Coordinating brain-distributed network activities in memory resistant to extinction

Highlights
- A brain-distributed beta-band (~20 Hz) activity pattern reports robust memory expression
- 4-Hz-paced VTA glutamatergic neurons actuate this transient cross-network coordination
- 4-Hz phase-driven VTA glutamate neuron silencing blocks post-extinction memory renewal

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In brief
The contribution of distributed brain activities to persistent memories was investigated by monitoring multiple neuronal regions during recall and renewal of cocaine-paired memory. A neural pattern actuated by the ventral tegmental area was revealed to coordinate and pace brain-distributed activities, whose suppression prevented the re-emergence of drug-biased behavior.
Coordinating brain-distributed network activities in memory resistant to extinction

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SUMMARY

Certain memories resist extinction to continue invigorating maladaptive actions. The robustness of these memories could depend on their widely distributed implementation across populations of neurons in multiple brain regions. However, how dispersed neuronal activities are collectively organized to underpin a persistent memory-guided behavior remains unknown. To investigate this, we simultaneously monitored the prefrontal cortex, nucleus accumbens, amygdala, hippocampus, and ventral tegmental area (VTA) of the mouse brain from initial recall to post-extinction renewal of a memory involving cocaine experience. We uncover a higher-order pattern of short-lived beta-frequency (15–25 Hz) activities that are transiently coordinated across these networks during memory retrieval. The output of a divergent pathway from upstream VTA glutamatergic neurons, paced by a slower (4-Hz) oscillation, actuates this multi-network beta-band coactivation; its closed-loop phase-informed suppression prevents renewal of cocaine-biased behavior. Binding brain-distributed neural activities in this temporally structured manner may constitute an organizational principle of robust memory expression.

INTRODUCTION

Memories typically support adaptation to the world we live in, allowing past experience to inform choices that are most appropriate to ongoing circumstances.1 However, not all memories adequately serve behavior this way. Notably, drugs of abuse such as cocaine produce powerful memories that associate drug experience with surrounding information, linking spatial context, discrete stimuli, and behavioral actions.2 Re-exposure to drug-associated stimuli evokes the memory recall of experiences involving cocaine, either directly or through indirectly related events. Thus, retrieval of cocaine-paired memories can invigorate drug-biased behavior even in contexts where the drug was never encountered. In such contexts, behavioral patterns of drug-biased actions eventually extinguish. But despite extinction, the chain of events linked to cocaine is not forgotten, and drug-paired signals can elicit the recovery of cocaine-biased behavior, a process known as renewal.3–8 This cocaine-paired memory retrieval, from initial recall to post-extinction renewal, increases the likelihood of further maladaptation. Previous studies have identified how cocaine alters molecular signaling, synaptic plasticity, and cellular activity (e.g., see references9–15). However, the brain-network-level mechanisms of cocaine-paired memory retrieval remain uncertain.

A central difficulty in identifying the network mechanisms of cocaine-paired memory retrieval could pertain to the distributed nature of this process, recruiting neurons from not one but multiple regions. Previous studies focusing on one brain region, or pairs of regions at most, have reported that important nodes in this distributed organization include the medial prefrontal cortex (PFC), the nucleus accumbens (NAc), the amygdala (Amy), and the hippocampus (Hpc).16–27 Neural dynamics in all these regions support memory-guided behaviors.28–35 In addition, the ventral tegmental area (VTA) has consistently been described as a central region for internal processing of reward/drug experience and memory.36–47 The VTA circuitry involves diverse cell types genetically defined to express dopamine, glutamate, GABA, or a combination of these neurotransmitters.48,49 Whether (and if so, how) the brain ultimately coordinates the neural activities distributed across these regions into a coherent, larger-scale (meta-) ensemble of neurons that collectively influence behavior remains unknown. In vivo electrophysiological recordings and theoretical work have highlighted that network oscillations report effective communication channels between distributed populations of neurons.50–57 Accordingly, here we investigate whether the dynamic retrieval of cocaine-paired memory involves selective patterns of network oscillatory activities that functionally couple neurons dispersed across multiple
A. Schematic representation of tetrodes and brain regions.

B. Wide-band local field potentials from different brain regions.

C. Illustration of the experimental setup for pre-test, conditioning, recall, extinction, and renewal.

D. Diagram showing the activity during different phases of the experiment.

E. Maps showing activity changes with different substances.

F. Graphs depicting the change in activity with cocaine minus saline LED activations over time.

G. Statistical analysis of cocaine-induced changes in LED triggering rates across different phases.

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regions, from initial recall to post-extinction renewal. Under this scenario, our overarching objective is to identify the cellular implementation of such cross-network coordination, leveraging this knowledge to deploy fine-grained feedback interventions for preventing re-emergence of cocaine-biased behavior.

To proceed, we developed a paradigm that models stages of cocaine-paired memory in mice while performing quintuple-brain-region recordings of network oscillations and neuronal ensembles from PFC, NAc, Amy, CA1 Hpc, and VTA. We designed an unsupervised analysis framework to detect and track activity patterns of multiregional coordination ("barcodes"). This unveiled a higher-order pattern of short-lived beta-band (15–25 Hz) neural co-activities during memory recall and renewal. This cross-network coordination is paced by the phase of a slower (4-Hz) oscillation in VTA, involving VTA glutamatergic neurons that project to the other regions. Activating VTA glutamatergic cells entrained neuronal spiking and induced cross-regional beta-band responses, orienting behavior toward place-cue stimuli paired with animal-position-controlled 4-Hz self-stimulation. Real-time closed-loop silencing of VTA glutamatergic neurons at their 4-Hz preferred phase disrupted cross-network beta-band coordination and prevented renewal of cocaine-biased behavior. Applying 4-Hz phase-informed silencing to the subset of VTA glutamatergic/dopaminergic neurons was sufficient to disrupt beta-band coordination and block memory renewal. These findings identify the oscillatory structure and anatomical pathway of a neural pattern that transiently binds neuronal populations distributed across brain regions in robust memory expression.

RESULTS

Modeling behavioral dynamics of cocaine-paired memory retrieval

To explore the network-level signature of cocaine-related memory, we developed a paradigm that models the dynamic expression of drug-biased behavior while permitting quintuple-brain-region recordings of both local field potentials (LFPs) and ensemble spike trains simultaneously from PFC, NAc, Amy, CA1 Hpc, and VTA (Figures 1A, 1B, S1A, and S1B). Our task leverages the conditioned place preference paradigm, measuring the propensity of the animal to self-activate a short-lived sensory cue (conditioned stimulus) by visiting a discrete location that is not itself directly linked to drug experience (Figures 1C–1G). This conditioned cue-place preference task uses a five-stage layout spanning multiple days through which mice acquire an indirect association between their ability to trigger an LED in one environment and an LED-paired cocaine experience in another environment.

In the first stage (Figure 1C; “pre-test,” 2 days), mice explored a bow-tie-shaped arena (“test enclosure”; Figure 1D) that contained two opposite zones of four unmarked quadrants. Using a place-controlled LED activation system, this stage allowed mice to learn two sets of place-LED associations where they could momentarily trigger one of two distinct LED wall displays (green L-shape versus orange circle61,62; mean LED activity duration = 392 ms; interquartile range = 42–634 ms) by moving within their corresponding zone (Figures 1C and 1D). We defined the last 20 min of this stage as the “pre-test” session, where we identified the initial (pre-conditioning) preference that each animal had for one LED while in the test enclosure and logged the corresponding behavioral pattern of place-controlled LED activations.

In the second stage (Figure 1C; “conditioning,” 3 days), mice associated each LED with either saline or cocaine in another square-shaped arena (“conditioning enclosure”). Despite being spatially distinct and not equipped with the place-LED controller, this enclosure shared some contextual features with the test enclosure. On each conditioning day, having received a saline injection, the animal explored this enclosure where the preferred LED now automatically flickered, emulating the behavioral pre-test LED activation pattern. The animal then received cocaine and re-explored the conditioning enclosure, where the non-preferred LED would automatically emulate the behavioral pre-test activation pattern (Figures 1E and S1C).

On the day following the last conditioning session, mice returned to the test enclosure (containing the place-LED controller) for the remaining three stages: “recall,” “extinction,” and “renewal” (Figure 1C; 1 day). Throughout these tests, we measured the animal’s propensity to activate each LED. During...
recall, mice inverted their pre-test preference for the cocaine-paired LED (Figures 1E–1G). This post-conditioning inversion was not likely explained by an LED novelty bias, since the place-LED preference initially expressed by mice remained similar across the 2-day pre-test stage (Figure S1D). This cocaine-conditioned LED preference evidenced that mice inferred across enclosures a second-order relationship between their ability to directly trigger each LED and their separate pairing with a substance.

Contextual alterations during transitions from stimulus-cued drug availability to unavailability decrease drug-motivated behavior, a phenomenon known as extinction. However, following extinction, re-exposure to discrete stimuli present within the drug-associated context revives drug-oriented responses, a phenomenon known as renewal. We modeled these phenomena in the final two tests. We altered the test enclosure to un-match the environmental features of the extinction stage to those experienced in pre-test and recall (i.e., different arena floor and no background white noise). Over time, the cocaine-paired LED preference decayed (Figure 1F). We defined the last 20 min of this session as extinction to permit comparable analyses with the other tests. For the renewal stage, the test enclosure was re-installed with all its original contextual features, and cocaine-biased behavior immediately recovered (Figures 1E–1G). Thus, mice renewed an extinguished association between their ability to activate an LED and a separate cocaine experience. Applying factor analysis to assess covariation between measures of zone-biased exploration further suggested that a common factor underlay recall and renewal (Figures S1E and S1F). Combining this conditioned cue-place preference task with electrophysiological recordings allowed us to determine whether specific patterns of brain network activity report cocaine-paired memory retrieval.

Multiregional beta-band coordination during memory retrieval

We developed an unsupervised analysis framework to identify neural patterns of higher-order coordination, which we refer to as cross-network activity “barcodes.” We extracted the region-tailored series of oscillatory signals composing each brain network’s LFPs (Figures 2A, S2A, and S2B). To focus on transient activities, we used signals spanning the 12–125 Hz frequency range and computed the time course of their instantaneous amplitudes (Figure 2A). These amplitude signals allowed us to detect moment-by-moment co-occurrence of short-lived activities, which were sampled over all animals to detect the barcodes (Figures S2C–S2I). Having validated the consistency of barcode detection across animals (Figures S2I and S2J), we computed each barcode strength over the time course of the multiregional LFPs throughout task stages (Figures 2A and S2G). With this, we computed for each barcode a cocaine-paired memory retrieval score to report the effect size for the difference in cocaine-versus-saline LED-zone strength with respect to changes in LED preference from pre-test to recall, extinction, and renewal (Figures 2B and S2K).

Remarkably, the barcodes that were significantly stronger in the cocaine-paired LED zone during recall and renewal (i.e., barcodes #3, #7 and #16; Figure 2C) all featured beta-band (15–25 Hz) signals (Figures 2A, 2D, and S3A). We confirmed this enhanced beta-band barcode strength with a linear mixed model that regressed out the effects of individual animals (barcode #3: p < 0.05; barcodes #7 and #16: p < 0.01; mixed-effects ANOVA model for zone + animal ~ strength, with Bonferroni correction for multiple comparisons [30 barcodes x 4 stages = 120]). A mixed-effects ANCOVA model further regressing out speed showed similar results. This speed- and animal-variation-accounted analysis revealed a significant cocaine-LED-zone-selective enhancement of beta barcodes during recall and their significant reduction in the post-recall session that progressed toward extinction when the learned drug-LED association was finally extinguished (recall: barcodes #3, #7, and #16, p < 0.05; extinction: barcodes #3 and #7, p < 0.001; barcode #16, p < 0.05; mixed-effects ANCOVA models for zone + animal + speed ~ strength, with Bonferroni correction for multiple comparisons; Figures S3B–S3D). When including theta-band signals in the barcode analysis, beta-band signals continued to feature most of the barcodes with significantly stronger expression in the cocaine-paired LED zone during recall and renewal (Figures S3E and S3F). Moreover, the amplitude of beta-band signals consistently increased in all regions during recall and renewal (i.e., stronger beta amplitudes in cocaine- than in saline-paired LED zones), a cross-regional change not observed for the lower (theta) nor higher (gamma) frequency bands (Figure S3G). These results showed that a cross-network activity pattern in the beta-frequency range reports cocaine-paired memory expression.

Cross-network beta-band modulation of neuronal spiking

We next investigated whether the identified beta-band network activity related to neuronal firing modulation. We detected bouts of high-amplitude beta in each region (Figure 3A; mean and standard deviation of beta-bout durations, 420.61 ± 21.50 ms; 491.80 ± 14.03 ms; 438.68 ± 7.44 ms; and 481.10 ± 16.67 ms for PFC, NAc, Amy, and Hpc, respectively). By using in turn each regional beta signal as a reference, we observed temporal modulation of individual neuron spiking with respect to beta-bout troughs (Figure 3B; using a single trough per beta-bout, ensuring trough separation by at least 250 ms; see also Figure S4A). Task-relevant information was tuned to beta-paced activity. We trained linear models (Linear Discriminant Analysis) on population vectors of spike discharge during recall to predict the active place-LED set (Figure 3C; mean number of neurons per model, 64.13; interquartile range, 51.00–83.50). Each model consisted of a cell-wise vector of coefficients whose magnitudes reflected the decoding contribution of each neuron (Figure 3D). Population decoding was significantly above chance during recall (Figure 3E; p = 2.393 × 10^-4, 1-sample t test), identifying with up to 79.31% accuracy the ongoing place-LED set (interquartile range 64.91%–76.96%; median 74.06%; chance level of 50%). Decoding accuracy remained above chance in the high behavioral cocaine-LED preference windows during the post-recall session progressing toward extinction, when decoding then fell to chance level as the learned place-LED association was extinguished (Figures S3B, S4B, and S4C). Recall-trained models performed above chance again during renewal with

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Figure 2. Cross-network beta-band activity barcodes report memory retrieval

(A) Example LFP traces (top) and their high-frequency (12–125 Hz) signals (middle, with instantaneous amplitudes as filled envelopes; color-coded, beta-band; gray, slow, mid, and fast gamma-bands). Corresponding cross-regional barcode strength time courses (bottom; color-coded and bold, barcodes #3, #7, and #16; fainter lines, all other barcodes). See also Figure S2.

(B) Barcode cocaine-paired memory retrieval scores. A higher score indicates that the barcode strength reflects memory recall and renewal (see Figure S2K). Data points, individual mice; bar charts, mean ± SEM over mice. *p < 0.05; 1-sample t test.

(C) Cross-test changes in cocaine minus saline zone modulation strength for the three barcodes with the highest memory retrieval scores in (B). Top, beeswarm plots of barcode strength during exploration (speed > 2 cm s⁻¹; 1 s windows) of saline-paired (gray dots) versus cocaine-paired (colored dots) LED activation zones across tests; gray lines, mean zone strengths for each animal; black squares/lines, distribution average. Bottom, corresponding mean difference. Black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval; open circles, animal mean differences. ***p < 0.001; **p < 0.01; permutation test, with Bonferroni correction for multiple comparisons (n = 4 stages).

(D) Barcode visualization. Note that beta-band signal interactions dominate these barcodes.
Figure 3. Local neuronal spiking is coupled to cross-regional beta oscillations

(A) Example beta-band signals from PFC, NAc, Amy, and Hpc (gray lines) with instantaneous amplitude envelopes (colored lines) and high-amplitude bouts (colored filled parts).

(B) Example beta-modulated cells for each region (one cell per row), with firing relative to beta-bout troughs (one trough per bout) in either PFC, NAc, Amy, or Hpc CA1 (one region beta-trough per column; 1.5-ms bins). Note that triggering neuronal spiking by beta troughs (separated by at least 250 ms) shows at least three peaks of rhythmic firing modulation matching beta frequency.

(C–F) Relationship between neuronal instantaneous spiking, beta-band coupling, and place-LED-substance association assessed using linear discriminant analysis models (see also Figures S4B–S4D). Each model was fitted to identify the active place-LED (set 1 versus set 2) from the ongoing population vectors of spike counts (100-ms windows) during recall (C), yielding a weight vector (example shown in D) with individual cell contribution to population decoding. Each model was applied in recall, extinction, and renewal (E); *p < 0.05, paired mean difference distribution estimation versus extinction, with Bonferroni correction for multiple comparisons; n = 2 stages. Cells with stronger contribution to population decoding exhibited higher spike-beta coherence (F).

(G) Time course of PFC, NAc, Amy, Hpc CA1, and VTA population firing relative to beta-bout troughs (NAc reference; 1.5-ms time bins).

(H) Example VTA cells with firing relative to troughs of PFC, NAc, Amy, or Hpc CA1 beta signals (25-ms bins). Note the rhythmic fluctuation of VTA spiking with a ~250-ms timescale.
cocaine-LED preference re-expression (Figure 3E; p = 1.364 × 10^{-2}, 1-sample t test; see also Figure S4D). This suggested that the preservation of spiking activity structure from recall to renewal related to the mnemonic features of cocaine experience and not to a contextual (sensory) distinction. We further found that the decoding contribution of individual cells positively correlated with their spiking coherence to cross-regional beta signals (Figure 3F; PFC, r = 0.405, p = 4.19 × 10^{-61}; NAc, r = 0.335, p = 2.75 × 10^{-45}; Amy, r = 0.360, p = 9.26 × 10^{-46}; Hpc, r = 0.360, p = 5.87 × 10^{-46}). Beta-paced neural activities thus appeared relevant to the goal-directed behavior in this task, and we sought to identify the biological implementation of this cross-network coordination.

By examining the population firing time course of each region, we observed that VTA cells exhibited a marked increase in firing before beta bouts (Figure 3G). When triggering individual VTA cell spiking with respect to the trough of the beta signals of the other regions, VTA firing further showed modulation over a slower (~250-ms window, i.e., ~ 4 Hz) timescale (Figure 3H). The VTA has consistently been reported as central to reward/ drug-oriented behavior and memory.36–46 This temporally structured firing relationship suggested that the VTA can organize the transient beta-paced activities distributed across the other brain regions.

**VTA 4-Hz oscillations pace distributed beta-band network activities**

We thus assessed how beta-band activities detected in individual regions are collectively orchestrated. Large-scale synchrony between multiple transient activities can be reported by slower oscillations.34,49–52,64–66 Notably, cross-regional phase-amplitude coupling by 4-Hz oscillations synchronizes faster (gamma-band) activities across hippocampal CA1, PFC, and VTA during spatial working memory.67 Here, we extracted 4-Hz signals from Hpc and VTA LFPs (Figures 4A and S2B). We thus evaluated the phase modulation exerted by these 4-Hz signals on each beta amplitude signal. PFC, NAc, and Amy betas were more strongly phase-modulated by VTA than Hpc 4-Hz (Figure 4B; p < 0.001, paired permutation test for VTA-minus-Hpc 4-Hz phase modulation scores, n = 4 regions). The amplitude of these beta oscillations shared the same phase preference for VTA, but not Hpc, 4-Hz (Figure 4B). While more strongly modulated by local 4-Hz, Hpc beta was still well-modulated by VTA 4-Hz, exhibiting an earlier phase preference compared to that shared by PFC, NAc, and Amy beta signals (Figure 4B). In fact, the region-pairwise difference between the VTA 4-Hz preferred phases of PFC, NAc, Amy, and Hpc betas decreased across conditioning days (Figures S5A–SSC), a phase alignment indicating synchronization of brain-distributed activities. VTA 4-Hz exerted preferential modulation of beta over the other faster (gamma) signals, which was not the case for Hpc 4-Hz (Figures S5D–SSG). Moreover, cross-regional spiking tended to be more modulated by VTA than Hpc 4-Hz (Figure 4C). Beta power relative to VTA 4-Hz troughs was also stronger for cocaine- than saline-paired LED activations during recall compared to pre-test, a contrast that disappeared in extinction but re-emerged during renewal (Figures 4D and 4E). A similar effect of VTA 4-Hz trough-related beta-band power occurred with another drug (morphine) in our place-LED task (Figures S6A and S6B), indicating that the identified pattern of beta coordination is not unique to cocaine. Interestingly, we observed a similar but weaker activity profile in the cocaine-conditioned place preference task (Figures S6C–S6F), suggesting that the described cross-network pattern of transient beta-band coordination is linked to the discrete sensory cues controlled by the animal’s behavior in our place-LED task. Altogether, these results relate VTA 4-Hz with the large-scale orchestration of brain-distributed beta-band activities.

VTA 4-Hz-paced glutamatergic cell spiking reflects multiregional beta coordination

We thus explored the cellular substrates that would allow VTA to engage in cross-network beta coordination. The VTA hosts diverse neuronal populations where dopaminergic and glutamatergic cells both influence downstream regions.37,40,48,68,69 To start assessing whether a particular VTA population supports cross-network coordination, we used the spike waveforms of individual VTA neurons as feature data to perform unsupervised cell clustering (Figure 5A). Extracellular recordings alone do not allow the direct identification of molecularly defined cell types. We thus also considered the spike waveforms of neurons recorded from the VTA of dopamine transporter (DAT)-Cre and vesicular-glutamate transporter-2 (VGLUT2)-Cre mouse lines, further optogenetically identified (optotagged) as dopaminergic and glutamatergic cells, respectively (Figures S7A–S7H). VTA optotagged VGLUT2 cells fell into one neuronal cluster, which...
we inferred to be the putative glutamatergic population (Figure 5A; cluster #2). VTA optotagged DAT cells spanned two putative dopaminergic clusters (Figure 5A; clusters #1 and #3).

Leveraging this neuronal clustering, we observed that the putative glutamatergic VTA population showed the strongest spike-phase coherence to VTA 4-Hz (Figure 5B), exhibiting the highest firing probability just before 4-Hz troughs (Figure 5C). Using the LFPs of each region, we then computed the spectral response to the spike times of each VTA population (controlling for the animal’s speed). Natural VTA glutamatergic spiking preceded transient increases of beta-band power in PFC, NAc, Amy, and Hpc (Figures 5D and 5F; cluster #2). VTA glutamatergic neurons, paced at 4-Hz, could thus play a central role in the coordination of brain-distributed beta-band activities.

Stimulation of VTA glutamatergic cells drives multiregional beta coordination and orientates exploratory behavior

To test this, we assessed the cross-network-level effects of VTA glutamatergic neuron activation. We transduced the VTA of VGlut2-Cre mice with the blue light (473 nm)-driven activator channelrhodopsin-2 (ChR2). In these VTA VGlut2::ChR2 mice, we performed parallel PFC, NAc, Amy, Hpc, and VTA ensemble recordings with brief VTA light-delivery (Figures 6A and 6B; 20-ms pulses). Remarkably, activating VTA VGlut2::ChR2 cells entrained neuronal spiking in all regions (Figure 6C), which suggested local computation of incoming inputs. This remote spike entrainment occurred with transiently enhanced beta-band power (Figures 5D and 5F; medium latency of beta power peak: PFC, 26.4 ms; NAc, 38.4 ms; Amy, 31.2 ms; Hpc, 32.8 ms). These brain-distributed spiking and spectral responses indicated that VTA glutamatergic cells could constitute a diverging pathway for the actuation of cross-network beta-band coordination.

We further evaluated the behavioral effect of VTA glutamatergic cell activation using 4-Hz-patterned optogenetic stimulation directly controlled by the animal’s location in the test enclosure (Figures 6E and 6F). For this, we identified the initial preference of each mouse for one place-LED set before any VTA light delivery. Mice next continued to explore the enclosure across additional sessions. In “VTA light-On” sessions, they could trigger 4-Hz-patterned stimulation of VTA VGlut2::ChR2 neurons by selectively entering the zone paired with their initially non-preferred LED (Figure 6E). These sessions alternated with “VTA light-Off” sessions with no stimulation (laser inactivated). Remarkably, in sessions with 4-Hz self-stimulation, mice switched their preference to the associated place-LED (Figure 6F). This showed that mimicking 4-Hz activity of VTA glutamatergic neurons is sufficient to reproduce the cocaine-biased behavior marking the recall and renewal stages of our conditioned cue-place preference task.

VTA 4-Hz phase-informed suppression of cross-network beta coordination prevents renewal of cocaine-paired memory

To determine whether glutamatergic VTA projections form a divergent pathway exerting a cross-brain-network influence, we transduced VTA VGlut2 neurons with the yellow light (561 nm)-driven silencer Archaierhodopsin-T (ArchT; Figures 7A and 7B). In these VTA VGlut2::ArchT mice, VGlut2-expressing VTA axonal projections innervated many regions including those recorded here (Figure 7C). Retrograde viral vector-mediated tract tracing confirmed that the VTA VGlut2 neurons projecting to one region (e.g., the Hpc) can also project to the others (Figures S7K–S7Q). These findings showed that VTA glutamatergic neurons instantiate a diverging (one-to-many-region) pathway.

To directly evaluate the role of this pathway in beta coordination during memory renewal, we deployed closed-loop optogenetic silencing of VTA VGlut2 neurons according to real-time tracking of VTA 4-Hz phase (Figure 7D). We applied this to both VTA and control VTA::GFP mice (where VTA glutamatergic neurons expressed the GFP-only construct), monitoring PFC, NAc, Amy, Hpc, and VTA activity combined with VTA light delivery at either the preferred or the non-preferred (opposite) 4-Hz phase subspace of VTA glutamatergic neuron firing (Figure 5C) during our task (Figures 7E–7G). By operating this feedback during the recall stage (without light delivery), we confirmed real-time detection of the highest versus the lowest VTA 4-Hz phase preference of VTA glutamatergic
closed-loop intervention with actual 561-nm light delivery during renewal significantly suppressed beta amplitudes in downstream (PFC, NAc, Amy, and Hpc) regions when applied at the preferred but not the non-preferred 4-Hz phase (Figure 7F). Critically, preventing beta-band coordination by closed-loop suppression of VTA glutamatergic cells selectively at their preferred 4-Hz firing phase prevented renewal of cocaine-biased behavior in VTAVglut2::ChR2 mice (Figure 7G). Closed-loop VTA light delivery at the preferred 4-Hz phase in VTAvglu2::GFP control mice did not prevent the re-emergence of cocaine-biased behavior (Figure 7G). These results showed that suppressing PFC–NAc–Amy–Hpc beta coordination by 4-Hz phase-informed silencing of the VTA glutamatergic population prevented post-extinction renewal of cocaine-biased behavior.

A large number of studies have implicated dopamine in memory (e.g., see references 38,39,46,71–73). Our results showed that cross-network beta-band activity reports cocaine-paired memory recall and renewal. Yet, by computing the LFP spectrograms of individual regions, we could not relate VTA dopamine population spiking to clear changes in beta-band power (Figure 5D). Our findings nevertheless indicated that the activity of some neurons (Figure 7E). Closed-loop intervention with actual VTA 561-nm light delivery during renewal significantly suppressed beta amplitudes in downstream (PFC, NAc, Amy, and Hpc) regions when applied at the preferred but not the non-preferred 4-Hz phase (Figure 7F). Critically, preventing beta-band coordination by closed-loop suppression of VTA glutamatergic cells selectively at their preferred 4-Hz firing phase prevented renewal of cocaine-biased behavior in VTAVglut2::ChR2 mice (Figure 7G). Closed-loop VTA light delivery at the preferred 4-Hz phase in VTAvglu2::GFP control mice did not prevent the re-emergence of cocaine-biased behavior (Figure 7G). These results showed that suppressing PFC–NAc–Amy–Hpc beta coordination by 4-Hz phase-informed silencing of the VTA glutamatergic population prevented post-extinction renewal of cocaine-biased behavior.
Figure 7. Phase-informed silencing of VTA glutamatergic neurons prevents renewal of cocaine-paired memory

(A–C) VTA VGlut2 diverging pathway. VTA of VGlut2-Cre mice transduced with ArchT-GFP (A). ArchT-GFP expression in medial VTA neurons (B; with tyrosine hydroxylase staining) and their projections in PFC, NAc, Amy, and Hpc (C). DAPI-stained nuclei. Bottom right, ArchT and VGlut2 co-expressing axon terminals in dorsal CA1.

(D) Quintuple-brain-region recording with VTA 4-Hz phase-dependent bilateral VTA yellow-light delivery targeting glutamatergic neurons expressing either ArchT-GFP or GFP-only.

(E) Laser activation as a function of VTA 4-Hz phase. Light (125 ms) pulse delivered with highest probability at the non-preferred versus the preferred phase subspace of VTA glutamatergic neurons (Figure 5C).

(F and G) Changes in beta power and cocaine-LED-oriented behavior.

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dopamine neurons is coupled to VTA 4-Hz (Figure 5C), which phase-relates to beta signal modulation (Figure 4). VTA dopamine neurons constitute a heterogeneous population, with some of its members co-expressing glutamate.48,74–78 We thus used an intersectional targeting strategy to manipulate the subset of genetically defined VTA glutamate neurons that are also dopaminergic (Figure 7H). We transduced the VTA of double-transgenic VGlut2-Cre; DAT-Flp mice with a viral construct for ArchT-eYFP expression conditional to the presence of both Cre and Flp recombinases (Figure 7I). We used these VTA\textsuperscript{VGlut2\textsuperscript{DAT}\textsuperscript{}} mice with quintuple-brain-site recordings and 4-Hz phase-locked light delivery during renewal, as before. Closed-loop suppression of VTA VGlut2-DAT neurons reduced the power of downstream beta-band signals in VTA\textsuperscript{VGlut2\textsuperscript{DAT}\textsuperscript{}} mice by 37.6% of that in VTA\textsuperscript{VGlut2::ArchT} mice (Figure 7J), along with preventing renewal of cocaine-biased behavior (Figure 7K). This showed that 4-Hz-paced VTA glutamate-dopamine neuron activity is necessary in post-extinction renewal of cocaine-paired memory expression.

**DISCUSSION**

In this study, we uncover a neural pattern of brain network activities that is actuated by a VTA diverging pathway and reports the initial recall and post-extinction renewal of cocaine-paired memory. This pattern couples a set of brain-distributed regions within a short (beta-band) timescale using 4-Hz-paced VTA glutamatergic neurons, whose closed-loop suppression prevents the re-emergence of cocaine-biased behavior.

**Cross-network coordination in retrieval of learned drug-cue-place association**

The renewal of cocaine-biased actions highlights the persistent nature of drug memories. This robustness could reflect the distributed nature of this type of memory, engaging neuronal populations from not one but multiple brain regions. Accordingly, we modeled cocaine-biased behavior from initial recall to extinction to renewal while recording network oscillations and neuronal ensembles from five regions (Figure 1). This selection was informed by previous work that dissected region- and pathway-specific contributions to separable components of cocaine-paired memory. For example, pharmacological manipulations have implicated Amy\textsuperscript{22} and NAc\textsuperscript{79} in cue-elicited drug-seeking and PFC\textsuperscript{80,81} and Hpc\textsuperscript{50,82,83} in context-dependent modulation of drug-paired memory retrieval. Numerous studies have implicated VTA in various aspects of drug-seeking behavior.15,42,84 Combining behavioral assays of drug-place conditioning with Hpc (and downstream NAc) recordings has recently begun to describe how drugs of abuse alter internal dynamics relevant to drug-associated memory.7,27,28,85 Our findings build upon these studies by revealing that the neuronal activities localized in each of these regions are organized into a coherent (meta-) ensemble, whose tight temporal structure serves memory expression.

By investigating higher-order patterns of coordination, our findings show that the structure of neural activity during recall and renewal of cocaine-paired memory resonates in the beta-frequency range (Figure 2). This could report various transient operations associated with goal-oriented behavior, including the processing of a global reward-seeking signal, the retrieval of learned drug-cue association, or the translation to a motivated behavior readout. By training machine learning models to identify the active place-LED set from population spiking vectors, we show that neurons representing task-relevant information are more coupled to beta signals (Figure 3). Neuronal spiking is modulated with reference to beta-frequency signals in either PFC, NAc, Amy, or Hpc CA1, in line with the observation that these signals are tightly coordinated. This beta-band coordination is consistent with theoretical work suggesting that neural synchrony over longer conduction delays (i.e., between distally connected regions) is more readily supported by the beta (rather than gamma) timescale.86,87 Our observation that cocaine-paired memory implicates cross-network beta coordination is also in line with the hypothesis that enhanced beta activity can report a cognitive state detrimental to flexible behavior.87 On the (brain) spatial scale, this preference for beta (fast, yet slower than gamma) coordination suggests the recruitment of larger neuronal ensembles.52,64,66 This could exert greater influence on an animal’s behavior, biasing it toward salience-associated stimuli. In line with this, past work showed that neuronal responses to reward-predicting stimuli increase with reward magnitude.86,87 The cross-network pattern identified here might thus report an “incentive salience” signature.2,30,91 Together, these findings support the view that the robustness of connected data points reporting the beta amplitude during renewal (laser-powered) compared to recall (laser-not-powered); lower panel, effect size for changes in beta power during renewal versus recall. **"p < 0.001, paired permutation test.**

(G) This intervention prevented renewal of cocaine-LED-oriented behavior in ArchT-GFP mice when performed at the preferred 4-Hz phase (ArchT Pref.). Upper panel, distribution of individual data points, with each set of four connected points reporting the difference in cocaine-paired minus saline-paired LED activations for one mouse across tests, compared to extinction (black-filled/unfilled, GFP-only/wild-type mice [Figure 1G]; red/green, ArchT-GFP; **"p < 0.01; **p < 0.05; repeated ANOVA versus extinction, with Bonferroni correction for multiple comparisons). Lower panel, effect size with paired mean difference for each test compared to extinction; black dot, mean difference; filled curve, distribution of mean differences; black lines, 95% confidence interval.

(H–K) VTA VGlut2; DAT neuron suppression reduced downstream beta amplitude and prevented renewal of cocaine-biased behavior.

(H) Intersectional strategy for VTA glutamatergic/dopaminergic neuron targeting in double-transgenic VGlut2-Cre; DAT-Flp mice injected with a Cre- and Flp-dependent ArchT-eYFP construct (top), used with quintuple-brain-region recordings and phase-informed bilateral VTA light delivery (bottom).

(I) VTA coronal section (at different magnifications) showing neuronal co-expression of ArchT-eYFP (green) and TH (magenta).

(J and K) This intervention reduced downstream beta power (J) and blocked cocaine-LED-oriented behavior (K) in renewal. **"p < 0.01, paired permutation test (J) and repeated ANOVA versus extinction, with Bonferroni correction for multiple comparisons (K).**

(L) Summary schematic. Retrieving a learned (drug-cue-place) association involves transient coordination of multiple short-lived beta-paced network activities that are co-enhanced in recall, decrease in extinction, and re-occur in renewal. This brain-distributed pattern is actuated by 4-Hz-paced VTA glutamatergic cells forming a one-to-many-region pathway. This cross-network coordination could yield a cohesive (meta-) ensemble of neurons serving robust memory expression.
of cocaine-paired memory expression relates to the temporally patterned coordination of distributed networks into a cohesive (beta-tuned) spiking structure. Via such distributed yet coordinated activities, downstream neurons would receive multiple inputs converging at a timescale that is fast enough to provide an effective neural syntax for memory and its translation into a behavioral output.50,57,92–94 Accordingly, identifying means to disrupt this cross-network organization would allow the prevention of renewal of drug-biased behavior.

A neural substrate of cross-regional communication

Cross-region communication is more likely facilitated by slower frequencies, which are coherent across brain structures and report rhythmic fluctuations of large-scale neural activity.34,55,57,84,96 Notably, 4-Hz phase modulation of gamma activities in PFC, Hpc, and VTA is enhanced during working memory demand.10 Our data-driven method extracted 4-Hz signals in VTA and Hpc (Figure 4). We observed strong VTA 4-Hz phase modulation of beta signal amplitude in PFC, NAc, Amy, and Hpc, which was also selective for beta over gamma signals in PFC, NAc, and Amy (Figures S5D–S5G). We also observed tighter preferred-phase alignment of PFC, NAc, and Amy beta signals for VTA compared to Hpc 4-Hz phase (Figure 4B). Moreover, VTA population firing increases prior to strong instances of beta-bout activity in the other regions (Figure 3G) and shows 4-Hz modulation when individual VTA cell responses are triggered by beta troughs (Figure 3H). These results suggested that VTA plays a leading role in the transient organization of cross-network activities during memory retrieval. This is consistent with work relating VTA to reward prediction signals.95–98 VTA also co-targets a large array of brain regions,99 including those recorded here. Notably, single-cell axon tracing revealed that individual VTA neurons can project to both cortical and limbic structures.100 These observations indicated the existence of a VTA diverging pathway actuating cross-regional coordination of short-lived beta activities with the phase of a slow (4-Hz) rhythm.

Identifying the cellular substrate of this network coordination remained to be elucidated. The VTA hosts diverse populations of molecularly defined neurons that can be distinguished by their spike waveforms.36,67,72,101–104 Accordingly, we leveraged an optogenetically validated VTA cell classification suggesting the glutamatergic population plays a central role in the actuation of multiregional beta coordination (Figure 5). Putative VTA glutamatergic neurons are most strongly coupled to local 4-Hz, with preferred firing toward the end of the descending phase (Figure 5C), just prior to the 4-Hz phase with highest beta amplitude (Figure 4B). Following natural spiking of VTA glutamatergic neurons, beta power markedly increased in PFC, NAc, Amy, and Hpc (Figure 5D). These observations could explain the phase alignment of cross-regional beta amplitudes to VTA 4-Hz (Figure 4B). Moreover, stimulating VTA VGlut2 neurons entrained neuronal spiking in all recorded regions and increased beta power (Figures 6A–6D). Prior work showed that VTA VGlut2 cells can shape goal-oriented behavior.37,40,105–108 Activating VTA VGlut2 is indeed behaviorally potent: 4-Hz patterned self-stimulation of VTA VGlut2 neurons was sufficient to orientate mice toward the associated place-LED set (Figures 6E and 6F), mimicking the behavioral bias observed during the recall and renewal of our task. Prior studies reported that mice prefer VTA VGlut2 stimulation at a higher (40-Hz) frequency.40,107 While this behavioral effect was not related to natural in vivo electrophysiological monitoring of VTA VGlut2 neurons, the reported preference for faster frequencies could reflect an inherently rewarding effect of VTA VGlut2 stimulation,40,105,108 rather than the actuation of a large-scale oscillatory substrate for mnemonic retrieval of rewarding experiences. Future studies could investigate these facets of reward-paired memory retrieval, their inherent relationships, and how they may be supported across electrophysiological, molecular, and anatomical domains. Interestingly, the VTA 4-Hz phase preference of the Hpc beta signal amplitude occurred earlier than those of PFC, NAc, and Amy beta signals (Figure 4B). This suggests that while VTA 4-Hz oscillations orchestrate multiregional beta coordination, part of this induction could be encouraged by hippocampal beta activity, perhaps broadcasting spatio-contextual information to downstream reader neurons. This may explain the tendency for stronger hippocampal beta signals in the recall and renewal stages of the conditioned place preference task (Figure 5F). In the conditioned place preference task, animals are not exerting direct control over the environmental cues. This difference between the two behavioral paradigms could influence the degree of reliance on short-lived binding of brain-distributed activities.

VTA VGlut2 cells form diverging projections (Figures 7A–7C and S7J–S7P).77,105 We thus hypothesized that a one-to-many-region VTA glutamatergic pathway actuates the large-scale coordination of beta activities during memory expression. We found that VTA 4-Hz phase-triggered silencing of VTA glutamatergic neurons suppressed beta signals in PFC, NAc, Amy, and Hpc and prevented re-emergence of cocaine-biased behavior (Figures 7D–7G). This outcome was obtained when silencing VTA VGlut2 neurons specifically at their preferred 4-Hz phase. This supports a 4-Hz-orchestrated brain-distributed network mechanism of robust memory retrieval actuated by the VTA glutamatergic population. Interestingly, we noted beta-band correlates of VTA putative dopaminergic population spiking (Figure 5D; VTA clusters #1 and #3), albeit weaker than that of VTA putative glutamatergic population (Figure 5D; VTA cluster #2). Numerous studies have implicated dopamine in drug experience and memory, and an increasing number of studies show that VTA dopamine neurons can co-express multiple neurotransmitters.48,49,74–78,99,109–111 By targeting the subset of VTA glutamate neurons expressing dopamine with 4-Hz phase-locked silencing, we found that the VTA glutamate-dopamine neuron subpopulation is required for memory renewal (Figures 7H–7K). Previous work showed that VTA dopamine cells receive local inputs from VTA glutamate cells.105,108 Our VTA ensemble recordings also showed that stimulation of VTA glutamate neurons entrains spiking activity of other local neurons (Figure 6C), many of which are likely to be dopaminergic.108 The reward-predicting properties of VTA dopamine neurons are well-established,36,39,41,98,112–115 with their firing activity being sustained on omission of expected reward following cocaine experience.42 These findings indicate that while the VTA glutamatergic population coordinates cross-regional targets during memory retrieval, it could further define a window wherein
short-timescale coordination of multiple networks is influenced by concomitant dopamine release. Theoretical work has predicted that glutamatergic inputs arriving from remote upstream circuits preferentially synchronize over the beta timescale. This timescale would permit the recruitment of distributed neuronal ensembles whose large spatial scale could be promoted by dopamine.

Limitations of the study
Our study identifies a brain-distributed pattern of short-lived beta-band activities that is actuated by VTA glutamatergic neurons and paced by a slower (4-Hz) oscillation. Our findings raise the exciting prospect that VTA neurotransmitter co-release at multiple temporally coordinated targets would provide fine control of transient cross-network ensemble activity and thus the complexity of moment-by-moment internal processing of mnemonic information. Yet, the precise relationships between VTA dopamine/glutamate release dynamics, cross-regional oscillatory coordination, internal processing of learned associations, and behavioral action to gain rewards/drugs remain to be elucidated. Future work aiming to delineate these relationships would require novel methodological approaches. Further, we considered five regions, but more are engaged by drug experience and memory. Whether additional networks participate in the described pattern of VTA 4-Hz-paced beta-band coordination remains to be assessed. This could leverage higher-density recordings. Additional regions would be expected to receive VTA glutamatergic projections. This study involved adult male mice able to carry a multiregional electrode drive for brain-distributed ensemble recordings. Whether the described pattern of cross-network coordination is generalizable to adult female mice and young mice remains to be assessed. By focusing on drug experience, our study supports the idea that binding together distinct networks in this temporally structured manner may constitute an organizational principle of robust memory expression. The extent to which this applies to other memories remains unevaluated. Moreover, drug-related memories can precipitate the transition toward loss of behavioral control and addiction. Our paradigm was confined to ~1 week. Monitoring brain networks over a much longer period in animals self-administering a drug would provide important insights into the neural dynamics of addiction.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.cell.2023.12.018.

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AUTHOR CONTRIBUTIONS

C.J.C.-W. and D.D. designed the study; C.J.C.-W., V.L.d.S., P.V.P., R.T., C.G.M., A.S., and D.D. developed the methodology; C.J.C.-W., L.L., D.B., R.R., A.A.C., K.H., and P.V.P. performed the experiments and acquired the data; C.J.C.-W. and D.D. analyzed the data; C.J.C.-W. and D.D. wrote the manuscript; and D.D. supervised the project. All authors discussed the results and commented on the manuscript.

DECLARATION OF INTERESTS

C.G.M. and A.S. are inventors on a pending patent application related to the phase-tracking algorithm used in this paper.

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### STAR METHODS

#### KEY RESOURCES TABLE

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### RESOURCE AVAILABILITY

#### Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, David Dupret (david.dupret@bndu.ox.ac.uk).

#### Materials availability
This study generated viral constructs for the Cre-dependent and Flp-dependent expression of ArchT-eYFP in small quantities and can be requested from david.dupret@bndu.ox.ac.uk while our stocks last, with priority to replication projects, and those entering into collaboration with us.
Data and code availability

- The electrophysiology dataset reported in this study is not yet openly available, as it is being used in ongoing projects. We welcome enquiries for sharing this as part of a collaboration, please contact david.dupret@bndu.ox.ac.uk.
- All original code has been deposited at Zenodo and is publicly available as of the date of publication. DOIs are listed in the key resources table.
- Any additional information required to reanalyze the data reported in this work paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

Animals

These experiments used adult (4–6 months old) male C57BL/6J wild-type mice (Charles River Laboratories, UK) and transgenic mice heterozygous for the transgene expressing the Cre recombinase under the control of either the vesicular-glutamate transporter-2 (VGlut2) or the dopamine transporter (DAT) promoter (Jackson Laboratories; obtained from C57BL/6J crossed with VGlut2-ires-cre B6.J.129S6(FVB)-Slc17a6tm1(cre)Nisw/J, stock number 028863, RRID: IMSR_JAX:028863; or with DAT-ires-cre B6.SJL-Slc6a3tm1.1(cre)Bkmn/J, stock number 006660, RRID: IMSR_JAX:006660) or for the transgene expressing the optimized Flp recombinase under the control of the DAT promoter (Jackson Laboratories; obtained from C57BL/6J crossed with DAT-P2A-Fip0 Slc6a3em1(cre)Yeo/J, stock number 005532, RRID: IMSR_JAX:005532) or with DAT-ires-cre B6.SJL-(VGlut2) or the dopamine transporter (DAT) promoter (Jackson Laboratories; obtained from C57BL/6J crossed with VGlut2-ires-heterozygous for the transgene expressing the Cre recombinase under the control of either the vesicular-glutamate transporter-2 (VGlut2) or the dopamine transporter (DAT) promoter (Jackson Laboratories). These experiments used adult (4–6 months old) male C57BL/6J wild-type mice (Charles River Laboratories, UK) and transgenic mice heterozygous for the transgene expressing the Cre recombinase under the control of either the vesicular-glutamate transporter-2 (VGlut2) or the dopamine transporter (DAT) promoter (Jackson Laboratories; obtained from C57BL/6J crossed with VGlut2-ires-cre B6.J.129S6(FVB)-Slc17a6tm1(cre)Nisw/J, stock number 028863, RRID: IMSR_JAX:028863; or with DAT-ires-cre B6.SJL-Slc6a3tm1.1(cre)Bkmn/J, stock number 006660, RRID: IMSR_JAX:006660) or for the transgene expressing the optimized Flp recombinase under the control of the DAT promoter (Jackson Laboratories; obtained from C57BL/6J crossed with DAT-P2A-Fip0 Slc6a3em1(cre)Yeo/J, stock number 005532, RRID: IMSR_JAX:005532).

Animals

These experiments used adult (4–6 months old) male C57BL/6J wild-type mice (Charles River Laboratories, UK) and transgenic mice heterozygous for the transgene expressing the Cre recombinase under the control of either the vesicular-glutamate transporter-2 (VGlut2) or the dopamine transporter (DAT) promoter (Jackson Laboratories; obtained from C57BL/6J crossed with VGlut2-ires-cre B6.J.129S6(FVB)-Slc17a6tm1(cre)Nisw/J, stock number 028863, RRID: IMSR_JAX:028863; or with DAT-ires-cre B6.SJL-Slc6a3tm1.1(cre)Bkmn/J, stock number 006660, RRID: IMSR_JAX:006660) or for the transgene expressing the optimized Flp recombinase under the control of the DAT promoter (Jackson Laboratories; obtained from C57BL/6J crossed with DAT-P2A-Fip0 Slc6a3em1(cre)Yeo/J, stock number 005532, RRID: IMSR_JAX:005532). Animals were housed with their littermates up until the start of the experiment. All mice held in IVCs, with wooden chew stick, nestlets and free access to water and food ad libitum in a dedicated housing facility with a 12/12 h light/dark cycle (lights on at 07:00), 19°C–23°C ambient temperature and 40–70% humidity. Experimental procedures performed on mice in accordance with the Animals (Scientific Procedures) Act, 1986 (United Kingdom), with final ethical review by the Animals in Science Regulation Unit of the UK Home Office.

METHOD DETAILS

Surgical procedure

All surgical procedures were performed under deep anesthesia using isoflurane (0.5–2%) and oxygen (2 L/min), with analgesia provided before (0.1 mg/kg ketoprofen) and after (5 mg/kg ketamine) surgery. To generate expression of either ChR2-eYFP, ArchT-GFP or GFP-only in VTA neurons, we used a Cre-loxP approach by injecting the Cre-inducible recombinant adeno-associated viral vector rAAV2-EF1a-DIO-hChR2(H134R)-eYFP-WPRE (from K. Deisseroth at UNC Vector Core), rAAV2-CAG-FLEX-ArchT-GFP (from E.S. Boyden at UNC Vector Core), or rAAV2-CAG-FLEX-GFP (from E.S. Boyden at UNC Vector Core), respectively. Viral injections were targeted bilaterally to VTA using stereotaxic coordinates (−3.1 mm anteroposterior from bregma, ±0.4 mm lateral from bregma, and −3.85 mm ventral from the brain surface; 150 nL per site) at a rate of 100 nL/min using a glass micropipette lowered to the target site and held in place for 5 min after virus delivery before being withdrawn. For electrophysiological recordings, mice were implanted with a single microdrive containing all (12 or 14) independently movable tetrodes. Tetrodes were constructed by twisting together four insulated tungsten wires (12 μm diameter, Doric Lenses, Canada) which were briefly heated to bind them together into a single bundle. Each tetrode was loaded in one cannula attached to a 6 mm long M1.0 screw to enable its independent manipulation of depth. The microdrive layout was organised on the horizontal plane of the brain so that the tetrode-holding cannulas were positioned according to the coordinates of region-centred craniotomies (see below), with individual tetrodes cut at the appropriate length so as to implant each tetrode ~300 μm above its target region on the vertical plane of the brain. The microdrive was implanted under stereotaxic control with reference to bregma. For dorsal hippocampal CA1, tetrodes were implanted by first identifying central coordinates −2.0 mm anteroposterior from bregma, +1.7 mm lateral from bregma as a reference to position each individual tetrode contained in the microdrive, initially implanting tetrodes above the pyramidal layer (−1.1 mm ventral from brain surface). A similar approach was used for tetrodes aimed at the medial prefrontal cortex using central coordinates +1.7 mm anteroposterior from bregma, +0.3 mm lateral from bregma, and an initial −1.5 mm ventral from brain surface; at the ventral tegmental area using central coordinates −3.1 mm anteroposterior from bregma, +0.4 mm lateral from bregma, and an initial −3.85 mm ventral from brain surface; at the nucleus accumbens using central coordinates +1.4 mm anteroposterior from bregma, +1.0 mm lateral from bregma, and an initial −4.0 mm ventral from brain surface; and at the amygdala using central coordinates −1.65 mm anteroposterior from bregma, +2.75 mm lateral from bregma, and an initial −3.8 mm ventral from brain surface. The distance between neighboring tetrodes inserted in each brain region was 350 μm. Following the implantation, the exposed parts of the tetrodes were covered with paraffin wax, after which the drive was secured to the skull using dental cement. For extra stability, four stainless-steel anchor screws were inserted into the skull before the drive was implanted. Two of the anchor screws, inserted above the cerebellum, were attached to 50 μm tungsten wires (California Fine Wire) and further served as a ground and reference electrodes during the recordings. For the recordings, each tetrode was lowered along the vertical axis to reach its target region, using the rotations applied to its tetrode cannula-holding screw and the spectral content of its LFPs, with final depth position subsequently confirmed by histology of anatomical tracks. For optogenetic manipulations, two optic fibers (230 μm diameter, Doric Lenses, Canada) were incorporated into the microdrive designed to bilaterally deliver light to VTA and implanted 10 days after viral injections.
To further assess whether the VTA glutamatergic neurons that project to one brain region also project to the other recorded regions, we performed retrograde viral vector-mediated tract tracing using VGlt2-cre mice injected with 50 nL of AAVrg-EF1a-DIO-FLPo-WPRE-hChR2-hGHpA (from Li Zhang Addgene viral prep # 87306-AAVrg; http://n2t.net/addgene:87306; RRID: Addgene_87306) in dorsal CA1 hippocampus (AP -2, ML ±1.2 and AP -2.3, ML ±1.4 mm from bregma, and DV -1.2 from the pia) and AAV5-EF1a-DIO-eYFP-WPRE (from K. Deisseroth at UNC Vector Core) in VTA (AP -3.2, ML +0.35 from bregma, and 4.02 mm from the pia under a 5° angle in the lateral to medial direction) (Figures S7J–S7P).

To assess whether the VTA neurons that are genetically defined to co-express glutamate and dopamine are required for cross-network beta-band signaling and renewal of cocaine-paired memory, we produced and injected a viral construct allowing the Cre-dependent and Flp-dependent expression of ArchT-eYFP in the VTA of double-transgenic VGlt2-Cre; DAT-Flp mice (Figures 7H and 7I). The corresponding pAAV-hSyn-Con/Fon-ArchT-eYFP construct was cloned in two stages. First, pAAV-hSyn-Coff/Fon-ArchT-eYFP has been cloned. Plasmid vectors pAAV-CamKII-ArchT-GFP (a gift from Edward Boyden, Addgene plasmid #37807; http://n2t.net/addgene:37807; RRID: Addgene_37807), pAAV-hSyn-Coff/Fon-hChR2(H134R)-eYFP (a gift from Karl Deisseroth, Addgene plasmid #55648; http://n2t.net/addgene:55648; RRID: Addgene_55648) and the combination of the PCR products was used to assemble two inserts that were then subcloned into the pAAV-hSyn Coff/Fon-hChR2(H134R)-eYFP vector to substitute the corresponding hChR2-eYFP coding exons with the ArchT-eYFP ones. Primers and the template plasmid DNA for the first insert (exon 1): GTTTCTGCTAGCAACCCCGACACTTACCTTAGCCAGCAGGGCCAG, GTTTCTGAGCTCGCCACCATGGACCC

Five-stage conditioned cue-place preference task
Following full recovery from the surgery, each mouse was first handled in a dedicated handling cloth, connected to the recording system, and exposed to an open-field enclosure to be familiarized with the recording procedure over a period of one week prior to the start of the experiment itself. During this period, tetrododes were gradually lowered to their corresponding target region using their estimated depth location, local field potentials and neuronal spike waveforms. An electrical commutator (Imetronic, Bordeaux, France) was installed above the dimly lit behavioral apparatus to hold the recording cable and allow its full rotation, preventing its twisting while the animal explored the enclosure.

The first stage of our ‘conditioned cue-place preference’ task occurred in the test enclosure (Figures 1C–1E; “pre-test”). This enclosure had a bow-tie shape (outer dimension: 46 cm width; 38 cm height) and contained two LED wall-displays placed on two opposing walls (green L-shape and orange circle-shape; Figure 1D). Along each wall holding an LED display, we defined two spatial zones of four equally sized, un-marked quadrants. We used this test enclosure with an open-loop system (Imetronic, Bordeaux, France) to track the animal’s position in real time and generate TTL-pulses that activate in turn the corresponding LED display for up to 1.5 s. To activate an LED again, the animal had to move into one of the neighboring zone quadrants. If the mouse moved to a neighboring quadrant while the LED was still active, this movement would briefly inactivate the LED (zone quadrant exit) before re-activating it (zone quadrant entry). Mice experienced the two sets of place-LED associations (Figure 1C) while exploring the test enclosure for two days (60 min on the first day exposure; 20 min on the second day exposure). The second exposure was used to quantify the baseline (pre-test) LED activation preference of each mouse.

The second stage of this task occurred in the conditioning enclosure (Figure 1C; “conditioning”), commencing 30 min after the pre-test and was repeated on two successive days. To proceed, the LED display that each mouse had preferentially activated during the pre-test was attached to one of the walls of the conditioning enclosure (square shape arena; 46 cm width; 38 cm height) and contained two LED wall-displays placed on two opposing walls (green L-shape and orange circle-shape; Figure 1D). Along each wall holding an LED display, we defined two spatial zones of four equally sized, un-marked quadrants. We used this test enclosure with an open-loop system (Imetronic, Bordeaux, France) to track the animal’s position in real time and generate TTL-pulses that activate in turn the corresponding LED display for up to 1.5 s. To activate an LED again, the animal had to move into one of the neighboring zone quadrants. If the mouse moved to a neighboring quadrant while the LED was still active, this movement would briefly inactivate the LED (zone quadrant exit) before re-activating it (zone quadrant entry). Mice experienced the two sets of place-LED associations (Figure 1C) while exploring the test enclosure for two days (60 min on the first day exposure; 20 min on the second day exposure). The second exposure was used to quantify the baseline (pre-test) LED activation preference of each mouse.

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solution (200 µL; obtained from Martindale Pharmaceuticals Ltd) to evaluate whether the observed cross-network pattern of VTA 4Hz-paced modulation of beta-band signals was specific to cocaine experience (Figures S6A and S6B).

The last three task stages occurred in the test enclosure (Figure 1C; “recall”, “extinction” and “renewal”), commencing 23 h after the last cocaine conditioning session and being all separated by 5 min (whilst the floor-contexts were changed; see below). In all these stages, the test enclosure was equipped with the place-LED controller, allowing mice to activate again the two LED displays as per the pre-test. In each of these three test stages, we quantified cocaine-biased behavior of each mouse using their LED trigger preference as the difference between the cocaine-paired LED activation rate minus the saline-paired LED activation rate. Both the recall and renewal stages lasted 20 min, with the test enclosure equipped with the floor used during pre-test. The extinction stage consisted of 10-min blocks during each of which the animal’s preference for the cocaine-paired versus the saline-paired LED was measured. To promote behavioral extinction of cocaine-biased actions, we replaced the floor used during the pre-test, recall and renewal stages by another floor, thereby introducing a contextual change within the test enclosure while preserving its spatial layout.\(^{132}\) Thus, during the course of this protocol each mouse encountered two floors in the test enclosure: a rough texture, light color floor and a smooth texture, dark color floor. One of these two floors was used specifically for the pre-test, recall and renewal stages; whereas the other was used for the extinction stage. The floor type assigned to the extinction context was counterbalanced across animals. To further promote behavioral extinction of cocaine-biased actions, a white noise was played during all these task stages but not during extinction. During this post-recall session, we considered that the learnt drug-cue-place association was extinguished when the animal exhibited for a minimum of 30-min a stable reduction in its preference for activating the cocaine-paired LED compared to the preference score measured during the recall stage and the first extinction block (mean number of extinction blocks required = 6.55; IQR = 4.00–7.00). For all analyses, the final 20-min period (last two 10-min blocks) defined the extinction stage and was used for comparisons with the 20-min pre-test, recall and renewal stages.

**Conditioned place preference task**

In a separate group of mice, we evaluated whether the identified pattern of transient coordination of cross-network beta signals could be generalized to the conditioned place preference (CPP) task. In this task, the animal performance is scored by the difference in time spent within the initially non-preferred (later cocaine-paired) compartment minus the time spent within the initially preferred (later saline-paired) compartment, over their sum. The CPP apparatus consisted of two compartments, the dimension of each of which was equal to that of the bow-tie shape test enclosure used in our conditioned cue-place preference task (outer dimension: 46 cm width; 38 cm height). The two compartments were connected via a bridge (8-cm length, 5-cm width), during the pre-test, recall, extinction, and renewal stages. As for the conditioned cue-place preference task, following full recovery from the surgery, each mouse was first handled in a dedicated handling cloth, connected to the recording system, and exposed to an open-field enclosure to be familiarized with the recording procedure prior to the start of the CPP experiment itself. During this period, tetrodes were gradually lowered to their corresponding target region using their estimated depth location, local field potentials and neuronal spike waveforms.

On the first CPP day, mice explored the entire apparatus for 20 min (pre-test) to determine their initial preference for one of the two compartments. Then, the bridge was removed and mice were conditioned for three days with two pairing sessions each day. Conditioning was performed with respect to the initial preference of each animal (as identified during the pre-test) for one of the two compartments.\(^{25}\) In the first session, mice received intraperitoneal injection of saline solution (200 µL), before exploring the preferred compartment for 30 min (saline-paired compartment). In the second session, 4 h later, mice received a 12 mg/kg cocaine solution (200 µL) before exploring the non-preferred compartment for 30 min (cocaine-paired compartment). On the day after (23 h after) the last conditioning session, we replaced the bridge between the two compartments and assessed CPP by allowing mice to re-explore the entire apparatus for 20 min (recall). The extinction stage next consisted of 10-min blocks during each of which the animal’s preference for the cocaine-paired versus the saline-paired compartment was measured, as in our conditioned cue-place preference task. To promote behavioral extinction of CPP behavior, we changed the floor used during the pre-test, recall and renewal stages by another floor while preserving its spatial layout, thereby introducing a contextual change similar to the one used within the conditioned cue-place preference task. Thus, during the course of this CPP protocol each mouse encountered two floors in each CPP compartment: a rough texture, light color floor and a smooth texture, dark color floor. One of these two floors was used specifically for the pre-test, recall and renewal stages; whereas the other was used for the extinction stage. The floor type assigned to the extinction context was also counterbalanced across animals. To further promote behavioral extinction of CPP behavior, a white noise was played during all these task stages but not during extinction. We identified the extinction stage as the last two 10-min blocks marked by a stable reduction in the animal’s preference for exploring the cocaine-paired compartment compared to the CPP score measured during the recall stage, as we did for the conditioned cue-place preference task. This final 20-min period defined the extinction stage and was used for comparisons with the 20-min pre-test, recall and renewal stages. We replaced the original floors and activated the white noise to conduct the renewal stage.

**Multichannel data acquisition, position tracking and light delivery**

Recordings were performed at the same time (i.e., 10.00 a.m. for Pre-test and Saline sessions; 2.00 p.m. for Cocaine and Post-test sessions) for all mice. The extracellular signals from each tetrode channel were amplified, multiplexed, and digitized using a single integrated circuit located on the head of the animal (RHD2164, Intan Technologies; http://intantech.com/products_RHD2000.html; pass band 0.09 Hz to 7.60 kHz). The amplified and filtered electrophysiological signals were digitized at 20 kHz and saved to disk.
along with the synchronization signals (transistor-transistor logic digital pulses) reporting the animal’s position tracking, LED display activations and laser activations. The location of the animal was tracked using three differently colored LED clusters attached to the electrode casing and captured at 39 frames per second by an overhead color camera (https://github.com/kevin-allen/positrack/wiki).

In addition, online location of the animal was used to trigger the corresponding LED displays in an open-loop manner, using the real time movement of the animal into the trigger quadrants (Imetronic, Bordeaux, France). This place-LED controller system was also used to activate 473-nm VTA light delivery for the 4-Hz patterned stimulation of VTA glutamatergic neurons in the VTA^Vglut2::ChR2 mice (Figures 6E and 6F). The LFPs were down-sampled to 1250 Hz for all subsequent analyses. For optogenetic interventions, 473-nm and 561-nm diode pumped solid-state lasers (Crystal Laser, models CL473-100 and CL561-100; distributor: Laser 2000, Ringstead, UK) were used to deliver light bilaterally to the VTA (~3–5 mW) via a 2-channel rotary joint (Doric Lenses Inc.).

To test for the required contribution of VTA glutamate cells in cocaine-paired memory renewal and cross-network beta coordination, we performed closed-loop optogenetic experiments to deliver VTA 561-nm light pulses in VTA^Vglut2::ArchT, VTA^Vglut2::DAT::ArchT, or VTA^Vglut2::GFP mice using dynamic tracking of ongoing VTA 4-Hz phase (Figures 7D, 7E, and 7H). The real-time phase estimate was obtained using the OscillTrack algorithm (https://doi.org/10.5287/bodleian:qa9ngXrzr) implemented in the field-programmable gate array of the Intan Technologies interface board. In these experiments, phase detection was obtained by continuously operating on the data stream coming from an input channel containing the VTA LFPs. This input channel used as the phase reference was high-pass filtered using a 1st order digital infinite impulse response filter with a corner frequency of 0.4 Hz to remove amplifier offset and electrode drift, then down-sampled 125-fold to a rate of 160 Hz for processing. The phase estimation was operated with a loop-gain of 0.0625 at a center frequency of 4 Hz. Stimulation was triggered with a phase lead, such that the target phase aligned with the middle of the 125-ms light pulse.

Spike detection and unit isolation

Spike sorting and unit isolation were performed with an automated clustering pipeline using Kilosort (https://github.com/cortex-lab/KiloSort) via the SpikeForest framework (https://github.com/flatironinstitute/spikeforest). To apply Kilosort to data acquired using tetrodes, the algorithm restricted templates to channels within a given tetrode bundle, while masking all other recording channels. All sessions recorded on a given day were concatenated and cluster cut together to monitor cells throughout the experiment day. The resulting clusters were verified by the operator using cross-channel spike waveforms, auto-correlation histograms and cross-correlation histograms. Each unit used for analyses showed throughout the entire recording day stable spike waveforms, clear refractory period in their auto-correlation histogram, and absence of refractory period in its cross-correlation histograms with the other units. Principal cells and interneurons were identified by their auto-correlograms, firing rates, and spike waveforms. In total, this study includes n = 2,602 neurons from quintuple-brain-site multichannel recording (PFC n = 421, NAc n = 292, Amy n = 247, Hpc n = 477, VTA n = 132 for the spiking analyses presented in Figures 3, 4, and 5; PFC n = 260, NAc n = 177, Amy n = 217, Hpc n = 308, VTA n = 70 for the spiking analysis presented in Figure 6).

QUANTIFICATION AND STATISTICAL ANALYSIS

Behavioral factor analysis

For each mouse, we collected a set of behavioral metrics (n = 11) to represent the multivariate nature of its cocaine-biased behavior during the recall, extinction, and renewal stages of our cocaine-paired memory task (see Table S1; Figures S1E and S1F). For each of these behavioral metrics, we then subtracted the corresponding baseline value measured in the pre-test stage. These pre-test adjusted metrics formed a feature matrix (test stages x behavioral metrics). Each metric column of this feature matrix was then normalized by a Z score transform such that factor analysis would not be biased toward metrics with higher numerical values. We then applied factor analysis using the scikit-learn 0.19.1 package (https://pypi.org/project/scikit-learn/).

LFP decomposition of individual brain regions into oscillatory components

For each mouse, we applied masked Empirical Mode Decomposition (mEMD) to break down the LFP raw trace recorded in each brain region during active exploration (training the algorithm using LFP traces from each animal corresponding to the continuous 5-min window which had the highest proportion of locomotion speeds above 2 cm s⁻¹) into its oscillatory components, namely the Intrinsic Mode Functions (IMFs; Figure S2) using the emd 0.2.0 package (https://emd.readthedocs.io/en/stable/). As an unsupervised sifting process, EMD successively extracts the highest frequency components of a time series until only a slow residual remains. In the mEMD approach, a masking signal is added at each sifting step to further prevent mixing of intermittent oscillatory components. Here, we used an iterative method, “tailored mEMD” (tmEMD; https://github.com/cjcw/tmEMD), to obtain in an unsupervised manner the set of masking frequencies used for each brain region in each animal, using the tetrode channel whose (z-scored LFP) power spectral density had the smallest Euclidean distance to the median power spectral density computed over all tetrodes across animals for that region. In the first iteration, we randomly generated 200 sets of masking frequencies. We then applied mEMD on the LFPs with each set of masking frequencies and computed a mixing score, defined as the absolute maximum of all pairwise correlations between the resulting IMFs. For each subsequent iteration, the ranges for the next sets of masking frequencies were restricted to the ranges observed in the top 10 current mixing scores. This iterative process converged to a set of masking frequencies that yielded the lowest mixing score for each brain region. As well as measuring mode mixing, the tmEMD...
algorithm also computed the between-animal consistency scores. Following the completion of the algorithm, each set of mask frequencies was given a score, defined as the mean of its mode mixing and consistency ranks. This score was then used to define the region-tailored mask frequencies to be used for mEMD. To assess the spectral consistency of the LFP signals across individual tetrodes, the PSDs of each tetrode IMF were combined to form a feature vector for each tetrode. We then applied Principal Component Analysis on the corresponding [tetrode x feature elements] matrix. To the first four principal components of this matrix (>97.6% of the variance explained), we applied t-distributed stochastic neighbor embedding (tSNE) to visualize the spectral similarity of tetrode oscillatory components across animals and regions (Figure S1A).

Cross-network activity barcode detection, tracking, and cocaine-paired memory score
To identify motifs of coordinated oscillatory activities within and between brain regions, we extracted network patterns of high-order (i.e., beyond one pair) IMF co-engagement (https://github.com/cjcw/barcode). To focus on short-lived, transient processes, we considered the IMFs with main frequencies between 12 and 125 Hz (Figure S2B). This selection yielded four signals for PFC, NAc, Amy and Hpc (i.e., 4 IMFs per region) and three IMF signals for VTA (Figure S2B; n = 19 total IMFs). We computed the time course of each IMF amplitude (Figure S2C, upper panel) using the Hilbert transform (scipy.signal.hilbert) and normalized each instantaneous amplitude by its standard deviation. We thus obtained a time-series matrix where the amplitude of each IMF had unitary variance (Figure S2C, lower panel). At each given time \( t \), we defined vectors \( \mathbf{a}_t \) that contained the normalized instantaneous amplitudes of these IMFs spanning the five brain regions (Figure S2D). We represented the set of instantaneous IMF amplitude co-modulation at time \( t \) by taking the outer product of its vectors \( \mathbf{a}_t \), only considering entries above and off the main diagonal such that all IMF pairs subsequently used were unique (i.e., each cross-IMF pair was only used once) and bipartite (i.e., no pairs were formed by a single IMF with itself) (Figures S2D and S2E). We used these entries obtained from the outer product of vectors \( \mathbf{a}_t \) as the elements of vector \( \mathbf{x}_t \), which therefore contained the set of instantaneous IMF co-engagement values (n = 171 cross-IMF pairs) at time \( t \) (Figures S2D and S2F).

To detect cross-network motifs of coordinated oscillatory activities that consistently occur over time (the cross-network “barcodes” of co-engaged oscillatory activities), we applied Independent Component Analysis on a feature matrix consisting of \( \mathbf{x}_t \) vectors sampled every 250 ms (Figures S2D and S2F). Each extracted independent component is a weight vector with individual elements corresponding to the amplitude co-modulation for a given pair of IMFs (Figure S2F). The strength of a given barcode at time \( t \) was then defined as the dot-product between the weight vector of its independent component and the instantaneous IMF co-engagement vector \( \mathbf{x}_t \). This computation was performed on the co-engagement time-series matrix (derived from the non-normalized amplitude matrix) at original time resolution (1.25 kHz) to track the instantaneous barcode strength (Figure S2G). For visualization purposes, each barcode weight vector was converted to a square matrix to depict its corresponding network motif of IMF co-engagement (Figures S2H and S2I). The barcodes obtained in this study were detected across all animals, and cross-validated to assess between-animal consistency (Figure S2J).

To identify the barcodes whose strength modulation showed the strongest signatures of cocaine-paired memory retrieval (i.e., increased expression strength in recall and renewal compared to pre-test and extinction), we computed a barcode cocaine-paired memory retrieval (CPMR) score (Figures 2B and S2K). For each barcode, we calculated the time course of its z-scored expression strength in each test stage using 1-s time bins while the animal explored the test enclosure (e.g., Figure 2A, bottom panel; Figure S2G). We then calculated the cocaine-minus-saline LED zone effect size between the obtained barcode strength distributions for each of the four task stages in the test enclosure (i.e., pre-test, recall, extinction and renewal) during active locomotion (speed>2 cm s\(^{-1}\)) bins. This procedure yielded a stage-wise vector containing the four barcode expression strength effect sizes for the difference between cocaine-versus saline-paired LED activation zones. The corresponding barcode CPMR score was obtained as the dot-product between this 4-element barcode strength vector and the 4-stage binary CPMR vector representing the task stages in the test enclosure (where element 0 corresponds to no cocaine-paired memory retrieval in the pre-test and extinction stages and element 1 corresponds to cocaine-paired memory retrieval in the recall and renewal stages). To measure the contribution of a particular frequency band to a given barcode (Figure S3E), a binary barcode vector mask was defined, where any pairwise interaction element involving that frequency band corresponded to a value of 1; and all other elements, 0. The contribution was then taken as the dot product between this mask vector and the barcode vector, divided by the sum of the mask vector.

Beta bouts and amplitude modulation strength
To detect bouts of high-amplitude beta in each recorded brain region, we used the extracted beta-band IMFs to isolate individual beta cycles. Each single cycle was defined by its ascending zero-crossing, positive-peak and descending zero-crossing; being in between two beta troughs, and with an additional ascending zero-crossing being detected after the second trough, before the next peak. The mean amplitude of each detected beta cycle was calculated from its instantaneous amplitude time course. We defined a beta bout as a chain of at least 5 consecutive cycles that all have a mean amplitude value above the 50th percentile of cycle amplitude values. The start-trough of the first cycle in the chain defined the beta bout start, and the end-trough of the last cycle in the chain defined the beta bout end.

To compare the modulation strength of the amplitude of a given time series by the phase of 4-Hz oscillations (Figure 4), we calculated the average z-scored amplitude within 32 equally spaced phase bins. The modulation strength was then defined as the amplitude difference between the most- and least-preferred bin indices. These phase modulation scores were used for statistical analysis relating to Figure 4B and to compare the modulation of the beta-band and (slow, mid, fast) gamma-band signals in Figures S5D–SSG.
To assess the effect of closed-loop VTA optogenetic silencing on cross-regional beta amplitudes (Figures 7F and 7J), each animal-region beta amplitude signal was z-scored and the average response with respect to the phase-triggered pulse onset was computed and compared between the recall and the renewal stages. Note that there was no actual light delivery during the recall stage (i.e., laser not powered), but an equivalent TTL pulse timestamp used for this comparison. The average amplitude was taken from 0 to 125 ms (TTL pulse-on) and from −125 to 0 ms (TTL pulse-off) windows. The difference between pulse-on minus pulse-off amplitudes is reported.

**Neuronal spiking activity**

To assess spiking correlates of beta-band signals, the instantaneous firing activity of recorded neurons was triggered by the troughs of the beta bouts detected in either PFC, NAc, Amy, or CA1 Hpc (using reference troughs separated by at least 250 ms; Figure 3B). To initially gather further evidence for cross-regional beta-band modulation of firing activity, we calculated for each neuron its spike-phase coherence (mean phase vector length) to detected beta signals during beta-bouts as: \( \text{abs}(\text{mean}(\text{e}^{j\theta})) \), where \( \theta \) is the phase of the spike-sampled phases (in radians) and \( j = \sqrt{-1} \). This coherence value was compared to a null distribution of coherence values, obtained by shuffling the beta bout windows, preserving both bout durations and inter-bout intervals. With this approach, a cell was deemed to be strongly beta-modulated if its beta bout coherence value was above the 95th percentile of the null distribution (PFC n = 23/421, NAc n = 25/292, Amy n = 3/247, Hpc n = 13/477, and VTA n = 19/132 cells). We then used the Rayleigh test (astropy 2.0.2) to assess modulation significance. The proportions of cells that are significantly beta-modulated across regions is reported in Figure S4A.

To explore the heterogeneity of the VTA cell population, we computed the average waveform for each recorded neuron using the tetrode channel with the highest spike waveform amplitude. This average waveform was then normalized by its maximum absolute value to construct a feature matrix (cells x waveforms). This matrix contained the spike waveforms of VTA cells recorded in wild-type mice during non-optogenetic experiments (i.e., where the molecular identity of recorded neurons was not optogenetically tested), along with the spike waveforms of both dopaminergic and glutamatergic VTA neurons recorded during light-off periods in DAT-Cre and VGluT2-Cre mice (where we further combined neuronal recordings with optogenetic stimulation for optogenetic identification; see also Figures S7A–S7H). We applied Principal Component Analysis on this feature matrix and used the first seven principal components (explaining >95% of the variance) as feature elements to Uniform Manifold Approximation and Projection (UMAP) for subsequent non-linear, dimensionality reduction. The projections of each VTA cell onto this UMAP embedding are shown in Figure 5A, and we obtained the VTA cell clusters using k-means (scikit-learn 0.24.4).

To assess whether the firing activity of a given cell was optogenetically modulated by light delivery (Figures 5, 6, and S7), the laser pulse times were randomly split into two-halves (half-1 and half-2). For each half-set of laser pulse times, we computed the spiking response of each VTA cell with respect to the onset of light delivery, averaging over all laser pulses and using 2-ms time bins. For a given cell, the time bin corresponding to the peak firing response in half-1 was used to sample the binned firing response in half-2. For a VTA cell to be determined as opto-tagged, this firing response had to be above the 99th percentile of binned firing response values observed between −1000 and −500 ms for both laser pulse sets, and the time corresponding to the maximum response evoked by half-2 had to be within 6 ms of the laser onset. For a cell to be considered as entrained by optogenetic activation of VTA VGluT2 cells, its maximum half-2 firing response (sampled by the peak response for half-1) had to be above the 95th percentile of values −1000 to −50 ms. Number of significantly entrained cells: PFC: 40/260; NAc: 57/177; Amy: 25/217; Hpc: 48/309; VTA 41/70 (9 of which were opto-tagged).

To compare the firing coupling of individual neuron members of a given VTA cell cluster (Figure 5A) to VTA 4-Hz oscillations, we computed population coherence percentile plots (Figure 5B). For each VTA cluster, this analysis allows visualizing the relative representation of spike-phase coherence values (see above) to the phase of VTA 4-Hz oscillations (y axis) above a given percentile threshold (x axis). To do so, we first calculated the whole-population percentile values corresponding to each single-neuron spike coherence observation in that cluster, then calculating the proportion of spike coherence values above a given whole-population percentile threshold. To test the significance of the over- or under-representation along the obtained curve, we computed a series of 1,000 “null curves” by repeatedly shuffling the group labels of all cells. For each VTA cell cluster, the corresponding population coherence percentile curve was deemed to represent significant over- or under-representation if falling outside of the 99% confidence limits of the null distribution. For each VTA cell cluster, the colored thicker line in its population coherence percentile curve represents significant over- or under-representation (Figure 5B).

To assess cross-regional spiking entrainment by VTA glutamatergic cell activation, we transduced VTA VGluT2-expressing neurons with the Cre-dependent viral construct encoding the blue light (473 nm)-driven neural activator channelrhodopsin-2 (ChR2) in mice expressing the Cre-recombinase under the control of the vesicular-glutamate transporter-2 (VGluT2) promoter (Figure 6). Optogenetic stimulation of these VTAVGluT2::ChR2 neurons was then obtained using 473-nm light 20-ms pulse delivery to VTA at random intervals (between 1 and 10 s) while performing ensemble recordings in PFC, NAc, Amy, Hpc and VTA. Neuronal spiking responses to light delivery were assessed as described above. The spiking responses of these cells (triggered by VTA light onset) were z-scored and used to construct the histogram and raster plots in Figure 6C.

**LFP spectrograms**

For triggered spectrogram analyses (Figures 4D, 5D, 6D, S6B, and S6F), the LFP of a given region was first z-scored and the spectrogram then computed using a set of complex Morlet wavelets (using scipy.signal.morlet function from the scipy 1.3.1 python module)
with main frequencies from 2 to 200 Hz, with 1-Hz steps up to 100 Hz and 5-Hz steps up to 200 Hz. To assess the changes in beta power relative to VTA 4-Hz troughs between LED activation zones across task stages, for each animal and test-stage, the resulting triggered spectral responses for saline- and cocaine-LED activation zones were z-scored for each frequency (taking the mean and S.D. over both zone spectrograms) before averaging over animals. Changes in beta power were then quantified by taking the median time of maximal beta across animal-zone responses (for each stage) to reference the z-scored beta power for each frequency within the beta range (Figures 4D, 4E, S6B, and S6F). To explore changes in cross-regional spectral power following spiking incidences of a given VTA cell cluster (Figure 5D), we used the spike train of each cell member of that cluster, considering both spikes discharged in isolation and the first spike discharged in a burst. A minimum of 20 spikes were required for this spike train to be used for triggering the average spectrogram responses centered around each spike for each of the recorded regions. As a control, the instantaneous speed of the animal was binned into 100 percentile bins. Each of the selected spikes were then randomly assigned a new speed bin-matched time, ensuring this did not correspond to a real spiking event for that cell. For each cell type, the corresponding triggered spectrograms of the real and speed-controlled spikes were averaged to compute the mean spectral response, and then z-scored for each frequency wavelet to assess changes with reference to spike times. For each cell, the speed-controlled spectrogram response was then subtracted from the original to produce the spike-triggered spectrogram responses, which were averaged over cells in that cluster.

For the light pulse-triggered spectrograms (Figure 6D), the average of each spectral response, centered by the onset of each pulse, was taken and averaged over animals. All spectrograms were plotted using the matplotlib.pyplot.contourf function.

**Anatomy**

Mice were deeply anesthetized with isoflurane/pentobarbital and transcardially perfused with PBS followed by cold 4% PFA dissolved in PBS. The brains were extracted, kept in 4% PFA for 24 h, and sliced into 50 μm thick coronal sections. Free-floating sections were rinsed extensively in PBS with 0.25% Triton X-100 (PBS-T) and blocked for 1 h at room temperature in PBS-T with 10% normal donkey serum (NDS). Sections were then incubated at 4 °C for 48 h with primary antibodies (chicken anti-GFP, 1:1000, Aves Labs, cat# GFP-1020, RRID: AB_10000240; guinea pig anti-TH, 1:2000, Synaptic Systems, cat# 213-104, RRID: AB_2619897; rabbit anti-Vglut2, 1:1000, Synaptic Systems, cat# 135–403, RRID: AB_887883) diluted in PBS-T with 3% NDS (“blocking solution”). After that, sections were rinsed three times for 10 min in PBS and incubated for 24 h at 4 °C in secondary antibodies (donkey anti-chicken Alexa Fluor 488, 1:500, Jackson ImmunoResearch, cat# 703-545-155, RRID: AB_2340375; donkey anti-guinea pig Cy3, 1:1000, Jackson ImmunoResearch, cat# 706-165-148, RRID: AB_2340460; donkey anti-rabbit Alexa Fluor 647, 1:500, Jackson ImmunoResearch, cat# 711-605-152, RRID: AB_2492288) diluted in PBS-T with 1% NDS. This step was followed by three rinses for 15 min in PBS. Sections were then incubated for 1 min with 4′,6-diamidino-2-phenylindole (DAPI; 0.5 μg/mL, Sigma-Aldrich, cat# D8417) diluted in PBS to label cell nuclei, before undergoing three additional rinse steps of 10 min each in PBS. Sections were finally mounted on slides, cover-slipped with Vectashield mounting medium (Vector Laboratories) and stored at 4 °C. Images were acquired on a laser scanning confocal microscope (LSM 880/Axio Imager, Zeiss).

**Data and statistical analyses**

Data and statistical analyses were performed in Python 3.6 (https://www.python.org/downloads/release/python-363/), using the packages scipy, **n**umpy, **m**atplotlib, **s**eaborn, **p**andas, **s**cikit-learn. **B**ar plots report group mean ± s.e.m., unless stated otherwise. All statistical tests related to a symmetric distribution were performed using two-sided using Gardner-Altman plots (to compare 2 groups) and Cumming plots (for more groups) as described in the Data Analysis with Bootstrap-coupled ESTimation (DABEST) statistics framework.144 These Difference Estimation plots allow visualizing the effect size by plotting the data as the mean or median difference between one of the groups (the leftmost group of each plot, used as group-reference) and the other groups (to the right, along the x axis of each plot). To estimate the effect size of unpaired observations from distributions A and B, the empirical mean difference is first calculated: \( \Delta = A - B \). For unpaired analyses (Figures 2, S3D, S3E, S6B, and S6F), the group mean difference \( \Delta \) is then subtracted from a series of null mean difference estimates, obtained by shuffling the group labels and then bootstrapping (n = 5,000; unless stated otherwise) from each of the shuffled distributions. This shuffling operation is repeated 10,000 times to obtain a probability distribution estimate of the true mean difference. Paired mean difference estimation plots (Figures 1, 3, 4, 6, 7, S1, S3C, S4, S5, S6A, and S6E) are computed in a similar fashion whereby for each shuffle, paired labels are randomly swapped or preserved with equal chance. For each estimation plot: (i) the upper panel shows the distribution of raw data points for the entire dataset, super-imposed on bar-plots reporting group mean ± SEM, unless stated otherwise; and (ii) the lower panel displays the difference between a given group and the (leftmost) group-reference, computed from 5,000 bootstrapped resamples and with difference-axis origin aligned to the mean or the median of the group-reference distribution. For each estimation plot: **black-dot**, mean (for normal distributions) or median (for skewed distributions) as indicated; **filled-curve**: bootstrapped sampling-error distribution; **black-lines**, error bars representing 95% confidence intervals (i.e., the 2.5th and 97.5th percentiles). For two-sided statistical tests, the percentile thresholds used are 2.5; 0.05; and 0.0005 for \( p < 0.05 \); \( p < 0.01 \); and \( p < 0.001 \), respectively. If a Bonferroni correction was used, then these percentiles were divided by the number of comparisons.
Figure S1. Consistency of tetrode LFP signals and behavior in the conditioned cue-place preference task, related to Figure 1
(A and B) Assessment of tetrode position. For the recordings, we used the spectral content of the LFPs from each tetrode as an electrophysiological readout of their position, adjusting it to the target region by using the tetrode cannula-holding screw that enabled their independent movement along the vertical axis.
(A) The spectral content of the LFP signals was consistent for tetrodes targeting the same region across animals (data visualized using t-distributed stochastic neighbor embedding [tSNE], with each color-coded data point representing the tetrode targeting a given region in one animal; see STAR Methods).
(B) After the recording experiment, tetrode positions were confirmed by histology of anatomical tracks (coronal sections imaged using dark field transmitted light, with additional staining for tyrosine hydrolase on the VTA section; yellow thick dashed lines show tetrode tips; yellow thin dashed lines show neighboring tracks with tips ending in other sections).
(C) Additional example animal paths (gray) during saline (left) versus cocaine (right) sessions of the conditioning stage. The green and orange dots overlaid on each path represent the animal’s position when the saline- and the cocaine-paired LEDs were automatically replayed, respectively.
(D) Stability of the initial place-LED preference across the first two days of the task. The difference in cocaine- minus saline-paired LED activations per minute is shown for the first 2-day stage (day 0 and pre-test) along with that in the recall stage. Negative numbers report preferred saline-paired LED activation; positive numbers report preferred cocaine-paired LED activation. Note that mice maintained across the first two days prior to conditioning their initial preference for one of the two place-LED sets. They then reverted this initial preference for the cocaine-paired LED during the recall stage following conditioning. The dataset is represented using a difference estimation plot to visualize the effect size for this cocaine-biased behavior. Upper panel, raw data points, with each set of three connected points reporting the difference between cocaine- minus saline-paired LED activation for one mouse across the first task session (day 0; before pre-test), pre-test, and recall; bar charts, average (mean ± SEM) LED activation preference across animals. Lower panel, corresponding effect size, using the mean difference estimation for each day compared to day 0. Black dot, mean difference; filled curve, distribution of mean differences; black lines, 95% confidence interval; *p < 0.01 paired permutation test compared to day 0, with Bonferroni correction for multiple comparisons.
(E and F) Factor analysis of mouse behavior during recall, extinction, and renewal. To evaluate the existence of a common process underlying the multivariate nature of cocaine-biased behavior in the recall, extinction, and renewal stages of our task, we applied factor analysis for dimensionality reduction of a set of behavioral metrics related to the animal’s response with respect to triggering cocaine- versus saline-paired LEDs in the test enclosure (see Table S1). (E) Factor analysis score and loading biplot. Each data point represents the score of the second factor versus the score of the first factor for one animal, color-coded according to recall, extinction, and renewal stages. Each arrow represents the loading of one behavioral metric on the factors. (F) Estimation plots to visualize the effect size in changes in animal factor loadings across recall, extinction, and renewal. Each set of three connected points in the upper panels show the raw loadings for each animal across the three task stages. The bottom panels show the paired mean difference estimation (effect size) compared to recall. Black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval.
Figure S2. Cross-brain-network activity barcode detection and consistency, related to Figure 2
(A) Example raw LFP traces (colored) with decomposed oscillatory components (intrinsic mode functions, IMFs) extracted using tailored masked Empirical Mode Decomposition shown underneath (gray) for each region.
(B) Power Spectral Density for each extracted IMF. IMFs with a main frequency within the 12–150 Hz range (colored) were selected for barcode detection.
(C) Instantaneous amplitude time courses for each of the selected IMFs from each region (top). These amplitudes were normalized by their standard deviation to produce the amplitude time-series matrix (bottom).
(D–F) Amplitude vectors ${a}_t$ were sampled every 250 ms (D) and converted to a co-engagement matrix by means of their outer product (E). Elements within the off-diagonal of this matrix were used to define the vector ${x}_t$, which represented all unique pairwise co-engagements of IMFs. These ${x}_t$ vectors were used to construct a feature matrix (F, top) to which Independent Component Analysis (ICA) was applied, resulting in 30 independent components (“barcodes”) (F, bottom).
(G) The instantaneous strength of a given barcode vector (e.g., barcode #9) was calculated by projecting its weight onto the non-normalized co-engagement time series.
(H) Each independent component vector was converted into a square matrix to visualize the co-engagement pattern it represented (e.g., barcode #9).
(I) Visualization of all barcodes detected across all animals (as in H), which were used for the analyses in Figure 2.
(J) Consistency of barcode detection between animals. Barcodes were detected for each animal individually to examine the consistency of barcode detection between animals. This is visualized using the tSNE algorithm. Open circles, barcodes detected for individuals; colored numbers, barcodes detected across the group data, as in (H) and (I).
(K) Schematic depiction of cocaine-paired memory retrieval (CPMR) score formula. For a given animal, the barcode strengths are sampled over 1-s windows of active (speed > 2 cm/s) exploratory behavior when the animal is in the saline- versus the cocaine-paired LED trigger zone in each test stage (pre-test, recall, extinction, and renewal). The effect size (difference of means, divided by standard deviation of all data) is then calculated, producing a test-stage-wise vector representing the four-stage barcode expression strength differences as cocaine-minus-saline effect sizes. To calculate the corresponding barcode CPMR score, the dot-product is taken of this 4-element barcode strength vector against the 4-stage binary vector representing the task stages in the test enclosure (i.e., the 4-element reference vector [0.0; 1.0; 0.0; 1.0] where element 0.0 corresponds to no cocaine-paired memory retrieval in the pre-test and extinction stages and element 1.0 corresponds to cocaine-paired memory retrieval in the recall and renewal stages).
Figure S3. Beta-band signals, barcode strength during extinction, and cross-network amplitude modulation through test stages, related to Figure 2

(A) Example bouts of beta-band rhythmic signals directly observable from the raw wide-band LFP traces of PFC, NAc, Amy, and CA1 Hpc. For each example, raw LFP traces (top, color-coded according to the region) and their corresponding spectrograms (bottom, with horizontal white dashed lines marking the beta-frequency range) are shown.

(B–D) Changes in beta-band barcode strength during the post-recall session progressing toward extinction. Shown is an example time course of cocaine-paired LED activation bias across the four stages in the test enclosure for one mouse (B) with test sessions labeled using gray fill along with the high- and low-preference epochs in the post-recall session progressing toward extinction, the corresponding assessment of the cocaine-minus-saline LED trigger preference (C), and the estimation plot (D) for changes in cocaine-minus-saline zone modulation strength for the three barcodes (# 3, #7, and #16) with the highest cocaine-paired memory retrieval scores (see Figure 2). (D) Top, beeswarm plots of barcode strength observations from 1-s windows when the mouse was active (speed > 2 cm/s) in the saline-paired (gray dots) or cocaine-paired (colored dots) LED zones; gray lines, mean zone strengths for each animal; bottom, corresponding cocaine-minus-saline mean difference estimation distribution for each stage, obtained from the beeswarm data points above; black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval; open circles, animal mean differences. ***p < 0.001; *p < 0.05; permutation test.

(E and F) Most of the LFP barcodes with significantly stronger expression in the cocaine-paired LED zone during recall and renewal feature high beta-band contributions, even when including theta-band signals in their detection. (E) Scatterplot showing the contribution of beta-band signals versus that of theta-band signals to each detected barcode (individual circles). Barcodes with significant CPMR scores (akin to Figure 2B; see Figure S2K for details) are shown as larger, red-filled circles. Note that only 1 barcode with high theta contribution has a significant CPMR score, while 4 barcodes with high beta contribution have significant CPMR scores. These significant barcodes are shown in (F).

(G) Estimation plots showing the cocaine-minus-saline LED zone difference in amplitude modulation for the oscillatory signals in the beta-band, (slow, mid, fast) gamma-band, and theta-band oscillations (rows) of each region (columns) through test stages. Changes in the amplitude of each signal computed when mice enter in the cocaine- minus the saline-paired LED activation zones for each task test stage (pre-test, recall, extinction, and renewal) using 1-s time windows of active exploratory behavior. In line with the barcode analysis, the only significant amplitude modulation consistently observed across regions during both recall and renewal concerns beta oscillations (i.e., stronger beta amplitudes in cocaine-paired zones for all regions in both recall and renewal). While theta amplitudes can increase in cocaine-paired zones during recall or renewal for some regions, this enhancement can also occur during extinction. ***p < 0.001; **p < 0.01; *p < 0.05; permutation test, with Bonferroni correction for multiple comparisons (n = 25).
Figure S4. Neuronal spiking coupling to oscillatory signals and decoding of active place-LED set, related to Figure 3

(A) Spike locking to frequency signals detected across brain regions. To compare the extent to which cells from each region were locked to individual oscillations, we applied the Rayleigh test for non-uniformity to the spike-sampled phases of theta, beta, slow-gamma, mid-gamma, and fast-gamma signals detected in each region. Numbers indicate the percentage of cells that are significantly modulated. For a cell to be included in this analysis, a minimum threshold of 200 spikes was imposed. To mitigate the effect of spike-leakage on coherence values, for local cell coupling to fast-gamma signals, the mean p value across local reference tetrode not associated with that cell was used.

(B–D) The relationship between neuronal spiking and active place-LED set during the post-recall session progression toward extinction (B; see also Figure S3B) was assessed using linear discriminant analysis (C) and polynomial support vector machine (D) models. Each model was fitted to identify the active place-LED (set 1 versus set 2) from the ongoing population vectors of spike counts (using 100-ms windows) during recall and then applied in the high- and low-preference epochs of the post-recall session progressing toward extinction, the extinction session, and the renewal session. Top row (B), ***p < 0.001; **p < 0.01; *p < 0.05, repeated ANOVA for pairwise stage-stage (versus extinction); top row (C and D), % % % % p < 0.01; % % p < 0.01; % p < 0.05; $ p = 0.051; 1-sample t test versus chance (50%). Bottom row (B–D): ***p < 0.001; **p < 0.01; *p < 0.05; paired permutation test versus the extinction stage, with Bonferroni correction for multiple comparisons.
Figure S5. Enhanced coordination of beta phase tuning to VTA 4-Hz under cocaine and preferential modulation of cross-regional beta signals by VTA 4-Hz, related to Figure 4

(A–C) The VTA 4-Hz coordination of beta activities detected in individual (PFC, NAc, Amy, and CA1 Hpc) regions improved across the 3-day cocaine conditioning. For each pair of brain regions, we computed the VTA 4-Hz preferred phase differences of corresponding regional beta amplitude signals. This was done for each animal, conditioning day, and conditioning session (saline versus cocaine). The VTA 4-Hz phase at which the beta amplitude was highest was used to sample the preferred phase, and then the differences between these preferred phases for each pairwise region combination were calculated to measure the strength of phase coordination by VTA 4-Hz for saline or cocaine (A).

(B) Cumulative distributions of these preferred phase differences. Note the sigmoid is steeper for cocaine observations in conditioning day 2/3, suggesting tighter phase-coordination under cocaine.

(C) Top, paired preferred phase difference observations for each animal-region pair, with the box-and-whisker plot indicating the minimum and maximum values, interquartile range, and median with the paired mean difference estimation plot shown below. Black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval. ***p < 0.001; **p < 0.01; determined from the cocaine- minus-saline paired mean difference estimation of region pairwise VTA 4-Hz preferred phase differences.

(D–G) Phase modulation of beta, slow-, mid-, and fast-gamma IMF amplitudes from PFC (D), NAc (E), Amy (F), and Hpc (G) by Hpc (left) and VTA (right) 4-Hz. Top of each panel, IMF amplitude as a function of 4-Hz phase; thick line, group mean; shaded error, ±SEM. Middle and bottom of each panel, paired Gardner-Altman estimation plot showing differences in phase modulation; middle, distribution of the raw data points, with each set of four connected points reporting the phase amplitude coupling for a given animal; lower, effect size, using the mean difference estimation for each signal amplitude compared to beta signal. Black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval. ***p < 0.001; **p < 0.01; *p < 0.05; determined from the paired mean difference estimation of phase modulation scores versus beta, with Bonferroni correction for multiple comparisons.
Figure S6. Changes in VTA 4-Hz-related beta-band power in two additional paradigms, related to Figure 4

A and B) Morphine conditioned cue-place preference paradigm.
(A) Behavioral place-LED preference (same task layout as with cocaine; Figure 1C). Estimation plot showing the effect size for changes in morphine-biased behavior across test stages (as in Figure 1G). Upper panel, raw data points, with each set of four connected points reporting the difference between morphine-minus saline-paired LED activation for one mouse in each stage; bar charts, average (mean ± SEM) LED activation preference. ## and ***p < 0.01; repeated ANOVA for pairwise stage-stage (#, vs. pre-test; *, vs. extinction) interactions, with Bonferroni correction for multiple comparisons. Lower panel, corresponding effect size, using the mean difference estimation for each stage compared to pre-test.
(B) Changes in VTA 4-Hz trough-triggered beta-band power between morphine-paired minus saline-paired LED activations over tests (as in Figure 4E). Top, each data point represents the beta amplitude triggered by individual VTA 4-Hz troughs for a given region (columns) for the saline-paired (gray) or morphine-paired (colored) LED zones across stages. Datapoints were bootstrapped (n = 10,000) for each animal-zone-stage distribution. Gray lines, means for each animal; black squares/lines, distribution average. Bottom, corresponding morphine minus saline mean difference estimation distribution. Black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval. ***p < 0.001; **p < 0.01; permutation test, with Bonferroni correction for multiple comparisons; n = 4 stages.

C–F) Conditioned place preference (CPP) paradigm.
(C) Task layout using five stages.
(D) Example animal paths during each stage on a given day (numbers indicate CPP scores, measured as the difference in time spent in the cocaine- minus the saline-paired compartments over the sum).
(E) Estimation plot showing CPP score across test stages. Upper panel, raw data points, with each set of four connected points reporting the CPP score for one mouse in each stage; bar charts, average (mean ± SEM) CPP score. ##p < 0.01; # and *p < 0.05; repeated ANOVA for pairwise stage-stage (#, vs. pre-test; *, vs. extinction) interactions, with Bonferroni correction for multiple comparisons. Lower panel, corresponding effect size, using the mean difference estimation for each stage compared to pre-test.
(F) Changes in VTA 4-Hz trough-triggered beta-band power between cocaine-minus saline-paired compartments over test stages. Top, each data point represents the beta amplitude triggered by individual VTA 4-Hz troughs for a given region (columns) for the saline-paired (gray) or morphine-paired (colored) LED zones across stages. Data points were bootstrapped (n = 10,000) for each animal-zone-stage distribution. Gray lines, means for each animal; black squares/lines, distribution average. Bottom, corresponding cocaine minus saline mean difference estimation distribution. Black dot, mean difference estimate; filled curve, distribution of mean difference estimates; black lines, 95% confidence interval. $p < 0.05, before Bonferroni correction; ***p < 0.001; permutation test, before and with Bonferroni correction for multiple comparisons; n = 4 stages.
Figure S7. Identification of VTA cell clusters and anatomical evidence for a divergent VTA VGlut2 neuronal pathway, related to Figures 5, 6, and 7

(A–F) We recorded the spike waveforms of optogenetically identified VTA dopaminergic (A–C) and glutamatergic (D–F) cells by transducing them with the Cre-dependent viral construct encoding the blue light (473 nm)-driven neural activator channelrhodopsin-2 (ChR2) in mice expressing the Cre-recombinase under the control of the dopamine transporter (DAT) or the vesicular-glutamate transporter-2 (VGlut2), respectively; subsequently implanting VTA tetrodes combined with VTA optic fibers for blue light delivery. (B) and (E) show VTA coronal sections with ChR2-eYFP expression in VTA cell bodies (along with tyrosine hydroxylase staining) in a DATVTA::ChR2 mouse (B) and a VGlut2VTA::ChR2 mouse (E). Scale bars, 200 microns. (C) and (F) show an example optotagged VTA cell recorded in a DATVTA::ChR2 mouse (C) and a VGlut2VTA::ChR2 mouse (F). Each example shows the histogram of the total spike count with respect to VTA light onset (top) and the corresponding raster plot showing the spiking incidences over individual light pulse events (bottom). At the top-right of each histogram is the spike waveform of the optotagged cell during the laser pulse (blue lines) and when the laser was off (black lines).

(G) Electrophysiological clustering of VTA neurons. The shape of spike waveforms was used in a dimensionality reduction algorithm (UMAP) to identify clusters of VTA neurons. This approach was applied to VTA neurons recorded in quintuple-brain-region-implanted wild-type mice (each ○ represents one wild-type VTA neuron) and to VTA neurons recorded and optogenetically identified (ChR2 optotagged) as dopaminergic (△) and glutamatergic (□) cells. The two optotagged cells colored in black (△ and □) correspond to the examples shown in (C) and (F).

(H) Spike waveforms of each VTA cluster. For each VTA cluster, each colored line represents the average spike waveform shape of a given VTA cell member of that cluster; the thick black line represents the waveform average of that cluster; and the dashed black lines represent the standard deviation. Spike waveforms normalized to maximum absolute amplitude. The spike waveform of optogenetically identified VGlut2 glutamatergic cells matched that of wild-type cells in cluster #2, which we thus designated as the VTA putative glutamatergic cell cluster. The spike waveform of optogenetically identified DAT dopaminergic cells matched that of cells in clusters #1 and #3, which we thus designated as VTA putative dopaminergic cell clusters.

(I) For each individual region (PFC, NAc, Amy, Hpc, and VTA), the average spectrograms constructed from the raw LFPs (top row for each region display) are shown with their corresponding speed-matched control spectrograms (bottom row of each region display), used to construct the spectrograms shown in Figure 5D. All spectrograms with respect to the discharge of action potentials by neurons from each VTA cluster (one cluster per column).

(J) Average LFP traces relative to optogenetic stimulation of VTA VGlut2:ChR2 neurons. (K and L) Intersectional approach for viral vector-mediated retrograde tract tracing of VTA VGlut2 neuron projections. The neuronal expression of the eYFP fluorescent report protein is made conditional to the activity of two Cre and FLPo recombinases (K). First, an AAVrg construct containing the Cre-dependent FLPo transgene (cDIO-FLPo) is injected in one brain region (here, the Hpc) of VGlut2-cre mice for retrograde transduction of the Cre-expressing VGlut2 neurons projecting to this (AAVrg-injected) target region (K and L). The VTA of the same mice was then injected with the second AAV5 construct (fDIO-eYFP) for FLPo-dependent eYFP expression (L).

(M–Q) Coronal sections from the VTA (M; along with tyrosine hydroxylase staining), PFC (N), NAc (O), Amy (P), and Hpc (Q) from the same VGlut2-Cre mouse injected as described in (K) and (L). Using this approach, we observed that targeting the dorsal Hpc allowed retrograde transduction of VGlut2 neurons located in the medial VTA (M; see also Figures 7A–7C). These Hpc-projecting VTA VGlut2 neurons also send axonal projections to PFC (N), NAc (O), and Amy (P); axonal projections are also shown for Hpc (Q). This viral tract tracing thus shows that individual VGlut2 glutamatergic VTA neurons can project to multiple brain region targets, thereby constituting a divergent pathway that reaches all recorded brain regions from VTA. Cell nuclei stained with DAPI. Anterior cerebral artery, aca; basolateral/dorsolateral amygdala, BLA/LaDL; stratum oriens, s.o.; stratum pyramidale, s.p.; stratum radiatum, s.r.; stratum lacunosum-moleculare, s.l.m.; granule cell layer, gcl.