Distinct Neural Signatures of Outcome Monitoring After Selection and Execution Errors

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Abstract
Losing a point in tennis could result from poor shot selection or faulty stroke execution. To explore how the brain responds to these different types of errors, we examined feedback-locked EEG activity while participants completed a modified version of a standard three-armed bandit probabilistic reward task. Our task framed unrewarded outcomes as the result of either errors of selection or errors of execution. We examined whether amplitude of a medial frontal negativity (the feedback-related negativity [FRN]) was sensitive to the different forms of error attribution. Consistent with previous reports, selection errors elicited a large FRN relative to rewards, and amplitude of this signal correlated with behavioral adjustment after these errors. A different pattern was observed in response to execution errors. These outcomes produced a larger FRN, a frontocentral attenuation in activity preceding this component, and a subsequent enhanced error positivity in parietal sites. Notably, the only correlations with behavioral adjustment were with the early frontocentral attenuation and amplitude of the parietal signal; FRN differences between execution errors and rewarded trials did not correlate with subsequent changes in behavior. Our findings highlight distinct neural correlates of selection and execution error processing, providing insight into how the brain responds to the different classes of error that determine future action.

INTRODUCTION
When an action fails to produce the desired goal, there is a “credit assignment” problem to resolve: Did the lack of reward occur because the wrong course of action was selected, or was it because the selected action was poorly executed? Consider a tennis player who, midgame, must determine whether losing the last point was the result of selecting the wrong action or executing the action poorly. The player might have attempted a lob rather than the required passing shot, an error in action selection. Alternatively, a lob might have been appropriate but hit with insufficient force, an error in motor execution.

Reinforcement learning presents a framework for understanding adaptive behavior through trial-and-error interactions with the environment. According to numerous models (e.g., temporal difference learning; Sutton & Barto, 1998), the discrepancy between expected and actual outcomes, the reward prediction error (RPE), provides a learning signal that allows an agent to refine its predictions and update its action selection policy. However, what happens when a negative prediction error could arise from either poor action selection or poor response execution?

To address this question, McDougle et al. (2016) used a “bandit” task in which participants chose between two stimuli to maximize reward. In one condition, choices were made using a standard button-press method, a situation in which the negative prediction errors on unrewarded trials were attributed to poor action selection (given the negligible demands on motor execution). In a second condition, choices were made by reaching to the desired bandit. Here, unrewarded trials were attributed to movement execution errors. In the latter condition, participants strongly discounted the negative prediction errors on unrewarded trials relative to the former condition. The authors hypothesized that errors credited to the motor execution system block value updating in the action selection system. Consistent with this hypothesis, McDougle et al. (2019) reported that RPE coding in the human striatum was attenuated after execution errors, relative to selection errors. Differences between responses to selection and execution errors have been attributed to a greater sense of “agency” in the latter, with participants’ choice biases indicating a belief that they can reduce execution errors by making more accurate movements (Parvin, McDougle, Taylor, & Ivry, 2018).

A window into the processes that underlie outcome monitoring is offered through the discovery of the feedback-related negativity (FRN), a negative deflection in the EEG first identified after the presentation of
feedback indicating incorrect responses (Miltner, Braun, & Coles, 1997). After its identification, the component quickly became the subject of intense investigation as a marker signaling gains and losses (Gehring & Willoughby, 2002) and outcomes that are worse than expected (Holroyd, Hajcak, & Larsen, 2006). The most prominent explanation of its significance, the “reinforcement learning theory of the error-related negativity” (Holroyd & Coles, 2002), holds that the component (and its response-locked variant, the error-related negativity [ERN]) indexes the activity of signals from the midbrain dopamine that are conveyed to the ACC for adaptive modification of behavior (Holroyd & Umemoto, 2016; Holroyd & Coles, 2002). Recent developments reveal that much of the variation in this component is driven by a positive going component (a reward positivity [RewP]) responding to outcomes that are better than expected (Praudht, 2015; Poti, Weinberg, Dien, & Hajcak, 2011; Holroyd, Pakzad-Vaezi, & Krigolson, 2008). Irrespective of whether this signal is framed as a feedback negativity or RewP (here, we refer to this component as the FRN—the most widely used label), there is a consensus, as indicated by a meta-analysis of 55 data sets (Sambrook & Goslin, 2015), that it is sensitive to RPE.

The FRN's sensitivity to errors of action is more contentious. A series of experiments (Krigolson, Holroyd, Van Gyn, & Heath, 2008; Krigolson & Holroyd, 2006, 2007a) contrasting high-level (goal-attainment) errors, variously operationalized as a failure to reach a target (Krigolson et al., 2008; Krigolson & Holroyd, 2007a), avoid a collision (Krigolson & Holroyd, 2006, 2007b), and the erroneous selection of the wrong hand or force (de Bruijn, Hulstijn, Meulenbroek, & Van Galen, 2003) with low-level errors (i.e., mismatch between actual and intended motor commands), concluded that the latter do not elicit an FRN. Instead, reflecting a hierarchical error processing system (Krigolson & Holroyd, 2006), these motor errors are proposed to be mediated within posterior parietal cortex (Diedrichsen, 2005; Desmurget et al., 1999, 2001). Further elaborations indicated that the FRN may only be generated for action errors that cannot be corrected (Krigolson et al., 2008; Krigolson & Holroyd, 2007a), indicating a binary high-level coding of outcomes in the FRN (i.e., signaling whether the goal was achieved or not). In line with this, a recent experiment isolating reward-based and sensory error-based motor adaptation reported an FRN in response to binary reward feedback, but not sensory error feedback, which instead generated a P300 (Palidis, Cashaback, & Gribble, 2019). Previous work on the P300’s sensitivity to “low-level” motor execution errors led to the proposal that this later parietalement distributed component might reflect the revision of an internal forward model in posterior parietal cortex (Krigolson & Holroyd, 2007a).

A contrasting set of results suggest that the FRN (and its response-locked variant, the ERN) may in fact be sensitive to motor errors and reflect more than binary coding of outcomes, with evidence showing that it scales with the magnitude of error during sensorimotor adaptation (Anguera, Seidler, & Gehring, 2009) and correlates with the size of hand-path deviations after external perturbation to target reaches (Torrecillos, Albouy, Brochier, & Malfait, 2014). These findings are more in line with a growing body of work suggesting that the FRN indexes a general salience prediction error (Torrecillos et al., 2014; Oliveira, McDonald, & Goodman, 2007). A computational model attempting to unify a broad range of findings on medial pFC function (Alexander & Brown, 2011) proposes that this region is responsible for tracking discrepancies between expectations and outcomes, which are reflected in the FRN. Viewed in this way, the processing of execution and selection error may share a common neural network that signals a mismatch between the outcome and expectations in the service of behavioral adaptation (Torrecillos et al., 2014; Cavanagh, Zambrano-Vazquez, & Allen, 2012).

To test whether outcome errors of action and selection can be dissociated in the medial frontal cortex, we recorded feedback-locked ERPs while participants engaged in a modified bandit task where choices were selected via rapid arm movements. Unrewarded trials were either framed as errors in choosing the wrong bandit (a selection error) or the result of an inaccurate movement (an execution error). Following a large body of evidence reporting that the FRN is sensitive to RPE (Sambrook & Goslin, 2015), we expected that unrewarded outcomes attributed to selection error would elicit an FRN response. If this medial frontal monitoring system also tracks general action–outcome discrepancies, then we should expect a deflection after errors of action execution too. However, should the recently proposed movement-dependent account of reinforcement learning hold, the FRN response should be attenuated when errors can be ascribed to the motor system. We would expect P300 amplitude, a putative index of internal forward model revision (Krigolson & Holroyd, 2007a), to be largest for execution errors.

In addition to these predictions, we also examined the relationship between the FRN and behavioral modifica-
tion. Specifically, we predicted that participants who exhibited a larger change in the FRN would be more likely to switch between the different options. Notably, we expected this brain–behavior relationship would hold for selection errors, but not for execution errors. Reasoning that action errors may instead be encoding information about the size of the execution error, with this feedback used to correct discrepancies between the planned and actual outcomes, we explored the possibility that these signals may be correlated with the magnitude of error and subsequent change in motor response.

**METHODS**

**Participants**

Using an effect size estimate derived from our previous work on the FRN (η² = .167; Mushtaq, Wilkie, Mon-Williams, & Schaefer, 2016), with a desired statistical power of 0.8 and alpha criterion set at .05, we set a
Design and Procedure

We employed a novel three-armed bandit task (Figure 1) where the absence of reward on a given trial could be the product of a poorly executed action or an error in action selection (McDougle et al., 2019). Following EEG setup, the participant was seated in a chair approximately 50 cm away from a 24-in. ASUS monitor (53.2 × 30 cm [2560 × 1600 pixels], 100-Hz refresh rate). The participant was instructed to make a choice by making a reaching movement, sliding his or her right arm across a graphics tablet (49.3 × 32.7 cm; Intuos 4 XL, Wacom) while holding a digitizing pen encased inside a customized air hockey paddle. The tablet was placed below the monitor on the table and between an opaque platform that occluded the hand.

The experimental session comprised 400 trials, with opportunity for self-paced breaks. To initiate each trial, the participant made a reaching movement, sliding his or her right arm to position a white cursor (diameter of 0.5 cm) inside the home position, indicated by a solid white circle at the center of the screen. After maintaining this position for 400 msec, the start circle turned green and the participant made a reaching movement, sliding his or her right arm across the graphics tablet (hand passed through the bandit), the cursor was displayed within the spatial extent of the bandit. On these trials, there were two possible outcomes: (1) The bandit could turn green, indicating that a reward would be earned for the trial (reward outcome), or (2) the bandit would turn red, indicating that, although the movement was accurate, no reward would be given on that trial (selection error). If the movement missed the bandit, a cursor would appear indicating the position when the hand was at the radial distance of the bandits and thus indicate if the execution error was clockwise or counterclockwise relative to the target. The bandit would turn yellow, further signaling an execution error. Participants were informed of the three possible outcomes before the start of the experiment and presented with demonstrations of the three outcomes.

Following McDougle et al. (2019), each bandit had its own fixed probabilities for the three trial outcomes. All bandits had a 40% reward outcome, and thus, the expected values for the three bandits were identical. However, the frequency of selection error and execution error trials varied. For one bandit, 50% of the trials resulted in execution errors and 10% resulted in selection errors. We refer to this as the “high execution/low selection error” bandit. A second bandit resulted in execution errors on 10% of trials and 50% resulted in selection errors (a “low execution/high selection error” bandit). A third, “neutral” bandit produced an equal number (30%) of execution and selection errors.

Figure 1. Experimental task.
(A) Participants moved a stylus on a tablet to make rapid shooting movements (i) through one of three bandits (large circles) at 90°, 210°, and 330° relative to the home position (small circle). After a 1000-msec delay (not pictured), pseudoveridical feedback (white cursor) was provided indicating if the outcome was a reward (ii), a selection error (iii), or an execution error (iv). (B) The hand was occluded throughout, and stimuli were presented on a monitor positioned in front of the participants at approximately eye level.
where a score of 7 indicated strongly agree and 1 indicated strongly disagree. From 21 respondents, a mean score of 5.57 (SD = 1.6), which was statistically significantly different to the midpoint (neither agree nor disagree) on the scale, t(20) = 4.41, p < .001, indicated general agreement with the intended experimental manipulation.

The experimental task was programmed using the Psychophysics Toolbox (Kleiner et al., 2007; Brainard, 1997) and lasted approximately 35 min, with an additional 25–30 min of technical setup for EEG data acquisition.

Electrophysiological Data Recording and Preprocessing

EEG data were recorded continuously from 64 scalp locations at a sampling rate of 1024 Hz using a BioSemi Active-Two amplifier. Four EOGs—above and below the left eye and at the outer canthi of each eye—were recorded to monitor eye movements. Two additional electrodes were placed on the left and right mastoids. The CMS and DRL active electrodes placed close to the Cz electrode of the International 10–20 system served as reference and ground electrodes, respectively. EEG preprocessing was performed using the EEGLAB (Delorme & Makeig, 2004) and Fieldtrip (Oostenveld, Fries, Maris, & Schoffelen, 2011) toolboxes, combined with in-house procedures running using MATLAB (The MathWorks, Inc.).

All data were first rereferenced offline to the average of all channels and downsampled from 1024 Hz to 256 Hz. The continuous time series data were filtered using a high-pass filter with a cutoff at 0.1 Hz (Kaiser windowed-sinc FIR filter, beta = 5.653, transition bandwidth = 0.2 Hz, order = 4638) and a low-pass filter with a cutoff at 30 Hz (Kaiser windowed-sinc FIR filter, beta = 5.653, transition bandwidth = 10 Hz, order = 126). A second filtering of the data was performed for subsequent independent component analysis using a high-pass filter cutoff at 1 Hz (Kaiser windowed-sinc FIR filter, beta = 5.653, transition bandwidth = 2 Hz, order = 4666). Independent component analysis typically attains better decompositions on data with a 1-Hz high-pass filter (Winkler, Debener, Muller, & Tangermann, 2015). The data were segmented into epochs beginning 1 sec before and lasting 1 sec after the onset of feedback.

Infomax independent component analysis, as implemented in the EEGLAB toolbox, was run on the 1-Hz high-pass-filter epoched data, and the resulting component weights were copied to the 0.1-Hz high-pass-filter epoched data. All subsequent steps were conducted on the 0.1-Hz high-pass-filtered data. Potentially artifactual components were selected automatically using SASICA (Chaumon, Bishop, & Busch, 2015), based on low autocorrelation, high channel specificity, and high correlation with the vertical and horizontal eye channels. The selections were visually inspected for verification purposes and adjusted when necessary. After removal of artifactual components, the Fully Automated Statistical Thresholding for EEG artifact Rejection plug-in for EEGLAB (Nolan,
(Whelan, & Reilly, 2010) was used for general artifact rejection and interpolation of globally and locally artifact contaminated channels, supplemented by visual inspection for further periods of nonstandard data, such as voltage jumps, blinks, and muscle noise.

After artifact removal, 93.5% of total trials were available for analysis. There was no difference in the percentage of trials removed across conditions, $F(2, 56) = 2.09, p = .133$. However, as a product of the experimental design, there was a difference in the total number of trials between the conditions, $F(2, 56) = 85.2, p < .001$, with more reward trials ($\mu = 150, \pm 9$) available for analysis relative to execution error ($\mu = 114, \pm 12$, $t(28) = 12.21, p < .001$, and selection error ($\mu = 110, \pm 11$, $t(28) = 13.89, p < .001$, trials. There was no difference in trial counts for the two types of errors, $t(28) = 0.82, p = .693$. To increase the reliability of our conclusions by addressing potential problems of distribution abnormalities and outliers, averaged waveforms were constructed for each individual by taking the bootstrapped ($n = 100,000$) means from the EEG time series epochs. The waveforms were baseline corrected using a 200-msec time window before feedback onset.

**ERP Quantification**

Given that we had specific hypotheses, we focused our analysis on two locations. First, meta-analyses (Sambrook & Goslin, 2015; Walsh & Anderson, 2012) have shown the feedback-locked FRN effect to be maximal over the frontocentral region of the scalp. As such, we averaged activity across three frontocentral electrodes: FC1, FCz, and FC2. Second, given that the P300 (specifically, the P3b subcomponent) is commonly present in feedback-locked ERPs and typically maximal over parietal electrodes (Polich, 2007), we averaged over electrodes P1, Pz, and P2. Averaging across electrodes improves the signal-to-noise ratio of the ERP measures (Oken & Chiappa, 1986).

To test whether our results might be biased by the specific configurations of electrodes included in the averaged cluster and use of bootstrapped waveforms, we calculated the similarity between four different approaches to calculating the ERPs: (i) grand-averaged activity from the raw waveforms in the clustered electrodes, (ii) grand-averaged activity from the bootstrapped waveforms in the clustered electrodes, (iii) grand-averaged activity from raw waveforms from a single electrode (FCz for frontocentral analysis and Pz for parietal), and (iv) grand-averaged activity from bootstrapped means extracted from a single electrode. An intraclass correlation coefficient indicated a high level of agreement between all four approaches (frontocentral ICC = .995, 95% CI [0.989, 0.997]; parietal ICC = .996, 95% CI [0.994, 0.997]). Clustered bootstrapped averaged ERP waveforms are reported here.

With growing evidence that most of the variation in the FRN is driven by a RewP, we decided to make use of difference waveforms for our analysis to detect differences irrespective of whether they were driven by positive or negative deflections in the ERP (Krigolson, 2018). A difference waveform procedure has the added benefit of more easily isolating the FRN from components that precede (P2) and follow (a large P3 component comprising a frontal P3a and a parietal P3b), eliminating activity in common between two conditions (Kappenman & Luck, 2017). Most research on the FRN has typically computed RPE difference waveforms, derived by subtracting error/loss trials from reward trials (Sambrook & Goslin, 2015). Here, we created a “selection error” difference waveform by subtracting the average activity associated with selection error trials from the average activity related to all reward trials and an “execution error” difference waveform by subtracting the average activity associated with execution error trials from the average activity associated with reward trials. Finally, we directly contrasted execution and selection error ERPs by subtracting the execution error waveform from the selection error waveform to create an “error sensitivity” difference waveform. For statistical analysis, the parent waveform outcome trials were subjected to a one-way ANOVA, and where main effects emerged, one-sample $t$ tests were conducted to identify where these difference waveforms were significantly different to zero.

To reduce the number of false positives (Luck & Gaspelin, 2017), the ERP data were downsampled to 250 Hz and only activity between 150 and 500 msec (spanning the P2, FRN, and P3 ERPs) was analyzed. For each analysis, $p$ values were corrected by applying a false discovery rate control algorithm (Lage-Castellanos, Martínez-Montes, Hernández-Cabrera, & Galán, 2010; Benjamini & Hochberg, 1995). The Benjamini–Hochberg correction approach was adopted as previous studies have shown it to reliably control the false discovery rate when data are correlated, even when the number of comparisons is relatively small (Hemmelmann, Horn, Siusse, Vollandt, & Weiss, 2005). This method is also ideally suited for the exploration of focally distributed effects (Groppe, Urbach, & Kutas, 2011).

To aid the interpretation of the difference waveforms, we first visualized the grand-averaged ERPs related to each outcome. For every statistically significant contrast, we present the mean amplitude from the cluster for each parent waveform. Differences between relevant conditions at each electrode site are also visualized through topographical maps to support the interpretation of underlying components: Predicated on previous research (Walsh & Anderson, 2012), we anticipated that the FRN should show a frontocentral topography and, after an early frontocentral peak, there would be a subsequent posterior maximum corresponding to the P3b subcomponent of the P300 (Holroyd & Krigolson, 2007).

**Brain–Behavior Relationships**

A key question in this study is whether electrophysiological signatures of different types of outcomes correlate with
the participants’ choice behavior (see San Martín, 2012, for a review). On the basis of a reinforcement learning account of the FRN (Holroyd & Coles, 2002), we would expect the amplitude of the FRN to scale with the degree of behavioral adjustment: Large differences in the FRN should be more likely to lead to changes in choice behavior compared with small differences in the FRN. Here, we can ask this question with respect to both selection and execution errors.

To examine brain–behavior correlations, we calculated a behavioral adjustment score, or “switch bias” rate, for each participant (operationalized as the ratio of the percentage of trials that the participant switched after an error to the percentage of switching after a reward). This served as an intuitive index of how much participants favored one outcome over another. Mean amplitudes from the statistically significant clusters of EEG activity were then correlated with these behavioral adjustment scores.

Rather than signaling a need to switch from one target to another, feedback from execution errors might be more readily used to modify a motor plan for future action. To quantify the magnitude of cursor error, we calculated the angular deviation of the cursor relative to the center of the selected target. Hand error was calculated as the position of the hand relative to the center of the selected target and was different to cursor error only on trials with perturbed outcomes. The degree of motor correction was examined on a subset of data where participants selected the same target on consecutive trials and quantified as the degree of angular change in hand position relative to cursor position on the previous outcome. Mean cursor error and motor correction scores were correlated with mean amplitudes from the previously identified statistically significant clusters of EEG activity.

Statistical Analysis

For reporting purposes, time points are rounded to the nearest millisecond, amplitude (in microvolts) to two decimal places and p values to three decimal places. The range for the scalp maps was time-interval specific and determined by the 1st and 99th percentile values across all electrodes. Spearman’s rho (r_s) was used to examine correlations between amplitude and behavior. For correlations between behavior and neural activity, peak and mean amplitudes were extracted. Both are reported, and the strongest correlations are visualized. Where appropriate, pairs of correlations were directly compared with Hittner, May, and Silver’s (2003) modification of Dunn and Clark’s (1969) approach, using a back-transformed average Fisher’s z procedure as implemented in the R package Cocor v. 1.1–3 (Diedenhofen & Musch, 2015). The statistical significance threshold was set at p < .05. Generalized eta squared (η^2_g) is used as a measure of effect size for repeated-measures ANOVAs. This measure was selected over eta squared and partial eta squared because it provides comparability across between- and within-subjects designs (Bakeman, 2005; Olejnik & Algina, 2003); we considered η^2_g = .02 to be small, η^2_g = .13 to be medium, and η^2_g = .26 to be a large effect size. All statistical analyses were performed using R (R Core Team, 2015).

RESULTS

Behavioral Responses

A one-way ANOVA revealed a significant difference in Bandit Preference, F(2, 56) = 8.27, p < .001, η^2_g = .23, with participants exhibiting bias toward the high execution/low selection error bandit. Overall, this bandit was chosen, on average, on 39% (SE = 2%) of the trials, which was significantly greater than the low execution/high selection error bandit (M = 29%, SE = 1%), t(28) = 4.03, p = .001, and neutral bandit (M = 32%, SE = 2%), t(28) = 2.58, p = .046, with no difference for the latter two, t(28) = 1.07, p = .877. Consistent with previous work, when expected value is equal, the data show that participants prefer choices in which unrewarded trials are attributed to errors in movement execution rather than errors in action selection (Parvin et al., 2018; Green, Benson, Kersten, & Schrater, 2010; Wu, Delgado, & Maloney, 2009).

We then examined the effect of the different outcomes on the subsequent choice, asking how they influenced switching behavior (Figure 2A). Participants exhibited high switching rates overall (54%), but the rate differed according to outcome type, F(2, 56) = 10.23, p < .001, η^2_g = .11. Switching was highest after selection errors (M = 66%, SE = 5%) and markedly lower after execution errors (M = 42%, SE = 5%), t(28) = 5.22, p < .001. This difference is consistent with the hypothesis that motor errors attenuate value updating, perhaps because participants believe they have more control to correct for execution errors (Parvin et al., 2018).

Interestingly, switch rates after rewarded trials fell between the other two outcome types (M = 55%, SE = 6%). There was no difference between switch rates after reward relative to selection errors, t(28) = 1.85, p = .227, or execution errors, although the latter approached significance, t(28) = 2.46, p = .062 (following Bonferroni correction). The fact that many participants (18 of 29) were so prone to switching after a rewarded outcome and even more so (numerically) than after an execution error was unexpected. The high switching rates would suggest a bias toward exploratory behavior in this task, which might have been promoted by the relatively low rewards and/or the highly probabilistic nature of the outcomes (Cohen, McClure, & Yu, 2007; Daw, O’Doherty, Dayan, Dolan, & Seymour, 2006). Notably, there were very large individual differences in the treatment of the outcomes: Switch rates ranged from 3% to 98% after rewards, 7%–99% after selection errors, and 4%–81% after execution errors.
ERP Responses

Our primary aim was to examine whether selection and execution errors could be reliably distinguished in outcome-locked ERPs. To start, we ran an exploratory 3 (Bandit Type: High Execution/Low Selection Error vs. Low Execution/High Selection Error vs. Neutral) × 3 (Outcome: Reward vs. Selection Error vs. Execution Error) ANOVA at each time point for the frontocentral and parietal clusters. The main effect of Bandit Type was not significant ($p_s ≥ .702$), and there was no Bandit Type × Outcome interaction ($p_s ≥ .671$). Thus, we collapsed across the three bandits in our primary analyses of the three outcomes, allowing us to avoid increasing the FWE rate.

The grand-averaged ERPs related to each outcome are shown in Figure 2B and C. F tests revealed two significant clusters in the frontocentral region between 156–180 and 210–336 msec and three clusters in the parietal region (176–196, 218–239, and 355–438 msec). Descriptively, the first cluster in the frontocentral region was driven by a delay in the onset of an initial P200-like signal after an execution error, and the second cluster incorporated FRN deflections after selection and execution errors, along with subsequent positive deflections, likely reflecting the P3a subcomponent of the P300 signal (Polich, 2007). The early two clusters in the parietal region reflect shifts in the latency and amplitude of the execution error ERP, with the third cluster driven by the attenuation of the P3b subcomponent of the P300 after selection errors.

Figure 3A depicts the selection error difference waveform, derived by subtracting the selection error waveform from reward ERPs for the frontocentral cluster (shown in Figure 2B), and shows a statistically significant cluster of time points between 242 and 336 msec (one-sample $t$ tests of the difference wave against zero). An examination of the scalp topography of the first (242–289 msec) and second (289–336 msec) half of this window indicated a clear frontocentral maximum in the early phase, followed by a shift toward centroparietal maximum in the later part of the window (Figure 3B).

In line with the reinforcement learning account of the FRN, there was a relationship between neural activity and behavior. Specifically, amplitude (mean: $r_s = -.483$, $p = .009$; peak: $r_s = -.36$, $p = .052$; Figure 3C) from the early part of the cluster (capturing the FRN) negatively correlated with behavioral adjustment: The larger the difference waveform (i.e., greater negative deflection for selection errors relative to rewards), the greater the bias for the participant to switch to a different bandit after a selection error outcome relative to a reward outcome. We note that one participant had a switch rate score of $-0.87$, which was 2.97 SDs away from the mean. Rerunning the analysis without this participant showed a weaker relationship, but the pattern remained statistically significant (mean: $r_s = -.39$, $p = .042$; peak: $r_s = -.34$, $p = .074$).

The topographical map (Figure 3C, inset) demonstrates that this effect was localized to the frontocentral region. We found no evidence for such a relationship in the later, P3a, part of the time window ($r_s = -.08$, $p = .672$; Figure 3D). The mean FRN and P3a correlations were marginally different from one another ($z = 1.96, p = .05$), providing support that the FRN, but not the P3a, is a reliable correlate of behavior change.

Execution Errors

To examine the electrophysiological correlates associated with unrewarded outcomes attributed to motor execution errors, we performed similar analyses but now focus on
the comparison between execution error trials and reward trials (the execution error difference waveform—the result of subtracting the execution error ERP from reward ERPs in the frontocentral cluster shown in Figure 2B). This comparison revealed two statistically significant clusters—one ranging from 156 to 180 msec and a second between 207 and 325 msec (Figure 4A).

The first cluster showed an amplitude reduction in response to execution errors relative to reward trials. Similar to the selection error waveform result, we expected the second cluster would be contaminated by a P3a signal. Thus, we followed the same protocol, splitting this cluster into two equal intervals: (i) an early phase marked by the time interval of 207–266 msec and (ii) a later phase for activity between 266 and 325 msec. There was a clear frontocentral distribution for the early phase and, in the later time window, a shift toward centroparietal electrodes (Figure 4B).

We next examined the relationship between these three epochs (156–180, 207–266, and 266–325 msec) and behavioral adjustment (Figure 4C and E). The peak amplitude difference in the earliest interval (156–180 msec) correlated positively ($r_s = .37, p = .05$) with switching rates after an execution error relative to reward. After execution errors, smaller peaks in the 156–to 180-msec time window were associated with a lower tendency to switch. Note that this pattern is opposite to that observed between the amplitude of the FRN and behavioral adjustments after selection errors. The mean amplitude measure had a similar pattern of results but was not significant ($r_s = .35, p = .065$). An examination of topography revealed this correlation to be maximal in the frontocentral cluster, suggesting
that smaller amplitudes in response to execution errors early in the feedback processing stream are associated with a higher tolerance to this outcome.

In contrast to the results for selection errors, the FRN captured in the 207- to 266-msec time window did not correlate with behavioral adjustment ($r_s = .07, p = .722$). We tested, and confirmed, that this correlation was reliably different to the correlation observed for selection errors in the FRN time interval ($z = 2.40, p = .016$). There was no correlation between the execution error waveform amplitude (shown on the ordinate, where positive values indicate larger amplitude for execution errors relative to reward) positively correlated with an increase in the switch bias score (shown on the abscissa, where positive values indicate more switching after execution errors relative to reward) in this early time window, but there were no correlations in the later time windows (D and E).

We conducted the same analysis for the execution error waveform in the parietal cluster of electrodes. Execution errors elicited smaller amplitude responses relative to rewards in an early time window (176–196 msec) but elicited larger amplitude responses at 218–239 msec after feedback. In the later time window, there was a positive correlation between amplitude and behavior ($r_s = .47, p = .01$) in the posterior region, suggesting a shift from frontocentral to parietal regions in the processes driving behavioral adjustment (Dhar & Pourtois, 2011; Overbeek, Nieuwenhuis, & Ridderinkhof, 2005). Interestingly, and unexpectedly, the amplitude of the P3b subcomponent of the P300 signal—proposed to reflect the revision of internal forward models in posterior parietal cortex (Krigolson & Holroyd, 2007a), showed no difference in the processing of execution errors and rewards (see Figure 2C), and there was no relationship with behavioral adjustment ($r_s = -.01, p = .946$).

**Error Sensitivity Difference Waveform**

As described in the previous two sections, when using a common baseline (rewarded trials), we observed differences in both the ERP results and correlational analysis between unrewarded trials that were attributed to failures in movement execution or action selection. We
performed a direct comparison between these two types of unreWARDED outcomes by analyzing an error sensitivity difference waveform, subtracting the ERP for selection errors from the ERP for execution errors (see Figure 2B for the parent waveforms).

In the frontocentral cluster, there was a significant difference in the range of the FRN (222–250 msec; Figure 5A and B). We had anticipated that the amplitude of the FRN would be attenuated after execution errors, assuming a lower response would be reflective of reduced value updating (McDougle et al., 2019). However, the observed effect was in the opposite direction: Execution errors elicited a larger FRN deflection, relative to selection errors.

We also examined whether the magnitude of this difference correlated with the “switch bias” rate. For this measure, the proportion of switches after execution errors was subtracted from the number of switches made after selection errors. Note that these values range from 0 to −0.91, because of the fact that no participants produced more switches after execution errors relative to selection errors. Although the parent waveforms for this correlation are included in the previous analyses, the EEG activity in this analysis is specific to the range of 220–250 msec, the window in which the error outcome ERPs differed significantly.

There was no relationship between mean amplitude in this window and switch bias ($r_s = .23, p = .23$). However, the peak negative amplitude revealed a positive correlation with switch bias ($r_s = .41, p = .026$; Figure 5C). Participants who had relatively similar switching rates to the two unrewarded outcomes had smaller FRN differences, whereas individuals with a large negative bias (i.e., less switching after execution errors) also exhibited larger FRN amplitudes for motor execution errors relative to selection errors. This correlation was maximal in frontocentral sites (Figure 5C, inset).

Examining the parietal cluster revealed no differences in the earliest interval (176–196 msec). However, differences emerged in the 218- to 239-msec and 359- to 445-msec epochs, with larger positive amplitudes for execution errors relative to selection errors. The mean amplitude across each of these clusters (218–239 and 359–445 msec) was not correlated with the behavioral adjustment scores ($r_s \leq .179, ps \geq .352$).

**Kinematic Analysis**

To gain a deeper understanding of the relationship between brain activity and task performance, we examined correlations between task kinematics and the statistically significant periods identified in the time series analysis in the frontocentral and parietal difference waveforms. We reasoned that, in contrast to selection errors, where there was a relationship between FRN amplitude and choice selection, the execution error FRN may instead be encoding information about cursor position and subsequent movement correction.

In the first analysis, we examined whether there was a relationship between cursor error (the presented position of the cursor shown to participants at the end of the movement) magnitude and ERP activity. There were no reliable correlations between the mean activity of the statistically significant clusters in the difference waveforms and...
corresponding differences in cursor error magnitude (execution error: \( r_s = .228, \ p < .001 \); selection error: \( r_s = .176, \ p < .05 \); error sensitivity: \( r_s = .152, \ p < .20 \)).

In the second analysis, we asked whether ERP amplitude on the current trial would correlate with the degree of motor correction on subsequent trials. Here, we restricted analysis to the subset of trials in which participants chose the same target consecutively. The amount of motor correction in response to feedback (computed as the mean absolute change in end-point veridical hand position relative to the cursor position on the previous trial), varied as a function of Feedback, \( F(2, 56) = 75.37, \ p < .001 \). As both outcomes indicated a successful movement, we expected, and found, no difference, \( t(28) = 0.47, \ p > .999 \), in the subsequent degree of correction for selection error \( (M = 3.73°, \ SE = 0.15°) \) and reward \( (M = 3.64°, \ SE = 0.17°) \) trials. In contrast, execution error, signaling a need to change one’s motor response to hit the target \( (M = 6.55°, \ SE = 0.22°) \), had higher rates of correction relative to both selection error, \( t(28) = 8.95, \ p < .001 \), and reward, \( t(28) = 8.95, \ p < .001 \). Outcomes. Despite these behavioral differences, there were no correlations between the error sensitivity difference waveform and cursor error magnitude from 150 to 500 msec.

We found no correlations between ERP difference waveforms and cursor error in the frontocentral \( (ps \geq .45) \) or parietal \( (ps \geq .75) \) sites after correction. We also note, with a degree of caution given the corrected \( p \) values were not significant, that there was one statistically significant pattern before correction—a positive correlation between the error sensitivity difference waveform and cursor error \( (r_s = .43, \ 406 \text{msec}) \). In correlating motor correction rates with ERP amplitude, we found no significant relationships in the frontocentral cluster \( (ps \geq .45) \). Here, we noted that the strongest relationship \( (r_s = .456) \) was a positive one between motor correction and the error sensitivity difference waveform at 164 msec—a pattern that was sustained across 156–174 msec. As participants made larger degrees of correction after execution errors relative to selection errors, they also exhibited greater amplitude. In the parietal cluster, we found no reliable patterns of activity after \( (ps \geq .97) \) or before \( (ps \geq .1) \) correction.

### Perturbation Awareness

In a final set of explorations, we examined whether participants were sensitive to the feedback manipulation that had been applied to control the frequency of our three outcomes. In almost half the trials \( (M = 47.8%, \ SE = 0.01%) \), we delivered perturbed instead of veridical feedback \( (M = 52.2%, \ SE = 0.01%) \). We had taken measures to minimize the likelihood of participants becoming aware of these changes (e.g., no online movement feedback was provided, and end-point feedback was presented 1 sec after the stylus had passed the bandit), and in a postexperimental survey, participants indicated that they believed execution error outcomes to be the result of poor reaches, suggesting no explicit awareness of the manipulation. Nevertheless, we did find differences in cursor error (Figure 6A), as revealed through a 3 (Outcome: Reward vs. Selection Error vs. Execution Error) × 2 (Veracity: Veridical vs. Perturbed) interaction, \( F(2, 56) = 27.4, \ p < .001 \). In all cases, cursor error was largest in the veridical trials, but the effect was greatest for reward \( (M = 1.68°, \ SE = 0.02°) \); perturbed \( (M = 0.98°, \ SE = 0.01°) \), \( t(28) = 26.83, \ p < .001 \), and selection error \( (M = 1.72°, \ SE = 0.02°) \); perturbed \( (M = 0.97°, \ SE = 0.02°) \), \( t(28) = 30.95, \ p < .001 \). For execution error, there was a visual difference of \( 0.27° \) \( (M = 5.99°, \ SE = 0.07°) \); perturbed \( (M = 5.72°, \ SE = 0.04°) \), \( t(28) = 3.5, \ p = .045 \).

In examining hand error (position of the hand relative to the center of the target), we found a Veracity × Outcome interaction, \( F(2, 56) = 4770.99, \ p < .001 \). Veridical execution error trials \( (M = 5.99°, \ SE = 0.07°) \) were not statistically significantly different to perturbed selection error \( (M = 5.90°, \ SE = 0.07°) \), \( t(28) = 1.08, \ p = .886 \), and perturbed reward \( (M = 5.93°, \ SE = 0.07°) \), \( t(28) = 1.09, \ p = .881 \). Similarly, there was no difference in hand error for perturbed execution error trials \( (M = 1.75°, \ SE = 0.02°) \) compared with veridical selection error \( (M = 1.72°, \ SE = 0.02°) \), \( t(28) = 0.998, \ p = .915 \), and veridical reward \( (M = 1.68°, \ SE = 0.02°) \), \( t(28) = 2.41, \ p = .188 \). Participants did not alter their behavioral strategy in response to feedback perturbations \( (Veracity: F(1, 28) = 0.899, \ p = .351, \ eta^2 = .01; \ Veracity \times Outcome: F(2, 56) = 1.42, \ p = .251, \ eta^2 = .01) \). However, a suggestion that they might have been implicitly sensitive to these differences is indicated by the degree of motor correction after veridical and perturbed feedback (Figure 6D). One participant had no stay trials after perturbed feedback in this subset of data and was excluded from this analysis. In the remaining participants, we observed an Outcome × Veracity interaction, \( F(2, 54) = 4.49, \ p = .016, \ eta^2 = .04 \). There were no differences in the degree of motor correction after execution error \( (Veridical: M = 6.3°, \ SE = 0.19°) \); perturbed \( (M = 6.84°, \ SE = 0.32°) \), \( t(27) = 2.07, \ p = .047 \), but greater corrections (reward: \( M = 2.92°, \ SE = 0.13° \); perturbed: \( M = 4.28°, \ SE = 0.26°) \), \( t(27) = 4.56, \ p < .001 \); selection error: \( M = 3.02°, \ SE = 0.20° \); perturbed: \( M = 4.62°, \ SE = 0.17°) \), \( t(27) = 6.30, \ p < .001 \) after false hits' trials. These positively surprising outcomes (real reaches had missed the target on these trials, hence the perturbation) may
have prompted overcompensation as participants sought to calibrate their movements to task feedback.

Given these differences, we explored the extent to which the ERP signal was sensitive to the veracity of the feedback. We reran the ERP time-series analysis, performing a 3 × 2 (Outcome: Reward vs. Selection Error vs. Execution Error) × 2 (Veracity: Veridical vs. Perturbed) at each time point for the frontocentral and parietal clusters. There were no statistically significant main effects of Veracity ($F$s $\leq$ 6.99, $p$s $\geq$ .397) and no Outcome × Veracity interactions ($F$s $\leq$ 2.55, $p$s $\geq$ .79) in the frontocentral cluster and, similarly, no main effects ($F$s $\leq$ 5.42, $p$s $\geq$ .853) or Veracity × Outcome interactions ($F$s $\leq$ 1.83, $p$s $\geq$ .986) in the parietal cluster.

We then explored whether there were any differences in the relationship between ERP activity and kinematic adjustment as a function of feedback veracity. As perturbed feedback elicited larger corrective movements than veridical feedback, we speculated that an ERP signal sensitive to positive surprise may scale in response to this behavior for selection and execution error trials. To explore this idea, a difference wave subtracting perturbed ERP amplitude from veridical was computed. The amplitude of this "perturbation difference" waveform was correlated with (i) the mean difference in cursor error for veridical and perturbed feedback per outcome and (ii) the mean difference in degree of correction after veridical relative to perturbed feedback per outcome.

In analyzing the relationship between the perturbation difference waveform and cursor error in the frontocentral cluster, we found no correlations that survived correction for multiple comparisons ($p$s $\geq$ .616). However, in the

Figure 6. Feedback perturbation and awareness. (A) Cursor error was larger for veridical feedback relative to perturbed. (B) There was no difference in the magnitude of hand error for perturbed selection and reward error trials relative to veridical execution error trials and no difference between perturbed execution error trials compared to veridical selection error and reward trials. (C) Despite smaller cursor error, participants made larger corrections in response to perturbed feedback, with the pattern most pronounced for false hits. (D) Perturbed feedback did not impact on the likelihood of switching bandits. (E) Amplitude differences between perturbed and veridical feedback in the parietal cluster for selection errors at 273 msec (shown on the ordinate, where positive values indicate larger amplitude for veridical relative to perturbed outcomes) correlated with magnitude of the difference in cursor error for these outcomes (shown on the abscissa, where positive values indicate larger veridical cursor errors relative to perturbed). deg = degrees.
parietal cluster, the selection error waveform strongly correlated with perturbation difference amplitude at 275 msec ($r_s = -0.62$, $p = .011$; Figure 6E), indicating a sensitivity to discrepancies between actual and presented hand position. Specifically, this correlation shows that, for participants with larger veridical errors, perturbed feedback elicited larger positive amplitudes in a manner consistent with the P300 signaling surprise (Nassar, Bruckner, & Frank, 2019; Donchin, 1981). The error sensitivity difference waveform showed a similar pattern but did not reach the significance threshold after correction ($r_s = -0.47$ at 343 msec). The pattern for execution error was reversed, with the strongest correlation observed later ($r_s = 0.45$ at 492 msec)—with amplitude highest when both cursor error and amplitude were higher in the veridical condition relative to the perturbed condition. However, this too was not significant after correction.

In terms of the relationship between perturbation amplitude differences and the degree of motor correction, there were no significant effects in the frontocentral ($p_s \geq .120$) or parietal ($p_s \geq .82$) clusters. With the same note of caution for nonsignificant correlations offered above, two patterns suggest a further dissociation in the processing of selection and execution errors: In the time frame of the FRN, there was a relationship between frontocentral amplitude of the perturbation difference waveform and motor correction ($r_s = -0.542$ at 289 msec). Here, greater corrective movements in response to perturbed feedback correlated with larger differences in the FRN, and (ii) later in the window, the perturbation difference waveform for execution errors positively correlated ($r_s = 0.52$ at 355 msec) with the degree of motor correction, indicating that larger cursor error corrections in response to perturbed feedback have correspondingly larger amplitudes for perturbed feedback in the time range of the P3a. Despite the finding that selection error, like reward, resulted in adaptation after perturbed relative to veridical outcomes, no relationship was observed, with the strongest effect at 420 msec ($r_s = -0.299$).

Finally, as an alternative to averaging over perturbed and veridical trials, we correlated the degree of perturbation on a single trial, computed as the difference between hand error and cursor error (which was zero on veridical trials, a positive value on trials where the cursor was shown to be closer to the target than the hand position, and a negative value when the cursor position was shown to be further away from the target relative to hand position), with amplitude in the frontocentral and parietal clusters at each time point in the ERP per outcome for every participant. We did not find any general patterns to indicate a sensitivity to perturbation magnitude. In the frontocentral cluster, one participant showed a positive correlation between perturbation and the processing of reward (between 152–172 and 254–289 msec), another showed a correlation for execution error trials (between 70–86, 110–137, 188–204, and 289–500 msec), and two participants showed positive correlations for selection error. The first had a positive correlation between 453 and 457 msec, and the second had a positive correlation in multiple clusters across the whole time series (between 4–11, 31–90, 117–188, 258–277, and 460–477 msec). In the parietal cluster, no relationships emerged for reward or execution error, with two participants showing positive correlations between the degree of perturbation and the processing of selection error: one between 340 and 356 msec and a second participant between 289–317 and 382–500 msec.

DISCUSSION

Adaptive behavior necessitates distinguishing between outcomes that fail to produce an expected reward because of either the selection of the wrong action plan or poor motor execution. Although most decision-making research, in neuroscience as well as economics, has focused almost exclusively on the former, a few studies have shown that failed outcomes attributed to sensorimotor errors can markedly bias choice behavior (McDougle et al., 2016, 2019; Green et al., 2010). Here, we examined this issue by asking how an ERP signature of reinforcement learning, the FRN/RewP, varied in response to selection and motor errors. Predicated on the theory that the FRN is a scalp-related prediction error (Holroyd & Coles, 2002), we tested the hypothesis that errors attributed to failures in execution should lead to an attenuation in the FRN.

Consistent with our expectations, selection errors elicited a larger FRN relative to reward outcomes. Moreover, in line with a reinforcement learning account, the amplitude of the FRN after selection errors was negatively correlated with the probability that participants switched between the response options after feedback. Behaviorally, participants showed lower switch rates after execution errors, a pattern consistent with the hypothesis that the reinforcement learning system discounts these errors (McDougle et al., 2019). However, contrary to the prediction that FRN amplitude would be attenuated after execution errors, these errors actually produced the largest FRN. A striking difference between the ERPs in response to selection and execution errors was that the amplitude of the FRN after selection errors was predictive of behavioral biases and learning, whereas this ERP response after execution errors did not correlate with these variables.

Although almost all participants were more likely to switch after a selection error compared with an execution error, the differential response (i.e., difference in switch rates) to these two error outcomes varied considerably across participants. Moreover, this behavioral difference was correlated with the neural response to the two types of feedback: The more similarly participants treated the two outcomes at a behavioral level, the smaller the difference in FRN amplitude in response to these outcomes. These findings could be reconciled by considering the top-down mechanisms that may modulate how execution errors are processed. Behavioral experiments have shown...
that a sense of agency related to the perceived ability to correct for motor errors biases choice behavior (Parvin et al., 2018). In the present experiment, the finding that participants persevered with a bandit after execution errors but switched more often after selection errors also points toward differences in agency. Previous work on the FRN has shown that outcomes that can be controlled lead to a more negative FRN than those that cannot (Sidarus, Vuorre, & Haggard, 2017) and the FRN is attenuated in the absence of actively performed actions (Donkers, Nieuwenhuis, & van Boxtel, 2005; Donkers & van Boxtel, 2005). The finding that execution errors produced a larger FRN relative to selection errors is consistent with the presumed greater sense of agency associated with this type of unrewarded outcome.

A recent fMRI experiment using a three-arm bandit task, similar to that employed here, revealed an attenuation of the signal associated with negative RPE in the striatum after execution failures (McDougle et al., 2019). Our observation of a larger negative deflection for execution error trials in the FRN may appear contrary to these previously reported striatal results. However, the fMRI investigation did show increased ACC activity in response to execution errors compared with selection errors, suggesting that the former have their own neural signature.

With regard to the EEG response, there have been a number of studies reporting FRN deflections in response to execution error (Torrecillos et al., 2014; Anguera et al., 2009; Krigolson et al., 2008). These studies, in line with the prediction–response outcome model of medial frontal cortex function (Alexander & Brown, 2011), point to the existence of a general monitoring system that responds to violation of expectations. However, an important aspect of these tasks is that errors in movement execution typically resulted in high-level goal errors (e.g., failure to reach or remain on target in a manual tracking task) and/or involved the introductions of perturbations during the movement phase (Krigolson et al., 2008). This makes it difficult to rule out the contribution of cognitive control and response inhibition processes, which are known to generate an N200 component that shares similar spatial and temporal characteristics to the FRN signal (Holroyd et al., 2008; Holroyd, 2004). A recent study separating reward and sensory prediction errors in a motor adaptation task showed that the FRN responds to the former, but not the latter (Palidis et al., 2019). The present findings, indicating qualitatively different relationships between the two medial frontal negativities with behavioral modification, add weight to the possibility that execution error processing may be distinct from dopamine-related reinforcement learning processes.

We also observed two distinct patterns of activity in time windows preceding and after the FRN that provide further support for the claim of differential processing of execution and selection errors. First, smaller amplitude responses were observed after execution errors relative to rewards in frontocentral sites 156–180 msec after feedback, and the amplitude of this component correlated with switch rates. Second, in parietal sites (218–239 msec), larger amplitude responses occurred after execution errors relative to reward, and this difference was also correlated with switch rates. Importantly, in a reversal of the FRN pattern, magnitude differences in these early fronto-central and late parietal signals correlated with behavioral adjustment linked to execution errors. This pattern points toward the existence of distinct error monitoring systems operating at different levels of behavioral control (Yordanova, Falkenstein, Hohnsbein, & Kolev, 2004).

Exploratory analysis on the relationship between ERP amplitude and task showed that the degree of motor correction after execution errors relative to selection errors correlated with amplitude differences in an early fronto-central cluster (156–174 msec). The time course of this cluster closely mirrored that of the earliest difference between execution error and reward—where amplitude differences correlated with switch rates. Given that we had no a priori expectations for such a result and that this specific result did not survive correction for multiple comparisons, interpretations must be treated with caution and require further robustly powered replication work to confirm. Should future work replicate this pattern, it would add weight to the idea that the need to make a behavioral modification after an error in the motor system precedes the generation of the FRN.

A pertinent question of the present task and data is the extent to which participants were aware of the perturbations applied to the feedback to control outcome frequencies. Participants did not have access to online feedback, and end-point cursor information was presented with a 1-sec delay to minimize the likelihood of participants becoming aware of the perturbations. In a postexperiment survey, participants indicated that they had attributed execution errors to poor motor control. Consistent with this, we found that during the task, perturbed feedback did not alter choice strategy, nor did it result in any significant differences in the ERP. However, participants did, on average, make larger corrective movements after perturbed feedback—this was despite these outcomes showing smaller cursor errors than veridical feedback. In exploratory analysis, we did not find any relationships between amplitude and perturbation magnitude at a trial level for most of the participants, but we did find a correlation between amplitude differences and cursor error when averaging across perturbed and veridical trials. This correlation manifested in the parietal cluster at 273 msec, which likely reflected the onset of the P300. Here, the positive amplitude of this signal reduced as the amount of veridical error increased. That the P300 shows sensitivity to discrepancies between actual and presented hand position is consistent with the theory that the signal is generated through the active updating of an internal model of the environment (Donchin & Coles, 1988). The P300 is also notable for being a putative marker of conscious perception (Rutiku, Martin, Bachmann, & Aru, 2015). If participants
did indeed have access to this information during the task, it may be that these perturbations were not sufficiently large enough to signal a need to change strategy.

These findings also raise a broader question of whether the present results might be specific to outcomes that are framed as execution errors or extend to any endogenous or exogenous event that results in an unrewarded trial in which the outcome does not provide information about the reward probability associated with the selected object (Green et al., 2010). For example, if an unexpected gust of wind blew a tennis lob out-of-bounds, would that be treated as an “execution error”? Or, if after pulling the lever on a slot machine, a power failure caused the game to terminate without a payoff, would this affect how the choice is judged? A future study could test endogenous execution errors (e.g., reaching error) and exogenous errors (e.g., the task screen goes blank randomly before an outcome is delivered) more explicitly than the perturbations applied here. If similar results are found in both settings, elements of the early activity observed in frontocentral sites may indicate the establishment of a sensory “state,” representing that the intended action plan was not properly implemented, irrespective of whether this mismatch was because of endogenous or exogenous factors, even before the prediction error is evaluated. This echoes the sequential ordering in models of temporal difference learning, where first the agent perceives its state and then computes RPEs relevant to that state (Sutton & Barto, 1998).

Limitations and Future Directions

Although we have hypothesized that execution errors impact choice behavior, either by attenuating the operation of reinforcement learning processes or via an enhanced sense of agency, it is also important to consider alternative hypotheses. In the behavioral data, we observed a high base rate for switching between bandits. The highly probabilistic nature of the outcomes, coupled with the relatively low reward rate increased, made the task of determining the optimal choice difficult (although each bandit has different frequencies of execution and selection errors, they all had the same expected value). This may have biased participants toward an exploration strategy to reduce uncertainty by focusing on gathering more information about the reward likelihood of each bandit for later exploitation (Cohen et al., 2006). Viewed in this way, repetition of target selection after execution errors might not be because of increased agency or reinforcement learning discounting but may instead reflect a failure to acquire information on the reward probability of the chosen target on the previous trial and a drive to reduce uncertainty. Future work could disentangle these explanations by, for instance, assigning a lower expected value to high execution/low selection error bandits and/or through the presentation of fictive outcomes for motor errors.

Conclusion

We observed a robust FRN in response to both selection and execution errors, but only the former correlated with behavioral adjustment. In contrast, the amplitude of a positive deflection in the ERP, both before and after the FRN, correlated with choice behavior after execution errors. These results indicate a need for a more nuanced interpretation of what the FRN represents and how it may be shaped by contextual information. More generally, the results provide insight into how the brain discriminates between different classes of error to determine future action.

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Diversity in Citation Practices

Retrospective analysis of the citations in every article published in this journal from 2010 to 2021 reveals a persistent pattern of gender imbalance: Although the proportions of authorship teams (categorized by estimated gender identification of first author/last author) publishing in the *Journal of Cognitive Neuroscience* (JoCN) during this period were M(an)/M = .407, W(oman)/M = .32, M/W = .115, and W/W = .159, the comparable proportions for the articles that these authorship teams cited were M/M = .549, W/M = .257, M/W = .109, and W/W = .115, and W/W = .159, the comparable proportions encourage all authors to consider gender balance explicitly when selecting which articles to cite and give them the opportunity to report their article’s gender citation balance.

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