# Stephen M. Fleming, Louise Whiteley, Oliver J. Hulme, Maneesh Sahani and Raymond J. Dolan

J Neurophysiol 103:3238-3247, 2010. First published Mar 31, 2010; doi:10.1152/jn.01084.2009

# You might find this additional information useful...

Supplemental material for this article can be found at:

http://jn.physiology.org/cgi/content/full/jn.01084.2009/DC1

This article cites 63 articles, 21 of which you can access free at:

http://jn.physiology.org/cgi/content/full/103/6/3238#BIBL

Updated information and services including high-resolution figures, can be found at:

http://jn.physiology.org/cgi/content/full/103/6/3238

Additional material and information about Journal of Neurophysiology can be found at:

http://www.the-aps.org/publications/jn

This information is current as of June 10, 2010.

# Effects of Category-Specific Costs on Neural Systems for Perceptual Decision-Making

Stephen M. Fleming,<sup>1</sup> Louise Whiteley,<sup>2</sup> Oliver J. Hulme,<sup>1,3</sup> Maneesh Sahani,<sup>2</sup> and Raymond J. Dolan<sup>1</sup>

<sup>1</sup>Wellcome Trust Centre for Neuroimaging, University College London; <sup>2</sup>Gatsby Computational Neuroscience Unit, University College London; and <sup>3</sup>Laboratory of Neurobiology, UCL Institute of Ophthalmology, London, United Kingdom

Submitted 10 December 2009; accepted in final form 25 March 2010

Fleming SM, Whiteley L, Hulme OJ, Sahani M, Dolan RJ. Effects of category-specific costs on neural systems for perceptual decision-making. J Neurophysiol 103: 3238–3247, 2010. First published March 31, 2010; doi:10.1152/jn.01084.2009. Perceptual judgments are often biased by prospective losses, leading to changes in decision criteria. Little is known about how and where sensory evidence and cost information interact in the brain to influence perceptual categorization. Here we show that prospective losses systematically bias the perception of noisy face-house images. Asymmetries in category-specific cost were associated with enhanced blood-oxygen-level-dependent signal in a frontoparietal network. We observed selective activation of parahippocampal gyrus for changes in category-specific cost in keeping with the hypothesis that loss functions enact a particular task set that is communicated to visual regions. Across subjects, greater shifts in decision criteria were associated with greater activation of the anterior cingulate cortex (ACC). Our results support a hypothesis that costs bias an intermediate representation between perception and action, expressed via general effects on frontal cortex, and selective effects on extrastriate cortex. These findings indicate that asymmetric costs may affect a neural implementation of perceptual decision making in a similar manner to changes in category expectation, constituting a step toward accounting for how prospective losses are flexibly integrated with sensory evidence in the brain.

# INTRODUCTION

Visual perception has long been considered a process of inference about the most likely explanation of the stimulus-of inferring the state of the world most likely to have caused the pattern of photons hitting the retina (Helmholtz 1856). However, in an ecological context, perceptual categorization needs to take into account not just probabilities but also gains and losses (Bohil and Maddox 2001; Davison and Tustin 1978; Kersten et al. 2004). Think of a radiologist trying to diagnose whether a tumor is present or not in an X-ray. The sensory data may only weakly signal the probability of a tumor, but the potential costs of making a false alarm (further investigation of the occasional healthy person) are far less than the costs of missing a real tumor. In these circumstances, the perceptual judgment may be biased by the prospective loss, creating more false alarms than misses. These shifts in decision criteria are clearly important for survival: for example, in the North American forest, brown bears are more dangerous than black bears. If perception is impoverished, it is better to assume a particular bear-like object is a brown bear. Such scenarios are widespread in perception and raise the question of how sensory evidence and prospective losses interact in the brain to bias

Address for reprint requests and other correspondence: S. Fleming, Wellcome Trust Centre for Neuroimaging, University College London, 12 Queen Square, London WC1N 3BG (s.fleming@fil.ion.ucl.ac.uk).

perceptual categorization in humans (Gold and Shadlen 2002; Heekeren et al. 2008).

Evidence from psychophysics suggests that prospective costs have strong effects on human perceptual decision criteria (Green and Swets 1966; Landy et al. 2007; Whiteley and Sahani 2008). Changes in value linked to particular regions of space are thought to alter intermediate representations between sensory coding and motor planning (Liston and Stone 2008) and to modulate spatially selective regions of early visual (Serences 2008) and somatosensory (Pleger et al. 2008) cortex, potentially via recruitment of fast attention-like mechanisms (Maunsell 2004; Serences 2008). However, it is unclear whether costs associated with particular categorical outcomes, such as tumor present/not present, or, as in this study, the presence of a face or house, are similarly mediated via category-sensitive visual areas (Fleming 2009). An alternative suggestion is that losses and gains are taken into account in frontoparietal regions thought to compare category evidence against a particular decision criterion (Heekeren et al. 2004; Ho et al. 2009; Philiastides and Sajda 2007; Philiastides et al. 2006; Pleger et al. 2006; Ploran et al. 2007; Thielscher and Pessoa 2007; Tosoni et al. 2008; but see McKeef and Tong 2007). This suggestion is supported by recent single-unit recording evidence showing that inducing shifts in decision criteria through changing a learned category boundary (the speed of moving dots) modulates neural firing in the frontal eye fields (Ferrera et al. 2009). A third possibility is that changes in the payoff matrix create a particular "task-set" in frontoparietal regions, which then acts to bias category-specific representations in visual cortex (cf. Summerfield et al. 2006a). This suggestion is in accord with the tight linkage between activity in category-specific ventral visual regions and subjective reports of perceiving faces or houses (McKeeff and Tong 2007; Summerfield et al. 2006b) even when the stimulus remains constant (Tong et al. 1998).

To examine how prospective losses bias perceptual categorization, we manipulated the costs associated with visual categories (faces and houses) while obtaining brain data using functional magnetic resonance imaging (fMRI). We predicted that if biases are expressed through changes in classically defined object representations, we should observe asymmetries in the activity of face- and house-selective regions located in fusiform and parahippocampal gyri in inverse proportion to the loss associated with a particular category. Alternatively, if losses solely bias evidence accumulation, effects of category-specific cost may be constrained to fronto-parietal regions known to compare evidence for perceptual decisions. Finally, if the "task set" hypothesis is correct, we would predict that the

categorical decision emerges out of an interaction between fronto-parietal and ventral visual activity.

#### METHODS

# Subjects

Nineteen right-handed subjects participated in the psychophysics session (7 male; 19–44 yr; mean age, 25.0 yr). All had normal or corrected-to-normal vision, and no history of psychological or neurological illness. Of these participants, 16 were scanned. One participant was excluded at this stage due to a change of response strategy in the scanner that led them to disregard the face/house image, leaving 15 subjects (5 male; 19–27 yr; mean age, 23.9 yr) in the analysis. Samples of this size have been shown to be highly sensitive to true underlying effects in a classical inference framework, assuming, as in the present study, a relatively large number of scans per subject (Friston et al. 2002). The study was approved by the Institute of Neurology (University College London) Research Ethics Committee.

#### Stimuli

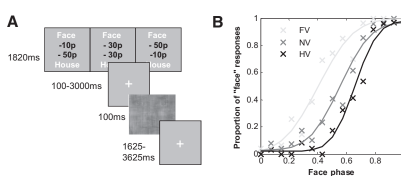
We used 10 neutral faces (5 male, 5 female) from the Karolinska Directed Emotional Faces face set (Lundqvist et al. 1998) and 10 houses (photographed by the 1st author). The stimuli were all cropped to be of equal size and converted to grayscale. To create a stimulus continuum, we adapted a technique used by Heekeren et al. (2004). Fourier transforms (FT) of each image were computed, producing 20 magnitude and 20 phase matrices. The average magnitude of all house and face stimuli was then stored. On each trial, a linear combination of one randomly selected house and face phase matrix was computed, plus a constant proportion (0.35) of a stored white noise matrix (see Supplementary Fig. S1). The resulting phase matrix was then recombined with the average magnitude matrix of the whole stimulus set using an inverse FT. Finally each image was normalized to have average luminance equal to that of the screen background and constant RMS contrast.

Face/house images were presented for 100 ms on a gray background using Cogent 2000 (www.vislab.ucl.ac.uk/cogent.php) running in MATLAB. In the psychophysics experiment, stimuli were presented using a 20.1 in Dell 2001 FP monitor running at a refresh rate of 60 Hz, situated in a dimly lit room. All images subtended 4° of visual angle at a viewing distance of 60 cm. During the fMRI experiment, stimuli were presented using an NEC LT157 LCD projector viewed by subjects via an adjustable mirror. At the beginning of each scanning session, a custom-written Cogent routine adjusted stimulus size and position to match that used in the psychophysics.

## **Psychophysics**

The experiment was divided into two separate sessions. The first session involved acquiring psychophysics data outside the scanner;

<sup>&</sup>lt;sup>1</sup> The online version of this article contains supplemental data.



the second session repeated the same task during fMRI data acquisition. Participants were not informed of the image continuum but were instead asked to categorize each briefly presented stimulus as either a noisy face or house. Participants found this task natural and were unaware of any blend between the two image categories when debriefed. Before introducing a monetary component to the task, each participant completed 540 trials of simple face/house discrimination using the same stimulus timings as in the main experiment. Face and house responses were made using left and right-hand key presses respectively. There were 15 stimulus levels, spaced in equal steps from 100% house to 100% face phase, enabling us to plot out each individual's psychometric function. The point of subjective equality (PSE) for each subject was then used to define face and house categories for the category-specific cost task.

The category-specific cost task involved further face/house discrimination under asymmetric losses for incorrect responses. There were three levels of the cost factor: face value (FV; -50p for an incorrect "house" response, -10p for an incorrect "face" response), neutral value (NV; -30p for an incorrect house response, -30p for an incorrect face response), and house value (HV; -10p for an incorrect house response, -50p for an incorrect face response). Before each block of trials, subjects were given an endowment of £10 and informed that they would keep any money they did not lose on the task. We used losses rather than gains as for the small amounts of money used here, losses were hypothesized to engender a greater behavioral impact on decision criteria than gains (Kahneman and Tversky 1979). Cumulative feedback screens displaying the current total were provided only every 15 trials to avoid incremental learning of decision strategy via trial-by-trial adjustments (Whiteley and Sahani 2008).

Image phase (15 levels) was randomized and orthogonal to the cost factor, which was signaled to participants prior to the face/house stimulus on every trial (Fig. 1A). The cost level changed every two trials. Subjects completed nine experimental blocks of 140 trials each, spanning a single session lasting around 3 h including breaks. Note that when the penalty for answering house incorrectly is greater, a reasonable strategy is to answer face more often when uncertain of the answer.

# fMRI experiment

The fMRI experiment took place within a week of the psychophysics and employed the same task with minor alterations. Subjects completed four runs of 105 trials. The initial endowment for each block was £12, and feedback was given every 10 trials. Each trial began with a cost cue presented for 1,820 ms, followed by a variable interval of 100–3,000 ms during which a fixation cross was presented. The face/house image was then presented for 100 ms, and subjects were able to respond immediately following the onset of the face-house image. Following the offset of the face-house image, a fixation cross was presented for a variable interval of 1,625–3,625 ms prior to the start of the next trial. The buttons indicating face and house responses were switched halfway through the session so that each

FIG. 1. Perceptual decision task and example psychophysics data. *A*: experimental procedure. Subjects viewed a cost signal screen informing them of the potential losses for an incorrect face or house categorization at the start of each trial. They were then asked to categorize an image randomly drawn from the face-house phase continuum as a "face" or "house." Timings shown are for the fMRI experiment. *B*: illustrative psychophysics data from 1 subject (*LB*). Crosses show choice probabilities for each stimulus phase and cost combination; lines show psychometric curves fit to the data.

subject made face and house decisions with both left and right button presses. To avoid switch costs, a short training run was given with the new response mapping without any imaging data being collected.

Stimuli were presented in a permuted randomized fashion, so that the full phase range was covered every seven trials. Similarly, the three cost levels were cycled every six trials (changing every 2 trials, as in the out-of-scanner psychophysics), while keeping stimulus phase and cost orthogonal. This cycling over ~30 s was chosen to match the filter properties of the canonical hemodynamic response function (HRF), maximizing power for estimating the cost- and stimulus-related parameters in our event-related analysis.

# fMRI acquisition

Images were acquired using a 3T Allegra scanner (Siemens, Erlangen, Germany). BOLD-sensitive functional images were acquired using a gradient-echo EPI sequence (48 transverse slices; TR, 3.12 s; TE, 65 ms;  $3 \times 3$  mm in-plane resolution; 2 mm slice thickness; 1 mm gap between adjacent slices; z-shim, +0.6 mT/m; positive phase encoding direction; slice tilt,  $-45^{\circ}$ ) optimized for detecting changes in the parahippocampal region and fusiform gyrus (Weiskopf et al. 2006). Four runs of 213 volumes were collected for each subject, followed by a T1-weighted anatomical scan and local field maps.

## Behavioral data analysis

Subjects' psychophysical responses outside the scanner were modeled using a cumulative normal psychometric function incorporating a random lapse term (Wichmann and Hill 2001), assuming binomial response counts (see Whiteley and Sahani 2008 for full details of the mathematical model). The curve for each cost condition (indexed by j) had three free parameters: the mean  $(\mu_j)$  reflecting the PSE, the slope  $(\rho_j)$  reflecting a subject's uncertainty over the whole stimulus range, and the lapse rate  $(\varepsilon_j)$  reflecting motor errors and lapses of attention. In the equation below,  $CP_{ij}$  gives the probability of answering face for each given stimulus phase combination,  $x_i$ , in the jth cost condition

$$CP_{ij} = (1 - \varepsilon_j) \cdot \frac{1 + erf\left[\sqrt{\pi \cdot \rho_j \cdot (x_i - \mu_j)}\right]}{2} + \frac{I}{2}\varepsilon_j$$

where

$$\operatorname{erf}(z) = \frac{2}{\sqrt{\pi}} \int_{0}^{z} e^{-t^{\tau}} dt$$

We used gradient descent algorithms to find the parameter values that produced optimal curve fits to the observed data. We additionally implemented a Bayesian model comparison to determine whether sharing each of the parameters  $\mu$ ,  $\rho$ , and  $\varepsilon$  between cost conditions gave better fits to the data than allowing each to be optimized separately. We fitted curves to the in-scanner data in the same manner as for the psychophysics data for visualization purposes.

To define category uncertainty for a given stimulus phase, we rectified each individual's psychometric function under the neutral cost condition around the PSE and normalized the result such that the range varied from 0 to 1.

$$U_i = \frac{\left| \left| 10.5 - \mathrm{CP}_i \right| - 0.5 \right|}{0.5}$$

Following Grinband et al. (2006), this procedure defines the PSE as having maximal categorical uncertainty and 100% house/face phase as having minimal categorical uncertainty. Note that this use of the term "uncertainty" refers to the difficulty of categorization for a particular face-house phase composition as opposed to the overall uncertainty about the task expressed in the slope of the psychometric function across the full phase axis. The latter might be expected to

change with, for example, practice, stimulus duration, or lighting conditions.

The psychometric function fits to the in-scanner data were not a robust basis for inference given the lower number of trials per data point compared with the psychophysics session. Consequently, we conducted further behavioral analysis using the framework of signal detection theory (SDT) (Green and Swets 1966). Stimuli were classified as being faces or houses, depending on which side of the PSE they fell, yielding a classic  $2 \times 2$  stimulus-response table for each cost condition. This approach implicitly approximates the stimulus continuum as being drawn from two overlapping Gaussian distributions, one for each category. This allowed us to compute subject-specific measures of sensitivity (d') and criterion (c) separately for each cost condition. Despite being a cruder measure of behavioral performance than the psychometric function fitting described in the preceding text, this method provides a useful index of whether cost primarily affects sensory discrimination or decision/response criteria (Macmillan and Creelman 2005) and circumvents the problem of fewer trials in the scanner leading to unreliable psychometric function fits.

#### fMRI data preprocessing and analysis

Functional data were analyzed using SPM5 (Statistical Parametric Mapping; www.fil.ion.ucl.ac.uk/spm). The first five volumes of each run were discarded to allow for T1 equilibration. Using the FieldMap toolbox (Andersson et al. 2001), field maps were estimated from the phase difference between the images acquired at the short and long TE. The EPI images were then realigned and unwarped using the created field map, and slice-timing correction applied to align each voxel's time series to the acquisition time of the middle slice. Each subject's T1 image was segmented into gray matter, white matter and cerebrospinal fluid (CSF), and the segmentation parameters were used to warp the T1 image to the SPM Montreal Neurological Institute template. The resulting normalization parameters were then applied to the functional data. Finally, the normalized images were spatially smoothed using an isotropic 8 mm full-width half-maximum Gaussian kernel.

fMRI time series were regressed onto a composite general linear model (GLM) containing delta (stick) functions representing the onsets of the cost cue, stimulus, response, and cumulative feedback (see Supplementary Table S1). These delta functions were convolved with the canonical HRF, and low-frequency drifts were excluded with a high-pass filter (128 s cutoff). Short-term temporal autocorrelations were modeled using an AR(1) process. The stimulus delta functions were separated into three regressors dependent on the cost condition on each trial (FV, face value; NV, neutral value; and HV, house value). Each stimulus onset was parametrically modulated by two subject-specific functions. The first was the choice probability (CP) curve fitted to the out-of-scanner psychophysics data in the neutral value condition. The second was the categorical uncertainty function (U), again derived from the out-of-scanner psychophysics data, and orthogonalized with respect to choice probability (see preceding text for mathematical definitions). The cumulative feedback stick function was also modulated with the amount of money lost on the previous 10 trials. To investigate interactions of value and response hand, the response delta function was separated by cost, decision and response hand, giving a 3 (cost; FV vs. NV vs. HV) × 2 (decision; face vs. house) × 2 (response; left vs. right) factorial combination. Motion correction regressors estimated from the realignment procedure were entered as covariates of no interest.

# Statistical inference

Statistical significance was assessed using linear compounds of the model parameters (regression coefficients of the trial-specific stimulus functions in the preceding text) for each subject. These contrast images were then entered into a second-level random effects analysis

using a one-sample t-test against zero to assess group-level significance. Cluster-based statistics (Friston et al. 1994) were used to define significant activations based both on their intensity and spatial extent. Clusters were defined using a height threshold of P < 0.001 and corrected for multiple comparisons across the whole brain using family-wise error correction (FWE) and a threshold of P < 0.05. Images are displayed at the cluster-defining threshold of P < 0.001using MRIcron (http://www.sph.sc.edu/comd/rorden/mricron/). Smallvolume correction (SVC) was applied to category-specific responses by using anatomical masks for fusiform and parahippocampal gyri as specified in the PickAtlas toolbox (Maldjian et al. 2003). Percentage signal change was extracted from clusters of interest for further analysis by averaging over subjects and sessions using MarsBar (Brett et al. 2002). Estimated time courses within clusters are plotted at seven TRs following stimulus onset using a finite impulse response (FIR) model. We note that time courses are plotted for illustration purposes only, inference having first been carried out using appropriate adjustments for multiple comparisons within SPM.

#### RESULTS

We asked subjects to categorize randomly presented noisy images as either faces or houses (Fig. 1A and Supplementary Fig. S1). Before each block of trials, subjects received a financial endowment and were informed that they could keep whatever amount they did not lose in penalties for incorrect answers. On each trial, variable monetary losses were imposed for *incorrect* face and house decisions: either the face response had lower cost (FV condition), the house response had the lower cost (HV condition), or they were balanced (NV condition). In our fMRI experiment, subjects performed the perceptual categorization task inside the scanner, enabling us to obtain measures of regional brain activity associated with task performance over time. Face and house responses were made with both left and right button presses, allowing us to further investigate whether changes in cost interacted with responsespecific brain activity, by decoupling the latter from the decision category.

Behavioral results—psychophysics

Subjects' average PSE in the face-house discrimination pretest was  $53.9 \pm 9.15\%$  face phase. Categorization probability data from a representative subject's psychophysical results are shown in Fig. 1B. To explore the effects of asymmetric cost on choice probability, we fit psychometric functions to the data with either shared or separate mean  $(\mu_j)$ , slope  $(\rho_j)$ , and lapse rate  $(\varepsilon_j)$  parameters for the three cost conditions (indexed by j). We then carried out Bayesian model comparison, thereby revealing which of the eight possible parameter structures (single vs. shared mean  $\times$  single vs. shared slope  $\times$  single vs. shared lapse rate) best accounted for the effects of manipulating asymmetric cost on choice probabilities.

All subjects consistently shifted their responses toward the category carrying lower cost as expected (Fig. 2B and Supplementary Fig. S2). Paired-sample t-test confirmed that average shifts were significantly different from NV for both FV  $[t_{(18)} =$ 5.95, P < 0.0001] and HV trials [ $t_{(18)} = 4.98$ , P < 0.001]. There were also small, but significant, differences in psychometric function slope between cost conditions [white markers in Fig. 2B; 1-way ANOVA, significant effect of cost: F(2,36) =4.61, P < 0.05]. Consistent with these results, Fig. 2A shows that the model with both separate means and slopes was the best model of the data despite the Occam's razor-like penalty for greater model complexity inherent in Bayesian model comparison. However, the magnitude of the difference between the summed log model evidences for shared and separate slopes is rather small, making firm conclusions about differences in slope between conditions difficult.

#### Behavioural results—fMRI

As fewer trials precluded fitting reliable psychometric curves to the choice data in the scanner, we carried out a signal detection analysis (Green and Swets 1966) to characterize

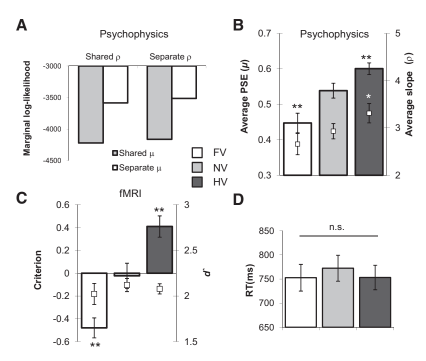
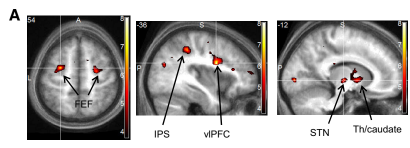


FIG. 2. Behavioral results. A: Bayesian model comparison was used to show that the best model for the psychophysics data were one with separate mean and slope parameters for each cost condition. The chart shows Laplace approximation to the total log marginal likelihood across subjects and across shared and separate error parameters (it seems possible that attentional lapses would vary with cost condition, which does not bear on the hypothesis of interest)—a smaller negative value indicates a better model. Note that each unit difference in log likelihood corresponds to an e-fold ratio of model probabilities. B and C: average parameters of the psychometric function fits to the psychophysics data, n = 19 (B), and a corresponding signal detection analysis of the in-scanner data, n = 15 (C). Bars represent the point of subjective equality (PSE)/criterion in face, neutral, and house value (FV, NV, and HV, respectively) conditions. White markers indicate the average slope/d' parameter in each cost condition for comparison. D: mean RTs averaged across changes in stimulus information for each cost condition. In all panels, error bars denote SEM; \*\*, P < 0.005; \*, P < 0.05 in comparison with NV.



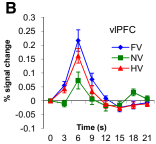


FIG. 3. Cost-related brain activity. A: axial (z = 54) and saggital (x = -36 and -12) sections showing brain activations reflecting the main effect of asymmetric cost [(FV + HV) >2NV], averaged over category. Shown are significant clusters in left ventrolateral prefrontal cortex (vIPFC), intraparietal sulcus (IPS), bilateral frontal eye fields (FEFs), and subcortical regions (STN, subthalamic nucleus region; Th, thalamus; see also Table 1). Labeled activations are significant at P < 0.05, cluster family-wise error (FWE) whole-brain corrected. B: hemodynamic response time courses aligned to stimulus onset for the 3 different cost conditions, plotted for the significant cluster in vIPFC.

in-scanner behavior, collapsing stimuli into either face or house categories based on each individual subject's PSE. This analysis confirmed that asymmetric cost led to deviations of the decision criterion in the predicted direction, relative to the neutral value condition [c; FV,  $t_{(14)} = 5.82$ , P < 0.0001; HV,  $t_{(14)} = 5.78$ , P < 0.0001] but did not change category discriminability [d'; F(2,28) = 0.41, P > 0.5; Fig. 2C].

Importantly, mean reaction times (RTs) did not differ across cost conditions [psychophysics, F(2,36) = 0.70, P > 0.4; in-scanner, F(2,28) = 1.67, P > 0.2], suggesting that any bias-related differences we find in brain activity are not driven by systematic differences in task difficulty (Fig. 2*D*). RT was however significantly correlated with categorical uncertainty (see METHODS and Supplementary Fig. S3; psychophysics, mean  $r = 0.79 \pm 0.092$ , n = 19; in-scanner, mean  $r = 0.56 \pm 0.21$ , n = 15).

#### fMRI results

COST-RELATED REGIONS. We first identified regions involved in processing the extra demand of integrating asymmetric cost by computing the cost > neutral contrast (FV + HV >2NV). A frontoparietal network (Fig. 3 and Table 1) was consistently active for both types of asymmetric cost condition compared with neutral (P < 0.05, whole-brain corrected), suggesting its involvement in the biasing of perceptual decisions as a function

of category-specific cost. In addition to frontoparietal areas, we found increased activity in a cluster spanning the subthalamic nucleus (STN) region, thalamus and caudate nucleus (P < 0.05, whole-brain corrected). Recent modeling work has shown the type of onset function (stick vs. epoch) can affect activation patterns in decision-making tasks even when decision times are short (Grinband et al. 2008). To check that our cost-related activations were robust to the type of onset function used, we constructed a second design matrix in which the decision period was modeled using a variable epoch. Similar activations were observed in both models with the overlap shown in Supplementary Fig. S5.

STIMULUS-SELECTIVE REGIONS. We next identified functional regions of interest (ROIs) sensitive to stimulus category information. In our group analysis, a cluster in right fusiform gyrus (FG; P < 0.05, SVC; Fig. 4A) correlated with the probability that the image was categorized as a face, averaging over cost condition. Conversely, activity correlating with the probability of an image being categorized as a house was expressed in bilateral parahippocampal gyrus (PHG; right, P < 0.05, SVC; left, P < 0.05, SVC; Fig. 4B). No other clusters correlating with the categorical choice probability survived whole-brain correction.

To test our first hypothesis, that category-specific costs directly affect responses in the ventral visual stream, we

TABLE 1. Summary of significant activations for the cost > neutral contrast reported in the main text

Label	Voxels at $P < 0.001$	Peak Z Score	P (Cluster FWE Corrected)	Peak Voxel MNI Coordinates	Laterality
IFG (p. opercularis)	106	4.92	< 0.001	-36, 3, 24	L
FEF	36	4.56	0.004	-27, -3, 54	L
Caudate/thalamus/STN	93	4.28	< 0.001	-15, -18, 0	L
FEF	42	4.10	0.002	26, -9, 54	R
IPS	49	4.26	0.001	-36, -39, 45	L
IFG (p. triangularis)	31	3.94	0.009	-45, 21, 0	L
Insula/putamen	33	3.90	0.007	36, 15, -6	R
pMTG	43	4.04	0.001	-36, -72, 21	L

FWE, family-wise error; MNI, Maltreal Neurological Institute; IFG, inferior frontal gyrus; IPS, intraparietal sulcus; FEF, frontal eye fields; pMTG, posterior middle temporal gyrus; STN, subthalamic nucleus region.

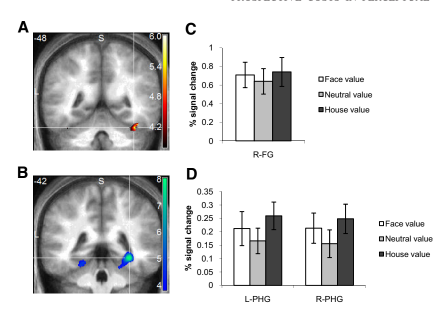


FIG. 4. Stimulus-selective brain activity. A: coronal (y = -48) section showing parametric effects of the probability an image was classified as a face in right fusiform gyrus (FG; 39, -48, -24; z-score = 4.07; P < 0.05, small-volume corrected). B: coronal (y = -42) section showing parametric effects of the probability an image was classified as a house in bilateral parahippocampal gyrus (PHG; left: -21, -42, -15; z-score = 3.68; right: 33, -42, -9; z-score = 5.26; both P < 0.05, small-volume corrected). C and D: percentage signal change as a function of cost condition in stimulus-selective ROIs defined from clusters shown in A and B.

computed the signal change for each cost condition in each stimulus-selective ROI identified in the preceding text. Oneway ANOVAs (FV vs. NV vs. HV) revealed effects of cost on right PHG [F(2,28) = 4.80, P = 0.002] but not left PHG or right FG [F(2,28) < 2.5, P > 0.1]. Further investigation of the pattern of differences in right PHG revealed increases in the HV condition compared with NV [ $t_{(14)} = 3.09$ , P = 0.008], a trend for increases in the FV condition  $[t_{(14)} = 1.77, P = 0.10],$ but no evidence for differences between the category-specific FV and HV conditions [ $t_{(14)} = 1.26$ , P = 0.23]. A similar trend for nonspecific increases under asymmetric cost conditions can be seen in all three ventral visual areas (Fig. 4, C and D), and an omnibus ANOVA in which region was included as a separate factor indicated an overall significant effect of cost  $[F_{(2.28)} = 5.95, P = 0.007]$ . Together, these analyses indicate that asymmetric cost has a general driving effect on ventral stimulus-selective regions, but in a manner that does not appear to discriminate between stimulus categories.

# Category-specific effects of cost

The account presented thus far indicates that categoryselective stimulus information is to some degree represented independently of category-specific biases induced by changes in the payoff matrix. In other words, payoff asymmetries lead to only general, not category-specific, increases in the signal in voxels sensitive to stimulus (face or house) information. However, the mechanism by which asymmetric value information brings about a change in perceptual decision (such as a bias toward responding face) remains unclear. To address this question, we computed the category-specific value contrasts (FV > HV; HV > FV). A cluster in left parahippocampal gyrus (-33, -36, -15; P < 0.05, SVC) responded specifically to increases in house value (HV > FV; decrease in cost for responding house), lateral and anterior to the stimulus-selective cluster we characterized previously (Fig. 5A). No significant clusters were evident in the opposite (FV > HV) contrast even at a liberal (P < 0.01) defining threshold. Together, these findings indicate that category-specific costs exert effects on the ventral visual pathway (at least for the bias toward responding house).

We noted that while the direction of decision criterion shifts under asymmetric cost was consistent across subjects (Fig. 2C), individual differences in the size of this shift were evident in the behavioral data (Fig. 5B). To explore whether subjects who exhibit greater decision criterion shifts also show greater activity within regions that are the putative sources or targets of these shifts, we regressed the category-specific value contrasts (FV > HV; HV > FV) against between-subjects covariates encoding the amount of behavioral bias in the relevant asymmmetric value condition (FV criterion shift; HV criterion shift). Subjects who displayed greater criterion shifts in the HV condition tended to activate the anterior cingulate cortex (ACC; 6, 36, 30) more than subjects who shifted to a lesser degree (Fig. 5, C and D; P < 0.05, whole-brain corrected). Again, as for the simple main effect of FV > HV, no significant correlations were found with individual differences in the FV criterion (P > 0.001, uncorrected).

# Categorical uncertainty

To examine brain regions responsive to categorical uncertainty, we regressed a parameter that essentially measures how close to chance the subject is in deciding whether the stimulus is a face or a house (see METHODS) onto the fMRI signal at the time of choice. Dorsal medial frontal (paracingulate) cortex (dMFC; 6, 12, 51) and right anterior insula (42, 24, -3)showed positive correlations with categorical uncertainty (both P < 0.05, whole-brain corrected; Fig. 6A). We were interested in establishing whether this uncertainty-related BOLD signal was independent or overlapping with the frontoparietal regions found to be active under conditions of asymmetric cost. By exclusively masking the cost > neutral contrast for regions correlating with categorical uncertainty at a liberal (P < 0.05, uncorrected) threshold, we found that left vIPFC, left caudate/ thalamus/STN, and bilateral FEF were specifically active under conditions of asymmetric cost, independent of changes in categorical uncertainty (Supplementary Table S2). Conversely, dMFC activity was independent of changes in category value

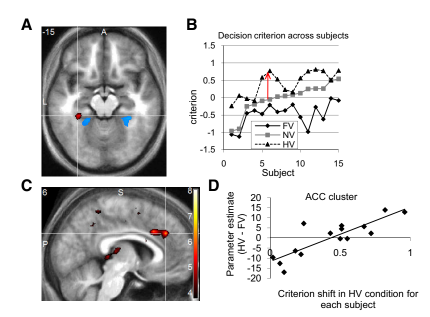


FIG. 5. Category-specific effects of asymmetric cost. A: axial (z = -15) showing the region in left parahippocampal gyrus (red) active during decreased cost (increased value) for houses (HV > FV; -33, -36, -15; z-score = 3.84; P < 0.05, small-volume corrected). Shown in blue are clusters selective for house stimulus information (Fig. 4B) for comparison. B: intersubject variation in decision criteria with subjects ordered by their decision criterion in the NV condition. The arrow shows the difference (extent of behavioral shift under HV) used as a covariate for the relevant contrast testing for HV-specific effects of asymmetric value shown in A. C: saggital (y = 6) and axial (z = 30) sections showing a region in the anterior cingulate (ACC) that shows greater activity in subjects who show greater behavioral shifts in the HV condition (6, 36, 30; z-score = 4.29; P < 0.05, whole-brain corrected). D: averaged HV > FV beta within the ACC cluster shown in C plotted against the criterion shift in the HV condition across subjects. Inference was carried out using appropriate corrections for multiple comparisons in the SPM framework; this plot is simply provided for illustration purposes.

(Fig. 6*B*), indicating a partial dissociation in the brain between regions encoding changes in categorical uncertainty and prospective costs during perceptual categorization. The negative effect of the categorical uncertainty regressor (testing for greater activity in "easy", certain decisions) was seen in ventromedial prefrontal cortex (vmPFC; 9, 33, -6; P < 0.05, whole-brain corrected; Fig. 6*A*). Interestingly, vmPFC was also active in proportion to the amount of money saved at each intermittent feedback screen (Supplementary Fig. S4), suggesting a neural relation between perceptual certainty and reward (cf. Maddox and Bohill 2003).

#### Interaction of cost with motor planning

The previous analyses identified brain regions that responded preferentially to a particular direction of bias (toward responding house). Our design further allowed us to ask whether any bias effects are expressed at the level of the motor system, given that response hand (left or right) was orthogonal

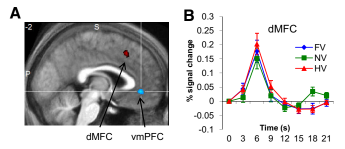


FIG. 6. Effects of categorical uncertainty. A: saggital section (y=-2) showing positive (red) and negative (blue) correlations with a regressor indexing categorical uncertainty (see METHODS). Dorsal medial frontal cortex (dMFC; 6, 12, 51; z score = 3.70; P < 0.05, whole-brain corrected) and insula (not shown; 42, 24, -3; z score = 4.11; P < 0.05, whole-brain corrected) showed positive correlations with uncertainty. A cluster in ventromedial prefrontal cortex (vmPFC) showed increased activity for easier perceptual decisions (6, 36, 30; z score = 4.20; P < 0.05, whole-brain corrected). B: hemodynamic response timecourses for the 3 different cost conditions, plotted for the significant cluster in dMFC. While showing strong correlations with the categorical uncertainty regressor, this region was insensitive to changes in category value (cf. Fig. 3B).

to decision (face or house). Interactions of cost asymmetry with response hand were computed by coding each trial as to whether the left or right hand was assigned to a high or low cost response (face or house) and examining the interaction of cost condition with response hand (see METHODS). No effects were found (P > 0.001, uncorrected), suggesting that the biasing effects of asymmetric value occur upstream of effector-specific response planning.

#### DISCUSSION

Perceptual judgments are often affected by the potential costs associated with different categorizations; for example, during a radiologist's assessment of an X-ray image or deciding whether a dangerous or benign animal is hidden in the trees. Here we investigated the brain mechanisms that integrate prospective costs and sensory evidence during categorization. Our behavioral manipulation systematically biased the perception of a noisy image using asymmetric costs, leading to shifts in decision criteria. These shifts functioned to reduce monetary losses, by biasing decisions toward the category with lower cost when the participant was unsure of the answer. Using fMRI, we asked whether category-specific shifts were reflected by changes in frontoparietal areas known to accumulate evidence leading to perceptual categorization (e.g., Heekeren et al. 2004), in ventral visual regions known to encode categoryspecific information about faces and houses, and/or via recruitment of regions thought to communicate information about the currently relevant task set (Serences et al. 2004; Summerfield et al. 2006a). Our data best fit the latter, "task set" hypothesis. The requirement to integrate asymmetric cost information into perceptual decisions robustly activated a frontoparietal network despite conditions being closely matched for expected value and reaction time. In addition, a cluster in the thalamus/ caudate STN was active under asymmetric cost, consistent with subcortical loops being important for the setting of decision criteria (Fleming et al. 2010; Lauwereyns et al. 2002; Lo and Wang 2006; Simen et al. 2006.) A specific effect on ventral visual areas (parahippocampal gyrus) was found under decreasing cost for houses, anatomically adjacent to the integration of stimulus information. Finally, subjects who showed greater shifts in decision criteria toward houses demonstrated greater activation of the anterior cingulate cortex (ACC), a region thought to be pivotal in the adjustment of decision strategy (Behrens et al. 2009; Botvinick et al. 2001).

# Sources of category-specific bias

The dorsal frontoparietal network active under asymmetric cost is similar to that commonly activated in studies of transient allocation of attention (Corbetta and Shulman 2002; Corbetta et al. 2008; Yantis et al. 2002), and has been recently implicated in the modulation of early visual cortical activity by rewards tied to particular locations in visual space (Serences 2008). It is plausible that changes in category-specific costs co-opt a similar network. Low-level changes in arousal or task difficulty are unlikely to be explanations for this increase in activity as RTs and potential gains/losses were matched across conditions. Instead our findings indicate that frontoparietal (ventrolateral prefrontal cortex, insula, intraparietal sulcus, frontal eye fields) and subcortical (anterior thalamus/STN) regions are recruited when information about payoffs needs to be incorporated into a perceptual decision. Bilateral activation was found at the junction of the precentral and superior frontal sulci, consistent with the location of the FEFs (Lobel et al. 2001), which are known to have a causal role in the modulation of sensory cortex (Ekstrom et al. 2008) and encode shifts in decision criteria (Ferrera et al. 2009). In addition, activation in ventrolateral prefrontal regions including insular cortex is consistent with involvement both in the accumulation of sensory evidence (Ho et al. 2009; Romo et al. 2004) and the incorporation of gains and losses in decision-making (Leon and Shadlen 1999; Watanabe and Sakagami 2007). Indeed the network outlined in the preceding text may be recruited more generally when shifts in decision criteria are induced by manipulations other than asymmetric payoffs. This view is supported by previous findings of modulation of subcomponents of this network when decision criteria are shifted through changes in category boundary—specifically, BOLD signal in anterior thalamus/caudate and insula/vlPFC (Grinband et al. 2006; Li et al. 2009) and single-unit activity in FEF (Ferrera et al. 2009).

The present analyses cannot pin down the source of the bias toward houses and faces as effects of category-specific bias were not observed in the network discussed in the preceding text. However, it is possible that local neural subpopulations within these areas encode biases toward face and house categories. This suggestion is supported by a recent study by Rorie et al. (2010) in monkeys, demonstrating that asymmetric payoffs in a perceptual decision task bias the initial firing rate of individual neurons in the intraparietal sulcus coding for a saccadic response to one of two particular targets. Similarly, using fMRI, distributed patterns in the inferior frontal gyrus/ insula have been found to discriminate the direction of criterion shifts in a visual categorization task (Li et al. 2009). Our finding that the ACC tracks individual differences in the extent of a signed criterion shift is also consistent with categoryspecific payoff information being represented in frontal cortex. Differential effects of house and face cost

When the signal change in stimulus-selective ROIs was calculated, general but not selective effects of asymmetric cost were observed. In contrast, a parahippocampal region anterior to the stimulus-selective areas was significantly more active when houses rather than faces were more valuable. Due to the potential impact of our smoothing and thresholding procedures, we are cautious in attributing separate locations to the stimulus- and cost-sensitive regions of the PHG. However, we note evidence suggesting local separation of task- and stimulus-driven regions using neural stimulation coupled with high resolution imaging (Ekstrom et al. 2008); similarly, differences between stimulus- and task-driven localizations have been reported in the fusiform gyrus during perceptual decision making (Philiastides and Sajda 2007).

Category-specific biases were seen in ventral visual regions for increases in house, but not face, value. Furthermore, across subjects, a correlation with decision criteria was seen in the ACC for bias toward houses but not faces. While we are cautious about overinterpreting null results, we note that previous studies examining attentional and decisional biases toward faces and houses have also found asymmetric effects of the two categories. Specifically, Summerfield et al. (2006b) found that mistaken categorizations of houses as faces were accompanied by increases in fusiform gyrus (FG) activity but that the opposite mistake did not increase PHG responses. In contrast, Serences et al. (2004) found that shifts in object-based attention toward houses recruited parietal and frontal regions to a greater degree than shifts toward faces. Both these findings and the asymmetry in the results of the present study can be reconciled by assuming that subjects have a dominant prior to respond "face." This hypothesis is supported by informal debriefing—some subjects in our study commented that they performed the task by responding "house" whenever evidence for a face was scant (see also Summerfield et al. 2006b). In the case of our data, increases in value for houses would lead to shifts in PHG activity to overcome this implicit prior toward responding face, but the converse may not be necessary. Whether visual phenomenology also changes under such shifts in decision criteria is an open question, one that could potentially be addressed by eliciting detailed reports from subjects under biased and unbiased conditions (Jack and Roepstorff 2002).

# Responses to categorical uncertainty

Consistent with previous reports, we found that activity in dorsal medial frontal (paracingulate) cortex (dMFC) and anterior insula correlates with categorical uncertainty (Grinband et al. 2006; Philiastides and Sajda 2007; Preuschoff et al. 2008). There has been recent debate about the functional role of the medial frontal/paracingulate cortex in perceptual decisions (Heekeren et al. 2008). Here we report preliminary evidence for segregation of networks responding to changes in categorical uncertainty and category value. Dorsal paracingulate activity correlated with increases in categorical uncertainty, independent of changes in value; conversely, the FEFs and caudate/thalamus/STN were specifically active during decisions requiring integration of category value information. As uncertainty was correlated with RT, we are unable to dissociate

the contributions of decision time to these activations (cf. Grinband et al. 2006, 2008). Interestingly, however, the dMFC region lies just dorsal to the ACC, which responded to the degree of decision criterion shift across individuals. Given that such shifts are only required when subjects are uncertain about the sensory data (Maddox and Bohil 2003), the close anatomical relationship between these regions may be optimal for integration of categorical uncertainty during shifts in decision criteria.

In contrast, the opposite contrast (examining brain activity that increases for "easy," certain choices) revealed a cluster in ventromedial prefrontal cortex that was close to that responding to the explicit delivery of monetary reward information. This finding supports recent evidence that the ventromedial prefrontal cortex may signal a perceptual "match" between observed and predicted stimulus information (Summerfield and Koechlin 2008) and suggests that perceptual accuracy itself may act as a reinforcer (Bohil and Maddox 2001). However, in our experiment, this signal could also be related to the ongoing assessment of the expected value of the current decision (Boorman et al. 2009; O'Doherty 2004), as both perceptual accuracy and potential rewards are highly correlated on any given trial.

#### Costs and priors

A recent theoretical suggestion holds that cost functions and priors perform a common role in perceptual inference (Friston et al. 2009), relying on backward communication of expectations. This idea has broad historical precedent in psychology, beginning with the "New Look" school of perception in the 1950s (Bruner 1957; Bruner and Goodman 1947). These studies emphasized the role of needs and desires in altering visual perception and have been echoed in recent studies showing that, for example, being motivated to receive a particular outcome leads to perceptual biases (Balcetis and Dunning 2006). Two elegant studies of changes in categorical priors have demonstrated increased backward connectivity from frontal cortex to stimulus-selective visual areas as a function of changes in expectation (Summerfield and Koechlin 2008; Summerfield et al. 2006), and changes in expectation of face/house stimuli activate similar frontoparietal regions to those identified here (Puri et al. 2009). Further work is needed to explicitly examine the neural overlap between cost- and prior-induced biases in visual decision making. We also note that the manipulation of category value in our study was achieved through the prospect of asymmetric losses in keeping with the psychometric tradition of using loss functions to manipulate behavior (Green and Swets 1966; Landy et al. 2007; Whiteley and Sahani 2008). However, given evidence that the brain may process the prospect of losses and gains differently (Liu et al. 2007; Yacubian et al. 2006), it would be useful to establish whether losses and gains differentially impact on mechanisms for perceptual decision making.

# Summary

To conclude, our findings extend previous reports that costs attributed to perceptual decision outcomes have consistent effects on stimulus categorization with subjects acting to minimize prospective losses. We show that this effect of cost on perceptual decisions is robustly associated with BOLD signal increases in a frontoparietal network, in keeping with the hypothesis that loss functions enact a particular task set. When the cost for responding "house" decreased, we observed selective activation within the parahippocampal gyrus. Across subjects, greater shifts in decision criteria were associated with greater activation of the medial frontal cortex (ACC). These findings are consistent with the hypothesis that asymmetric costs alter an intermediate representation between perception and action, expressed via general effects on frontal cortex, and selective effects on extrastriate cortex. Returning to our radiologist's difficult decision, the implicit costs associated with different diagnoses may set up top-down priors that subtly bias sensorimotor dynamics, leading to more false alarms and less misses. Our results constitute a step toward accounting for how prospective costs are flexibly integrated with sensory evidence during effective perceptual decision making.

#### ACKNOWLEDGMENTS

We thank T. Sharot and K. Friston for comments on an earlier version of this manuscript and N. Wright for helpful discussions.

Present address of S. M. Fleming: Wellcome Trust Centre for Neuroimaging, 12 Queen Square, London WC1N 3BG, UK.

#### GRANTS

This work was supported by Wellcome Trust Programme Grant to R. J. Dolan, MRC funding within the UCL 4 Year PhD Programme in Neuroscience to S. M. Fleming, a Wellcome Trust PhD Studentship to L. Whiteley, and a Gatsby Charitable Foundation Grant to M. Sahani.

#### DISCLOSURES

No conflicts of interest are declared by the authors.

#### REFERENCES

Andersson JL, Hutton C, Ashburner J, Turner R, Friston K. Modeling geometric deformations in EPI time series. *NeuroImage* 13: 903–919, 2001.
Balcetis E, Dunning D. See what you want to see: motivational influences on visual perception. *J Pers Soc Psychol* 91: 612–625, 2006.

**Behrens TEJ, Woolrich MW, Walton ME, Rushworth MFS.** Learning the value of information in an uncertain world. *Nat Neurosci* 10: 1214–1221, 2009.

**Bohil C, Maddox W.** Category discriminability, base-rate, and payoff effects in perceptual categorization. *Percept Psychophys* 63: 361–376, 2001.

**Boorman ED, Behrens TEJ, Woolrich MW, Rushworth MFS.** How green is the grass on the other side? Frontopolar cortex and the evidence in favor of alternative courses of action. *Neuron* 62: 733–743, 2009.

**Botvinick MM, Braver TS, Barch DM, Carter CS, Cohen JD.** Conflict monitoring and cognitive control. *Psychol Rev* 108: 624–652, 2001.

Brett M, Anton J, Valabregue R, Poline J. Regions of Interest Analysis Using an SPM Toolbox. 8th International Conference on Functional Mapping of the Human Brain, Sendai, Japan, 2002.

Bruner J. On perceptual readiness. Psychol Rev 64: 123–152, 1957.

**Bruner J, Goodman C.** Value and need as organising factors in perception. *J Abnorm Soc Psych* 64: 33–44, 1947.

**Corbetta M, Patel G, Shulman GL.** The reorienting system of the human brain: from environment to theory of mind. *Neuron* 58: 306–324, 2008.

**Corbetta M, Shulman GL.** Control of goal-directed and stimulus-driven attention in the brain. *Nat Rev Neurosci* 3: 201–215, 2002.

Davison M, Tustin R. The relation between the generalized matching law and signal-detection theory. J Exp Anal Behav 29: 331–336, 1978.

Ekstrom LB, Roelfsema PR, Arsenault JT, Bonmassar G, Vanduffel W. Bottom-up dependent gating of frontal signals in early visual cortex. *Science* 321: 414–417, 2008.

**Ferrera VP, Yanike M, Cassanello C.** Frontal eye field neurons signal changes in decision criteria. *Nat Neurosci* 12: 1458–1462, 2009.

- **Fleming SM.** Shaping what we see: pinning down the influence of value on perceptual judgements. *Front Hum Neurosci* 3: 9, 2009.
- Fleming SM, Thomas CL, Dolan RJ. Overcoming status quo bias in the human brain. *Proc Natl Acad Sci USA* 107: 6005–6009, 2010.
- Friston KJ, Glaser DE, Henson RNA, Kiebel S, Philips C, Ashburner J. Classical and Bayesian inference in neuroimaging: applications. *NeuroImage* 16: 484–512, 2002.
- Friston KJ, Daunizeau J, Kiebel SJ. Reinforcement learning or active inference? PloS One 4: , 2009.
- Friston KJ, Worsley KJ, Frackowiak RSJ, Mazziotta JC, Evans AC. Assessing the significance of focal activations using their spatial extent. *Hum Brain Mapp* 1: 214–220, 1994.
- **Gold J, Shadlen M.** Banburismus and the brain: decoding the relationship between sensory stimuli, decisions, and reward. *Neuron* 36: 299–308, 2002.
- **Green D, Swets J.** Signal Detection Theory and Psychophysics. New York: Wiley, 1966.
- **Grinband J, Hirsch J, Ferrera V.** A neural representation of categorization uncertainty in the human brain. *Neuron* 49: 757–763, 2006.
- Grinband J, Wager TD, Lindquist M, Ferrera VP, Hirsch J. Detection of time-varying signals in event-related fMRI designs. *Neuroimage* 43: 509– 520, 2008.
- **Heekeren H, Marrett S, Bandettini PA, Ungerleider LG.** A general mechanism for perceptual decision-making in the human brain. *Nature* 431: 859–862, 2004.
- **Heekeren H, Marrett S, Ungerleider L.** The neural systems that mediate human perceptual decision making. *Nat Rev Neurosci* 9: 467–479, 2008.
- Helmholtz H. Treatise on Physiological Optics. Thoemmes Continuum, London, UK, 1856.
- **Ho TC, Brown S, Serences JT.** Domain general mechanisms of perceptual decision making in human cortex. *J Neurosci* 29: 8675–8687, 2009.
- Jack AI, Roepstorff A. Introspection and cognitive brain mapping: from stimulus-response to script-report. Trends Cog Sci 6: 333–339, 2002.
- Kahneman D, Tversky A. Prospect theory—analysis of decision under risk. Econometrica 47: 263–291, 1979.
- Kersten D, Mamassian P, Yuille A. Object perception as Bayesian inference. Annu Rev Psych 55: 271–304, 2004.
- Annu Rev Psych 55: 271–304, 2004. Landy MS, Goutcher R, Trommershauser J, Mamassian P. Visual estima-
- tion under risk. J Vis 7: 4: 1-15, 2007. Lauwereyns J, Watanabe K, Coe B, Hikosaka O. A neural correlate of
- response bias in monkey caudate nucleus. *Nature* 418: 413–417, 2002. **Leon M, Shadlen M.** Effect of expected reward magnitude on the response of neurons in the dorsolateral prefrontal cortex of the macaque. *Neuron* 24: 415–425, 1999.
- Li S, Mayhew SD, Kourtzi Z. Learning shapes the representation of behavioral choice in the human brain. *Neuron* 62: 441–452, 2009.
- **Liston D, Stone L.** Effects of prior information and reward on oculomotor and perceptual choices. *J Neurosci* 28: 13866–13875, 2008.
- Liu X, Powell DK, Wang H, Gold BT, Corbly CR, Joseph JE. Functional dissociation in frontal and striatal areas for processing of positive and negative reward information. J Neurosci 27: 4587–97, 2007.
- **Lo** C, Wang X. Cortico-basal ganglia circuit mechanism for a decision threshold in reaction time tasks. *Nat Neurosci* 9: 956–963, 2006.
- Lobel E, Kahane P, Leonards U, Grosbras M, Lehericy S, Le Bihan D, Berthoz A. Localization of human frontal eye fields: anatomical and functional findings of functional magnetic resonance imaging and intracerebral electrical stimulation. *J Neurosurg* 95: 804–815, 2001.
- Lundqvist D, Flykt A, Ohman A. The Karolinska Directed Emotional Faces KDEF, 1998.
- **Macmillan N, Creelman C.** Detection Theory: A User's Guide. New York: Erlbaum, 2005.
- Maddox W, Bohil C. A theoretical framework for understanding the effects of simultaneous base-rate and payoff manipulations on decision criterion learning in perceptual categorization. J Exp Psychol Learn Mem Cogn 29: 307–320, 2003.
- Maldjian JA, Laurienti PJ, Kraft RA, Burdette JH. An automated method for neuroanatomic and cytoarchitectonic atlas-based interrogation of fMRI data sets. *NeuroImage* 19: 1233–1239, 2003.
- **Maunsell JH.** Neuronal representations of cognitive state: reward or attention? *Trends Cog Sci* 8: 261–265, 2004.

- McKeeff T, Tong F. The timing of perceptual decisions for ambiguous face stimuli in the human ventral visual cortex. *Cereb Cortex* 17: 669–678, 2007
- **O'Doherty JP.** Reward representations and reward-related learning in the human brain: insights from neuroimaging. *Curr Opin Neurobiol* 14: 769–776, 2004.
- Philiastides MG, Ratcliff R, Sajda P. Neural representation of task difficulty and decision making during perceptual categorization: a timing diagram. J Neurosci 26: 8965–8975, 2006.
- Philiastides MG, Sajda P. EEG-informed fMRI reveals spatiotemporal characteristics of perceptual decision making. J Neurosci 27: 13082–13091, 2007
- Pleger B, Blankenburg F, Ruff CC, Driver J, Dolan RJ. Reward facilitates tactile judgments and modulates hemodynamic responses in human primary somatosensory cortex. *J Neurosci* 28: 8161–8168, 2008.
- Pleger B, Ruff CC, Blankenburg F, Bestmann S, Wiech K, Stephan KE, Capilla A, Friston KJ, Dolan RJ. Neural coding of tactile decisions in the human prefrontal cortex. *J Neurosci* 26: 12596–12601, 2006.
- Ploran E.J., Nelson SM, Velanova K, Donaldson DI, Petersen SE, Wheeler ME. Evidence accumulation and the moment of recognition: dissociating perceptual recognition processes using fMRI. *J Neurosci* 27: 11912–11924, 2007
- Preuschoff K, Quartz SR, Bossaerts P. Human insula activation reflects risk prediction errors as well as risk. J Neurosci 28: 2745–2752, 2008.
- Puri AM, Wojciulik E, Ranganath C. Category expectation modulates baseline and stimlus-evoked activity in human interotemporal cortex. *Brain Res* 1301: 89–99, 2009.
- Rorie AE, Gao J, McClelland JL, Newsome WT. Integration of sensory and reward information during perceptual decision-making in lateral intraparietal cortex of the macaque monkey. PLoS ONE 5: 2010.
- **Romo R, Hernandez A, Zainos A.** Neuronal correlates of a perceptual decision in ventral premotor cortex. *Neuron* 41: 165–173, 2004.
- Serences J. Value-based modulations in human visual cortex. *Neuron* 60: 1169–1181, 2008
- Serences JT, Schwarzbach J, Courtney SM, Golay X, Yantis S. Control of object-based attention in human cortex. Cereb Cortex 14: 1346–1357, 2004.
- Simen P, Cohen JD, Holmes P. Rapid decision threshold modulation by reward rate in a neural network. *Neural Networks* 19: 1013–1026, 2006.
- Summerfield C, Egner T, Greene M, Koechlin E, Mangels J, Hirch J. Predictive codes for forthcoming perception in the frontal cortex. *Science* 314: 1311–1314, 2006a.
- Summerfield C, Egner T, Mangels J, Hirsch J. Mistaking a house for a face: neural correlates of misperception in healthy humans. *Cereb Cortex* 16: 500–508, 2006b
- Summerfield C, Koechlin E. A neural representation of prior information during perceptual inference. *Neuron* 59: 336–347, 2008.
- **Thielscher A, Pessoa L.** Neural correlates of perceptual choice and decision making during fear-disgust discrimination. *J Neurosci* 27: 2908–2917, 2007
- **Tong F, Nakayama K, Vaughan JT, Kanwisher N.** Binocular rivalry and visual awareness in human extrastriate cortex. *Neuron* 21: 753–759, 1998.
- **Tosoni A, Galati G, Romani GL, Corbetta M.** Sensory-motor mechanisms in human parietal cortex underlie arbitrary visual decisions. *Nat Neurosci* 11: 1446–1453, 2008.
- **Watanabe M, Sakagami M.** Integration of cognitive and motivational context information in the primate prefrontal cortex. Cereb Cortex 17 *Suppl* 1: i101–i109, 2007.
- Weiskopf N, Hutton C, Josephs O, Deichmann R. Optimal EPI parameters for reduction of susceptibility-induced BOLD sensitivity losses: a whole-brain analysis at 3 T and 1.5 T. *NeuroImage* 33: 493–504, 2006.
- Whiteley L, Sahani M. Implicit knowledge of visual uncertainty guides decisions with asymmetric outcomes. J Vis 8: 2–15, 2008.
- Wichmann F, Hill N. The psychometric function: I. Fitting, sampling, and goodness of fit. *Percept Psychophys* 63: 1293–1313, 2001.
- Yacubian J, Glascher J, Schroeder K, Sommer T, Braus DF, Buchel C. Dissociable systems for gain- and loss-related value predictions and errors of prediction in the human brain. *J Neurosci* 26: 9530–7, 2006.
- Yantis S, Schwarzbach J, Serences JT, Carlson RL, Steinmetz MA, Pekar JJ, Courtney SM. Transient neural activity in human parietal cortex during spatial attention shifts. *Nat Neurosci* 5: 995–1002, 2002.