

Title: Dynamic refinement of behavioral structure mediates dopamine-dependent credit assignment

Dopamine initially reinforces spatially similar and temporally proximal actions to actions that trigger dopamine release, and drives a gradual refinement of the entire behavioral repertoire to home-in on reward-producing actions.

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Abstract

Animals exhibit a diverse behavioral repertoire when exploring new environments and can learn which actions or action sequences produce positive outcomes. Dopamine release upon encountering reward is critical for reinforcing reward-producing actions^{1–3}. However, it has been challenging to understand how credit is assigned to the exact action that produced dopamine release during continuous behavior. We investigated this problem with a novel self-stimulation paradigm in which specific spontaneous movements triggered optogenetic stimulation of dopaminergic neurons. We uncovered that dopamine self-stimulation rapidly and dynamically changes the structure of the entire behavioral repertoire. Initial stimulations reinforced not only the stimulation-producing target action, but also actions similar to the target and actions that occurred a few seconds before stimulation. Repeated pairings led to gradual refinement of the behavioral repertoire leading animals to home in on the target action. Reinforcement of action sequences revealed further temporal dependencies of behavioral refinement. Action pairs that tend to be spontaneously separated by long time intervals promoted a stepwise credit assignment, with early refinement of actions most proximal to stimulation and subsequent refinement of more distal actions. Thus, a retrospective reinforcement mechanism promotes gradual refinement of the entire behavioral repertoire to assign credit to specific actions and action sequences that lead to dopamine release.

Main Text

Background

Animals spontaneously transition amongst a repertoire of movements when exploring new environments. Movements or movement sequences that produce positive outcomes are reinforced and increase in frequency to maximize the obtainment of those outcomes^{4,5}. However, it is still not completely clear how animals assign credit to the exact action that produce reward in the context of a continuous behavioral space. This credit assignment problem^{2,6-9} during spontaneous behavior poses at least two main challenges. First, it is unclear how animals come to preferentially perform a specific reward-producing action or action sequence above other possibilities in the behavioral repertoire. Second, it is unclear how animals derive contingency between a reward-producing action and reward if there can be variable delays between action performance and reward delivery.

Dopamine (DA) has been proposed to mediate credit assignment^{6,10}. At the cellular level, DA can facilitate synaptic plasticity in corticostriatal synapses¹¹ within a critical time window that is behaviorally relevant¹²⁻¹⁴. Still, it is unknown how DA changes the dynamics of spontaneous behavior to mediate credit assignment. We therefore developed a paradigm to investigate how DA shapes the evolution of continuous behavior during action learning to gain insights into the process of credit assignment.

Conventional operant conditioning paradigms^{5,15-19} have helped derive principles of behavioral reinforcement, but they are not ideal for studying action credit assignment. In general, such paradigms do not permit the clean isolation of actions as the trigger for reward versus particular

locations or objects. In such paradigms, animals are also required to perform a series of consummatory actions, such as approaching and interacting with reward-delivering devices to retrieve reward. These requirements make it difficult to investigate how credit is assigned to a specific action or action sequence in the behavioral repertoire during continuous behavior.

Until recently, technological and conceptual limits have made it difficult to study how the entire structure of continuous behavior evolves as naive animals come to associate specific action or action sequences with reward. To address previous limitations, we developed a new approach to study action credit assignment. This approach directly reinforces specific spontaneous action(s) by triggering dopaminergic neuron (DA neuron) excitation and DA release upon action performance. It combines wireless inertial sensors, unsupervised clustering of continuous behavior^{20,21} and optogenetics²² into a closed-loop system linking specific action performance to immediate phasic DA release (Methods; Fig. 1a-f). This paradigm permits action detection and reinforcement without requiring an animal to approach or interact with a place/object/cue, or to perform consummatory behavior. These combined features overcome the aforementioned caveats associated with conventional paradigms.

Rapid reinforcement of actions via closed-loop dopamine stimulation

To implement the action detection component of the closed loop system, we first classified the entire behavioral repertoire of individual mice²³ mice in a grey-walled open field using inertial sensors and unsupervised affinity propagation clustering^{20,21} (Fig. 1d). Self-paced behavior was monitored using a novel, wireless inertial sensor system (WEAR; Methods) that allows minimal movement restraints, high resolution behavior monitoring and fast data transmission to open-

source hardware and software for online experimentation (Fig. 1b, Extended Data Fig. 1a).

Affinity propagation clustering is particularly well suited to cluster an unknown number of clusters²⁰, is computationally efficient²⁴, and easily outputs similarity between clusters.

Clustering begins by processing accelerometer and gyroscope data to extract 4 features discriminating postural changes, movement momentum, head and head-body rotations, and total body accelerations. Feature values from 300 ms long segments of behavior were discretized into histograms, upon which pairwise similarity comparisons could be made using a Earth-Mover's Distance (EMD)²⁵ metric. The similarity matrix of all possible pairwise comparisons were fed into an unsupervised affinity propagation clustering algorithm²⁰ (Methods), identifying naturally occurring repertoire of 300 ms long behavioral clusters²¹, or “actions” (Fig. 1c, Extended Data Fig. 1b). The choice of 300 ms long movements was informed by previous studies^{21,26}. Using these parameters, we identified over 30 clusters of spontaneous behavior per individual (34.3 +/- 2.1 and 35.6 +/- 2.5 total actions per ChR2-YFP and YFP mice, respectively; mean +/- standard deviation, 15 ChR2-YFP and 10 YFP mice). We chose particular clusters of actions to be reinforced (hereby named target action A).

To implement closed-loop reinforcement, we used Cre-dependent AAV viruses (EF1a-DIO-expression cassette) to express channelrhodopsin ChR2-YFP²² or the control protein YFP bilaterally in DA neurons of the ventral tegmental area (VTA)^{27,28} of DAT-Cre mice (Fig. 1a, Extended Data Fig. 2a-c). Using the wireless inertial sensor, we tracked behavior continuously in a white open field and used the similarity metric to match ongoing 300 ms behavioral segments to exemplars representing each mouse's repertoire of actions (Fig. 1d-e). Upon a match to a defined target action (target action A), a 25 hz, 600 ms long train of

optogenetic stimulation was delivered to DA neurons of the VTA parabrachial pigmented area (PBP) (30-60 ms delay, Fig. 1e). These target action As were different for different animals, and were dispersed across a behavioral space (Fig. 1g). To evaluate whether stimulation parameters triggered DA release similar in magnitude to that triggered by sucrose reward in food restricted mice, we delivered random optogenetic stimulations to ChR2-YFP- or YFP-expressing VTA DA neurons while monitoring DA release with the GRAB rDA1m sensor²⁹ in both ventral and dorsal striatum (Fig. 1f). We also measured DA release in the same animals upon delivery of sucrose while they were food deprived. Sucrose presentation led to a sharp increase in DA release in both ventral and dorsal striatum (Fig. 1f). Interestingly, optogenetic stimulation of DA neurons in VTA with the parameters described above, resulted in a similar phasic increase in DA not only in ventral striatum but also in dorsal striatum (Fig. 1f). This is consistent with emerging evidence showing the existence of dorsal striatum-projecting VTA neurons^{30,31}. Thus, our optogenetic stimulation triggered DA release similar in decay and spatial localization to that triggered by sucrose reward in food restricted mice (Fig. 1f), offering us a suitable approach to interrogate how pairing DA release with specific action performance leads to credit assignment.

Closed loop reinforcement for a specific action occurred over a 3-day, 60-90 minute/session protocol designed to probe both intra- and inter-session changes in behavior (Fig. 1h-m, Extended Data Fig. 3). Optogenetic stimulation of VTA DA neurons upon execution of a particular target action (action A) resulted in significant increase in the frequency of action A for ChR2-YFP, but not YFP mice (Fig. 1h, Extended Data Fig. 3b). Increased action A in ChR2-YFP animals depends on optogenetic stimulation, as removal of closed-loop stimulations resulted in progressive extinction of action A (Fig. 3h, Extended Data Fig. 3d). Resuming paired stimulation

led to rapid re-instatement of action A (Fig. 1h, Extended Data Fig. 3c,e). Interestingly, during extinction, ChR2-YFP animals kept performing exploratory unrewarded bursts of action A, which could explain rapid reinstatement (Extended Data Fig. 3e,f). This paradigm revealed that just a few pairings with DA leads to rapid reinforcement, as changes in multiple parameters including decreased trigger latency, increased action A frequency and increased average behavioral similarity towards action A become significant following 10-15 stimulations (Fig. 1i, Extended Data Fig. 4a-b).

We next examined if only action A changed in frequency or if other non-stimulated actions also changed with closed-loop reinforcement of action A. We calculated baseline-normalized frequency of all actions in the repertoire and ordered them as a function of similarity to the target action (Fig. 1j). Earth-Mover's Distance (EMD)^{21,25} was used to measure each action exemplar's similarity to the target exemplar (Methods), with lower EMD value indicating increased similarity. Surprisingly, we observed that optogenetic stimulation resulted in a dramatic change in the entire behavioral repertoire. We observed that early in training actions most similar to target tended to also increase in frequency (Fig. 1j-l, Extended Data Fig. 4c) whereas actions most dissimilar to target tended to decrease in frequency. Repeated pairing led to refinement of the actions that were performed at high frequency, and by late stages action A became the predominant action being performed, with a sharp drop-off of non-target action frequencies as similarity to target decreased (Fig. 1k-l). Such effects were not observed in YFP controls (Extended Data Fig. 4d-e). These data suggested that early reinforcement results in rapid reshaping of the entire behavioral repertoire, biasing animals towards actions similar to the target

action, and continued pairing resulted in gradual refinement and assignment of credit to the specific target action.

Dynamics of behavioral refinement during reinforcement

To better describe individual action dynamics during reinforcement, we categorized actions (511 actions, $n=15$ ChR2-YFP animals) by the trajectories of their changes in frequency throughout learning (Methods). Three meaningful types of trajectories were categorized, comprising over 94% of all actions. These types were characterized by either initial increase that remained stable (Sustained Increase), initial increases that decreased over time (Transient Increase) and initial decreases that remained stable (Decreased) (Fig 1m, Extended Data Fig. 5-6). We again confirmed that the frequency dynamics type of each particular action was related to its similarity to target, regardless of whether actions were sorted based on their raw or percentile similarity scores (Extended Data Fig. 6b-c). Actions most similar to target were predominately Sustained Increase types, while moderately similar actions mostly comprised of Sustained Increase or Transient Increase types and more dissimilar actions are more of the Decreased type (Extended Data Fig. 6b-c). Taken together, these finer resolution analyses indicate again that the dynamics of action frequency are related in great part to the similarity to target action.

Reinforcement and refinement after reversal of action-reward contingencies

Next, we asked if animals could follow changes in contingency between action and closed-loop DA stimulation. We therefore chose a different action, action B, which is clearly distinct from the action A for each animal (Methods, Fig. 2a, Extended Data Fig. 1c) and started delivering DA neuron optogenetic stimulation after action B. Chosen action A/B pairs were relatively

dissimilar in the context of entire action similarity distributions (Fig. 2b). Upon reinforcement, previously trained ChR2-YFP, but not YFP animals showed increased action B performance over time (Fig. 2c-e, Extended Data Fig. 7). In contrast, action A frequency changes clearly moved in the opposite direction from that of action B over time (Fig. 2c). Maintenance of action B performance depended on continual reinforcement (Fig. 2c, Extended Data Fig. 7d-e). Similar to action A, action B credit assignment unfolds by initially biasing the entire repertoire, i.e., increasing the frequency of similar actions and reducing the frequency of dissimilar actions. This was again followed by gradually refining for action B relative to similar actions as pairing progressed (Fig. 2d-e, Extended Data Fig. 7f). To confirm that action learning is contingent on action B appearing before reinforcement, we subjected trained animals to a contingency degradation protocol in which we delivered a similar number of random stimulations uncoupled to action B performance. Action B performance decreased following contingency degradation and could be re-instated upon resuming the action B-stimulation contingency (Fig. 2f, Extended Data Fig. 7g). These experiments indicate that animals can follow changes in the contingency between actions and DA release and assign credit to a new action through a similar process of behavioral repertoire refinement.

Although animals show similar patterns of behavioral refinement for actions A and B, animals that previously credited an action (action A) for DA release did initially respond to reinforcement of a new action (action B) differently from naïve animals (Fig. 2g-j). Whereas naïve animals responded to initial reinforcements for target action A by significantly increasing action A performance relative to the non-target action B (Fig. 2g,i,left graph), animals with a history of reinforcement on action A animals responded to initial reinforcements of action B by increasing

non-target action A performance (Fig. 2g,i,right graph). This trend reverses later such that target action B becomes significantly increased over the non-target action A (Fig. 2g,i,right graph). YFP control animals showed no such trends (Fig. 2h,j). Thus, DA reinforcement does not simply reinforce the recently performed, temporally contiguous action, but trigger previously credited actions in the face of a new action-reward contingency that is not yet learned. This suggest again that animals learned the contingency between action performance and DA release.

Temporal constraints of DA-dependent reinforcement

The contingency degradation results above indicate that the temporal relation between target action and DA phasic activity is important for reinforcement (Fig. 2e). Reinforcement is thought to occur on behavior that precedes reward in time^{10,12,14,19}, and while temporal contiguity between action and reinforcement has long been recognized³²⁻³⁴, it is not clear how the position of an action relative to the time of DA phasic activity influences its subsequent frequency. We investigated if in addition to behavioral similarity, the temporal relationship between action and stimulation influenced the dynamics of behavioral repertoire evolution during reinforcement and credit assignment.

We observed that the median inter-target action interval decreased with stimulation in ChR2-YFP mice (Fig. 3a,b). We therefore examined the distribution of the action dynamic types categorized above (Sustained Increase, Transient Increase, Decreased) according to both an action's similarity to target and the median time of that action's performance leading into target during baseline, before reinforcement protocol began (Fig. 3c-e). Action dynamic types showed distinct distribution patterns for these two dependent variables (similarity and time). Further,

these two dependent variables were not significantly collinear (Methods). Thus, action similarity to target as well as baseline temporal proximity to target should together predict action dynamic type upon reinforcement better than either factor alone. To test this idea, we performed multinomial logistic regression to assess whether 1- or 2-factor models best fit the observed dynamics pattern that an action would follow upon reinforcement (Fig. 3f,g). The two-factor model outperformed either one-factor models, and prediction of action dynamics type with this model was significantly above chance as assessed by precision-recall curves, which is suitable for evaluating datasets with imbalanced categories³⁵ (Fig. 3g). The beta coefficients indicated that increased similarity to target and decreased median time to target increases prediction of Sustained Increase and Transient Increase dynamic types relative to Decreased types (Supplementary Table). These results suggest that DA may reshape behavioral repertoire by reinforcing not only actions similar to the target action but also actions that happen to be performed temporally close to the reinforcer, as suggested before^{10,12,14,19}.

To more rigorously test whether DA reinforcement acts in a retrospective or prospective manner, we increased the resolution of analysis by examining 1st order action transitions leading into and out of stimulation (Fig. 3h-j). By focusing analysis on action transitions enriched within specific 1.2 second moving windows, one could distinguish more clearly behavior that occurred leading up to, during, and after DA stimulation. Our analyses showed that action transitions enriched in windows up to 1.2 seconds prior to stimulation onset, as well as during stimulation, are reinforced early on (Fig. 3i). However, this did not occur to action transitions following stimulation, suggesting an asymmetric process. Indeed, action transitions enriched in windows leading into stimulation were also preferentially reinforced relative to those enriched in windows

after stimulation (Fig. 3j). Thus, DA stimulation promotes reinforcement of behaviors occurring during stimulation and a few seconds before stimulation.

Credit assignment for action sequences

In the real world, when animals are spontaneously shifting between actions in their repertoire, outcomes are often not the result of a single action but rather of a sequence of actions performed at variable intervals. We therefore investigated the dynamics of reinforcement when the release of DA is contingent upon the performance of a sequence of 2 actions (target action 1 and 2, T1 and T2). We applied closed loop optogenetics to ask whether naïve animals can learn a T1→T2 reinforcement rule, where the delays between T1 and T2 are governed by the spontaneous behavior of the animals and not experimentally controlled (n=15 ChR2-YFP and 10 YFP mice, Fig. 4a, Extended Data Fig. 2a,d-e, Extended Data Fig. 8-10). Various T1/T2 pairs were sampled, with focus on sequences sharing general commonalities in movement order across animals (Extended Data Fig. 1d,f-g). Overall, mice learned to increase the performance of a sequence of two actions to obtain DA stimulation. Some animals showed a ChR2-dependent increase in reinforcement within 5 sessions, but others experienced a lag in learning (Fig. 4b). We hypothesized that this could relate to the initial time distance between T2 trigger and the closest distal T1 (T1→T2 interval). Indeed, animals reinforced for action pairs with initially long interval values tended to show slower learning curves (Fig. 4c-d). To capture a learning time point whereby individuals reach similar rising phase in their respective learning curves, a criterion frequency was set (Methods). 14 of 15 trained animals eventually reached criterion (Fig. 4e; Extended Data Fig. 8a-c). Sequence performance depended on continuing DA reinforcement (Fig. 4f,g). Learning was also revealed by decreases in the median T1→T2 time

intervals (Fig.4h-i) and convergence of T1-to-T2 frequency ratio towards 1 (Fig. 4j). To quantify the specific credit assignment of T1 and T2 we used a refinement index that compares the median frequency of actions uniquely similar to T1 with those uniquely similar to T2, with the frequencies normalized by either that of T1 or T2 (Methods). Values lower than 1 indicate that the target actions are being performed even more frequently than similar actions, and thus indicate greater refinement (Methods). By the end of learning, T1 and T2 became credited as the reward-producing actions relative to their similar counterparts (Fig. 4k). YFP controls did not show any of these trends (Fig.4c-d,4g-h). Thus, closed loop reinforcement promoted learning of a two-action sequence rule in freely moving mice starting from a naïve state.

Importantly, the initial median T1→T2 interval performed by ChR2-YFP animals was inversely related to the eventual number of sessions required for each animal to reach criterion frequency (Fig. 4l). A sigmoidal curve was fit to the data, showing that animals with longer open field T1→T2 intervals beyond the sigmoidal midpoint tended to face sudden increase in sessions to reach criterion frequency (Fig. 4l). ChR2-YFP animals were divided according to the half-maximum point of the sigmoidal curve into 'Fast Learners' and 'Slow Learners'. Fast Learners quickly reached criterion frequency and low T1→T2 time intervals, whereas Slow Learners experienced a time lag in reaching criterion frequency and low T1→T2 intervals. Slow Learners tended to suddenly increase the frequency of sequence performance in sessions that showed a drop in the median T1→T2 interval to below 2-4 seconds (Fig.4d,h). In contrast, there was no stable sigmoidal relationship between T1-T2 action similarities and sessions to criterion frequency (Extended Data Fig. 8d). Thus, the initial median time distances between distal action

T1 and proximal action T2(which produced DA stimulation) modulated how fast animals learned to effectively perform the reinforced action sequence.

If DA is acting retrospectively to reinforce actions performed earlier in time, we hypothesized that the action most proximal to reinforcement, T2, should experience earlier refinement relative to the distal action, T1. We again used the median target normalized frequencies of actions uniquely related to T1 or T2 as refinement indices (Methods). Proximal T2 clearly refines towards its most refined level earlier than the distal T1, at least in some animals (Fig. 5a). By subtracting the area under the refinement curve for T1 from the curve for T2, one could calculate differential refinement between the two actions. Positive values indicate refinement preferentially favoring T2, and vice versa. A linear relationship was found between open field median T1→T2 interval and differential refinement between T1 and T2 (Fig. 5b). This suggests for longer T1→T2 median intervals, the proximal action T2 spends more sessions being more refined than the distal action T1. In contrast, there was no significant linear relationship between the initial intervals between the execution of the proximal action that led to reward and the next initiation of the sequence (T2→T1) or of the similarity between T1 and T2 actions, and the dynamics of differential refinement between T1 and T2 (Fig. 5b, right graph, Extended Data Fig. 9a).

We next investigated if the differential refinement between T1 and T2 was different for slow and fast learners. We analyzed changes in T1-T2 refinement curves relative to ‘Starting Points’ at which the refinement indices of T1 and T2 are most similar or are biased towards the distal T1 rather than the proximal T2 action (Methods). All Slow Learners showed a pattern where they

initially refine the repertoire of T2 from these Starting Points, and after reaching a maximum Turning Point, they start showing a bias towards greater T1 refinement (Fig. 5c). Notably, by these Turning Points the median intervals of T1→T2, but not T2→T1 events had decreased significantly relative to initial values (Fig. 5d, Extended Data Fig. 9b). Therefore, the median T1→T2 interval decrease occurred before a decrease in the interval to perform the next sequence (T2→T1), which started decreasing after the Turning Point (Fig. 5e). Using these learning landmarks, we asked more rigorously how animals homed in on T1 vs T2 over time (Fig. 5f, Extended Data Fig.10a). We found that animals initially refined the action proximal to DA stimulation (T2, between Starting Point and Turning Point), whereas T1 refinement occurred several sessions later, after the Turning point (Fig. 5f, Extended Data Fig.10a). Indeed, the Turning Point coincided with an increased probability of the T1 being found within 3.6 secs before T2 and reinforcement (Fig. 5g-h). These results indicate that animals can assign credit to sequences of actions that lead to reinforcement, following similar retrospective dynamics that were observed for single actions, whereby the actions most proximal to reinforcement are refined earlier and the actions more distal to reinforcement refined later, when they probabilistically start to occur within a few seconds of DA release.

Discussion

Our results demonstrate that DA reinforcement promotes single action credit assignment from a naïve state through a dynamic process whereby the entire behavioral repertoire is restructured. During the initial stages of reinforcement both actions similar to the target action and actions that were performed in close temporal proximity of the target action increase in frequency, while very dissimilar actions decrease in frequency. With repeated reinforcement there is a process of

gradual refinement that homes in on the action that produces DA release. In the case of action sequences, we observe a similar gradual refinement process whereby credit assignment for the action sequence is accomplished by early refinement for the actions most temporally proximal to reinforcement, followed by later refinement for the more temporally distal actions.

Previous synaptic and cellular studies^{36,37} proposed that DA reinforcement may act retrospectively to reinforce behavior. By utilizing the closed loop system, we rigorously tested this prediction. Since retrospective reinforcement of behavior is not confined to the target action alone, it facilitates credit assignment to a stimulation-producing action even when reinforcement is delayed; stimulation-producing action pairs that tend to be performed close together in time were learned much faster than pairs that tended to be performed far apart in time. Intriguingly, animals eventually learned to assign credit to distal stimulation-producing actions even in the latter scenario. This is characterized by a gradual process whereby early on, the median time interval between distal and proximal target actions decreased and the repertoire proximal to reinforcement was preferentially refined to favor the performance of the proximal target action. As the distal target action became significantly more likely to occur within second timescale distance prior to reinforcement, retrospective reinforcement of the correct stimulation-producing sequences became increasingly likely, resulting in whole behavioral refinement for the distal target as well, hence increasing sequence performance (Fig. 5g).

It has been suggested that retrospective reinforcement of behavior is mediated by DA modulation of an eligibility trace left by action potential-triggered synaptic plasticity¹⁰. Studies of DA action at the striatal synaptic level^{36,37} indicate that the timescale within which retrospective reinforcement may occur is on the order of a few seconds, but the behavioral consequences have

remained elusive until now. Our behavioral findings are consistent with cellular studies in that behavior occurring within a few seconds leading into DA stimulation are reinforced. It is also noteworthy that distal T1 refinement in two action reinforcement occurs after the closest T1 to DA stimulation has become more probable within a few seconds of stimulation. The cutoff of retrospective reinforcement by phasic DA activities within a few seconds could explain the sudden increase in sessions required to reach criterion frequency amongst animals that were reinforced for action pairs with initially longer median time separations. Retrospective behavioral reinforcement may be mediated by DA modulation of Ca²⁺ influx left by earlier spiking activities. Ca²⁺ influx triggered by NMDA receptors would increase adenosine 3',5'-cyclic monophosphate at thin distal dendrites of medium spiny neurons, leading to transient and localized protein kinase A activity specifically within the retrospective time window, as regulated by high phosphodiesterase activity¹⁴. Similar actions have more similar and overlapping striatal neural ensemble activities²¹. Arrival of DA upon activation of action-specific ensembles may reinforce not only a specific action, but also similar actions. As striatal ensembles specific to actions are activated and a trail of eligibility traces is left temporally, DA arrival could set the stage for retrospective reinforcement of a spatially graded repertoire of actions within a few seconds, resulting in the observed behavioral learning patterns. Future studies testing these ideas would clarify how synaptic plasticity and cellular ensemble activities integrate to produce a dynamic refinement process, resulting in the behavioral principles for credit assignment revealed here.

END OF MAIN TEXT

Methods

Animals: All experiments were approved by the Portuguese DGAV and Champalimaud Centre for the Unknown Ethical Committee and performed in accordance with European guidelines. They were also performed according to National Institutes of Health (NIH) guidelines and approved by the Institutional Animal Care and Use Committee of Columbia University. 3-5 months old DAT-Cre male mice in the C57/BL6J background²³ were used.

Sample Sizes, randomization, and blinding. For sample size, we applied a power of 0.8, significance of $p < 0.05$, and standard variation of 20% of the mean. We determined sample sizes of 4-8 mice per group for different mean-based tests (matched pairs, 2 groups). No formal method of randomization was used; littermates were equally divided among the groups being compared. The experimenter was not blinded of the experimental groups. Optogenetic manipulations were performed automatically via a computer algorithm and not manually by the experimenter.

Recombinant adeno-associated viral vectors, stereotaxic injections, and implants. 750 nl of rAAV.EF1a.DIO.hChR2(H134R).eYFP or rAAV.EF1a.DIO.eYFP ($3-4 \times 10^{12}$ vg/ml, AAV5, University of North Carolina Vector Core; $1-2 \times 10^{13}$ vg/ml, AAV1, Addgene, 27056-AAV1 and 20298-AAV1) were injected into each hemisphere of the VTA of 3-4 month old DAT-Cre mice. For viral injections, the coordinates are AP - 3.52 mm, ML - ± 0.35 mm, DV - 4.3 mm. Injections were made at 0.2 Hz pulses. Each pulse injects 4.6 nl volume. Injected needles were kept in place in the injection site for ~15 minutes before withdrawal. For each mouse, a dual optic fiber cannula (200/240 μ m diameter, 6 mm length, 0.7 mm center-to-center FLT, 0.22 NA;

Doric, DFC_200/240-0.22_6mm_DF0.7_FLT) was placed 200 µm above the injection site and fixed to the skull. Next, a 4-position receptacle connector (Harwin Inc., M52-5000445) was fixed anteriorly to the dual optic fiber cannula, with its posterior edge set at -0.6 mm. Skull implants are then fixed with dental cement. A 4-position connector (Harwin Inc., M52-040023V0445) with pins removed from one end was used to cap the receptacle connector.

For photometry experiments, 3-5 month old DAT-Cre males were used. The conditions used for VTA injections and implants were as above. Additionally, 1 µl and 500 nl of AAV9-hSyn-GRAB-rDA1m (2×10^{13} vg/ml; Addgene, 140556-AAV9) were injected into the dorsal striatum (AP 0.5 mm, ML +2.1 (right), DV 2.3 (from brain surface)) and ventral striatum (AP 1.15mm, ML +1.65 (right), DV 4.2 (from Bregma)) , respectively. For photometry fiber implants, mono fiberoptic cannula were used (400/430 µm diameter, 4 mm length (dorsal striatum) and 6 mm length (ventral striatum), 0.37 NA, 1.25 diameter ferrule, flat; Doric, MFC_400/430-0.37_6mm_MF1.25_FLT (ventral striatum) and MFC_400/430-0.37_4mm_MF1.25_FLT (dorsal striatum)). Implants were inserted at a 22 degrees angle. For dorsal striatum implantation, the cannula entered the skull at AP 0.5 mm and ML 3.03 mm at 22-degree angle. The angled implant penetrated the brain from its surface for 1.92 mm. For ventral striatum implantation, the cannula entered the skull at AP 2.85 mm at 22 degrees angle, ML 1.65 mm. The angled implant penetrated the brain from its surface for 4.25 mm.

WEAR motion sensor system. The WEAR motion sensor family was developed by the Champalimaud Hardware platform and Costa lab as a wired or wireless solution to obtain self-centered 9-axis motion data based on 3-axis accelerometer, gyroscope, and magnetometer

(<https://www.cf-hw.org/harp/wear>). The wired version is a very small and extremely lightweight device (200mg) that can sample motion data up to 500 Hz and at the same time provide current up to 500mA that can be used to power LEDs for optogenetic experiments or stimulating electrodes. The wireless version is small and lightweight (~1.8g) and can sample motion data up to 200 Hz while having the ability to provide up to 50 mA that can be used to power LEDs for optogenetic experiments or stimulating electrodes. The battery of the wireless WEAR allows recordings up to 4 h at 200 Hz sampling rate and even more at lower sampling rates. These devices communicate with the computers through a base station based on the HARP design developed by the Champalimaud Hardware Platform, which can be accessed through a software GUI to easily change sensor parameters to best fit the experimental needs. The base stations have several important hardware features such as 2 digital inputs and outputs, an analog input, 2 outputs for camera triggering, and a clock sync input and output that provides hardware-based synchronization. The sensor can be started or stopped by software or pin. The WEAR motion sensor family and base station are all open source (repository at <https://bitbucket.org/fchampalimaud/workspace/projects/HP>). Moreover, the WEAR devices are compatible with the Bonsai visual reactive programming software (<https://bonsai-rx.org/>), also open source, and allow the integration and synchronization of the streams of data being collected using the WEAR sensor with other data sources such as cameras.

Taking these specs and features together, the WEAR allows researchers to acquire high-resolution motion data wirelessly and for long periods of time, without being computationally very demanding. The 9-dimensional motion data acquired through WEAR is simple to process, easy to connect to analysis software, which allowed the fast online behavior classification that was fundamental for the experiments described in this paper.

459

460 **Open field experiment.** One-month post-surgery, mice were habituated to head-mounted

461 equipment over 2 days. On day 1, an actual or mock wireless inertial sensor (~2.5 cm H x 1 cm L

462 x 0.5 cm W with ~ 2.5-3.0 cm antennae, ~1.8 g weight) glued to the 4-position connector

463 (Harwin Inc., M52-040023V0445) was attached to the implanted receptacle connector on the

464 skull cap. Individual mice roamed freely in the home cage for 1 hour. On day 2, an actual

465 wireless inertial sensor and mono fiberoptic patchcord (200/220 µm diameter, 0.22 NA; Doric

466 DFP_200/220/900-0.22_2m_DF1.0-2FC) was attached to the skull cap via a mating sleeve.

467 Patchcords were attached to 1x2 fiber-optic rotary joint (intensity division, 0.22 NA; Doric,

468 FRJ_1x2i_FC-2FC) and mice roam freely in home cage for 1 hour. On open field recording day,

469 sensor/patchcord habituated mice were anesthetized by isoflurane, attached to equipment,

470 subjected to calibration protocol described below, and individually placed in an open field box

471 inside a sound insulated chamber. The open field box is made of 410 x 400 mm grey opaque

472 acrylic walls and a 410 x 400 mm white matte acrylic base. Individual mice were allowed to

473 behave freely inside the box for 75 minutes. The wireless inertial sensor (~1.8 g in weight,

474 WEAR wireless sensor v1.1; Champalimaud Scientific Hardware Platform) conveys motion

475 information sampled at 200 hz (set on WEAR v1.3.2 software; Champalimaud Scientific

476 Hardware Platform) to a receiver base-station (Harp basestation v1.1 or v. 1.2, Assembly v0,

477 Harp v1.4, Firmware v1.5; Champalimaud Scientific Hardware Platform), which conveys the

478 information to the experimental computer running a Bonsai script (Bonsai³⁸ editor v2.3.1) to

479 capture and record motion data and video information. Video was captured with a camera (Flea3

480 FL3-U3-I3Y3M(17450451), Point Grey Research) coupled to a 1/2" format lens (NMV-6WA,

481 Navitar).

482

483 **Calibration.** To ensure sensor stability within sessions, several approaches were employed.

484 First, a coated mating sleeve was attached to the dual optic fiber cannula that sits immediately
485 posterior to the sensor. The sleeve was thickened with black tape to a desired outer diameter such
486 that it stabilized the sensor in the anterior-posterior direction. Second, the metal pins in the 4-
487 position connector glued to the sensor were thickened with solder to stabilize their fit inside the
488 receptacle connector in the skull cap. This protects against displacement in all directions. Third,
489 stretchable black tape was wound around the base of the attached sensor and sleeve-covered
490 cannula, further protecting against shifts in sensor positioning.

491

492 To control for possible variation in sensor positioning across sessions, a calibration approach was
493 developed. Wireless inertial sensor was attached to individual isoflurane-anesthetized mice and
494 the sensor was secured with the above strategies. Next, individual mice was placed in a custom-
495 made calibration rig. The essential element of the rig is a vertical stainless-steel pole suspended
496 above a stably secured table. In the setup used, the vertical pole was fixed to the horizontal edge
497 of a vertically reversed “L” shape, stainless steel post assembly mounted on a breadboard
498 (Thorlabs). The space between the lower end of the vertical pole and the table is enough for an
499 individual mouse to slide underneath. The lower end of the vertical pole is fixed to a custom-
500 made connector that resembles the connecting end of the fiberoptic patchcord. To perform
501 calibration, individual isoflurane-anesthetized mice was securely attached to the vertical pole via
502 a mating sleeve bridging the connection to the mouse’s cannula implant. Next, replicate readings
503 of the immobilized inertial sensor were made on Bonsai. Next, mice were attached to the
504 experimental patchcord and allowed to recover in home cage for 20 minutes or until individual

mice are clearly recovered and behaviorally active. Individual mice were then placed in open-field box for experimentation.

Calibration involves rotating all accelerometer and gyroscope readings from the inertial sensor by a rotation matrix such that the final gravitational field vector of the stationary sensor, when mounted on the mouse and fixed to the calibration rig, is in a universal frame of reference whereby there is zero vertical tilt. In other words, the only non-zero acceleration is on the universal z-axis (pointing down). To accomplish this, the accelerometer pitch and roll orientation angles of the fixed stationary accelerometer were determined and then applied to calculate the rotation matrix. The rotation matrix is multiplied by the sensor accelerometer and gyroscope readings to remove the stationary vertical tilt from the sensor. To account for possible drift in gyroscope baseline over time, a daily reading of stationary gyroscope baseline was made with a mock cement skull cap attached to the sensor just before the start of each experimental day. The baseline gyroscope readings were subtracted from all gyroscope values before the rotation matrix is applied to sensor data.

Action Selection. After open field run in the grey-walled box, off-line behavioral clustering was performed on calibrated sensor data. To identify the natural action repertoire of individual mice, we quantified behavior using acceleration and gyroscope time series features in a similar fashion as described previously²¹. For the ground truth analysis, we used: 1.) Gravitational acceleration (GA) along the anterior-posterior (A-P) axis for the discrimination of postural changes - GAap. 2.) Raw sensor acceleration along the dorsal-ventral (D-V) axis to quantify movement

momentum – ACCdv. 3.) D-V axis of gyroscope to extract head head-body rotational information – GYRdv. 4.) Total body acceleration to differentiate resting state from movement.

Total body acceleration (TotBA) was defined as:

$$\text{TotBA} = \sqrt{\text{BA}_{\text{ap}}^2 + \text{BA}_{\text{ml}}^2 + \text{BA}_{\text{dv}}^2},$$

where BA_{ap}, ml and dv represent the body acceleration of the anterior-posterior, medio-lateral and dorsal-ventral axis, respectively. We calculated each individual BA component by median-filtering the raw acceleration signals followed by a fourth-order Butterworth high-pass (0.5Hz) filter. For the gravitational acceleration (GA) axis, the BA components were subtracted from the median filtered raw signal axis.

All four time series features were binned into non overlapping 300 ms long window segments²⁶. The values of each bin and per feature were then discretized, using fixed thresholds, producing a summary distribution of each segment. For GA_{ap} and ACCdv we used 10 equal size threshold values, plus two added bins between the limits and infinity to capture an approximated distribution of values within each window bin. For GYRdv we used 5 thresholds (0, ±50, ±100) to discriminate left and right turns. For TotBA, a single threshold was used to separate moving from resting. The threshold was kept constant for all experiments and was set to the average value separating the bimodal distribution of logTotBA (natural logarithm of TotBA feature). For each 300-ms window segment we get four resulting histograms, one for each feature. The feature

histograms were individually normalized to obtain probability distributions and used to calculate the pairwise similarities between segments.

We used the "earth mover's" (EM) distance as a measure of similarity²⁵:

$$S = -(dEM/4)^2$$

where dEM is the sum of the normalized EM distances for the 4 features (GAap, ACCdv, GYRdv and TotBA) defined above. The bin normalizations constrain S values within the range [-1,0], specifically, -1 and 0 define the maximum dissimilarity and identity between the two probability distributions, respectively. Finally, to produce a continuous unbiased classification of behavioral states, the similarity measures were clustered using affinity propagation²⁰, with the preference parameter set to the minimal value of the similarity matrix; this particular value was used for its stable number of behavioral clusters within its range.

Using the behavioral clusters identified by affinity propagation clustering of the grey open field behavior¹³ as a ground truth for the true identity of each 300 ms histogram, we were able to simulate and evaluate the precision with which the Earth Mover's Distance (EMD) metric^{21,25} could be applied for cluster matching online. Notable difference between the EMD metric used here is the use of the 4 features mentioned above rather than the 3 features used previously²¹, as well as the multiplication of the similarity score by -1 such that the range of possible scores from maximal identity to dissimilarity is 0 to 1, respectively. Although the EMD cluster matching outcome correlates strongly with affinity propagation clustering, some false positive and false

negatives may occur. Several filters were set to optimize cluster selection for reinforcement: 1.) We selected for clusters that show low false positive rate ($<5.5\%$) and below the 60th percentile false positive rate amongst all clusters per animal. 2.) We selected against clusters with high false negative rates ($> 90^{\text{th}}$ percentile of clusters per animal). 3.) We selected against clusters that tend to be performed serially within a short time interval. We calculated the probability that a target cluster or its top 5 most similar clusters (determined by EMD score) would reappear 3-18 seconds after the first occurrence of the target cluster. Clusters that tend to be repeated either by itself or have a high probability of having similar clusters appear within this 15 second window ($> 90^{\text{th}}$ percentile for median and range of probabilities of cluster appearing in window) were removed from selection pool. 4.) We filtered against clusters whose matching by EMD would be more sensitive to anterior-posterior shifts of the inertial sensor (although we already protected against this possibility with the safeguards above) ($> 90^{\text{th}}$ percentile for percent deviation from original cluster matching after shifts of accelerometer reading in the anterior or posterior direction). For each cluster, percent deviation is calculated first by summing up the total absolute cluster matching changes from original cluster matching data in the anterior and posterior shifted datasets. Next, the sum of deviation in the two altered datasets is divided by two and then divided by the total of cluster calls from the original dataset, and multiplied by 100 to get percent deviation from original cluster matching result. 5.) We selected for clusters that show fully accelerating movement (cluster exemplar value of less than the maximum value of 1 in the body acceleration feature bin of histogram). To choose dissimilar clusters per animal, an algorithm was written filtering clusters of each animal's repertoire based on the feature histogram values of each cluster's representative, or exemplar. Thresholds were set along the GAap and GYRdv features to divide cluster exemplars based on the distribution of values within these feature

histograms. For each repertoire, all histogram values from all cluster exemplars are pooled to create a pooled histogram. The range of bins with non-zero values for each feature are identified. The algorithm then filters cluster exemplars in the repertoire for non-zero values in the high, medium, low, or high+low value bins. For example, action A identification occurs by selecting for a cluster exemplar with median counts falling in the high GAap and GYRdy value bins. action B would then be selected by filtering for an exemplar with median counts falling in the low GAap and GYRdy value bins. This results in actions that are highly dissimilar. For example, EMD similarity scores comparing action A to action B almost always, except for 1 Chr2-YFP animal, fall in the more dissimilar end of a distribution of scores created by comparing action A to all actions in each animal. Hereafter, clusters will be referred to as actions.

Closed-Loop Optogenetics. For close loop optogenetics, a computer running a Bonsai script captured and recorded wireless sensor motion data and video information as described above in grey-walled open-field experiment. Here, data is also streamed to a custom MATLAB code which analyzes action composition changes over the course of action reinforcement, we used the EMD metric²¹ to label individual 300 ms motion histograms with an action ID. For each arriving 300-ms segment we calculate the EMD distance between each cluster exemplar (or representative) of the ground truth cluster library from the grey open field behavior recording. The motion features histogram is assigned to the action for which comparison with the exemplar gave the lowest EMD score (most similar to target) amongst all comparisons. Decision making for stimulation has a range of 35-55 ms time gap between action performance and sent decision for stimulation. To trigger optogenetics, a Multi-Pulse Width Modulation (PWM) generator (Harp Multi-PWM Generator hardware v1.1, Assembly v1, Harp v1.4, Firmware v1.1; Harp

Multi-PWM Generator software v2.1.0; Champalimaud Scientific Platform) converts each decision to trigger laser into electrical signals for 15 light pulses of 10 ms pulse duration at 25 Hz, with each train of pulses occurring over 600 ms and at 25% duty cycle. The multi-PWM signal is passed through a 12 V, 7.2 W amplifier (Champalimaud Scientific Platform) and fixed frequency driver (Opto-electronic, MODA110-D4-30 (2001.320220)) to control the activities of a 473 nm, blue low noise laser (Shanghai Dream Lasers Technology, Co, Ltd. SDL-473-200T), which was sent through an acousto-optic modulator (Opto-electronic, MTS110-A3-V1S (1001 / 330433)). The laser component that is modulated is then reflected by a mirror and funneled to a mono fiberoptic patchcord, which is then coupled to a commutator. The output laser is then passed through a dual-optic fiber patchcord and connected to the implant cannula. Power adjustment out of the tip of patchcord was made so that ~5mW was emitted from each end of the dual optic fiber cannula. To ensure common time stamps from different channels, a clock synchronization device (Harp Clock Sync v1.0; Champalimaud Scientific Platform) was performed between the basestation and multi-PWM device.

Single action sequence selection. Mice were placed in a white open field box for closed loop reinforcement protocol. Individual mice were subjected to a single session of protocol each day, with sessions following each other on consecutive days. The white open field box is made of 410 x 400 mm white matte acrylic walls and a 410 x 400 mm white matte acrylic base. To acquire baseline behavior, individual mice were allowed to behave freely inside the box for 30 minutes on the first action A selection session. Closed loop reinforcement by blue laser stimulation of VTA DA neurons were made available for 60 minutes. 90 minutes of closed loop reinforcement were made available for individual mice during sessions 2 and 3. For session 4, an

extinction protocol was carried out comprising of 20-minute maintenance of reinforced behavior with laser availability, followed by 60 minutes of extinction of reinforced behavior without laser availability, followed by 20-minute re-acquisition of reinforced behavior with laser availability. To select for action B, a repeat of the protocol described above for action A was performed starting on the day following extinction protocol of action A. Upon completion of the reinforcement and extinction protocols for action B, a contingency degradation protocol was performed comprising of 20-minute maintenance of action B with laser availability, followed by 60 minutes of contingency degradation of reinforced behavior by triggering laser randomly, followed by 40-minute re-acquisition of reinforced behavior with laser availability for action B performance.

Photometry experiment. One-month post-surgery, mice were habituated to head-mounted equipment for 2 days. On day 1, habituation was made to wireless inertial sensor as described above. On day 2, a multi-fiber bundled patch cord (3 fiber bundle, 400/440 μm diameter for a maximum of inner diameter at 900 μm , 0.37 NA, 3.5 m long, 1.25 mm fiber tip diameter, low-autofluorescence; Doric, BBP(3)_400/440/900-0.37_3.5m_FCM-3xMF1.25_LAF) was attached to individual mice in addition to the wireless sensor and optogenetic patchcord. Individual mice were allowed to habituate to the equipment for 1 hour in its home cage. On photometry recording day, mice were subjected to 30 frames per second photometry recording (Neurophotometrics), with 75-150 μW 560 nm LED illuminating rDA1m, and equivalent closed loop optogenetic parameters described above were used. To test for DA release in the context of closed loop optogenetic setup, an average of 30 hits of blue light were delivered randomly within the span of 30 minutes. To evaluate DA release in the context of food reward, mice were placed on food

deprivation protocol and kept within 85% of original weight. Mice were placed in an operant chamber with a nosepoke linked to a lick detector (PyControl). Each lick detection triggers dispensing 2 μ l 10% sucrose. Since animals tend to accidentally trigger lick detector at the beginning of sessions, between 40-50 sucrose dispensing events were gathered per animal and rDA1m activities associated with the last 35 rewards of the session were used for analysis.

Two action sequence selection. Two action sequence selection occurs as follows: after sensor/patchcord habituation and grey open field behavior recording, offline behavioral clustering and action filtering were performed as for single action selection. For each animal, median time intervals between all possible pairs of actions during open field were calculated as described above. Across animals, T1/T2 pairs with median T1→T2 interval values varying between 2 and 10 seconds, and with the feature of going from a head down(T1) to a head up(T2) movement, were chosen for reinforcement.

On the first reinforcement session, a 30-minute baseline was taken when laser stimulation was not available for reinforcement. Laser became available for reinforcement in all subsequent sessions until extinction experiment. During reinforcement periods, when closed-loop system detects performance of the proximal action (T1) of interest, the algorithm enters a state where laser is triggered upon performance of the distal action (T2), regardless of the amount of time that has elapsed between the latest T1 and T2. On Session 1, 60 minutes of laser availability was given while in all subsequent reinforcement sessions, 90 minutes of laser availability was given.

Histology and Immunohistochemistry. After behavioral sessions were completed, mice were deeply anesthetized with isoflurane and perfused transcardially in PBS and then 4% PFA/PBS. Dissected brains with skulls attached were perfused in 4% PFA in PBS at 4 degrees Celsius overnight. The next day, brains were rinsed 3 times in PBS. Next, brain regions including VTA and implants were sectioned by vibratome into 50 or 100 μ m slices. Slices are then subjected to immunohistochemistry using the reagents below. Standard immunohistochemistry protocols were applied to stain for the following reagents - Rabbit anti-GFP 488 conjugate (1:1000; Molecular Probes A21311). Mouse Anti-TH (1:5000; Immunostar Th 22941) with Goat Anti-Mouse - IgG (H+L) Highly cross-adsorbed secondary antibody - Alexa Fluor647 (1:1000; ThermoFisher, A-21236), DAPI (1:1000 of 20 mg/mL stock; Sigma, D9542).

Imaging. Zeiss Axio Imager M2 microscope was used to acquire brain section pictures. 10x tiled images were taken through the relevant fluorescent channels. The M2 is equipped with a fast Colibri.7 LED illumination for excitation of fluorophores. Images are captured with a high-sensitivity monochromatic sCMOS camera (Hamamatsu Orca Flash 4.0 v2). The objective used for the images is a ZEISS Plan-ApoChromat 10x/0.45, which allows to resolve up to 577 nm when using a wavelength of observation of 520nm and it is fully corrected for chromatic and spherical aberrations. Implant locations were determined using standard mouse atlas³⁹.

Single action selection analyses. For target action frequency analysis, we analyzed frequencies within 25-minute windows at 4 time points: Baseline (before first reinforcement trigger), Early (after first reinforcement trigger in Session 1 (action A) or 5 (action B)), Mid (after 2-minute mark in Session 2 (action A) or 6 (action B)), Late (after 2-minute mark in Session 3 (action A)

or 7 (action B)). For 3D action repertoire plots, baseline normalized frequencies were plotted and actions whose time series include NaN or Infinity values were discarded from the plot. (Plotted actions: 509 of 514 actions, 15 ChR2YFP animals (action A); 427 of 443 actions, 13 ChR2YFP animals (action B); 355 of 356 actions, 10 YFP animals (action A); 341 of 356 actions, 10 YFP animals (action B)).

Three parameters were assessed for rapid behavioral adaptation following cumulative closed loop reinforcements: latency between Target A triggered reinforcements, Target A frequency and average behavioral similarity to Target A. To calculate the latency parameter, the average latency between 10 consecutive Target A triggered reinforcements following a specified number of cumulated reinforcements were taken and then normalized by the average latency taken over the final 10 baseline Target A instances that in simulations would have triggered reinforcement. To calculate the frequency parameter, the frequency of Target A triggered reinforcements over the course of 1 minute following a specified number of cumulated reinforcements were taken and then normalized by frequency of the final 10 baseline Target A instances that in simulations would have triggered reinforcement. To calculate the behavioral similarity parameter, the average behavioral similarity (EMD score) to Target A between 10 consecutive Target A triggered reinforcement events following a specified number of cumulated reinforcements were taken and then normalized by the corresponding value taken over the final 10 baseline Target A instances that in simulations would have triggered reinforcement.

rDA1m Fiber Photometry Analyses. To evaluate DA release in the context of food reward, the delta F/Fo signal was plotted for rDA1m signal aligned to lick detection/reward trigger. The

baseline F_0 value was taken as the median rDA1m raw fluorescence signal of the 10 time points (333.33 milliseconds) preceding the trigger event. To test whether DA release is triggered in the context of the closed loop system, the activity of the rDA1m sensor was quantified. Delta F/F_0 was calculated by subtracting baseline value from each fluorescent rDA1m value of a smoothened time series (smooth function, default moving average filter, MATLAB), and then dividing the outcome by the baseline value. To account for control ChR2-independent effects, the average delta F/F_0 trace of ChR2-YFP animals were subtracted from the corresponding average trace of YFP animals, giving the differential delta F/F_0 used for the plots. The standard deviation of ChR2-YFP minus YFP curves were obtained by taking the square root of the sum of squared variances of ChR2-YFP and YFP delta F/F_0 curves.

Categorizing behavioral actions by temporal dynamics. To categorize behavioral actions by temporal dynamics, moving mean of action counts was used as input. Various window sizes were examined; 2.5-minute windows moving at 300 ms steps were found suitable for analyses. The baseline frequency (f_0) was the average of 5 minutes of moving mean data preceding the first reinforcement event. Early frequency rate (f_1) was the average of 30 minutes moving means immediately following the first reinforcement event. Mid- and Late frequency rates were taken from Day 2 (f_2) and Day3 (f_3), respectively. f_2 and f_3 rates were calculated from the beginning 30 minutes period after moving windows has accumulated enough bins (2.5 minutes) following the start of the session. Significant positive modulation above baseline was judged if in 500 consecutive moving windows (2.5 minutes period) in Early/Mid or Late stages the frequency rate of all bins were greater than the 99th percentile bin of baseline frequency. Significant negative modulation below baseline was judged if in 500 consecutive moving windows (2.5-minute

period) in Early/Mid or Late stages the frequency rate of all bins were less than or equal to the 5th percentile bin of baseline frequency. Actions that showed both significantly positive and negative modulation at Early/Mid or Late stages when compared to baseline were delegated to positive modulation group. For figure plotting, time-course median frequencies of action dynamic types were downsampled 10-fold. To investigate the relationship between target similarity and frequency, two approaches were taken. To perform multiple comparison statistics, actions were binned by their percentile ranking in terms of similarity to target (EMD). This is because action distribution based on raw EMD binning was not even. Percentile binning allowed for even distribution of actions amongst the groups. To examine the distribution of action dynamic type frequencies in terms of target similarity, a binning by raw EMD score (0.5 score binwidth) was used because this allowed for clear visualization of the relationship between target similarity and frequency. Alternatively, percentile binning of EMD score was also used and gave similar trends.

Criterion for action dynamic types. Action dynamics were grouped according as follows: 1.) Increasing actions showed significant increase in f0 to f1/2 and f1 to f2/3 comparisons and showed either significant increase or unchanged frequency in f1/2 to f3 comparisons. 2.) Sustained actions showed significant increase in f0 to f1/2 comparisons, and unchanged frequency in f1 to f2/3 and f1/2 to f3 comparisons. 3.) Transient actions showed significant increase in f0 to f1/2 comparisons, and significant decrease in f1/2 to f3 comparisons. 4.) Decreasing actions showed significant decrease in f0 to f1/2 and f0 to f3 comparisons. 5.) Other actions were all remaining actions that did not fall in the above groups. In the main figure only dynamic subtypes with more than 10 members are shown.

Extinction analyses. 10 minutes portions from different time windows along the extinction protocols (Session 4 for action A and Session 8 for action B) were chosen. Early maintenance (M^1) starts from the first instance of target action performance in the session. Late maintenance (M^2) is the portion preceding the first performance of target upon extinction. Early extinction (E^1) begins at the first instance of target performance upon extinction. Late extinction (E^3) is the portion preceding the first performance of target upon re-acquisition. Mid extinction (E^2) begins at the midpoint between the starts of E^1 and E^3 . Early re-acquisition (R^1) starts at the first performance of target upon re-acquisition condition. Late re-acquisition (R^2) is the final portion of the extinction protocol.

Action burstiness analysis. To evaluate action burstiness, or dispersion, we used Fano factor (variance/mean) as a measure. A survey of moving mean frequencies of reinforced actions across animals suggest that actions are more dispersed during the extinction phase, but the timescale with which this may occur is variable. To identify a suitable timescale to detect dispersion across reinforced actions, we screened a range of window sizes (600 ms to 5 minutes windows in 600 ms steps) with which to calculate moving window frequencies, and then calculate Fano factor in varying time segments. We chose a moving window of 15 seconds (50 x 300 ms action units) to construct moving mean frequencies. This window size consistently gave decreased Fano factor in baseline vs. maintenance session across animal, a result that would be expected as reinforcement led to stable target action performance.

Single action reinforcement, inter-target, and inter-action interval analyses. To quantify inter-target action intervals, the median amount of time that transpired between the start of successive target actions over the course of a time window was calculated. The time periods analyzed were: 1.) Baseline from the start of Day 1 (Sessions 1 and 5 for action A and B, respectively) until the first reinforcement event. 2-4.) Days 1 to 3 reinforcement. For reinforcement periods, behavior from the start of the first reinforcement event of that session until the end of session were analyzed. We considered the possibility that including the time interval between consecutive repeating of target actions (resulting in an inter-target action interval of 300 ms) would greatly affect the result. To test this, we removed values collected from consecutively repeating target actions. However, this did not affect result interpretations. Thus, we included intervals from consecutively repeating target actions in the presented analyses. For single action reinforcement, the median amount of time between the closest occurring action of interest and target action was calculated for both pre-target and post-target intervals.

Multinomial logistic regression predicting action dynamic types. To test whether intrinsic and baseline action properties are predictive of classifiable action dynamics during single action reinforcement from naïve state, two factors were considered. The factors are Earth Mover's Distance (EMD) similarity of action to target and median time interval of closest action of interest prior to target appearance at baseline condition.

To perform multinomial logistic regression, data from both dependent variables were log-transformed after addition of a constant value of 1. Transformed data were tested for collinearity

by examining scatter plots, Pearson’s correlation coefficients, Variance Inflation Factors (VIF) and condition indices. The two variables showed some correlation, but the coefficient value was not above typical thresholds^{40,41} and direct collinearity diagnostics did not show significant collinearity (Pearson’s correlation: $0.67 < 0.8^{40}$, VIFs: $1.82 < 5 \cdot 10^{42}$, condition indices: $6.6 < 10 \cdot 30^{43}$). Multinomial logistic regression was performed using MATLAB functions `mnrfit` and `mnrval`. Non-Target A actions from all animals from reinforcement of action A were included except those whose reinforcement dynamics were previously classified as “Other” types ($n = 30$ actions from a total of 514 actions, 15 ChR2-YFP animals). Decreasing dynamics type actions were used as the reference group. Model accuracies were assessed using a 20-repeat, 10-fold cross-validation approach for a total of 200 unique models for Real data, and 10,000 unique models from 50 shuffled datasets.

To evaluate multinomial logistic regression, the deviance measure was used to judge model fitting. Model performances were judged by area under precision-recall curve as this criterion is suitable for imbalanced categories in the data³⁵. A model containing both dependent variables was found to outperform that of any single variable, even after consideration for penalties for an extra factor (Akaike Information Criterion). The lack of significant collinearity between dependent variables was supported by the stability of two relevant parameters, beta-coefficient directions and significant p-values, across 200 cross-validation models and single- and double-factor regression conditions (See Supplementary Information for tables).

Dopamine retrospective window analysis. To analyze whether DA reinforces actions proximal to target, baseline rates of action transitions occurring close to reinforced action were examined.

First, a matrix tabulating 300 ms action counts from 2.4 seconds before to 2.4 seconds after each theoretical target-triggered laser stimulation (600 ms in length) during baseline condition was constructed. Next, all possible 600 ms action transitions (ex. $X \rightarrow Y$) for each animal were then counted using the above matrix, resulting in an action transition type (row) vs. time bin (column) matrix where the counts of each action transition type occurring in specific 600 ms transition windows (ex. $X \rightarrow Y$) were recorded (sum across rows). This will be called the count matrix. Next, the relative enrichment of each action transition type in a specific transition window against all transition windows was calculated by dividing the action transition count matrix by the total number of action transitions per type (probability across rows). Next, action transition probability within a sliding 1.2 second transition window (containing a total of three action transitions) relative to surrounding temporal environment (3.6 seconds) was derived by subtracting the total number of action transitions per type within the surrounding 3.6 second window from the total number of action transitions per type within the 1.2 second sliding window of interest. This will be called the differential probability matrix. Next, action transition types that showed greater than a threshold of 0.001 relative probability within sliding 1.2 second windows of interest over the corresponding surrounding windows were filtered and kept for the next step. Next, for each sliding 1.2 second window, the count matrix from above was analyzed to select for action transition types that occurred between 2 to 6 times during the 30 minutes baseline period (0.067 to 0.2 action transitions per minute). The count range was chosen to filter out single events while selecting for action transitions with low initial frequencies over the baseline period and analysis time range. Since the range of probabilities of specific action transition types could vary greatly between different sliding 1.2 second windows, filtering as above also balances the distribution of action transition probabilities amongst all action transition

types analyzed across sliding 1.2 second transition windows. The above process results in a list of action transition types enriched for each sliding 1.2 second transition window, and baseline normalized frequencies of these action transition types upon reinforcement in subsequent sessions were calculated. Note that baseline normalized frequencies were calculated from all occurrences of specific action transition types, regardless of their time distance in relationship to target occurrence. Baseline normalized frequencies of individual action transition types were averaged within animals and the means between animals are averaged to produce animal-balanced results. Identical data trends and conclusions could be reached even if baseline normalized frequencies of all action transitions were used for analyses.

Two action sequence experiment analyses. Two action sequence frequency was quantified in terms of laser triggers per minutes. To assess learning across animals, the baseline frequency was subtracted from frequencies of all reinforcement sessions. A criterion baseline subtracted frequency of 3.2 triggers per minute was set after considering the range of baseline subtracted frequencies observed in the open field and reinforcement sessions all animals. The criterion is set such that it is > 20 % above the highest baseline-subtracted frequency value seen at open field condition. The criterion point consistently falls above the open field frequencies of all animals and marks the rising phase of all reinforcement frequency curves.

T1→T2 intervals were quantified as the time distance between the end of the latest distal action (T1) and the end of the proximal action (T2) that triggers laser. T2→T1 intervals were quantified as the time distance between the end of T2 that triggers laser and the end of the next closest T1.

To produce equivalent measures in open field and baseline conditions, laser trigger events were simulated by scanning across the data as if reinforcement was available.

Significance testing was performed on 14 of 15 ChR2-YFP animals that reached criterion frequency (ChR2-YFP Criterion). The lone animal that did not reach criterion frequency was removed because the T1→T2 median interval was still very high after session 10. This animal was subsequently subjected to single action reinforcement protocol to assess its ability to learn T1 and subsequently T2. Next, the animal was again subjected to T1→T2 reinforcement protocol. These results indicate that this animal was capable of action learning for both T1 and T2 separately, and for T1→T2 sequence after learning of each individual action.

Reinforcement sessions for the 14 ChR2-YFP animals that reached beyond criterion frequency continued until the T1→T2 interval has been decreased to below at least a median of 2 seconds. As YFP animals do not decrease the T1→T2 median interval over sessions, we stopped reinforcement at session 20.

Two action sequence extinction. Extinction session begins with a 25-minute maintenance period for two action-sequence reinforcement, followed by a 40-minute extinction period when laser was inactive, followed by a 25-minute re-acquisition period whereby reinforcement was made available again. To quantify performance for plotting, frequency was calculated over 5 minutes bins and then normalized to the last 5 minutes bin of the maintenance condition. For significant testing, raw frequencies were analyzed at the last 5 minutes of maintenance, extinction, and re-acquisition conditions.

914

915 **Two action sequence refinement.** To measure refinement for T1 and T2 in the two-action

916 sequence, actions that were uniquely related to one but not the other were identified. Actions

917 performed by each animal in their open field repertoires were ranked by their EMD similarity

918 scores to T1 or T2. The top-12 actions (within action repertoires ranging between 30-40 actions)

919 most similar to either T1 or T2 were identified. Actions common to both T1 and T2 in these lists

920 were removed, leaving actions uniquely similar to T1 or T2. We required at least 3 non-target

921 actions to be uniquely related to each of T1 and T2. One of the animals did not meet this

922 requirement, because less than 3 actions were uniquely similar to each of T1 and T2 when

923 considering the top-12 actions related to T1 or T2. For this animal, we relaxed the stringency by

924 considering actions that uniquely belong as the top-9 actions most similar to either T1 or T2. We

925 took the median target-normalized frequency of these uniquely similar actions to T1 or T2 as the

926 refinement index. A refinement index of above or around 1 indicates little to no refinement of

927 uniquely related actions to target. Refinement index below 1 indicates refinement relative to

928 target; the lower the score the more refinement. Refinement curves were smoothed using the

929 Savitzky-Golay filter to improve visualization of trends. To better compare the progress of

930 refinement between T1- and T2-related actions, refinement indices were scaled such that the

931 minimum value amongst all sessions for individual animals would be zero and target-normalized

932 median frequency of 1 would remain at a scaled value of 1.

933

934 **Relationship between T1→T2 interval and sessions to criterion frequency.** To describe the

935 trend in a T1→T2 interval vs. sessions to criterion frequency scatter plot, non-linear sigmoidal fit

936 was tested against a 4th order polynomial fit. A linear fit was also tested. Sigmoidal fitting gave

the best result. The same fitting was tested for T2 → T1 interval vs. sessions to criterion frequency, but the fit was poor and midpoint was unstable. For the T1 → T2 sigmoidal curve, half-maximum was 2.59 sessions to criterion frequency and midpoint was 4.69 seconds of open field median interval. The half-maximum value was used to divide ChR2-YFP animals into slow (above half-max) and fast (below half-max) learners.

Differential refinement analyses. The difference in area between T1 and T2 scaled refinement curves over sessions was used to assess the relative refinement status between T1 and T2 over sequence learning. The difference in areas were summed up using the trapezoid method across sessions until the session when both T1 and T2 has or had reached minimal scaled refinement. Next, the relationship between open field median interval and average difference in area under T1 – T2 refinement curves per session was tested. Linear regression proved most suitable for fitting (Goodness-of-fit: $R^2 = 0.66$). The fit for T1 → T2 linear line was $y = 0.1893x - 0.7050$. Slope was significantly non-zero ($p = 0.0004$). The same fitting was tested for T2 → T1 interval vs. difference in area under T1 – T2 refinement curves per session ($y = 0.00736x + 0.1356$), but the fit was poor, and goodness of fit was low (Goodness-of-fit: $R^2 = 0.07$). The slope was not significantly non-zero ($p = 0.7063$).

Starting Point identification for evaluating progression of differential T1/T2 refinement. To more precisely examine whether proximal action (T2) refinement precedes that of distal action (T1) in Slow Learners, it was important to consider refinement progression of T1 relative to T2. To rule out any bias towards proximal refinement because of initial bias towards proximal T2 refinement, a specific session was chosen as a Starting Point for analysis for each animal. This

Starting Point is defined by an early session in which T1 and T2 were relatively similar in refinement levels or when the distal action T1 was more refined than proximal T2. To identify these Starting Points, a scan was made retrospective from the session for which the T1→T2 time interval is close to final value (less than or equal to a median of 3 seconds). Using this approach, we identified earlier sessions in which distal T1 refinement was equal to or greater than proximal T2 (T2 – T1 refinement curve area less than or equal to 0). The latest such session was set as the Starting Point for analysis. If at no point early in learning did an animal have a session where proximal (T1) action is most refined relative to distal (T2) action, an early session of closest T1 and T2 refinement was used as the Starting Point. The initial T2-T1 refinement curve area difference calculated from the Starting Point to next session was subtracted from all T2-T1 area differences calculated in subsequent sessions. This value is called the Starting Point subtracted refinement difference. This made it possible to clearly track the change in relative refinement of distal(T1) vs. proximal(T2) actions over time (Values above zero indicate T2>T1 refinement, and values below zero indicate T1>T2 refinement). To identify the Turning Points for each animal, sessions carrying the local maximum value of the Starting Point subtracted refinement difference were identified for each animal. To calculate Starting Point subtracted refinement, scaled refinement values from sessions of interest were subtracted from that of the Starting Point session defined above.

Odds ratio analysis. For odds ratio calculation, the total amount of open field → Turning Point session (second of two consecutive sessions used to calculate the refinement difference at Turning Point as mentioned above) and Turning Point → session of criterion frequency median interval changes were summed up for T1→T2 and T2→T1 intervals, respectively. Next, the

proportion of total interval change stemming from the open field condition→Turning Point period, and from Turning Point→session reaching criterion frequency period, were calculated. Next, the proportion of open field→Turning Point interval change was divided by the proportion of Turning Point → session reaching criterion frequency period interval change for T1→T2 and T2→T1 interval types, respectively. This gives the odds ratio.

T1 probability rank and refinement change across time bins from T2 trigger. For every actual or simulated trigger for T1→T2 performance, the first occurrences of every action before or after T2 triggers were counted at specific 300 ms time bins for up to 6 seconds before and after T2 trigger. This was done for the specific conditions of baseline, Starting Point, Turning Point, session passing criterion frequency, and last session. The probability of an action occurring at a specific 300 ms time bin was calculated for all actions in the repertoire, and the values were used to determine probability rank in terms of percentiles (100 percentile is most probable action relative to all actions at a specific 300 ms time bin). To assess total T1 probability rank change within 0.3-1.8 or 2.1-3.6 second time bins, the area under the curve was determined and values were normalized by subtraction from each animal's corresponding baseline values. Refinement change was calculated by first taking the median probability rank of actions most uniquely related to T1 at varying time distances before or after T2 trigger. This value is then normalized by T1 probability rank to arrive at a refinement index. The area under the curve was determined and values were normalized by subtraction from each animal's corresponding baseline values. Decreasing values from Starting Point indicate increasing refinement.

Statistical Analysis:

Standard statistical analyses were performed on Prism (GraphPad Software, Inc.) and permutation/bootstrap analyses were performed on MATLAB (MathWorks Inc.). To determine appropriate tests for comparisons, datasets were assessed for normality using Anderson-Darling, D'Agostino & Pearson, Shapiro-Wilk and/or Kolmogorov-Smirnov tests whenever applicable. Datasets were also visualized for normality using QQ plots and assessed for equal variance by examining the Residual plot (Residuals vs. Predicted Y). Parametric or non-parametric tests were chosen based on the combination of these analyses. Data were transformed logarithmically (with or without addition of a constant prior to transformation) whenever it was appropriate to promote normality and equal variance. Unless specified, sphericity was not assumed, and Geisser-Greenhouse correction was applied in all ANOVA tests. The appropriate post hoc multiple comparisons tests were applied to compare between the means of specific conditions wherever applicable. Significance was set at $\alpha = 0.05$. For bootstrap analysis, significance was determined by asking whether the original target action mean Fano factor was greater or less than the 95% confidence interval of the bootstrap distribution. Permutation test was applied in the comparisons between regression models because of the large sample size discrepancy between groups. Bonferroni p adjustment was used to account for multiple comparisons in this case. For detailed description of statistical procedures please refer to Supplementary Information.

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Author Contributions:

J.C.Y.T and R.M.C. designed the study, interpreted results and wrote the paper. J.C.Y.T. performed and analyzed experiments. J.C.Y.T, V.P., F.C. designed close loop optogenetic system. J.C.Y.T., F.C. and A.S. executed assembly of the closed loop optogenetic system. F.C. and A.S. designed and assembled software and hardware. A.S. designed and assembled wireless inertial sensor and hardware. A.K. contributed Earth Mover's Distance code and was involved in early conceptions of the closed loop system. J.A.d.S., F.C. and A.S. designed and assembled the WEAR system. R.M.C. supervised the project. All authors edited the paper.

Competing Interests: F.C. is the Director of Open Ephys Production Site.

1052

1053 **Additional Information:** Supplementary Information is available for this paper.

1054

1055 **Code availability.** MATLAB (MathWorks) codes used for data analysis are available from the
1056 corresponding author.

1057

1058 **Data availability.** Source Data are available from the corresponding author upon reasonable
1059 request.

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Figures and Figure Legends:

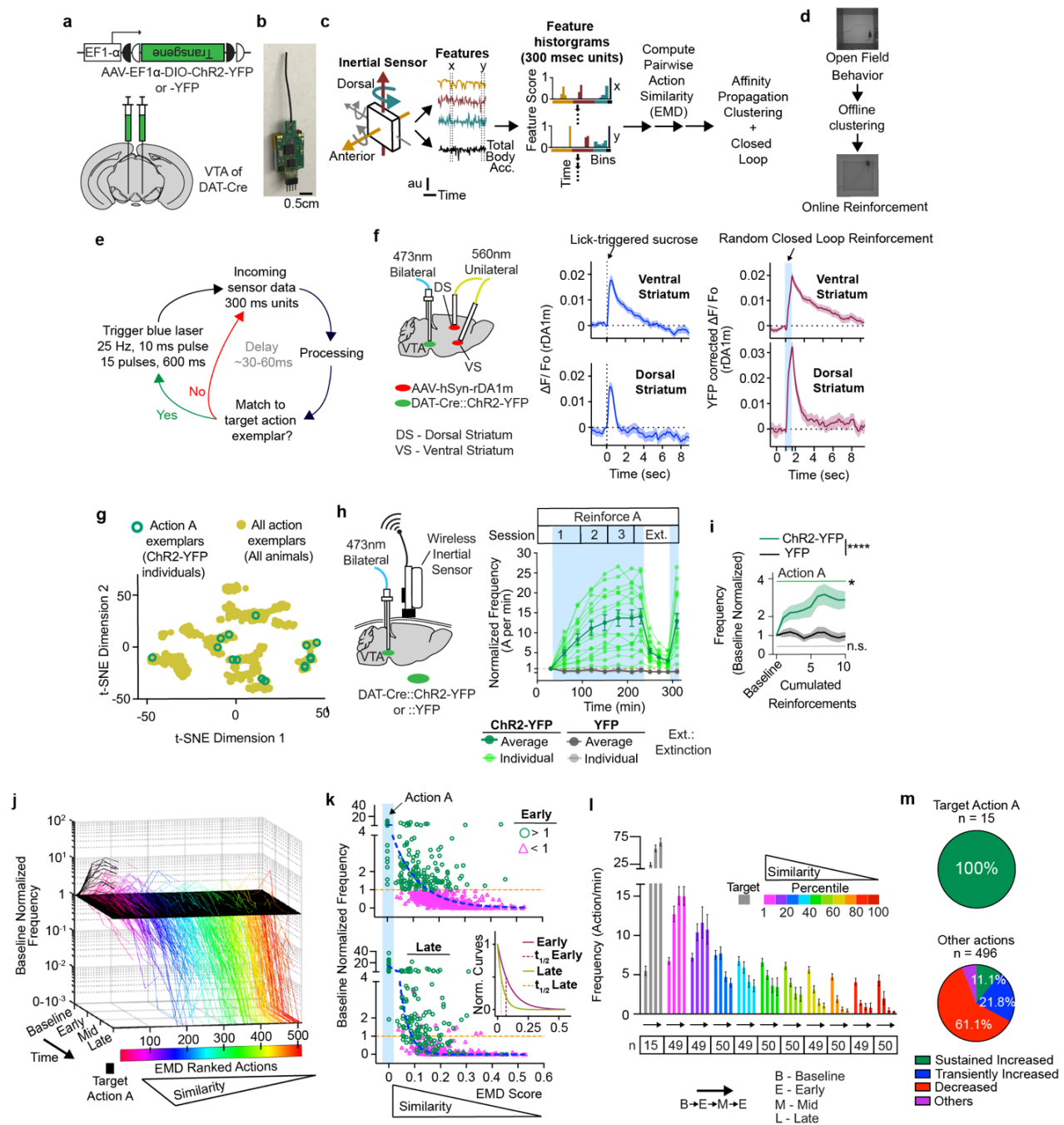


Fig. 1. Learning of a single action from the naïve state as mediated by closed loop

optogenetics. **a**, Injection scheme. **b**, Wireless inertial sensor. **c**, Sensor data processing. **d**, Open field behavioral clustering and action reinforcement. **e**, Closed loop schematic. **f**, Dopamine release in dorsal and ventral striatum (n = 70 sucrose rewards, 2 ChR2-YFP mice; n = 66 and 65 random

stimulations, 2 ChR2-YFP and 2 YFP animals, respectively). Plots were mean, S.E.M. **g**, Action A exemplar locations in behavioral space. **h-m**, ChR2-dependent reinforcement of Action A ($n = 15$ ChR2-YFP animals (green). $n = 10$ YFP animals (grey)). Plots were mean, S.E.M. **h**, Left: Head-mount setup. Right: Light green/grey lines represent individual ChR2-YFP/YFP animals, respectively. **i**, Rapid increase in target action performance in response to close-loop reinforcements. Significant Time x Group Interactions (Supplementary Information). Plots were mean, S.E.M. **j**, Evolution of pooled behavior repertoire ($n = 509$ actions, ChR2-YFP mice) across learning. **k**, Early/Late cross-sectional views of (**j**) (Early: baseline normalized frequency >1 , green circles, < 1 , magenta triangles). Blue dashed lines - single phase log decay fits. Bottom inset graph shows Early/Late fitted lines normalized to 1 at EMD=0. **l**, Raw frequencies across learning and target similarity percentile groups. Plots were mean, S.E.M. Two-way mixed effects statistics in Supplementary Information. **m**, Pie chart summarizing distribution of actions according to their dynamics within reinforced Action A (left) or other actions (right). Asterisks: **** $p < 0.0001$. *** $p < 0.001$. ** $p < 0.01$. * $p < 0.05$. n.s. – not significant. See Supplementary Information for statistical/sample details.

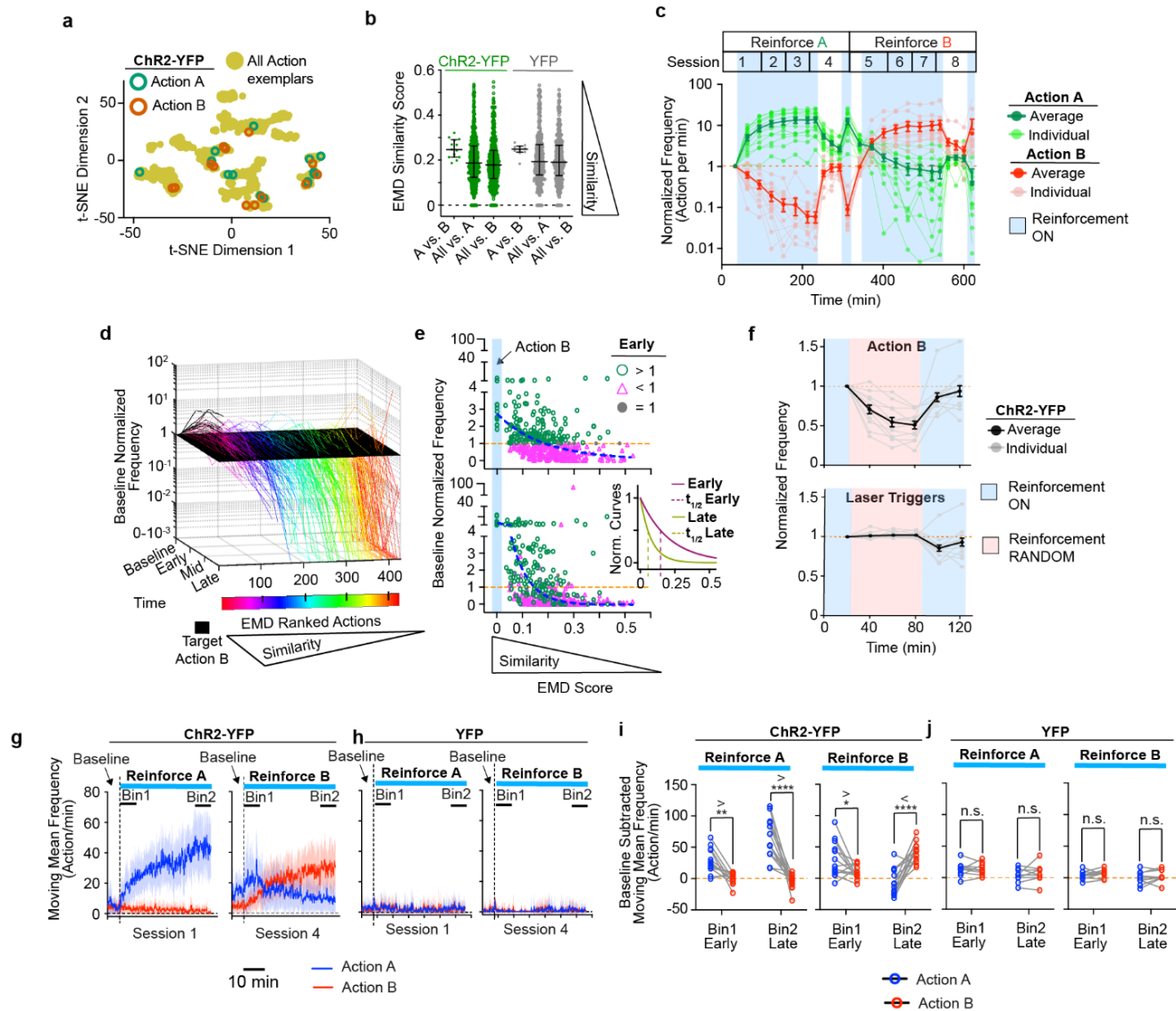


Figure 2. Transitioning from learned action to reinforcing new action. a-j, Animals reinforcing for Action A ($n = 15$ ChR2-YFP) to Action B ($n = 13$ of 15 ChR2-YFP). $n = 10$ YFP animals. **a**, Action A and B exemplar locations in behavioral space. **b**, Action similarity comparisons (A vs. B; $n = 15/10$, ChR2-YFP/YFP; All vs. A; $n = 514/356$, ChR2-YFP/YFP) or Action B (All vs. B; $n = 443/356$, ChR2YFP/YFP). Plot indicates median/interquartile range. **c**, Reinforcement for Action A and B in ChR2-YFP animals. Plot indicates mean/S.E.M. **d**, Evolution of pooled action repertoire ($n = 427$ ChR2-YFP actions) reinforced for Action B. **e**, Early/Late cross-sectional views of (d). Blue dashed lines indicate fitted decay curve. Bottom inset graph shows normalized Early/Late fitted

curves. **f**, Contingency degradation of Action B. Target random laser triggers frequencies (bottom) is based on initial Action B performance prior to contingency degradation. Plots indicate mean/S.E.M. **g-j**, Action A (blue) induced by reinforcement for Action B in experienced ChR2-YFP animals. **g-h**, Moving mean frequencies over reinforcement for Action A or B. Dashed, vertical line mark first reinforcement. Plots are mean/S.E.M (colored fill). Bin1/Bin2 are time bins for (**i-j**). **i-j**, Frequency measures within time bins noted in (**g,h**). Repeated measures two-way ANOVA reveal significant difference across time and actions A/B frequencies (not shown). Šidák's post hoc comparisons. Asterisks except in (**h**): **** $p < 0.0001$. ** $p < 0.01$. * $p < 0.05$. n.s. – not significant. See Supplementary Information for statistical/sample details.

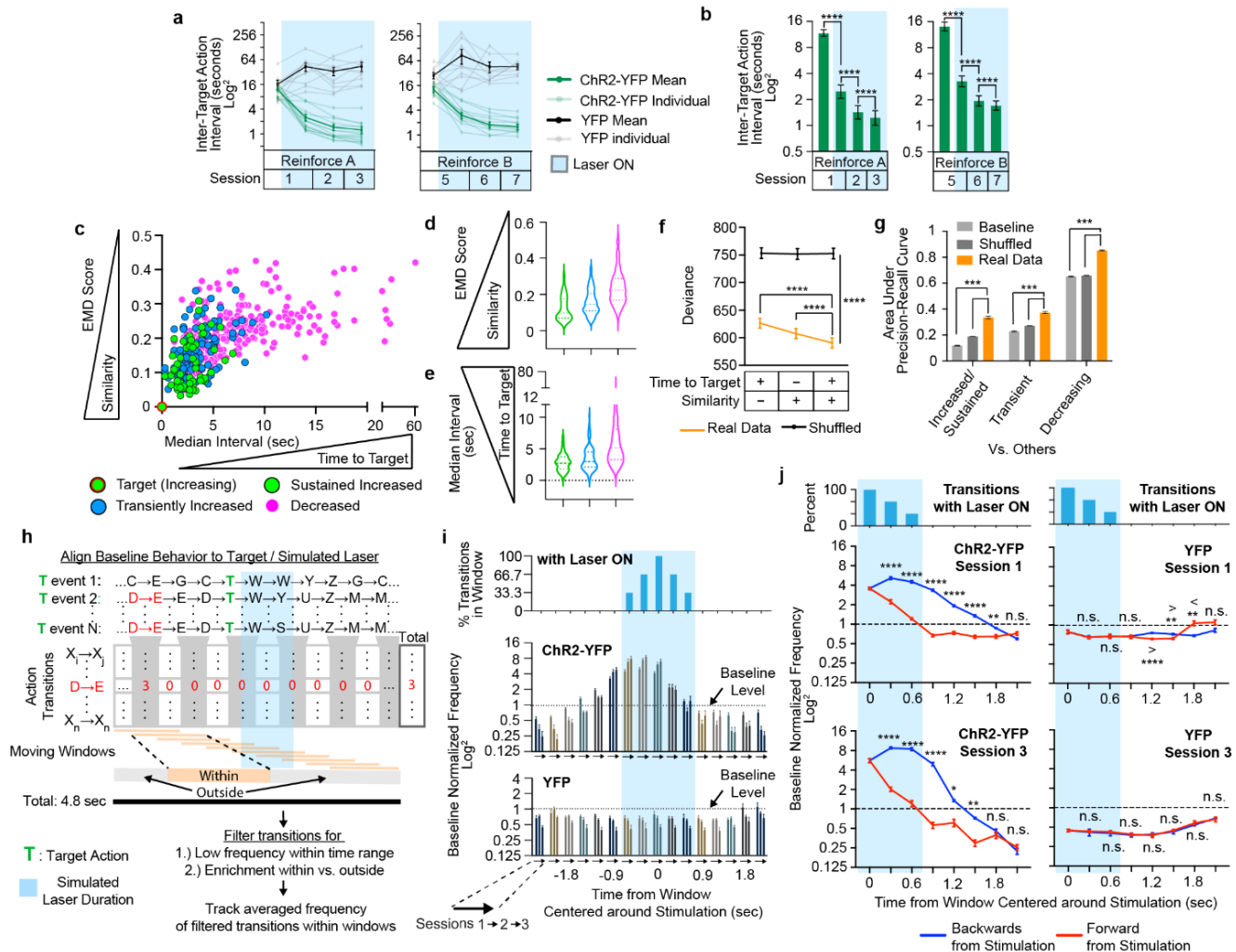


Figure 3. Dopamine mediates retrospective reinforcement of freely moving behavior. a-b,

ChR2-dependent reinforcement decrease inter-action intervals for Action A ($n = 15$ ChR2-YFP) and B ($n = 13$ of 15 ChR2-YFP). $n = 10$ YFP animals. Plots are mean/S.E.M (a-b). Significant

difference across time and ChR2-YFP/YFP (Mixed Effect Model. Action A: $F(3,69) = 72.26$, $p <$

0.0001. Action B: $F(3,62) = 33.78$, $p < 0.0001$.) b, Post-hoc Tukey's multiple comparisons of (a). c-

d, Distribution of action dynamic types ($n = 464$ actions, 15 ChR2-YFP animals) according to target

similarity (c,d), median time to target (c,e). d-e, Violin plots show median/quartiles. Two-tailed

permutation tests with Bonferroni-adjusted p-values. f-g, Multinomial logistic regression of all

factor combinations in Real data (200 models) versus Shuffled data (10,000 models). f. Groups

differ across combinations (repeated measures, two-way ANOVA. $F(2,30594) = 518.2$, $p <$

0.0001.). Post-hoc Dunnett multiple comparisons. Plots are mean/std. **g**, Performance of double-factor regression model measured with area under the precision-recall curves (AUPRC). Two-tailed permutation test with Bonferroni-adjusted p-value. Plots are mean/S.E.M. **h**, Identifying moving window-enriched action transitions. **i**, ChR2-dependent reinforcement for Action A increases action transitions prior to and within stimulation window. Plots indicate mean/S.E.M. **j**, Quantification of **(i)**. Significant difference across time and Retrospective/Forward reinforcement directions (Mixed Effect Modeling. ChR2-YFP Session1: $F(6,168) = 114.8$, $p < 0.0001$. ChR2-YFP Session 3: $F(6,168) = 46.62$, $p < 0.0001$, YFP Session1: $F(6,108) = 10.52$, $p < 0.0001$. YFP Session 3: $F(6,168) = 0.8992$, $p = 0.4984$). Post-hoc Šidák multiple comparisons. **** $p < 0.0001$, *** $p < 0.001$, ** $p < 0.01$, * $p < 0.05$, n.s. – not significant. See Supplementary Information for statistical/sample details.

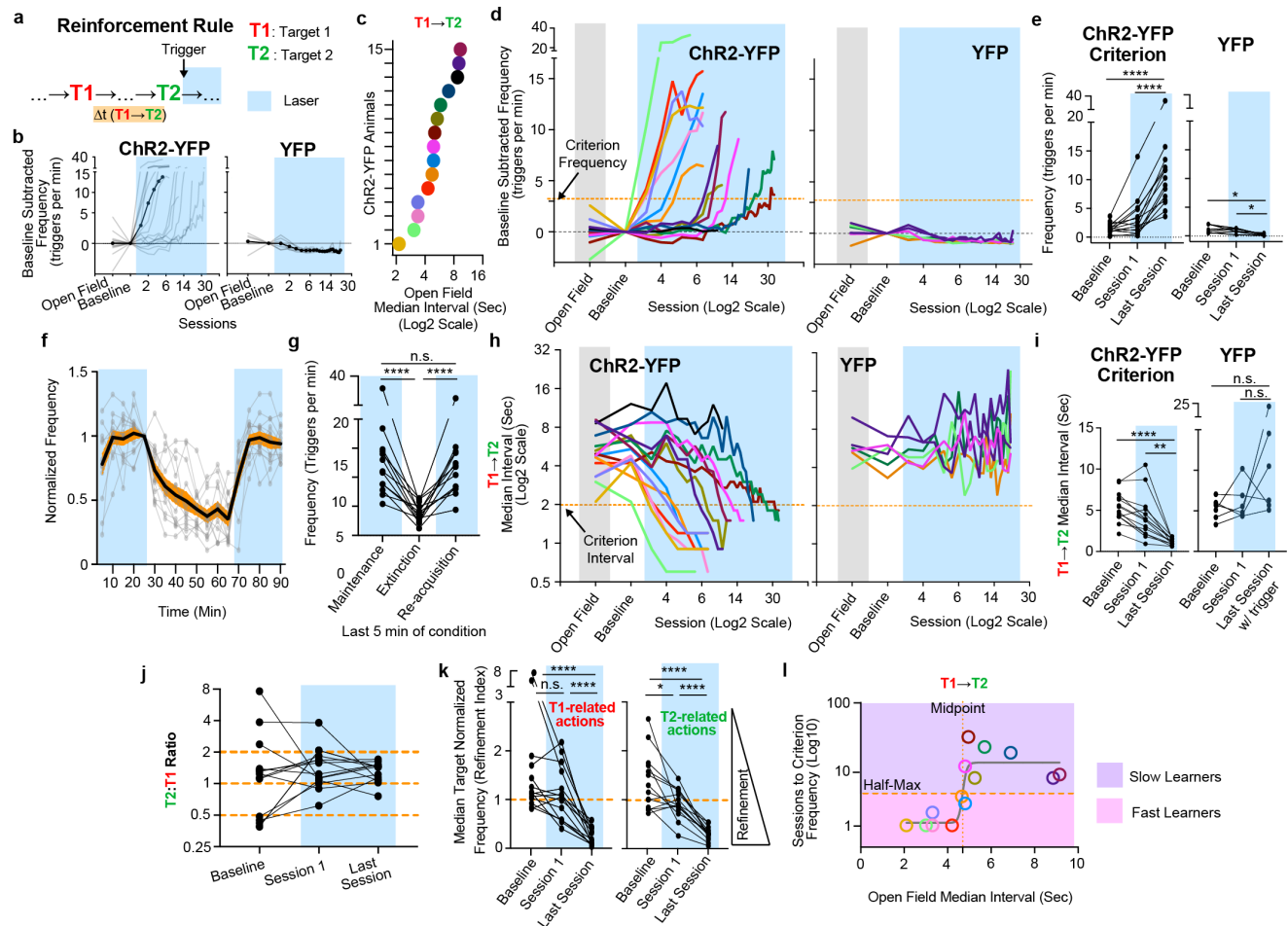


Figure 4. Relationship between pre-reinforcement inter-action intervals and learning of a two-action sequence. **a**, Schema. **b-l**, $n = 15$ (**b,d,h**) or 14 (**e,g,i-l**) ChR2-YFP, 6 YFP animals.

Repeated measures one-way ANOVA, post hoc Šidák tests applied in (**e,g,i,k**). Plots of individuals in (**d-e**). **b**, ChR2-dependent increase in $T1 \rightarrow T2$ triggers (no laser during open field / baseline). **c**, Open field inter-action intervals of $T1/T2$ pairs chosen. Same color codes in (**d,h**). **d**, Individual learning curves labeled by color codes in (**c**). **e**, Frequency changes over conditions ($F(1.911, 24.85) = 51.02$, $p < 0.0001$). **f-g**, Extinction of $T1 \rightarrow T2$ sequence (ChR2-YFP). **f**, Plot shows mean(black)/S.E.M.(orange fill)/individuals(grey). **g**, Frequency changes over extinction conditions ($F(1.073, 12.87) = 52.96$, $p < 0.0001$). **h-i**, ChR2-dependent decrease in $T1 \rightarrow T2$ intervals. ($F(1.377, 17.90) = 35.95$, $p < 0.0001$) (**i**). **j**, $T2:T1$ frequency ratios (ChR2-YFP) **k**, Target refinement shown by median target normalized frequencies of related actions. ($T1$: $F(1.237, 16.08) = 43.38$. $T2$:

$F(1.171, 15.22) = 48.74$. Both $p < 0.0001$). Individual color code as in (c,g). **I**, Sigmoidal relationship between open field $T1 \rightarrow T2$ interval and sessions to criterion frequency.

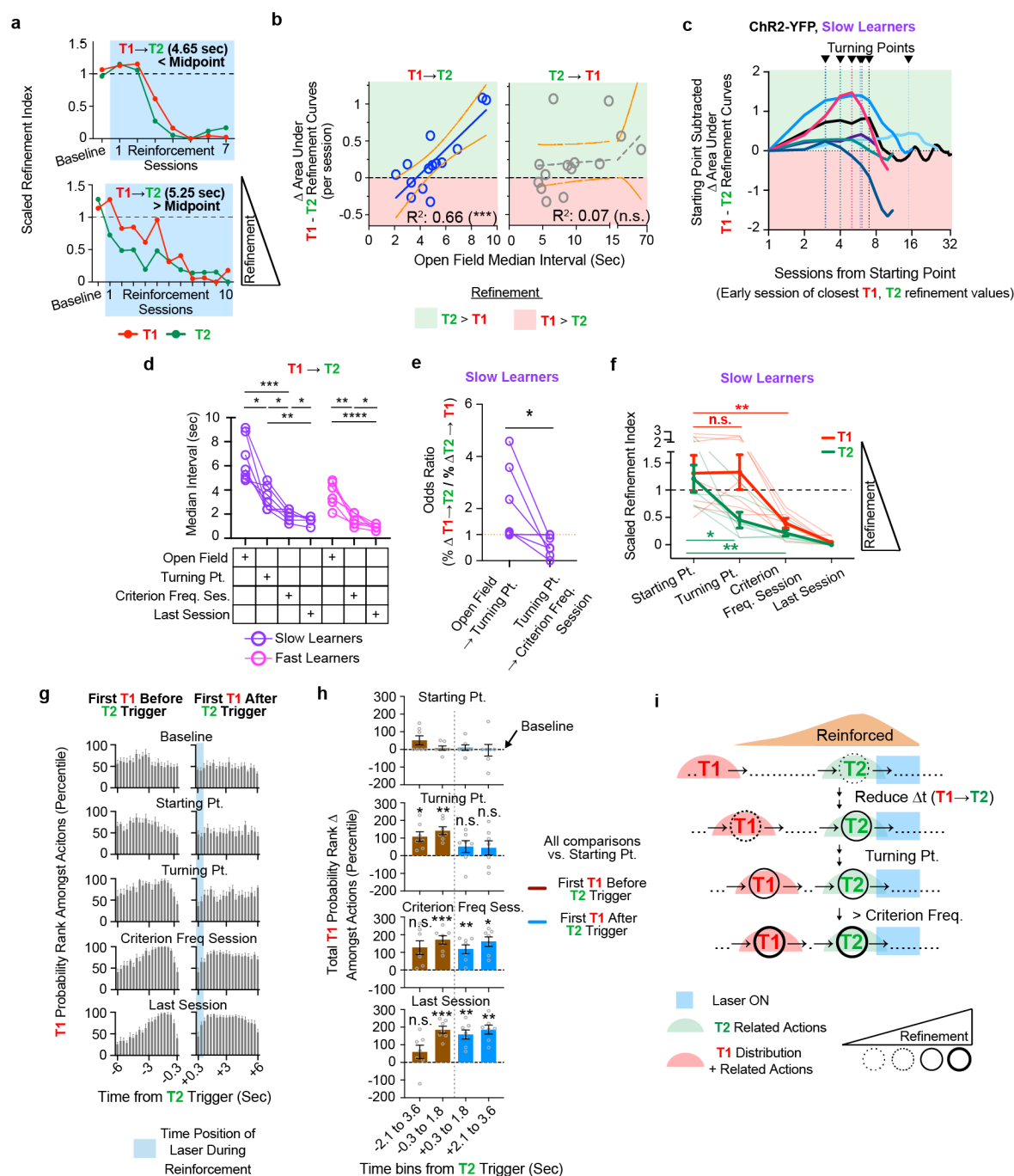


Figure 5. Behavioral process underlying learning of a two action sequence. $n = 14$ ChR2-YFP (7 Slow Learners). **a**, T1/T2 refinements in two ChR2-YFP individuals. **b**, Linear relationship between initial T1→T2 interval and differential T1-T2 refinement. Non-zero slope significance: T1→T2, $p = 0.0004$, T2→T1, $p = 0.7063$. **c**, Progression of differential T1-T2 refinement from Starting Point in Individual Slow Learners. **d**, T1→T2 interval significantly decreased by Turning Point in Slow

Learners. Repeated-measures 2-way ANOVA. Post hoc Tukey's test. **e**, Odds ratio of T1→T2 / T2→T1 interval changes. Paired Wilcoxon test ($p = 0.0312$, $n = 7$ animals). **f**, Preferential refinement of T2 relative to T1 by Turning Point in Slow Learners. Raw scaled refinement indices. Repeated measures, mixed effects model. Significant main effects. Time ($F(2.184, 26.20) = 54.21$, $p < 0.0001$). Post-hoc Šidák test. **g**, First occurrences of T1 before (left) and after (right) T2 triggers across learning stages. **h**, Quantification of pooled time bins from (**g**). Repeated measures, 2-way ANOVA for learning stage vs. rank change. First T1 Before and After T2 Trigger groups differ across learning stage and total T1 rank change. (Proximal bins (0.3-1.8 sec): $F(3,36) = 3.126$. $p=0.0376$. Distal bins (2.1 to 3.6 sec): $F(3,36) = 7.701$. $p<0.001$). Post-hoc Šidák relative to Starting Point values. **g**, Model for learning initially distantly separated T1→T2 sequences. Time not drawn to scale. **** $p < 0.0001$. *** $p < 0.001$. ** $p < 0.01$. * $p < 0.05$. n.s. – not significant. All bar plots indicate mean +/- S.E.M. See Supplementary Information for statistical/sample details.