Prefrontal cortex output circuits guide reward seeking through divergent cue encoding

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The prefrontal cortex is a critical neuroanatomical hub for controlling motivated behaviours across mammalian species1–3. In addition to intra-cortical connectivity, prefrontal projection neurons innervate subcortical structures that contribute to reward-seeking behaviours, such as the ventral striatum and midline thalamus4. While connectivity among these structures contributes to appetitive behaviours5–11, how projection-specific prefrontal neurons encode reward-relevant information to guide reward seeking is unknown. Here we use in vivo two-photon calcium imaging to monitor the activity of dorsomedial prefrontal neurons in mice during an appetitive Pavlovian conditioning task. At the population level, these neurons display diverse activity patterns during the presentation of reward-predictive cues. However, recordings from prefrontal neurons with resolved projection targets reveal that individual corticostriatal neurons show response tuning to reward-predictive cues, such that excitatory cue responses are amplified across learning. By contrast, corticothalamic neurons gradually develop new, primarily inhibitory responses to reward-predictive cues across learning. Furthermore, bidirectional optogenetic manipulation of these neurons reveals that stimulation of corticostriatal neurons promotes conditioned reward-seeking behaviour after learning, while activity in corticothalamic neurons suppresses both the acquisition and expression of conditioned reward seeking. These data show how prefrontal circuitry can dynamically control reward-seeking behaviour through the opposing activities of projection-specific cell populations.

Neurons in the prefrontal cortex (PFC) respond diversely to reward-predictive cues14–17, although how this cue encoding fits into a broader circuitry to guide reward seeking is unknown. To address this, we designed a Pavlovian conditioning task that allows two-photon imaging of deep cortical tissue during behaviour. Head-fixed mice were trained to associate one conditioned stimulus (CS\textsuperscript{+}) but not another (CS\textsuperscript{−}), with sucrose (Fig. 1a, b). Following multiple training sessions, mice behaviourally discriminated between the cues by displaying anticipatory licks to the CS\textsuperscript{+} but not CS\textsuperscript{−} (Fig. 1c), confirming that the cue-reward contingencies had been established by the later sessions (Fig. 1d, e; Extended Data Fig. 1). To monitor neural activity during this task, we injected a virus into dorsomedial PFC for delivery of a calcium indicator18 under the control of the calcium/calmodulin-dependent protein kinase II alpha (Camk2a) promoter (AAVdj-CaMK2a-GCaMP6s; Fig. 1f), which putatively targets cortical excitatory neurons19. Ex vivo brain slice recordings revealed that fluorescent deflections of GCaMP6s-expressing PFC neurons reliably tracked elevations and reductions in action potential frequency, whereas hyperpolarization from rest alone did not influence GCaMP6s-mediated fluorescence (Extended Data Fig. 2). Next, we implanted optical cannulae approximately 2.2 mm beneath the surface of the brain, allowing chronic optical access to hundreds of dorsomedial PFC neurons in each awake, behaving mouse (Fig. 1g–i; Supplementary Video 1). Collectively, we recorded from GCaMP6s-expressing PFC neurons before learning (n = 1,473) and after learning (n = 1,569), and found that while many of these neurons displayed increased activity in response to reward-predictive cues, other neurons exhibited inhibitory cue responses (Fig. 1j, k, n, o). These responses were most prevalent during presentation of the CS\textsuperscript{+}, but not the CS\textsuperscript{−}, after learning (Fig. 1l, m, p, q; Extended Data Figs 3a, b and 4a–c). Thus, the responses of many individual PFC neurons could be used to decode whether the CS\textsuperscript{+} or CS\textsuperscript{−} was presented on any given trial after learning (Extended Data Fig. 4d).

We found that many dorsomedial PFC neurons encoded cues after learning; however, both inhibitory and excitatory responses were common. Considering this, it is possible that distinct populations of neurons in the PFC have opposing cue-encoding properties. Two major projection targets of the PFC are the nucleus accumbens (PFC–NAc) and paraventricular nucleus of the thalamus (PFC–PVT)4,20. Furthermore, studies examining reward seeking in rodent addiction models have found that PFC–NAc neurons express immediate early genes following cue-induced relapse21,22, and inhibition of these cells can prevent drug seeking6,8,12. Additionally, reward-predictive cues may influence the activity of PVT neurons23, and such activity is thought to modulate cue salience24 and conditioned behaviours25,26. Thus, we hypothesized that PFC–NAc and PFC–PVT neurons encode reward-predictive stimuli to orchestrate appetitive learning. To test this hypothesis, we first determined whether PFC–NAc and PFC–PVT neurons are distinct or overlapping populations of neurons. We injected retrograde cholera toxins conjugated to different fluorophores into NAc and PVT of the same mice. Electrophysiological recordings and histological analysis revealed that PFC–NAc and PFC–PVT neurons are physiologically distinct and anatomically segregated within dorsomedial PFC (Extended Data Fig. 5), such that PFC–NAc neurons are in layers II/III and V (46% layer II/III, 51% layer V, 4% layer VI) whereas PFC–PVT neurons are predominantly in layer VI (0% layer II/III, 20% layer V, 79% layer VI). These data are consistent with anatomical studies, as corticostriatal projection neurons reside in layers II/III and V in rats, whereas layer VI is specific for corticothalamic neurons27,28. Furthermore, using a retrograde rabies tracing strategy to sparsely label PFC–NAc or PFC–PVT neurons as well as their afferent inputs, we found that these cell populations have both shared and unshared afferent connectivity (Extended Data Fig. 6). Finally, PFC–NAc and PFC–PVT neurons express CaMK2a (Extended Data Fig. 7a–d), collectively revealing that these neurons make up two non-overlapping subpopulations of CaMK2a-expressing projection neurons.

We next targeted PFC–NAc and PFC–PVT neurons for in vivo two-photon calcium imaging. Cre-inducible GCaMP6s (AAVdj-DIO-GCaMP6s) was injected into dorsomedial PFC, and in the same surgery a retrogradely transported virus, canine adenovirus 2 encoding
Cre-recombinase (Cav2-Cre), was injected into either the NAc or PVT (Fig. 2a, g). This resulted in projection-specific GCaMP6s expression in PFC–NAc and PFC–PVT neurons (Fig. 2b, h; Extended Data Fig. 7e–j). Next, mice underwent Pavlovian conditioning with simultaneous optogenetic manipulation by injecting Cav2-Cre into the NAc or PVT, and Cre-inducible channelrhodopsin-2 (AAV5-DIO-Chr2-eYFP), halorhodopsin (AAV5-DIO-eNpHR3.0-eYFP), or control (AAV5-DIO-eYFP) into dorsomedial PFC (Fig. 4a–c, j–l). Next, we optogenetically manipulated these cells during cue delivery throughout Pavlovian conditioning (sessions 1–8; Fig. 4d–f, m–o), and tested the effects on acquisition of this task during a subsequent test without laser. Data reveal that activation (in PFC–NAc::Chr2 mice) or inactivation (in PFC–NAc::eNpHR3.0 mice) of PFC–NAc neurons did not influence CS+–evoked anticipatory licking during the no-laser test (Fig. 4g–i), implying that these cells do not control the acquisition of anticipatory licking. By contrast, optogenetic activation of PFC–PVT neurons (in PFC–PVT::Chr2 mice) reduced CS+–evoked anticipatory licking during the no-laser test, whereas optogenetic inactivation of PFC–PVT neurons (in PFC–PVT::eNpHR3.0 mice) increased anticipatory licking responses after learning (Fig. 3e, f). Thus, the responses of PFC–NAc, but not PFC–PVT, neurons after learning could be predicted on the basis of responses before learning (Fig. 3c, f, correlations). Collectively, corticostriatal and corticothalamic neurons showed distinct functional plasticity across appetitive learning, such that PFC–NAc neurons adjusted their activity to allow cue discrimination, whereas PFC–PVT neurons gradually acquired responses for cue discrimination.

Our data reveal that projection-specific PFC neurons show distinct cue encoding properties across appetitive learning. However, whether this activity controls the acquisition of conditioned reward seeking is unclear. To test this, we targeted PFC–NAc or PFC–PVT neurons for optogenetic manipulation by injecting Cav2-Cre into the NAc or PVT, and Cre-inducible channelrhodopsin-2 (AAV5-DIO-Chr2-eYFP), halorhodopsin (AAV5-DIO-eNpHR3.0-eYFP), or control (AAV5-DIO-eYFP) into dorsomedial PFC (Fig. 4a–c, j–l). Next, we optogenetically manipulated these cells during cue delivery throughout Pavlovian conditioning (sessions 1–8; Fig. 4d–f, m–o), and tested the effects on acquisition of this task during a subsequent test without laser. Data reveal that activation (in PFC–NAc::Chr2 mice) or inactivation (in PFC–NAc::eNpHR3.0 mice) of PFC–NAc neurons did not influence CS+–evoked anticipatory licking during the no-laser test (Fig. 4g–i), implying that these cells do not control the acquisition of anticipatory licking. By contrast, optogenetic activation of PFC–PVT neurons (in PFC–PVT::Chr2 mice) reduced CS+–evoked anticipatory licking during the no-laser test, whereas optogenetic inactivation of PFC–PVT neurons (in PFC–PVT::eNpHR3.0 mice) increased anticipatory licking.
Figure 2 | PFC projection neurons have opposing responses to reward-predictive cues. a, Viral strategy (a) allowed recordings of PFC–NAC::GCaMP6s neurons (early, n = 84 neurons; late, n = 101 neurons; n = 4 mice) in vivo (b). c, Population heat plots showing responses for all PFC–NAC::GCaMP6s neurons averaged across CS− trials (c) and CS+ trials (d) after learning. e, Population data of all PFC–NAC::GCaMP6s neurons showing no difference in CS− versus CS+ responses during early sessions (top; χ2(1) = 0.38, P > 0.6); however, these responses were different during late sessions (bottom; χ2(1) = 41.06, P < 0.001). f, Cumulative distribution frequency (CDF) plots showing that the dynamics of individual PFC–NAC::GCaMP6s neurons could be used to accurately decode whether the CS+ or CS− was presented in late conditioning sessions (compared to late shuffled; Welch’s t159.66 = 5.63, P < 0.001), but not in the early conditioning sessions (compared to early shuffled; Welch’s t125.47 = 1.13, P > 0.2). g, h, Viral strategy (g) allowed recordings of PFC–PVT::GCaMP6s neurons (early, n = 4 mice) after learning. i, j, Population heat plots showing responses for all PFC–PVT::GCaMP6s neurons averaged across CS− trials (i) and CS+ trials (j) after learning. k, Population data of all PFC–PVT::GCaMP6s neurons showing no difference in CS− versus CS+ responses during early sessions (top; χ2(1) = 2.02, P > 0.35); however, these responses were different during late sessions (bottom; χ2(1) = 43.86, P < 0.001). l, CDF plots showing that the dynamics of individual PFC–PVT::GCaMP6s neurons could be used to accurately decode whether the CS+ or CS− was presented during late conditioning sessions (compared to late shuffled; Welch’s t165.47 = 6.03, P < 0.001) but not during the early conditioning sessions (compared to early shuffled: Welch’s t165.47 = –0.56, P > 0.5). Vertical dotted lines refer to timing of sucrose delivery. Scale bars, 100 μm; EarlySh, early shuffled; LateSh, late shuffled.

Figure 3 | PFC projection neurons show distinct functional plasticity across learning. a, Representative images show the same PFC–NAC::GCaMP6s neurons tracked from early (left) to late (right) sessions (n = 37 neurons; n = 4 mice). b, Traces from individual example neurons averaged across trials during early (left) and late (right) sessions. c, Cue responses of PFC–NAC::GCaMP6s neurons during early conditioning sessions could be used to predict responses during late conditioning sessions (CS+, r = 0.73, P < 0.001; CS−, r = 0.70, P < 0.001). auROC is the area under receiver operating characteristic, and auROC − 0.5 is a statistic that compares the probability that GCaMP6s fluorescence dynamics are the same during the CS+ versus CS− (auROC = 0), higher for the CS+ versus CS− (auROC > 0), or less for the CS+ versus CS− (auROC < 0). Inset, bar graphs showing that most PFC–NAC::GCaMP6s neurons showed elevated (grey) GCaMP6s fluorescence in response to the CS+ across learning, and reduced (white) GCaMP6s fluorescence in response to the CS− across learning (χ2(1) = 8.07, **P < 0.005). d, Representative images show the same PFC–PVT::GCaMP6s neurons tracked from early (left) to late (right) sessions (n = 61 neurons; n = 3 mice). e, Traces from individual example neurons averaged across trials during early (left) and late (right) sessions. f, Cue responses of all PFC–PVT::GCaMP6s neurons during early Pavlovian conditioning sessions could not be used to predict subsequent responses during late conditioning sessions (CS+, r = 0.08, P > 0.05; CS−, r = 0.24, P > 0.05). Inset, bar graphs showing that most PFC–PVT::GCaMP6s neurons showed reduced (white) GCaMP6s fluorescence in response to the CS+ across learning, whereas equivalent numbers of neurons showed elevated (grey) and reduced (white) GCaMP6s fluorescence in response to the CS− across learning (χ2(1) = 6.73, **P < 0.01). Vertical dotted lines refer to timing of sucrose delivery. Scale bars, 25 μm.

During the no-laser test (Fig. 4p-r). Thus, inhibitory cue encoding in PFC–PVT neurons, but not PFC–NAC neurons, contributes to the acquisition of conditioned reward seeking. Although activity in PFC–NAC neurons did not control the acquisition of conditioned reward seeking, differences in licking behaviour were apparent between groups during the laser conditioning sessions (sessions 1–8; see Fig. 4g, h). Thus, one possibility is that PFC–NAC neurons control the expression of conditioned reward seeking, rather than acquisition. To test this directly, we next performed the optogenetic manipulations after conditioning in separate groups of mice.
We found that opticogenetic activation of PFC–NAc neurons during the CS\(^+\) increased anticipatory licking (Fig. 5a, c, f), whereas inactivation of PFC–NAc neurons reduced anticipatory licking (Fig. 5b, e, i). By contrast, opticogenetic activation of PFC–PVT neurons during the CS\(^+\) reduced anticipatory licking (Fig. 5g, j, k) whereas inactivation of PFC–PVT neurons had no effect (Fig. 5h, l, k). The lack of effect for PFC–PVT inactivation is probably due to CS\(^-\)-evoked inhibition of these cells (see Figs 2 and 3), as neuronal hyperpolarization from sub-threshold potentials should not result in further adjustment of neuronal output. Collectively, activation of PFC–NAc neurons and inhibition of PFC–PVT neurons supports the expression of conditioned reward seeking.

In addition to encoding the reward-predictive stimulus after learning, some PFC–NAc and PFC–PVT neurons displayed CS\(^-\) encoding (see Fig. 2). Considering this, we also evaluated how projection-specific cells influenced licking during the CS\(^-\). We found that opticogenetic perturbations of PFC–NAc and PFC–PVT neurons during the CS\(^-\) did not influence the acquisition of CS\(^-\) licking (Extended Data Fig. 8a–f). By contrast, inactivation but not activation of PFC–NAc neurons reduced the expression of CS\(^-\) licking, whereas opticogenetic manipulations of PFC–PVT neurons had no effect on the expression of CS\(^-\) licking (Extended Data Fig. 8g–i). Moreover, opticogenetic manipulation of PFC–NAc or PFC–PVT neurons during random time epochs outside of cue delivery had no effect on licking (Extended Data Fig. 9), nor did opticogenetic manipulations produce a real-time place preference or aversion (Extended Data Fig. 10). These results suggest that activity in PFC–NAc and PFC–PVT neurons specifically during the cue period is critical for conditioned reward seeking.

Here we found that the opposing population dynamics of prefrontal output circuits orchestrate conditioned reward seeking. In addition to divergent population encoding, subsets of cells in both corticostriatal and corticothalamic circuits have response profiles that are opposite to the overall population (that is, some PFC–NAc neurons have inhibitory CS\(^+\) responses, whereas some PFC–PVT cells have excitatory CS\(^+\) responses). Although the cause and function of such heterogeneity is unclear, it is possible that subpopulations of cells within each group may have distinct cell-type-specific targets, afferent inputs, and spatial locations. Thus, while it remains unclear how all cells in PFC function to orchestrate reward seeking, our data suggest that projection-specific and anatomically segregated prefrontal neurons can have opposing activity dynamics, plasticity profiles, and functional control of conditioned reward seeking.

**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.
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METHODS

Subjects and surgery. Adult male C57BL/6J mice (Jackson Laboratories) or Ai9 reporter mice (Cg-Gt(Rosa)26Sorjtm1(CAG-diTomato)HrJ; Jackson Laboratories) were group-housed (25–35 g, 6–8 weeks old) with littermates until surgery. For all experiments, mice underwent surgery during which they were anesthetized with 0.8–1.5% isoflurane vaporized in pure oxygen (1 min−1) and placed within a stereotactic frame (David Kopf Instruments). Ophthalmic ointment (Akorn) and a topical anesthetic (2% lidocaine; Akorn) were applied during surgeries, and subcutaneous injections of sterile saline (0.9% NaCl in water) were administered to prevent dehydration. During surgeries, virus injections were administered unilaterally (for two-photon microscopy experiments) or bilaterally (for optogenetic or anatomical experiments) targeting dorsal medial PFC (specifically prelimbic cortex; 500 nl per side; relative to bregma: AP, +1.85 mm; ML, ±0.60 mm; DV, −2.50 mm), bilaterally targeting NAc (500 nl per side; relative to bregma: AP, +1.42 mm; ML, ±0.73 mm; DV, −4.80 mm), and/or on the midline targeting PVT (300 nl; relative to bregma: AP, −1.46 mm; ML, −1.13 mm; DV, −3.30 mm; 20° angle). The UNC Vector Core packaged all viruses except canine adenovirus 2 encoding Cre (Cav2-Cre; Institut de Génétique Moléculaire de Montpellier). For two-photon imaging experiments, an optical cannula (Inscoptix) was implanted above the PFC injection site (relative to bregma: AP, +1.85 mm; ML, −0.8 mm; DV, −2.2 mm; see ref. 29 for details of using similar surgical protocols for imaging experiments). For optogenetic experiments, custom-made optical fibres5 were implanted bilaterally approximately 0.5 mm above the PFC injection sites (relative to bregma: AP, +1.85 mm; ML, ±0.83 mm; DV, −1.93 mm; 10° angle). For experiments involving head-fixed behaviour, a custom-made ring (stainless steel; 5 mm ID, 11 mm OD) was attached to the skull during surgery to allow head fixation (see Fig. 1a). Following surgeries, mice received acetaminophen in their drinking water for two days, and were allowed to recover with access to food and water ad libitum for at least 21 days. After recovery, mice were water restricted (water bottles taken out of the cage), and 0.6 ml of water was delivered every day to a dish placed within each home cage. Behavioural experiments began when mice weighed less than 90% of free drinking weight (around 10 days for all experiments). To ensure good health and weight maintenance, mice were weighed and handled daily. This protocol resulted in weight stabilization between 85–90% of free-drinking weight during each experiment. No mouse was given more or less than 0.6 ml of water for weight concerns during water restriction procedures, nor did any health problems related to dehydration arise at any point from these protocols. All experimental procedures were performed in accordance with the Guide for the Care and Use of Laboratory Animals (National Institute of Health), and were approved by the Institutional Animal Care and Use Committee at the University of North Carolina prior to experiments commencing.

Head-fixed behaviour. Following recovery from surgery, mice were habituated to head fixation for 3 days, during which unpredictable drops of sucrose (10% sucrose in water; 2.0–2.5 μl) were delivered intermittently for one hour (approximately 60 drops per hour) through a gravity-driven, solenoid-controlled lick tube. Once the mice displayed sufficient licking (>1,000 licks per session), they underwent Pavlovian conditioning. During each conditioning session, two cues (3 kHz pulsing or 12 kHz constant tones, 2s, 70 dB) were randomly presented 50 times before the delivery of sucrose (CS+; 10% sucrose in water; 2.0–2.5 μl) or no sucrose (CS−), such that there was a one second trace interval between delivery of the CS+ and sucrose (see Fig. 1b). The cue contingencies were counterbalanced across cohorts of mice to ensure that mice acquired conditioned licking in response to either tone when paired with sucrose. The inter-trial interval between the previous reward delivery (CS+) or withholding time (CS−) and the next cue was chosen as a random sample from a uniform distribution bounded by 40s and 80s. Cue discrimination was quantified using the area under a receiver operating characteristic (auROC) formed by the number of baseline-subtracted licks during the CS+ versus CS− sessions. For both active and passive experiments, we classified sessions as ‘early’ or ‘late’ in learning, defined by both behavioural performance (early, auROC < 0.65; late, auROC > 0.66) and session number (early, sessions 1–5; late, sessions 7 or later). These criteria were used as post hoc analysis revealed that an auROC > 0.66 approximates high performance in a phase space formed by behavioural performance across sessions. Finally, behavioural data are displayed and analysed throughout the manuscript as the change in lick rate between each 3-s cue period and 1-s baseline period (baseline period is immediately before each cue). In addition, we show raw lick rates during both the cue and baseline periods for all imaging experiments (see Extended Data Fig. 1). Baseline lick rates remained relatively low across all experiments, and therefore the optogenetic studies only change the lick rate in shown and analysed (see Figs 4, 5 and Extended Data Figs 8 and 9).

Two-photon microscopy. Experimental design. Two-photon microscopy was used to visualize activity dynamics of PFC neurons in vivo. A virus encoding the calcium indicator GCaMP6s18 (AAVd2-CaMK2a-GCaMP6s; 5.3 × 1012 infectious units per ml) was injected into PFC (see Subjects and surgery). For imaging projection-specific neurons, a virus encoding the Cre-dependent calcium indicator GCaMP6s (AAVd2-Cfi-CIO-DIO-GCaMP6s; 3.1 × 1013 infectious units per ml; from Karl Deisseroth) was injected into PFC, and the retrogradely transported canine adenovirus encoding Cre-recombinase31,32 was injected into either NA or PVT (Cav2-Cre; 4.2 × 1013 infectious units per ml). After a minimum of 8 weeks to allow virus transport and infection, mice underwent Pavlovian conditioning, during which GCaMP6s-expressing neurons were visualized using two-photon microscopy.

Data acquisition, signal extraction, and analysis. A two-photon microscope (FVMP-E R5) was equipped with the following to allow imaging of PFC in vivo: a hybrid scanning core set with galvanometers and fast resonant scanners (up to 30 Hz frame-rate acquisition; set to 2.5 Hz), multi-alkali PMT and GaAsP-PMT photo detectors with adjustable voltage, gain, and offset features, a single green/red NDD filter cube, a long working distance 20× air objective designed for optical transmission at infrared wavelengths (Olympus, LCP LN20XIR, 0.45 NA, 8.3 mm WD), a software-controlled modular syringe stage loaded on a manual z-deck, and a tunable Mai–Tai Deep See laser system (Spectra Physics, laser set to 955 nm, ∼100 fs pulse width) with automated four-axis alignment. Before each condition session, a particular field of view (FOV) was selected by adjusting the imaging plane (z-axis), and each FOV was spaced at least 50 μm from one another to prevent visualization of the same cells across multiple FOVs. During each condition session, two-photon scanning was triggered for each trial 1 s before cue delivery, and a 20 s video was then collected for each trial. Data were both acquired and processed using a computer equipped with FluoView (Olympus, FV1200) and cellSens (Olympus) software packages. Following data acquisition, videos were motion corrected using a planar hidden Markov model (SIMA v1.3)33 and regions of interest (ROIs) were hand drawn around each cell using the standard deviation projection of the motion-corrected video using ImageJ. Next, calcium transient time series data were extracted with SIMA and analysed using custom Python data analysis pipelines written in the laboratory (by V.M.K.N.). For analysis, data were split into two groups (early and late) that were defined based on behavioural performance and the day of conditioning (see Head-fixed behaviour). Next, each recorded neuron was defined as having an excitatory response, inhibitory response, or no response. Significant responses represent significant two-tailed auROC comparing average fluorescence (Δf/f) of the trace interval (1 s after CS onset) versus baseline (1 s before CS onset) where P < 0.05 after Benjamini–Hochberg false discovery rate correction. Each P value for auROC was defined by calculating the P values for the corresponding Mann–Whitney U statistic. χ2 tests were then used to compare the number of CS+ responders to CS− responders for each group. For additional decoding analysis (for example, Fig. 2f, l), we tested whether the identity of the cue on any given trial could be decoded from the mean trace interval response on that trial using support vector machines. To this end, we used the Python module, scikitlearn, with GridSearchCV and a support vector classifier (SVC) estimator with a radial basis function kernel, optimizing across the following parameters: γ ∈ {10−2, 10−1, 100, 104}, C ∈ {10−2, 10−1, 101, 104}. Quantification of performance was done using tenfold validation and the R2 metric (note that this metric can be infinitely negative, indicating arbitrarily poor performance, but is bounded on the positive end at 1, indicating perfect decoding). We found that as a population, the number of anticipatory licks during the trace interval could not be decoded in the late sessions in CaMK2a-expressing neurons (mean R2 = −1.21), PFC–NAc neurons (mean R2 = −0.92) or PFC–PVT neurons (mean R2 = −0.39). These negative numbers
reflect the absence of a relationship between licking and calcium activity in each cell population.

**Optogenetics.** Behavioural optogenetics were performed as described in detail previously35. In brief, during surgery a virus encoding Cre-inducible channelrhodopsin-2 (AAV5-ef1α-DIO-hChR2(H134R)-eYFP; 5.0 × 10^{12} infectious units per ml), halorhodopsin (AAV5-ef1α-eNPHR3.0-eYFP; 8.0 × 10^{13} infectious units per ml), or control (AAV5-ef1α-eYFP; 6.0 × 10^{12} infectious units per ml) was injected into PFC; and Cav2-Cre (ref s 31, 32) (4.2 × 10^{13} infectious units per ml) was injected into either NAc or PVT. After a minimum of 8 weeks to allow sufficient virus transport and infection, mice underwent Pavlovian conditioning.

For acquisition experiments (for example, Fig. 4), mice underwent eight daily conditioning sessions with laser followed by a test session (no laser). For photo-activation manipulations in ChR2 or control mice, the laser (473 nm; 8–10 mW) was turned on for 5-ms pulses (20 Hz) during 80% of the cue trials, starting at the cue onset and ending at the reward delivery. For photoinhibition manipulations in eNPHR3.0 or control mice, the laser (532 nm; 8–10 mW) did not pulse. Because the laser had no effect in the control mice, these data were collapsed across PFC–NAc and PFC–PVT groups. For expression experiments (for example, Fig. 5), after mice reached high performance criterion (late, auROC > 0.66), they underwent six daily conditioning sessions. Furthermore, every other session was selected for optogenetic manipulations, during which the laser was presented for 3 s during either the cue and trace interval or at random time epochs outside of cue or reward delivery. Because there was no effect of laser in the ChR2 or eNPHR3.0 control mice, these data were collapsed for PFC–NAc groups and PFC–PVT groups. In addition, for expression experiments subsets of control mice were used twice, once as ChR2 controls (blue light), and again as eNPHR3.0 controls (green light). Following experiments, histological verification of fluorescence and optical fibre placements were performed as described previously35.

Behavioural data (change in lick rate, see above) was analysed based on a priori comparisons of interest (effect of laser on ChR2/eNPHR3.0 animals versus effect of laser in eYFP animals). For acquisition experiments (Fig. 4; Extended Data Fig. 8a–f), we analysed data from the no-laser test day only, and specifically compared the change in lick rate between the ChR2 or eNPHR3.0 groups versus the eYFP group. To correct for the double comparison (ChR2 or eNPHR3.0 versus eYFP), we performed a Benjamini–Hochberg multiple comparisons correction. For expression experiments (Fig. 5; Extended Data Figs 8g–1 and 9), in each pair of sessions (no laser, laser) we calculated the difference in mean lick rate between the two in order to obtain a statistical measure of the ‘effect of laser’ per session pair. Next, we compared the effects of laser from the ChR2 or eNPHR3.0 groups versus the corresponding effect of laser in the eYFP group. To correct for the double comparison (ChR2 or eNPHR3.0 versus eYFP), we again performed a Benjamini–Hochberg multiple comparisons correction. Considering this, for optogenetic experiments all P values (which are two-tailed throughout the manuscript) have been corrected for multiple comparisons.

**Retrograde tracing.** The anatomy and electrophysiological properties of PFC–NAc and PFC–PVT neurons were evaluated through retrograde tracing34. Specifically, during surgeries the retrograde tracer cholera toxin subunit B conjugated to Alexa Fluor (ChB–488, ChB–594; Molecular Probes) was injected bilaterally into NAc (500 nl per side) and on the midline in PVT (300 nl; colour counterbalanced across mice). Ten days following surgery, animals were killed for histology (n = 3 mice) or slice electrophysiology (n = 3 mice). For anatomical experiments, a student blind to all experiments (E.P.M.) and conditions counted the number of ChB–488 positive, ChB–594 positive, and double-positive neurons in prelimbic medial prefrontal cortex (a subregion of dorsal medial PFC). The distance of each cell from the midline and the layer specificity of each cell were then measured using ImageJ. For electrophysiological experiments, mice were euthanized ten days following surgery for patch-clamp recordings ex vivo (see below for details).

**Rabies tracing.** The monosynaptic afferents to PFC–NAc and PFC–PVT neurons were identified using a glycoprotein-deleted rabies strategy17 in combination with Cav2-Cre targeting of projection-specific neuron populations. Specifically, during the first surgery a cocktail containing the Cre-dependent starter viruses encoding the G-protein and Tva were injected into PFC (3:1 of AAV5-FLEX-RG and AAV5-FLEX-TVA-mCherry; 300 nl per side), and Cav2-Cre was injected into either NAc (500 nl per side) or PVT (300 nl). Five weeks later, mice were given a second surgery in which the G-deleted rabies virus was injected into PFC (1.5 diluted rabies–GFP). Finally, 8 days after the rabies injection each mouse (n = 3 per group) was euthanized for histology and cell quantification. Our rabies protocol led to sparse labelling of PFC-projecting neurons, allowing quantification of individual cells in each brain section (40 μm thick). Each ROI was selected based on previous PFC tracing experiments36, as well as the fluorescence intensity observed in our experiments. Next, out of all tissue collected for each ROI in each mouse, we selected the three sections containing the most cells per region, and used confocal microscopy to get cellular-resolution images of all cells in each of those sections. For each section, we quantified all individual input neurons (GFP+ and starter cells (both GFP+ and mCherry+)). Considering that the anterior cingulate cortex (ACC) was close to the PFC injection site, some sections containing ACC also had starter cell labelling. Thus, because we were interested in long-range inputs from ACC only, sections that did not have mCherry labelling were used for ACC input quantification. Finally, rabies-tracing data were analysed by comparing the number of cells in each section across groups (raw neuron count), and by comparing the percentage of input neurons per starter cell for each particular mouse.

**Patch-clamp electrophysiology.** Mice were anaesthetized with pentobarbital (50 mg kg−1) before transcardial perfusion with ice-cold sucrose cutting solution containing the following (in mM): 225 sucrose, 119 NaCl, 1.0 NaH2PO4, 4.9 MgCl2, 0.1 CaCl2, 26.2 NaHCO3, 1.25 glucose, 305 mOsM. Brains were then rapidly removed, and coronal sections 300 μm thick were made using a vibratome (Leica, VT 1200). Sections were then incubated in aCSF (32 °C) containing the following (in mM): 119 NaCl, 2.5 KCl, 1.0 NaH2PO4, 1.3 MgCl2, 2.5 CaCl2, 26.2 NaHCO3, 15 glucose, approximately 306 mOsM. After an hour of recovery, slices were constantly perfused with aCSF (32 °C) and visualized using differential interference contrast through a 40× water-immersion objective mounted on an upright microscope (Olympus BX51WI). Whole-cell recordings were obtained using borosilicate pipettes (3–5 MΩ) back-filled with internal solution containing the following (in mM): 130 K gluconate, 10 KCl, 10 HEPES, 10 EGTA, 2 MgCl2, 2 ATP, 2 GTP (pH 7.35, 270–285 mOsM).

Current-clamp recordings were obtained from GCaMP6s-expressing neurons to identify how action potential frequency correlated with GCaMP6s fluorescence. Specifically, to determine how elevations in action potential frequency influence GCaMP6s fluorescence, a 1 s train of depolarizing pulses (2 nA, 2 ms) was applied at a frequency of 1, 2, 5, 10 or 20 Hz. To determine how attenuations in action potential frequency influence GCaMP6s fluorescence, a 3 s pause was applied after a 10 s baseline train of depolarizing pulses (2 nA, 2 ms; 1, 2, 5, 10 or 20 Hz). Finally, to determine if hyperpolarization influences GCaMP6s fluorescence in the absence of action potential frequency modulation, a 3 s hyperpolarizing step (150 pA) was applied in neurons that were held either above or below resting membrane potential. During electrophysiological recordings, GCaMP6s fluorescence dynamics were visualized using a mercury lamp (Olympus, U-RFL-T) and microscope-mounted camera (QImaging, optiMOS). Imaging data were acquired using Micro-Manager, and extracted through hand-drawn ROIs for each recorded neuron using ImageJ.

Current-clamp recordings were also obtained to identify the intrinsic properties of PFC–NAc and PFC–PVT neurons in retrograde tracing experiments, as previously described35. First, action potential firing was examined by applying a series of long depolarizing sweeps (800 ms) at +25 pA steps (0–450 pA). Next, rheobase (the minimum amount of current required for an action potential to fire) was measured by applying a series of short depolarizing sweeps (50 ms) at +10 pA steps (starting at 0 pA) until the recorded neuron fired an action potential. For all patch-clamp experiments, data acquisition occurred at 1 kHz sampling rate through a MultiClamp 700B amplifier connected to a Digidata 1440A digitizer (Molecular Devices). Data were analysed using Clampfit 10.3 (Molecular Devices).

**Data collection.** The nature of all imaging and behavioural experiments yields high-power datasets, as we can test responses to reward-predictive cues hundreds of times within a single session. Thus, although the experiments themselves require rigorous experimentation, the number of mice that are required for each experiment is generally 3–6 per group, depending on the effect size (which was not predetermined for these experiments). Mice were randomly picked for each group in each experiment, by alternating the surgery for each mouse in a cage. During data collection, investigators were only blind to the conditions for rabies tracing cell counting and ChR2 cell counting. The only mice excluded from final analysis were those that died before or during the experiments (n = 3). For optogenetic experiments, mice were excluded if histology confirmed ectopic virus expression outside of PFC (n = 1), or if optical-fibre placements were not in dorso medial PFC (n = 0). For data analysis, equal variance was not assumed for behavioural optogenetics or imaging datasets. Equal variance was assumed for cell counting experiments and electrophysiological experiments.

**Code and data availability.** We used Python (codes written by V.M.K.N.) to analyse imaging and optogenetic datasets included in this manuscript (see Figs 1–5). That data, as well as the codes used for analysis, are openly available (https://github.com/stuberlab). All other data are available upon request from the corresponding author.
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Extended Data Figure 1 | Mice used for imaging experiments acquired cue-specific anticipatory licking across conditioning. a, Average lick rate during the 1-s baseline period (immediately before each cue delivery) for all imaging experiments (early, n = 30 sessions; late n = 30 sessions). b, Average lick rate during each cue (rather than the change in lick rate presented in the main figures) for all imaging experiments (early, n = 30 sessions; late n = 30 sessions). c, Individual behavioural discrimination (licking during CS⁺ versus CS⁻; auROC – 0.5) scores during early and late conditioning sessions for all imaging sessions used in this manuscript (early, n = 30 sessions; late, n = 30 sessions; t₅₈ = 17.6, P < 0.001). Line graphs represent the mean ± s.e.m. These data are presented in a summarized form in Fig. 1d, e.
Extended Data Figure 2 | Elevations and reductions in GCaMP6s fluorescence track action potential frequency but not voltage per se.

a, Virus injections of AAVdj-CaMK2a-GCaMP6s into dorsomedial PFC allowed subsequent whole-cell patch-clamp slice recordings of GCaMP6s-expressing neurons. Coronal cartoon redrawn based on ref. 40.
b, c, Example traces showing GCaMP6s fluorescence (b) during current-clamp recordings (c) in the absence (left) and presence (right) of action potentials (n = 8 neurons; n = 2 mice).
d, Grouped data revealing that hyperpolarization resulted in negative GCaMP6s fluorescence deflections in the presence of baseline action potentials, but not in the absence of baseline action potentials (interaction: F1,14 = 20.0; ***P < 0.001; post hoc tests: baseline action potentials, P > 0.4; no baseline action potentials, P < 0.001).
e, Example traces showing a series of depolarizing pulses (1–20 Hz) applied in current-clamp mode to drive trains of action potentials (bottom), during which GCaMP6s fluorescence was tracked in recorded neurons (top; n = 12 neurons; n = 2 mice).
f, Action potential generation resulted in linear elevations in GCaMP6s fluorescence (r = 0.776, ***P < 0.001), such that a single action potential was detectable (red waveform; peak = 12.6 ± 4.0% Δf/f0).
g, A series of baseline depolarizing pulses (1–20 Hz) was applied before and after a 3 s pause (n = 7 neurons; n = 2 mice).
h, The pause in action potentials resulted in linear reductions in GCaMP6s fluorescence (r = −0.656, ***P < 0.001), such that a 1 Hz reduction in firing frequency was detectable (red waveform; peak = −8.4 ± 2.0% Δf/f0). Line graphs represent the mean ± s.e.m. AP, action potential; ILc, infralimbic cortex; PLc, prelimbic cortex.
Extended Data Figure 3 | Cue responses in PFC CaMK2a-expressing neurons, PFC–NAc neurons, and PFC–PVT neurons before appetitive learning. a, b, Population heat plots showing average responses for all individual PFC CaMK2a-expressing neurons (n = 1,473 neurons; n = 8 mice) across all CS− trials before learning (a) and all CS+ trials before learning (b). c, d, Population heat plots showing average responses for all individual PFC–NAc neurons (n = 84 neurons; n = 4 mice) across all CS− trials before learning (c) and all CS+ trials before learning (d). e, f, Population heat plots showing average responses for all individual PFC–PVT neurons (n = 92 neurons; n = 3 mice) across all CS− trials before learning (e) and all CS+ trials before learning (f). Data shown here are from neurons presented in Figs 1 and 2. Vertical dotted lines refer to the time of sucrose delivery in CS+ trials or the equivalent time for CS− trials.
Extended Data Figure 4 | Cue discrimination in PFC CaMK2α-expressing neurons before and after appetitive learning.

a-c, Histograms for all recorded CaMK2α-expressing PFC neurons (early, \( n = 1,473 \) neurons; late, \( n = 1,569 \) neurons; \( n = 8 \) mice), showing CS+ responses (a), CS− responses (b), and CS+ /CS− discrimination (c) during both early and late Pavlovian conditioning sessions. d, CDF plot showing that the dynamics of individual CaMK2α-expressing neurons could be used to accurately decode whether the CS+ or CS− was presented in early conditioning sessions (compared to early shuffled: Welch’s \( t_{2,925.61} = 7.30, P < 0.001 \)), as well as in late conditioning sessions (compared to late shuffled: Welch’s \( t_{2,727.06} = 24.84, P < 0.001 \)). Data shown here are from neurons presented in Fig. 1. EarlySh, early shuffled; LateSh, late shuffled.
Extended Data Figure 5 | See next page for caption.
Extended Data Figure 5 | Corticostriatal and corticothalamic projection neurons are anatomically and electrophysiologically distinct. 

a, b, CtB-488 and CtB-594 were injected (n = 3 mice) into either NAc or PVT (scale bars, 250 μm) (a), resulting in retrograde labelling of NAc-projecting and PVT-projecting neurons in dorsomedial PFC (scale bars, 50 μm) (b). c–e, Coronal sections of anterior (c), middle (d), and posterior (e) dorsomedial PFC revealing spatial separation of PFC–NAc and PFC–PVT neurons (scale bars, 250 μm). f–h, Population histograms from all mice (n = 3) showing cell counts from anterior (f), middle (g), and posterior (h) dorsomedial PFC reveal that NAc-projecting neurons (red) are in more superficial layers as compared to PVT-projecting neurons (blue). Black refers to overlap (medial–lateral axis) between red and blue bars, and purple refers to double-labelled neurons (that is, both NAc- and PVT-projection neurons). i, Current-clamp recordings from dorsomedial PFC CtB-labelled neurons projecting to either NAc (top; n = 9 neurons; n = 3 mice; scale bars = 25 μm) or PVT (bottom; n = 10 neurons; n = 3 mice; scale bars = 25 μm). j, k, Representative waveforms (j) and averaged data (k) showing that PFC–NAc neurons fired fewer action potentials (spikes) as compared to PFC–PVT neurons during somatic depolarization (interaction: F_{16,272} = 16.6, ***P < 0.001). l, m, Representative waveforms (l) and averaged data (m) revealing no differences in rheobase (the minimum current required to evoke an action potential) between PFC–NAc and PFC–PVT neurons (t_{17} = 1.22, P > 0.2 (n.s., not significant)). n, o, Representative waveforms (n) and averaged data (o) showing that PFC–NAc neurons had larger peak afterhyperpolarizations (AHPs) as compared to PFC–PVT neurons (t_{17} = 4.67, ***P < 0.001). p, The maximum number of action potentials (spikes) in each neuron was correlated with the peak afterhyperpolarization (r = 0.80, P < 0.0001).

Line and bar graphs represent the mean ± s.e.m. cc, corpus callosum; ILc, infralimbic cortex; MO, medial orbitofrontal cortex; PLC, prelimbic cortex; VO, ventral orbitofrontal cortex.
Extended Data Figure 6 | Corticostriatal and corticothalamic projection neurons have distinct monosynaptic inputs. a–d, Viral strategy for rabies tracing experiments in which the monosynaptic inputs to PFC–PVT (a, b) and PFC–NAc (c, d) neurons were evaluated (n = 3 mice per group). e–g, Example images showing mCherry+ cells (TV A expression) (e), RV-GFP+ (rabies virus encoding GFP) cells (f), and overlap revealing mCherry+/RV-GFP+ cells (starter cells; mCherry−/RV-GFP+ cells are considered local inputs) (g). h, The number of local input neurons (nonstarter; only RV-GFP+ cells per section) to each projection population, as quantified by raw neuron count and by the per cent of starter cells for each mouse, was equivalent for PFC–NAc and PFC–PVT neurons (raw neuron count: t16 = 0.59, P = 0.56; per cent starter cells: t16 = 0.13, P = 0.90). i, Representative image showing RV-GFP but not mCherry expression in the anterior cingulate cortex (ACC). j, The number of input neurons from ACC was higher for PFC–PVT neurons as compared to PFC–NAc neurons (raw neuron count: t16 = 3.51; **P = 0.003; per cent starter cells: t16 = 3.31, **P = 0.004). k, Representative image showing RV-GFP but not mCherry expression in the lateral pre-optic area (LPO). l, The number of input neurons from the LPO was equivalent for PFC–NAc and PFC–PVT cells (raw neuron count: t16 = 1.77, P = 0.10; per cent starter cells: t16 = 0.20, P = 0.84). m, Representative image showing RV-GFP but not mCherry expression in the ventral hippocampus (vHipp). n, The number of input neurons from the ventral hippocampus was higher for PFC–NAc neurons as compared with PFC–PVT neurons (raw neuron count: t16 = 4.44; ***P < 0.001; per cent starter cells: t16 = 4.00, ***P = 0.001). o, Representative image showing RV-GFP but not mCherry expression in the ventral tegmental area (VTA). p, The number of input neurons from the VTA was equivalent for PFC–NAc and PFC–PVT cells (raw neuron count: t16 = 0.56; P = 0.59; per cent starter cells: t16 = 0.09, P = 0.93). Bar graphs represent the mean ± s.e.m. Note that no RV-GFP+ neurons were detected in any nucleus of the amygdala for either projection group.
Extended Data Figure 7 | Corticostriatal and corticothalamic projection neurons express CaMK2a and have distinct basal activity dynamics. a–d, Injections of AAV5-CaMK2a-eYFP into dorsomedial PFC and the retrograde tracer CtB-594 into NAc (a) or PVT (c) resulted in expression of eYFP in CtB-labelled PFC–NAc neurons (b) and PFC–PVT neurons (d). These data reveal that PFC–NAc and PFC–PVT are subpopulations of CaMK2a-expressing neurons (n = 2 mice per group). e, f, In a9 reporter mice, (e) AAVdj-DIO-GCaMP6s injections in dorsomedial PFC and Cav2-Cre injections into PVT (f) result in expression of GCaMP6s and tdTomato (marker for Cre-recombinase), which have spatial overlap in PFC (n = 2 mice). These data reveal that GCaMP6s expression is specific to the projection cells of interest.

g, Example traces revealing spontaneous calcium dynamics from in vivo two-photon imaging in GCaMP6s-expressing PFC–NAc neurons (top; n = 69 neurons; n = 4 mice) and PFC–PVT neurons (bottom; n = 63; n = 3 mice) in awake, head-fixed mice. Red and blue dots refer to detected events. h–j, Averaged data reveal no differences in event amplitude (h) or event duration (i); however, PFC–NAc neurons had significantly shorter inter-event intervals (j) as compared to PFC–PVT neurons (amplitude: t\_130 = 1.10, P > 0.2; duration: t\_130 = 0.68, P > 0.4; interval: t\_130 = 2.30, *P < 0.05). Bar graphs represent the mean ± s.e.m.
Extended Data Figure 8 | Effects of corticostriatal and corticothalamic optogenetic manipulations on acquisition and expression of CS-licking. 

a, Line graph showing average CS-lick rate during conditioning (with laser) and test (no laser) from PFC–NAc::ChR2 (n = 5), PFC–NAc::eNpHR3.0 (n = 6), and PFC–NAc::eYFP mice (n = 10). b, c, CDF plots and bar graphs showing CS-lick rate during conditioning (b) and test (c). No differences were observed between PFC–NAc groups during the no-laser test (ChR2 versus eYFP: auROC = 0.47, Benjamini–Hochberg corrected P = 0.45; eNpHR3.0 versus eYFP: auROC = 0.47, P = 0.45).

d, Line graph showing average CS-lick rate during conditioning (with laser) and test (no laser) from PFC–PVT::ChR2 (n = 6), PFC–NAc::PVT (n = 5), and PFC–PVT::eYFP mice (n = 10). e, f, CDF plots and bar graphs showing CS-lick rate during conditioning (e) and test (f). No differences were observed between PFC–PVT groups during the no-laser test (ChR2 versus eYFP: auROC = 0.53, Benjamini–Hochberg corrected P = 0.44; eNpHR3.0 versus eYFP: auROC = 0.35, P = 0.36). g–i, CDF plots and bar graphs showing CS-lick rates for PFC–NAc::ChR2 mice (versus PFC–NAc::eYFP: auROC = 0.43, P = 0.26), although there was an effect of laser for PFC–NAc::eNpHR3.0 mice (versus PFC–NAc::eYFP: auROC = 0.23, *P = 0.006). j–l, CDF plots and bar graphs showing CS-lick rates for PFC–PVT::ChR2 (n = 5), PFC–PVT::eNpHR3.0 (n = 5), and PFC–PVT::eYFP mice (n = 8). There were no significant differences in CS-lick rate for PFC–PVT::ChR2 mice (versus PFC–PVT::eYFP: auROC = 0.35, P = 0.15) or PFC–PVT::eNpHR3.0 mice (versus PFC–PVT::eYFP: auROC = 0.55, P = 0.31). Line and bar graphs represent the mean ± s.e.m. NL, no-laser test.
Extended Data Figure 9 | Effects of corticostriatal and corticothalamic optogenetic manipulations are timing dependent. a–c, CDF plots (top) and bar graphs (bottom) show anticipatory licking rates for PFC–NAc::ChR2 (n = 5) or PFC–NAc::eNpHR3.0 (n = 5) versus PFC–NAc::eYFP mice (n = 6) during sessions in which the laser was randomly presented outside of cue delivery. There were no significant differences in anticipatory lick rate for PFC–NAc::ChR2 mice (versus PFC–NAc::eYFP: auROC = 0.56, Benjamini–Hochberg corrected P = 0.30) or PFC–NAc::eNpHR3.0 mice (versus PFC–NAc::eYFP: auROC = 0.63, P = 0.23). d–f, CDF plots (top) and bar graphs (bottom) show anticipatory licking rates for PFC–PVT::ChR2 (n = 5) or PFC–PVT::eNpHR3.0 (n = 5) versus PFC–PVT::eYFP (n = 8) mice during sessions in which the laser was randomly presented outside of cue delivery. There were no significant differences in anticipatory lick rate for PFC–PVT::ChR2 mice (versus PFC–PVT::eYFP: auROC = 0.42, P = 0.21) or PFC–PVT::eNpHR3.0 mice (versus PFC–PVT::eYFP: auROC = 0.36, P = 0.19). g–i, CDF plots (top) and bar graphs (bottom) show CS− lick rates for PFC–NAc::ChR2 (n = 5) or PFC–NAc::eNpHR3.0 (n = 5) versus PFC–NAc::eYFP mice (n = 6) during sessions in which the laser was randomly presented outside of cue delivery. There were no significant differences in CS− lick rate for PFC–NAc::ChR2 mice (versus PFC–NAc::eYFP: auROC = 0.41, P = 0.19) or PFC–NAc::eNpHR3.0 mice (versus PFC–NAc::eYFP: auROC = 0.40, P = 0.19). j–l, CDF plots (top) and bar graphs (bottom) show CS− lick rates for PFC–PVT::ChR2 (n = 5) or PFC–PVT::eNpHR3.0 (n = 5) versus PFC–PVT::eYFP (n = 8) mice during sessions in which the laser was randomly presented outside of cue delivery. There were no significant differences in CS− lick rate for PFC–PVT::ChR2 mice (versus PFC–PVT::eYFP: auROC = 0.39, P = 0.12) or PFC–PVT::eNpHR3.0 mice (versus PFC–PVT::eYFP: auROC = 0.36, P = 0.12). Bar graphs represent the mean ± s.e.m.
Extended Data Figure 10 | Optogenetic manipulations of corticostriatal and corticothalamic neurons are not appetitive, aversive, and do not affect movement. Mice underwent a freely moving real-time place preference assay for 20 minutes to assess the rewarding or aversive value of PFC–NAc or PFC–PVT stimulation or inhibition (laser and group parameters are identical to head-fixed optogenetics experiments; see Methods). a, Tracking data from single example mice showing that PFC–NAc::ChR2 (left; \( n = 5 \)) and PFC–NAc::eNpHR3.0 (right; \( n = 5 \)) mice spent equivalent time in chambers that were paired with laser (PFC–NAc::eYFP mice, \( n = 8 \)). b, Grouped data show that laser stimulation in PFC–NAc mice did not lead to a real-time place preference (interaction: \( F_{2,30} = 2.15, P > 0.13 \)). c, Grouped data show that optogenetic manipulations in PFC–NAc mice did not influence velocity of movement (interaction: \( F_{2,30} = 0.12, P > 0.88 \)). d, Tracking data from single example mice during real-time place preference experiments showing that PFC–PVT::ChR2 (left; \( n = 5 \)) and PFC–PVT::eNpHR3.0 (right; \( n = 5 \)) mice spent equivalent time in chambers that were paired with laser (PFC–PVT::eYFP mice, \( n = 5 \)). e, Grouped data show that laser stimulation in PFC–PVT mice did not lead to a real-time place preference (interaction: \( F_{2,24} = 0.15, P > 0.86 \)). f, Grouped data show that optogenetic stimulation in PFC–PVT did not influence velocity of movement (interaction: \( F_{2,24} = 0.10, P > 0.90 \)). g, h, Coronal plates (redrawn from ref. 40) located 1.98, 1.78, and 1.54 mm anterior to bregma illustrate the placements of optical fibre tips in PFC–NAc mice (g) and PFC–PVT mice (h). Bar graphs represent the mean ± s.e.m. NoStim, no laser stimulation; Stim, laser stimulation.