Learning of distant state predictions by the orbitofrontal cortex in humans

Abbreviated title: OFC in distant state predictions

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Abstract

Representations of our future environment are essential for planning and decision making. Previous research in humans has demonstrated that the hippocampus is a critical region for forming and retrieving associations, while the medial orbitofrontal cortex (OFC) is an important region for representing information about recent states. However, it is not clear how the brain acquires predictive representations during goal-directed learning. Here, we show using fMRI that while participants learned to find rewards in multiple different Y-maze environments, hippocampal activity was highest during initial exposure and then decayed across the remaining repetitions of each maze, consistent with a role in rapid encoding. Importantly, multivariate patterns in the OFC-VPFC came to represent predictive information about upcoming states approximately 30 seconds in the future. Our findings provide a mechanism by which the brain can build models of the world that span long-timescales to make predictions.
Introduction

Making predictions about the future has been proposed to be a core organizing principle of brain function (Rao and Ballard, 1999; Friston, 2005; Clark, 2013). Over short timescales, sensory and motor systems enable the prediction of immediate stimuli and consequences of our actions (Rao and Ballard, 1999; den Ouden et al., 2012; Summerfield and de Lange, 2014). Over longer timescales, predictions about the future state of the world – and the associated positive and negative consequences of our potential actions – are essential for goal-directed decision making and planning for the future. While the neural mechanisms of prediction in sensory and motor systems have been the subject of extensive research, we know little about the neural systems involved in learning to make predictions from experiences that extend beyond several seconds.

Neural systems including the hippocampus and the orbitofrontal cortex (OFC) have emerged as potential substrates for representing both spatial and abstract maps of the environment (Eichenbaum and Cohen, 2001; Wilson et al., 2014; Behrens et al., 2018). Building on the original discovery of place representations in hippocampal neurons (O'Keefe and Nadel, 1978), the hippocampus has been proposed to encode relations between stimuli that can combine to form a cognitive map of the environment (Eichenbaum and Cohen, 2001). Supporting a role for hippocampal representations in active prediction, research in animals has shown that when freely navigating an environment, activation of ‘place cell’ sequences can predict future movement trajectories (Pfeiffer and Foster, 2013; Olafsdottir et al., 2018). In humans, the hippocampus has also been shown to support the reactivation of visual associations originally separated by a few seconds (Wimmer and Shohamy, 2012). Other fMRI studies have found evidence for visual cortex reactivation for similar short-timescale
associations (Doll et al., 2015; Ekman et al., 2017), however, especially over longer timescales, such predictive knowledge must be supported by a still undetermined learning system outside of the sensory cortex (Summerfield and de Lange, 2014).

Research on the orbitofrontal cortex has recently found support for related cognitive functions as the hippocampus. While the OFC has long been viewed as critical for value representation and flexible learning, a unifying account proposes that the OFC represents a cognitive map of relevant states in the environment (Wilson et al., 2014; Bradfield et al., 2015; Stalnaker et al., 2015; Schuck et al., 2016; Schuck et al., 2018). For example, using fMRI in humans, Schuck et al. (2016) recently demonstrated that the OFC represented recently-observed hidden contextual information that was necessary for task performance. Interestingly, results from a recent study in rodents indicate that the OFC and the hippocampus interact to represent task-relevant structure and guide behavior (Wikenheiser and Schoenbaum, 2016; Wikenheiser et al., 2017). However, it is not known whether and how the hippocampus and/or OFC acquire and represent predictions during goal-directed behavior.

From a computational perspective, learning longer timescale predictions poses difficulties for reinforcement learning models that track only the value of states. Temporal difference learning models are constrained by the slow propagation of errors back through preceding states (Sutton and Barto, 1998), and neurally, the reward prediction error signal carried by midbrain dopamine neurons loses its fidelity beyond 5-10 s after feedback (Fiorillo et al., 2008; Kobayashi and Schultz, 2008). Alternative computational models that increase learning speed have been proposed, including eligibility traces (Sutton and Barto, 1998) as well as models that explicitly represent state transitions such
as successor representations and model-based learning (Daw et al., 2005; Stachenfeld et al., 2017; Gershman, 2018). Positive evidence for learning of long-timescale prediction in the hippocampus or OFC would support a neural mechanism underlying model-based or successor representation accounts of learning in multi-stage decision problems (Huys et al., 2012; Lee et al., 2014; Hassabis et al., 2017).

To investigate learning mechanisms supporting the prediction of distal future states, we focused on initial learning as participants navigated eight unique Y-maze environments, each of > 40 s duration. Participants learned to select the correct left or right door in an initial room (state 1) that led down one of two arms to a delayed reward (versus a loss) after moving through two intervening forced-choice states (Figure 1a, c). Critically, state 3 was represented by a unique and different category stimulus than state 1, allowing us to examine the development of predictive representations of state 3 when participants make their choice at state 1. The four repetitions of each maze were separated by at least 4 min, a delay that minimizes short-term maintenance of choice strategy and likely requires a contribution of longer-term memory for successful learning (Collins and Frank, 2012; Wimmer et al., 2018). By studying the initial learning process in relatively long-duration experiences, our goal was to better understand learning in a situation like those outside the lab where rapid learning is evident after only one or a few experiences.

Using fMRI in humans, we find that activity in the hippocampus is higher during initial exposures to each maze, potentially reflecting rapid encoding of a representation of the environment. Critically, across learning we also find that predictions of distal states
~ 30 s in the future come to be reflected in patterns of activity in the medial OFC / ventromedial PFC (OFC-VMPFC).
Figure 1. Learning phase task design and behavioral results. (a) Learning phase maze trial structure. Across 8 unique maze sequences, participants learned which action to make in state 1 in order to receive a deterministic reward at the end of the maze. Participants made a left or right choice in the first state and then proceeded through instructed left or right choices in state 2 and state 3, followed by reward or loss in the feedback state. Maze-unique stimuli from 3 categories (faces, scenes, and objects) were presented in state 1 and state 3. The delay between state 1 and state 3 was on average greater than 30 s, while repetitions of unique mazes were separated by at least 4 min. Critical decoding analyses focused on representations of information about state 3 at the onset of state 1 (represented by “Decoding” in the blue box). (b) Mean learning phase performance across four repetitions of each maze. Half of initial repetitions ended in reward and half in loss, and the resulting mean 50 % level of performance is indicated by the open square at repetition 1. Error bars represent standard error of the mean. (c) Illustration of the participant’s screen view, a cartoon room with potential left and right door options in each state followed by a path through a hallway between states. In state 2 and state 3, the instructed choice was indicated by a shift in the central stimulus to the instructed door. A localizer phase, which was used to derive classifiers for faces, scenes, and objects, followed the learning phase.
Results

Behavioral results. In the navigation learning experiment, behaviorally, participants showed a rapid increase in mean performance across repetitions of the 8 unique mazes, rising to 76% correct at the second exposure (CI [70.0 83.6]; t(34) = 22.81, p < 0.0001, t-test versus chance; Figure 1b). Initial feedback also exerted an effect on learning: if a reward was received on the first exposure, performance on the next repetition was significantly higher than performance following an initial loss (repetition 2, initial reward, 82.1% CI [76.0 88.2]; p < 0.0001, t-test; repetition 2, initial loss, 71.4% CI [62.1 80.8]; p < 0.0001, t-test; t(34) = 2.77, CI [2.8 18.6]; p = 0.009, t-test). Supporting the involvement of multiple cognitive systems in learning, we found that individual differences in working memory capacity as measured by the OSPAN task significantly correlated with mean accuracy over the learning phase (r = 0.39, p = 0.026, Pearson correlation; OSPAN performance mean 38.3, range 0-86; arithmetic component performance mean 89.2%).

The relationship between performance and working memory is likely related to the maintenance of state 1 information throughout the long delay until feedback; in contrast, it is unlikely to be related to working memory maintenance of choice policy across repetitions, as repetitions of the 8 individual mazes were separated by 4 min on average.

While episodic memory and working memory are often viewed as a distinct functions, the hippocampus is also known to be involved in critical aspects of working memory maintenance (Olsen et al., 2012).

fMRI univariate results. Our fMRI analysis first examined univariate learning-related brain responses. At the first state in a maze, state 1, we expected to find a learning...
signal reflected in an initially high and then (exponentially) decreasing response across repetitions in regions supporting memory and associative encoding. We indeed found a significant decreasing effect of exposure number (repetition) on BOLD activity in the bilateral hippocampus (right: $z = 4.42; p = 0.004$ SVC; 18 -10 -21; left: $z = 4.05; p = 0.02$ SVC; -18 -14 -21; Figure 2a, Supplementary Figure 1) as well as the right parahippocampal gyrus ($p<0.05$ whole-brain FWE-corrected; Supplementary Table 1). This effect was relatively selective, with only three other above-threshold clusters of activity in this contrast: two in the occipital lobe and one in the left dorsal premotor cortex (whole-brain FWE-corrected; Supplementary Table 1). The pattern of decreasing activity across repetitions was also selective to state 1; we found only a single significant cluster in the precuneus at state 2 and no significant clusters of activity in state 3 (Supplementary Table 1). Overall, the pattern of initially high and then decreasing activity over time in the hippocampus and parahippocampal cortex supports a role for the hippocampus and broader MTL in rapid episodic or relational learning.

Next, we confirmed expected responses to delayed reward feedback. We found that reward versus loss feedback, arriving ~ 40 s after the state 1 choice for each maze, was correlated with BOLD activity in a cluster including the ventral striatum and OFC-VMPFC (Figure 2b; Supplementary Figure 2). We also found reward-related activation in the left hippocampus ($z = 3.84; p = 0.028$ SVC; -30 -18 -15; this effect was also part of a larger significant cluster in the whole-brain analysis), in line with predictions from research on fMRI responses to relatively delayed feedback (~ 7 s; Foerde and Shohamy, 2011). Note that these results revealed some clusters of apparent activation on the edge of the grey matter which could be due to residual noise or motion; it is possible that such
effects could be removed with more advanced motion- and noise-correction algorithms (Pruim et al., 2015; Glasser et al., 2016). Interestingly, we found that neural processing related to the effect of a reward versus loss persisted through the 16 s rest period following feedback. During rest, feedback continued to differentially affect activity in a cluster including the OFC-VMPFC and ventral striatum, as well as two clusters in the right posterior hippocampus (OFC-VMPFC and ventral striatum, p < 0.05 whole-brain FWE-corrected; right hippocampus z = 4.75; p = 0.001 SVC; 32 -34 -3; z = 4.22; p = 0.007 SVC; 30 -18 -12; Supplementary Figure 2, Supplementary Table 1).
Figure 2. Learning-related univariate fMRI responses. (a) Across repetition of mazes during the learning phase, activity in the hippocampus significantly decreased, supporting a role for the hippocampus in maze encoding (right and left hippocampus, p < 0.05 SVC; image threshold p < 0.005 unc.) (b) At feedback, beginning > 40 s after the start of a maze, activation in the ventral striatum and OFC - ventromedial PFC, among other regions, was significantly activated by the receipt of reward versus loss. (Images p < 0.05 whole-brain FWE-corrected; full maps available at: https://neurovault.org/images/100616/ and https://neurovault.org/images/100619/.)

fMRI multivariate results. We next turned to multivariate analysis methods to pursue our primary question of whether learning mechanisms in the OFC-VMPFC and hippocampus acquire predictive information about future states. We examined an OFC
region of interest based on Schuck et al. (2016) (Figure 3a) and a hippocampal ROI based on an anatomical atlas. As a control, we also analyzed responses in visual category-selective regions in the occipital and ventral temporal cortex.

We first verified that a classifier trained on multivoxel patterns of activity from the post-learning localizer phase was able to generalize to accurately classify stimuli at state 1 and state 3 during the learning phase (state 2 did not contain any image stimuli; Figure 1a). Averaging across the three categories (faces, scenes, and objects), we found significant classification in all regions of interest (OFC-VMPFC mean AUC 9.88 CI [6.45 13.31]; t(30) = 5.88, p < 0.001, one-sample two-tailed t-test versus zero; hippocampus 10.63 CI [7.68 13.57]; t(30) = 7.37, p < 0.001, t-test; visual ROIs 48.21 CI [47.61 48.85]; t(30) = 158.42, p < 0.001, t-test). The classification of current state information in the OFC-VMPFC was comparable or higher than previous reports (e.g. Schuck et al. 2016).

For interpretation, it is important to note that conducting statistical inference on informational measures derived from cross-validation is problematic (see Methods) (Allefeld et al., 2016). However, our classification decision values are derived from a different method, cross-classification, where training is conducted on the post-task localizer and testing on the learning phase. Moreover, because our learning effect of interest tests for changes in information over time, our primary measures are completely isolated from this concern.

We then used the trained classifier to test for evidence of prospective future state representation at the onset of state 1 in each maze, ~ 30 s before the expected future state became visible. Our method tests how well patterns of activity at state 1 were related to the actual current stimulus category on the screen as well as the future state 3
stimulus category. The future state 3 information would be initially experienced upon the first rewarded repetition of a maze. In our analyses, we refer to this effect as ‘correct repetitions’; this variable is coded from 0 (no experience with the future state) to 3 (maximum possible experiences with the future state). Using raw classifier decision values derived from patterns of BOLD activity at state 1, which reflect the strength of evidence for each tested category, we analyzed whether decision values were explained by the current state (state 1) and future state (state 3), and whether this changed with learning across repetitions (see Methods). Note that the use of a regression analysis provides a measure of effects versus null (zero) but does not yield a percent correct measure as reported in many classification studies. Importantly, the application of the category classifiers to the learning phase also will not be affected by any learned associations between reward and the stimuli. The 3 categories appeared (at state 1 and state 3) in rewarded and non-rewarded paths in approximately equal numbers, which would lead to similar value associations across categories after learning. Further, any change in representation across learning cannot by definition be due to the applied classifier which is the same for all repetitions.

As expected, distributed patterns of activity in the OFC-VMPFC accurately discriminated the stimulus visible at the current state (including all participants; 0.092 CI [0.050 0.134]; t\(_{(30)}\) = 4.48, p < 0.001, t-test; in this test and all following, one-sample two-sided t-tests were used; Figure 3b). We found an effect of future state overall (0.052 CI [0.001 0.103]; t\(_{(28)}\) = 2.08, p = 0.047, t-test; in this and following results, analyses exclude participants with poor current state classification performance), although this effect is difficult to interpret. We also found a negative effect of correct repetition (-0.016 CI [-
0.029 -0.003]; \( t_{(28)} = -2.48, p < 0.02, t\)-test). Note that the above caveat about interpreting main effects of across-phase decoding also applies to the above effects of current and future state.

Critically, we found a significant increase in the representation of the future state across repetitions in the OFC-VMPFC (interaction between future state and correct repetition 0.051 CI [0.01 0.093]; \( t_{(28)} = 2.53, p = 0.017, t\)-test; Figure 3b). With a medium effect size (\( d = 0.47 \) CI [0.01 0.85]), a replication of the OFC-VMPFC result aiming for 80% power to detect an effect would require a minimum of 38 participants (two-tailed, uncorrected; \( n = 30 \), one-tailed). This effect of future state representation was selective, as the interaction between current state and correct repetition was near zero (0.007 CI [-0.041 0.055]; \( t_{(28)} = 0.30, p = 0.77, t\)-test). The current state by repetition effect was also statistically equivalent to a null effect, as indicated by an equivalence test using the TOST procedure (Lakens, 2017): the effect was within the bounds of a medium effect of interest (Cohen’s \( d = \pm 0.55 \), providing 80% power with 29 participants in the current analysis; \( t_{(28)} = -2.66, p = 0.007 \)), allowing us to reject the presence of a medium-size effect.
Figure 3. Multivariate OFC-VMPFC responses at state 1 onset related to the representation of anticipated future states. (a) Outline of the OFC-VMPFC region of interest shown over the across-participant average anatomical MRI. (b) Regression coefficients for decoded activity in the OFC-VMPFC for the current state (state 1), future state (state 3), the repetition since performance was correct, the interaction of current state with correct repetition, and the critical interaction of future state with correct repetition. Distribution density is represented by violin plot width; individual participant datapoints are in black. (* p < 0.05, t-test; error bars represent standard error of the mean; statistical comparisons were not completed on the current state representation in (b) because the plotted data represents results after exclusion of 2 participants with poor decoding during learning.) (c) Breakdown of the future state by correct repetition interaction, for illustration. Future state classification is plotted separately for each correct maze repetition; these effects were derived from control models examining current and future state separately for each correct repetition bin.

In the hippocampus, in contrast, while we observed evidence for representing the current state (0.0263 CI [0.0079 0.0446]; t(30) = 2.93, p = 0.007, t-test), we did not find any evidence for increasing representation of future states (interaction 0.008 CI [-0.018 0.035]; t(26) = 0.64, p = 0.53, t-test, Supplementary Figure 6). This effect was statistically equivalent to a null effect (t(26) = 2.32, p = 0.014, TOST). The future state interaction effect in the hippocampus was non-significantly weaker than the effect in the OFC-VMPFC (t(25) = 1.57; CI [-0.0931 0.126]; p = 0.129, paired t-test). While we
observed significant representations of the current state in our control regions in the posterior and ventral occipital cortex that responded to the categories of visual stimuli (0.318 CI [0.292 0.345]; t(30) = 24.62, p < 0.001, t-test), we did not find evidence for increasing representation of future states (interaction -0.005 CI [-0.0213 0.012]; t(30) = -0.5722, p = 0.57, t-test; Supplementary Figure 6). This effect was statistically equivalent to a null effect (t(30) = 2.38, p = 0.012, TOST). The future state interaction effect in the visual cortex was also significantly weaker than the effect in the OFC-VMPFC (t(28) = 2.76; CI [0.014 0.098]; p = 0.01, paired t-test).

Multivariate control analyses. We conducted several additional analyses to support the finding of future state representation across learning in the OFC-VMPFC. First, in a control analysis modeling each correct repetition bin separately, we found no pattern of change over time in current (state 1) representation in the OFC-VMPFC (Supplementary Figure 4). As expected, we observed an increase in future (state 3) information over time (Figure 3c) such that the strength of representation of the current visible state and the state anticipated > 30 s in the future was no longer significantly different by the last learning repetition (t(28) = 1.91; CI [-0.005 0.152]; p = 0.066, t-test).

We also confirmed that the increase in future state information in the OFC-VMPFC across learning was not sensitive to our exclusion of two participants who did not meet the orthogonal requirement of above-zero classification of both state 1 and state 3 on-screen stimuli. Across the full 31 participants, we continued to find a significant future state by correct repetition interaction (0.046 CI [0.006 0.086]; t(30) = 2.33, p = 0.027, t-test; Supplementary Figure 3). Next, in two control regression models, we removed the
current state or the future state from the model and found similar results for the included variables as in the full model above (Supplementary Figure 4), indicating that the result was not due to unexpected interactions between included variables. We then examined whether the pattern of future state representation was found across all three stimulus categories that make up the combined analysis or whether it was driven by particular categories. Indeed, we found a positive effect of the future state by correct repetition interaction across the separate face, scene, and object analyses that did not differ between categories (Supplementary Figure 5).

In a final control analysis, we examined the specificity of the OFC-VMPFC future by correct repetition effect. We examined 25 different ROIs covering the PFC based on functional coactivation patterns across published neuroimaging studies (Chang et al., 2018) (Supplementary Figure 10) plus 4 additional hippocampal subregions. Across all of the PFC sub-regions, one lateralized sub-part of the left dorsolateral PFC exhibited an effect at the 5 % level ($p < 0.0051$, t-test; Supplementary Table 2). Given the large number of exploratory regions considered and the lack of multiple comparisons correction, we do not interpret this result further. In the hippocampus, as an additional check of specificity, we further examined whether significant representations of future states were present but hidden by using the bilateral ROI. We separately analyzed the right anterior, left anterior, right posterior, and left hippocampus; across these 4 additional hippocampal ROIs, we found no future by correct repetition interaction effects ($p$-values $> 0.40$, t-test; Supplementary Table 2). The lack of general effects supports our a priori focus on the OFC-VMPFC, indicating relative selectivity. In turn, this selectivity indicates that this result is not likely to be due to a general confound in the
design or analysis that would produce a result frequently irrespective of an underlying true effect.

We next examined whether information representation at subsequent states in the maze trials reflected future expectations or recent experience. At state 2, which follows state 1 after approximately 7 s but does not include any stimuli on the screen, we found a positive but non-significant change in the representation of the future state 3 across learning in the OFC-VMPFC (interaction 0.040 CI [-0.002 0.082]; t(28) = 1.95, p = 0.062, t-test; Supplementary Figure 7). This state 2 interaction was in the same direction as the effect at state 1, potentially indicating that OFC-VMPFC representations of future states across learning continued through state 2. We found no interaction effects in the hippocampus and visual control regions (Supplementary Figure 7).

At state 3, we did not find any evidence for the representation of preceding states (information reflecting state 1 during state 3) in the OFC-VMPFC (Supplementary Figure 8). However, we did find that information about the current stimulus decreased across learning (interaction -0.060 CI [-0.108 -0.012]; t(28) = -2.56, p = 0.016, t-test; Supplementary Figure 8), which could reflect suppression of responses to presented stimuli as they become increasingly predicted by this circuit. A similar negative effect was observed in the visual control regions (-0.023 CI [-0.040 -0.007]; t(28) = -2.95, p = 0.006, t-test). Finally, during reward feedback, we found no significant representation of past states in any ROI (Supplementary Figure 9).
fMRI connectivity results. We finally returned to univariate analyses to examine whether the OFC-VMPFC and hippocampus may interact. In an initial general PPI analysis using the OFC-VMPFC region as a seed, we found robust positive connectivity between these two regions across different maze states, including state 1, the feedback state, and rest ($p < 0.0001$ SVC; Supplementary Figure 11). A conjunction of results across these three states showed that in addition to the hippocampus, activity in only a few clusters including the posterior cingulate and dorsomedial PFC remained at stringent uncorrected thresholds.

In a PPI model specifically testing for learning effects, we examined changes in OFC-VMPFC connectivity with the rest of the brain across repetitions of mazes that were successfully learned. We found no significant positive or negative main effects of learning repetition on OFC-VMPFC connectivity. However, we found a significant positive relationship between the strength of OFC-VMPFC connectivity and individual differences in the OFC-VMPFC acquisition of future state representation in a cluster including the right hippocampus, midbrain, and thalamus (Supplementary Figure 12; Supplementary Table 3). This indicates that increases in the OFC-VMPFC representation of future states across learning were related to concurrent increases in functional connectivity between the OFC-VMPFC and hippocampus. Additional clusters included the putamen as well as the dorsal medial and lateral PFC (Supplementary Figure 12; Supplementary Table 3). As a control, we conducted the same PPI analysis on later states in a maze trial (state 2, state 3, and the feedback state); here we found no significant relationship between OFC-VMPFC connectivity and the OFC-VMPFC future state interaction effect (Supplementary Table 3).
Discussion

Learning to predict the future is a critical aspect of goal-directed decision making. While previous research has focused on the relatively short-timescale prediction of sensory and motor events (Rao and Ballard, 1999; den Ouden et al., 2012; Summerfield and de Lange, 2014), we do not yet know how the brain acquires predictive representations of distal future states during goal-directed learning. In a simple learning task with extended sequences of experience, we found that participants learned over only a few repetitions to make correct choices. Using fMRI, we demonstrated that this rapid learning across repetitions – each separated by minutes – was related to initially high and then decreasing activity in the hippocampus and parahippocampal cortex. Critically, as learning progressed, we found an increasing representation of to-be-visited future states in multivariate patterns of activity in the OFC-VMPFC. By the last learning repetition, information representing a hidden state ~ 30 s in the future was of similar strength as the state currently visible on the screen. Our results indicate that a learning mechanism in the OFC-VMPFC may enable the human brain to represent the distant and unseen future consequences of goal-directed actions.

Our finding that the human OFC-VMPFC acquires distal future state representations provides novel support for the proposal that the OFC represents a cognitive map of the environment (Wilson et al., 2014; Stalnaker et al., 2015; Schuck et al., 2016; Schuck et al., 2018). Computationally, the representation of future state information in the OFC-VMPFC also lends support to reinforcement learning models that explicitly represent state-to-state associations, such as model-based or successor representation models (Daw et al., 2005; Stachenfeld et al., 2017; Gershman, 2018) over
additions to model-free temporal difference algorithms such as eligibility traces (Sutton and Barto, 1998). However, we cannot rule out a contribution of a cooperative model-free eligibility trace learning mechanism that could assist in state and action value learning by maintaining preceding state information until feedback (Gmaz et al., 2018; Maggi et al., 2018). While our experimental paradigm was designed to be relatively simple in order to isolate effects of interest during initial learning, our OFC-VMPFC findings have the potential to inform studies of more complex multi-state problems or decision trees (Huys et al., 2012; Lee et al., 2014; Pezzulo et al., 2015; Hassabis et al., 2017), especially in problems where the delay between a choice at one state and an outcome in a distal state is relatively long.

Previous experimental work in humans has shown that recent past (hidden) state information that is necessary for task performance is represented in the medial OFC (Schuck et al., 2016). In contexts that allow for predictions, other work has also revealed that the OFC may represent information about the identity of immediately upcoming rewards (Howard et al., 2015; Howard and Kahnt, 2018). Our results indicate that the medial OFC / VMPFC acquires predictive information over time, allowing the OFC to represent critical future state information for goal-directed decision making. The > 40 s duration of mazes in our experiment, while relatively short in comparison to many experiences outside the lab, also goes well beyond the timescale of common human fMRI studies of decision making. Other studies have found evidence for visual cortex reactivation for relatively short timescale predictive associations (Wimmer and Shohamy, 2012; Doll et al., 2015; Ekman et al., 2017), including during reward-based decision making (Doll et al., 2015). However, when state predictions extend beyond several
seconds, predictive knowledge must be supported by a learning system outside of the sensory cortex (Summerfield and de Lange, 2014). Our results indicate that a mechanism in the OFC-VMPFC is capable of fulfilling this role.

In the hippocampus, we found that initially high activity rapidly decreased across learning repetitions. This pattern is consistent with a rapid episodic encoding mechanism, where activity decreases as repetitions contain less new information (Kaplan et al., 2014). Notably, in the current experiment, repetitions of mazes were separated by at least 4 min, a delay that likely requires a contribution of longer-term memory for successful learning. A recent study from our group replicated the finding of a role for working memory in the maintenance of recently-learned value associations (Wimmer et al., 2018). This study used learning from immediate feedback to examine the distinct but related question of spacing of learning sessions across days instead of minutes. While the vast majority of previous human studies have examined memory for brief episodes (e.g. static pictures), our results extend what is known about learning-related hippocampal effects by showing that hippocampal activity decreases across learning selectively at the beginning of goal-directed episodes. This finding, however, contrasts with a previous report which found increases in event-onset hippocampal activity over repeated viewing of passive movies (Ben-Yakov et al., 2014). These different effects could be related to the goal-directed learning component of the current study, where participants are incentivized to predict future consequences. Behaviorally, the rapid learning we observed after even a single repetition fits well with recent research showing that humans can make value-based decisions based on information provided by specific past experiences (Gluth et al., 2015; Wimmer and Buechel, 2016; Bornstein et al., 2017).
For goal-directed decision making, this rapid encoding of maze experiences may work together with other learning mechanisms, potentially in the OFC, to allow us to seek out positive outcomes or avoid negative outcomes.

While we found that the OFC-VMPFC represented distal future states, we did not find evidence for future state representation in the hippocampus. Such a null effect is surprising, as research in rodents has demonstrated that the hippocampus represents future paths (Johnson and Redish, 2007; Pfeiffer and Foster, 2013). However, we did find a significant correlation between the increase in functional connectivity between the OFC-VMPFC and hippocampus across learning and the increase in OFC-VMPFC representation of future states across learning. Previous research in humans has found that hippocampal activity during reward learning co-varies with the activation of associated items in the visual cortex (Wimmer and Shohamy, 2012) and that hippocampal activity patterns represent retrieved information about currently cued target locations (Brown et al., 2016). Recent papers have reported reactivation of hippocampal activity patterns during rest periods following learning (Schapiro et al., 2018; Schuck and Niv, 2018) and that hippocampal activity co-varies with the reactivation of visual stimuli (Momennejad et al., 2018). Interestingly, in recent work using MEG, an imaging method with much faster temporal resolution, Kurth-Nelson et al. (2016) reported that fast sequences of activity reflected potential trajectories in a non-spatial environment. While no correlations between MEG activity and behavior were identified in that study, such rapid sequence-related activity, likely decoded primarily from the visual cortex, could be driven by the hippocampus in humans, similar to findings in rodents (Pfeiffer and Foster, 2013; Olafsdottir et al., 2018).
One limitation of the design was that our focus on question of relatively long timescales of future representation limited the design to include only 8 unique maze structures. For comparison, Schuck et al. report a stronger effect when examining OFC representation of the relevant task context (indicated by the previous and current trial stimuli) in a rapid event-related design with more than 10 times as many trials. Importantly, our OFC-VMPFC result is supported by numerous additional analyses that indicate specificity in time and consistency across different categories. The effect was also selective to the OFC-VMPFC, such that at an uncorrected level only 1 of 25 additional prefrontal regions of interest exhibited an effect. However, it will be important to replicate these findings in future studies and related paradigms.

The brain has been proposed to be a predictive machine (Rao and Ballard, 1999; Friston, 2005; Clark, 2013). Our results provide new support for how the brain is able to make relatively long-timescale predictions that are critical for goal-directed decision making. We find that as models of the environment are initially established, a learning mechanism in the OFC-VMPFC is capable of making long-timescale predictions about future states. At the same time, overall activity in the hippocampus correlated with maze exposure across learning. Our complementary results in the OFC-VMPFC and hippocampus, combined with their connectivity relationship, suggest that these regions may interact to support learning and decision making in goal-directed tasks (Wikenheiser and Schoenbaum, 2016; Wikenheiser et al., 2017). Further understanding of these mechanisms and their interaction will be important for understanding deficits in both learning the structure of the environment and in using these representations for goal-directed decisions that are found in various psychiatric disorders and addiction.
Methods

Participants. A total of 38 subjects participated in the experiment. Participants were recruited via a list of prior participants in MRI studies who had consented for re-contact, postings on an online university bulletin board, and referral from other participants. Participants were fluent German speakers with normal or corrected-to-normal vision and no self-reported neurological or psychiatric disorders or depression as indicated by questionnaire. Three participants were excluded based on behavior: one for previous participation in a related experiment, one for sleeping during the task, and one for chance-level performance. Thus, data from 35 participants were included in the behavioral analysis (21 female; mean age, 25.7 years; range, 19-31 years). After additional fMRI-based exclusion (detailed below), data from 32 remaining participants (18 female) were included in the univariate fMRI analysis. One additional participant was excluded from the multivariate analysis due to an error in the localizer phase. All participants were remunerated for their participation, and participants were additionally compensated based on the reward received during maze learning. The Ethics committee of the Medical Chamber Hamburg approved the study and all participants gave written consent.

Learning phase. The navigation learning task consisted of a learning phase followed by a localizer phase (Figure 1a, c). In the learning phase, across four repetitions for each of eight Y-maze sequences (plus two final non-analyzed mazes), participants attempted to learn the correct initial choice at state 1 in order to progress through the correct state 3 and reach a reward at the terminal feedback state. Maze- and state-unique stimuli from 3
categories (faces, scenes, and objects) were presented at state 1 and state 3 to enable decoding of state representations across learning. For the sequence design, we utilized an adapted Y-maze structure with a choice at state 1 and instructed choices at states 2 and 3 to both allow for rapid learning (by limiting the structure to two branches) and relatively long maze durations without excessive scan duration.

In each maze, the participant navigated through an illustrated series of rooms (states) with hallways in-between states (Figure 1a, c). At each stage in a maze, focal stimuli were presented in the center wall of the room. Two walls with space for doors extended to the left and right of the center wall. In state 1 and state 3, a maze- and state-unique stimulus was presented on the center wall. In state 2, a gray square was presented on the central wall. In state 1 and 3, both the left and right door options were initially visible. In state 2, only the single available pseudo-randomly determined (and fixed throughout learning) left or right door option was visible. In state 3, the non-available door was hidden after a short duration. In the reward state, no door options were visible. Between states, an illustrated hallway was presented with an occasional central upward-pointing arrow indicating to participants to press a key to continue advancing to the next state.

Maze duration, from state 1 until the end of the feedback state, was on average 48.5 s for the eight mazes of interest (range 38.3-77.4 s). The mean time between state 1 onset and reward feedback indicating the accuracy of the state 1 choice was 42.8 s (range 32.9-71.4 s). This delay from stimulus and action to feedback is well past the short range of time where dopamine responses show fidelity to a reward prediction error signal (Fiorillo et al., 2008; Kobayashi and Schultz, 2008). The mean time between state
1 onset and state 3 onset was 31.2 s (range 22.1-58.9 s). For the first participant, total maze duration was ~10 s shorter due to abbreviated inter-state periods (36.7 s mean duration), leading to shorter durations between state 1 and state 3 or reward. The delays between repetitions of the same maze were on average more than 4 min. Specifically, the delay between repetition 1 and repetition 2 was 4.4 min (mean 266.0 s, range 104.4 - 430.2). The delay between repetition 2 and repetition 3 was 4.6 min from (mean 278.5 s, range 102.2 - 1294.6). The delay between repetition 3 and repetition 4 was 6.9 min (mean 414.0 s, range 250.5 - 1266.0). This large delay strongly reduces the likelihood of between-repetition working memory maintenance as an explanation of learning performance (Collins and Frank, 2012; Wimmer et al., 2018), a common problem in reward learning paradigms where learning repetitions are separated only by several seconds on average.

At state 1, an illustration of a room with a superimposed central stimulus was presented for 4 s while a small white fixation cross was shown over the stimulus. When the fixation cross disappeared, participants were allowed to make their choice. After making a choice, the selected door opened slightly and the central stimulus moved to the selected door for 1.5 s. Next, the screen changed to a hallway phase for the inter-state period (described below). In states 2 and 3, the room illustration and central stimulus were shown for 2 s (state 2) or 4 s (state 3), after which the stimulus moved to the computer-determined (and still closed) door. Next, participants selected the indicated correct left or right button (for a maximum 30 s duration). A warning appeared if the participant was too late or if they selected the incorrect door. The selected door then opened slightly and the central stimulus moved to the door for 1.5 s. The screen then
changed to the hallway view. When participants reached the feedback state, the center wall was blank. They were required to press the bottom button to reveal the feedback. If the correct choice was made (or for the first repetition, if the maze had been assigned to be rewarded), a euro coin was displayed. Alternatively, a 50-cent euro coin with an overlaid red ‘X’ was displayed to indicate loss feedback. The feedback image was shown for 5 s. During the inter-state period between all states, after a 0.25 s blank screen, an illustration of a hallway was presented. An upward-pointing arrow appeared after 0.25 s. Participants progressed to the next state by pressing the top button when the arrow appeared. A series of two arrows, separated by a short delay, followed the first and second state, while one arrow followed the third state. The average duration from start to the end of a maze experience was 48.5 s. Trials were followed by a mean 16 s (range 15-18.5) inter-trial-interval during which a central white fixation cross was presented; the fixation cross changed to black for the 1 s preceding the start of the next maze.

For the eight mazes of interest, reward feedback was given on the first exposure for 4 mazes (independent of the state 1 left or right choice of the participant) and miss feedback was given on the first repetition for the other 4 mazes. Based on the initial state 1 left or right choice of the participant and whether the maze was assigned to end in an initial reward or loss, the ‘correct’ and ‘incorrect’ choice was assigned for subsequent repetitions for each maze. The mazes were presented in a pseudo-random order with staged introduction of new mazes as follows. The first four mazes were shown through three repetitions, such that each maze was repeated one time before another maze could be repeated again. Intermixed with the fourth repetition, two mazes from the second set were interleaved. At the fourth repetition for the second set of mazes, two
additional mazes were interleaved. These additional mazes always ended in a loss in order to better balance the reward and loss associations for probe questions in the subsequent localizer. During scanning, the experiment was divided into four blocks of 10 maze trials per block.

To measure internal activation of future state representations, each maze in the experiment included three categories of stimuli: faces, scenes, and objects. These were assigned to state 1 and the two alternative state 3 states. Color pictures of female faces set on a gray background were drawn from the Stanford Face Database, while scenes were drawn from an internal database; both sets of stimuli have previously been utilized to identify internal reactivation of state representations (Wimmer and Shohamy, 2012). Color pictures of objects were drawn from a previously used set of images compiled via internet search (Wimmer et al., 2014), composed of medium-sized items that can be manually interacted with (e.g. a book, water bottle, or guitar) set on a white background. Three pseudo-random orderings of the maze- and state-unique picture stimuli were used for counterbalancing. To indicate reward or loss feedback, a picture of a 1 euro coin was used for reward feedback while a picture of a 50 cent euro coin with a red ‘X’ drawn through it was used for miss feedback. Additionally, for reward feedback, outwardly-moving gold-colored dots were shown behind the coin to allow us to examine a potential motion-sensitive cortical response to rewards and anticipated rewards. To achieve this effect, a rapidly presented sequence of 12 static images started with only a few visible dots around the coin and then progressed as these and additional dots moved outward, giving the impression of a burst of dots moving away from the coin. These 12 images were repeated in sequence 3 times during the 5 s feedback period.
Before entering the MRI scanner, participants completed 1 trial of a practice maze. This practice maze was repeated again inside the MRI before the start of the experiment to ensure participants were familiarized with the screen and button box.

**Localizer phase.** To measure patterns of activity associated with the stimulus categories seen at states 1 and 3 in the learning phase, we next collected a “localizer” phase in the MRI scanner. During each trial, participants viewed either a stimulus from the learning phase, a scrambled object stimulus, or moving dot stimulus. For the face, scene, and object stimuli from the learning phase, participants were instructed to try to remember whether that stimulus was followed by reward or loss during learning so that they could perform well on occasional probe questions.

The localizer phase included 155 total trials, composed of 120 face, scene, and object trials, 20 scrambled object trials, and 15 “motion” trials with a moving dot background. The motion stimuli were composed of a gray hexagon with black outline and blue dots shifting across 12 static images, modeled after the reward feedback background gold dots seen during the learning phase. On each trial, static stimuli were presented for 2 s. For motion trials, the stimulus duration was 3 s. If a memory probe followed the picture, the text “Reward? ^ ” and “Loss? v ” appeared above and below the stimulus picture, respectively. Participants indicated their response with corresponding the top and bottom buttons. After a 0.5 s delay followed by a 0.25 s fixation period, a confidence rating scale appeared along with the text “How certain?”. The rating scale was composed of 4 levels of confidence: “not at all”, “a little”, “medium”, and “very”, which participants navigated using the left and right buttons on the box, confirming their
answer with the bottom button. After a 0.25 s pause, the task continued with the inter-
trial-interval indicated by a fixation cross (mean 2.5 s duration; range, 1.5-4.5 s).

The localizer phase was composed of 3 mini-blocks, each containing 40 face, 
scene, and object stimuli, 4-8 scrambled object stimuli, and 5 motion stimuli. Face, 
scene, and object trials were pseudo-randomly ordered, while scrambled object stimuli 
were shown at the end of the mini-block in two sets of 4, preceding and following a set of 
5 motion stimulus trials. Probe questions that assessed memory for the reward or loss 
associated with that stimulus during learning were included after 25% of the face, scene, 
and object trials to increase participant attention and to assess memory for the reward or 
loss association. The base list was modified for each participant, such that stimuli that 
were not seen during the learning phase were replaced with an additional 2 s inter-trial-
interval (mean, 9.9; range, 3-12). One participant was excluded from localizer analyses 
and multivariate analyses because an error in list creation led to an exclusion of 57 trials, 
yielding too few trials for accurate category classification.

After exiting the MRI scanner, participants completed two additional measures. 
We first collected the Beck Depression Inventory (BDI). The second measure we 
collected was the operation-span task (OSPAN) which was used as an index of working 
memory capacity (Lewandowsky et al., 2010; Otto et al., 2013). In the OSPAN, 
participants made accuracy judgments about simple arithmetic equations (e.g. ‘2 + 2 = 
5’). After a response, an unrelated letter appeared (e.g. ‘B’), followed by the next 
equation. After arithmetic-letter sequences ranging in length from 4 to 8, participants 
were asked to type in the letters that they had seen in order, with no time limit. Each 
sequence length was repeated 3 times. In order to ensure that participants were fully
practiced in the task before it began, the task was described with in-depth instruction slides, followed by 5 practice trials. Scores were calculated by summing the number of letters in fully correct letter responses across all 15 trials (Otto et al., 2013; Wimmer et al., 2018).

The experimental tasks were presented using Matlab (Mathworks, Natick, MA) and the Psychophysics Toolbox (Brainard, 1997). The task was projected onto a mirror above the participant’s eyes. Responses during fMRI scanning were made using a 4-button interface with a “diamond” arrangement of buttons (Current Designs, Philadelphia, PA). At the end of the experiment, participants completed a paper questionnaire querying their knowledge of the task instructions and learning strategy. For all parts of the experiment, verbal and on-screen instructions were presented in German; for the methods description and task figures, this text has been translated into English.

**fMRI Data Acquisition.** Whole-brain imaging was conducted on a Siemens Trio 3 Tesla system equipped with a 32-channel head coil (Siemens, Erlangen, Germany). fMRI measurements were performed using single-shot echo-planar imaging with parallel imaging (GRAPPA, in-plane acceleration factor 2; TR = 1240 ms, TE = 26 ms, flip angle = 60; 2 x 2 x 2 mm voxel size; 40 axial slices with a 1 mm gap) (Griswold et al., 2002) and simultaneous multi-slice acquisitions (“multiband”, slice acceleration factor 2) (Feinberg et al., 2010; Moeller et al., 2010; Xu et al., 2013) as described in (Setsompop et al., 2012). The corresponding image reconstruction algorithm was provided by the University of Minnesota Center for Magnetic Resonance Research. Slices were tilted approximately 30° relative to the AC–PC line to improve signal-to-noise.
ratio in the orbitofrontal cortex (Deichmann et al., 2003). Head padding was used to minimize head motion. Three participants were excluded for excessive head motion (a total of 5 or more > 2.0 mm framewise displacement translations from TR to TR per learning phase block).

During the learning phase, four functional runs of an average of ~10 min and 45 s were collected, each including 10 maze trials. During the localizer phase, one functional run of an average of ~15 min was collected. For each functional scanning run, four discarded volumes were collected prior to the first trial to allow for magnetic field equilibration and to collect reference data for multiband reconstruction.

Structural images were collected using a high-resolution T1-weighted magnetization prepared rapid acquisition gradient echo (MPRAGE) pulse sequence (1 x 1 x 1 mm voxel size) between the learning phase and the test phase.

**Behavioral Analysis.** Behavioral analyses were conducted in Matlab 2016a (The MathWorks, Inc., Natick, MA). Results presented below are from the following analyses: t-tests vs. chance for learning performance, within-group (paired) t-tests comparing differences in reward- and loss-associated stimuli across conditions, Pearson correlations, and Fisher z-transformations of correlation values. We additionally tested whether non-significant results were weaker than a moderate effect size using the Two One-Sided Test (TOST) procedure (Schuirmann, 1987; Lakens, 2017) as implemented in the TOSTER library in R (Lakens, 2017). We used bounds of Cohen’s $d = 0.53$, where power to detect an effect in the included group of 32 participants was estimated to be 80% ($d$ was adjusted accordingly when analyzing subsets of participants).
**fMRI Data Analysis.** Preprocessing was conducting using FMRIPREP version 1.0.0-rc2 (http://fmriprep.readthedocs.io) on openneuro.org, which is based on Nipype (Gorgolewski et al., 2011). Slice timing correction was disabled due to short TR of the input data. Each T1 weighted volume was corrected for bias field using N4BiasFieldCorrection v2.1.0 (Tustison et al., 2010), skullstripped using antsBrainExtraction.sh v2.1.0 (using the OASIS template), and coregistered to skullstripped ICBM 152 Nonlinear Asymmetrical template version 2009c (Fonov et al., 2009) using nonlinear transformation implemented in ANTs v2.1.0 (Avants et al., 2008). Cortical surface was estimated using FreeSurfer v6.0.0 (Dale et al., 1999).

Functional data for each run was motion corrected using MCFLIRT v5.0.9 (Jenkinson et al., 2002). Distortion correction for most participants was performed using an implementation of the TOPUP technique (Andersson et al., 2003) using 3dQwarp v16.2.07 distributed as part of AFNI (Cox, 1996). Functional data was coregistered to the corresponding T1 weighted volume using boundary based registration with 9 degrees of freedom implemented in FreeSurfer v6.0.0 (Greve and Fischl, 2009). Motion correcting transformations, T1 weighted transformation and MNI template warp were applied in a single step using antsApplyTransformations v2.1.0 with Lanczos interpolation. Framewise displacement (Power et al., 2014) was calculated for each functional run using Nipype implementation. For more details of the pipeline see http://fmriprep.readthedocs.io/en/1.0.0-rc2/workflows.html. The data were not resampled; voxel size remained at 2 x 2 x 3. For univariate analyses, images were then smoothed with a 6 mm FWHM Gaussian kernel.
General linear model analyses were conducted using SPM (SPM12; Wellcome Trust Centre for Neuroimaging). Prior to modeling, we applied a dilated mean anatomical mask to the functional data in order to provide a reasonable constraint on whole-brain multiple comparisons correction at the second level. Data from the four learning phase blocks was then concatenated, using a method developed by T. Sommer and adapted from Schultz et al. (2012). An autoregressive model, AR(1) (adapted to account for the concatenated sessions), was used to model serial autocorrelations in the data. fMRI model regressors were convolved with the canonical hemodynamic response function and entered into a general linear model (GLM) of each participant’s fMRI data. Six scan-to-scan motion parameters (x, y, z dimensions as well as roll, pitch, and yaw) produced during realignment were included as additional regressors in the GLM to account for residual effects of participant movement.

A univariate GLM was constructed to examine BOLD correlates of learning across repetitions and responses to reward feedback. We predicted that brain regions involved in encoding the stimuli and the associations in the maze episode would show a rapid decrease (or decay) in activity across repetitions. To capture this decrease, we created a decaying encoding predictor that decreased exponentially across repetitions. Specifically, a geometric decay of 50% was applied across the four repetitions, with values for the four repetitions of: 1, 0.50, 0.25, and 0.125. This encoding decay regressor was entered as a separate parametric modulator of activity at states 1, 2, and 3, although our primary prediction was about activity at the start, in state 1. At the reward state, reward and loss feedback were modeled as 1 and -1. We additionally modeled reward feedback during the subsequent rest period to test for lasting effects of reward on...
BOLD. States 1 and 3 and the feedback state were modeled as 5 s duration events. State 2 was modeled as a 4 s duration event. The rest period was modeled as a 12 s duration event. The 5 s duration of the State 1 and 3 primary events of interest captures the 4 s presentation of the central stimulus plus 1 additional second where the stimulus had shifted to the left or right door and matches the duration of the modeled feedback state events. Responses to the 8 mazes of interest in addition to the first exposure to maze 9 and 10 were included in the model. (Later repetitions of maze 9 and 10 were excluded because both choices were predetermined to lead to a loss.)

Our navigation learning task, where reward associations remained fixed across repetitions, did not allow for us to examine effects of choice value at state 1 or feedback due to collinearity with learning and reward variables. As a consequence, we could not examine whether responses to delayed reward feedback were better captured by reward magnitude versus reward prediction error, as suggested by neurophysiological studies in non-human primates (Fiorillo et al., 2008; Kobayashi and Schultz, 2008). Specifically, reward and choice value, the components of reward prediction error, were highly positively correlated (median r = 0.57) and thus we could not include them together in a model at the same time period. At stage 1, decaying encoding across repetitions and choice value were also highly negative correlated (median r = -0.71) and could not be included together in a model. At feedback, while the correlation between decaying encoding and reward was lower than the above correlations (median r = -0.46), we opted to not model decaying encoding at feedback or post-feedback rest.

Next, we analyzed the localizer phase to identify univariate patterns of activity that discriminated between the stimulus categories. These contrasts were used to identify
regions of interest for multivariate analyses. The initial GLM included separate regressors for faces, scenes, objects, scrambled objects, and motion stimuli. Static localizer stimuli were modeled as 2 s events while motion stimuli were modeled as 3 s events. From these first-level results, we computed contrasts with each of the three categories compared versus either of the two alternatives as well as each category versus the combination of both alternatives.

We additionally conducted psychophysiological interaction (PPI) analyses to examine functional connectivity between the OFC-VMPFC and the hippocampus. A general PPI was estimated first, looking at main effects of connectivity at different states in the maze trials. Each state was modeled as a 6 s event and rest was modeled as a 12 s event. The OFC-VMPFC ROI was as described above. Three general PPIs were estimated for state 1, the feedback state, and rest state. We computed a conjunction of these results to examine common OFC-VMPFC connectivity across multiple states during maze navigation.

A second PPI specifically examined changes in connectivity at state 1 across repetitions for successfully learned mazes. The psychological term in the PPI was a contrast between repetitions 2 and 3 (late learning) and repetitions 0 and 1 (early learning). Mazes where above-chance performance was not achieved were omitted from this contrast. Our goal in this analysis was to examine changes in connectivity that may relate to learning, so we additionally entered the OFC-VMPFC future by correct repetition effect as a covariate. This analysis tested for increases in OFC-VMPFC connectivity across learning that positively relate to increases in OFC-VMPFC future state.
representation across learning. Additional PPI models were estimated for state 2, state 3, and the feedback state as controls.

**Multivariate fMRI analyses.** Our tests of predictive representations relied on multivariate analyses because univariate analyses are not able to analyze representation content across multiple categories in a single region. In the learning phase we estimated a mass-univariate GLM using the non-smoothed BOLD data. Each stage as well as the rest period were each modeled with individual regressors, yielding 200 regressors of interest for the learning phase. The learning phase event durations were as above: 5 s for states 1 and 3 and the reward state, 4 s for state 2, and 12 s for the rest period. In the localizer phase we estimated a separate mass-univariate GLM using the non-smoothed BOLD data. Each trial was modeled as a 2 s event with an individual regressor, yielding 155 total regressors. Models included the six motion regressors; for the learning phase, block regressors were additionally included as effects of no interest.

In general, multivariate analyses were necessary to test our hypotheses for several reasons, including the following: when focusing on a single region, univariate analyses are insensitive to distributed patterns of information in cases where average signal does not differ between stimulus categories. Moreover, univariate analyses of a single region are not able to uniquely disambiguate between potential explanations for changes in responses over time, as any change could be due to multiple factors including attention, changes in the processing of the on-screen stimulus, and increases or decrease in response to one distal state option or the other distal state option.
Multivariate analyses were conducting using The Decoding Toolbox (Hebart et al., 2014). Classification utilized a L2-norm learning support vector machine (LIBSVM; Chang and Lin, 2011) with a fixed cost of $c = 1$. The classifier was trained on the full localizer phase data, which was divided into 3 mini-blocks. In the case of imbalanced numbers of face, scene, and object trials per mini-block (due to omitted stimuli not seen during learning) other trials were randomly left out to achieve balance. The trained classifier was then tested on the full learning phase data. Note that for the primary across-phase classification analysis, no cross-validation is necessary for training because no inferences are drawn and no results are reported on the localizer phase data. Based on the strength of discriminability in the localizer phase (using cross-validation), classification of faces was based on the comparison of faces versus scenes. Classification of scenes was based on the comparison of scenes versus faces. Classification of objects was based on the comparison of objects versus faces. While scrambled objects were included in the localizer phase to compare to objects, classification performance for objects versus scrambled objects was worse than that for objects versus faces. For objects in general, across OFC-VMPFC, hippocampal, and visual ROIs, classification performance was lower than that for faces or scenes. Learning phase classification is reported as area under the curve (AUC) which uses graded decision values and better accounts for biases in classification that may arise due to the different processes engaged by the localizer and learning phases.

The representation of current and future state information was analyzed by applying classifiers trained on the localizer phase to the patterns of activity evoked during maze learning, with a focus on the state 1 onset of each maze repetition. As
described above, one classifier for each category was trained based on the localizer phase data. These trained classifiers were applied to each stage of each maze trial to derive decision values, a parametric variable ranging from 0 to 1 representing how well the pattern of multivoxel activity in regions of interest at each state was described by the classifier. Importantly, because the classifier was trained after learning and all 3 categories would end with approximately the same average positive expected reward value, discrimination of localizer patterns would not be systematically influenced by learned value. Given this balance of value in the training data, it is not possible for any changes in classification over learning to be affected by shifts in value alone across learning.

The classifier decision values for each classifier and for each state were then entered into regression models to measure representations of current and future state. We computed three regression models, one for each category, to examine the information present in classifier decision values at state 1. These models estimated whether decision values were related to state 1 category, state 3 category, the number of correct maze repetitions (derived from behavior), and the two interactions between state 1 and state 3 category and the number of correct repetitions. State 1 and state 3 category were represented as binary variables, where 1 indicated that the state matched the category of interest and 0 otherwise. The number of correct repetitions variable was based on learning performance for each maze, focusing on whether the state 3 stimulus of interest had been reached during learning. For the first repetition, the value was set to 0. The value remained at 0 until the repetition following when the participant reached the correct state 3 and was able to observe the stimulus (and category) present in that state.
Subsequently, the value incremented by 1 for every subsequent repetition. The value was incremented for every repetition after the critical future state 3 had been visited, which allowed for this state information to be integrated into decision making at state 1. We made no assumptions about whether future predictions would be stronger or weaker if participants returned again to the incorrect choice. An alternative model where the correct repetition variable was only incremented on correct trials yielded qualitatively similar results. It is important to note that the use of a regression analysis across decision values for all trials provides a regression coefficient which we compare to a null (zero). However, this analysis does not yield a percent correct measure such as those reported in many classification studies, and consequently the resulting values should not be interpreted as a percentage measure. Note that while the localizer followed learning, any significant effects of learning on future state representation cannot be due to the classifier detecting activation of learned associations at test: any such confounding effects would only affect the main effect of future state representation, not the change of this representation across learning.

Within-participant, the results from the above regression models were averaged across the three categories to allow for group-level analyses (via $t$-tests). This approach was used instead of a single hierarchical mixed-effects model because each trial would be included three times in a single model, violating independence of observations. Participants’ data for individual categories were excluded from averaging across models if the classifier failed to generalize across task phases and classify learning phase state 1 and state 3 on-screen stimuli. Performance was calculated as the regression coefficient between (concatenated) decision values for state 1 and state 3 and the
contrast between the presence of the actual category of interest (e.g. faces) versus the comparison category (e.g. scenes) at state 1 and state 3. For 2 participants, the classifiers failed to generalize to any category during the learning phase and data from these participants were excluded from further analysis.

It has been shown that it is not valid to conduct statistical inference specifically on cross-validated classification accuracy measures of information using $t$-tests (Allefeld et al., 2016). In part, as informational measures cannot be below zero, assumptions underlying the $t$-test are violated for cross-validation within the same dataset. Our training and testing was conducted on separate datasets (“cross-classification” between the localizer and the learning phase) which does allow for potential “true” below-zero values, a case not addressed by Allefeld et al. (2016). Further, we found that our cross-classification results for current and future state in the OFC-VMPFC follows a normal distribution (Anderson-Darling goodness-of-fit hypothesis test). While the above concern may still apply to inferences made about the main effects of current and future state in our regression approach, our primary hypothesis rests on the change in information about future states across learning repetitions and not on comparing whether information content is different than zero. Thus, the use of a $t$-test is valid.

A first control analysis examined whether the exclusion of 2 outliers affected the primary future state by correct repetition results. Other analyses examined current, future, and past state representations at different states in the task (state 2, state 3, and the feedback state). We did not explore timepoint-by-timepoint modeling of the rest period due to signal-to-noise concerns: the category decoding of current states was at a
relatively low level and the search through many timepoints of data was less constrained than the modeling of specific states as above.

Additional control analyses examined different versions of the primary model above. First, we examined four separate models, one for each correct repetition bin, with current state and future state as predictors. This allowed us to examine the pattern of change in current and future state representation over time without the interaction included in the primary model. Second, we examined two alternative models where either current state or future state was omitted from the model. This allowed us to examine whether the inclusion of both current and future state in the same model significantly affected the results. Next, we examined generality of the future state by repetition effect: as the primary results collapse across the 3 stimulus categories (faces, scenes, and objects) we examined whether results were driven by a particular category or whether the same pattern was consistently observed across categories.

Finally, we examined whether other parts of the PFC represented information about future states across learning. This analysis informs whether any pattern of effects found in the OFC-VMPFC, hippocampus, or visual cortex was unique or instead commonly found throughout other frontal cortical regions. A common approach to whole-region analyses is to use a “searchlight” analysis, based on classification accuracy in a spherical region around each voxel. Our primary multi-category and multiple regression analysis of state information representation change (detailed above), however, makes a common searchlight approach unwieldy. As an alternative, we examined regions of the PFC in an approximately tiled manner. This approach has a benefit over searchlight
analyses in that it pools voxels together that have a common functional architecture, thus respecting the boundaries of different functional regions.

We conducted the classification and regression analyses separately in the left and right components of the bilateral ROIs, and thus the supplemental ROI analysis included an additional 25 regions (described below). We label the ROIs in Supplementary Table 2, where the abbreviations refer to: anterior OFC (antOFC), perigenual anterior cingulate (pgACC), dorsomedial PFC (dmPFC), frontopolar PFC (fpPFC), lateral PFC (latPFC parts a and b), lateral OFC (latOFC), ACC (ACC parts a, b, and c), subgenual ACC (sgACC), inferior PFC (infPFC), dorsolateral PFC (dlPFC parts a, b, c, and d). An additional set of analyses investigated the right and left anterior hippocampus (antHipp) and posterior hippocampus (postHipp).

**Regions of interest.** For univariate results, linear contrasts of univariate SPMs were taken to a group-level (random-effects) analysis. We report results corrected for family-wise error (FWE) due to multiple comparisons (Friston et al., 1993). We conduct this correction at the peak level within small volume ROIs for which we had an a priori hypothesis or at the whole-brain cluster level (both after an initial thresholding of p < 0.005 uncorrected). For univariate and multivariate analysis, we created two regions of interest in the OFC-VMPFC and hippocampus. We constructed the a priori OFC-VMPFC ROI (x, y, z: 0 44 -14; radius 10mm) based on a previous finding of short-term preceding state representations in the OFC (Schuck et al., 2016). The bilateral hippocampus ROI was derived from the Harvard-Oxford atlas – which provided the best overlap with our mean anatomical image – at a threshold of 25 %. We focused on the hippocampus
instead of the combined hippocampus and parahippocampal cortex to limit the size of the ROI for multivariate analyses. As control regions, we also constructed ROIs based on univariate responses to faces, scenes, and objects in the localizer phase. The face ROI was constructed based on the group contrast of faces versus scenes (derived from smoothed data at the first level) at a threshold of p < 0.001, selecting clusters surrounding (±42 -42 -22, including z < -14 and y < -68; z < 0 and y > -68). The scene ROI was constructed based on the contrast of scenes versus faces at a threshold of p < 0.001, selecting clusters encompassing the bilateral parahippocampal gyrus (including z < 4 and y < -68; z < 16 and y > -68). The object ROI was constructed based on the contrast of objects versus faces in symmetric clusters in the posterior occipital lobe at a threshold of p < 0.025 uncorrected (centered at ±58 -58 -10).

Our supplemental PFC regions of interest were selected from a 50-region whole-brain parcellation map derived from coactivation patterns across more than 10,000 published studies in the Neurosynth database (Chang et al., 2018)(http://neurovault.org/images/39711). From this parcellation mask, we extracted 14 ROIs (5 medial and 9 bilateral) in the prefrontal cortex with clustered spatial distributions and sizes approximately matching our OFC-VMPFC ROI, plus one additional bilateral ventral OFC ROI in a region lacking parcellation coverage (Supplementary Figure 10). The added bilateral ventral OFC ROI was located in the anterior mid-OFC centered at ±22, 53, -21. A stretched spherical ROI was drawn using a 10mm radius sphere with centers from y = 48 through y = 54. The existing ROIs were subtracted from this addition ROI, and the result was masked by the group whole-brain mask to remove voxels clearly outside of the brain. The resulting mask including all the above regions plus our 3
primary ROIs can be found here: https://neurovault.org/images/122507/. In addition to the PFC ROIs, in this supplemental analysis we also separated the bilateral hippocampal ROI into 4 parts: right and left anterior hippocampus (defined by y > -20) and right and left posterior hippocampus (defined by y < -20).

All voxel locations are reported in MNI coordinates, and results are displayed overlaid on the average of all participants’ normalized high-resolution structural images using the xjView toolbox (http://www.alivelearn.net/xjview) and AFNI (Cox, 1996).

**Data availability.**

Behavioral data are available on the Open Science Framework (https://osf.io/gt5kz/).

Whole-brain fMRI results are available on NeuroVault (https://neurovault.org/collections/4420/) and the full imaging dataset is available on Openneuro (https://openneuro.org/datasets/ds001019/)

**Code availability.**

Code for the multivariate analysis of current and future state representations is available at https://github.com/gewimmer-neuro/ofc-prediction.
References

Allefeld, C., Gorgen, K., and Haynes, J.D. (2016). Valid population inference for information-based imaging: From the second-level t-test to prevalence inference. Neuroimage 141, 378-392.

Andersson, J.L., Skare, S., and Ashburner, J. (2003). How to correct susceptibility distortions in spin-echo echo-planar images: application to diffusion tensor imaging. Neuroimage 20, 870-888.

Avants, B.B., Epstein, C.L., Grossman, M., and Gee, J.C. (2008). Symmetric diffeomorphic image registration with cross-correlation: evaluating automated labeling of elderly and neurodegenerative brain. Med Image Anal 12, 26-41.

Behrens, T., Muller, T.H., Whittington, J.C.R., Mark, S., Baram, A.B., Stachenfeld, K.L., and Kurth-Nelson, Z. (2018). What is a cognitive map? Organizing knowledge for flexible behavior. bioRxiv.

Ben-Yakov, A., Rubinson, M., and Dudai, Y. (2014). Shifting gears in hippocampus: temporal dissociation between familiarity and novelty signatures in a single event. J Neurosci 34, 12973-12981.

Bornstein, A.M., Khaw, M.W., Shohamy, D., and Daw, N.D. (2017). Reminders of past choices bias decisions for reward in humans. Nat Commun 8, 15958.

Bradfield, L.A., Dezfooli, A., van Holstein, M., Chieng, B., and Balleine, B.W. (2015). Medial Orbitofrontal Cortex Mediates Outcome Retrieval in Partially Observable Task Situations. Neuron 88, 1268-1280.

Brainard, D.H. (1997). The Psychophysics Toolbox. Spat Vis 10, 433-436.

Brown, T.I., Carr, V.A., LaRocque, K.F., Favila, S.E., Gordon, A.M., Bowles, B., Bailenson, J.N., and Wagner, A.D. (2016). Prospective representation of navigational goals in the human hippocampus. Science 352, 1323-1326.
Chang, C.C., and Lin, C. (2011). LIBSVM: a library for support vector machines. ACM Trans. Intell. Syst. Technol. 2, 1-27.

Chang, L.J., Jolly, E., Cheong, J.H., Rapuano, K., Greenstein, N., Chen, P.A., and Manning, J.R. (2018). Endogenous variation in ventromedial prefrontal cortex state dynamics during naturalistic viewing reflects affective experience. bioRxiv.

Clark, A. (2013). Whatever next? Predictive brains, situated agents, and the future of cognitive science. Behav Brain Sci 36, 181-204.

Collins, A.G., and Frank, M.J. (2012). How much of reinforcement learning is working memory, not reinforcement learning? A behavioral, computational, and neurogenetic analysis. Eur J Neurosci 35, 1024-1035.

Cox, R.W. (1996). AFNI: software for analysis and visualization of functional magnetic resonance neuroimages. Comput Biomed Res 29, 162-173.

Dale, A.M., Fischl, B., and Sereno, M.I. (1999). Cortical surface-based analysis. I. Segmentation and surface reconstruction. Neuroimage 9, 179-194.

Daw, N.D., Niv, Y., and Dayan, P. (2005). Uncertainty-based competition between prefrontal and dorsolateral striatal systems for behavioral control. Nat Neurosci 8, 1704-1711.

Deichmann, R., Gottfried, J.A., Hutton, C., and Turner, R. (2003). Optimized EPI for fMRI studies of the orbitofrontal cortex. Neuroimage 19, 430-441.

den Ouden, H.E., Kok, P., and de Lange, F.P. (2012). How prediction errors shape perception, attention, and motivation. Frontiers in psychology 3, 548.

Doll, B.B., Duncan, K.D., Simon, D.A., Shohamy, D., and Daw, N.D. (2015). Model-based choices involve prospective neural activity. Nat Neurosci 18, 767-772.

Eichenbaum, H., and Cohen, N.J. (2001). From Conditioning to Conscious Recollection: Memory Systems of the Brain (New York: Oxford University Press).
Ekman, M., Kok, P., and de Lange, F.P. (2017). Time-compressed preplay of anticipated events in human primary visual cortex. Nat Commun 8, 15276.

Feinberg, D.A., Moeller, S., Smith, S.M., Auerbach, E., Ramanna, S., Gunther, M., Glasser, M.F., Miller, K.L., Ugurbil, K., and Yacoub, E. (2010). Multiplexed echo planar imaging for sub-second whole brain FMRI and fast diffusion imaging. PLoS One 5, e15710.

Fiorillo, C.D., Newsome, W.T., and Schultz, W. (2008). The temporal precision of reward prediction in dopamine neurons. Nat Neurosci 11, 966-973.

Foerde, K., and Shohamy, D. (2011). Feedback timing modulates brain systems for learning in humans. J Neurosci 31, 13157-13167.

Fonov, V.S., Evans, A.C., McKinstry, R.C., Almli, C.R., and Collins, D.L. (2009). Unbiased Nonlinear Average Age-Appropriate Brain Templates from Birth to Adulthood. Neuroimage 47.

Friston, K. (2005). A theory of cortical responses. Philos Trans R Soc Lond B Biol Sci 360, 815-836.

Friston, K.J., Worsley, K.J., Frackowiak, S.J., Mazziotta, J.C., and Evans, A.C. (1993). Assessing the significance of focal activations using their spatial extent. Human Brain Mapping 1, 210-220.

Gershman, S.J. (2018). The Successor Representation: Its Computational Logic and Neural Substrates. J Neurosci 38, 7193-7200.

Glasser, M.F., Smith, S.M., Marcus, D.S., Andersson, J.L., Auerbach, E.J., Behrens, T.E., Coalson, T.S., Harms, M.P., Jenkinson, M., Moeller, S., et al. (2016). The Human Connectome Project's neuroimaging approach. Nat Neurosci 19, 1175-1187.
Gluth, S., Sommer, T., Rieskamp, J., and Buchel, C. (2015). Effective Connectivity between Hippocampus and Ventromedial Prefrontal Cortex Controls Preferential Choices from Memory. Neuron 86, 1078-1090.

Gmaz, J.M., Carmichael, J.E., and van der Meer, M.A. (2018). Persistent coding of outcome-predictive cue features in the rat nucleus accumbens. Elife 7.

Gorgolewski, K., Burns, C.D., Madison, C., Clark, D., Halchenko, Y.O., Waskom, M.L., and Ghosh, S.S. (2011). Nipype: a flexible, lightweight and extensible neuroimaging data processing framework in python. Frontiers in neuroinformatics 5, 13.

Greve, D.N., and Fischl, B. (2009). Accurate and robust brain image alignment using boundary-based registration. Neuroimage 48, 63-72.

Griswold, M.A., Jakob, P.M., Heidemann, R.M., Nittka, M., Jellus, V., Wang, J., Kiefer, B., and Haase, A. (2002). Generalized autocalibrating partially parallel acquisitions (GRAPPA). Magn Reson Med 47, 1202-1210.

Hassabis, D., Kumaran, D., Summerfield, C., and Botvinick, M. (2017). Neuroscience-Inspired Artificial Intelligence. Neuron 95, 245-258.

Hebart, M.N., Gorgen, K., and Haynes, J.D. (2014). The Decoding Toolbox (TDT): a versatile software package for multivariate analyses of functional imaging data. Frontiers in neuroinformatics 8, 88.

Howard, J.D., Gottfried, J.A., Tobler, P.N., and Kahnt, T. (2015). Identity-specific coding of future rewards in the human orbitofrontal cortex. Proc Natl Acad Sci U S A 112, 5195-5200.

Howard, J.D., and Kahnt, T. (2018). Identity prediction errors in the human midbrain update reward-identity expectations in the orbitofrontal cortex. Nat Commun 9, 1611.
Huys, Q.J., Eshel, N., O’Nions, E., Sheridan, L., Dayan, P., and Roiser, J.P. (2012). Bonsai trees in your head: how the pavlovian system sculpts goal-directed choices by pruning decision trees. PLoS Comput Biol 8, e1002410.

Jenkinson, M., Bannister, P., Brady, M., and Smith, S. (2002). Improved optimization for the robust and accurate linear registration and motion correction of brain images. Neuroimage 17, 825-841.

Johnson, A., and Redish, A.D. (2007). Neural ensembles in CA3 transiently encode paths forward of the animal at a decision point. J Neurosci 27, 12176-12189.

Kaplan, R., Horner, A.J., Bandettini, P.A., Doeller, C.F., and Burgess, N. (2014). Human hippocampal processing of environmental novelty during spatial navigation. Hippocampus 24, 740-750.

Kobayashi, S., and Schultz, W. (2008). Influence of reward delays on responses of dopamine neurons. J Neurosci 28, 7837-7846.

Kurth-Nelson, Z., Economides, M., Dolan, R.J., and Dayan, P. (2016). Fast Sequences of Non-spatial State Representations in Humans. Neuron 91, 194-204.

Lakens, D. (2017). Equivalence tests: A practical primer for t-tests, correlations, and metaanalyses. Soc Psychol Personal Sci.

Lee, S.W., Shimojo, S., and O'Doherty, J.P. (2014). Neural computations underlying arbitration between model-based and model-free learning. Neuron 81, 687-699.

Lewandowsky, S., Oberauer, K., Yang, L.X., and Ecker, U.K. (2010). A working memory test battery for MATLAB. Behav Res Methods 42, 571-585.

Maggi, S., Peyrache, A., and Humphries, M.D. (2018). An ensemble code in medial prefrontal cortex links prior events to outcomes during learning. Nat Commun 9, 2204.
Moeller, S., Yacoub, E., Olman, C.A., Auerbach, E., Strupp, J., Harel, N., and Ugurbil, K. (2010). Multiband multislice GE-EPI at 7 tesla, with 16-fold acceleration using partial parallel imaging with application to high spatial and temporal whole-brain fMRI. Magn Reson Med 63, 1144-1153.

Momennejad, I., Otto, A.R., Daw, N.D., and Norman, K.A. (2018). Offline replay supports planning in human reinforcement learning. Elife 7.

O'Keefe, J., and Nadel, L. (1978). The Hippocampus as a Cognitive Map (New York: Oxford University Press).

Olafsdottir, H.F., Bush, D., and Barry, C. (2018). The Role of Hippocampal Replay in Memory and Planning. Current biology : CB 28, R37-R50.

Olsen, R.K., Moses, S.N., Riggs, L., and Ryan, J.D. (2012). The hippocampus supports multiple cognitive processes through relational binding and comparison. Front Hum Neurosci 6, 146.

Otto, A.R., Raio, C.M., Chiang, A., Phelps, E.A., and Daw, N.D. (2013). Working-memory capacity protects model-based learning from stress. Proc Natl Acad Sci U S A 110, 20941-20946.

Pezzulo, G., Rigoli, F., and Friston, K. (2015). Active Inference, homeostatic regulation and adaptive behavioural control. Prog Neurobiol 134, 17-35.

Pfeiffer, B.E., and Foster, D.J. (2013). Hippocampal place-cell sequences depict future paths to remembered goals. Nature 497, 74-79.

Power, J.D., Mitra, A., Laumann, T.O., Snyder, A.Z., Schlaggar, B.L., and Petersen, S.E. (2014). Methods to detect, characterize, and remove motion artifact in resting state fMRI. Neuroimage 84, 320-341.

Pruim, R.H.R., Mennes, M., van Rooij, D., Llera, A., Buitelaar, J.K., and Beckmann, C.F. (2015). ICA-AROMA: A robust ICA-based strategy for removing motion artifacts from fMRI data. Neuroimage 112, 267-277.
Rao, R.P., and Ballard, D.H. (1999). Predictive coding in the visual cortex: a functional interpretation of some extra-classical receptive-field effects. Nat Neurosci 2, 79-87.

Schapiro, A.C., McDevitt, E.A., Rogers, T.T., Mednick, S.C., and Norman, K.A. (2018). Human hippocampal replay during rest prioritizes weakly learned information and predicts memory performance. Nat Commun 9, 3920.

Schuck, N.W., Cai, M.B., Wilson, R.C., and Niv, Y. (2016). Human Orbitofrontal Cortex Represents a Cognitive Map of State Space. Neuron 91, 1402-1412.

Schuck, N.W., and Niv, Y. (2018). Sequential replay of non-spatial task states in the human hippocampus. bioRxiv.

Schuck, N.W., Wilson, R.C., and Niv, Y. (2018). A state representation for reinforcement learning and decision-making in the orbitofrontal cortex. In Goal-Directed Decision Making: Computations and Neural Circuits, R.W. Morris, A.M. Bornstein, and A. Shenhav, eds. (London: Academic Press), pp. 259-278.

Schuirmann, D.J. (1987). A comparison of the two one-sided tests procedure and the power approach for assessing the equivalence of average bioavailability. J Pharmacokinet Biopharm 15, 657-680.

Schultz, H., Sommer, T., and Peters, J. (2012). Direct evidence for domain-sensitive functional subregions in human entorhinal cortex. J Neurosci 32, 4716-4723.

Setsompop, K., Gagoski, B.A., Polimeni, J.R., Witzel, T., Wedeen, V.J., and Wald, L.L. (2012). Blipped-controlled aliasing in parallel imaging for simultaneous multislice echo planar imaging with reduced g-factor penalty. Magn Reson Med 67, 1210-1224.

Stachenfeld, K.L., Botvinick, M.M., and Gershman, S.J. (2017). The hippocampus as a predictive map. Nat Neurosci 20, 1643-1653.
Stalnaker, T.A., Cooch, N.K., and Schoenbaum, G. (2015). What the orbitofrontal cortex does not do. Nat Neurosci 18, 620-627.

Summerfield, C., and de Lange, F.P. (2014). Expectation in perceptual decision making: neural and computational mechanisms. Nat Rev Neurosci 15, 745-756.

Sutton, R.S., and Barto, A.G. (1998). Reinforcement Learning: an Introduction (Cambridge: MIT Press).

Tustison, N.J., Avants, B.B., Cook, P.A., Zheng, Y., Egan, A., Yushkevich, P.A., and Gee, J.C. (2010). N4ITK: improved N3 bias correction. IEEE Trans Med Imaging 29, 1310-1320.

Wikenheiser, A.M., Marrero-Garcia, Y., and Schoenbaum, G. (2017). Suppression of Ventral Hippocampal Output Impairs Integrated Orbitofrontal Encoding of Task Structure. Neuron 95, 1197-1207 e1193.

Wikenheiser, A.M., and Schoenbaum, G. (2016). Over the river, through the woods: cognitive maps in the hippocampus and orbitofrontal cortex. Nat Rev Neurosci 17, 513-523.

Wilson, R.C., Takahashi, Y.K., Schoenbaum, G., and Niv, Y. (2014). Orbitofrontal cortex as a cognitive map of task space. Neuron 81, 267-279.

Wimmer, G.E., Braun, E.K., Daw, N.D., and Shohamy, D. (2014). Episodic memory encoding interferes with reward learning and decreases striatal prediction errors. J Neurosci 34, 14901-14912.

Wimmer, G.E., and Buechel, C. (2016). Reactivation of reward-related patterns from single past episodes supports memory-based decision making. J Neurosci 36, 2868-2880.

Wimmer, G.E., Li, J.K., Gorgolewski, K.J., and Poldrack, R.A. (2018). Reward learning over weeks versus minutes increases the neural representation of value in the human brain. J Neurosci 38, 7649-7666.
Wimmer, G.E., and Shohamy, D. (2012). Preference by association: how memory mechanisms in the hippocampus bias decisions. Science 338, 270-273.

Xu, J., Moeller, S., Auerbach, E.J., Strupp, J., Smith, S.M., Feinberg, D.A., Yacoub, E., and Ugurbil, K. (2013). Evaluation of slice accelerations using multiband echo planar imaging at 3 T. Neuroimage 83, 991-1001.

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Author Contributions

G.E.W. and C.B. conceived the project and designed the experiment. G.E.W. collected and analyzed the behavioral and fMRI data. G.E.W. and C.B. wrote the manuscript.

Additional Information

Supplementary Information accompanies this paper.

Competing Interests: The authors declare no competing interests.
Supplementary Information

Learning of distant state predictions by the orbitofrontal cortex in humans

G. Elliott Wimmer and Christian Büchel
Supplementary Figures and Supplementary Tables

Supplementary Figure 1. Timecourse of hippocampal activity at state 1 illustrating a decrease across learning repetitions, as shown the whole-brain univariate analysis in Figure 2. Sequence onset is at time 0; expected peak of BOLD response to state 1 is between 6.2 – 8.7 s. (Repetition 0 in darkest aqua; repetition 3 in lightest aqua; error bars represent SEM.)
Supplementary Figure 2. Reward-related univariate responses at feedback and during post-feedback rest, related to the univariate analysis presented in Figure 2. (a) Reward versus loss feedback at the reward stage, including left hippocampus (top), and overlaid with the OFC-VMPFC ROI (bottom) (b) Reward versus loss feedback during the post-feedback rest period, including overlap with the OFC-VMPFC ROI in cyan (bottom). (Images p < 0.05 whole-brain FWE-corrected.) Full maps available at: https://neurovault.org/images/100619/ and https://neurovault.org/images/100620/
Supplementary Figure 3. OFC-VMPFC coding of task information including all participants. (a) OFC-VMPFC coding of task information as shown in Figure 3b for comparison. (b) OFC-VMPFC coding of task information including the 2 participants that were excluded from the primary analysis because of below-zero discrimination of actual state 1 and state 3 on-screen categories (individual datapoints marked in yellow). (c) OFC-VMPFC effects as in (a) shown in a standard column chart format instead of a violin plot. (Error bars represent SEM.)
Supplementary Figure 4. State representation in the OFC-VMPFC across repetitions and control regression models. (a) Effects of current and future state modeled separately for each bin of correct repetition, as in Figure 3c. Current state in aqua; future state in magenta. (b) Information representing current state, correct repetition, and interaction of current state and correct repetition in a control model omitting future state. Note that effects are similar to the full model shown in Figure 3b. (c) Information representing future state, correct repetition, and the critical interaction of future state and correct repetition in a control model omitting current state. (* p < 0.05; ** p < 0.01; *** p < 0.001; error bars represent SEM.)
**Supplementary Figure 5.** Current and future state representation at state 1 in the OFC-VMPFC, as in Figure 3b, plotted separately for the 3 stimulus categories. (a) Information coding for face stimuli only. (b) Information coding for scene decoding only. (c) Information coding for object stimuli only. Current state in aqua; future state in magenta. (Error bars represent SEM.)
Supplementary Figure 6. Current and future state representation at state 1 in the hippocampus (a) and in visual control regions (b), as in Figure 3b. (c) Results from an analysis conducted separately for each correct repetition bin for illustration in the hippocampus. Current state in aqua; future state in magenta. (d) As in (c) for the visual cortex. (* p < 0.05; ** p < 0.01; *** p < 0.001; statistical comparisons for the hippocampus were not completed on the current state representation because the plotted data represents results after exclusion of participants with poor current state decoding. Error bars represent SEM.)
Supplementary Figure 7. State representation at state 2 in the OFC-VMPFC (a), hippocampus (b), and visual control regions (c). Note that the future by correct repetition effect in the OFC-VMPFC was positive ($p = 0.062$), similar to the effect at state 1. (* $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$; error bars represent SEM.)
Supplementary Figure 8. State representation at state 3 in the OFC-VMPFC (a), hippocampus (b), and visual control regions (c). (* p < 0.05; ** p < 0.01; *** p < 0.001; statistical comparisons for OFC-VMPFC and hippocampus were not completed on the current state representation because the plotted data represents results after exclusion of participants with poor current state decoding; error bars represent SEM.)
Supplementary Figure 9. State representation at the feedback state in the OFC-VMPFC (a), hippocampus (b), and visual control regions (c). (* p < 0.05; ** p < 0.01; *** p < 0.001; error bars represent SEM.)
Supplementary Figure 10. Regions of interest for additional testing of task information coding in the prefrontal cortex. Regions outside of the OFC-VMPFC and anterior OFC were derived from a functional coactivation map created from Neurosynth (Chang et al., 2018). Results are shown in Table S2. Abbreviations: anterior OFC (antOFC), perigenual anterior cingulate (pgACC), dorsomedial PFC (dmPFC), frontopolar PFC (fpPFC), lateral PFC (latPFC parts a and b), lateral OFC (latOFC), ACC (ACC parts a, b, and c), subgenual ACC (sgACC), inferior PFC (infPFC), dorsolateral PFC (dLPFC parts a, b, c, and d). Full mask available at: https://neurovault.org/images/122507/
Supplementary Figure 11. Connectivity (PPI) between the OFC-VMPFC and the hippocampus: conjunction of effects at state 1, feedback state, and post-feedback rest (individual maps thresholded at p < 0.0001 unc.). Full maps available at: https://neurovault.org/images/123490/, https://neurovault.org/images/123491/, https://neurovault.org/images/123492/
**Supplementary Figure 12.** Connectivity between the OFC-VMPFC and other regions for late versus early repetitions in successfully learned sequences that correlated with individual differences in the OFC-VMPFC future state by repetition effect. (a) A cluster including the hippocampus, midbrain, and thalamus (p < 0.05 whole-brain FWE). Blue and purple represent overlap with decay effects in hippocampus as in Figure 2 (p<0.005;
non-overlapping voxels in cyan). (b) Hippocampal cluster in a sagittal view. (c) Illustration of OFC-VMPFC – hippocampal correlation with OFC-VMPFC future state by repetition effect. (d) Additional clusters included the putamen as well as the dorsal medial and dorsal lateral PFC. (Images p < 0.05 whole-brain FWE-corrected.) Full map available at: https://neurovault.org/images/123494/
**Supplementary Table 1.** Summary of univariate analysis results. Clusters of activity exceeding whole-brain p < 0.05 FWE-corrected (cluster-forming threshold p < 0.005). Within each cluster, the first 10 regions are listed that include ≥ 10 voxels in a cluster. For the reward versus non-reward contrast at feedback, the cluster-forming threshold was set to p < 0.0005 to allow for informative clusters.

| Contrast       | Regions                                      | Cluster size | x   | y   | z   | Peak z stat |
|----------------|----------------------------------------------|--------------|-----|-----|-----|-------------|
|                | L Superior Frontal Gyrus                     | 454          | -30 | 26  | 54  | 3.78        |
|                | R Middle Occipital Gyrus                     | 450          | 18  | 100 | 18  | 4.1         |
|                | R Middle Temporal Gyrus                      |              |     |     |     |             |
|                | R Superior Occipital Gyrus                   |              |     |     |     |             |
| Decay (State 1)| R Lingual Gyrus                              | 295          | 24  | -78 | -9  | 3.96        |
|                | R Fusiform Gyrus                             |              |     |     |     |             |
|                | R Middle Occipital Gyrus                     |              |     |     |     |             |
|                | R Parahippocampal Gyrus                      | 218          | 28  | -44 | -12 | 4.24        |
|                | R Fusiform Gyrus                             |              |     |     |     |             |
|                | R Occipital Gyrus                            |              |     |     |     |             |
|                | L Middle Occipital Gyrus                     | 126          | -34 | -86 | 18  | 3.42        |
|                | L Superior Occipital Gyrus                   |              |     |     |     |             |
| Decay (State 2)| R Precuneus                                  | 296          | 8   | -66 | 48  | 4.11        |
|                | L Precuneus                                  |              |     |     |     |             |
| Decay (State 3)| Middle Occipital Gyrus                       | 5642         | 7.01| 46  | -62 | 6           |
|                | Middle Temporal Gyrus                        |              |     |     |     |             |
|                | Lingual Gyrus                                |              |     |     |     |             |
|                | Cuneus                                       |              |     |     |     |             |
|                | L Superior Temporal Gyrus                    |              |     |     |     |             |
|                | Inferior Temporal Gyrus                      |              |     |     |     |             |
|                | Inferior Parietal Lobule                     |              |     |     |     |             |
|                | Inferior Occipital Gyrus                     |              |     |     |     |             |
|                | R Precuneus                                  |              |     |     |     |             |
|                | R Fusiform Gyrus                             |              |     |     |     |             |
|                | L Insula                                     |              |     |     |     |             |
| Reward         | Medial Frontal Gyrus                         | 1855         | 5.24| -14 | 56  | 9           |
|                | Anterior Cingulate                           |              |     |     |     |             |
|                | L Middle Frontal Gyrus                       |              |     |     |     |             |
|                | Superior Frontal Gyrus                       |              |     |     |     |             |
|                | L Inferior Frontal Gyrus                     |              |     |     |     |             |
|                | Ventral Striatum                             |              |     |     |     |             |
| Anatomical Structure                          | MNI Coordinates | Z Score | Y Score | X Score | archetype |
|----------------------------------------------|-----------------|---------|---------|---------|-----------|
| Caudate Head                                 |                 |         |         |         |           |
| Rectal Gyrus                                 |                 |         |         |         |           |
| Subcallosal Gyrus                            |                 |         |         |         |           |
| Cingulate Gyrus                              |                 |         |         |         |           |
| Paracentral Lobule                           |                 |         |         |         |           |
| Medial Frontal Gyrus                         | 1785            | 5.04    | -10     | 12      | 27        |
| Precentral Gyrus                             |                 |         |         |         |           |
| Precuneus                                    |                 |         |         |         |           |
| Postcentral Gyrus                            |                 |         |         |         |           |
| R Superior Temporal Gyrus                    |                 |         |         |         |           |
| R Inferior Parietal Lobule                   |                 |         |         |         |           |
| R Precentral Gyrus                           |                 |         |         |         |           |
| R Postcentral Gyrus                          | 1408            | 5.55    | 58      | -36     | 15        |
| R Insula                                     |                 |         |         |         |           |
| R Transverse Temporal Gyrus                  |                 |         |         |         |           |
| R Inferior Frontal Gyrus                     |                 |         |         |         |           |
| L Precentral Gyrus                           |                 |         |         |         |           |
| L Transverse Temporal Gyrus                  |                 |         |         |         |           |
| L Postcentral Gyrus                          | 239             | 4.55    | -44     | -24     | 9         |
| L Superior Temporal Gyrus                    |                 |         |         |         |           |
| L Insula                                     |                 |         |         |         |           |
| R Dorsal Caudate                             | 176             | 4.87    | 22      | 10      | 30        |
| White Matter                                 |                 |         |         |         |           |
| R Precentral Gyrus                           | 168             | 4.35    | 40      | -22     | 42        |
| R Postcentral Gyrus                          |                 |         |         |         |           |
| R Putamen                                    | 132             | 4.39    | 32      | 0       | 3         |
| R Insula                                     |                 |         |         |         |           |
| R Medial Frontal Gyrus                        | 118             | 4.45    | 16      | 54      | 3         |
| R Middle Frontal Gyrus                        |                 |         |         |         |           |
| R Superior Frontal Gyrus                      |                 |         |         |         |           |
| L Caudate Tail                               | 86              | 4.35    | -18     | -2      | 27        |
| L Posterior Putamen                          |                 |         |         |         |           |
| L Superior Temporal Gyrus                    | 82              | 4.71    | -64     | -10     | 3         |
| L Cingulate Gyrus                            | 51              | 4.41    | -8      | 42      | 30        |
| L Precuneus                                  |                 |         |         |         |           |
| Reward (State rest)                          |                 |         |         |         |           |
| L Middle Temporal Gyrus                       |                 |         |         |         |           |
| Posterior Cingulate                          |                 |         |         |         |           |
| Cuneus                                       |                 |         |         |         |           |
| Precuneus                                    |                 |         |         |         |           |
| L Middle Occipital Gyrus                     |                 |         |         |         |           |
| Cingulate Gyrus                              |                 |         |         |         |           |
| L Superior Occipital Gyrus                   |                 |         |         |         |           |
| L Lingual Gyrus                              |                 |         |         |         |           |
| Reward (State rest)                          |                 |         |         |         |           |

**Note:** The table lists various anatomical structures along with their corresponding MNI coordinates and Z scores. The Z scores indicate the statistical significance of the activations, with higher values indicating stronger activations. The MNI (Montreal Neurological Institute) coordinates are used to locate the activations in the brain. The table also includes the archetype or template used for comparison.
| Brain Region                                      | Coordinates | p-Value | MNI Z-Value | MNI X-Value | MNI Y-Value |
|--------------------------------------------------|-------------|---------|-------------|-------------|-------------|
| L Inferior Occipital Gyrus                       |             |         |             |             |             |
| L Parahippocampal Gyrus                          |             |         |             |             |             |
| L Superior Temporal Gyrus                        |             |         |             |             |             |
| Medial Frontal Gyrus                             |             |         |             |             |             |
| Anterior Cingulate                               |             |         |             |             |             |
| Ventral Striatum                                 |             |         |             |             |             |
| Caudate Head                                     | 1321        | 5.35    | 8           | 24          | 3           |
| Rectal Gyrus                                     |             |         |             |             |             |
| L Superior Frontal Gyrus                         |             |         |             |             |             |
| R Middle Occipital Gyrus                         |             |         |             |             |             |
| R Inferior Occipital Gyrus                       |             |         |             |             |             |
| R Cuneus                                         |             |         |             |             |             |
| R Middle Temporal Gyrus                          | 995         | 4.53    | 32          | -62         | 9           |
| R Inferior Temporal Gyrus                        |             |         |             |             |             |
| R Posterior Cingulate                            |             |         |             |             |             |
| R Lingual Gyrus                                  |             |         |             |             |             |
| L Cingulate Gyrus                                | 458         | 4.76    | -22         | -26         | 36          |
| White Matter                                     |             |         |             |             |             |
| R Precentral Gyrus                               | 333         | 4.25    | 30          | -38         | 72          |
| R Postcentral Gyrus                              |             |         |             |             |             |
| R Hippocampus                                    | 250         | 4.93    | 32          | -34         | 0           |
| R Parahippocampal Gyrus                          |             |         |             |             |             |
| L Parahippocampal Gyrus                          |             |         |             |             |             |
| L Fusiform Gyrus                                 |             |         |             |             |             |
| L Hippocampus                                    | 250         | 3.88    | -38         | -52         | -12         |
| L Middle Occipital Gyrus                         |             |         |             |             |             |
| L Middle Temporal Gyrus                          |             |         |             |             |             |
| R Precentral Gyrus                               | 207         | 4.3     | 56          | 0           | 18          |
| R Insula                                         |             |         |             |             |             |
| R Postcentral Gyrus                              |             |         |             |             |             |
| L Medial Frontal Gyrus                           | 181         | 4       | -16         | 42          | 24          |
| L Superior Frontal Gyrus                         |             |         |             |             |             |
**Supplementary Table 2.** Additional PFC region of interest state information analysis.

Regression results are shown for current state (state 1) and the future state (state 3) by correct repetition interaction. Results from the OFC-VMPFC are as shown in **Figure 3**. Number of included participants per ROI for the future by repetition analysis depends on exclusions for below-zero classification of state 1 and state 3. antHipp = anterior hippocampus; postHipp = posterior hippocampus.

| region    | current \(n = 31\) | CI         | \(t\)-stat | \(p\)-value | \(n\) | fut*rep | CI         | \(t\)-stat | \(p\)-value |
|-----------|----------------------|------------|-------------|-------------|-------|---------|------------|-------------|-------------|
| OFC       | 0.092                | 0.05 - 0.13| 4.48        | 0.0001      | 29    | 0.051   | 0.01 - 0.09| 2.53        | 0.0172      |
| R antOFC  | -0.001               | -0.04 - 0.03| -0.08       | 0.9384      | 22    | 0.011   | -0.05 - 0.07| 0.37        | 0.7176      |
| L antOFC  | 0.014                | -0.03 - 0.05| 0.70        | 0.4880      | 24    | 0.033   | -0.01 - 0.08| 1.46        | 0.1570      |
| pgACC     | 0.022                | -0.01 - 0.06| 1.27        | 0.2152      | 24    | 0.034   | -0.01 - 0.08| 1.57        | 0.1311      |
| dmPFC     | 0.138                | 0.1 - 0.17 | 7.81        | 0.0000      | 30    | 0.017   | -0.02 - 0.05| 0.90        | 0.3749      |
| fpPFC     | 0.097                | 0.06 - 0.13| 5.63        | 0.0000      | 30    | 0.001   | -0.04 - 0.04| 0.08        | 0.9371      |
| R latPFC a| 0.038                | 0.00 - 0.08| 1.95        | 0.0610      | 25    | 0.002   | -0.05 - 0.05| 0.08        | 0.9351      |
| L latPFC a| 0.021                | -0.01 - 0.06| 1.22        | 0.2324      | 21    | -0.007  | -0.05 - 0.04| -0.31       | 0.7588      |
| R latOFC  | 0.084                | 0.06 - 0.11| 7.84        | 0.0000      | 29    | -0.004  | -0.03 - 0.02| -0.30       | 0.7701      |
| L latOFC  | 0.038                | 0.01 - 0.07| 2.57        | 0.0153      | 27    | -0.008  | -0.04 - 0.03| -0.47       | 0.6439      |
| ACC a     | -0.007               | -0.04 - 0.03| -0.42       | 0.6808      | 23    | 0.014   | -0.05 - 0.07| 0.47        | 0.6401      |
| sgACC     | 0.017                | -0.01 - 0.05| 1.13        | 0.2688      | 27    | 0.008   | -0.03 - 0.04| 0.43        | 0.6739      |
| R latPFC b| 0.121                | 0.08 - 0.16| 5.78        | 0.0000      | 28    | -0.007  | -0.04 - 0.03| -0.38       | 0.7100      |
| L latPFC b| 0.093                | 0.04 - 0.14| 3.90        | 0.0005      | 29    | -0.026  | -0.09 - 0.04| -0.86       | 0.3952      |
| R infPFC  | 0.052                | 0.0 - 0.1  | 2.16        | 0.0393      | 29    | -0.003  | -0.06 - 0.06| -0.11       | 0.9134      |
| L infPFC  | 0.061                | 0.01 - 0.11| 2.57        | 0.0153      | 28    | 0.052   | -0.02 - 0.12| 1.45        | 0.1573      |
| ACC b     | 0.020                | -0.02 - 0.06| 0.98        | 0.3357      | 29    | -0.018  | -0.07 - 0.03| -0.76       | 0.4509      |
| ACC c     | 0.020                | -0.04 - 0.08| 0.70        | 0.4889      | 25    | 0.008   | -0.05 - 0.07| 0.27        | 0.7931      |
| R DLPFC a | 0.050                | 0.0 - 0.1  | 2.04        | 0.0507      | 27    | 0.050   | -0.01 - 0.11| 1.79        | 0.0858      |
| L DLPFC a | 0.025                | -0.01 - 0.06| 1.50        | 0.1451      | 25    | 0.027   | -0.06 - 0.11| 0.67        | 0.5115      |
| R DLPFC b | 0.027                | 0.0 - 0.06 | 1.88        | 0.0694      | 24    | 0.035   | -0.01 - 0.08| 1.58        | 0.1278      |
| L DLPFC b | 0.051                | 0.02 - 0.08| 3.33        | 0.0023      | 24    | 0.047   | 0.02 - 0.08 | 3.10        | 0.0051      |
| R DLPFC c | 0.021                | -0.07 - 0.02| -1.00       | 0.3232      | 26    | 0.025   | -0.04 - 0.09| 0.79        | 0.4377      |
| L DLPFC c | 0.076                | 0.02 - 0.13| 2.66        | 0.0125      | 21    | 0.041   | -0.03 - 0.11| 1.28        | 0.2164      |
| R DLPFC d | 0.056                | -0.02 - 0.13| 1.59        | 0.1220      | 25    | -0.030  | -0.13 - 0.07| -0.64       | 0.5267      |
| L DLPFC d | 0.062                | 0.01 - 0.11| 2.62        | 0.0136      | 27    | 0.070   | -0.01 - 0.15| 1.87        | 0.0725      |
|       | 0.031  | 0.00 0.06 | 2.40  | 0.0228 | 30 | 0.001 | -0.03 0.03 | 0.08 | 0.9370 |
|-------|--------|-----------|-------|--------|----|--------|-----------|------|--------|
| R antHipp | 0.012 | -0.04 0.07 | 0.45  | 0.6553 | 26 | -0.001 | -0.06 0.05 | -0.06 | 0.9564 |
| L antHipp | 0.038 | -0.00 0.08 | 1.87  | 0.0717 | 27 | -0.026 | -0.09 0.04 | -0.84 | 0.4091 |
| R postHipp | -0.013 | -0.06 0.04 | -0.55 | 0.5842 | 18 | -0.012 | -0.07 0.05 | -0.43 | 0.6745 |
Supplementary Table 3. Summary of OFC-VMPFC Late > Early repetition PPI analysis effects correlated with the OFC-VMPFC future state by correct repetition effect. State 1 was the timepoint of interest; later states were analyzed in control models. The PPI contrast of late versus early repetitions (3 and 4 versus 1 and 2) or the reverse was regressed against individual differences in the OFC-VMPFC future state by correct repetition regression coefficient (as depicted in Figure 3b). No main effects of the contrasts of late > early repetitions or early > late repetitions were found. Clusters of activity exceed whole-brain p < 0.05 FWE-corrected (cluster-forming threshold p < 0.005). Within each cluster, the first 10 regions are listed that include >= 10 voxels in a cluster.

| Contrast                      | Regions                                      | Cluster size | x    | y    | z    | Peak z stat | P-value |
|-------------------------------|----------------------------------------------|--------------|------|------|------|-------------|---------|
| State 1 OFC PPI Late > Early future*repcorr | Bilat. Middle Frontal Gyrus                  | 5712         | 42   | 12   | 39   | 4.93        | 0       |
|                               | Bilat. Inferior Frontal Gyrus                |              |      |      |      |             |         |
|                               | Bilat. Superior Frontal Gyrus                |              |      |      |      |             |         |
|                               | Medial Frontal Gyrus                         |              |      |      |      |             |         |
|                               | Cingulate Gyrus                              |              |      |      |      |             |         |
|                               | Precentral Gyrus                             |              |      |      |      |             |         |
|                               | L Putamen                                    |              |      |      |      |             |         |
|                               | Anterior Insula                              |              |      |      |      |             |         |
|                               | L Caudate Body                               |              |      |      |      |             |         |
|                               | R Putamen                                    | 275          | 24   | -2   | 0    | 4.54        | 0.006   |
|                               | R Pallidum                                   |              |      |      |      |             |         |
|                               | R Insula                                     |              |      |      |      |             |         |
|                               | R Amygdala                                   |              |      |      |      |             |         |
|                               | Midbrain                                     | 285          | -6   | -10  | -9   | 4.47        | 0.005   |
|                               | Thalamus                                     |              |      |      |      |             |         |
|                               | R Hippocampus                                |              |      |      |      |             |         |
|                               | L Postcentral Gyrus                          | 282          | -50  | -20  | 24   | 3.99        | 0.005   |
|                               | L Supramarginal Gyrus                        |              |      |      |      |             |         |
|                               | L Inferior Parietal Lobule                   |              |      |      |      |             |         |

State 1 OFC PPI Early > Late future*repcorr

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| State 2       | OFC PPI | Late > Early future\*repcorr | - |
|--------------|---------|-------------------------------|---|
| State 2      | OFC PPI | Early > Late future\*repcorr  | - |
| State 3      | OFC PPI | Late > Early future\*repcorr  | - |
| State 3      | OFC PPI | Early > Late future\*repcorr  | - |
| State reward | OFC PPI | Late > Early future\*repcorr  | - |
| State reward | OFC PPI | Early > Late future\*repcorr  | - |