The Orbitofrontal Cortex Represents Advantageous Choice in the Iowa Gambling Task

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The orbitofrontal cortex represents advantageous choice in the Iowa gambling task

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Abstract

A good-based model proposes that the orbitofrontal cortex (OFC) represents binary choice outcome, i.e., the chosen good. Previous studies have found that the OFC represents the binary choice outcome in decision-making tasks involving commodity type, cost, risk, and delay. Real-life decisions are often complex and involve uncertainty, rewards, and penalties; however, whether the OFC represents binary choice outcomes in such a decision-making situation, e.g., Iowa gambling task (IGT), remains unclear. Here, we propose that the OFC represents binary choice outcome, i.e., advantageous choice versus disadvantageous choice, in the IGT. We propose two hypotheses: first, the activity pattern in the human OFC represents an advantageous choice; and second, choice induces an OFC-related functional network. Using functional magnetic resonance imaging and advanced machine learning tools, we found that the OFC represented an advantageous choice in the IGT. The OFC representation of advantageous choice was related to decision-making performance. Choice modulated the functional connectivity between the OFC and the superior medial gyrus. In conclusion, the OFC represents an advantageous choice during the IGT. In the framework of a good-based model, the results extend the role of the OFC to complex decision-making when making a binary choice.

Keywords: orbitofrontal cortex, advantageous choice, multivoxel pattern analysis, psychophysiological interaction, fMRI
Introduction

The identified neurobiological mechanism underlying economic decision-making includes a valuation stage and a choice stage\(^1\). Decision makers evaluate the subjective values and characteristics of available options in the valuation stage. However, encoding subjective value and characteristics is not sufficient for making decisions, and one of the available options still needs to be chosen by decision makers at the choice stage\(^1\). At this stage, a good-based model, a central neurobiological model of economic decision-making, proposes that the orbitofrontal cortex (OFC) represents the binary choice outcome, i.e., the chosen good\(^4\). A good is defined by a group of determinants characterizing the conditions in which the commodity is offered, which can include commodity type, time delay, cost, risk, and ambiguity\(^4\).

Consistent with a good-based model, previous studies have found that the OFC represents the binary choice outcome in juice-choice tasks\(^5\) and decision-making tasks involving costs\(^6\), risks\(^7\), and delays\(^8\). For example, different OFC neurons respond when a monkey chooses between different juice types\(^5\). Some OFC neuronal responses in monkeys encode choosing a high-cost option versus choosing a low-cost option\(^6\). Some other OFC neurons in monkeys encode choosing a risky option versus choosing a nonrisky option\(^7\). The OFC activity pattern in the human brain can classify choosing smaller-but-immediate options versus choosing larger-but-delayed options\(^8\). However, real-life decisions are often complex and involve uncertainty, rewards, and penalties.

The inability to make choices in a complex decision-making situation, e.g., Iowa gambling task (IGT), is a symptom of several brain disorders, including borderline personality disorder\(^9\), attention-
deficit/hyperactivity disorder, anorexia nervosa, addiction, obsessive-compulsive disorder, and schizophrenia. In the IGT, reward value is a key decision-making parameter. Whether the OFC represents a binary choice outcome in the IGT, advantageous choice (i.e., choosing an option with a high reward value) versus disadvantageous choice (i.e., choosing an option with a low reward value), remains unclear.

A line of studies has implicated the OFC at the valuation stage, i.e., evaluating available options such as the value, risk, ambiguity, and environmental statistics. For example, Hare et al. and Kable et al. reported that OFC activity was correlated with high values versus low values. Both Bartra et al. and Clithero et al. have shown that the OFC is a key brain area related to high subjective values versus low subjective values of different types of rewards using meta-analysis. Some studies have also investigated the neural basis of high ambiguity versus low ambiguity in decision-making. For example, Levy et al. showed that OFC activity is correlated with ambiguity level. Hsu et al. also revealed that the OFC showed greater activation in response to the level of ambiguity. Huettel et al. found increased activation in the inferior frontal sulcus, insula, and parietal cortex when ambiguity was present. Bach et al. found that ambiguity is related to parietal cortex activity. Therefore, these studies have implicated the OFC in evaluating risk, ambiguity, and value.

Another line of neurobiological studies investigated advantageous versus disadvantageous choice in the IGT; however, they found that blood oxygenation level-dependent (BOLD) activation using individual voxel-based methods in the OFC was not significantly associated with advantageous choice versus disadvantageous choice. For example, Brevers et al. did not find any advantageous choice-related
activation in the OFC in healthy controls or poker gamblers in the IGT\textsuperscript{37}. One potential explanation for the finding that the OFC was not implicated in advantageous choice in the IGT in these studies is that ensembles of many voxels, rather than single voxel activation, are responsible for generating economic choices. For example, multiple voxels in the OFC were shown to contain information on the discrimination between choosing a larger-but-delayed option and choosing a smaller-but-immediate option in decision-making involving delays\textsuperscript{8}. Therefore, the individual-voxel-based methods used by previous studies might not be suitable. As multivoxel pattern analysis (MVPA) can detect fine-grained spatial patterns across multiple voxels that might discriminate between cognitive processes\textsuperscript{41}, it may be a potential method to detect advantageous choice in the OFC in the IGT. Therefore, in the present study, we hypothesized that the OFC represented advantageous choice in the IGT.
Methods

Participants. Fifty-five healthy participants were recruited in the study, and one participant was excluded after presenting with significant head motion (>3.0 mm) during functional magnetic resonance imaging (fMRI) scanning. The remaining fifty-four participants included 45 males and nine females [age: mean, 22.7 years; standard deviation (SD), 2.1 years; range, 19 to 27 years; education: mean, 16.3 years; SD, 1.8 years; range, 13 to 19 years]. All participants were free of psychiatric or neurological history and had normal or corrected-to-normal vision. The study was approved by the Human Research Ethics Committee of the University of Science and Technology of China. The methods and procedures used in this study were carried out in accordance with the approved guidelines. Written informed consent was obtained from all participants before the study, consistent with the Declaration of Helsinki guidelines.

Task paradigm. In the present study, we used the Iowa gambling task (IGT) (Fig. 1), a popular decision-making task for indexing real-life complex decision-making. In each trial, the participants selected a card from among four decks of cards. The four decks were labelled A, B, C, and D as presented from left to right. On each card, there were different numbers of gain and possible loss points, and the participant received the net (gain - loss) points for choosing that card. Participants did not know the expected reward and variability in the outcomes for all decks before engaging in the task. In the task, the participants were asked to maximize the points they gained. Specifically, for each selection from deck A or B (“low reward value decks”), participants would gain 100 points, but the losses were organized so that over 10 selections from the decks, the participants would have an overall loss of 250 points. Specifically, deck A provided -150, -200, -250, -300 and -350 (loss) points every ten selections, whereas deck B provided -1250 (loss) points in one out of ten selections. For each selection from deck C or D (“high reward value decks”), the participants would win 50 points, and the losses were organized so that if participants made over 10 selections from these decks, they would obtain an overall profit of 250 points. The two decks differed in the frequency and magnitude of the punishment. Similar to the previous two decks, deck C provided -25, -40, -50, -60 and -75 (loss) points every ten selections, whereas deck D provided -250 (loss) points once every ten selections. Decks A and B had negative reward expectations and were operationally defined as having a low reward value. In contrast, decks C and D had positive reward expectations and were defined as having a high reward value. Therefore,
choosing decks C and D was an advantageous choice, and choosing decks A and B was a
disadvantageous choice. Similar to previous studies\textsuperscript{42,43}, the IGT was extended to 180 trials from the
original 100 trials to facilitate rule learning\textsuperscript{14}. The IGT consisted of three scan runs, with three blocks
for each scan run and 20 trials for each block. The participants who had positive net winnings at the end
of the task would obtain extra money (10¥/1000 points). The final net winnings were defined as the
total score.

Behavioural analysis—reinforcement learning model. This procedure followed that of a previous
study\textsuperscript{43}. The reinforcement learning model\textsuperscript{44} was adapted to analyse the behavioural data. Reward
prediction errors (RPEs) were included in the model, according to the suggestion by Sutton and Barto\textsuperscript{45}.
An RPE ($\delta_t$) was defined as the difference between the actual reward $r_t$ and the predicted reward $\hat{v}_t$
at trial t. The formula for this definition was as follows:

$$\delta_t = r_t - \hat{v}_t.$$  \hspace{1cm} (1)

The RPE was used to update reward prediction in the model using the following formula:

$$\hat{v}_{t+1} = \hat{v}_t + \alpha \cdot \delta_t.$$  \hspace{1cm} (2)

where $\alpha$ is the learning rate for the RPE in the update formula\textsuperscript{44}. Then, maximum likelihood estimation
(MLE) was adopted to estimate the learning rate based on the samples. Here, $\pi_{it}$ was defined as the
probability of choice $i$ at trial $t$. We transformed the data with an exponential function when we
calculated the value of $\pi_{it}$ using the following formula:

$$\pi_{it} = \frac{e^{\hat{v}_{it}}}{\sum_{j=1}^{n} e^{\hat{v}_{jt}}}.$$  \hspace{1cm} (3)

The learning rate was estimated separately by maximizing the likelihood function for each participant:

$$\text{Maximum log - likelihood} = \max \sum_{t=1}^{M} \log \pi_{i_{t},t}$$  \hspace{1cm} (4)

where $i_{t}$ represents the deck selected at trial $t$, $i_{t} \in \{1, 2, 3, 4\}$, and $\pi_{i_{t},t}$ represents the probability
of selecting deck $i_{t}$ at trial $t$.

To test whether the participants’ decision-making performance was better than random chance, we
performed a random selection simulation 1000 times. We compared the learning rate from the
participants’ choices with that from the simulation using the t test. We tested group differences using t
test if data conformed normality and using Mann-Whitney test if data do not conform normality in the
present study. Cohen's $d$ values were calculated via G*Power 3.1 software\textsuperscript{46}. We calculated the total net good decks, which was the number of advantageous choices minus the number of disadvantageous choices in 180 trials.

**fMRI data acquisition and preprocessing.** Gradient echo-planar imaging data were acquired using a 3.0 T, 8-channel head coil Trio scanner (Siemens Medical Solution, Erlangen, Germany) with a circularly polarized head coil in Hefei. We restrained head motion with foam padding. A T2*-weighted echo-planar imaging sequence (FOV = 240 mm, TE = 30 ms, TR = 2000 ms, flip angle = 85°, matrix = 64 × 64) with 33 axial slices (no gaps, 3.7 mm thick) covering the whole brain was used to acquire the functional MR images. There were three runs of IGT, each of which contained 210 epochs. Furthermore, high-resolution T1-weighted spin-echo imaging data (1 mm isotropic voxel) were also acquired for anatomical overlay.

We preprocessed the imaging following the workflows proposed in a previous paper\textsuperscript{47}. All functional MR images were preprocessed using Analysis of Functional Neuroimages (Version AFNI_18.2.03) software\textsuperscript{48}. All fMRI data were corrected for temporal shifts between slices and motion and grand-mean scaled. Low-frequency signal drifts were filtered using a cutoff of 128 s. Volumes meeting the following criteria were removed: translation>0.3mm or rotation>0.3° between consecutive volumes\textsuperscript{49}. For each run, we dropped the first two volumes to enhance stability. Linear regression was also performed to remove linear trends. All functional volumes were non-linearly transformed to MNI space (resampled voxel size: $4 \times 4 \times 4$ mm$^3$) according to the spatial transformation between the anatomical data and the MNI space. Volumes were spatially smoothed with a Gaussian kernel (full-width at half-maximum = 8 mm) and were used for general linear model and psycho-physiological interaction (PPI) analysis. Unsmoothed data were used for MVPA.

**General linear model for value signals.** To illustrate the neural activations of the values, including RPE, gain, loss, and reward predictions for the four decks, a general linear model was used to examine the BOLD signals in which brain regions were correlated with these values. The general linear model was run for each value and included 1) an interest regressor, \textit{i.e.}, one-value regressor, defined as RPE, gain, loss, or reward prediction for the four decks during the epochs when feedback was presented and 0 for
other epochs, and 2) six noninterest regressors for head motion. Then, the parameter estimates were
extracted for each value and for each participant. We performed a group-level one-sample t test for
parameter estimates using family-wise error correction.

Whole brain searchlight-based multivoxel pattern analysis. We first used whole brain searchlight-based
MVPA to classify advantageous choice versus disadvantageous choice. We adapted the within-subject
MVPA methods from a previous study. We used the least squares-separate (LSS) method to extract
choice-related activations according to a previous study. LSS is the most effective method to estimate
choice activation and has been widely used in the field. According to the LSS method, a general
linear model was used to extract activation for each choice. There were 180 choices, including
advantageous choices and disadvantageous choices, for each participant. A general linear model was
run for each choice. For the i\textsuperscript{th} choice, the general linear model included two choice regressors. The
first was the choice regressor of interest. During a trial with choice \( C_i \), this regressor was defined as 1
during the epoch when a button press was made in the selection phase and 0 for the other epochs; during
trials with choices \( C_{1...i-1, i+1...180} \), this regressor was defined as 0 for all epochs. The other was the
choice regressor of nuisance. During a trial with choice \( C_i \), this regressor was defined as 0 for all epochs;
during trials with choices \( C_{1...i-1, i+1...180} \), this regressor was defined as 1 during the epoch when a
button press was made in the selection phase and 0 for the other epochs. The value of \( \beta \) for the choice
regressor of interest in the general linear model was the activation for choice \( C_i \). The general linear
model was repeated 180 times to extract activations for 180 choices for each participant. The general
linear model was performed using MATLAB’s \texttt{regstats} function (MATLAB v2019a, Mathworks Inc,
Natick, MA, PC).

We implemented two steps to control the effects of values, as choices can be expected to be related to
value signals, including RPE, gain, loss, and reward predictions for the four decks. For step 1, we used
the Gram-Schmidt orthogonalization algorithm to orthogonalize choices and values before
implementing the general linear model. Specifically, we orthogonalized choice and RPE, gain, loss,
and reward predictions for the four decks. For step 2, the orthogonalized choice regressor of interest,
the orthogonalized choice regressor of nuisance, the regressors for RPE, gain, loss, and reward
predictions for the four decks [those defined as RPE, gain, loss, or reward predictions for the four decks
during the epochs when feedback was presented and 0 for the other epochs], and six regressors of no
interest for head motion were included in each general linear model. We extracted \( \beta \), the activation of
the orthogonalized choice regressor of interest, for each voxel in the whole brain in each general linear
model. The extracted activations were grouped into two categories according to the choice type, i.e.,
advantageous choice versus disadvantageous choice, for each voxel and for each participant.

We performed whole brain searchlight-based MVPA that did not depend on a priori assumptions but
searched for predictive information across the whole brain. For each voxel \( v_i \), considering the local
patterns that contained the spatial correlation that might decode advantageous choice versus
disadvantageous choice, we constructed a spherical collection of voxels \( S_{I...N} \), with 33 voxels\(^{56}\) centred
on voxel \( v_i \). For each voxel \( S_{I...N} \) in the collection, we extracted \( \beta \); namely, \( V_{I...N} \). \( V_{I...N} \) were
normalized to the range from 0 to 1 for advantageous choice and disadvantageous choice separately to
give all voxels equal importance during classifier training\(^{57,58}\). The values \( V_{I...N} \) were then used to train
and test the classifier model, which was a support vector machine with a linear kernel. The decoding
accuracy of the central voxel \( v_i \) was acquired by five-fold cross-validation. The implementation of the
support vector machine and cross-validation were based on sklearn.svm.SVC in Python’s scikit-learn
toolbox (version 0.21.2)\(^{59}\). During training and testing of the classification model, random
undersampling was used to handle the imbalance in samples between advantageous choice and
disadvantageous choice. For example, if the number of advantageous choices was larger than that of
disadvantageous choices, advantageous choices were removed randomly to make the numbers the same
as the disadvantageous choices by the numpy.random.shuffle function in Python (version 3.6.8). Equal
numbers of both choices were labelled the original data sample, which was then randomly partitioned
into five equal sized subsamples for five-fold cross-validation. The same procedure was performed for
each voxel over the whole brain for each participant. The whole brain decoding accuracy was
normalized by subtracting the mean of the whole brain accuracy for each participant.

We performed a group-level one-sample t test for whole brain searchlight-based MVPA for decoding
accuracy using family-wise error correction.

We also tested whether choice-related activations were correlated with value signals, i.e., RPE, gain,
loss, and reward predictions for the four decks using both whole brain analysis and region of interest (ROI) analysis. Specifically, the extracted activations in the general linear model were grouped into two categories according to the median split of the values of the trials, i.e., high and low subgroups for RPE, gain, loss, and reward predictions for the four decks for each participant. We tested whether these subgroups showed differences for RPE, reward predictions for the four decks, gain, and loss separately using the t test in the whole brain with family-wise error correction. We further included the left and right OFC ROIs from the Anatomical Automatic Labeling atlas (AAL2)\(^60\). The left OFC ROI included OFCmed\_L, OFCant\_L, OFCpost\_L, and OFClat\_L and the right OFC ROI included OFCmed\_R, OFCant\_R, OFCpost\_R, and OFClat\_R. The extracted activations for values above were averaged in the left and right OFC ROIs separately, then, were fed into group comparisons using the t test with uncorrected \(p<0.05\).

ROI-based MVPA. We further tested whether the OFC represented choice using ROI-based MVPA. First, we included OFC ROIs from the AAL2 that showed overlapping areas with the peak voxel for significant clusters in the whole brain searchlight-based MVPA. Second, we extracted the activations associated with each choice for each OFC ROI. Activations were also normalized to the range from 0 to 1 for advantageous choice and disadvantageous choice separately\(^57,58\). The decoding accuracy for each OFC ROI was acquired by five-fold cross-validation. We tested whether the decoding accuracy was greater than chance level (0.5) for each OFC ROI using a one-sample t test. We tested whether the decoding accuracy was correlated with the learning rate, total score, and total net good decks using Pearson correlations.

To test whether the signal-to-noise ratio (SNR) affected the decoding results, Pearson correlations between the SNR and decoding accuracy for each ROI were determined.

**PPI analysis.** To investigate whether the functional connectivity of the OFC identified in ROI-based MVPA differed between advantageous and disadvantageous choices, we ran PPI analysis. First, we created a “seed” time series by extracting mean time courses for each OFC identified in ROI-based MVPA. Second, we computed the interaction terms between the “seed” and either the (1) advantageous...
choice regressor, defined as 1 during the epoch when a button press was made in the selection phase and 0 for other epochs during trials with advantageous choice or the (2) disadvantageous choice regressor, defined as 1 during the epoch when a button press was made in the selection phase and 0 for other epochs during trials with disadvantageous choice and as 0 for all epochs during trials with advantageous choice. Third, we estimated a PPI general linear model including the following regressors: (1) the advantageous choice regressor, (2) the disadvantageous choice regressor, (3) the OFC seed time course, (4) the interaction term between the “seed” and advantageous choice regressor, defined as advantageous choice PPI, (5) the interaction term between the “seed” and disadvantageous choice regressor, defined as disadvantageous choice PPI, (6) seven value regressors including RPE, gain, loss, and reward predictions for the four decks, defined as RPE, gain, loss, or reward predictions for the four decks, respectively, during the epochs when feedback was presented and 0 for the other epochs, and (7) six noninterest regressors for head motion. The PPI general linear model was performed using AFNI’s 3dDeconvolve.

We computed the first-level contrast for the disadvantageous choice PPI $\beta$ minus the advantageous choice PPI $\beta$. We performed a one-sample t test to identify significant differences in the contrast to identify PPI effects using family-wise error correction.

We also tested whether there were overlapping regions in the brain among the whole brain searchlight-based MVPA and PPI.

As a control analysis, we tested whether value signals modulated OFC functional connectivity. To achieve this, we performed PPI analysis for RPE, gain, loss, and reward predictions for the four decks separately. The PPI general linear model included the following regressors: (1) a value regressor, defined as RPE, gain, loss, or reward predictions for the four decks during the epochs when feedback was presented and 0 for other epochs, (2) the OFC seed time course, (3) the interaction term between the “seed” and value regressor, defined as the value PPI, and (4) six noninterest regressors for head motion. We computed the first-level contrast for PPI $\beta$ values and performed a one-sample t test to identify PPI effects using family-wise error correction.
Results

Summary of behavioural performance in the IGT

We found that the participants’ learning rate was significantly higher than the learning rate from the computer's random 1000 selections [Mann-Whitney test, Mann-Whitney U=53, $p < .001$, 95% confidence interval: [0.085, 0.116]]. The participants’ learning rate, response time, number of advantageous choices, number of disadvantageous choices, total score, and total net good decks are summarized in Table 1.

BOLD activity in the OFC is correlated with value signals

We found significant activations in the OFC, striatum, and posterior cingulate cortex for value signals, including RPE, gain, loss (Fig. 2 and Table 2), and reward predictions for the four decks (Supplementary Figure 1 and Supplementary Table 1). Therefore, the results are consistent with previous studies showing that the OFC is implicated in value evaluation.42,61,62

The OFC represents advantageous choice

As the OFC has been implicated in the representation of value signals, we next examined whether the OFC represented advantageous choice while controlling for value effects. Using whole brain searchlight-based MVPA, we found that the activity pattern in the OFC indeed represented advantageous choice (Fig. 3a and Table 3). Whole brain searchlight-based MVPA also revealed that activity in the frontal regions and the parietal regions represented advantageous choice (Fig. 3a and Table 3); thus, we replicated similar findings regarding the representation of choice in the frontoparietal network from previous studies.12

Are choice related activations in the OFC related to value signals? We found that there were no significant activations in the OFC between the high and low subgroups for RPE, gain, loss (Fig. 3b, 3c, 3d, and Supplementary Table 3), or reward predictions for the four decks (Supplementary Figure 2 and Supplementary Table 4). We further found that there were no significant differences in the left or right OFC ROIs between the high and low subgroups for RPE, gain, loss (all $p_s > 0.05$, uncorrected). The results suggest that choice-related activations in the OFC for MVPA were not confounded by value signals.
The peak voxels of significant clusters in the whole brain searchlight-based MVPA showed an overlapping area with OFCmed_R in AAL2; therefore, we further examined the choice representation in OFCmed_R using ROI-based MVPA. We found that OFCmed_R represented an advantageous choice [$t_{53} = 7.770, p < 0.001$, Cohen’s $d = 1.057$, 95% confidence interval: [0.075, 0.126]] (Fig. 4a). We found significant correlations between the decoding accuracy and learning rate [$r = 0.559, p < 0.001, N = 54$], total score [$r = 0.357, p = 0.008, N = 54$], and total net good decks [$r = 0.468, p < 0.001, N = 54$] (Fig. 4b, 4c, and 4d). The decoding accuracy showed no significant correlations with the SNR [$r = −0.119, p = 0.390, N = 54$] or censor rate [$r = 0.071, p = 0.610, N = 54$], suggesting that the decoding accuracy was not explained by either of these parameters.

**Choice modulates OFC functional connectivity with the superior medial gyrus**

PPI analysis revealed greater OFC connectivity with the superior medial gyrus when choosing disadvantageous options versus choosing advantageous options (Fig. 5a). Furthermore, the superior medial gyrus showed an overlapping area with brain regions representing advantageous choice revealed by whole brain searchlight-based MVPA (Fig. 5b). As a control analysis, we tested whether value signals modulated OFC functional connectivity. We found that there was no significant OFC functional connectivity in the whole brain for RPE, gain, loss, or reward predictions for the four decks (Fig. 5c), suggesting that choice-modulated OFC functional connectivity was not confounded by the value signals.
Discussion

Consistent with the proposal of a good-based model, the present study demonstrates that the OFC represents advantageous choice, which provides strong evidence to support the role of the OFC in binary choice in the IGT. Furthermore, IGT behavioural performances were correlated with the advantageous choice representation in the OFC. Third, the functional connectivity between the OFC and superior medial gyrus supports choice.

The OFC represents an advantageous choice in the IGT

In the present study, we demonstrated that the OFC represents binary choice in the IGT based on the distributed activity pattern. These results are supported by neurobiological studies with human\textsuperscript{14,63} as well as animal\textsuperscript{64,65} prefrontal lesions (including in the OFC), consistently indicating that the OFC plays a necessary role in decision-making.

Furthermore, beyond the OFC, the frontoparietal network was also implicated in choice in the present study, and the finding is consistent with previous studies\textsuperscript{1,2}. Both the OFC and the frontoparietal network represented choice in the present study; therefore, we expected to find functional connectivity between the regions for choice. Indeed, we identified a functional connectivity between the OFC and the superior medial gyrus for choice, but not for values, suggesting that the OFC is functionally coupled with the prefrontal cortex when humans make choices. As the frontoparietal network has been widely implicated various decision-making situations\textsuperscript{1,2}, the connectivity between the OFC and the superior medial gyrus would be helpful for choice under various decision-making contexts. We also found that OFC activity is related to value signals, e.g., RPE, gain, and loss. Therefore, the finding that both choice and value
were represented in the OFC may make it easier for individuals to make optimal choices with the help of frontoparietal network modulation during difficult decision-making situations that lack sufficient information.

A proposed role for the OFC: representation of choice-related complex information along a continuous spectrum

In the present study, the OFC was shown to represent advantageous choice. This finding is supported by a recently proposed cognitive map representing a state space. In the context of the cognitive map, the OFC is activated when the decision maker becomes cognizant of unobservable information and makes a correct choice; however, the OFC would not activate when the decision maker is not cognizant of unobservable information and makes an incorrect choice.

Interestingly, in the present study, even though the participants did not know the specific reward value for each deck, the OFC nevertheless represented advantageous choice. Integration of these findings shows that exact knowledge of complex information is not necessary for OFC activation. This may suggest that the OFC, in part, could play a role in unconscious influences, e.g., emotions, in complex decision-making.

We found that decoding accuracy in the OFC correlated with decision-making performance. We therefore propose a role for the human OFC based on the cognitive map idea: the OFC may represent choice-related complex information along a continuum, e.g., from a high decoding accuracy of advantageous choice if the decision maker exactly knows the complex information to a low decoding
accuracy if they do not. Our proposal further predicts that the OFC represents choice in choosing between other decision-making parameters, such as self-control and cost. This is important because humans often face choices that have unknown costs for effort control. It is beneficial to exert an appropriate level of effort for an appropriate choice. The OFC seems to be a candidate for the brain region used when making choices based on the aforementioned parameters in a complex context; this hypothesis should be investigated in future work.

Several shortcomings of the present study should be acknowledged. First, few female participants were recruited in the present study. We conducted ROI-based MVPA for males and females separately to test whether females and males showed a difference in decoding accuracy. We found consistent results (see Supplementary Table 5) between the sexes, suggesting that the percentage of females might not influence our results. Future work should include more female participants to substantiate our conclusion. Second, the IGT design convolves decision-making with uncertainty with learning. Good learning would presumably result in choosing from the high value deck and not from the other decks and would also presumably result in choosing from the high value deck more in the late IGT runs and less in the early IGT runs. Therefore, a reasonable assumption would be that changes in activation patterns between choosing high value versus choosing low value may be due to differences in choice probability. However, the choice probability is related to reward prediction, which we controlled for when we performed MVPA and PPI analysis. We found that choice-related activations in the OFC could represent choice in MVPA and that choice-related activations in the OFC were not related to reward predictions. We also found that choice, but not reward predictions, modulated OFC functional connectivity with the superior medial gyrus in PPI analysis. Therefore, our results suggest that neither
the choice probability nor the learning effect in the IGT confounds the decoding or PPI results in the present study. Future work will require a complex decision-making task without learning to substantiate our conclusion. Third, attention modulates the value signal in the OFC, thus, a reasonable assumption would be that choice-related signals in the OFC may be confounded with attention. However, the OFC signal has been related to value of attended option and we found that choice-related signals in the OFC were not related to all value signals, including RPE, gain, loss, or reward predictions for the four decks. Therefore, our results suggest that attention may not confound the decoding or PPI results in the present study. Future work will require a decision-making task with covert shift of attention to substantiate our conclusion.

Conclusions

In conclusion, our results demonstrate that the OFC represents advantageous choice in the IGT. Our data provide evidence to support the integration of knowledge in the OFC to make choices in a complex context, which may be helpful for survival. Decreased decoding accuracy in the OFC may be related to poor decision-making ability, and these findings may provide potential insight into understanding impulsive behaviours.
Credit authorship contribution statement

Rujing Zha: Conceptualization, Formal analysis, Methodology, Software, Data curation, Writing-Original draft preparation, Project administration; Peng Li: Data curation, Formal analysis, Methodology, Writing-Reviewing and Editing; Ying Li: Formal analysis, Validation, Visualization; Nan Li: Formal analysis; Meijun Gao: Formal analysis, Writing- Reviewing and Editing; Yiyang Fan: Formal analysis; Ruhuiya Aili: Formal analysis; Ying Liu: Methodology; Xiaochu Zhang: Supervision, Conceptualization, Methodology, Writing- Reviewing and Editing; Jun Li: Supervision, Conceptualization, Methodology, Writing-Reviewing and Editing.

Data availability

All data are available from the authors.

Code availability

Matlab codes are available from the authors. Code for fMRI orthogonalization are available at https://www.mathworks.com/matlabcentral/fileexchange/55881-gram-schmidt-orthogonalization. Code for MVPA analysis are available at https://scikit-learn.org/dev/modules/svm.html#svm-classification. Code for PPI analysis are available at https://afni.nimh.nih.gov/CD-CorrAna.

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Figure Legends

Figure 1. Experimental paradigm of the Iowa gambling task. Experimental paradigm of the Iowa gambling task. There were two phases for each trial. Four decks were presented in the first phase. Participants selected a card within 4 s in this phase (selection phase, 4 s); then, the outcome, including gain and loss, was presented in the second phase (feedback phase, 1 s).

Figure 2. BOLD activity in the OFC was correlated with value signals. BOLD signals in the OFC, striatum, and posterior cingulate cortex were correlated with value signals, including a) RPE, b) gain, and c) loss. Family-wise error at a cluster-level threshold of p < 0.05 (voxel-level threshold of p < 0.001, voxel size > 13 for RPE, 33 for gain, and 19 for loss). N = 54.

Figure 3. The OFC represents advantageous choice, and choice-related activations in the OFC are not correlated with value signals. a) Whole brain searchlight-based MVPA revealed that the OFC represents choice. Choice-related activations in the OFC are not significantly correlated with value signals, including b) RPE, c) gain, or d) loss. R, Right, L, Left. Family-wise error at a cluster-level threshold of p < 0.05 (voxel-level threshold of p < 0.001, voxel size > 4 for advantageous choice, 1 for RPE, 3 for gain, and 2 for loss). N = 54.

Figure 4. The OFC represents an advantageous choice, and the OFC decoding accuracy is correlated with behavioural performances. a) The OFCmed_R region in AAL2 represents an advantageous choice. The OFCmed_R decoding accuracy was correlated with the b) learning rate, c) total score, and d) total net good decks in the IGT. The dashed line in the panels shows the chance level (0.5), and the dashed area in the panels shows the 95% confidence interval. The error bar shows SE. N = 54.

Figure 5. The OFC is functionally connected with the superior medial gyrus for choice, but not for
a) Compared with advantageous choice, disadvantageous choice increased the OFC functional connectivity with the superior medial gyrus. Voxel size: 40; peak voxel coordinates: -10, -66, +4. R, Right, L, Left. Family-wise error at a cluster-level threshold of p < 0.05 (voxel-level threshold of p < 0.001, voxel size > 13). b) The overlapping area between the superior medial gyrus and the brain regions representing advantageous choice contained 16 voxels. c) There was no significant OFC functional connectivity across the whole brain for RPE, gain, loss, or reward predictions for the four decks. Family-wise error at a cluster-level threshold of p < 0.05 (voxel-level threshold of p < 0.001, voxel size > 1 for RPE, 3 for gain, 2 for loss, 16 for reward prediction for deck A, 18 for reward prediction for deck B, 16 for reward prediction for deck C, 15 for reward prediction for deck D). N = 54.
Table Legends

Table 1. Summary of behavioural performance in the Iowa gambling task.

$N = 54$.

Table 2. BOLD activity in the OFC is correlated with the value signals.

aThe coordinates of the peak voxel are shown in MNI space (+ left, - right; + posterior, - anterior; + superior, - inferior). Family-wise error at a cluster-level threshold of $p < 0.05$ (voxel-level threshold of $p < 0.001$, voxel size > 13 for RPE, 33 for gain, and 19 for loss). $N = 54$.

Table 3. Brain regions including the OFC that represent advantageous choice.

aThe coordinates of the peak voxel are shown in MNI space (+ left, - right; + posterior, - anterior; + superior, - inferior). Family-wise error at a cluster-level threshold of $p < 0.05$ (voxel-level threshold of $p < 0.001$, voxel size > 4). This table only displays the brain regions that formed a cluster of more than 20 voxels; all significant clusters are shown in Supplementary Table 2. $N = 54$. 
Figure 1. Experimental paradigm of the Iowa gambling task.
Figure 2. BOLD activity in the OFC was correlated with value signals.
Figure 3. The OFC represents advantageous choice, and choice-related activations in the OFC are not correlated with value signals.
Figure 4. The OFC represents an advantageous choice, and the OFC decoding accuracy is correlated with behavioural performances.
Figure 5. The OFC is functionally connected with the superior medial gyrus for choice, but not for values.
Table 1. Summary of behavioural performance in the Iowa gambling task.

|                           | Mean  | SD     | Min  | Max   |
|---------------------------|-------|--------|------|-------|
| Learning rate             | 0.152 | 0.119  | 0.012| 0.605 |
| Response time             | 0.655 | 0.240  | 0.273| 1.424 |
| The number of advantageous choices | 136.426 | 20.752 | 83.000 | 168.000 |
| The number of disadvantageous choices | 43.574  | 20.752 | 12.000 | 97.000 |
| Total score               | 5050.741 | 977.772 | 3075 | 6885  |
| Total net good decks      | 92.852 | 41.503 | -14.000 | 156.000 |
Table 2. BOLD activity in the OFC is correlated with the value signals.

| Brain regions                  | Voxels | x^a | y   | z   |
|-------------------------------|--------|-----|-----|-----|
| **RPE**                      |        |     |     |     |
| Right inferior frontal gyrus  | 2450   | -26 | -22 | +24 |
| Left superior frontal gyrus   | 43     | +22 | -42 | +48 |
| Left superior frontal gyrus   | 37     | +18 | -62 | +12 |
| Right cerebellum             | 20     | -42 | +78 | -36 |
| Left cerebellum              | 19     | +10 | +46 | -16 |
| **Gain**                     |        |     |     |     |
| Right lingual gyrus          | 11827  | -18 | +90 | -4  |
| Left hippocampus             | 337    | +26 | +10 | -16 |
| Left mid orbital gyrus       | 301    | +2  | -54 | -12 |
| Right medial temporal pole   | 124    | -50 | -14 | -28 |
| Right Rolandic operculum     | 80     | -42 | +14 | +20 |
| Right SMA                    | 75     | -6  | +10 | +52 |
| Right angular gyrus          | 42     | -58 | +66 | +28 |
| **Loss**                     |        |     |     |     |
| Right insula lobe            | 4508   | -34 | -22 | +0  |
| Right inferior parietal lobule| 3604  | -42 | +42 | +48 |
| Left paracentral lobule      | 531    | +6  | +30 | +64 |
| Left middle frontal gyrus    | 65     | +50 | -38 | +20 |
| Right cerebellum             | 28     | -42 | +82 | -36 |
| Left superior frontal gyrus  | 25     | +18 | -46 | +48 |
Table 3. Brain regions including the OFC that represent advantageous choice.

| Brain regions                  | Voxels | x  | y  | z  |
|-------------------------------|--------|----|----|----|
| Right inferior occipital gyrus| 497    | -38| +82| -16|
| Right superior medial gyrus   | 157    | -10| -66| +20|
| Left superior orbital gyrus   | 135    | +14| -62| -8 |
| Right superior orbital gyrus  | 130    | -14| -38| -20|
| Left superior temporal gyrus  | 72     | +54| +38| +20|
| Right middle temporal gyrus   | 59     | -54| +38| +4 |
| Left insula lobe              | 46     | +42| -6 | -12|
| Right paracentral lobule      | 45     | -2 | +38| +76|
| Right superior occipital gyrus| 42     | -30| +78| +40|
| Left inferior occipital gyrus | 39     | +46| +74| -8 |
| Right temporal pole           | 36     | -62| -2 | +0 |
| Right temporal pole           | 28     | -54| -14| -16|
| Right angular gyrus           | 27     | -50| +70| +36|
| Left superior parietal lobule | 27     | +22| +46| +64|
| Left superior parietal lobule | 26     | +30| +66| +64|
| Left inferior occipital gyrus | 22     | +18| +98| -8 |
| Left temporal pole            | 20     | +58| -10| -4 |
| Left postcentral gyrus        | 20     | +66| +14| +16|
Figures

Figure 1

Experimental paradigm of the Iowa gambling task.
Figure 2

BOLD activity in the OFC was correlated with value signals.
The OFC represents advantageous choice, and choice-related activations in the OFC are not correlated with value signals.
Figure 4

The OFC represents an advantageous choice, and the OFC decoding accuracy is correlated with behavioural performances.
Figure 5

The OFC is functionally connected with the superior medial gyrus for choice, but not for values.

Supplementary Files

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