Willing to wait: Elevated reward-processing EEG activity associated with a greater preference for larger-but-delayed rewards

Narun Pornpattananangkul\textsuperscript{a,b,*}, Robin Nusslock\textsuperscript{a}

\textsuperscript{a} Department of Psychology, Northwestern University, Evanston, IL, United States
\textsuperscript{b} Department of Psychology, University of Singapore, Singapore

**ABSTRACT**

While almost everyone discounts the value of future rewards over immediate rewards, people differ in their so-called delay-discounting. One of the several factors that may explain individual differences in delay-discounting is reward-processing. To study individual-differences in reward-processing, however, one needs to consider the heterogeneity of neural-activity at each reward-processing stage. Here using EEG, we separated reward-related neural activity into distinct reward-anticipation and reward-outcome stages using time-frequency characteristics. Thirty-seven individuals first completed a behavioral delay-discounting task. Then reward-processing EEG activity was assessed using a separate reward-learning task, called a reward time-estimation task. During this EEG task, participants were instructed to estimate time duration and were provided performance feedback on a trial-by-trial basis. Participants received monetary-reward for accurate-performance on Reward trials, but not on No-Reward trials. Reward trials, relative to No-Reward trials, enhanced EEG activity during both reward-anticipation (including, cued-locked delta power during cue-evaluation and pre-feedback alpha suppression during feedback-anticipation) and reward-outcome (including, feedback-locked delta, theta and beta power) stages. Moreover, all of these EEG indices correlated with behavioral performance in the time-estimation task, suggesting their essential roles in learning and adjusting performance to maximize winnings in a reward-learning situation. Importantly, enhanced EEG power during Reward trials, as reflected by stronger 1) pre-feedback alpha suppression, 2) feedback-locked theta and 3) feedback-locked beta, was associated with a greater preference for larger-but-delayed rewards in a separate, behavioral delay-discounting task. Results highlight the association between a stronger preference toward larger-but-delayed rewards and enhanced reward-processing. Moreover, our reward-processing EEG indices detail the specific stages of reward-processing where these associations occur.

**1. Introduction**

Choosing between receiving $400 today or $800 today is easy. Most people, if not everyone, will select $800 today. However, choosing between receiving $400 today or $800 in three years is more difficult, and different people will choose differently. This latter decision becomes harder and requires a stronger computational demand (Rangel et al., 2008) because the subjective value of $800 is devalued, or discounted, over time. For decades, economists, psychologists, and, more recently, cognitive neuroscientists have studied this so-called delay-discounting phenomenon (also known as temporal discounting or inter-temporal choices; Ainslie, 1975; Frederick et al., 2003; Kalenscher and Pennartz, 2008; Peters and Büchel, 2011; Samuelson, 1937; Schultz, 2010). While the phenomenon is well documented, it is clear that people vary in how much they discount future rewards. In fact, individual differences in delay-discounting are stable over time and are sometimes considered a personality trait (Kirby, 2009; Odum, 2011). Recently personality and cognitive-neuroscience research has shown that these individual differences in delay-discounting are correlated with several trait affective and cognitive variables (Civai et al., 2016; Hirsh et al., 2008; Mahalingam et al., 2014). Among these variables is reward-processing (Benningfield et al., 2014; Boettiger et al., 2007; Hariri et al., 2006), which relates to the value

**Abbreviations:** ANOVA, analysis of variance; COMT gene, Catechol-O-methyltransferase gene; EEG, electroencephalogram; ERSP, event-related spectral perturbation; FDR, false discovery rate; fMRI, functional magnetic resonance imaging; FMT, frontal-midline theta; FRN, feedback-related negativity; HEDG, horizontal electrooculography; IQR, inter-quartile range; ITI, inter-trial interval; L-OFC, left orbitofrontal cortex; MID task, Monetary Incentive Delay task; PCA, principle component analysis; VEOG, vertical electrooculography; VS, ventral striatum.

* Correspondence to: Department of Psychology, Faculty of Arts and Social Sciences, National University of Singapore, Block A54, #02-07, 9 Arts Link, Singapore 117570.

E-mail addresses: monnarun@u.northwestern.edu, psynn@nus.edu.sg (N. Pornpattananangkul).

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an individual places on potential rewards during both the expectation and receipt of that reward (McClure et al., 2004; Schultz et al., 2000). Yet, the exact nature of the relationship between delay-discounting tendencies and individual differences in reward-processing is still unclear, perhaps due to the multifaceted nature of reward-processing.

1.1. Reward-processing and individual differences in delay-discounting responses

An early study by Hariri et al. (2006) reported that individuals with elevated reward-related neural activation in the ventral striatum (VS) during an incentivized fMRI card-guessing task had a stronger preference toward smaller-but-immediate rewards, as indexed by a subsequent behavioral delay-discounting task. This finding suggests that enhanced reward-processing is related to a stronger preference toward smaller-but-immediate rewards. However, recent data suggests the opposite pattern, indicating that elevated reward-processing is associated with a preference for larger-but-delayed (as opposed to smaller-but-immediate) rewards. In line with this view, a recent fMRI study using the Monetary Incentive Delay (MID) task reported that elevated VS activation during reward-anticipation among adolescents was associated with a stronger preference for larger-but-delayed rewards on a subsequent behavioral delay-discounting task (Benningfield et al., 2014). This finding is consistent with other recent fMRI studies reporting that elevated activation in the VS is associated with a stronger preference toward larger-but-delayed rewards (Ballard and Knutson, 2009; Samanez-Larkin et al., 2011). This relationship is also in line with animal research showing that lesions to the VS lead to a preference for smaller-but-immediate choices (Cardinal et al., 2001). In addition to the VS, enhanced activity in another neural region implicated in reward processing, the lateral orbitofrontal cortex (L-OFC), has been associated with a stronger preference toward larger-but-delayed rewards during an fMRI delay-discounting task (Boettiger et al., 2007). Next, there is indirect evidence from research involving the Val158Met polymorphism of the catechol-O-methyltransferase (COMT) gene. The Met-allele of the COMT gene is associated with higher synaptic dopamine levels and carriers of this allele display enhanced VS activation in an fMRI reward task (Chen et al., 2004; Vucaburan et al., 2007). Critically, Met-allele carriers also show a preference toward larger-but-delayed rewards (Boettiger et al., 2007; Gianotti et al., 2012; Smith and Boettiger, 2012). More recently, research with Parkinson patients suggests that medications designed to elevate dopamine signaling are associated with a heightened preference for larger-but-delayed rewards (Foerde et al., 2016). Thus, taken as a whole, evidence from fMRI, animal, genetic, and pharmacological studies suggest that individuals with elevated reward-processing have a tendency to wait for larger rewards and forgo smaller-but-immediate rewards.\footnote{The relationship between elevated reward-processing and a stronger preference for larger-but-delayed rewards does not necessarily support or contradict one of the most cited theories in delay-discounting, called the visceral factor perspective by Loewenstein and colleagues (Frederick et al., 2003; Loewenstein, 1996, 2000). Loewenstein and colleagues proposed that visceral factors, or drive states (such as, hunger, thirst, emotions, moods, craving for drugs, etc.), may explain intra-individual variability in delay-discounting. These visceral factors are endogenous, and can be changed from moment to moment depending on one’s drive state (e.g., hunger level depending on how long ago a person ate food). To support his hypothesis, Loewenstein (1996) highlights research by Mischel (1974) and Mischel et al. (1989) showing that children prefer a smaller-but-immediate reward if the reward object (e.g., marshmallow) is placed in front of them. But children would prefer a larger-but-delayed reward if they see a photograph of the delayed reward object. Loewenstein (1996) interpreted these findings as suggesting that having the reward object in front of you enhances the drive state for the immediate rewards, and seeing the photograph of the delayed reward object enhances drive states for the delayed reward. The focus in the current study, however, is inter-individual relationships between reward-processing and delay-discounting. This is reflected in the use of separate reward-processing and delay-discounting tasks, and the fact that we did not manipulate the visceral factors of our delay-discounting task (see Section 2).}

The current study aimed to further test and substantiate the relationship between a stronger preference toward larger-but-delayed rewards and enhanced reward-processing by investigating the relationship at different temporal stages of reward-processing via electroencephalogram (EEG). Reward-processing is thought to be comprised of two temporal stages that are mediated by distinct neurobiological systems: reward-anticipation and reward-outcome (Berridge, 1996; Wise, 2008). The superior temporal resolution of EEG, compared to fMRI, allows researchers to more accurately dissociate neural-cognitive states that occur close to each other in time (Cohen, 2014; Luck, 2014), such as reward-anticipation and reward-outcome stages, as well as between different sub-stages within reward-anticipation itself (Brunia et al., 2011; Goldstein et al., 2006; McAdam and Seales, 1969). EEG, for instance, has been used to dissociate reward-anticipation from motor-preparation (Brunia et al., 2011; Hughes et al., 2013), which has been a challenge in previous fMRI research on the relationship between reward-processing and delay discounting tendencies. As noted by Benningfield et al. (2014), for example, the fMRI MID task does not isolate motor-preparation processes from reward-anticipation processes. Moreover, recent advancements in EEG time-frequency analyses allow researchers to investigate neural processes in ways that may not be available in other techniques, such as examining changes in neural activation (power) at a specific time windows and frequency bands (Cohen, 2014; Makeig et al., 2004).

1.2. The reward time estimation task

To identify different stages of reward-processing, we adapted a feedback-learning task (called the reward time estimation task) (e.g., Damen and Brunia, 1987; Kotani et al., 2003; Luft, 2014; Pornpattananangkul and Nusslock, 2015). Here participants were asked to estimate a specific time duration by pressing a button 3.5 s after the onset of a Reward or No-Reward cue. The Reward/No-Reward cue indicated whether the current trial was a Reward- or No-Reward trial. Participants received monetary-reward for accurate-performance (pressing close to 3.5 s) on Reward trials, but not on No-Reward trials. Two seconds after making the button press, participants received feedback regarding the accuracy of their time-estimation. This paradigm allowed us to parse EEG related to reward-processing into reward-anticipation and reward-outcome phases (i.e., before and after feedback onset, respectively). Within reward-anticipation, we further separated EEG activity into 1) a cue-evaluation stage, involving a period immediately following the Reward/No-Reward cue, and 2) a feedback-anticipation stage, involving a period right before the feedback while participants were waiting to see if their recent action was considered accurate (i.e., close to 3.5 s). In addition to EEG indices, the improvement of time-estimation accuracy as the task proceeds can also be used as a behavioral index for the effectiveness in learning through feedback (Luft et al., 2013a). When the task provides rewards based on performance, then this behavioral index may reflect motivated-learning, or how motivated people are in learning to improve their performance in order to maximize reward earning (Luft, 2014). This motivated-learning behavioral index can be used 1) in corroborating EEG activity as indices of reward processing, and 2) in and of itself as an indirect, behavioral measure for individual-differences in reward-processing.
1.3. EEG indices at each stage of reward-processing

During cue-evaluation, participants evaluated whether their immediate future action could lead to a reward. In previous research, a cue signaling the possibility of receiving a reward was associated with a stronger P3 ERP component (compared to a cue in No-Reward trials) (Broyd et al., 2012; Goldstein et al., 2006; Ramsey and Finn, 1997; Santesso et al., 2012). More recently, Cavagnah (2015) found an enhancement of EEG power (or synchronization) in the delta band (≈1–3 Hz, called cue-locked delta) to reward-related cues. This cue-locked delta power found predominantly at parietal sites approximately 100–500 ms following cue onset is relevant to the P3 ERP component. Moreover, Cavagnah (2015) showed that enhanced cue-locked delta power to reward-related cues predicted behavioral adjustments in a re-inforcement learning task, suggesting an important role of cue-locked delta power in the cue evaluation stage.

As for feedback-anticipation, the suppression (or desynchronization) of alpha (≈8–13 Hz) EEG power at parieto-occipital sites prior to stimulus onset appears to index anticipation-related processes toward upcoming visual stimuli. For instance, using a time-estimation task, Bastiaansen et al. (1999, 2002) and Bastiaansen and Brumia (2001) provided either visual or auditory feedback regarding the accuracy of time-estimation to participants. The authors reported alpha power suppression immediately preceding both types of feedback, but this alpha suppression was strongly distributed to occipital sites for the visual, but not auditory, feedback. This suggests the role of pre-stimulus alpha suppression in modality-specific anticipation processes. Given that alpha power reflects the functional inhibition of neural activity (Jensen and Mazaheri, 2010), alpha suppression during the pre-stimulus anticipatory period likely reflects the dis-inhibition of neural activity in sensory cortices to facilitate attentional allocation to the upcoming stimulus. Additionally, this pre-stimulus suppression of parieto-occipital alpha to visual stimuli has been found to track the location of where people anticipate the stimuli to appear (Thut et al., 2006). Stronger suppression of pre-stimulus alpha power is also associated with how well people perceive the preceding near-threshold stimulus (Hanslmayr et al., 2007). Collectively, these findings suggest a relationship between alpha-suppression and anticipatory attention. More recently, stronger alpha suppression has been reported following monetary-reward cues and preceding monetary-reward feedback (Hughes et al., 2013). This additional suppression of alpha power by reward motivational cues suggests that alpha suppression indexes enhanced attentional processes during the anticipation of reward-related stimuli/feedback. Consistent with this idea, van den Berg et al. (2014) recently investigated the role of reward-related cues in a Stroop task, and demonstrated an inter-individual relationship. Specifically, individuals who had particularly strong alpha suppression following reward-related cues were more likely to have better behavioral performance on reward trials. Therefore, pre-feedback alpha suppression should serve as a reliable index for individual differences in reward-processing during the feedback-anticipation stage of reward-anticipation.

The reward-outcome period in our time-estimation task involved participants receiving feedback for that particular trial. We focused on two types of feedback evaluation: reward-evaluation and performance-evaluation. During reward-evaluation, individuals evaluate the motivational value of the feedback (Luft, 2014). That is, regardless of whether their performance outcome is good or bad, people should be more motivated to learn the outcome of their performance when this performance can lead to monetary reward (i.e., during Reward trials compared to No-Reward trials). During performance-evaluation, individuals assess whether their prior action was good or bad in meeting their performance goal, e.g., whether or not their time-estimation was accurate (Cavanagh and Shackman, 2015; Miltnner et al., 1997). The concept of performance evaluation has been related to both prediction error and conflict resolution. Unfortunately, many EEG studies have lumped these two aspects of feedback evaluation (i.e., reward-evaluation and performance evaluation) together, making it difficult to interpret the specific cognitive processes underlying their EEG findings. Here we separate reward-evaluation and performance evaluation. Furthermore, we examine three separate EEG indices during the reward-outcome stage, each occurring at distinct frequency bands: feedback-locked delta, feedback-locked theta and feedback-locked beta. As outlined next, we argue that each of these feedback-locked EEG profiles index individual differences in reward-processing during the reward-outcome stage. First, similar to cue-locked delta, recent studies have started to document changes in feedback-locked delta (1–3 Hz) power at parietal sites during reward-outcome approximately 100–500 ms following feedback onset (Cavanagh, 2015; Foti et al., 2015; Leicht et al., 2013). Cavanagh (2015) reported that feedback-locked delta was associated with prediction error in a reinforcement-learning task. Foti et al. (2015) reported that feedback-locked delta activity was stronger to feedbacks indicating monetary gains, compared to losses. Nonetheless, because monetary-gain feedback in these previous studies indicated both good performance and a positive reward outcome, it remains unclear whether enhancement in feedback-locked delta activity to gain feedback is driven by performance-evaluation or reward-evaluation. In the present study, we use the time estimation task to dissociate neural processes associated with reward and performance evaluation, and we examine their respective relationships with individual differences in delay-discounting.

Next, the enhancement of feedback-locked theta (≈4–7 Hz) power at frontal-midline sites (i.e., frontal-midline theta, FMT) approximately 200–400 ms following feedback onset has been implicated in feedback/outcome evaluation (Cohen et al., 2011). Thought to be generated from the anterior-cingulate cortex (Cavanagh and Frank, 2014), enhanced feedback-locked theta has been associated with cognitive-control processes that incorporate feedback/outcome information to facilitate behavioral adjustment on subsequent trials in order to maximize performance (van de Vijver et al., 2011). Feedback-locked theta appears sensitive to both performance-evaluation and reward-evaluation (Luft, 2014). In contrast to feedback-locked delta, feedback-locked theta is reliably stronger for bad-performance (compared to good-performance) feedback (Cohen et al., 2007; for a review see Luft (2014)). Additionally, such enhancement to bad-performance feedback predicts behavioral adjustment on a subsequent trial (Cavanagh and Shackman, 2015). As for reward-evaluation, Van den Berg et al. (2012) employed the time-estimation task and focused on the Feedback-Related Negativity (FRN), an event-related potential (ERP) thought to reflect the phase/time-locked feature of feedback-locked theta (Cavanagh et al., 2012). They found an enhanced FRN to feedback during Reward-trials where Good-performance led to monetary reward, compared to feedback during No-Reward trials where performance had no monetary consequences. Similar to the FRN, other studies have shown the influence of reward-evaluation on feedback-locked theta. For instance, feedback-locked theta is modulated by reward expectation (Cohen et al., 2007) and is stronger following feedback indicating a higher magnitude of monetary reward (Leicht et al., 2013). Thus, we expected feedback-locked theta to be modulated by both reward and performance evaluation.

Lastly, in addition to feedback-locked delta and theta, several recent studies have focused on feedback-locked EEG in the beta band (≈15–25 Hz) for review, see Luft (2014)). Similar to feedback-locked delta (but opposite to feedback-locked theta),
researchers have consistently found stronger beta power to positive feedback (e.g., monetary gains), compared to negative feedback (e.g., monetary losses) (Cohen et al., 2007; De Pascalis et al., 2012; Haji Hosseini et al., 2012; Marco-Pallares et al., 2008; Marco-Pallarés et al., 2009). Given that a similar pattern of enhanced beta power has been reported in the ventral striatum of animals during a reward-processing task (Berke, 2009; Courtemanche et al., 2003), it has been proposed that feedback-locked beta power presents reward-related signals from this region. Parietal feedback-locked beta power is reduced in humans following feedback during reward-learning tasks, such as the time estimation task (Luft et al., 2013a, 2013b; van de Vijver et al., 2011). This reduction in feedback-locked beta power (desynchronization) is less pronounced when feedback indicates good performance, compared to bad performance. Additionally, a greater enhancement of feedback-locked beta power (i.e., less reduction/desynchronization) predicts more rapid learning of a time duration in the time estimation task (Luft et al., 2013a). Nonetheless, similar to feedback-locked delta, most previous studies of feedback-locked beta have either lumped performance evaluation and reward evaluation feedback together (e.g., monetary gain indicating both good performance and reward associated with the performance) or focused solely on performance-evaluation, making it hard to interpret the psychological meaning of the effects. The current study separated the two aspects of feedback evaluation, and assessed their relationships with delay-discounting tendencies.

1.4. Current study

In the present study, participants completed a behavioral delay-discounting task and then a separate EEG reward time-estimation task. Drawing on existing research (Benningfield et al., 2014; Boettiger et al., 2007; Foerde et al., 2016), we predict that enhanced reward-related neural activity will be associated with a greater preference for larger-but-delayed rewards. An important contribution of this study is that we examined the relationship between reward-related neural activity and delay discounting tendencies at different temporal stages of reward-processing based on EEG time-frequency characteristics. Within the reward-anticipation stage, elevated reward-processing was operationalized as 1) greater cued-locked delta activity during cue-evaluation and 2) greater pre-feedback alpha suppression during feedback-anticipation. Within the reward-outcome stage, elevated reward-processing was operationalized as greater feedback-locked delta, theta and beta activity. To help corroborate these EEG variables as indices of reward processing, we examined the relationship between EEG-related data and behavioral learning performance during the reward time estimation task, which reflects motivated learning (Luft et al., 2013a). We also expect this behavioral index for motivated learning to correlate with individual differences in delay-discounting in a manner that is similar to reward-processing EEG indices.

2. Methods

2.1. Participants

Thirty-seven right-handed < 18, Chapman Handedness Scale; Chapman and Chapman (1987) native English speakers (21 females; age M = 19.05 years, SD = 1.22) at Northwestern University received partial course credit for their participation. Participants also earned additional monetary bonus based on their performance in the reward time estimation task (see below). Data from nine additional participants were discarded due to an EEG equipment problem (n = 1), excessive EEG slow-frequency (i.e., sweat artifact; n = 2) or high-frequency noise (i.e., muscle artifact; n = 1), giving the same answer throughout the Delay-Discounting task (n = 2), or a model-fit index on the Delay-Discounting task (R-square, see below) that was 1.5 inter-quartile ranges (IQRs) away from the nearer quartile (n = 3). Participants had no neurological history of head injury and were not taking psychotropic medications at the time of the study. Participants provided informed consent before the experiment. Northwestern Institutional Review Board approved the study.

2.2. Behavioral measure of delay-discounting responses: the delay-discounting task

Before EEG setup, participants completed a computer-adaptive version of the Delay-Discounting task (Fig. 1a) to assess individual differences in delay-discounting tendencies (Ahn et al., 2011; Rachlin et al., 1991). Given previous studies reporting a similar pattern of responses in delay-discounting between hypothetical and real monetary rewards (Johnson and Bickel, 2002; Lagorio and Madden, 2005), we used hypothetical rewards in this task to allow us to assess a wide range of delay periods and reward amounts.

For each trial in the delay-discounting task, participants were told to choose between a smaller-but-immediate (e.g., "$400 now") or larger-but-delayed (e.g., "$800 in one year") reward (see Fig. 1 for an example of a trial). There were six blocks of trials. Each block involved the same distribution of six different delay periods: two weeks, one month, six months, one year, three years or ten years. The order of blocks was fully randomized across participants. During the first trial of each block, participants made a choice between "$400 now" vs. "$800 at a given delay." A mean of the upper and lower bounds according to the choice made on the current trial were used as a smaller-but-immediate choice on the subsequent trial (Ahn et al., 2011; Du et al., 2002; Green and Myerson, 2004). The larger-but-delayed choice was always fixed at $800 regardless of the choice made on the previous trial. For example, if "$400 now" was chosen over "$800 in one year," then the smaller-but-immediate choice on the next trial would have been $200 (i.e., the mean of upper ($400) and lower ($0) bounds). This is because, based on the current-trial choice, "$800 in one year" had a lower subjective value than "$400 now", which means that the subjective value of "$800 in one year" was between "$0 now" and "$400 now". Thus, the subsequent trial would assess whether "$800 in one year" had a lower or higher subjective value than "$200 now." Conversely, if "$800 in one year" was chosen over $400, the next smaller-but-immediate choice would have been "$600 now" (i.e., the mean of upper ($800) and lower ($400) bounds). Following procedures conventionally used in studies focusing on individual-differences (Ahn et al., 2011; Du et al., 2002), we continued the adaptation of the smaller-but-immediate choice to the sixth trial of each block. Accordingly, the mean of the upper and lower bounds of the sixth trial was considered participants' subjective value of $800 at a given delay (Ahn et al., 2011; Du et al., 2002). The subjective values of $800 at every delay from all of the blocks were then fit into a hyperbolic model, which represented...
individual differences in delayed-discounting tendencies (Green and Myerson, 2004; Mazur, 1987). The steepness of the slope within this hyperbolic model (known as the discounting rate, or \( k \) value) reflects the extent to which people prefer smaller-but-immediate (compared to larger-but-delayed) rewards (see data analyses below).

2.3. EEG measure of reward-processing: the reward time-estimation task

The reward time-estimation task, adapted from a previous EEG paradigm (e.g., Damen and Brunia, 1987; Kotani et al., 2003), was used to assess individual differences in reward-related EEG activity. Participants were instructed to press a button with their right index finger 3.5 s after seeing a cue (see Fig. 2 for a schematic representation of the task). During Reward trials, participants earned 20 cents for accurate time estimations and received no monetary reward for inaccurate estimates. During No-Reward trials, participants received no monetary reward irrespective of their performance. Thus, the difference in EEG activity between Reward trials and No-Reward trials was considered an index of reward-related neural activity and reward-processing more generally. Each trial began with a Reward/No-Reward anticipation cue that signaled whether the trial was a Reward or No-Reward trial. This cue was presented for 300 ms and involved either circle or square shapes (counterbalanced across participants). The circle and square shapes were matched for luminance, contrast, and spatial frequency using Shine toolbox (Willenbockel et al., 2010) in Matlab.

We considered accurate responses as estimations that fell within the correct time-window. To control for variance in time-estimation ability among participants, two procedures were used to determine the correct time window. First, the correct time window on a given trial was shortened (or lengthened) by 20 ms if the response on the previous trial was (or was not) within the correct time-window (Kotani et al., 2003; Ohgami et al., 2006). This method has been found to generate an accuracy-rate of approximately 50%. Second, we included three Control blocks, each comprised of 36 trials, which were administered prior to the Experimental blocks. During these Control blocks, participants were instructed to button-press 3.5 s after seeing a triangle-shaped cue. There was no reward involved during these Control blocks, and participants were not informed about any of the reward contingencies in the time estimation task until after they finished these blocks. Moreover, unlike the Experimental blocks, participants received no performance feedback during the Control blocks, except if they made an extremely fast (a response faster than 2 s, indicated by a “< 2” sign) or slow (a response slower than 5 s, indicated by a “> 5” sign) response. These Control blocks allowed us to generate a correct response time window that was then calibrated to each individual participant and to measure participant’s initial time-estimation ability prior to them learning through reward/performance feedback in the Experimental blocks (see below). The initial time window for a correct response during the Control blocks was ± 500 ms, centered at 3500 ms. Following the Control blocks, participants completed 30 practice trials. These practice trials resembled trials during the Experimental blocks, except there were no earnings. The individualized time window for a correct response obtained from the Control blocks was used in the first practice trial.

The Experimental blocks consisted of six blocks of 36 trials. Within each block, there was a random distribution of Reward and No-Reward trials with a 50/50 split for each trial type. During the Experimental blocks, two lines of feedback text appeared two seconds following the button-press in the middle of the screen for 1000 ms (see Fig. 2). The top-line indicated performance feedback information, which was separated into three categories: Good-Performance, Bad-Performance and Extremely Fast/Slow. Good-Performance corresponded to an accurate response, or a response within the correct time-window, and was indicated by a “=” sign on the top line. Bad-Performance corresponded to a response slower than 2 s and faster than 5 s, but not within the correct time-window, and was indicated by a “< 3.5” (for a response between 2 s and the lower end of the correct time-window) or “> 3.5” (for a response between the higher end of the correct time-window and 5 s) sign on the top line. Extremely Fast/Slow feedback corresponded to a response faster than 2 s or slower than 5 s and was indicated by a “< 2” (for a response faster than 2 s) or “> 5” (for a response slower than 5 s) sign on the top line. The bottom line of the feedback text indicated whether or not participants won money for that particular response and included the following: “$” indicated the participant won money (20 cents) for that trial, and “0” indicated the participant did not win money for that trial. Thus, during Reward trials, participants would see “$” for Good-Performance, and see “0” for both Bad-Performance and Extremely Fast/Slow estimation. For No-Reward trials, participants would see “0” regardless of their performance. Trials were terminated with a randomly distributed ITI between 1000 and 1150 ms. To incentivize participants’ continued attention on both Reward and No-Reward trials, they were told they would receive no earnings if they saw feedback indicating extremely fast/slow
responses (i.e., “<2” or “>5”) more than 15 times. This ensured that they avoided extremely fast or slow responses.

2.4. Procedure

Following consent, participants completed the delay-discounting task. EEG electrodes were then applied and participants next completed the Time-Estimation Task. To help familiarize participants with the time duration of 3.5-s, participants first listened to two beep sounds 3.5-s apart as many times as they desired. Participants then completed the Control blocks for the Time-Estimation Task with no knowledge of the upcoming Experimental blocks. Participants were then given instructions regarding the Experimental blocks and corresponding Reward and No-Reward cues and the different kinds of feedbacks for the Time-Estimation Task. Participants were tested on their comprehension of these cues and feedbacks. Each block was separated by breaks of participant-determined length. During these breaks, participants were informed of their earnings and reminded of the meaning of the Reward/No-Reward Cues.

2.5. Electrophysiological recording

Continuous EEG data with a sampling rate at 500 Hz (DC to 100 Hz on-line, Neuroscan Inc.) were collected from inside an electro-magnetic shielded booth. Twenty-four Ag/AgCl scalp electrodes were used (F7/3/z/4/8, FC3/z/4, C3/z/4, T3/4, CP3/z/4, P3/z/4, T5/6, O1/z/2). HEOG and VEOG were recorded with four separate eye electrodes. Recordings were referenced on-line to a left mastoid and re-referenced of linked mastoids. Impedance was kept below 5 kΩ and 10 kΩ for scalp and eye electrodes, respectively. During offline analyses, eye movement artifacts were first corrected with PCA algorithms implemented in NeuroScan EDIT (Neuroscan Inc.). Movement-related artifacts were removed manually. EEG data were offline high-pass-filtered at .01 Hz.

2.6. Data analyses for the delay-discounting task

Each participants’ subjective value of $800 at every delay were fit to the hyperbolic model in the form of $V = A/(1 + kD)$ (Green and Myerson, 2004; Mazur, 1987) using the Curve Fitting Toolbox in Matlab. $V$ was the subjective value, $A$ was a larger-but-delayed reward amount ($800), D$ was the delay interval in weeks, and $k$ was a free parameter, reflecting a discounting rate. The smaller the $k$, the stronger preference toward larger-but-delayed rewards. To normalize its distribution, the natural log of $k$, ln($k$), was used as an index for individual-differences in delay-discounting responses. R-square as a model-fit index was also calculated. R-square indicates the proportion of variance accounted for by the hyperbolic model, ranging from 0 to 1 (perfect fit).

2.7. Data analyses for the reward time-estimation task

2.7.1. Manipulation check

We computed an Inaccurate Estimation index as the standard deviation of the absolute difference between participants’ actual estimations (in milliseconds, ms) and the target time interval (3500 ms) (Luft et al., 2013a). Thus, higher Inaccurate Estimation reflects worse time estimation performance. To normalize the Inaccurate Estimation distribution, a natural-log (ln) transformation was applied. As a manipulation check for whether participants were more motivated during Reward trials, we compared Inaccurate Estimation during 1) Reward trials from all of the Experimental blocks, 2) No-Reward trials from all of the Experimental blocks and 3) Control trials from all of the Control blocks. We predicted lower Inaccurate Estimation during Reward (compared to No-Reward and Control) trials.

2.7.2. Behavioral indices for individual-differences in motivated learning

In addition to being used as a manipulation check, Inaccurate Estimation was also employed as a behavioral index for individual-differences in motivated learning. Specifically, Inaccurate
Estimation during the Control blocks (referred to as Control Inaccurate Estimation) served as an index for individual differences in estimation ability prior to learning in the Experimental blocks. Inaccurate Estimation during the last (i.e., 6th) Experimental block (referred to as Motivated-Learning Inaccurate Estimation) served as an index for how motivated participants were to learn through feedback over the course of the experiment (for a similar approach, see Luft et al. (2013a)). For individual differences in Motivated-Learning Inaccurate Estimation, we collapsed across Reward and No-Reward trials during the last experiment block. This is because, in the context of the current time-estimation task, learning about their estimation performance through feedbacks during the No-Reward trials also helped participants to earn more rewards during the Reward trials. Thus, toward the end of the experiment (i.e., 6th block), people who were more motivated to obtain rewards in the task would perform better in both Reward and No-Reward trials. Additionally, combining both Reward and No-Reward trials during the last experiment block led to more trials to be computed, thus increasing stability of Motivated-Learning Inaccurate Estimation as a measure for individual differences. This is important, given that we only had 36 trials (Reward and No-Reward trials combined) in each block. We then examined whether both In (k) and EEG indices have relationship with Control Inaccurate Estimation and Motivated-Learning Inaccurate Estimation.

2.7.3. EEG data
Extremely fast (less than 2 s, “<2”) or slow (more than 5 s, “>5”) trials were excluded from the EEG analyses as they may reflect a lack of attention to the task. EEGlab (Delorme and Makeig, 2004) was used to analyze EEG data. After a linear detrend was applied to EEG epochs, any epoch containing artifacts (±75 μV) were rejected. Time-frequency decomposition was then performed to compute event-related spectral perturbation (ERSP). ERS refers to changes in EEG power from the baseline period at specific frequency and time (Makeig et al., 2004). We used a linear space for both frequency (at every 1 Hz) and time (at every 2 ms).

We separated EEG activity into three phases of reward-processing: 1) cue-locked EEG corresponding to cue-evaluation during the reward-anticipation stage (i.e., cue evaluation during Reward vs. No-Reward trials), 2) pre-feedback EEG corresponding to feedback-anticipation during the reward-anticipation stage (i.e., anticipation while waiting for feedback outcome during Reward vs. No-Reward trials), and 3) feedback-locked EEG corresponding to the reward-outcome stage (i.e., evaluation of feedback that revealed Reward and Performance information). For cue-locked and feedback-locked EEG activity, we focused on ERSP power following the onset of the cue and feedback, respectively. Here, EEG data were epoched from −2500 to 3500 ms relative to cue/feedback onset. This relatively long epoch allowed us to investigate ERSPs from a low (1 Hz) to a relatively high (50 Hz) frequency, which covered EEG frequency bands of interest during these time periods (delta, theta and beta). A modified complex sinusoidal wavelet was used with a sliding window of 3342-ms wide (leading to 3 cycles at 1 Hz) and a wavelet factor of .92. To reduce the potential edge effects, we removed half of the sliding window (1671 ms) at the edges (i.e., the beginning and the end) of each epoch, leaving an epoch between −829 and 1927 ms after edge removal. The baseline for cue- and feedback-locked EEG was between −300 to −100 ms before the stimulus onset.

For pre-feedback EEG activity, we focused on ERSP power prior to feedback onset. Specifically, pre-feedback EEG activity was epoched from −2557 to 2557 ms relative to the button-press. Given that our focus during this period was specifically on the Alpha band (8–13 Hz), we computed pre-feedback ERP at a narrower frequency range of 3–30 Hz. Because the lowest frequency (3 Hz) for pre-feedback EEG was higher than it was in the cue-/feedback-locked EEG (1 Hz), we used a shorter sliding window of 1114-ms wide (leading to 3 cycles at 3 Hz) and a wavelet factor of .5. The removal of epoch edges (557 ms or half of a sliding window) shortened the epoch to be between −2000 and 2000 ms relative to each button-press, which ended at the onset of the feedback. EEG power during this period from −2000 to −1500 ms before the button-press was used as the baseline. This early baseline period was selected in order to avoid movement-related activity near the time of the button-press.

Two-tailed paired parametric tests were performed on ERSP time-frequency and topographic maps, as implemented in EEGLab. Multiple comparisons were corrected using False Discovery Rate (FDR) (Benjamini and Yekutieli, 2001). P-values based on these statistical tests were shown on time-frequency and topographic maps. These maps of p values indicate times, frequencies and electrodes at which there was a significant effect. For cue-locked and pre-feedback EEG activity, the statistical tests were used to assess whether there was an effect of Reward conditions (Reward-trial vs. No-Reward-trial) on ERSP power. For feedback-locked EEG activity, we separated feedbacks based on evaluation types: Reward Evaluation (Reward-trial vs. No-Reward-trial) x Performance Evaluation (Bad-Performance vs. Good-Performance). The main effects and interaction between evaluation types were tested on ERSP power.

2.7.4. Reward-processing ERSPs as indices for individual differences
We focused individual differences analyses on ERSPs that significantly varied between Reward and No-Reward trials according to main-effect analyses (see Section 3). Specifically, for cue-locked EEG, individual-differences analyses were centered at cue-locked delta (1–3 Hz) between 100 and 500 ms following the cue onset at the parietal CPz site, similar to a recent study on reward-processing to the cue (Cavanagh, 2015). For pre-feedback EEG, we used alpha (8–13 Hz) power within the 500-ms window right before feedback onset at Oz. Occipital pre-feedback alpha suppression (i.e., lower power) is related to anticipation processes toward upcoming visual stimuli (Bastiaansen et al., 1999; Hughes et al., 2013). Thus, the more negative the alpha-suppression, the stronger the anticipation-related processes. For feedback-locked EEG, we focused on three different ERSPs from three different frequency bands: feedback-locked delta, theta and beta. Similar to cue-locked delta, feedback-locked delta (1–3 Hz) was quantified as averaged EEG power between 100 and 500 ms following the feedback onset at CPz. This is similar to previous research on feedback-locked delta power (Foti et al., 2015). As for feedback-locked theta (4–7 Hz), we employed averaged EEG power between 200 and 400 ms after feedback onset at Fz, following previous research on frontal-midline theta and outcome/feedback-processing (Cohen et al., 2007; Marco-Pallares et al., 2008). Lastly, feedback-locked beta (15–25 Hz) was quantified through averaged EEG power between 400 and 600 ms after feedback onset at CPz following a recent time-estimation study (Luft et al., 2013a).

In addition to examining ERSPs at separate conditions (i.e., Reward-trial and No-Reward-trial ERSPs for cue-locked and pre-feedback EEG; Reward-trial, No-Reward-trial, Good-Performance and Bad-Performance ERSPs for feedback-locked EEG), we also computed ERSP difference scores by subtracting ERSP power during No-Reward trials from that during Reward trials (collapsing across Bad- and Good-Performance feedback for feedback-locked EEG). Referred to as ΔReward ERSPs, these difference scores allowed us to focus in on reward-processing EEG activity. For feedback-locked EEG, we also computed ΔPerformance ERSP difference scores by subtracting ERSP power during Good-Performance trials from that during Bad-Performance trials, collapsing across Reward- and No-Reward trials. The ΔPerformance ERSPs reflect individual differences in performance evaluation.
3. Results

3.1. Delay-discounting responses

Fig. 1b shows how subjective values of rewards were discounted as a function of delays. The mean of ln (k) was −4.72 (SD=1.45). R-square, as a model-fit index for the hyperbolic model used, had the median of .89 (IQR=.94−.81), similar to previous studies (de Wit et al., 2007; Hariri et al., 2006).

3.2. Reward time-estimation behavior

3.2.1. Manipulation check

During the time-estimation task, behavioral data from one participant was lost due to a technical error, leaving data from 36 (as opposed to 37) participants. Overall, during the Experimental blocks, participants estimated 3.5 s quite well (M_{RT}=3.52 s, SD=.11), and rarely made extremely fast (less than 2 s) or slow (more than 5 s) responses (M=2.58 trials out of 216 trials, SD=2.15). Given that the accuracy rate (M=50.3%, SD=3.06) closely matched 50%, it is clear that the time-adaptation algorithm worked effectively. Across blocks, there were differences in Inaccurate Estimation (i.e., SD of the absolute difference between actual estimations and the target time interval) among Reward and No-Reward trials during all of the Experimental blocks, F(1,14, 39,90)=84.33, p < .001, Greenhouse-Geisser corrected. As expected, Inaccurate Estimation was lower during Reward trials (M=5.44 ln (ms), SD=.38) compared to during No-Reward trials (M=5.54 ln (ms), SD=.39, p < .001), which was lower than during Control trials (M=6.22 ln (ms), SD=.33, p < .001).

3.2.2. Relationship between motivated learning inaccurate estimation and delay-discounting responses

Motivated-Learning Inaccurate Estimation (i.e., collapsing across all Reward and No-Reward trials during the last Experimental block) was positively correlated with ln (k), while Control Inaccurate Estimation (i.e., during the Control blocks) was not (see Table 1 and Fig. 3). Thus, participants with poorer estimation performance after learning throughout the task (i.e., higher Motivated-Learning Inaccurate Estimation) had higher k values (i.e., stronger preference toward small-but-immediate rewards). In other words, people who were more motivated to learn through feedbacks throughout the task were more likely to prefer larger-but-delayed rewards.

3.3. Cue-locked EEG activity

Fig. 4a and b show ERSP time-frequency and topographic maps of cue-locked EEG. Overall, following cue onset, there was an increase in ERSP power at the delta (1–3 Hz) and theta (4–7 Hz) bands and a decrease in ERSP power in the alpha/beta (8–25 Hz) band. More importantly, cue-locked ERSP power at the delta (but not theta or beta) band was significantly stronger for Reward trials.

### Table 1

<table>
<thead>
<tr>
<th>ERSPs and behavioral indices</th>
<th>Motivated-Learning Inaccurate Estimation</th>
<th>Control Inaccurate Estimation</th>
<th>Delay-discounting tendencies ln (k)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cue-locked</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reward-trial delta</td>
<td>−.377*</td>
<td>.047</td>
<td>−.076</td>
</tr>
<tr>
<td>No-reward-trial delta</td>
<td>−.069</td>
<td>.125</td>
<td>−.143</td>
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<tr>
<td>ΔReward delta</td>
<td>−.359*</td>
<td>−.090</td>
<td>.080</td>
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<tr>
<td>Pre-feedback</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Reward-trial alpha suppression</td>
<td>.375*</td>
<td>.209</td>
<td>.441*</td>
</tr>
<tr>
<td>No-reward-trial alpha suppression</td>
<td>.327*</td>
<td>.237</td>
<td>.254</td>
</tr>
<tr>
<td>ΔReward alpha suppression</td>
<td>.196</td>
<td>.035</td>
<td>.400*</td>
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<tr>
<td>Feedback-locked Reward-evaluation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reward-trial delta</td>
<td>−.507**</td>
<td>−.185</td>
<td>−.264</td>
</tr>
<tr>
<td>No-reward-trial delta</td>
<td>−.321†</td>
<td>−.178</td>
<td>−.194</td>
</tr>
<tr>
<td>ΔReward delta</td>
<td>−.466*</td>
<td>−.032</td>
<td>−.191</td>
</tr>
<tr>
<td>Reward-trial theta</td>
<td>−.536*</td>
<td>−.117</td>
<td>−.407†</td>
</tr>
<tr>
<td>No-reward-trial theta</td>
<td>−.302†</td>
<td>−.131</td>
<td>−.293†</td>
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<tr>
<td>ΔReward theta</td>
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<td>−.043</td>
<td>−.344*</td>
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<tr>
<td>Reward-trial beta</td>
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<td>−.427**</td>
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<tr>
<td>No-reward-trial beta</td>
<td>−.302†</td>
<td>.061</td>
<td>−.361*</td>
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<tr>
<td>ΔReward beta</td>
<td>−.237</td>
<td>.038</td>
<td>−.246*</td>
</tr>
<tr>
<td>Feedback-locked Performance-evaluation</td>
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<tr>
<td>Bad-performance delta</td>
<td>−.480**</td>
<td>−.205</td>
<td>−.196</td>
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<tr>
<td>Good-performance delta</td>
<td>−.338*</td>
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<tr>
<td>ΔPerformance delta</td>
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<td>.079</td>
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<tr>
<td>Bad-performance theta</td>
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<td>−.15</td>
<td>−.354†</td>
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<tr>
<td>Good-performance theta</td>
<td>−.454*</td>
<td>−.076</td>
<td>−.400*</td>
</tr>
<tr>
<td>ΔPerformance theta</td>
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<tr>
<td>Bad-performance beta</td>
<td>−.254</td>
<td>.149</td>
<td>−.303†</td>
</tr>
<tr>
<td>Good-performance beta</td>
<td>−.365*</td>
<td>−.013</td>
<td>−.436**</td>
</tr>
</tbody>
</table>

Note. For feedback-locked Reward-evaluation ERSPs, ERSPs were collapsing across Bad-Performance and Good-Performance feedbacks. ΔReward ERSPs were calculated by subtracting No-reward-trial ERSPs from Reward-trial ERSPs. Similarly, for feedback-locked Performance-Evaluation ERSPs, ERSPs were collapsing across Reward- and No-Reward-Trial feedbacks. ΔPerformance ERSPs were calculated by subtracting Good-Performance ERSPs from Bad-Performance ERSPs. More negative values of ΔReward Alpha-Suppression reflect the greater suppression of alpha activity during Reward trials (relative to No-Reward trials). Inaccurate Estimation refers to the deviation from the goal of estimating time (3.5 s), and therefore high values reflect worse performance. Motivated-Learning Inaccurate Estimation and Control Inaccurate Estimation refer to Inaccurate Estimation indices during the last block of the experimental session, and during the control blocks (i.e., before the learning occurred), respectively. *p < .01, †p < .05, ‡p < .10 (2-tailed). *Became significant (r(35)=.36, p=.03), after a bivariate outlier was removed (see Fig. 12).
suppression during anticipation-related activity reported in previous studies (Bastiaansen et al., 1999; Hughes et al., 2013; van den Berg et al., 2014). As predicted, alpha suppression was significantly stronger (i.e., higher reduction in alpha power) for Reward trials than for No-Reward trials starting approximately 500 ms before feedback onset. This effect was primarily significant at parieto-occipital sites (for \( p \) values, FDR corrected, see Fig. 5a). Thus, we employed pre-feedback alpha power during the 500-ms window prior to feedback onset at Oz as an EEG index for individual differences in feedback anticipation during the reward-anticipation stage.

Motivated-Learning Inaccurate Estimation had a significant, positive correlation with pre-feedback alpha during Reward trials, and a marginally significant, positive correlation with this ERSP during No-Reward trials (see Table 1 and Fig. 5c). Nonetheless, Motivated-Learning Inaccurate Estimation was not correlated with the \( \Delta \)Reward (Reward minus No-Reward-trial) pre-feedback alpha (see Table 1). Additionally, Control Inaccurate Estimation was not correlated with any pre-feedback alpha indices (see Table 1). This suggests that enhanced alpha suppression (i.e., lower alpha power) prior to the feedback onset was related to better adjustment of time-estimation performance (i.e., lower Inaccurate Estimation) throughout the task. However, the relationship between alpha suppression and time-estimation performance may not be limited to Reward trials.

With respect to delay-discounting tendencies, pre-feedback alpha power during Reward trials (but not during No-Reward trials) had a positive relationship with \( \ln (k) \) (see Table 1). Moreover, there was a significant positive relationship between \( \Delta \)Reward pre-feedback alpha and \( \ln (k) \) (see Table 1 and Fig. 10). These correlations suggest that participants who had stronger alpha suppression (i.e., lower power) during Reward (compared to No-Reward) trials (i.e., more negative \( \Delta \)Reward pre-feedback alpha) had smaller \( k \) values. In other words, individuals who had stronger feedback anticipation during Reward trials had a stronger preference toward larger-but-delayed, compared to smaller-but-immediate, rewards.

3.5. Feedback-locked EEG activity

ERP time-frequency and topographic maps following feedback onset are shown in Figs. 6–8. Overall, following feedback onset, EEG power was increased at the delta (1–3 Hz) and theta (4–7 Hz) bands and decreased at the alpha/beta (8–25 Hz) band. Moreover, the effect of Reward-Evaluation (Reward-trial vs. No-Reward-trial feedbacks) on feedback-locked EEG power was found at delta, theta and beta bands (for \( p \) values, FDR corrected, see Figs. 6b–8b).

3.5.1. Feedback-locked delta

Similar to cue-locked delta, feedback-locked delta (1–3 Hz) was enhanced approximately 100–500 ms following feedback onset at parietal sites. There were main effects of both Reward Evaluation and Performance Evaluation on feedback-locked delta (see Fig. 6a and c for ERSP power and Fig. 6b for \( p \) values, FDR corrected). Specifically, for Reward Evaluation, feedback-locked delta was stronger for Reward-trial, than No-Reward-trial, feedback at CPz and other parietal sites. For Feedback Evaluation, feedback-locked delta was significantly stronger for Good-Performance, than Bad-Performance, feedback at parietal sites. There was also a significant interaction. Yet, while the effect of Reward Evaluation (i.e., stronger feedback-locked delta during Reward trials than during No-Reward trials) was numerically larger for Good-Performance (than for Bad-Performance) feedback, the Reward Evaluation effect was significant for both Good-Performance and Bad-Performance feedback. Thus, we collapsed across Good-Performance and Bad-Performance feedback when employing the feedback-locked delta

3.4. Pre-feedback EEG activity

Fig. 5a and b show ERSP time-frequency and topographic maps during a period prior to feedback onset. There was a clear pattern of alpha (8–13 Hz) suppression (i.e., reduction in ERSP power) at parieto-occipital sites, peaking at the period right before feedback onset. This EEG pattern is similar to the profile of occipital alpha
as an index for individual differences in Reward Evaluation of the feedback.

Motivated-Learning Inaccurate Estimation had a significant, negative correlation with feedback-locked delta during Reward trials, and a marginally significant, positive correlation with this ERSP during No-Reward trials (see Table 1 and Fig. 6d). Similar to \( \Delta \)Reward cue-locked delta, \( \Delta \)Reward (Reward- minus No-Reward-trial) feedback-locked delta was also negatively correlated with Motivated-Learning Inaccurate Estimation (see Table 1 and Fig. 9b). Motivated-Learning Inaccurate Estimation, however, had no relationship with \( \Delta \)Performance (Bad minus Good-Performance) feedback-locked delta. Similarly, Control Inaccurate Estimation was not correlated with any feedback-locked delta indices (see Table 1). Altogether, this suggests that enhanced feedback-locked delta power to Reward-trial (compared to the No-Reward-trial) feedback was related to a better behavioral adjustment of time-estimation performance (reflected by smaller Motivated-Learning Inaccurate Estimation). Nonetheless, similar to cue-locked delta indices, feedback-locked delta indices did not significantly predict \( \ln(k) \) (see Table 1).
3.5.2. Feedback-locked theta
ERSP power in the theta frequency (4–7 Hz) was enhanced during the 200–400 ms window following feedback onset at frontal-midline sites. This EEG pattern is similar to the profile of frontal-midline theta (FMT) during outcome-processing reported in previous studies (Cohen et al., 2007; Luft, 2014). As expected, there was a main effect of both Reward Evaluation and Performance Evaluation (see Fig. 7a and c for ERSP power and Fig. 7b for p values, FDR corrected). At Fz and other frontal-midline sites, feedback-locked theta was significantly stronger for Reward-trial, than No-Reward-trial, feedback. In contrast to feedback-locked delta, feedback-locked theta was significantly stronger for Bad-Performance, than Good-Performance, feedback at Fz and other frontal-midline sites. There was no interaction between Reward Evaluation and Performance Evaluation on feedback-locked theta at any frontal-midline sites (for p values, FDR corrected, see Fig. 7b).

Similar to feedback-locked delta, Motivated-Learning Inaccurate Estimation was negatively correlated with feedback-locked theta during Reward trials, and there was a marginally

---

### Table 1

<table>
<thead>
<tr>
<th>Reward Trial ERSP (dB)</th>
<th>No-Reward Trial ERSP (dB)</th>
<th>P-values, FDR corrected</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1.png" alt="Image" /></td>
<td><img src="image2.png" alt="Image" /></td>
<td><img src="image3.png" alt="Image" /></td>
</tr>
</tbody>
</table>

0 ms = button press  
2000 ms = feedback onset

---

**Fig. 5.** Pre-feedback occipital alpha suppression. a. Shows event-related spectral perturbation (ERSP) time-frequency maps at the electrode Oz (Top) and topographic maps (the grand-average of alpha (8–13 Hz) ERSP power at the 500-ms window prior to feedback onset) (Bottom) depicting pre-feedback reward-anticipation EEG activity. Specifically, ERSP activity during Reward and No-Reward trials is depicted in the left and middle columns, respectively. The right column depicts paired t-tests’ (df = 36) p-values (FDR corrected) for which Reward trial ERSP power was significantly different from No-Reward trial ERSP power. For these p-value maps, the time-frequency map shows p values at each time and frequency at Oz, while the bottom topo map shows p values at the alpha (8–13 Hz) band at the 500-ms window prior to feedback onset at every electrode. The time scale in the time-frequency maps is relative to button-presses: 0 ms is when a button was pressed, and 2000 ms is when the feedback was presented. Note that we computed time-frequency maps from −2000 to 2000 ms (see text). However, for visual purposes, we only show 0–2000 ms here, given our focus on the time period right before feedback onset. Note also that heat-map scales for time-frequency and topographic maps are different. b. Depicts ERSP at the alpha band (8–13 Hz) at Oz. c. Shows a scatterplot between Motivated-Learning Inaccurate Estimation (i.e., during the last experimental block) and pre-feedback alpha ERSP (8–13 Hz at the 500-ms window pre-feedback) at Oz. In this scatterplot, pre-feedback alpha ERSP is separated to the ERSP during Reward Trials and No-Reward Trials. Gray areas indicate the confident interval of 95% around the regression line.
Fig. 6. The influence of reward-evaluation and performance-evaluation on parietal delta ERSP following feedback onset. a. Shows ERSP time-frequency maps at the electrode CPz and topographic maps (the grand-average of delta ERSP power at 1–3 Hz, 100–500 ms window following feedback onset). ERSP activity locked to feedback onset is separated into reward evaluation (Reward-Trial vs. No-Reward Trial feedback) and performance evaluation (Bad-Performance vs. Good-Performance feedback). Note that we computed time-frequency maps from 830 to 1828 ms (see text). However, for visual proposes, we only show 300 to 1000 ms relative to the feedback onset here. Note also that heat-map scales for time-frequency and topographic maps are different. b depicts p-values (FDR corrected) showing the main effect of Reward Evaluation (Reward Trial vs. No-Reward Trial), main effect of Performance Evaluation (Bad- vs. Good-Performance) and their interaction. For these p-value maps, the time-frequency map shows p values at each time and frequency at CPz, while the topographic map shows p values at the delta (1–3 Hz) band at the 100–500 ms window following feedback onset at every electrode. c. Shows grand-average of delta power (1–3 Hz) at CPz during 100–500 ms window following feedback onset, separated by conditions. Error bars represent standard error of the mean, corrected for repeated-measure comparison. d. Shows a scatterplot between Motivated-Learning Inaccurate Estimation (i.e., during the last Experimental block) and feedback-locked delta ERSP (1–3 Hz, 100–500 ms window following feedback onset). In this scatterplot feedback-locked delta ERSP was separated to the ERSP during the Reward trials and No-Reward trials, collapsing across Bad- and Good-Performance. Gray areas indicate the confident interval of 95% around the regression line.
Fig. 7. The influence of reward-evaluation and performance-evaluation on frontal-midline theta ERSP following feedback onset. a. Shows ERSP time-frequency maps at the electrode Fz and topographic maps (the grand-average of theta ERSP power at 4–7 Hz, 200–400 ms window following feedback onset). ERSP activity locked to feedback onset is separated into reward evaluation (Reward-trial vs. No-Reward-trial feedback) and performance evaluation (Bad-Performance vs. Good-Performance feedbacks). Note that we computed time-frequency maps from −830 to 1828 ms (see text). However, for visual proposes, we only show −300 to 1000 ms relative to the feedback onset here. Note also that heat-map scales for time-frequency and topographic maps are different. b. Depicts p-values (FDR corrected) showing the main effect of Reward Evaluation (Reward Trial vs. No-Reward Trial), main effect of Performance Evaluation (Bad- vs. Good-Performance) and their interaction. For these p-value maps, the time-frequency map shows p values at each time and frequency at Fz, while the topographic map shows p values at the theta (4–8 Hz) band at the 200–400 ms window following feedback onset at every electrode. c. Shows grand-average of theta power (4–7 Hz) at CPz during 200–400 ms window following feedback onset, separated by conditions. Error bars represent standard error of the mean, corrected for repeated-measure comparison. d. Shows a scatterplot between Motivated-Learning Inaccurate Estimation (i.e., during the last Experimental block) and feedback-locked theta ERSP (4–7 Hz, 200–400 ms window following feedback onset). In this scatterplot feedback-locked theta ERSP was separated to the ERSP during the Reward Trials and No-Reward Trials, collapsing across Bad- and Good-Performance. Gray areas indicate the confident interval of 95% around the regression line.
Fig. 8. The influence of reward-evaluation and performance-evaluation on parietal beta ERSP following feedback. a. Shows ERSP time-frequency maps at the electrode CPz and topographic maps (the grand-average of beta ERSP power at 15–25 Hz, 400–600 ms window following feedback onset). ERSP activity locked to feedback onset is separated into reward evaluation (Reward-trial vs. No-Reward-trial feedback) and performance evaluation (Bad-Performance vs. Good-Performance feedbacks). Note that we computed time-frequency maps from -830 to 1828 ms (see text). However, for visual proposes, we only show 300–700 ms relative to the feedback onset here. This is to highlight beta activity (15–25 Hz) between 400 and 600 ms. To further highlight ERSP power at the beta band, we also constrains the heat-map scales for the time-frequency maps to -2 to 2 dB. Note also that heat-map scales for the time-frequency and topographic maps are different. b. Depicts p-values, FDR corrected, showing the main effect of Reward Evaluation (Reward Trial vs. No-Reward Trial), main effect of Performance Evaluation (Bad- vs. Good-Performance) and their interaction. For these p-value maps, the time-frequency map shows p values at each time and frequency at CPz, while the topographic map shows p values at the beta (15–25 Hz) band at the 400–600 ms window following feedback onset at every electrode. c. Shows grand-average of beta power (15–25 Hz) at CPz during 400–600 ms window following feedback onset, separated by conditions. Error bars represent standard error of the mean, corrected for repeated-measure comparison. d. Shows a scatterplot between Motivated-Learning Inaccurate Estimation (i.e., during the last Experimental block) and feedback-locked theta ERSP (15–25 Hz) during 400–600 ms window following feedback onset at CPz. The feedback-locked theta ERSP is separated into the ERSP during the Reward Trials and No-Reward Trials, collapsing across Bad- and Good-Performance. Gray areas indicate the confident interval of 95% around the regression line.
significant negative correlation between Motivated-Learning Inaccurate Estimation and feedback-locked theta during No-Reward trials (see Table 1 and Fig. 7d). Similar to ΔReward cue-locked and feedback-locked delta, ΔReward feedback-locked (Reward minus No-Reward-trial) theta was also negatively correlated with Motivated-Learning Inaccurate Estimation (see Table 1 and Fig. 9c). There was, however, no relationship between ΔPerformance (Bad minus Good-Performance) feedback-locked theta and Motivated-Learning Inaccurate Estimation. Control Inaccurate Estimation was not predicted by any of the feedback-locked theta indices (see Table 1). Together, this suggests that enhanced feedback-locked theta power to Reward-trial (compared to the No-Reward-trial) feedback was associated with a better adjustment of time-estimation performance (reflected by smaller Motivated-Learning Inaccurate Estimation).

In contrast to cue-locked and feedback-locked delta, but similar to pre-feedback alpha, feedback-locked theta was related to delay-discounting tendencies. Specifically, ln (k) was negatively correlated with feedback-locked theta during Reward trials, and there was a marginally significant negative correlation between ln (k) and feedback-locked theta during No-Reward trials (Table 1). Moreover, while the relationship between the ΔPerformance feedback-locked theta and ln (k) was not significant, there was a significant negative relationship between the ΔReward feedback-locked theta and ln (k) (Table 1). Thus, participants with stronger feedback-locked theta power following Reward-trial, compared to No-Reward-trial, feedback (i.e., more positive ΔReward feedback-locked theta) had smaller k values (i.e., stronger preference toward larger-but-delayed rewards). To investigate the specificity of this relationship, we examined correlations between ln (k) and the ΔReward feedback-locked theta following Bad-Performance feedback separately from ΔReward feedback-locked theta following Good-Performance feedback. ln (k) was negatively correlated with the ΔReward feedback-locked theta following Good-Performance feedback, r(35) = .44, p = .006 (Fig. 11), but not with the ΔReward feedback-locked theta following Bad-Performance feedback, r(35) = .14, p = .39.

3.5.3. Feedback-locked beta

Feedback-locked beta power (15–25 Hz) was reduced starting approximately 200 ms after feedback onset at parietal sites (see Fig. 8a and c for ERSP power). This is similar to a pattern of beta

Fig. 9. Scatterplots of the correlation between Motivated-Learning Inaccurate Estimation (i.e., during the last experimental block) and ΔReward ERSPs (Reward-trial ERSPs minus No-Reward-trial ERSPs). Note that higher Motivated-Learning Inaccurate Estimation values reflect worse time-estimation performance during the last Experimental block. Potentials bivariate outliers were detected using Minimum Covariance Determinant (MCD), and represented by red data points. Ellipses contain non-outlying data. Pink shaded areas represent 95% bootstrapped confidence intervals around the linear regression line after the potential bivariate outliers were removed. Importantly, all of the correlations presented below remain significant, with or without bivariate outlier removal, suggesting the robustness of the relationship. a. Represents the correlation between Motivated-Learning Inaccurate Estimation and ΔReward cue-locked delta (Reward-trial cue-locked delta minus No-Reward-trial cue-locked delta). The correlation coefficient was significant both before (r(34) = 0.36, p = 0.03) and after (r(32) = 0.39, p = 0.02) bivariate outliers removal. b. Represents the correlation between Motivated-Learning Inaccurate Estimation and ΔReward feedback-locked delta (Reward-trial cue-locked delta minus No-Reward-trial cue-locked delta, collapsing across Bad- and Good-Performance feedback). The correlation coefficient was significant both before (r(34) = 0.47, p = 0.004) and after (r(31) = 0.65, p < .001) bivariate outliers removal. c. Represents the correlation between Motivated-Learning Inaccurate Estimation and ΔReward feedback-locked theta (Reward-trial cue-locked theta minus No-Reward-trial cue-locked theta, collapsing across Bad- and Good-Performance feedbacks). The correlation coefficient was significant both before (r(34) = 0.56, p < .001) and after (r(33) = 0.45, p = 0.007) bivariate outliers removal. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
desynchronization found in a recent time-estimation study (Luft et al., 2013a). Significant main effects of both Reward Evaluation and Performance Evaluation on feedback-locked beta were found at approximately 400–600 ms following feedback onset (see Fig. 8b for p values, FDR corrected). Specifically, for Reward Evaluation, feedback-locked beta was less reduced during Reward trials than No-Reward trials. Moreover, for Feedback Evaluation, feedback-locked beta was less reduced following Good-Performance feedback, compared to following Bad-Performance feedback. There was a trend for the Reward Evaluation by Performance Evaluation interaction, but this interaction did not survive FDR correction. Thus, we collapsed across Good-Performance and Bad-Performance feedback when employing feedback-locked beta as an index for individual differences in Reward Evaluation at this frequency band.

Similar to feedback-locked delta and theta, Motivated-Learning Inaccurate Estimation was negatively correlated with feedback-locked beta during Reward trials, and was marginally correlated with feedback-locked beta during No-Reward trials (see Table 1 and Fig. 8d). Nonetheless, neither ΔReward feedback-locked (Reward minus No-Reward-trial) beta or ΔPerformance feedback-locked (Bad minus Good-Performance) beta was correlated with Motivated-Learning Inaccurate Estimation (see Table 1). Additionally, Control Inaccurate Estimation was not predicted by any feedback-locked beta indices (see Table 1). Similar to pre-feedback alpha, this suggests that a smaller reduction in feedback-locked beta was related to better adjustment of time-estimation performance (i.e., smaller Motivated-Learning Inaccurate Estimation). However, the relationship between feedback-locked beta and Motivated-Learning Inaccurate Estimation does not appear to be unique to Reward trials.

Similar to pre-feedback alpha and feedback-locked theta, there was a relationship between feedback-locked beta and delay-discounting tendencies (see Table 1). Specifically, Ln(k) was negatively correlated with feedback-locked beta during both Reward and No-Reward trials. Thus, participants with stronger (i.e., less of a reduction in) feedback-locked beta power had smaller k values (i.e., stronger preference toward larger-but-delayed rewards), regardless of whether it was a Reward-trial or No-Reward-trial feedback. Moreover, while the correlation coefficient between Ln(k) and ΔReward (Reward-trial minus No-Reward-trial) feedback-locked beta was not significant, r(35) = −0.25, p = .13, the correlation became significant after one bivariate outlier was removed, r(34) = −.36, p = .03 (see Fig. 12). We employed the Minimum Covariance Determinant (MCD) estimator (Rousseeuw and Dierssen, 1999) to detect this bivariate outlier, using the Robust Correlation and LIBRA toolboxes in Matlab (Pernet et al., 2012; Verboven and Hubert, 2005). This so-called skipped correlation method is a robust parametric method designed for controlling for outliers while preserving statistical power compared to non-parametric method (Rousseeuw, 1984; Wilcox, 2012). Thus, participants with less of a reduction in feedback-locked beta power following Reward-trial, compared to No-Reward-trial, feedback (i.e., more positive ΔReward feedback-locked beta) had smaller k values. To investigate the specificity of this relationship, we removed the outlier and examined correlations between Ln(k) and the ΔReward feedback-locked beta following Bad-Performance feedback separately from the ΔReward feedback-locked beta following Good-Performance feedback. Ln(k) was marginally correlated with the ΔReward feedback-locked beta following Good-Performance feedback, r(34) = −.30, p = .07, but not with the ΔReward feedback-locked beta following Bad-Performance feedbacks, r(34) = −.11, p = .53. Additionally, the correlation between ΔPerformance (Bad minus Good-Performance) feedback-locked beta and Ln(k) was not significant.

3.6 Delay-discounting responses as predicted by multiple reward-processing EEG activity

Because multiple ΔReward (Reward minus No-Reward) EEG indices were correlated with Ln(k), an additional multiple-regression analysis was used to assess for shared vs. unique effects of these EEG indices (see Table 2). The ΔReward EEG indices included in this multiple-regression analysis were ΔReward pre-feedback alpha, ΔReward feedback-locked theta following Good-
Performance feedback and ΔReward feedback-locked beta (collapsing across Bad- and Good-Performance feedback). First, we examined the correlations among these indices (see Table 3), and found that ΔReward pre-feedback alpha and ΔReward feedback-locked theta following Good-Performance feedback were correlated with each other. However, ΔReward feedback-locked beta was not correlated with either ΔReward pre-feedback alpha or ΔReward feedback-locked theta following Good-Performance feedback. Then, the three ΔReward EEG indices were entered as simultaneous predictors with ln (k) as a criterion. In this model, while ΔReward pre-feedback alpha was not significant, ΔReward feedback-locked theta following Good-Performance feedback was significant, and ΔReward feedback-locked beta was marginally significant (p = .06). Because ΔReward pre-feedback alpha and ΔReward feedback-locked theta following Good-Performance feedback were correlated with each other (but not with ΔReward feedback-locked beta), these two EEG indices may account for similar variance in ln (k). However, this variance in ln (k) was unique from that of ΔReward feedback-locked beta.

### 4. Discussion

The current study tested the relationship between individual differences in delay-discounting tendencies and reward-processing at specific temporal stages. To operationalize individual differences in reward-processing, we examined EEG activity during a reward time estimation task. Our use of time- and frequency-specific EEG measures allowed us to separately investigate individual differences in reward-processing at the reward-anticipation stage (including, cued-locked delta during cue-evaluation and pre-feedback alpha suppression during feedback-anticipation) and at the reward-outcome stage (including, feedback-locked delta, theta and beta). All of our EEG indices were elevated during Reward, compared to No-Reward, trials. Moreover, these EEG indices significantly predicted behavioral performance during the time estimation task (reflected by smaller Motivated-Learning Inaccurate Estimation), highlighting their essential roles in modulating performance during a motivated-learning situation.

### 4.1. Reward-processing and individual differences in delay-discounting responses

Overall, results supported our primary hypotheses regarding the relationship between enhanced reward-processing and a stronger preference toward larger-but-delayed rewards among adolescents with higher delay-discounting tendencies. Higher theta and beta activity, which is associated with greater reward expectation, leads to a preference for smaller-but-immediate rewards over larger-but-delayed rewards. In addition, a stronger preference for large delayed rewards is related to weaker pre-feedback alpha suppression and stronger feedback-locked theta and beta activity. These findings suggest that an enhanced reward-processing of delayed rewards is associated with a stronger preference for delayed rewards. In contrast, a weaker preference toward delayed rewards, a stronger preference for immediate rewards, is related to a weaker reward-processing of delayed rewards. These findings are consistent with several previous studies that examined reward-related brain activity and delay-discounting tendencies.

### Table 3

Zero-order correlations among ΔReward (Reward-trial minus No-reward-trial) EEG indices that predicted ln (k).

<table>
<thead>
<tr>
<th>Index</th>
<th>r</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>ΔReward pre-feedback alpha</td>
<td>.30</td>
<td>.25</td>
</tr>
<tr>
<td>ΔReward feedback-locked theta following Good-Performance feedback</td>
<td>-.37</td>
<td>.18</td>
</tr>
<tr>
<td>ΔReward feedback-locked beta (collapsing across Bad- and Good-Performance feedback)</td>
<td>-.75</td>
<td>.39</td>
</tr>
<tr>
<td>ΔReward feedback-locked theta following Good-Performance feedback</td>
<td>-.30</td>
<td>.21</td>
</tr>
<tr>
<td>ΔReward feedback-locked beta (collapsing across Bad- and Good-Performance feedback)</td>
<td>-.23</td>
<td>.04</td>
</tr>
<tr>
<td>ΔReward feedback-locked theta following Good-Performance feedback</td>
<td>-.71</td>
<td>.71</td>
</tr>
<tr>
<td>ΔReward feedback-locked beta (collapsing across Bad- and Good-Performance feedback)</td>
<td>-.71</td>
<td>.14</td>
</tr>
</tbody>
</table>

Note. r² = .31 (p = .006); Collinearity statistics (Tolerance and VIF) indicate that multicollinearity between the predictors was not a concern.

* Became significant (p = .03) after a bivariate outlier was removed (see Fig. 12).
Third, psychiatric and health-related research has shown that dampened reward sensitivity and reduced reward-related brain function is a risk factor for high-risk and poor health-related behaviors (Johnson and Kenny, 2010; Stice et al., 2008). These data have been conceptualized within the Reward Deficiency Model of addiction, which proposes that persons with low reward sensitivity self-medicate negative emotions and/or attempt to elevate positive emotions through pursuing high-risk, short-term rewards (e.g., drugs, alcohol, high-fat diets) (Blum et al., 2000; Volkow et al., 2012). The logic of the Reward Deficiency Model is consistent with our results given that participants with reduced reward-related EEG activity in the present study had a stronger preference for smaller-but-immediate rewards. Additionally, individuals with the Met-allele of the COMT-gene, which has been associated with elevated reward-related brain function (Chen et al., 2004; Yacubian et al., 2007), show a preference for larger-but-delayed reward (Boettiger et al., 2007; Gianotti et al., 2012; Smith and Boettiger, 2012; Yacubian et al., 2007). Similarly, a recent study also found a stronger preference toward larger-but-delayed rewards as a result of medications designed to increase dopamine signaling in patients with Parkinson’s disease (Foerde et al., 2016). Thus, combined with results from the present study, we argue that there is growing evidence that elevated reward-processing is associated with a higher propensity to forgo smaller-but-immediate rewards, in favor of larger-but-delayed rewards.

4.2. EEG indices at each stage of reward-processing

Our use of EEG extends previous fMRI findings (Benningfield et al., 2014) by providing a more time-sensitive index of reward-related neural activity, allowing us to demonstrate the nature of the relationship between delay-discounting and reward-processing at specific temporal stages (e.g., during feedback-anticipation and reward outcome, but not cue-evaluation). Owing to the time-frequency decomposition of the EEG data, we also were able to show that these relationships were quite sensitive to specific neural frequency bands (e.g., theta, alpha and beta, but not delta).

4.2.1. Pre-feedback alpha-suppression during the reward time estimation task

With respect to feedback-anticipation EEG activity, we found a clear pattern of pre-feedback occipital alpha suppression peaking right before feedback onset. This pattern is consistent with previous reports suggesting that occipital alpha suppression prior to upcoming visual stimuli indexes anticipatory processes (Bastaansen et al., 1999; Hughes et al., 2013; Pfurtscheller and Aranibar, 1977). Also consistent with previous research (Hanslmayr et al., 2007; Hughes et al., 2013; van den Berg et al., 2014), occipital alpha suppression was modulated by reward conditions, as we observed greater alpha suppression during Reward, compared to No-Reward, trials. These findings are in line with the recent perspective that occipital alpha suppression indexes enhanced attentional processes during the anticipation of reward-related stimuli/feedback, and that this enhanced attentional processing is associated with individual differences in reward-processing (van den Berg et al., 2014). Moreover, our study is among the first to demonstrate that pre-feedback alpha suppression predicts enhanced motivated-learning during a reward task (reflected by smaller Motivated-Learning Inaccurate Estimation). This suggests enhanced attentional processing prior to feedback facilitates learning and behavioral adjustment to maximize rewards.

4.2.2. Pre-feedback alpha-suppression and delay-discounting responses

Most importantly, however, is the fact that stronger occipital alpha suppression, reflecting elevated feedback-anticipation neural activity, was associated with a greater preference toward larger-but-delayed rewards during the delay-discounting task. Given the temporal and frequency precision afforded by our EEG time-estimation paradigm, we were able to dissociate neural activity during reward feedback-anticipation from other reward-anticipation processes (e.g., cue-evaluation and motor-preparation). This degree of temporal precision is not feasible with fMRI studies of reward processing using paradigms such as the MID task (Benningfield et al., 2014), given the slow dynamics of the fMRI BOLD signal. We argue that the neurophysiological index of reward-anticipation neural activity related to delay-discounting responses in the present study (i.e., occipital alpha suppression) reflects the dis-inhibition of neural activity in sensory cortices that facilitate anticipatory attention to reward-related feedbacks (Hughes et al., 2013; Jensen and Mazaheri, 2010; van den Berg et al., 2014). Future research with alpha-suppression prior to feedback onset will have important implications for unpacking the nuances of reward-anticipation, particularly when aiming to control for other relevant processes, such as reward cue-evaluation or motor-preparation.

4.2.3. Reward-outcome EEG activity

With respect to reward-outcome EEG activity during the reward time-estimation task, we found that EEG activity in two frequency bands (feedback-locked theta and beta) were associated with individual differences in delay-discounting. Importantly, our study is among the first to separate the influence of reward-valuation from performance-valuation on individual differences in reward-outcome EEG activity (Leicht et al., 2013). Most studies of individual-differences of reward-outcome EEG data collapse across these two factors (e.g., Bress and Hajcak, 2013; Bress et al., 2012; Foti and Hajcak, 2009; Van den Berg et al., 2011). For instance, in several studies, good-performance feedback also indicates the winning of a monetary reward, while bad-performance feedback implies no-wining/losing of monetary reward. This makes it difficult to dissociate whether individual differences in outcome EEG data are driven by variability in reward-feedback evaluation, performance-feedback evaluation, or some combination of the two (Luft, 2014).

4.2.4. Feedback-locked theta during the reward time estimation task

For feedback-locked theta, our findings are in line with previous research suggesting that feedback-locked theta is sensitive to both reward- (Leicht et al., 2013) and performance- (Cohen et al., 2007; Luft, 2014) evaluation. Specifically, for reward evaluation, we observed greater feedback-locked theta during Reward (Catena et al., 2012; Morís et al., 2013), our study controlled for uncertainty. That is, participants with reduced re-

(footnote continued)
given that performance information during Reward trials indicated monetary-re-

ward earnings.
trials (compared to No-Reward trials). For performance evaluation, greater feedback-locked theta was found in response to Bad-Performance (compared to Good-Performance) feedback. These two main effects were independent of each other given the absence of a reward-by-performance evaluation interaction. Consistent with previous studies, feedback-locked theta predicted behavioral performance in our reward-learning task (Cavanagh and Shackman, 2015; van de Vijver et al., 2011). This confirms the role of feedback-locked theta in cognitive-control processes that incorporate feedback/outcome information (Luft, 2014). More importantly, by separating reward from performance evaluation, we demonstrated for the first time that reward evaluation (i.e., ΔReward feedback-locked theta), but not performance evaluation (i.e., ΔPerformance feedback-locked theta) predicted enhanced motivated-learning during a reward task (reflected by smaller Motivated-Learning Inaccurate Estimation). This suggests that the enhancement of cognitive-control during Reward (compared to No-Reward) trials facilitates one’s ability to learn through feedback in a reward situation to maximize reward receipt.

4.2.5. Feedback-locked theta and delay-discounting responses

Similarly, reward evaluation (but not performance evaluation) of feedback-locked theta predicted a greater preference to larger-but-delayed rewards. This may indicate that enhanced cognitive-control processes during Reward trials (reflected by reward evaluation), as opposed to prediction error or uncertainty resolution (reflected by performance evaluation), drove the relationship between reward-processing and delay-discounting responses. Moreover, subsequent analyses indicated that the relationship between ΔReward feedback-locked theta and delay-discounting responses was primarily driven by ΔReward feedback-locked to Good-Performance, but not Bad-Performance, feedback. This suggests that a preference toward larger-but-delayed rewards is particularly related to theta activity in the gain, but not the no-gain, domain, highlighting the fact that this relationship is specific to a reward-processing context.

4.2.6. Feedback-locked beta during the reward time estimation task

As for feedback-locked beta, our findings are consistent with studies showing stronger beta power (less reduction/desynchronization) to positive (compared to negative) feedback (for review, see Luft (2014)). By separating reward from performance evaluation, we extend previous research on feedback-locked beta power by showing the effect of both reward (stronger during Reward trials) and performance (stronger to Good-Performance feedback) evaluation. Additionally, greater enhancement of feedback-locked beta predicted enhanced motivated-learning during a reward task, replicating a previous time-estimation study (Luft et al., 2013a). Altogether, this pattern of feedback-locked beta is in line with the interpretation of this EEG index as a reward-specific signal (Cohen et al., 2007; De Pascalis et al., 2012; HajiHosseini et al., 2012; Marco-Pallares et al., 2008; Marco-Pallarés et al., 2009).

4.2.7. Feedback-locked beta and delay-discounting responses

Relating to our main focus on delay discounting, we demonstrate for the first time that reward evaluation (but not performance evaluation) of feedback-locked beta predicted a greater preference to larger-but-delayed rewards. This suggests that individual differences in delay-discounting may be related to the motivational saliency of the feedback (i.e., during Reward trials) as opposed to the positive behavioral performance signal from the feedback (i.e., a Good-Performance feedback).

4.2.8. Cue-locked and feedback-locked delta during the reward time estimation task

Although cue-locked delta and feedback-locked delta did not predict delay-discounting tendencies, the present study provides additional insight into their roles in reward-processing. During cue-evaluation in the reward-anticipation stage, cue-locked delta was stronger during Reward, compared to No-Reward, trials. This pattern is consistent with previous research focusing on a P3 component elicited by reward-related cues at similar time and topography (Broyd et al., 2012; Goldstein et al., 2006; Ramsey and Finn, 1997; Santesso et al., 2012). In line with a recent reinforcement study (Cavanagh, 2015), cue-locked delta predicted enhanced motivated-learning in our reward task (reflected by smaller Motivated-Learning Inaccurate Estimation). More importantly, motivated-learning was predicted by cue-locked delta power to the Reward cue (i.e., ΔReward cue-locked delta), suggesting an essential role of cue-locked delta in reward-processing, not just cue-evaluation in general. As for feedback-locked delta during the reward-outcome stage, similar to feedback-locked beta, we found an effect of both reward evaluation (stronger during Reward trials) and performance evaluation (stronger to Good-Performance feedbacks). This extends previous research that collapsed the two evaluation types together (Cavanagh, 2015; Foti et al., 2015; Leicht et al., 2013). More specifically, reward evaluation (ΔReward feedback-locked delta) of feedback-locked delta (not performance evaluation) predicted motivated-learning during the reward time-estimation task. This suggests that higher sensitivity to the motivational saliency of the feedback (reflected by stronger ΔReward feedback-locked delta) facilitates one’s ability to learn through feedback and adjust their performance to maximize gains. Taken together with other EEG indices, our study provides a comprehensive overview of reward-processing neural activity across specific temporal stages.

4.2.9. Relationship among reward-processing ERSP indices

We also report, for the first time, a relationship between reward-processing during feedback-anticipation (ΔReward pre-feedback alpha) and during reward-outcome (ΔReward feedback-locked theta following Good-Performance feedback) (see Table 3). ΔReward pre-feedback alpha, however, did not correlate with ΔReward feedback-locked beta. This suggests that enhanced attential processes prior to feedback (pre-feedback alpha suppression) was specifically related to enhanced cognitive-control processes during evaluation of reward-outcome (feedback-locked theta).

4.3. Limitations

This study is not without its limitations. First, similar to previous fMRI research on delay discounting and reward-related neural activity (Benningfield et al., 2014), our reward task did not include a punishment condition. Given that our focus was on individual differences in reward-processing, we were concerned that adding a punishment condition would unnecessarily lengthen the task. Task length is an important issue given fatigue has been shown to attenuate reward-related EEG data (Boksem and Tops, 2008). Future research involving a punishment condition is needed to examine whether the relationship between delay discounting and reward-related neural activity is primarily driven by a sensitivity to rewarding stimuli, or by a more general sensitivity to arousing or motivationally salient stimuli. Second, using undergraduate students may restrict the range of individual differences in impulsivity compared to other populations (Henrich et al., 2010). This is important, given that individual differences in impulsivity are associated with delay-discounting as well as reward-processing (e.g., Bjork et al., 2008; Hahn et al., 2009; for a review see Plichta and Scheres (2014)). Future research may employ other populations to further investigate whether the relationship between reward-related EEG activity and delay-discounting
manifests differently among individuals with high (e.g., hypomania) or low (e.g., depression) in impulsivity.

An important future direction is to examine the relationship between individual differences in reward-processing EEG activity at specific temporal stages to personality indices of reward sensitivity more generally. One of the most widely used personality theories of reward-sensitivity is Gray’s biopsychological theory of personality (Gray, 1987, 1989). In particular, Gray proposed that the behavioral activation system (BAS) underlies individual differences in appetitive motivation. Later personality researchers have separated the BAS into three subtypes: BAS Drive, Fun Seeking and Reward Responsiveness (Carver and White, 1994). The main difference between Gray’s BAS reward-sensitivity and reward-processing used in the current study is that Gray did not separate BAS into temporal reward-processing stages as is common in contemporary neuroscientific theories (Berridge, 1996; McClure et al., 2004; Schultz et al., 2000; Wise, 2008). Accordingly, it is unclear how Gray’s theory and the three proposed BAS subtypes map onto individual differences in reward-processing across each specific temporal stage of reward processing. Future research is needed to examine these relationships. Additionally, the time-estimation reward task used in the present study only provides immediate rewards, and does not provide delayed rewards. The delay-discounting task, by contrast, involves both immediate and delayed reward choices. Accordingly, we were only able to investigate the relationship between delay-discounting responses and neural activity to immediate-reward stimuli. Future research may wish to vary whether rewards are delivered immediately or following a delay, as in previous research (Cherniakowsky and Holroyd, 2013). Similarly, while the delay-discounting task in the present study employed hypothetical monetary rewards, the time-estimation reward task employed real monetary rewards. Although previous studies suggest a similarity between hypothetical and real monetary rewards in the delay-discounting task (M. W. Johnson and Bickel, 2002; Lagorio and Madden, 2005), future research should employ real monetary rewards in both tasks to test the generalizability of our findings.

5. Conclusions

In conclusion, one of the multiple factors that may modulate individual differences in delay-discounting responses is reward-processing. To comprehensively study individual differences in reward-processing, however, one needs to consider its heterogeneity in temporal dynamics. Here using EEG, we were able to separate reward-processing neural activity at each temporal stage into different indices based on time and frequency dimensions. In line with recent research (Benningfield et al., 2014; Boettiger et al., 2007; Foerde et al., 2016), we found that a stronger preference toward larger-but-delayed rewards was associated with enhanced reward-related neural activity at specific stages of reward-processing. Our findings not only substantiate the association between individual differences in delay-discounting responses and reward-processing, but also provide specific details about which stages of reward-processing drive these associations.

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