Accumulation of evidence during decision making in OCD patients

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Decision-making often entails the accumulation of evidence. Previous studies suggested that people with obsessive-compulsive disorder (OCD) process decision-making differently from healthy controls. Both their compulsive behavior and obsessive thoughts may influence the evidence accumulation process, yet the previous studies disagreed on the reason. To address this question, we employed a probabilistic reasoning task in which subjects made two alternative forced choices by viewing a series of visual stimuli. These stimuli carried probabilistic information toward the choices. While the OCD patients achieved similar accuracy to the control, they took longer time and accumulated more evidence, especially in difficult trials in which the evidence strength was low. We further modeled the subjects’ decision making as a leaky drifting diffusion process toward two collapsing bounds. The control group showed a higher drifting rate than the OCD group, indicating that the OCD group was less sensitive to evidence. Together, these results demonstrated that the OCD patients were less efficient than the control at transforming sensory information into evidence. However, their evidence accumulation was comparable to the healthy control, and they compensated for their decision-making accuracy with longer reaction times.

KEYWORDS
obsessive-compulsive disorder (OCD), decision-making, evidence accumulation, drift diffusion model, probabilistic reasoning

Introduction

Obsessive-compulsive disorder (OCD) is a common psychiatric disorder in which a patient experiences either uncontrollable intrusive thoughts, repetitive behaviors, or both (1). A broad spectrum of behavior anomalies have been reported in OCD patients, including but not limited to deficits in working memory and long-term memory (2–6), intolerance of uncertainty (7–18), and abnormal motor inhibition (19). Understanding the neural substrate of these behavior anomalies is essential, as neural modulation treatment targeting specific brain regions has been growing more common. Appropriate
behavior paradigms, especially those whose neural mechanisms have been carefully studied in humans and animals, may help reveal relevant neural circuitry in OCD patients’ altered behavior.

In particular, numerous studies have been done to advance the understanding of the decision-making process in OCD patients, yet the results were mixed. Some studies showed that OCD patients had lower decision accuracy than the controls (20, 21) while others showed the opposite (22, 23). Besides the choice accuracy, it has been reported that OCD patients were slow in committing to a decision relative to the controls and accumulated more evidence in the process (20–24). However, this oversampling behavior among OCD patients was not replicated in several other studies (25, 26). Some even observed impulsive behavior in OCD patients, whose decisions were often hasty, risky, and based on weak evidence (27).

The inconsistency among these studies reflects the complex nature of decision-making. The decision-making process could be segmented into several stages, including sensory-to-evidence transformation, evidence accumulation, decision threshold crossing, and motor planning and execution. Different cognition processes and their corresponding neural circuitries are engaged during each stage. It has been shown that the orbitofrontal cortex (Sirunyan et al.) computes and transforms visual sensory inputs into evidence, and a frontoparietal network, including the dorsolateral prefrontal cortex and the posterior parietal cortex, is involved in integrating evidence in decision making (28–31). For OCD, many studies have also pointed to a prominent role of the OFC in OCD symptoms. It has been suggested that an unbalanced activation of the orbitofrontal-striatal circuit underlies OCD (32). Deep brain stimulation (DBS) studies in OCD patients show that targeting fibers from the OFC to the striatum leads to better treatment effects (33). Furthermore, optogenetic stimulations of the Sapap3 mutant mice’s lateral OFC (IOFC) neurons successfully inhibited their OCD-like repetitive self-grooming behavior (34). These results lead to our speculation that the sensory-evidence transformation during decision-making in OCD patients is compromised.

To test this hypothesis, we used a probabilistic reasoning task adapted from earlier animal studies to investigate the evidence accumulation process in OCD patients’ decision-making (35, 36). This task paradigm is advantageous over typical perceptual decision-making paradigms, such as the random dots motion task (24, 37–39), which are based on the discrimination of noisy stimuli and less ideal for isolating the cognitive component of decision making. In the task, the subjects needed to accumulate probabilistic evidence based on a sequence of visual stimuli to perform a two-alternative forced-choice task. The stimuli were easily discriminable arrow shapes of different directions and contrasts. Each stimulus provided a piece of evidence toward the choices, and the subjects needed to accumulate the evidence for making appropriate decisions. We studied the OCD patients’ performance and how stimuli and their associated evidence contributed to decisions. We found that the OCD patients were less sensitive to individual pieces of evidence than the healthy controls, yet they achieved a similar overall accuracy by collecting more evidence. As a result, their reaction times were significantly longer than healthy controls but only in the difficult trials. These results, combined with the previous findings of the prefrontal circuitry underlying evidence accumulation, hint an altered prefrontal–orbitofrontal in particular—functions underly decision-making in OCD patients.

Method

Participants

We collected behavior data from 28 healthy controls and 33 OCD patients at Shanghai Mental Health Center. The OCD patients’ diagnosis, recruitment, and psychiatric tests (Yale-Brown Obsessive-Compulsive Scale (Y-BOCS) (40), Hamilton Anxiety Rating Scale (HAM-A) (41), and Hamilton Depression Rating Scale (HAM-D) (42) were performed by the psychiatric professionals from Shanghai Mental Health Center. The inclusion criteria of the OCD patients were: (a) between 18 and 54 years old, with no gender restrictions; (b) diagnosed as OCD based on DSM-IV criteria; (c) Y-BOCS ≥ 16 points; (d) not under any medication for at least 8 weeks; (e) education level no less than junior high school; (f) sufficient vision to accomplish the experiment, and (g) ability to provide the informed consent. The exclusion criteria were: (a) diagnosed with DSM-IV Axis I other than OCD; (b) severe physical disease, central nervous system disease, or substance abuse; (c) pregnancy or nursing; (d) prominent suicidal ideation. The inclusion criteria for the control group were: (a) no obsessive and compulsive symptoms; (b) no psychotropic medication history; (c) no relatives within three generations of direct or collateral blood had any psychiatric disorder history; (d) sufficient vision to accomplish the experiment and (e) ability to provide the informed consent. The exclusion criteria of the control group were: (a) diagnosed with the DSM-IV Axis I psychiatric disorders; (b) diagnosed with the DSM-IV Axis II compulsive personality disorders; (c) pregnancy or nursing; (d) prominent suicidal ideation. The healthy control participants were recruited through open advertisement. All procedures followed the protocol approved by the Ethics Committee of Shanghai Mental Health Center. (No. 2017-02) (Shanghai, China).

We calculated the accuracy of each subject and excluded those that were outside of the 95% confidence interval. With this exclusion criterium, 26 healthy controls and 32 OCD patients were included in the rest of the analyses. The accuracies of the two excluded healthy controls and the one OCD patient were 0.55, 0.53, and 0.47, respectively.
**Task paradigm**

The subjects made two-alternative forced choices by pressing a left or a right button with their fingers (Figure 1A). In each trial, a stream of visual stimuli appeared sequentially at the center of a computer screen. The stimuli were sampled from a stimulus set of 10 arrow shapes, which faced either left or right and had five different contrast levels (Figure 1B). The subjects were asked to judge, on average, which group of arrows had higher contrast by pressing the corresponding button.

Each arrow provides a piece of evidence toward the correct choice because its appearance probability depended on the choice. The probabilities of the arrows associated with the left choice were \([0.044, 0.083, 0.15, 0.17, 0.22, 0.18, 0.084, 0.048, 0.017, \text{ and } 0.006]\), and those with the right choice were \([0.006, 0.017, 0.048, 0.084, 0.18, 0.22, 0.17, 0.15, 0.083, \text{ and } 0.044]\). Thereby, the evidence associated with the arrows can be quantified by the evidence weights, which are defined as the natural log-ratio between its appearance probability associated with the right choice and that associated with the left choice. They were \([-0.9, -0.7, -0.5, -0.3, -0.1, 0.1, 0.3, 0.5, 0.7, \text{ and } 0.9]\) (Figure 1B). The five left arrows had negative weights; they were evidence favoring the left choice. The five right arrows had positive weights, which were in favor of the right choice. In addition, arrows with higher contrasts were associated with larger absolute evidence weights, indicating stronger evidence for their corresponding choices.

The trial sequence was as follows. A circle with a 25-pixel diameter appeared on the center of the screen indicating the trial start. After 0.5 s, two round targets with a diameter of 62.5 pixels appeared on the left and the right side of the central circle. After another 0.3 s delay, the central circle disappeared, and a serial
of arrows was displayed at the center of the screen sequentially. Each arrow was presented for 150 ms with no inter-stimulus interval. The maximal length of the stimulus sequence was 100. The participants were allowed to make choices throughout the stimulus viewing period by pressing the left or the right key. Trials were aborted if a choice was not indicated before the maximum number of stimuli was run out. A green check mark would appear on the screen center for 0.5 s if the choice were correct. Otherwise, a red cross sign would appear instead. The subjects first took twenty practice trials and then completed a hundred test trials. The test session typically took <20 min. The subjects received a small amount of cash (RMB 80 yuan) for their participation in the experiment. The amount of payment was unrelated to their performance.

**Behavioral analyses**

**Non-decision stimulus number**

The last few stimuli in each trial might not contribute to the decision due to the motor preparation and other downstream processes (35). To estimate the number of non-contributing stimuli, we modeled the subjects’ choices by segmenting the arrow sequence into two:

$$\text{choice} = \logit(\beta_0 + \beta_1 \sum_{i=1}^{N-n_{nd}} w_i + \beta_2 \sum_{i=1}^{n_{nd}} w_{N-n_{nd}+1}),$$

where $w_i$ is the weight of the $i$th stimulus, $n_{nd}$ is the number of non-contributing arrows, and $N$ is the length of the stimulus sequence ($N > n_{nd}$). We began with $n_{nd} = 1$ and increased $n_{nd}$ until $\beta_2$ reached the significant level in the model. We defined the non-decision stimulus number using the largest $n_{nd}$ without $\beta_2$ reaching significance and removed the last $n_{nd}$ stimuli from the subject's trial sequences in the following analysis. This non-decision time was independently estimated for each participant. We used $N^*$ to indicate the length of the adjusted stimulus sequence. Trials with length equal or $< N^*$ were excluded in the rest of the analyses.

**Psychometric curve and subjective value**

The psychometric curve was a logistic regression fitting to the subjects’ choice:

$$P(\text{Choice} = \text{Right}) = \frac{1}{1 + e^{-Q}},$$

$$Q = \beta_0 + \beta_1 \sum_{i=1}^{N^*} w_i,$$

where $w_i$ represents the weight of the arrow in epoch $i$, $\beta_0$ is the bias term, and $\beta_1$ is the estimate of how the total weight affected the choice. The estimate reflects the sensitivity of the subjects’ choice to the accumulated evidence.

We performed another logistic regression to assess how each type of arrow affected the choice:

$$P(\text{Choice} = \text{Right}) = \frac{1}{1 + e^{-Q}},$$

$$Q = \beta_0 + \sum_{i=1}^{10} \beta_i N_i,$$

where $N_i$ represents how many times arrow type $i$ appears in a trial, $\beta_0$ is the bias term, and $\beta_1$ are the estimates of how much leverage each type of arrow has on the subjects’ choices and termed the subjective weights of evidence (36).

**Trial difficulty**

To measure the evidence strength of a trial, we regressed the accumulated evidence at each presentation of the arrows against its index with a linear model:

$$\sum_{i=1}^{T} w_i = \beta_0 + \beta_1 T,$$

where $\sum_{i=1}^{T} w_i$ is the sum weight up to the $T$:th arrow ($T = 1. N^*$). Thus, $\beta_1$ is a measure of how fast the evidence is accumulated with which we defined the trial’s evidence strength. The evidence strength is a signed number. The sign indicates the choice that the evidence favors, and the absolute value indicates the strength.

We divided all trials into 8 bins based on their evidence strength for analyses in Figure 3. We used smaller bins for evidence strengths smaller than 0.2 and larger bins for those higher than 0.2, because the trials with evidence strength smaller than 0.2 were over-represented in the data. Thereby, each bin has enough trials, while it does not cover too broad a range of evidence strength (Supplementary Table 1). Our conclusions did not rely on the particular choice of this binning method.

**Drift diffusion model**

We fitted each subject’s choice and reaction time with the DDM using the pyddm toolbox (43). We included symmetric exponentially collapsing (rate $r$) bounds (bound height $B$) and a leaky (leaky rate $\lambda$) integrator in the model. To speed up the fitting, we used a constant drifting rate in each trial, which was proportional to the evidence strength, with a sensitivity constant $C$. A non-decision time parameter ($t_{nd}$) was also included. Additionally, we included a fixed lapse term $L$, which is the proportion of trials with choices generated from an evidence-independent Poisson process with the rate $= 1/sec.$
History effects

The influence of the trial history on the current choice (left/right) was studied with logistic regression:

$$choice = \text{logit}(\beta_0 + \beta_1 \sum_{i=1}^{N_i} w_i + \beta_2 C_i + \beta_3 R_i + \beta_4 C_i \times R_i),$$

(7)

where $\sum_{i=1}^{N_i} w_i$ is the accumulated evidence of the current trial, $C_i$ is the last trial’s choice (left/right), $R_i$ is the outcome of the last trial (correct/wrong), and the last term is the interaction between the two.

Data availability statement

The data that supports the findings of this study are available from the corresponding author upon reasonable request.

Results

We tested 28 healthy controls and 33 OCD patients with a probabilistic reasoning task (Figure 1A). The task, adapted from a previous study (35), required the subjects to make two-alternative choices based on a sequence of stimuli. In each trial, the computer first randomly decided on a correct answer. The subjects then watched a sequence of visual stimuli displayed on a computer screen. The stimuli were arrows that were sampled randomly with replacement from a pool of ten arrows, the distribution of which depended on the correct answer (Figure 1B). Thereby, each arrow $A_i$ was associated with two likelihoods ($P[A_i|\text{left}]$ and $P[A_i|\text{right}]$), describing how likely one may observe the arrow given the correct answer being the left or the right. The subjects reported their choice by pressing the left or the right button whenever they felt ready. The sequence was cut off at the 100th arrow, after which the trial would be aborted if the subjects did not make a response.

We defined the arrows’ weight as to the log ratio between the two likelihoods. The five right arrows had positive weights, indicating that they were evidence supporting the right choice. The five left arrows had negative weights and supported the left choice. Unlike the previous studies (35), in which the associations between the stimuli’s visual features and their evidence weights were arbitrary, we used the arrow directions and contrasts to indicate their weights of evidence: larger contrasts were assigned with weights of larger absolute values, indicating stronger evidence. This design reduced the training process in which the subjects had to learn the weights assigned to the stimuli. Subjects only needed to combine evidence from the arrows in a trial by calculating the sum of the weights associated with them. The sum of the weights is the log odds between the right choice and the left choice being the correct choice (Eqs. 2 and 3).

Most of the participants from both groups learned the task successfully. We excluded participants whose performance accuracy was outside of the 95% confidence interval. With this criterion, we included 26 healthy controls and 32 OCD patients in the following analysis. There were no significant differences between gender, age and the education of the two groups (Table 1; gender: $x^2(1, N = 58) = 0.0056, p = 0.94$; chi-square test; age: $t(56) = -0.98, p = 0.33$, education: $t(56) = 1.11, p = 0.27$, $t$-test). There was no significant difference in either the mean or the variance between the accuracies of the OCD and the healthy control group (Figure 2A, Control (mean±SEM): 0.72 ± 0.01, OCD: 0.76 ± 0.02; $t(56) = -1.72, p = 0.092$, $t$-test; effect size = −0.447; 95%CI: [−0.970, 0.087], Hedges’ g; Levene’s test, $p = 0.246$).

Next, we examined how the subjects accumulated evidence for decision-making. The evidence was furnished with a stream of stimuli in each trial. However, because the subjects made decisions while a continuous stream of stimuli was presented, the last few stimuli before the button pressing likely did not contribute to the decision-making due to the extra processing required by the brain after a decision was made. We used logistic regression to estimate the number of stimuli that did not contribute to decision making, we found that the OCD patients looked at significantly more stimuli than the healthy controls before they committed to their decisions (Figure 2B, OCD: 22.54 ± 1.96, Control: 13.61 ± 1.52; $t(56) = -3.47, p < 1e^{-3}$, $t$-test; effect size

### Table 1 Characteristics of the participants.

|          | Age  | Gender (M/F) | Education | FOA | Dur | YB-O  | YB-C  | YB-T  | HAM-D | HAM-A |
|----------|------|--------------|-----------|-----|-----|-------|-------|-------|-------|-------|
| Control  | 26.27 (7.78) | 16/10 | 16.12 (2.18) | n.a. | n.a. | 0.77 (1.07) | 0.50 (0.99) | 1.27 (1.78) | 1.35 (1.62) | 0.81 (0.85) |
| OCD      | 28.25 (7.53) | 20/12 | 15.38 (2.77) | 21.31 (7.25) | 7.28 (5.34) | 11.00 (2.99) | 9.91 (3.12) | 7.91 (5.22) | 9.91 (3.12) | 20.91 (4.54) | 7.91 (5.22) | 9.91 (3.12) |

Table above listed the mean and the standard deviations (in parentheses). FOA, first onset age; Dur, illness duration; YB-O, Y-BOCS-obession; YB-C, Y-BOCS-compulsion; YB-T, Y-BOCS-total; HAM-A, Hamilton Anxiety Rating Scale; HAM-D, Hamilton Depression Rating Scale.
Comparison between the OCD and the control group. (A) Accuracy. The boxes indicate the second and third quartiles, and the whiskers indicate a 99.3% interval. (B) Reaction time is measured with the number of stimuli used for decisions. (C) The sum weight of the stimuli used for decisions. (D) The choice was plotted as a function of sum weight. The data points and the error bars indicated the mean and the S.E.M. across subjects. The curves were based on the logistic regression fitting to all trials of each group’s subjects. (E) The logistic regression coefficient β1 of individual subjects. (F) The logistic regression coefficient β0 (intercept) of individual subjects. (G) The subjective weights plotted against the assigned weights for each type of arrow. The data points and the error bars indicated the mean and the S.E.M. across subjects. (H) The slopes of the fitted lines of the subjects’ subjective weights. (I) The logistic regression coefficient β0 (intercept) of individual subjects. The gray dots indicated subjects with slopes not significantly different from zero, who were excluded from the analysis **p < 0.01, ***p < 0.001.

The fact that the OCD patients accumulated more evidence to achieve similar performance to the control group pointed to a lower decision-making efficiency in the OCD patients. The decision efficiency difference is also evident in the psychometric curves, which were steeper in the control group than in the OCD group (Figure 2D). A logistic regression model of the subject’s choice revealed that the total evidence exerted less leverage on the OCD patients’ choices than those of the healthy controls (Figure 2E, OCD: 1.03 ± 0.14, Control: 1.73 ± 0.14; z = 3.12, p = 0.0018, Wilcoxon rank-sum test), which leads to a less steep psychometric curve for the OCD patients. The OCD patients’ performance was also more variable than the controls (Levene’s test, p = 0.0446). Neither group showed a significant choice bias (Figure 2F, OCD(mean±standard deviation): −0.02 ± 0.36, t(31) = −0.30, p = 0.77; Control: −0.10 ± 0.50,
difficulty became overwhelming (moderately difficult, but they collected less evidence when the evidence became weaker and the trials became harder except for the last bin). Therefore, each piece of evidence contributed less to the choices in the OCD patients than in the healthy controls, suggesting the patients were less sensitive to the evidence. Again, the subjects in neither group exhibited a choice bias (Figure 2I; OCD: 1.07 ± 0.07; control: 1.39 ± 0.08; p = 0.0028, t-test; Levene’s test, p = 0.280). Therefore, the OCD patients accumulated a comparable amount of evidence. The trial difficulty can be quantified by how fast the evidence grows in a trial, which we term evidence strength. We regressed the accumulated evidence weight against the stimulus index in each trial, excluding the non-contributing arrows, and defined the evidence strength as the slope of the regression line (Eq. 6). We grouped the trials by their evidence strengths. At each level of evidence strength, the choice accuracy did not show any significant difference between the two groups (Figure 3A). During the easiest trials, the OCD patients and the controls also took a similar amount of time to make their decisions (Figure 3B, last bin). However, the OCD patients spent significantly more time when the evidence strength was weak (Figure 3B). The difference became gradually larger as the evidence became weaker and the trials became harder except in the most difficult trials. When we looked at the amount of accumulated evidence at the time of the decision, both groups showed a non-monotonic pattern with the evidence strength increased. Subjects from both groups accumulated increasingly more evidence as the task difficulty elevated from easy to moderately difficult, but they collected less evidence when the difficulty became overwhelming (Figure 3C). During the easiest trials (the last bin in Figure 3C), the OCD and the control group accumulated a comparable amount of evidence. The trial numbers and participant numbers included in each were displayed in (Supplementary Table 1).

In addition, we observed no repetitive choice pattern from the OCD subjects. When we regressed the subjects’ choice against the previous trial’s choice, reward outcome, and their interactions (Eq. 7), we did not find any historical influences in either group at the group level (Supplementary Figure 2).

These results indicated that the OCD patients had longer reaction times. It could be because they required more evidence for making decisions, or it could also be each stimulus provided a smaller amount of evidence, or both. The former means a higher decision threshold, and the latter leads to a slower evidence accumulation rate. To gain further insights into a mechanistic explanation of the OCD patients’ decision-making, we used the drift-diffusion model (DDM) to infer the subjects’ evidence accumulation process during the decision-making (44). The DDM is a model for explaining both the choice and reaction time behavior in two-alternative choice tasks. In the DDM, the decision-making is modeled as a particle drifting between two bounds. The drifting rate depends on the strength of the evidence, and the decisions are made when the particle hits one of the two bounds. We modeled the subjects’ decision-making process using the drift-diffusion model (DDM) to infer the subjects’ evidence accumulation process during the decision-making (44). The DDM is a model for explaining both the choice and reaction time behavior in two-alternative choice tasks. In the DDM, the decision-making is modeled as a particle drifting between two bounds. The drifting rate depends on the strength of the evidence, and the decisions are made when the particle hits one of the two bounds.
with a DDM in which the evidence accumulation process is leaky and the bounds collapse over time (Figure 4A). To simplify the model and speed up the fitting, the drifting rate in each trial is set to be fixed, which is proportional to the evidence strength, with the evidence sensitivity being the proportionality constant. The bounds collapse exponentially with a time constant τ. The details of the fitting procedures were described previously (45).

The model fitting results supported the second hypothesis that the longer reaction time of the OCD group was due to a less efficient evidence accumulation process. We found that the decision bound was similar between the two groups (Figure 4B, OCD (mean±SEM): 1.11 ± 0.26, control: 1.14 ± 0.27; t(56) = 0.09, p = 0.931, two-sample t-test). The bound collapsing rate τ was also comparable (Figure 4C, OCD: 1.16 ± 0.19, control: 1.68 ± 0.38; z = 0.11, p = 0.91, Wilcoxon rank-sum test). What was significantly different between the two groups was the evidence sensitivity, which was larger in the control (Figure 4E, OCD: 17.03 ± 1.17, control: 20.91 ± 1.45; t(56) = 2.10, p = 0.04, two-sample t-test). Therefore, the evidence was accumulated more quickly and reached the decision bound faster in the control group, leading to a shorter reaction time. The accumulation leak rate showed no difference between the OCD patients and the control (Figure 4D, OCD: −5.63±1.67, control: −3.61 ± 2.34; t(56) = 0.72, p = 0.475, two-sample t-test). We also did not find a difference in the non-decision time (Figure 4F, OCD: 0.040 ± 0.008, control: 0.040 ± 0.009; t(56) = −0.17, p = 0.86, two-sample t-test), which accounted for the time for extra sensory and motor processing for decision making, suggesting the OCD patients and the controls were similar in sensory processing and action execution speed.

We further used DDM with constant bound to fit the subjects’ behavior (Supplementary Figure 3). As above in the model with collapsing bounds, the evidence sensitivity again showed a significant group difference (OCD (mean±SEM): 15.26 ± 0.98, control: 18.19 ± 1.04; t(56) = 2.03, p = 0.047, two-sample t-test). In addition, the leak rate was also found to be slightly but significantly smaller in the OCD group.
than in the control (OCD: 3.50 ± 0.84, control: 4.70 ± 1.96; z = 2.31, p = 0.021, Wilcoxon rank sum test). Bound height (OCD: 1.34 ± 0.13, control: 1.20 ± 0.16; t(56) = −0.69, p = 0.49, two-sample t-test) and non-decision time (OCD: 0.019 ± 0.006, control: 0.025 ± 0.007; t(56) = 0.63, p = 0.53, two-sample t-test) were comparable between the two groups. The DDM with collapsing bound was not significantly better than the DDM with constant bound in either the control or the OCD group (Supplementary Figure 4). (∆AICcontrol = 1.70, t(50) = −0.05, p = 0.96; ∆AICOCD = 3.62, t(62) = −0.15, p = 0.88; ∆BICcontrol = −0.89, t(50) = 0.03, p = 0.98; ∆BICOCD = 1.04, t(62) = −0.04, p = 0.97).

Finally, we studied how the individual variations of the OCD patients’ performance may correlate with their disease symptoms’ severity and characteristics (Supplementary Table 2). Each patient completed the tests of the Yale-Brown Obsessive-Compulsive Scale (Y-BOCS) (40), Hamilton Anxiety Rating Scale (HAM-A) (41), and Hamilton Depression Rating Scale (HAM-D) (42). We observed several trends that may indicate a correlation between the patients’ decision-making and their symptoms. In particular, the patients with higher compulsive symptoms, measured with Y-BOCS-C had less accuracy (Pearson’s r = −0.31, p < 0.1) and shorter reaction time (Pearson’s r = −0.31, p < 0.1). Similarly, the patients’ HAM-A scores were also negatively correlated with both the reaction time (Pearson’s r = −0.32, p < 0.1) and the sum weight (Pearson’s r = −0.32, p < 0.1), as well as a positive correlation between the HAM-A score and the decision bound’s collapsing rate (tau) (Pearson’s r = 0.4, p < 0.05). The results indicate that higher anxiety levels may lead to hasty decisions in OCD patients. However, none of these correlations survive after multiple-comparison corrections. One possible reason is that the OCD cohort had a small range of symptom severity level, which is not sufficient to reveal the correlation between the symptom severity and the decision-making deficits. In addition, the patients’ impaired sensory-evidence transformation may affect their symptoms via a complicated mechanism that cannot be detected via a simple correlation analyses. Further investigations are necessary to understand the relationship between the OCD symptoms and the altered cognitive functions.

**Discussion**

We studied and compared the evidence accumulation during decision-making in OCD patients and controls. Consistent with previous studies, the OCD patients had longer reaction times than the healthy controls but achieved similar overall accuracy by collecting more evidence (24). However, the difference in the reaction time was not evident when the trials were easy. We demonstrated that the longer reaction times were due to OCD patients’ less efficiency in using the evidence for decision making.

The behavior paradigm used in our study has several distinct features. First, unlike the random dots motion task, which has been extensively used in both primate and human studies (24, 37, 39), the stimuli were highly discriminable in the current experiment. Therefore, our results are not confounded by the perceptual limits. In addition, unlike a monkey study with a similar concept (35), the stimulus-evidence association was not entirely arbitrary. Instead, the arrows of different directions and contrast levels indicated the evidence strength and the correct target. Thereby, we lowered the memory demand and speeded up the learning of the task. This is particularly helpful in the patient study due to a stricter limit of trial numbers. Finally, the amount of evidence can be directly calculated from the weights of the stimuli that appear in each trial, allowing us to explore the evidence accumulation process during decision making.

In particular, the accumulated evidence at the time of the decision, measured with the sum of weights, is not the same as the decision bound determined by the DDM. While the decision bound reflects an internal threshold used for decision making, the evidence weight, and its sum is an objective measure of the amount of evidence collected. The evidence sensitivity determines how external evidence is transformed and used in the brain for decision making. We used a DDM to show that the evidence sensitivity differed between the OCD patient group and the healthy control group, suggesting that the same amount of evidence contributed less to decision-making in the OCD patients.

This deficiency in OCD patients does not necessarily mean that they were worse at perceiving visual stimuli. There is no strong evidence suggesting that OCD patients’ sensory perception is compromised. Instead, we believe that it reflects a deficit in the transformation from the sensory stimuli into the evidence in OCD patients. The same stimulus appears to be less informative to the patients than to the healthy controls, which may lead to their apparent indecisiveness.

By calculating the sum weight at the decision time, we found that the total accumulated evidence first increased with the trial difficulty but collapsed at extreme difficulty levels. This trend was not seen in the reaction time, which increased monotonically with the trial difficulty. At the easiest difficulty level, the OCD patients accumulated a similar amount of evidence to that of the healthy controls. Yet, the increase of the accumulated evidence with the trial difficulty was significantly higher among the OCD patients than among the healthy controls, reflecting that the OCD patients’ decision-making is more impaired under challenging situations.

Greater trial difficulties lead to more uncertain and less confident decisions. The estimation of confidence may allow the brain to adjust decision strategies dynamically during decision-making. It has also been proposed that confidence may
be estimated in a metacognition process based on the same evidence accumulation process during the decision making (46, 47). Also, a prefrontal network that includes the orbitofrontal cortex and the anterior cingulate cortex was implicated (48–50). In particular, a recent study using a similar task showed that the orbitofrontal cortex encodes evidence associated with each stimulus, while the dorsolateral prefrontal cortex transforms evidence into action value and accumulates evidence for making motor responses (28). Our results suggest a deficit in the orbitofrontal cortex among OCD patients, which has been demonstrated previously (51–53).

Therefore, we suspect that an abnormal interplay between the prefrontal metacognition circuitry and the decision-making circuitry, which includes the posterior parietal cortex and basal ganglia, underlies the observed decision-making behavior in OCD patients. Consistent with this idea, a recent study showed that capsulotomy, which affected cortical-basal-ganglia projections, could rectify some of the decision-making problems in OCD patients (54).

We only observed weak correlations between the OCD patients’ decision-making and the severity of their symptoms, which were measured with Y-BOCS, HAM-A, and HAM-D scores. These correlations did not point to a consistent behavior pattern. For example, while higher compulsivity seems to lead to shorter reaction times and worse accuracies, patients with higher anxiety levels, although also exhibiting shorter reaction times, were more sensitive to the evidence, and their accuracies were less affected. The link between the patients’ symptom and their decision-making behavior may be complex, and multiple factors may be in play. Further studies are necessary to confirm these findings.

In conclusion, we used a novel probabilistic reasoning task to explore the decision-making behavior in OCD patients. The results revealed that the deficiency in OCD patients’ decision-making is due to an inefficient accumulation of evidence. With decision bounds similar to those of the controls, the OCD patients, however, could compensate for this deficiency with longer reaction times to reach comparable accuracies.

Data availability statement

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

Ethics statement

The studies involving human participants were reviewed and approved by the Ethics Committee of Shanghai Mental Health Center (Shanghai, China). The patients/participants provided their written informed consent to participate in this study.

Author contributions

TY, ZW, QF, and YC: conceptualization. YL: data acquisition. TY, YC, QF, and YL: data analysis. TY and YC: drafting. TY, QF, and YC: final approval. TY and QF: funding and supervision. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpsyt.2022.980905/full#supplementary-material
References

1. American Psychiatric Association. Diagnostic and Statistical Manual of Mental Disorders: DSM-5. Washington, DC: American Psychiatric Association (2013).

2. Chamberlain SR, Blackwell AD, Fineberg NA, Robbins TW, Sahakian BJ. The neuropsychology of obsessive compulsive disorder: the importance of failures in cognitive and behavioural inhibition as candidate endophenotypic markers. Neurosci Biobehav Rev. (2005) 29:399–419. doi: 10.1016/j.neubiorev.2004.11.006

3. Fontenelle L, Mendowicz M, Mattosc P, Versiani M. Neuropsychological findings in obsessive-compulsive disorder and its potential implications for treatment. Curr Psychiatry Rev. (2006) 2:11–26. doi: 10.2174/157340060775101454

4. Hermans D, Martens K, De Cort K, Pieters G, Eelen P. Reality monitoring and metacognitive beliefs related to cognitive confidence in obsessive-compulsive disorder. Behav Res Ther. (2003) 41:383–401. doi: 10.1016/S0005-7967(02)00015-3

5. Kielcz AK, Hohagen F, Voderholzer U. Neuropsychological performance in obsessive-compulsive disorder: a critical review. Biol Psychiatry. (2004) 65:185–236. doi: 10.1016/j.bip.2003.07.007

6. Olson CA, Hale LR, Hamilton N, Powell JN, Martin LE, Savage CR. Altered source memory retrieval is associated with pathological doubt in obsessive-compulsive disorder. Behav Brain Res. (2016) 296:53–60. doi: 10.1016/j.bbr.2015.08.031

7. Dar R. Facilitating the mechanism of uncertainty and doubt in obsessive-compulsive checkers. J Behav Ther Exp Psychiatry. (2004) 35:153–63. doi: 10.1016/j.jbtep.2004.04.006

8. Dar R, Rush S, Herrmich H, Taub M, Fux M. Realism of confidence in obsessive-compulsive checkers. J Abnorm Psychol. (2000) 109:673–8. doi: 10.1037/0021-843X.109.4.673

9. Holaway RM, Heimberg RG, Coles ME. A comparison of intolerance of uncertainty in analogue obsessive-compulsive disorder and general anxiety disorder. J Anxiety Disord. (2006) 20:158–74. doi: 10.1016/j.janxdis.2005.01.002

10. Lazarov A, Dar R, Oded Y, Liberman N. Are obsessive-compulsive tendencies related to reliance on external proxies for internal states? Evidence from biofeedback-aided relaxation studies. Behav Res Ther. (2010) 48:516–23. doi: 10.1016/j.brat.2010.02.007

11. Lazarov A, Dar R, Liberman N, Oded Y. Obsessive-compulsive tendencies and undermined confidence are related to reliance on proxies for internal states in a false belief paradigm. J Behav Exp Psychiatry. (2012) 43:556–64. doi: 10.1016/j.jbep.2011.07.007

12. Nestadt G, Kamath M, Maher BS, Krasnow J, Nestadt P, Wang Y, et al. Increased decision thresholds enhance information gathering performance in juvenile Obsessive-Compulsive Disorder (OCD). PLoS Comput Biol. (2017) 13:e1005440. doi: 10.1371/journal.pcbi.1005440

13. Volano PJ. Styles of decision-making and probability appraisal in selected obsessional and phobic patients. Br J Soc Clin Psychol. (1976) 15:305–17. doi: 10.1111/j.2044-8260.1976.tb00388.x

14. Banca P, Vestergaard MD, Rankov V, Baek K, Mitchell S, Lapa T, et al. Evidence accumulation in obsessive-compulsive disorder: the role of uncertainty and monetary reward on perceptual decision-making thresholds. Neuropsychopharmacology. (2015) 40:1192–202. doi: 10.1038/npp.2014.303

15. Chamberlain SR, Fineberg NA, Blackwell AD, Clark L, Robbins TW, Sahakian BJ. A neuropsychological comparison of obsessive-compulsive disorder and trichotillomania. Neuropsychologia. (2007) 45:654–62. doi: 10.1016/j.neuropsychologia.2006.07.016

16. Jacobsen P, Freeman D, Salkovskis P. Reasoning bias and belief conviction in obsessive-compulsive disorder and delusions: jumping to conclusions across disorders? Br J Clin Psychol. (2012) 51:84–99. doi: 10.1111/j.2044-8260.2011.02014.x

17. Grassi G, Pallanti S, Righi L, Figer M, Mantione M, Denys D, et al. Think twice: Impulsivity and decision making in obsessive-compulsive disorder. J Behav Addict. (2015) 4263–72. doi: 10.1037/bja0000039

18. Lin Z, Nie C, Zhang Y, Chen Y, Tang T. Evidence accumulation for value computation in the prefrontal cortex during decision making. Proc Natl Acad Sci U S A. (2020) 117:30728–37. doi: 10.1073/pnas.2019077117

19. Roitman JD, Shadlen MN. Response of neurons in the lateral intraparietal area during a combined visual discrimination reaction time task. J Neurosci. (2002) 22:9475–89. doi: 10.1523/JNEUROSCI.22-21-09475.2002

20. Shadlen MN, Kiani R. Decision making as a window on cognition. Neuron. (2013) 80:791–806. doi: 10.1016/j.neuron.2013.10.047

21. Padoa-Schioppa C. Neurobiology of economic choice: a good-based model. Annu Rev Physiol. (2013) 74:333–59. doi: 10.1146/annurev-physiol-021113-113844

22. Gunaydin LA, Kreitzer AC. Cortico-basal ganglia circuit function in psychiatric disease. Annu Rev Physiol. (2016) 78:327–50. doi: 10.1146/annurev-physiol-021115-105355

23. Alonso P, Cuadras D, Gabriels L, Denys D, Goodman W, Greenberg BD, et al. Deep Brain stimulation for obsessive-compulsive disorder: a meta-analysis of treatment outcome and predictors of response. PLoS ONE. (2015) 10:e0133591. doi: 10.1371/journal.pone.0133591

24. Burguiere E, Monteiro P, Pong G, Graybiel AM. Optogenetic stimulation of lateral orbitofronto-striatal pathway suppresses compulsive behaviors. Science. (2013) 340:1243–6. doi: 10.1126/science.1232380

25. Kira S, Yang T, Shadlen MN. A neural implementation of Wald’s sequential probability ratio test. Neuron. (2015) 85:861–73. doi: 10.1016/j.neuron.2015.01.007

26. Yang T, Shadlen MN. Probabilistic reasoning by neurons. Nature. (2007) 447:1075–80. doi: 10.1038/nature05852

27. Britten KH, Shadlen MN, Newsome WT, Movshon JA. The analysis of visual motion: a comparison of neuronal and psychophysical performance. J Neurosci. (1992) 12:4745–65. doi: 10.1523/JNEUROSCI.12-12-04745.1992

28. Erhan C, Balci F. Obsessive compulsive features predict cautious decision strategies. Q J Exp Psychol (Hove). (2017) 70:179–90. doi: 10.1177/1747021815513077

29. Ratcliff R, McKoon G. The diffusion decision model theory and data for two-choice decision tasks. Neural Comput. (2008) 20:873–922. doi: 10.1162/neco.2008.12-6-420

30. Goodman WK, Price LH, Rasmussen SA, Mazure C, Fleischmann RL, Hill CL, et al. The Yale-Brown Obsessive Compulsive Scale. I Development, use, and reliability. Arch Gen Psychiatry. (1989) 46:1006–11. doi: 10.1001/archpsyc.1989.01811101048007

31. Hamilton M. The assessment of anxiety states by rating. Br J Med Psychol. (1959) 32:50–5. doi: 10.1111/j.2044-8341.1959.tb04647.x

32. Hamilton M. The hamilton rating scale for depression. In: Assessment of Depression. Berlin, Heidelberg: Springer (1986). doi: 10.1007/978-3-642-08468-4_14

33. Shinn M, Lam NH, Murray JD. A flexible framework for simulating and fitting generalized drift-diffusion models. Elife. (2020) 9. doi: 10.7554/eLife.56938.sa2

Chen et al. 10.3389/fpsyg.2022.980905
44. Ratcliff R. A theory of memory retrieval. *Psychol Rev.* (1978) 85:59–108. doi: 10.1037/0033-295X.85.2.59

45. Shinn M, Ehrlich D, Lee D, Murray JD, Seo H. Confluence of timing and reward biases in perceptual decision-making dynamics. *J Neurosci.* (2020) 40:7326–42. doi: 10.1523/JNEUROSCI.0544-20.2020

46. Fetsch CR, Kiani R, Newsome WT, Shadlen MN. Effects of cortical microstimulation on confidence in a perceptual decision. *Neuron.* (2014) 83:797–804. doi: 10.1016/j.neuron.2014.07.011

47. Kiani R, Shadlen MN. Representation of confidence associated with a decision by neurons in the parietal cortex. *Science.* (2009) 324:759–64. doi: 10.1126/science.1169405

48. Bang D, Fleming SM. Distinct encoding of decision confidence in human medial prefrontal cortex. *Proc Natl Acad Sci U S A.* (2018) 115:6082–7. doi: 10.1073/pnas.1800795115

49. Beer JS, Lombardo MV, Bhanji JP. Roles of medial prefrontal cortex and orbitofrontal cortex in self-evaluation. *J Cogn Neurosci.* (2010) 22:2108–19. doi: 10.1162/jocn.2009.21359

50. Lak A, Costa GM, Romberg E, Koulakov AA, Mainen ZF, Kepecs A. Orbitofrontal cortex is required for optimal waiting based on decision confidence. *Neuron.* (2014) 84:190–201. doi: 10.1016/j.neuron.2014.08.039

51. Baxter LR Jr, Phelps ME, Mazzotta JC, Gune BH, Schwartz JM, Selin CE. Local cerebral glucose metabolic rates in obsessive-compulsive disorder: A comparison with rates in unipolar depression and in normal controls. *Arch Gen Psychiatry.* (1987) 44:211–8. doi: 10.1001/archpsyc.1987.01800150017003

52. Crespo-Facorro B, Cabranes JA, Lopez-Ibor Alcocer MI, Paya B, Fernandez Perez C, Encinas M, et al. Regional cerebral blood flow in obsessive-compulsive patients with and without a chronic tic disorder. A SPECT study. *Eur Arch Psychiatry Clin Neurosci.* (1999) 249:156–61. doi: 10.1007/s004060050081

53. Whiteside SP, Port JD, Abramowitz JS. A meta-analysis of functional neuroimaging in obsessive-compulsive disorder. *Psychiatry Res.* (2004) 132:69–79. doi: 10.1016/j.pscychresns.2004.07.001

54. Zhang C, Chen Y, Tian S, Wang T, Xie Y, Jin H, et al. Effects of anterior capsulotomy on decision making in patients with refractory obsessive-compulsive disorder. *Front Psychol.* (2017) 8:1814. doi: 10.3389/fpsyg.2017.01814