Learning in visual regions as support for the bias in future value-driven choice

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10 Abstract

Learning biases decision-making towards higher expected outcomes. Cognitive theories describe this through the tracking of value and outcome evaluations within striatum and prefrontal cortex. Decisions, however, first require processing of sensory input and to-date far less is known about the learning perception interplay. This fMRI study (N=49), relates visual BOLD responses to value-beliefs during choice, and, signed prediction errors after outcomes. To understand these relationships, which co-occurred in striatum, we next evaluated relevance with the prediction of future value-based choices, using a separate transfer-phase with learning already established. We decoded choice outcomes with a 69% accuracy given unseen trials with a machine learning algorithm that combined trial-by-trial BOLD from prefrontal, striatal, and visual regions. Importantly, this classification of value-driven choice outcomes again showed an important role for visual activity. These results raise the intriguing possibility that value learning in visual cortex is supportive for the striatal bias towards valued options.

Keywords: Reinforcement learning, perceptual learning, decoding, Bayesian hierarchical modelling, random forest machine learning

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$_{\scriptscriptstyle 25}$ Introduction

In decision-making, our value beliefs bias future choices. This bias is shaped by the outcomes of similar decisions made in the past where the action, or stimulus chosen, becomes associated with a positive or negative outcome ('value beliefs'). The evaluation of value after an outcome, or the comparison of value in decisions, is traditionally associated with activity in the prefrontal cortex and striatum¹⁻⁷.

To underset the bias in action selection midbrain dopamine neurons are thought to send a teaching signal towards the striatum and prefrontal cortex after an outcome $^{8-10}$. In the striatum, future actions are facilitated 31 by bursts in dopamine after positive outcomes or discouraged by dopamine dips after negative outcomes. The 32 dorsal and ventral parts of the striatum are known to receive differential, but also overlapping, inputs from midbrain neurons^{7,11}. Ventral and dorsal striatum have also been ascribed a differential role during learning 34 by reinforcement learning theories. Here, the ventral parts of the striatum are involved with the prediction of future outcomes through the processing of prediction errors, whereas the dorsal striatum uses the same information to maintain action values as a way to bias future actions towards the most favored option^{4,12,13}. 37 Intriguingly, however, before many of these value-based computations can take place, stimuli first have to be parsed from the natural world, an environment where most reward predicting events are perceptually complex. This suggests that sensory processing might be an important integral part of optimized value-based decision-making.

Here, we investigate whether choice outcomes can modulate the early sensory processing of perceptually complex stimuli to help bias future decisions. Recent neurophysiological studies find visually responsive neurons in the tail of the caudate nucleus, which is part of the dorsal striatum^{14,15}. These neurons encode and differentiate stable reward values of visual objects to facilitate eye movements towards the most valued target, while at the same time inhibiting a movement towards the lesser valued object¹⁶. Critically, differential modulations are also observed in the primary visual cortex where stronger cortical responses are seen for objects with higher values^{17,18}, which is consistent with the response of visual neurons in the caudate. As visual cortex is densely connected to the striatum^{19,20}, prioritized visual processing of high-value stimuli could aid the integration of information regarding the most-valued choice in the striatum^{21–24}. To understand these visual-striatal interactions, we focus on a more detailed parsing of the underlying computations.

Specifically, we explored two questions by reanalyzing fMRI data from a probabilistic reinforcement learning task using faces as visual stimuli²⁵ (Figure 1a). First, we focus on the interplay between learning and visual activity in the fusiform face area (FFA) and occipital cortex (OC). Here, with the use of a Bayesian hierarchical reinforcement learning model (Figure 1b) we outline how trial-by-trial estimates of action values (Q-value) and reward prediction errors (RPE) relate to the BOLD response of visual regions and the striatum^{26,27} (Figure 1c). Second, we analyze data from a follow-up transfer phase, where the learning of value was already established. In our analysis, the importance of visual brain activity in the prediction, or decoding, of future value-based decisions is evaluated by using a supervised Random Forest (RF) machine learning algorithm^{28,29}. Specifically, transfer phase single-trial BOLD estimates from anatomically defined visual, prefrontal, and subcortical regions are combined by RF to predict, or decode, choice outcomes for unseen trials. We focus on classification accuracy, and the relative importance of each brain region in the correct classification of future value-based decisions.

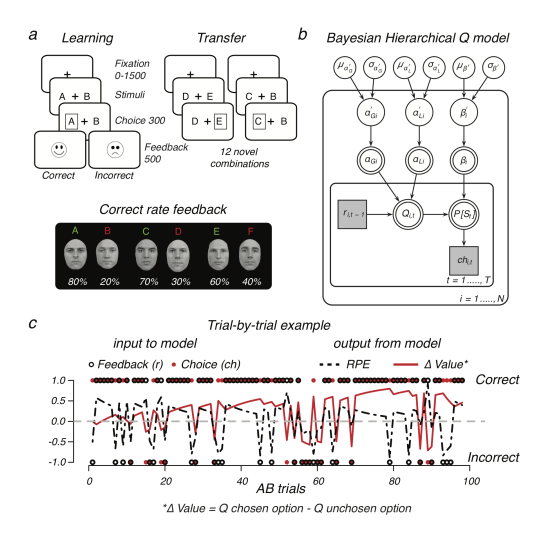


Figure 1: **Design and Model. a**) Reinforcement learning task using faces. During learning, two faces were presented on each trial, and participants learned to select the optimal face identity (A, C, E) through probabilistic feedback (% of correct is shown beneath each stimulus). The learning-phase contained three face pairs (AB, CD, ED) for which feedback was given. In a follow-up transfer phase these faces were rearranged into 12 novel combinations to asses learning. These trials were identical to learning trials, with the exception of feedback. b) Graphical Q-learning model with hierarchical Bayesian parameter estimation. The model consists of an outer subject $(i = 1, \ldots, N)$, and an inner trial plane $(t = 1, \ldots, T)$. Nodes represent variables of interest. Arrows are used to indicate dependencies between variables. Double borders indicate deterministic variables. Continuous variables are denoted with circular nodes, and discrete with square nodes. Observed variables are shaded in grey (see methods for details about the fitting procedure). c) Illustration of the observed trial-by-trial input (i.e., the choice made, and feedback received), and output (i.e., Q for the chosen and unchosen stimulus, Q and RPE) of the model given the estimated variability in learning rates from either positive Q or negative Q or negative Q for the tendency to exploit Q higher values Q in the endency of exploit Q higher values Q in the endency of exploit Q higher values Q in the endency of exploit Q higher values Q in the endency of exploit Q higher values Q in the exploit Q higher values Q in the exploit Q higher values Q in the explosion of the exploit Q higher values Q in the explosion of the explosion

64 Results

- To understand how value learning relates to the activity pattern in perceptual regions we reanalyzed the
- behavioral and fMRI recordings of a recent study²⁵. In this study, BOLD signals were recorded while
- participants performed a reinforcement learning task using male or female faces, and a stop-signal task (which

was discussed in 25). The fusiform face area (FFA) was localized using a separate experimental run. 49 young adults (25 male; mean age = 22 years; range 19-29 years, 43 analyzed, see Methods) participated in the study. As shown in Figure 1a, in the reinforcement learning task participants learned to select among choices with different probabilities of reinforcement (i.e., AB 80:20, CD 70:30, and EF 60:40). A subsequent transfer phase, where feedback was omitted, required participants to select the optimal option among novel pair combinations of the faces that were used during the learning phase (Figure 1a).

74 Model and Behavior

In the learning phase, subjects reliably learned to choose the most optimal face option in all pairs. For each pair the probability of choosing the better option was above chance (p's < .001), and the effect of learning decreased from AB (80:20) and CD (70:30) to the most uncertain EF (60:40) pair (F(2, 84) = 13.74, p < .0001).

At the end of learning, value beliefs differentiating the optimal (A, C, E) from the sub-optimal (B, D, F) action were very distinct for the AB and CD face pairs but decreased with uncertainty (F(2, 84) = 39.70, p < 0.0001, Figure 2a). Value beliefs were estimated using the individual subject parameters of the Q-learning model that best captured the observed data (Figure 2b-e; reproduced from 25 to show performance).

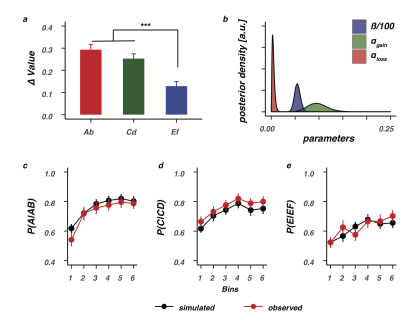


Figure 2: Value differentiation and model performance. a) Value differentiation (Δ Value) for the selection of the optimal (A,C,E) stimuli over the suboptimal (B,D,F) stimuli decreased as a function of feedback reliability, and was smallest for the most uncertain EF stimuli. *** = p < 0.0001, Bonferroni corrected. b) Group-level posteriors for all Q-learning parameters. The bottom row shows model performance, where data was simulated with the estimated individual subject parameters and evaluated against the observed data for the AB (c), CD (d), or EF (e) pairs. Bins contain +/- 16 trials. Error bars represent standard error of the mean (SEM).

82 BOLD is modulated by reliable value differences between faces in striatal and visual regions

For each pair of faces presented during the learning phase (AB, CD, EF) we asked how the BOLD signal time-course in striatal and visual regions relates to trial-by-trial value beliefs about the two faces presented as a choice. First, as a reference, we focused on the activity pattern of three striatal regions. Results showed BOLD responses in dorsal (caudate, putamen) but not ventral (accumbens) striatum to be differentially modulated by the estimated value beliefs of the chosen face (Q_{chosen}), in comparison to value beliefs about the face that was not chosen ($Q_{unchosen}$). Thus, BOLD responses in the dorsal striatum were modulated more strongly by value beliefs about the chosen stimulus (Q_{chosen} ; Figure 3a bottom row). Critically, this differential modulation was only observed with the presentation of AB faces where value differences were most distinct because of the reliable feedback scheme. Next, we evaluated the relationship between value and BOLD in the FFA, and OC. Again, only with the presentation of the AB face option, trial-by-trial BOLD fluctuations were differentially modulated by values of the chosen versus not chosen face option (Figure 3b bottom row). These evaluations highlight how the BOLD response in striatal and perceptual regions is especially sensitive to values of the (to-be) chosen stimulus when belief representations are stable and distinct.

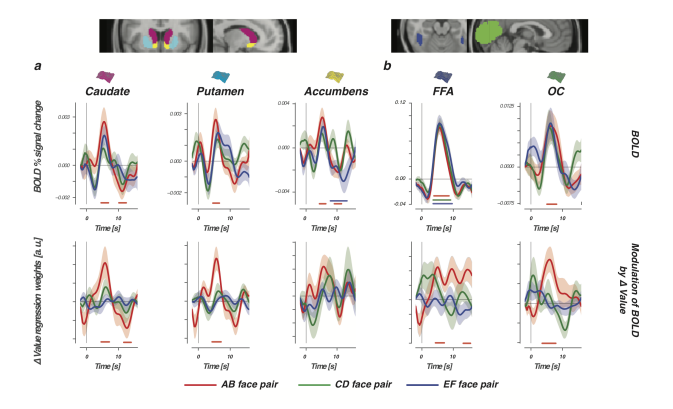


Figure 3: **BOLD** and the modulation of Δ Value in the learning phase. Top row shows the BOLD signal time course, time-locked to presentations of AB (80:20, red lines), CD (70:30, green lines), and EF (60:40, blue lines) face pairs, for three striatal regions (a) and two perceptual regions (b). Bottom row displays differential modulation by value (Δ Value = modulation Q_{chosen} - modulation $Q_{unchosen}$). Horizontal lines show the interval in which modulation was significantly stronger for Q_{chosen} . With the presentation of AB faces, BOLD responses in the dorsal striatum (caudate and putamen) and visual regions (FFA and OC) were modulated more by values of the chosen stimulus when compared to values of the unchosen stimulus. Differential AB value modulation was not significant in the ventral striatum (i.e., accumbens). Nor did we observe any differential value modulations with the presentation of the more uncertain CD and EF pairs. Confidence intervals were estimated using bootstrap analysis across participants (n = 1000), where the shaded region represents the standard error of the mean across participants (bootstrapped 68% confidence interval).

Reward prediction errors in striatal and visual regions

Our findings so far described relationships between BOLD and value time-locked to the moment of stimulus presentation – i.e., when a choice is requested. Learning occurs when an outcome is different from what was expected. We therefore next focused on modulations of the BOLD response when participants received feedback. Learning modulations were explored by asking how trial-by-trial BOLD responses in perceptual and striatal regions relate to either signed (outcome was better or worse than expected) or unsigned (magnitude of expected violation) reward prediction errors³⁰. Consistent with the literature, BOLD responses in all striatal regions were modulated by signed RPEs, with larger responses after positive RPEs or smaller responses after negative RPEs (Figure 4a bottom row). Activity in the accumbens (ventral striatum) was additionally tied to unsigned RPEs in the tail of the BOLD time-course, with larger violations (either positive or negative) tied to smaller dips. Consistently, estimated BOLD responses in both visual regions were modulated by the

signed RPE, and once more mirrored the striatal modulations with stronger positive RPEs eliciting stronger BOLD responses (Figure 4b bottom row). FFA BOLD responses were additionally modulated by unsigned RPEs. However, in contrast to the relationship found between unsigned RPEs and the accumbens, the FFA modulation was positive and co-occurred with the modulation of the signed RPE. That is, bigger violations and more positive outcomes each elicited a stronger response in the FFA.

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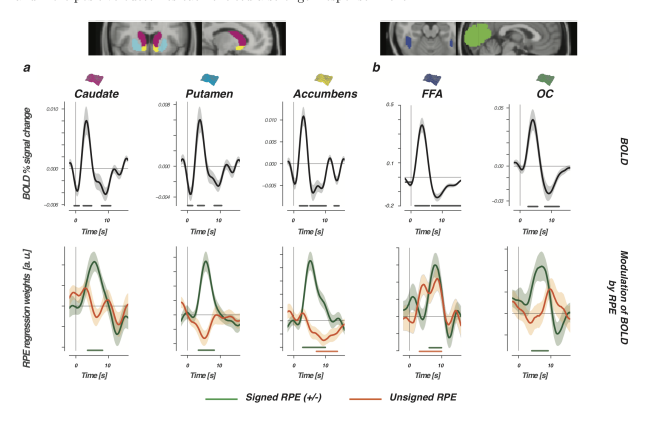


Figure 4: Reward prediction errors modulate BOLD in striatal and visual regions. The top row shows the FIR-estimated BOLD signal time-course, which was time-locked to the presentation of choice feedback and evaluated for three striatal regions (a) and two perceptual regions (b). Bottom row displays modulations of the estimated BOLD time-course by singed (green lines), or unsigned (orange lines) RPEs. The horizontal lines represent the interval in which signed or unsigned RPEs contributed significantly to the modulation of BOLD in the multiple regression. Note that both variables were always evaluated simultaneously in one GLM.

Can past learning in visual regions support the prediction of future value-based decisions?

Stable value representations and reward prediction errors both modulated the activity of visual and striatal regions. These modulations in the striatum are described to bias future actions towards the most favored option (the dorsal striatum), or to predict future reward outcomes (the ventral striatum). To better understand the value and RPE modulations observed in visual regions, we next assessed the importance of these visual regions alongside the striatum in the correct classification (decoding) of future value-driven choice outcomes. Here, activity of prefrontal regions was added to the importance evaluation based on our previous work with

this data in the transfer phase²⁵ (please see supplementary Figures 1&2 for the evaluation of these regions during learning).

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In the transfer phase, participants had to make a value-driven choice based on what was learned before, i.e., during the learning phase. To specify the relevance of visual regions in the resolve of value-driven choice outcomes, in the transfer phase, a random forest (RF) classifier was used^{28,29} (Please see Figure 5a-c for the procedure). The RF classifier relies on an ensemble of decision trees as base learners, where the prediction of each trial outcome is obtained by a majority vote that combines the prediction of all decision trees (Figure 6a). To achieve controlled variation, each decision tree is trained on a random subset of the variables (i.e. subset of columns shown in Figure 5a), and a bootstrapped sample of data points (i.e. trials). Importantly, we ensured that the forest was not simply learning the proportion of optimal choices in the transfer phase by training all models on balanced draws from the training set with equal numbers of optimal and sub-optimal choices.

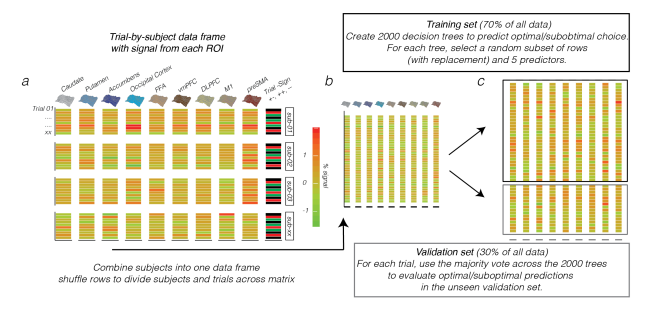


Figure 5: Random Forest input and data-structure. a) Trial-by-subject data matrix with the % signal change drawn for each choice trial in the transfer-phase (rows) from 9 a-priori defined regions of interest (columns). In addition to the ROI data, the matrix contained a column with the identity of participants (sub-01, etc) and Trial Sign, which specified a choice between two positives (+/+; AC, AE, CE), negatives (-/-, BD, BF, DF), or between a negative and positive option (+/-, e.g., AD, CF, etc) given the feedback scheme in the learning-phase. b) The individual subject data frames were then combined into one matrix, in which the rows were subsequently shuffled to randomly distribute trials and subjects across the rows. c) This matrix was then divided into a training set (2/3 of the data) for the creation of 2000 decision trees of which the majority vote on each trial is then used to evaluate the predictive accuracy of optimal/suboptimal choices in a separate validation set (1/3 of the data).

Evaluation of all participants resulted in a classification accuracy of 65% (AUC = 0.75) using the trial-by-trial BOLD estimates from the ROIs and increased to 69% with the evaluation of the good learners (AUC = 76; N = 34, criteria: accuracy > 60% across all three learning pairs). Hence, in 65 (all participants) or 69 (good learners) out of 100 trials the forest correctly classified whether participants would pick the option with the highest value (optimal choice) or not (sub-optimal choice) in the validation set. RF predictions were substantially lower when labels of the validation set were randomly shuffled (accuracy: all participants= 52%;

good learners = 56%).

The improvement of accuracy with the evaluation of only the good learners is remarkable because the classifier 139 was given less data to learn the correct labelling (fewer subjects/trials) and implied that the 2000 decision trees were picking up information related to past learning. Further support for this important observation 141 was found by asking how the uncertainty of each prediction (defined as the proportion of agreement in the 142 predicted outcome among the 2000 trees for each trial) relates to the difference in value beliefs (Δ Value) about the two options presented on each trial (computed using the end $Q_{beliefs}$ of participants at the end 144 of learning about face A-to-E). As plotted in Figure 6c, the uncertainty in predicting that a trial choice 145 outcome is optimal – defined as the proportion of disagreement among the 2000 decision trees - decreased with larger belief differences in the assigned values (please see supplementary Figure 3 for the evaluation of 147 all participants). 148

Besides providing insights into how BOLD responses in the transfer-phase contribute to predict value-driven 149 choice outcomes (i.e., whether participants would choose the option with the highest value given past learning) the RF algorithm additionally outputs a hierarchy, thereby ranking the contribution of each region in the 151 achieved classification accuracy. Figure 6d shows the ranking of all ROIs for good learners where the model 152 had the highest predictive accuracy. First, regions in the dorsal striatum were most important, which aligned 153 well with both the literature and the BOLD modulations we found by Δ Value and RPE during the learning 154 phase. These regions were next followed by the preSMA. Evaluation of this region during the learning phase 155 showed no modulations by Δ Value or RPE on BOLD (supplementary Figure 1&2). Nevertheless, this region 156 is typically associated with choice difficulty/conflict and might be essential in the resolve of a choice when 157 value differences are small. Remarkably, the third region in this hierarchy was the FFA. In a task where 158 participants pick the most valued face based on past learning, this ranking of the FFA just above the vmPFC 159 and accumbens (ventral striatum) implies that the Δ Value and RPE modulations of BOLD observed during 160 learning could function to strengthen the recognition of valuable features. Note, however, that with the 161 evaluation of all participants – including some who were less good in learning – the ranking of both the FFA 162 and vmPFC was much lower (please see supplementary Figure 3b), which might be caused by more noise 163 across the group in learning. We will return to this point in the discussion. 164

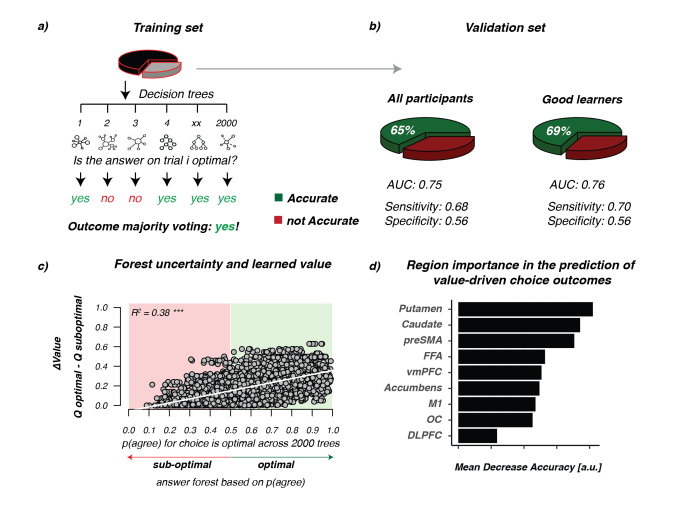


Figure 6: Random Forest performance and importance ranking. The prediction of value-driven choice outcomes in the transfer phase using trial-by-trial BOLD responses from striatal, perceptual, and prefrontal cortex regions. a) Overview of the Random Forest approach where the training-set is used to predict choice outcomes for each trial by using the majority vote of 2000 different decision trees. Each tree is build using a different set, or sample, of trials and predictors from the training set. The forest is trained on a training set sampled from all participants (N=43), or only 'the good learners' (N=34). b) Shows the classification, or decoding, accuracy (green) given the separate unseen validation sets, for all participants and good learners. c) Plotted relationship between forest uncertainty (i.e., proportion of agreement across 2000 trees), on each prediction/trial (x-axis) and Δ Value (y-axis) for the model with the highest accuracy (i.e., the good learners). Δ Value was computed for each trial in the transfer phase by using the end beliefs (Q) that participants had about each stimulus (A-to-F) at the end of the learning phase. Forest uncertainty is defined as the proportion of trees saying 'yes! the choice on this trial was optimal'. When this ratio is bellow 0.5 the forest will predict 'no' (sub-optimal), otherwise the prediction is 'yes! the choice on this trial was optimal' (optimal). R^2 =adjusted R^2 . Note that, the same pattern was found for all participants ($R^2 = 0.41^{***}$, please see supplementary Figure 3). d) Plotted ranking of the ROI's in their contribution to the predictive accuracy of the best performing model (i.e., good learners).

Discussion

This study provides novel insights into how reinforcements modulate visual activity and specifies its potential in the prediction of future value-driven choice outcomes. First, by focusing on how participants learn, we find 167 BOLD in visual regions to change with trial-by-trial adaptations in value beliefs about the faces presented, 168 and then to be subsequently scaled by the signed RPE after feedback. Next, the relevance of these observed value and feedback modulations was sought by exploring the prediction of future value-driven choice outcomes 170 in a follow-up transfer phase where feedback was omitted. Our machine learning algorithm here shows a 171 classification accuracy of 69% for participants who were efficient in learning by combining trial-by-trial BOLD estimates from perceptual, striatal, and prefrontal regions. The evaluation of region importance in these 173 predictions ranked the FFA just after the dorsal striatum and the preSMA, thereby showing an important 174 role for visual regions in the prediction of future value-driven choice outcomes in a phase where learning is established. 176

In a choice between two faces, BOLD responses in both the dorsal striatum and perceptual regions were
affected more by values of the chosen face, relative to the unchosen face. Across three levels of uncertainty,
we only observed the differential modulation of value on BOLD when belief representations were stable.
This specificity aligns with neuronal responses to perceptual stimuli in the caudate tail¹⁶, visual cortex^{31,32},
and imaging work across sensory modalities^{17,18,33–35}, where it fuels theories in which the learning of stable
reward expectations can develop to modulate, or sharpen, the representation of sensory information critical
for perceptual decision making^{31,35}.

After a choice was made, feedback modulations of signed ('valence') and unsigned ('surprise') RPEs³⁰ were 184 evaluated on BOLD responses, by using an orthogonal design where the unsigned and signed RPE compete to 185 explain BOLD variances. Both visual and striatal regions respond to prediction errors³⁶. In the striatum both 186 valence and surprise are thought to optimize future action selection in the dorsal striatum, or the prediction of 187 future rewards in the ventral striatum. In perceptual regions, a mismatch between the expected and received 188 outcome is often explained as surprise where a boost in attention or salience changes the representation of 189 an image without a representation of value per se. We found positive modulatory effects of signed RPEs 190 in all striatal regions, as well as, in the FFA and OC. Concurrently, modulations of unsigned RPEs were 191 only observed in the accumbens (ventral striatum) and FFA, where notably the direction of modulation 192 was reversed. We speculate that this contrast arises from the differential role of the regions. In the FFA, 193 specialized and dedicated information processing is essential to quickly recognize valuable face features. Complementary boosts of surprise and valence here could prioritize attention towards the most rewarding 195 face feature to strengthen the reward association in memory, or help speed up future recognition^{37–39}. In 196 the accumbens, boosted effects of positive valence on BOLD were dampened by larger mismatches. Large mismatches in what was expected are rare in stable environments. We therefore reason that in the accumbens 198 the contrast between valance and surprise could function as a scale to refine learning, eventually leading to 199 more reliable predictions of future rewards.

Whereas BOLD in the ventral striatum was shaped by both signed and unsigned RPEs, the dorsal striatum was sensitive to differential value up-to a choice and signed RPEs with the presentation of feedback ^{34,40–43}.

The concurrent modulation of differential value in the primary motor cortex (please see M1 in supplementary Figure 1) associates the dorsal striatum with the integration of sensory information ^{16,44–46}, where increased visual cortex BOLD responses to faces with the highest value could potentially help bias the outcome of a

value-driven choice.

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We explored this line of reasoning with the prediction of value-driven choice outcomes in a follow-up transfer 207 phase after leaning. In recent years, machine learning approaches have become increasingly important in 208 neuroscience^{47–50}, where the ease of interpretation has often motivated a choice for linear methods above 209 non-linear methods^{49,51}. Despite the latter being less constrained and able to reach a better classification 210 accuracy by capturing non-arbitrary, or unexpected relationships⁵². Value-driven choices after a phase of initial learning are influenced by the consistency of past learning, memory updating, and attention. All 212 of these processes are affected by both linear and non-linear neurotransmitter modulations 53-56. Our RF 213 approach was unconstrained by linearity with classification accuracies well above chance and improved with 214 the evaluation of only the good learners; despite substantial decreases in data given to the algorithm to 215 learn the correct labelling. Critically, we additionally found that the uncertainty of trial-by-trial predictions 216 made by RF is tied to the differentiability of value beliefs – an index that we could compute for the novel pair combination in the transfer phase by using the value (Q) beliefs that participants had about each face 218 at the end of learning. These results showcase how trial-by-trial BOLD fluctuations in striatal, prefrontal, 219 and sensory regions can be combined by machine learning, or decoding, algorithms to reliably predict the 220 outcome of a value-driven choice. Where we refine the interpretation of non-linear predictions by combining 221 the RF output with cognitive computational modelling. With this combiantion we essentially show how the 222 uncertainty of RF predictions is tied to value beliefs acquired with learning in the past. 223

An important evaluation intended with our machine learning approach was the ranking of regions by their contribution to the predictive (decoding) accuracy in the transfer phase. After the observed modulations of BOLD in the learning phase this explorative analysis sought the relevance of learning-BOLD relationships in the resolve of future choices. Here, the ranking made by RF first identified signals from the dorsal striatum (putamen and caudate) as most important followed by the preSMA, and then most notably, visual regions. That is, when the quality of leaning was high across participants, FFA ranked just above traditional regions such as the vmPFC and the accumbens^{2,5,6,57}. Notably, FFA was replaced by OC in ranking with the evaluation of all participants (please see supplementary Figure 3b). This difference could occur because the quality of learning was more variable across all participants, or because RF predictions based on the 232 heterogeneous data from all participants were less accurate. In general, the shift in ranking implies that when learning is less consistent choice outcomes are better predicted by fluctuations in OC - perhaps with the identification of rewarding low-level features. With better or more consistent learning, however, participants should increasingly rely on memory and specialized visual areas. Thus, search for specific face features associated with high value by recruiting the FFA in the visual ventral stream. Consistent with this reasoning recent neuronal recordings show rapid visual processing of category-specific value cues in the ventral visual stream. These specific value cues are only seen for well-learned reward categories, and critically, precede the processing of value in prefrontal cortex⁵⁹.

We note that although BOLD fluctuations in the preSMA ranked second in the prediction of value-driven 241 choice outcomes, no reliable modulations of BOLD were observed by either differential value or RPEs in the learning phase. The preSMA is densely connected to the dorsal striatum and consistently associated with 243 action-reward learning⁶⁰, or choice difficulty⁶¹. The lack of associations in this study might result from our 244 noisier estimates of the BOLD response that is typical for regions in the prefrontal cortex 62,63, the anatomical masks selected, or smaller variability across trials in the learning phase (i.e., 3 pairs in learning-phase vs 15 246 pairs in transfer-phase). Nevertheless, the importance indicated by RF, combined with our previous analysis of this transfer phase data²⁵, implies an important role for the preSMA in the resolve of value-driven choices in concert with the striatum. More research with optimized sequences to estimate BOLD in PFC is required to clarify the link between learning and transfer.

To summarize, we find an important role for perceptual regions in the prediction of future value-driven choice outcomes, which coincides with the sensitivity of BOLD in visual regions to differential value and signed feedback. These findings imply visual regions to learn prioritize high value features with the integration of feedback, to support and fasten, optimal response selection via the dorsal striatum in future encounters.

$_{255}$ Methods

256 Participants

All participants had normal or corrected-to-normal vision and provided written consent before the scanning session, in accordance with the declaration of Helsinki. The ethics committee of the University of Amsterdam approved the experiment, and all procedures were in accordance with relevant laws and institutional guidelines. In total, six participants were excluded from all analyses due to movement (2), incomplete sessions (3), or misunderstanding of task instructions (1).

262 Reinforcement learning task

Full details of the reinforcement learning task are provided in 25. In brief, the task consisted of two phases 263 (Figure 1a). In the first learning phase, three male or female face pairs (AB, CD, EF) were presented in a random order, and participants learned to select the most optimal face (A, C, E) in each pair solely through 265 probabilistic feedback ('correct': happy smiley, 'incorrect': sad smiley). Choosing face-A lead to 'correct' on 266 80% of the trials, whereas a choice for face-B only lead to the feedback 'correct' for 20% of the trials. Other 267 ratios for 'correct' were 70:30 (CD) and 60:40 (EF). Participants were not informed about the complementary 268 relationship in pairs. All trials started with a jitter interval where only a white fixation cross was presented 269 and had a duration of 0, 500, 1000 or 1500ms to obtain an interpolated temporal resolution of 500ms. Two 270 faces were then shown left and right of the fixation-cross and remained on screen up to response, or trial end 271 (4000ms). If a response was given on time, a white box surrounding the chosen face was then shown (300ms) 272 and followed (interval 0-450ms) by feedback (500ms). Omissions were followed by the text 'miss' (2000ms). 273 The transfer-phase contained the three face-pairs from the learning phase, and 12 novel combinations, in which 274 participants had to select which item they thought had been more rewarding during learning. Transfer-phase trials were identical to the learning phase, with the exception that no feedback was provided. All trials had 276 a fixed duration of 4000ms, where in addition to the jitter used at the beginning of each trial, null trials 277 (4000ms) were randomly interspersed across the learning (60 trials; 20%) and transfer (72 trials; 20%) phase. 278 Each face was presented equally often on the left or right side, and choices were indicated with the right-hand 279 index (left) or middle (right) finger. Before the MRI session, participants performed a complete learning 280 phase to familiarize with the task (300 trials with different faces). In the MRI scanner, participants performed two learning blocks of 150 trials each (300 trials total; equal numbers of AB, CD and EF), and three transfer 282 phase blocks of 120 trials each (360 total; 24 presentations of each pair). All stimuli were presented on a 283 black-projection screen that was viewed via a mirror-system attached to the MRI head coil.

285 Reinforcement learning model

Trial-by-trial updating in value beliefs about the face selected in the learning phase, and reward prediction errors (signed expectancy violations) were estimated with a variant of the computational Q-learning algorithm^{27,64,65} that is frequently used with this reinforcement learning task and contains two separate learning rate parameters for positive (α_{gain}) and negative (α_{loss}) reward prediction errors^{4,23,25,58}. Q-learning assumes participants to maintain reward expectations for each of the six (A-to-F) stimuli presented during the learning phase. The expected value (Q) for selecting a stimulus i (could be A-to-F) upon the next presentation is then updated as follows:

$$Q_{i}(t+1) = Q_{i}(t) + \begin{cases} \alpha_{Gain}[r_{i}(t) - Q_{i}(t)], & \text{if } r = 1\\ \alpha_{Loss}[r_{i}(t) - Q_{i}(t)], & \text{if } r = 0 \end{cases}$$

Where $0 \le \alpha_{gain}$ or $\alpha_{loss} \le 1$ represent learning rates, t is trial number, and r = 1 (positive feedback) or r = 0 (negative feedback). The probability of selecting one response over the other (i.e., A over B) is computed as:

$$P_A(t) = \frac{\exp(\beta * Q_t(A))}{\exp(\beta * Q_t(B)) + \exp(\beta * Q_t(A))}$$

With $0 \le \beta \le 100$ known as the inverse temperature.

296 Bayesian hierarchical estimation procedure

To fit this Q-learning algorithm with two learning rate parameters we used Bayesian hierarchical estimation procedure. The full estimation procedure is explained in 25 . To summarize, this implementation assumes that probit-transformed model parameters for each participant are drawn from a group-level normal distribution characterized by group level mean and standard deviation parameters: $z \sim N(\mu_z, \sigma_z)$. A normal prior was assigned to group-level means $\mu_z \sim N(0,1)$, and a uniform prior to the group-level standard deviations $\sigma_z \sim U(1,1.5)$. Model fits were implemented in Stan, where multiple chains were generated to ensure convergence.

Image acquisition

The fMRI data for the Reinforcement learning task was acquired in a single scanning session with two learning and three transfer phase runs on a 3-T scanner (Philips Achieva TX, Andover, MA) using a 32-channel head coil. Each scanning run contained 340 functional $T2^*$ -weighted echo-planar images for the learning phase, and 290 $T2^*$ -weighted echo planar images for the transfer phase (TR = 2000 ms; TE = 27.63 ms; FA = 76.1°; 3 mm slice thickness; 0.3 mm slice spacing; FOV = 240 × 121.8 × 240; 80 × 80 matrix; 37 slices, ascending slice order). After a short break of 10 minutes with no scanning, data collection was continued with a three-dimensional T1 scan for registration purposes (repetition time [TR] = 8.5080 ms; echo time [TE] = 3.95ms; flip angle [FA] = 8°; 1 mm slice thickness; 0 mm slice spacing; field of view [FOV] = 240 × 220 × 188), the fMRI data collection using a stop signal task (described in 25), and a localizer task with faces, houses, objects, and scrambled scenes to identify FFA responsive regions on an individual level (317 $T2^*$

weighted echo-planar images; TR = 1500 msec; TE = 27.6 msec; $FA = 70^{\circ}$; 2.5 mm slice thickness; 0.25 mm slice spacing; $FOV = 240 \times 79.5 \times 240$; 96×96 matrix; 29 slices, ascending slice order). Here, participants viewed a series of houses, faces, objects as well as phase-scrambled scenes. To sustain attention during functional localization, subjects pressed a button when an image was directly repeated (12.5% likelihood).

³¹⁹ fMRI analysis learning phase

The interplay between learning and perceptual activity was examined by evaluating how trial-by-trial computations of value-beliefs, and reward prediction errors relate to BOLD responses in the occipital cortex 321 (OC) and fusiform face area (FFA). To compare perceptual responses with the more traditional literature, we 322 first show how value-beliefs and RPEs relate to the activity pattern of the dorsal (i.e., caudate, or putamen) 323 or ventral (i.e., accumbens) parts of the striatum. Regions of interest (ROI) templates were defined using 324 anatomical atlases available in FSL, or the localizer task for FFA. For this purpose, the localizer scans were 325 preprocessed using motion correction, slice-time correction, and pre-whitening⁶⁶. For each subject, a GLM 326 was fitted with the following EVs: for FFA, faces > (houses and objects), for PPA, houses > (faces and 327 objects) and for LOC, intact scenes > scrambled scenes. Higher-level analysis was performed using FLAME 328 Stage 1 and Stage 2 with automatic outlier detection⁶⁷. For the whole-brain analysis Z (Gaussianized T/F) 329 statistic images were thresholded using clusters determined by z > 2.3 and p < .05 (GRFT) to define a 330 group-level binary FFA region. Templates used for the caudate [center of gravity (cog): (-) 13, 10, 10], 331 putamen [cog: (-) 25, 1, 1], and Nucleus accumbens [cog: (-)19, 12, -7] were based on binary masks. Because 332 participants were asked to differentiate faces, for each participant, we multiplied the binary templates of OC 333 (V1) [cog: 1, -83, 5], FFA [cog: 23, -48, -18] with the individual t-stats from the localizer task contrast faces 334 > (houses and objects). All anatomical masks, and the localizer group-level FFA mask can be downloaded 335 from github (see acknowledgements). 336

Deconvolution analysis learning phase

To more precisely examine the time course of activation in the striatal and perceptual regions, we performed finite impulse response estimation (FIR) on the BOLD signals. After motion correction, temporal filtering 339 (3rd order savitzky-golay filter with window of 120 s) and percent signal change conversion, data from each 340 region was averaged across voxels while weighting voxels according to ROI probability masks, and upsampled 341 from 0.5 to 3 Hz. This allows the FIR fitting procedure to capitalize on the random timings (relative to 342 TR onset) of the stimulus presentation and feedback events in the experiment. Separate response time 343 courses were simultaneously estimated triggered on two separate events: stimulus onset, feedback onset. FIR time courses for all trial types were estimated simultaneously using a penalized (ridge) least-squares fit, as 345 implemented in the FIRDeconvolution package⁶⁸, and the appropriate penalization parameter was estimated 346 using cross-validation. For stimulus onset events (i.e., onset presentation of face pairs) response time courses were fit separately for the AB, CD and EF pairs, while also estimating the time courses of signal covariation 348 with chosen and unchosen value for these pairs. For these events, our analysis corrected for the duration of 349 the decision process. For the feedback events, the co-variation response time course with signed and unsigned prediction errors were estimated. These signal response time courses were analysed using across-subjects 351 GLMs at each time-point using the statsmodels package⁶⁹. The α value for the contributions of Q or RPE 352 was set to 0.0125 (i.e. a Bonferroni corrected value of 0.05 given the interval of interest between 0 and 8 s).

Random Forest classification

To specify the relevance of perceptual regions in the resolve of future value-driven choices a random forest 355 (RF) classifier was used²⁸. The RF classifier relies on an ensemble of decision trees as base learners, where 356 the final prediction (e.g., for a given trial is the choice going to be correct/optimal? or incorrect/suboptimal? 357 given past learning) is obtained by a majority vote that combines the prediction of all decision trees. To 358 achieve controlled variation, each decision tree is trained on a random subset of the variables (i.e. regions 359 of interest chosen), and a bootstrapped sample of data points (i.e. trials). In the construction of each tree 360 about 1/3 of all trials is left out - termed as the out-of-bag sample - and later used to see how well each tree 361 preforms on unseen data. The generalized error for predictions is calculated by aggregating the prediction for 362 every out-of-bag sample across all trees. An important feature of the RF classification method is the ease to measure the relative importance of each variable (i.e., region), in the overall predictive performance. That is, 364 it allows for the ranking of all regions evaluated in the prediction of future value-based decisions. 365

366 ROI selection and Random Forest procedure

This study used the 'Breiman and Cutler's Random Forests for Classification and Regression' package in R. termed randomForest. RF evaluations relied on the fMRI data recorded during the transfer phase, in a set 368 of 9 regions of interest (ROIs). These ROIs included all templates from the learning phase (i.e., caudate, 369 putamen, accumbens, OC, and FFA), as well as, the ventromedial prefrontal cortex (vmPFC), dorsolateral 370 prefrontal cortex (DLPFC), pre-supplementary motor area (preSMA), and the primary motor cortex (M1). 371 The selection of these additional anatomical templates was inspired by our previous analysis of this data 372 with those templates focusing on networks^{25,62,70}. From each ROI a single parameter estimate (averaged 373 normalized β estimate across voxels in each ROI) was obtained per trial, per subject. All, pre-processing steps 374 to obtain single-trial images are described in 25. Single-trial activity estimates were used as input variables 375 in RF to predict choice outcomes (optimal/sub-optimal) in the transfer phase. Here, participants choose the 376 best/optimal option based on values learned during the learning phase. We defined optimal choices as correct 377 (i.e., when participants choose the option with the higher value), and sub-optimal choices as incorrect. Misses 378 were excluded from RF evaluations.

By design, the transfer-phase contained 360 trials including 15 different pairs (12 novel), where each pair was 380 presented 24 times with the higher value presented left in 12 of the 24 presentations, and on the right for the 381 other half. With so many subtle value differences across the options presented and only one BOLD estimate per trial/region the prediction of future choices is under powered (Figure 5a). Therefore, assuming that all 383 participants come from the same population, a fixed effects approach was taken for evaluations with RF. Here, 384 the trial*region activity matrices for all participants were combined into one big data matrix (Figure 5b) and subsequently shuffled across the rows, so that both participants and trials were re-arranged in a random 386 order across rows. Besides the single trial BOLD estimates from the 9 ROI's, this shuffled matrix contained 387 two additional columns, which specified subject id (to which subject does each trial belong), and Trial Sign i.e., is the choice between the two faces about two positive (+/+; AC, AE, CE), negative (-/-; BD, BF, DF), 389 or a positive-negative (+/-; e.g. AD, CF etc.) associations given the task manipulation during learning. 390 Subject id was included to control for different BOLD fluctuations across participants, whereas Trial Sign was added because both BOLD and choice patterns differ across these options (please see 25). The shuffled 392 fixed effect matrix was divided into a separate training (2/3) of whole matrix, and validation (1/3) set, to 393

be used for RF evaluations (Figure 5c). Learning was based on the training set, using 2000 trees with the number of variables (regions) used by each tree optimized with the tuneRF function in R, and accordingly set to 5. For the construction of each tree about 1/3 of all trials is left out - termed as the out-of-bag sample – and later used to see how well each tree preforms on unseen data. The generalized error for predictions is calculated by aggregating the prediction for every out-of-bag sample across all trees. Besides this out-of-bag approximation we evaluated the predictive accuracy of the whole RF on the separate unseen validation-set. The single trial data used as input, the RF evaluation codes, and ROI templates can all be downloaded from the github link provided in acknowledgements.

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405 Author contribution

SJ and TK developed the questions and analysis plan for the re-analysis. SJ and TK contributed novel methods and analyzed the data. SJ wrote the first draft of the MS with edits from TK. JT commented on the final draft.

409 Competing interests

410 The authors declare to have no competing interests.

111 Data availability

The code and preprocessed files for behavioral and decoding analyses can be download from: https://github.com/sarajahfari/Pearl3T.git, and fMRI preprocessing and deconvolution analysis code are available at https://github.com/tknapen/pearl_3T. The raw data can be downloaded from openfMRI in BIDS after acceptance of this MS.

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