intraparietal cortex. *Nature*. 1998:395(6701):500–503.
- Snow JC, Strother L, Humphreys GW. Haptic shape processing in visual cortex. J Cogn Neurosci. 2014:26(5):1154–1167.
- Stilla R, Sathian K. Selective visuo-haptic processing of shape and texture. *Hum Brain Mapp.* 2008:29(10):1123–1138.
- Striem-Amit E, Amedi A. Visual cortex extrastriate body-selective area activation in congenitally blind people "seeing" by using sounds. Curr Biol. 2014:24(6):687–692.
- Striem-Amit E, Cohen L, Dehaene S, Amedi A. Reading with sounds: sensory substitution selectively activates the visual word form area in the blind. *Neuron*. 2012:76(3):640–652.
- Tang S, Lee TS, Li M, Zhang Y, Xu Y, Liu F, Teo B, Jiang H. Complex pattern selectivity in macaque primary visual cortex revealed by large-scale two-photon imaging. *Curr Biol.* 2018:28(1):38–48.e3.
- Ungerleider LG, Mishkin M. Two cortical visual systems. Anal Vis Behav. 1982:549–586.
- Wang X, Peelen MV, Han Z, He C, Caramazza A, Bi Y. How visual is the visual cortex? Comparing connectional and functional fingerprints between congenitally blind and sighted individuals. J Neurosci. 2015b:35(36):12545–12559.
- Wolbers T, Klatzky RL, Loomis JM, Wutte MG, Giudice NA. Modalityindependent coding of spatial layout in the human brain. Curr Biol. 2011:21(11):984–989.
- Wurm MF, Caramazza A. Distinct roles of temporal and frontoparietal cortex in representing actions across vision and language. *Nat Commun.* 2019:10(1):289.
- Xu Y. The posterior parietal cortex in adaptive visual processing. Trends Neurosci. 2018:41(11):806–822.
- Xu Y, Chun MM. Dissociable neural mechanisms supporting visual short-term memory for objects. Nature. 2006:440(7080):91–95.
- Xu Y, Vignali L, Sigismondi F, Crepaldi D, Bottini R, Collignon O. Supramodal shape representation in the human brain. bioRxiv. 2022–11. https://doi.org/10.1101/2022.11.22.517472.
- Yang J, Molfese PJ, Yu Y, Handwerker DA, Chen G, Taylor PA, Ejima Y, Wu J, Bandettini PA. Different activation signatures in the primary sensorimotor and higher-level regions for haptic three-dimensional curved surface exploration. *NeuroImage*. 2021:231:117754.
- Yue X, Pourladian IS, Tootell RBH, Ungerleider LG. Curvatureprocessing network in macaque visual cortex. Proc Natl Acad Sci. 2014:111(33):E3467–E3475.
- Yue X, Robert S, Ungerleider LG. Curvature processing in human visual cortical areas. NeuroImage. 2020:222: 117295.