Striatal action-value neurons reconsidered

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Abstract

It is generally believed that during economic decisions, striatal neurons represent the values associated with different actions. This hypothesis is based on a large number of electrophysiological studies, in which the neural activity of striatal neurons was measured while the subject was learning to prefer the more rewarding action. Here we present an alternative interpretation of the electrophysiological findings. We show that the standard statistical methods that were used to identify action-value neurons in the striatum erroneously detect the same action-value representations in unrelated neuronal recordings. This is due to temporal correlations in the neuronal data. We propose an alternative statistical method for identifying action-value representations that is not subject to this caveat. We apply it to previously identified action-value neurons in the basal ganglia and fail to detect action-value representations. In conclusion, we argue that there is no conclusive evidence for the generally accepted hypothesis that striatal neurons encode action-values.

Key words

action-value; striatum; temporal correlations;

There is a long history of operant learning experiments, in which a subject, human or animal, repeatedly chooses between actions and is rewarded, often stochastically, according to its choices. A popular theory posits that the subject's decisions in these tasks utilize estimates of the different action-values. These action-values correspond to the expected reward associated with each of the actions, and actions associated with a higher estimated action-value are more likely to be chosen¹. In recent years, there is a lot of interest in the neural mechanisms underlying this computation^{2,3}. In particular, based on electrophysiological experiments⁴⁻¹⁵, it is now widely accepted that a population of neurons in the striatum represents these action-values, adding sway to this actionvalue theory. To identify neurons that represent the internal values of the different actions, researchers have searched for neurons whose firing rate is significantly correlated with the average reward associated with exactly one of the actions. There are several ways of defining the average reward associated with an action. For example, the average reward can be defined by the reward schedule: in a multi-armed bandit task with binary rewards, the average reward associated with an action can be defined as the corresponding probability of reward. Alternatively, one can adopt the subject's perspective, and use the subject-specific history of rewards and actions in order to estimate the average reward. In particular, the Rescorla-Wagner model (equivalent to the standard ones-state Q-learning model) has been used to estimate action-values^{4,6}. In this model, the value associated with an action i in trial t, termed $Q_i(t)$, is an exponentially-weighted average of the rewards associated with this action in past trials:

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$$Q_i(t+1) = Q_i(t) + \alpha (R(t) - Q_i(t))$$
 if $a(t) = i$ (1)

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$$Q_i(t+1) = Q_i(t)$$
 if $a(t) \neq i$

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- In the framework of a two-alternative task with binary rewards, $i \in \{1,2\}$, $a(t) \in \{1,2\}$ and $R(t) \in \{1,2\}$
- 46 $\{1,0\}$ are the possible actions, choice and reward in trial t, respectively, α is the learning rate and
- 47 Q_i is the action-value associated with action i.
- 48 It is typically assumed that the probability of choosing an action is a sigmoidal function, typically
- softmax, of the difference of the action-values (see also¹⁶):

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$$\Pr(a(t) = 1) = \frac{1}{1 + e^{-\beta(Q_1(t) - Q_2(t))}}$$
 (2)

- where β is a parameter that determines the tradeoff between exploration and exploitation (the bias
- towards the action associated with the higher action-value). The parameters of the model, α and
- β , can be estimated from the behavior, allowing the researchers to compute Q_1 and Q_2 on a trial-
- 54 by-trial basis.
- By measuring neural activity while the subject is performing the operant task, computing the
- regression of the trial-by-trial spike counts of the neurons on the latent variables $Q_i(t)$ and
- 57 identifying neurons for which this regression is statistically significant, one can identify the
- 58 neurons that represent action-values.
- 59 Using this framework, several electrophysiological studies in the past decade have found that the
- firing rate of a substantial fraction of striatal neurons (12%-40% for different significance
- 61 thresholds) is significantly correlated with the average reward associated with one of the actions,
- regardless of whether the action was chosen. These and similar results were considered as evidence
- that neurons in the striatum represent action-values $^{4-10,12}$.
- In this paper we point out that this literature has widely ignored a known caveat in regression
- 65 analysis it can result in erroneous identification of neurons as representing action-value if the

firing rates are temporally correlated. After a systematic literature search we conclude that this caveat has not yet been fully addressed. We maintain that the conclusion that there is representation of action-value in the striatum must await new evidence that is not prone to this caveat. Three clarifications are required. First, although this paper discusses a methodological problem that may also be of relevance in other fields of neuroscience, we focus on a single scientific claim, namely that a representation of action-values in the striatum is an established fact. Second, our criticism is restricted to the representation of action-values, and we do not make any claims regarding the possible representations of other decision variables, such as policy, chosen-value or reward-prediction-error. Third, we focus on the striatum and do not make claims about the possible representations of action-values elsewhere in the brain. The paper is organized in the following way. We commence by describing a standard method for identifying action-value neurons. Then, we demonstrate that this method erroneously identifies action-value neurons when they do not exist in a mathematical model, as well as in unrelated neuronal recordings from the motor cortex of a monkey, the auditory cortex of anaesthetized rats and the basal ganglia of behaving rats. Finally, we conduct a systematic literature search and show that all alternative approaches for identifying action-value neurons that were previously used can also lead to the erroneous identification of action-value neurons. We conclude by proposing a different method for identifying action-value neurons, that is not subject to this caveat and applying it to basal ganglia recordings, in which action-value neurons were previously identified. Using this new method, we fail to detect any action-value representations.

Results

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Identifying action-value neurons

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We commence by examining the standard methods for identifying action-value neurons using a simulation of an operant learning experiment. We simulated a task, in which the subject repeatedly chooses between two alternative actions, which yield a binary reward with a probability that depends on the action. Specifically, each session in the simulation was composed of four blocks such that the probabilities of rewards were fixed within a block and varied between the blocks. The probabilities of reward in the blocks were (0.1,0.5), (0.9,0.5), (0.5,0.9) and (0.5,0.1) for actions 1 and 2, respectively (Fig. 1a). The order of blocks was random and a block terminated when the more rewarding action was chosen more than 14 times within 20 consecutive trials^{4,10}. To simulate learning behavior, we used the Q-learning framework (Eqs. (1) and (2) with $\alpha = 0.1$ and $\beta = 2.5$ (taken from distributions reported in⁶) and initial conditions $Q_i(1) = 0.5$). As demonstrated in Fig. 1a, the model learned, such that the probability of choosing the more rewarding alternative increased over trials (black line). To model the action-value neurons, we simulated neurons whose firing rate is a linear function of one of the two O-values and whose spike count in a 1 sec trial is randomly drawn from a corresponding Poisson distribution. The firing rates and spike counts of two such neurons, representing action-values 1 and 2, are depicted in Fig. 1b in red and blue, respectively. One standard method for identifying action-value neurons is to compare the firing rates after learning by comparing the spike counts at the end of the blocks (horizontal bars in Fig. 1b). Considering the red-labeled Poisson neuron, the spike count in the last 20 trials of the second block, in which the probability of reward associated with action 1 was 0.9, was significantly higher than that count in the first block, in which the probability of reward associated with action 1 was 0.1 (p < 0.01; rank sum test). By contrast, there was no significant difference in the spike counts between the third and fourth blocks, in which the probability of reward associated with action 1

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was equal (p = 0.91; rank sum test; Fig. 1b, red). This is consistent with the fact that the red-labeled neuron was an action 1-value neuron: its firing rate was a linear function of the value of action 1. Similarly for the blue labeled neuron, the spike counts in the last 20 trials of the first two blocks were not significantly different (p = 0.92; rank sum test), but there was a significant difference in the counts between the third and fourth blocks (p < 0.001; rank sum test). These results are consistent with the probabilities of reward associated with action 2 and the fact that in our simulations, this neuron's firing rate was modulated by the value of action 2 (Fig. 1b, blue). This approach for identifying action-value neurons is limited, however, for several reasons. First, it considers only a fraction of the data, the last 20 trials in a block. Second, action-value neurons are not expected to represent the block average probabilities of reward. Rather, they will represent a subjective estimate, which is based on the subject-specific history of actions and rewards. Therefore, it is more common to identify action-value neurons by regressing the spike count on Qvalues, estimated from the subject's history of choices and rewards^{4-6,10-12}. Note that when studying behavior in experiments, we have no direct access to these estimated action-values, in particular because the values of the parameters α and β are unknown. Therefore, following common practice, we estimated the values of α and β from the model's sequence of choices and rewards using maximum likelihood, and used the estimated learning rate (α) and the choices and rewards to estimate the action-values (thin lines in Fig. 1c, see Materials and Methods). These estimates were similar to the true action-value, which underlay the model's choice behavior (thick lines in Fig. 1c). Next, we regressed the spike count of each simulated neuron on the two estimated action-values. As expected, the t-values of the regression coefficients of the red-labeled action 1-value neuron was significant for the estimated Q_1 $(t_{18} \pm Q_1) = 4.05$) but not for the estimated Q_2

 $(t_{18} \not Q_2) = -0.27)$. Similarly, the t-values of the regression coefficients of the blue-labeled action 2-value neuron was significant for the estimated Q_2 ($t_{18} \not Q_2$) = 3.05) but not for the estimated Q_1 ($t_{18} \not Q_1$) = 0.78).

A population analysis of the t-values of the two regression coefficients is depicted in Fig. 1d,e. As expected, a substantial fraction (42%) of the simulated neurons in the simulation were identified as action-value neurons. Only 2% of the simulated neurons had significant regression coefficients with both action-values. Such neurons are typically classified as state or policy (preference) neurons, if the two regression coefficients have the same or different signs, respectively 10 . Note that despite the fact that by construction, all neurons were action-value neurons, not all of them were detected as such by this method. This failure occurred for two reasons. First, the estimated Q-values are not identical to the true action-values, which determine the firing rates. This is because of the finite number of trials and the stochasticity of choice (note the difference, albeit small, between the thin and thick lines in Fig. 1c). Second and more importantly, the spike count

Identifying "action-value" neurons in the absence of value (model)

The identification of the simulated neurons in Fig. 1d,e as action-value neurons relied on the interpretation that a large t-value is highly improbable under the null hypothesis that the firing rate of the neuron is not modulated by action-values. However, when computing the significance threshold for rejection of the null hypothesis it was implicitly assumed that the different trials are independent. To see why this assumption is essential, we consider a case in which it is violated. Fig. 2a depicts the firing rates and spike counts of two simulated Poisson neurons, whose firing rates follow a bounded Gaussian random-walk process:

in a trial is only a noisy estimate of the firing rate because of the Poisson generation of spikes.

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$$f(t+1) = [f(t) + z(t)]_{+}$$
 (3)

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where f(t) is the firing rate in trial t (we consider epochs of 1 second as "trials"), z(t) is a diffusion variable, randomly and independently drawn from a normal distribution with mean 0 and variance $\sigma^2 = 0.01$ and $[x]_+$ denotes a linear-threshold function, $[x]_+ = x$ if $x \ge 0$ and 0 otherwise. These random-walk neurons are clearly not action-value neurons. Nevertheless, we tested them using the analyses depicted in Fig. 1. To that goal, we randomly matched the "trials" in the simulation of the random-walk neurons to the trials in the simulation depicted in Fig. 1a, and considered the spike counts of the random-walk neurons in the last 20 trials of each of the four blocks in Fig. 1a. Considering the top neuron in Fig. 2a and utilizing the same analysis as in Fig. 1b, we found that its spike count differed significantly between the first two blocks (p < 0.01, rank sum test) but not between the last two blocks (p = 0.28, rank sum test), similar to the simulated action 1-value neuron of Fig. 1b (red). Similarly, the spike count of the bottom random-walk neuron matched that of a simulated action 2-value neuron (compare with the blue-labeled neuron in Fig. 1b; Fig. 2a). Moreover, we regressed each vector of spike counts for 20,000 random-walk neurons on randomly matched estimated Q-values from Fig. 1e and computed the t-values (Fig. 2b). This analysis classifies 42% of these random-walk neurons as action-value neurons (see Fig. 2c). In particular, the top and bottom random-walk neurons of Fig. 2a were identified as action-value neurons for action 1 and 2, respectively (squares in Fig. 2b). To further quantify this result, we computed the fraction of random-walk neurons erroneously classified as action-value neurons as a function of the diffusion parameter σ (Fig. 2d). When σ =0, the spike counts of the neurons in the different trials are independent and the number of random-

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walk neurons classified as action-value neurons is slightly less than 10%, as expected from a significance criterion of 5% and two statistical tests, corresponding to the two action-values. The larger the value of σ , the higher the probability that a random-walk neuron will pass the selection criterion for at least one action-value and thus be erroneously identified as an action-value, state or policy neuron. The excess action-value neurons in Fig. 2 emerged because the statistical analysis was based on the assumption that the different trials are independent from each other. In the case of a regression of a random-walk process on an action-value related variable, this assumption is violated. The reason is that in this case, both predictor (action-value) and the dependent variable (spike count) slowly change over trials, the former because of the learning and the latter because of the random drift. As a result the statistic, which relates these two signals, is correlated between temporallyproximate trials, violating the independence-of-trials assumption of the test. Because of these dependencies, the expected variability in the statistic (be it average spike count in 20 trials or the regression coefficient), which is calculated under the independence-of-trials assumption, is an underestimate of the actual variability. Therefore, the fraction of random-walk neurons classified as action-value neurons increases with the magnitude of the drift, which is directly related to the magnitude of correlations between spike counts in proximate trials (Fig. 2d). Importantly, the Gaussian random-walk process is just one example in which the firing rate is nonstationary. Other processes, in which the firing rate is non-stationary (e.g., oscillatory or trend following) and thus the independence-of-trials assumption is violated may also lead to an erroneous identification of neurons as action-value neurons. For example, it has been suggested that synaptic plasticity that stochastically implements or approximates direct policy gradient learning underlies some forms of operant learning 17-22. In general, there will be no explicit or implicit representation of action-values when these algorithms are implemented. However, because neural activity in these algorithms slowly varies over trials, the methods described above for identifying action-value neurons may erroneously identify action-value representations in implementations of these algorithms. To test this, we studied learning mediated by covariance-based synaptic-plasticity^{21,23–25} in the learning task of Fig. 1a (Supplementary Information). Indeed, not only did this algorithm successfully learn to prefer the better alternative, when considering the spike counts of the simulated neurons in this algorithm, 43% of these neurons were erroneously identified as action-value neurons (Fig. S1).

Identifying "action-value" neurons in the absence of value (experiments)

In the previous section we demonstrated that the standard analysis depicted in Fig. 1 may lead to the erroneous identification of neurons as action-value neurons if the firing rate is sufficiently non-stationary. To test whether this theoretical finding is relevant to electrophysiological experiments, we considered the spike count of 89 single neurons recorded extracellularly from the motor cortex of a monkey. This was a brain-machine-interface (BMI) experiment composed of 600 identical trials (Materials and Methods). For the purpose of our analysis, we considered as a "trial" the spike count of the neuron in the last 1 sec of each inter-trial-interval in the original experiment.

As in the analyses in Fig. 2, every spike count sequence of a motor cortex neuron was randomly paired with a pair of estimated Q-values from one of the simulations of the operant task depicted in Fig. 1 (truncating the number of experimentally-measured trials in accordance with the number of trials in the simulation). Fig. 3a depicts two estimated Q-values from two sessions (lines), imposed on the spike counts (dots) of two motor cortex neurons. Similar to the random-walk neurons (Fig. 2a), we compared the spike count in the last 20 trials of each of the four blocks and found that the sequence of spike counts of the Top neuron in Fig. 3a matched that of an action 1-

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value neuron. The sequence of spike counts of the Bottom neuron in Fig.3a matched that of an action 2-value neuron. For the population analysis, we regressed all vectors of motor-cortex spike counts on the estimated Q-values of Fig. 1. Similarly to the results of the simulations of the random-walk neurons, 36% of the motor cortex neurons in this experiment were classified as action-value neurons (fig. 3b,c). These results demonstrate that the magnitude of non-stationarity in standard electrophysiological recordings is sufficient to result in an erroneous identification of neurons as representing actionvalues. To test whether this limitation of the analysis is restricted to extracellular recordings, we considered intracellular recordings of 39 auditory cortex neurons in 125 sessions (of which 29 sessions were excluded in all repetitions of the analysis due to low spike count, see Materials and Methods) in anaesthetized rats, responding to auditory stimuli²⁶ (Materials and Methods). In short, the animals were exposed to a long sequence of pure tones, presented every 300-1000 msec. Depending on the session, trials in our analysis were taken to be 300 msec or 500 msec long (trial length remained the same throughout a session) and included a single pure tone. Repeating the same analysis on these auditory cortex neurons (Fig. 4), 23% of the neurons passed the selection criterion for action-value neurons (Fig. 4b,c; see individual examples in Fig. 4a). Identifying representations of unrelated "action-value" in the basal ganglia To test whether the erroneous identification of action-value neurons in the motor and auditory cortices is relevant to the statistics of firing in the striatum, we considered recordings from the nucleus accumbens (NAc) and ventral pallidum (VP) of rats in an operant learning experiment⁷. The experiment was a combination of a tone discrimination task and a reward-based free-choice

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task. We considered only the free choice trials, in which the appropriate response of the animal was to perform a nose poke in either the left or right hole after exiting the center hole. The experimental session was composed of 4-11 blocks. The reward schedule in each block was pseudo randomly chosen from the ones presented in Fig. 1a. Blocks changed when the subject chose the higher-valued action at least 80% of the trials within 20 trials. As in⁷, we considered the spike count in three 1 sec phases of the trial - before nose poking in the central hole, 1 sec following initiation of choice-instruction tone and last 1 sec of nose poke in central hole. In what follows we aggregate the three phases. For each recording, we simulated the Q-learning model with a random sequence of blocks. This sequence of blocks was independent of the actual sequence of blocks used in that session both in the reward probabilities and in the timing of transition between blocks. To allow for regression of the entire spike sequence (mean and standard deviation of number of trials was 518 and 122, respectively) on the estimated Qs, longer sessions than those of Fig. 1 were simulated. These simulated sessions consisted of 3 random repetitions of the 4 blocks of used in Fig. 1 and were then truncated to fit the length of the spike sequence. As before, we used the results of this simulation to extract estimates of the two O-values and we regressed the sequence of spike counts on these randomly assigned estimated Q-values. The t-values of 642 regressions (214 neurons in three sessions) are presented in Fig. 5a. The standard analysis identified 43% of the neurons as action-value neurons, despite the fact that these action-values were completely unrelated to the experimental session in which these neurons were recorded (Fig. 5b).

Alternative approaches for identifying action-value neurons

So far, our population analysis was based on fitting Eqs. (1) and (2) to the sequence of actions and rewards and using the resultant Q-values as estimates of the action-values (Thin lines in Fig. 1c).

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Our analyses in Figs. 2-5 clearly demonstrate that this approach can lead to the erroneous identification of action-values neurons. However, other studies have concluded that action-value is represented in the striatum when utilizing alternative approaches. In order to challenge the finding of action-value neurons in the striatum, we conducted a literature search to find all the alternative approaches used to identify action-value representation in the striatum (see Materials and Methods). We identified 22 papers that directly related neural activity in the striatum to actionvalues. These papers included reports of single-unit recordings, functional magnetic resonance imaging (fMRI) experiments and manipulation of striatal activity. Of these, 3 papers have used the term action-value to refer to the value of the *chosen* action (also known as chosen-value)^{27–29} and therefore we will not discuss them any further. A second group of 11 papers did not distinguish between action-value and policy representations^{5,6,9,12–14}, or reported policy representation^{15,30–33}. While action-value representation is often implied from policy representation, it is well-known that policy representation can emerge in the absence of action-value representation. For example, in computer science, direct policygradient methods that do not entail values are routinely used³⁴. In neuroscience, several studies have proposed neuronal mechanisms that approximate direct policy reinforcement learning and decision making by means of reward-modulated synaptic plasticity (e.g., ^{17–25}). All these models will result in policy representation in the absence of action-value representation. For this reason, these findings do not necessarily imply action-value representation in the striatum. In 2 additional papers, it was shown that the activation of striatal neurons changes animals' behavior, and the results were interpreted in the action-value framework^{35,36}. However, a change in policy does not entail an action-value representation because, as noted above, a policy can be learned (or preference emerge) and be modulated by reinforcers without any action-value

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computation. Therefore, these papers are not a strong support to the striatal action-value representation hypothesis. Finally, 6 papers correlated action-values, separately from other decision variables, with neuronal activity in the striatum^{4,7,8,10,11,37}. Five used electrophysiological recordings of single units in the striatum and one was an fMRI study. All used block-design experiments where action-values are temporally correlated. In addition to the regression of the spike count on estimated Q-values described in Figs. 1-5 and S1, some of them considered alternative approaches. However, as described below, these alternatives are subject to the same caveat. A standard alternative approach to estimating action-values, which is model-free, is to use the average reward associated with the block as a measure of the action-value and regress the spike count at the end of the block on it^{4,7}. This is similar to the analysis in the individual examples of Figs. 1b, 2a, 3a, 4a and S1b (in which two rank sum tests, and not regression, were used). However, because this analysis is also based on the assumption of independence-of-trials, applying it to the random-walk neurons, as well as to the experimentally measured neurons, we identify a comparable number of action-value neurons to the one reported in the striatum (Fig. S2). In principle, temporal dependencies in the firing rates could result from trends. Indeed, detrending has been applied to the spike count¹⁰. In detrending, trial number is added to the regression model as an additional variable. However, this does not remove many of the temporal correlations. Indeed, we find a comparable number of erroneously-identified action-value neurons to that found in the striatum¹⁰ when applying this analysis to the random-walk model, the motor cortex, the auditory cortex and the basal ganglia neurons (Fig. S3).

It has also been noted that the significance analyses depicted above are biased towards classifying neurons as action-value neurons, at the expense of state or policy neurons⁸. The reason is that the former class requires a single significant regression coefficient whereas the latter require two significant regression coefficients (Figs. 1d, 2b, 3b, 4b, 5a, S1b, S2 and S3). Therefore, an unbiased alternative has been proposed⁸. However, for the same reasons (neural activities in consecutive trials are correlated), this analysis yields a comparable number of erroneously-identified action-value neurons in the random-walk, the motor cortex, the auditory cortex and the basal ganglia neurons to that reported in the striatum (Fig. S4).

Taken together, we conclude that previous reports on action-value representation in the striatum could reflect the representation of other decision variables or temporal correlations in the spike count that are not related to action-value learning.

Attempts to account for the non-stationarity

The mean length of the sessions used in the analysis in Figures 1-4 was 174 trials (standard deviation 43 trials). It is tempting to believe that adding more trials in a block or adding more blocks to the experiment may solve the problem of erroneous identification of action-value neurons. The idea is that the larger the number of trials, the less likely it is that a neuron that is not modulated by an action-value (e.g., a random-walk neuron) will have a large regression coefficient on one of the action-values. However surprisingly, this intuition is wrong. Specifically, we increased the number of trials by simulating the random-walk neurons in an eight-block design (as opposed to 4 blocks in Fig. 1), where the four blocks from fig 1a were repeated twice (both times in random permutation). The resulting mean length of the sessions was 347 trials (standard deviation 65 trials). We found that 45%±1.6% of the random-walk neurons were classified as action-value neurons. Similarly, when the spike counts of neurons from the motor cortex were

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regressed on estimated Q-values from the 8-blocks design, 37%±5% of these neurons were classified as action-value neurons. The same was done for spike counts of neurons from the auditory cortex (324 estimated Q-values with more than 370 trials did not participate in this analysis) and 24%±4.7% of these neurons were classified as action-value neurons. We did not perform this analysis on the basal ganglia neurons because we were limited by the length of these recordings. In fact, our results suggest that increasing the number of blocks can result in a larger fraction of erroneously-identified action-value neurons. This is because the estimated variance of regression coefficients is proportional to the inverse of the number of degrees of freedom, which increases with the number of trials. As a result, the significance threshold decreases with the number of trials. Adding blocks can be useful, however, if the reward schedules in the different blocks are independent, and the number of assumed degrees of freedom in the statistical analysis depends on the number of blocks and not on the number of trials. For example, the single-neuron statistical analysis in Figs. 2a, 3a and 4a is flawed because the variance in the mean spike count (in the last 20 trials of the block) is estimated assuming that the spike counts in consecutive trials are independent of each other. A correct analysis would have considered this mean as a single data point, in which the variance cannot be estimated. However, this is experimentally difficult because it requires a substantially larger number of blocks and thus trials in an experiment, than is typically used. Trial design experiments are not prone to the caveat we discuss here because by construction, the different trials are independent from each other, so the predictors in consecutive trials are not correlated. However, learning the values of actions requires that reward probabilities (or

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magnitudes) in consecutive trials will be strongly correlated, which is not possible in trial design. Several studies used tasks, in which cues mark the reward-probability 14,15,33. This way it is possible to use a trial design, in which the expected rewards associated with an action in consecutive trials are independent. However, these studies did not distinguish between values and policy representations. It should be noted that in this design, the learning is the association of cue and reward (cue-values), and not the association of action and reward (action-values)³⁸. Two studies noted that processes such as slow drift in firing rate may violate the independence-oftrials assumption of the statistical tests and suggested unique methods to deal with this issue in a block-design^{6,9}. Although in these studies action-value representation was not differentiated from policy representation, we repeated the methods described in them to see if they are subject to the same caveat described above. As shown below, we report erroneous detection of action-value neurons even when these methods are applied. In the first study⁶, a permutation test was proposed, in which the spike count, permuted within each block is also regressed on the estimated O-values for a large number of different permutations. A neuron is considered as an action-value modulated neuron if the t-value of the regression coefficient of the original spike count is large (in absolute value) relative to the distribution of tvalues of the permuted spike counts. Using a slightly modified reward schedule⁶, it was identified that 10%-13% of striatal neurons were significantly modulated by at least one of the action-values. However, this method may erroneously identify action-value neurons because the permutation reduces the correlations between the firing rates in consecutive trials. As a result, if there are temporal correlations in the original sequence of spike counts, the regression coefficients for the permuted spike counts are expected to be smaller than that of the original spike count. Indeed, conducting this analysis on the random-walk, the motor cortex, the auditory cortex and the basal

ganglia neurons, we found action-value neurons in a comparable number to that reported for the striatum (Fig. S5)

In the second study⁹, the spike-counts in the last 3 trials were also used as predictors in the regression model. However, this method does not address longer term dependencies and conducting this analysis using the same reward schedule as in⁹ on the random-walk, the motor cortex, the auditory cortex and the basal ganglia neurons, we found action-value neurons in a comparable number to that reported for the striatum (Fig. S6).

A possible solution to statistical significance in non-stationary time-series

The concerns about the statistical tests described above result from the possibility that non-stationarity of the spike count in striatal neurons is not the result of action-value learning. In principle, if the statistical structure of this non-stationarity is known, it may be possible to construct a statistical test, such as generalized least squares (GLS)³⁹ that decorrelates the trials. Alternatively, it may be possible to compare the number of identified action-value neurons to the expected number of erroneously detected "action-value" neurons. However, both approaches require an accurate model of the learning-independent non-stationarity, which is absent, and cannot be extracted from global measures such as the autocorrelation. This is because the non-stationarity could be due to many different processes, including oscillations, trends, random-walks and their combinations.

Therefore, we propose a non-parametric permutation test that does not make specific assumptions about the learning-independent non-stationarity. In contrast to the previously-described methods, this test allows us to estimate the probability of an erroneous detection of an action-value neuron

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under the null hypothesis. Importantly, this method can also be used to reanalyze the activity of previously-recorded striatal neurons^{4–10}. We propose a permutation test, in which we seek neurons that are more correlated with the actionvalue that was estimated from the session in which the neuron was recorded than with surrogate action-values that were estimated from other sessions. This is illustrated in Fig. 6. First, we computed the t-values of the regression coefficients of the spike counts of the two simulated actionvalue neurons in Fig. 1b on each of the estimated Q-values from all relevant sessions (see below). The two distributions of t-values, one for each simulated neuron, are depicted in Fig. 6a. Note that the 5% significance boundaries, which are exceeded by exactly 5% of t-values in each distribution, are substantially larger (in absolute value) than 2 (1.96 is 97.5th percentile in a t-distribution). There are two reasons for these wide distributions of t-values. First, there are trial-to-trial correlations both in the estimated Q-values and in the spike counts of the simulated action-value neurons. As a result, the effective number of degrees of freedom is substantially smaller than the number of trials, leading to larger t-values than expected when trials are independent. Second, some of the surrogate sessions corresponded to an identical or opposite sequence of reward probabilities, resulting in surrogate estimated O-values that are highly correlated (or anti-correlated) with the O-value that is estimated from the neuron's session. We posit that a regression coefficient is significant if the tvalue of the regression on the Q-value that is estimated from the neuron's session exceeds the significance boundaries derived from the permutations. Indeed, when considering the Top (red) simulated action 1-value neuron, we find that its spike count is significantly correlated with the estimated Q_1 from its session (red arrow) but not with that of estimated Q_2 (blue arrow). Because the significance boundary exceeds 2, this approach is less sensitive than the original one (Fig. 1) and indeed, the regression coefficients of the Bottom simulated neuron (blue) do not exceed the

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significance level (red and blue arrows) and thus this analysis fails to identify it as an action-value neuron. Considering the population of simulated action-value neurons of Fig. 1, this analysis identified 29% of the action-value neurons of Fig. 1 as such (Fig.6b, black), demonstrating that this analysis can identify action-value neurons. When considering the random-walk neurons (Fig. 2) or two of the experimentally measured neurons (Figs. 3 and 4), this method defines only approximately 10% of the neurons as action-value neurons, as predicted by chance (Fig. 6b). One technical point of caution is that the number of trials can affect the distribution of t-values. Therefore, we only considered in our analysis the first 170 trials of the 504 sessions longer or equal to 170 trials. We used this new method to consider action-value representation in the basal ganglia. To that goal, we considered the recordings reported in ⁷. That paper utilized the model-free method depicted in Fig. S2 to identify action-value neurons. They reported that 13% and 10% of the neurons × phases represent the left and right action-values, respectively (with p<0.01), suggesting that approximately 23% of the striatal neurons represent action-values at different phases of the experiment. As a first step in our analysis, we applied the standard model-based approach presented in Figs. 1d, 2b, 3b, 4b, 5a: we used the behavior of the animals to estimate the Q-values and regressed the spike counts in the three phases of the experiment on the estimated Q-values. This analysis yielded that approximately 32% of the neurons represent action values (16% $(103/(214 \times 3))$ and 16% $(100/(214 \times 3))$ of the neurons × phases represent the left and right action-values, respectively with p<0.01), a number that is slightly higher than the result of the model-free approach (Fig. 6c). Next, we applied the permutation analysis. Remarkably, this

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anlalysis yielded that only 3.6% of the neurons $(1.9\% (12/(214 \times 3)))$ and $1.7\% (11/(214 \times 3))$ of the neurons × phases) have a significantly higher regression coefficient with their corresponding left or right action-values, respectively, than with surrogate action-values (Fig. 6c). These results further challenge the hypothesis of action-value representation in the striatum. It is worth pointing out that the fraction of action-value neurons reported in is low relative to other publications^{4,10}, a difference that has been attributed to the location of the recording in the striatum (ventral as opposed to dorsal). It would be interesting to apply this method to other striatal recordings^{4,8,10}. Two points are noteworthy regarding this alternative analysis. First, Fig. 6a demonstrates that the distribution of the t-values of the regression of the spike count of a neuron on all action-values depends on the neuron. Similarly, the distribution of the t-values of the regression of the spike counts of all neurons on an action-value depends on the action-value (not shown). Therefore, the analysis could be biased in favor (or against) finding action-value neurons if the number of neurons per session is different between sessions. Second, this analysis is still biased towards classifying neurons as action-value neurons at the expense of state or policy neurons, as noted above⁸. Therefore, it may erroneously identify neurons whose activity is correlated with other decision variables, such as state or policy, as action-value neurons (Fig. S7). To prevent this, it is useful to apply the correction suggested in⁸. **Discussion** In this paper, we performed a systematic literature search to discern the methods that have previously been used to infer the representation of action-values in the striatum. We show that

none of the methods that have been proposed to distinguish between action-value representation

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and other decision variables are able to overcome a serious statistical caveat: temporal dependencies in the firing rates of neurons may result in their erroneous identification as representing action-values. Specifically, we considered a particular example of a violation of the independence-of-trials assumption by simulating neurons whose firing rates follow a bounded random-walk process. We erroneously identified apparent action-value representations in these simulated neurons. Moreover, these methods also erroneously identified neurons recorded in unrelated experiments in different cortical regions, as well as in the basal ganglia, as representing action-values. We propose an alternative method of analysis that is not subject to this limitation, which can be utilized to reanalyze data from previous experiments. When applying this novel method to basal ganglia recordings in which apparent action-value neurons were previously identified, we failed to detect action-value representations. It is important to note that we do not take these results to imply that erroneous detection of actionvalue representation may occur in every brain region and in any epoch of the trial. On the contrary, neurons in different brain areas, and even within the same brain area, differ according to their degree of non-stationarity and the time-constants of the modulations that cause this nonstationarity; features that both affect the probability of erroneous detection of action-value representation. Indeed, the fraction of erroneously identified action-value neurons differed between the auditory and motor cortices (compare Figs. 3 and 4). Considering the ventral striatum, our analysis indicates that the identification of action-value representations there may have been erroneous, resulting from non-stationary firing rates (Figs. 5, 6c). We were unable to directly analyze recordings from the dorsal striatum because relevant raw data is not publically available. However, previous studies have shown that the firing rates of dorsal-striatal neurons change slowly over time^{40,41}. As a result, spike counts are temporally correlated, and violate the independence-

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of-trials assumption which underlies all previous attempts to identify action-value neurons there^{4,8,10,11}. The potential statistical pitfalls associated with non-stationarity of neural activity are relevant to attempts to identify any neural correlate of a slowly changing variable, be it the spike count of a neuron, EEG signal from a sensor, or BOLD signal in a voxel. Our focus here was neural representations of action-value, but other variables associated with gradual learning are also likely to vary slowly and hence identifying them using any measure of neural activity will pose a similar challenge. Another situation in which such a problem may arise is in the estimation of noise correlations. For example, a population of neurons whose firing rates follow independent random-walk processes (or any other temporally correlated process), may appear to be endowed with significant noise correlations – again due to the violation of the independence-of-trials assumption, which leads to an underestimation of the variance of the correlation statistic under the null-hypothesis. Similar issues may arise when studying correlations in BOLD fluctuations between voxels when assessing resting state functional connectivity (see^{42,43} for discussions of autocorrelation in fMRI analyses). Any statistical test performed in these cases should consider the possibility of irrelevant nonstationarity. Returning to the question of action-value representations in the striatum, it has been previously noted that identifying a specific neuronal correlate of value is difficult, because it is hard to disentangle value from other variables, such as salience, the outcome's sensory properties or information about the properties of the task⁴⁴. It is also difficult to disentangle action-value representation from choice representation, as shown in Fig. S7.

To our knowledge, all studies that have claimed to provide direct evidence that neuronal activity in the striatum is specifically modulated by action-value were either susceptible to the statistical caveat demonstrated in this paper^{4,7,8,10,11,37}, or reported results in a manner indistinguishable from policy, which does not necessarily imply value representation (as shown in Fig. S1)^{14,15,33}. Indeed, many studies were susceptible to both of these confounds^{5,6,9,12,13}. Furthermore, it should be noted that not all studies investigating the relation between striatal activity and action-value representation have reported positive results. Several studies have reported that striatal activity is more consistent with direct policy learning than with action-value learning 45,46 and one noted that lesions to the dorsal striatum do not impair action-value learning⁴⁷. The fact that the basal ganglia in general and the striatum in particular play an important role in operant learning, planning and decision-making is not in question^{3,35,48–52}. However, our results show that special caution should be applied when relating activity in neurons there with specific variables, derived from reinforcement learning algorithms, which vary slowly over time. The prevailing belief that neurons in the striatum represent action-values must await further tests that can account for the potential caveats discussed here.

Materials and Methods

Literature search

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Key words "action-value" and "striatum" were searched for in Web-of-Knowledge, Pubmed and Google Scholar, returning 43, 21 and 980 results, respectively. In the first screening stage, we excluded all publications that did not report new experimental results (e.g., reviews and theoretical papers), focused on other brain regions, or did not address value-representation or learning. In the remaining publications, the abstract of the publication was read and the body of the article was

searched for "action-value" and "striatum". After this step, articles in which it was possible to find description of action-value representation in the striatum were read thoroughly. The search included PhD theses, but none were found to report new relevant data, not found in articles. Overall, we identified 22 papers that reported new evidence in support of action-value representation in striatal neurons ^{4–15,27–33,35–37} and these were considered in this manuscript.

- The action-value neurons model
- To model neurons whose firing rate is modulated by an action-value, we considered neurons whose
- 544 firing rate changes according to:

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$$f(t) = B + K \cdot r \cdot (Q_i(t) - 0.5)$$
 (4)

- Where f(t) is the firing rate in trial t, B = 2.5Hz is the baseline firing rate, $Q_i(t)$ is the action-
- value associated with one of the targets $i \in \{1,2\}$, K = 2.35Hz is the maximal modulation and r
- denotes the neuron-specific level of modulation, drawn from a uniform distribution, $r \sim U[-1,1]$.
- The spike count in a trial was drawn from a Poisson distribution, assuming a 1 sec-long trial.
- *Estimation of Q-values from model choices and rewards*
- To imitate experimental procedures, we regressed the spike count on estimates of the Q-values,
- rather than the Q-values that underlied behavior (to which the experimentalist has no direct access).
- For that goal, for each session, we assumed that $Q_i(1) = 0.5$ and found the set of parameters $\hat{\alpha}$
- and $\hat{\beta}$ that yielded the estimated Q-values that best fit the sequences of actions in each experiment
- by maximizing the likelihood of the sequence. Q-values were estimated from Eq. (1), using these
- estimated parameters and the sequence of actions and rewards. Overall, the estimated values of the
- parameters α and β were comparable to the actual values used: on average, $\hat{\alpha} = 0.12 \pm 0.09$
- (standard deviation) and $\hat{\beta} = 2.6 \pm 0.7$ (compare with $\alpha = 0.1$ and $\beta = 2.5$).

Exclusion of neurons

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Following standard procedures, a sequence of spike-counts, either simulated or experimentally measured was excluded due to low firing rate if the mean spike count in all blocks was smaller than 1. This procedure excluded 0.02% (4/20,000) of the random-walk neurons. 34% (42/125) of the auditory cortex neurons were excluded on average and 23% (29/125) were excluded in all 40 repetitions. 20% (126/(214×3)) of basal ganglia neurons were excluded on average and 11% were excluded in all 40 repetitions (74/(214×3)). None of the simulated action-value neurons (0/20,000) or the motor cortex neurons (0/89) was excluded.

- 567 Statistical analyses
- The computation of the t-values of the regression of the spike counts on the estimated Q-values
- was done using *regstats* in MATLAB. The following regression model was used:

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$$s(t) = \beta_0 + \beta_1 Q_1(t) + \beta_2 Q_2(t) + \epsilon(t)$$

- Where s(t) is the spike count in trial t, $Q_1(t)$ and $Q_2(t)$ are the estimated action-values in trial t,
- 572 $\epsilon(t)$ is the residual error in trial t and β_{0-2} are the regression parameters.
- To find neurons whose spike count in the last 20 trials is modulated by reward probability (Figs.
- 1b, 2a, 3a, 4a), we executed the Wilcoxon rank sum test, using *ranksum* in MATLAB. All tests
- were two-tailed.
- 576 The motor cortex recordings
- 577 The data was recorded from one female monkey (Macaca fascicularis) at 3 years of age, using a
- 578 10x10 microelectrode array (Blackrock Microsystems) with 0.4mm inter-electrode distance. The
- array was implanted in the arm area of M1, under anesthesia and aseptic conditions.

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Behavioral Task: The Monkey sat in a behavioral setup, awake and performing a BMI and sensorimotor combined task. Spikes and LFP were extracted from the raw signals of 96 electrodes. The BMI was provided through real time communication between the data acquisition system and a custom-made software, which obtained the neural data, analyzed it and provided the monkey with the desired visual and auditory feedback, as well as the food reward. Each trial began with a visual cue, instructing the monkey to make a small hand move to express alertness. The monkey was conditioned to enhance the power of beta band frequencies (20-30Hz) extracted from the LFP signal of 2 electrodes, receiving a visual feedback from the BMI algorithm. When a required threshold was reached, the monkey received one of 2 visual cues and following a delay period, had to report which of the cues it saw by pressing one of two buttons. Food reward and auditory feedback were delivered based on correctness of report. The duration of a trial was on average 14.2s. The inter-trial-interval was 3s following a correct trial and 5s after error trials. The data used in this paper, consists of spiking activity of 89 neurons recorded during the last second of intertrial-intervals, taken from 600 consecutive trials in one recording session. Pairwise correlations were comparable to previously reported⁵³, $r_{SC} = 0.047 \pm 0.17$ (SD), $(r_{SC} = 0.037 \pm 0.21)$ for pairs of neurons recorded from the same electrode). Animal care and surgical procedures complied with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and with guidelines defined by the Institutional Committee for Animal Care and Use at the Hebrew University. The auditory cortex recordings The auditory cortex recordings are described in detail in²⁶. In short, membrane potential was

experimental session were considered. Each session consisted of 370 50 msec tone bursts,

recorded intracellularly from 39 neurons in the auditory cortex of anesthetized rats. 125

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presented every 300-1000 msec. For each session, all trials were either 300 msec or 500 msec long. Trial length remained identical throughout a session and depended on smallest interval between two tones in each session. Trials began 50 msec prior to tone burst. For spike detection, data was high pass filtered with a corner frequency of 30Hz. Maximum points that were higher than 60 times the median of the absolute deviation from the median were classified as spikes. The Basal ganglia recordings The basal ganglia recordings are described in detail in⁷. In short, rats performed a combination of a tone discrimination task and a reward-based free-choice task. Extracellular voltage was recorded in the behaving rats from the NAc and VP using an electrode bundle. Spike sorting was done using principal component analysis. In total, 148 NAc and 66 VP neurons across 52 sessions were used for analyses (In 18 of the 70 sessions there were no neural recordings). Data Availability The data of the basal ganglia recordings is available online at https://groups.oist.jp/ncu/data and was analyzed with permission from the authors. Other data are available upon request. Code Availability Custom MATLAB scripts used to create simulated neurons and to analyze data are also available upon request. **Author Contributions** L.E.D. and Y.L. designed the analysis and wrote the paper Acknowledgements

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References

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- 632 1. Sutton, R. S. & Barto, A. G. Reinforcement Learning: An Introduction. (MIT Press, 1998).
- 633 2. Louie, K. & Glimcher, P. W. Efficient coding and the neural representation of value. *Ann.*
- 634 *N. Y. Acad. Sci.* **1251,** 13–32 (2012).
- 635 3. Schultz, W. Neuronal Reward and Decision Signals: from Theories to Data. *Physiol. Rev.*
- **95,** 853–951 (2015).
- 637 4. Samejima, K., Ueda, Y., Doya, K. & Kimura, M. Representation of Action-Specific
- Reward Values in the Striatum. *Science* **310**, 1337–1340 (2005).
- 5. Lau, B. & Glimcher, P. W. Value Representations in the Primate Striatum
- during Matching Behavior. *Neuron* **58**, 451–463 (2008).
- 641 6. Kim, H., Sul, J. H., Huh, N., Lee, D. & Jung, M. W. Role of Striatum in Updating Values
- of Chosen Actions. J. Neurosci. **29**, 14701–14712 (2009).
- 7. Ito, M. & Doya, K. Validation of Decision-Making Models and Analysis of Decision
- Variables in the rat basal ganglia. *J. Neurosci.* **29**, 9861–9874 (2009).
- 8. Wang, A. Y., Miura, K. & Uchida, N. The dorsomedial striatum encodes net expected
- return, critical for energizing performance vigor. *Nat. Neurosci.* **16**, 639–47 (2013).
- 647 9. Kim, H., Lee, D. & Jung, M. W. Signals for Previous Goal Choice Persist in the
- Dorsomedial, but Not Dorsolateral Striatum of Rats. *J. Neurosci.* **33**, 52–63 (2013).
- 10. Ito, M. & Doya, K. Distinct Neural Representation in the Dorsolateral, Dorsomedial, and
- Ventral Parts of the Striatum during Fixed- and Free-Choice Tasks. J. Neurosci. **35**, 3499–
- 651 3514 (2015).

- 11. Ito, M. & Doya, K. Parallel Representation of Value-Based and Finite State-Based
- Strategies in the Ventral and Dorsal Striatum. *PLoS Comput. Biol.* **11,** 1–25 (2015).
- 654 12. Funamizu, A., Ito, M., Doya, K., Kanzaki, R. & Takahashi, H. Condition interference in
- rats performing a choice task with switched variable- and fixed-reward conditions. *Front.*
- 656 *Neurosci.* **9,** 1–14 (2015).
- 657 13. Her, E. S., Huh, N., Kim, J. & Jung, M. W. Neuronal activity in dorsomedial and
- dorsolateral striatum under the requirement for temporal credit assignment. Sci. Rep. 6, 1–
- 659 11 (2016).
- 660 14. Cai, X., Kim, S. & Lee, D. Heterogeneous Coding of Temporally Discounted Values in
- the Dorsal and Ventral Striatum during Intertemporal Choice. *Neuron* **69**, 170–182 (2011).
- 662 15. Kim, S., Cai, X., Hwang, J. & Lee, D. Prefrontal and striatal activity related to values of
- objects and locations. Front. Comput. Neurosci. 6, 1–13 (2012).
- 664 16. Shteingart, H. & Loewenstein, Y. Reinforcement learning and human behavior. *Curr*.
- *Opin. Neurobiol.* **25,** 93–98 (2014).
- 666 17. Seung, H. S. Learning in Spiking Neural Networks by Reinforcement of Stochastic
- Synaptic Transmission. *Neuron* **40**, 1063–1073 (2003).
- 668 18. Fiete, I. R., Fee, M. S. & Seung, H. S. Model of Birdsong Learning Based on Gradient
- Estimation by Dynamic Perturbation of Neural Conductances. J. Neurophysiol. 98, 2038–
- 670 2057 (2007).
- 671 19. Urbanczik, R. & Senn, W. Reinforcement learning in populations of spiking neurons. *Nat.*
- 672 *Neurosci.* **12,** 250–252 (2009).

- 673 20. Darshan, R., Leblois, A. & Hansel, D. Interference and Shaping in Sensorimotor
- Adaptations with Rewards. *PLoS Comput. Biol.* **10,** 1–20 (2014).
- 675 21. Neiman, T. & Loewenstein, Y. Covariance-Based Synaptic Plasticity in an Attractor
- Network Model Accounts for Fast Adaptation in Free Operant Learning. J. Neurosci. 33,
- 677 1521–1534 (2013).
- 678 22. Fremaux, N., Sprekeler, H. & Gerstner, W. Functional Requirements for Reward-
- Modulated Spike-Timing-Dependent Plasticity. *J. Neurosci.* **30**, 13326–13337 (2010).
- 680 23. Loewenstein, Y. & Seung, H. S. Operant matching is a generic outcome of synaptic
- plasticity based on the covariance between reward and neural activity. *PNAS* **103**, 15224–
- 682 15229 (2006).
- 683 24. Loewenstein, Y. Robustness of Learning That Is Based on Covariance- Driven Synaptic
- 684 Plasticity. *PLoS Comput. Biol.* **4,** 1–10 (2008).
- 685 25. Loewenstein, Y. Synaptic theory of Replicator-like melioration. Front. Comput. Neurosci.
- **4,** 1–12 (2010).
- 687 26. Hershenhoren, I., Taaseh, N., Antunes, F. M. & Nelken, I. Intracellular Correlates of
- Stimulus-Specific Adaptation. J. Neurosci. **34**, 3303–3319 (2014).
- 689 27. Pasquereau, B. et al. Shaping of Motor Responses by Incentive Values through the Basal
- 690 Ganglia. **27,** 1176–1183 (2007).
- 691 28. Day, J. J., Jones, J. L. & Carelli, R. M. Nucleus accumbens neurons encode predicted and
- ongoing reward costs in rats. *Eur. J. Neurosci.* **33**, 308–321 (2011).
- 693 29. Seo, M., Lee, E. & Averbeck, B. B. Article Action Selection and Action Value in Frontal-

- 694 Striatal Circuits. *Neuron* **74,** 947–960 (2012).
- 695 30. Kim, Y. B. et al. Encoding of Action History in the Rat Ventral Striatum. J Neurophysiol
- **98,** 3548–3556 (2007).
- 697 31. Stalnaker, T. A., Calhoon, G. G., Ogawa, M., Roesch, M. R. & Schoenbaum, G. Neural
- 698 correlates of stimulus response and response outcome associations in dorsolateral
- 699 versus dorsomedial striatum. Front. Integr. Neurosci. 4, 1–18 (2010).
- 700 32. Guitart-Masip, M. et al. Go and no-go learning in reward and punishment: Interactions
- between affect and effect. *Neuroimage* **62**, 154–166 (2012).
- 702 33. Fitzgerald, T. H. B., Friston, K. J. & Dolan, R. J. Action-Specific Value Signals in
- Reward-Related Regions of the Human Brain. J. Neurosci. 32, 16417–16423 (2012).
- 704 34. Mongillo, G., Shteingart, H. & Loewenstein, Y. The misbehavior of reinforcement
- 705 learning. *Proc. IEEE* **102**, 528–541 (2014).
- 706 35. Tai, L.-H., Lee, A. M., Benavidez, N., Bonci, A. & Wilbrecht, L. Transient stimulation of
- 707 distinct subpopulations of striatal neurons mimics changes in action value. *Nat. Neurosci.*
- 708 **15,** 1281–9 (2012).
- To 36. Lee, E., Seo, M., Monte, O. D. & Averbeck, B. B. Injection of a Dopamine Type 2
- Receptor Antagonist into the Dorsal Striatum Disrupts Choices Driven by Previous
- Outcomes, But Not Perceptual Inference. J. Neurosci. 35, 6298–6306 (2015).
- 712 37. Wunderlich, K., Rangel, A. & O'Doherty, J. P. Neural computations underlying action-
- based decision making in the human brain. *PNAS* **106**, 17199–17204 (2009).
- 714 38. Padoa-Schioppa, C. Neurobiology of Economic choice: A Good-Based Model. *Annu. Rev.*

- 715 *Neurosci.* **34,** 333–359 (2011).
- 716 39. Montgomery, D. C., Peck, E. A. & Vining, G. G. Introduction to Linear Regression
- 717 *Analysis.* (WILEY, 2006).
- 718 40. Gouvea, T. S. et al. Striatal dynamics explain duration judgments. Elife 4, e11386 (2015).
- 719 41. Mello, G. B. M., Soares, S. & Paton, J. J. A Scalable Population Code for Time in the
- 720 Striatum. *Curr. Biol.* **25,** 1113–1122 (2015).
- 42. Woolrich, M. W., Ripley, B. D., Brady, M. & Smith, S. M. Temporal autocorrelation in
- univariate linear modeling of FMRI data. *Neuroimage* (2001).
- 723 43. Arbabshirani, M. R. et al. Impact of autocorrelation on functional connectivity.
- 724 *Neuroimage* (2014).
- 725 44. O'Doherty, J. The problem with value. *Neurosci Biobehav Rev* 43, 259–268 (2014).
- 726 45. Li, J. & Daw, N. D. Signals in Human Striatum Are Appropriate for Policy Update Rather
- 727 than Value Prediction. *J. Neurosci.* **31,** 5504–5511 (2011).
- 728 46. FitzGerald, T. H. B., Schwartenbeck, P. & Dolan, R. J. Reward-Related Activity in
- 729 Ventral Striatum Is Action Contingent and Modulated by Behavioral Relevance. *J.*
- 730 *Neurosci.* **34,** 1271–1279 (2014).
- 731 47. Vo, K., Rutledge, R. B., Chatterjee, A. & Kable, J. W. Dorsal striatum is necessary for
- stimulus-value but not action-value learning in humans. *Brain* **137**, 3129–3135 (2014).
- 733 48. McDonald, R. . J. & White, N. . M. A Triple Dissociation of Memory Systems:
- Hippocampus, Amygdala, and Dorsal Striatum. *Behav. Neurosci.* **107**, 3–22 (1993).

- 735 49. O'Doherty, J. et al. Dissociable Roles of Ventral and Dorsal Striatum in Instrumental
- 736 Conditioning. *Science* **304**, 452–454 (2004).
- 737 50. Thorn, C. A., Atallah, H., Howe, M. & Graybiel, A. M. Differential Dynamics of Activity
- Changes in Dorsolateral and Dorsomedial Striatal Loops During Learning. *Neuron* **66**,
- 739 781–795 (2010).
- 740 51. Yarom, O. & Cohen, D. Putative cholinergic interneurons in the ventral and dorsal regions
- of the striatum have distinct roles in a two choice alternative association task. *Front. Syst.*
- 742 *Neurosci.* **5,** 1–9 (2011).
- 743 52. Ding, L. & Gold, J. I. Caudate encodes multiple computations for perceptual decisions. J.
- 744 *Neurosci.* **30,** 15747–15759 (2010).
- 745 53. Cohen, M. R. & Kohn, A. Measuring and interpreting neuronal correlations. *Nat.*
- 746 *Neurosci.* **14,** 811–819 (2011).

Figures

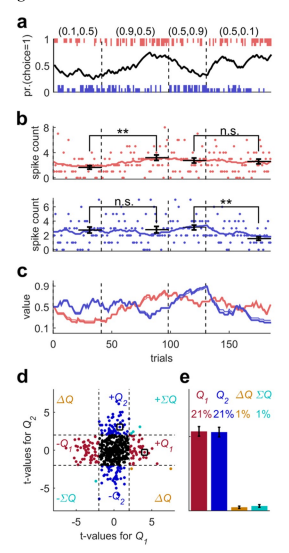


Figure 1 Model of action-value neurons (a) Behavior of model in example session, composed of four blocks (separated by a dashed vertical line). The probabilities of reward for choosing actions 1 and 2 are denoted by the pair of numbers above the block. Black line denotes the probability of choosing action 1; vertical lines denote choices in individual trials, where red and blue denote actions 1 and 2, respectively, and long and short lines denote rewarded and unrewarded trials, respectively. (b) Neural activity. Firing rate (line) and spike-count (dots) of two example simulated action-value neurons in the session depicted in (a). The red and bluelabeled neurons represent Q_1 and Q_2 , respectively. Black horizontal lines denote the mean spike count in the last 20 trials of the block. Error bars denote the standard error of the mean. The two asterisks denote p<0.01 (rank sum test). (c) Values. Thick red and blue lines denote Q_1 and Q_2 , respectively. Note that the firing rates of the two neurons in (b) are a linear function of these values. Thin red and blue lines denote the estimations of Q_1 and Q_2 , respectively, based on the choices and rewards in A. The similarity between the thick and thin lines indicates that the parameters of the model can be accurately estimated from the behavior (see also Materials and Methods). (d) and (e) Population

analysis. (d) Example of 500 simulated action-value neurons from randomly chosen sessions. Each dot corresponds to a single neuron and the coordinates correspond to the t-values of regression of the spike counts on the estimated values of the two actions. Color of dots denote significance: dark red and blue denote significant regression coefficient only on one estimated action-value, action 1 and action 2, respectively; light blue – significant regression coefficients on both estimated action-values with similar signs (ΣQ), orange - significant regression coefficients on both estimated action-values with opposite signs (ΔQ). Black – no significant regression coefficients. The two simulated neurons in (b) are denoted by squares. (e) Fraction of neurons in each category, estimated from 20,000 simulated neurons in 1,000 sessions. Error bars denote the standard error of the mean.

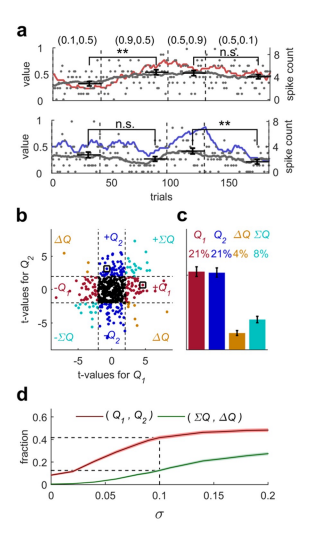


Figure 2 Erroneous detection of actionvalue representation in random-walk neurons (a) Two example random-walk neurons presented as if the sequence of spike counts of these neurons corresponds to the sequence of trials in figure 1. Gray lines and gray dots denote the neurons' firing rates and the spike counts, respectively. Black horizontal lines denote the mean spike count in the last 20 trials of the block. Error bars denote the standard error of the mean. The two asterisks denote p<0.01 (rank sum test). The red and blue lines denote the estimated action-values 1 and 2, respectively (same as in Fig. 1c). These are presented here because the t-value of the regression of the spike counts on the corresponding estimated action-values was larger than 2. (b) and (c) Population analysis, same as in Figs.1d and 1e for the random-walk neurons. The two randomwalk neurons in (a) are denoted by squares in (b). (d) Fraction of random-walk neurons classified as action-value neurons (red), and classified as state neurons (ΣQ) or policy neurons (ΔQ) (green) as a function of the magnitude of the diffusion parameter of random-walk (σ). Light red and light green

are standard error of the mean. Dashed lines mark the results for σ =0.1, which is the value of the diffusion parameter used in Fig. 2a-c. Initial firing rate for all neurons in the simulations is f(1) = 2.5Hz.

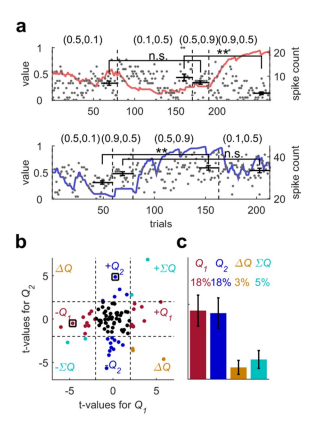


Figure 3 Erroneous detection of action-value representation in motor cortex (a) Two example motor cortex neurons recorded in a BMI task, presented as if the sequence of spike counts of these neurons corresponds to the sequence of trials in two sessions (one for each neuron) of operant learning used for the population analysis in Fig. 1e. Gray dots denote the spike-counts. Black horizontal lines denote the mean spike counts in the last 20 trials of the assigned blocks. Error bars denote the standard error of the mean. The two asterisks denote p<0.01 (rank sum test). Each session was associated with two estimated action-values and for each neuron. we computed the t-values of the regression of the spike counts on the two corresponding estimated action-values. The red and blue lines denote those action-values whose tvalue exceeded 2 (in absolute value). (b) and (c) Population analysis. (b) The t-values of 89 neurons regressed on the estimated

action-values of randomly selected 89 sessions (same as Fig. 1d). The neurons in (a) are denoted by squares. (c) Fraction of neurons classified in each category, estimated by regressing each of the 89 motor cortex neurons on 80 different estimated action-values from 40 randomly selected sessions. Error bars denote the standard error of the mean.

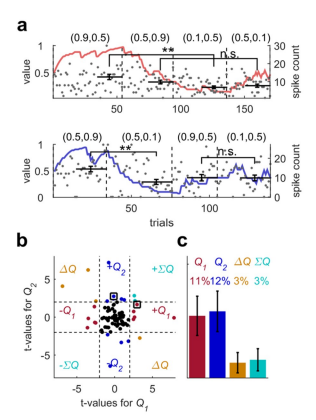


Figure 4 Erroneous detection of action-value representation in auditory cortex (a) Same as in Fig. 3a for two auditory cortex neurons in an anesthetized rat responding to the presentation of pure tones. (b) and (c) Population analysis. (b). The t-values of 82 neurons regressed on the estimated actionvalues of randomly selected 82 sessions (same as Fig. 3b). The neurons in (a) are denoted by squares. (c) Fraction of neurons classified in each category, estimated by regressing 125 auditory cortex neurons on 80 different estimated action-values from 40 randomly selected sessions (in each session, 34% of neurons were excluded on average, see Materials and Methods). Error bars denote the standard error of the mean.

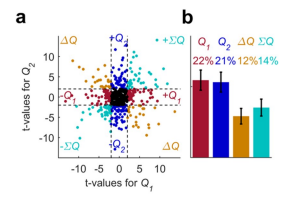


Figure 5 Erroneous detection of irrelevant action-value representation in basal ganglia (a) and (b) Population analysis. (a) The t-values of 214 neurons in three different phases regressed on the estimated action-values from randomly selected 642 simulated sessions (same as Fig. 4b). (b) Fraction of neurons classified in each category, estimated by regressing 214 neurons in three different phases on 80 different estimated action-values from 40 randomly selected sessions (in each session, 20% of neurons were excluded on average,

see Materials and Methods). Error bars denote the standard error of the mean.

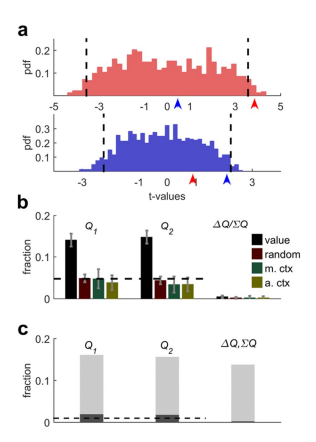


Figure 6 Permutation analysis (a) Red and blue figures correspond to red and blue labeled neurons in fig. 1b, respectively. For each neuron, we computed the t-values of the regressions of its spike-count on both estimated Q-values from all sessions in Fig. 1e (excluding sessions shorter than 170 trials) and used these t-values to compute probability distribution functions of the t-values. Dashed black lines denote the 5% significance boundary. Red and blue arrows denote the tvalues from regressions on the estimated Q_1 and Q_2 , respectively, from the session in which the neuron was simulated (depicted in Fig. 1a). **(b)** Fraction of neurons classified in each category using the permutation analysis for the action-value neurons (black, Fig. 1), randomwalk neurons (maroon, Fig. 2), motor cortex neurons (green, Fig. 3) and auditory cortex neurons (dark yellow, Fig. 4). Dashed line denotes chance level for action-value 1 or 2 classification. Error bars denote the standard

error of the mean. (c) Light gray, fraction of basal ganglia neurons classified in each category when regressing the spike count of basal ganglia neurons on the estimated Q-values associated with their experimental session. Dark gray, fraction of basal ganglia neurons classified in each category when applying the permutation analysis. Dashed line denotes significance level of p<0.01.