1 Control of Parallel Hippocampal Output Pathways by Amygdalar Long-Range Inhibition.

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7 ABSTRACT

Projections from the basal amygdala (BA) to the ventral hippocampus (vH) are proposed to provide information 9 about the rewarding or threatening nature of learned associations to support appropriate goal-directed and 10 anxiety-like behaviour. Such behaviour occurs via the differential activity of multiple, parallel populations of 11 pyramidal neurons in vH that project to distinct downstream targets, but the nature of BA input and how it 12 connects with these populations is unclear. Using channelrhodopsin-2-assisted circuit mapping in mice, we show 13 that BA input to vH consists of both excitatory and inhibitory projections. Excitatory input specifically targets BA-14 and nucleus accumbens-projecting vH neurons, and avoids prefrontal cortex-projecting vH neurons; while 15 inhibitory input preferentially targets BA-projecting neurons. Through this specific connectivity, BA inhibitory 16 17 projections gate place-value associations by controlling the activity of nucleus accumbens-projecting vH neurons. Our results define a parallel excitatory and inhibitory projection from BA to vH that can support goal-18 directed behaviour. 19

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22 INTRODUCTION

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The hippocampus is key for episodic memory, learning and spatial navigation, as well as motivation, affect and 24 anxiety (Gray and McNaughton, 2003; O'Keefe and Nadel, 1978; Strange et al., 2014; Wikenheiser and 25 Schoenbaum, 2016). At almost every level of investigation - from gene expression, to afferent and efferent 26 connectivity, and behavioural function - the hippocampus is organised as a gradient along is dorsal to ventral 27 (posterior to anterior in humans) axis (Fanselow and Dong, 2010; Strange et al., 2014). Within this axis the most 28 dorsal portion is proposed to be involved in learning and utilising fine-grained spatial and temporal structure. 29 whereas the most ventral pole is thought to be involved in affect and motivation, and has a key role in value 30 based and reward driven decision making and anxiety-like calculations (Fanselow and Dong, 2010; Strange et 31 al., 2014). 32

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A distinguishing factor that separates the ventral from the dorsal hippocampus is dense input from the 34 corticobasolateral nuclear complex of the amygdala (basal amygdala, BA; McDonald and Mott, 2016; Strange 35 et al., 2014). The BA is comprised of a diverse set of nuclei including the basolateral amygdala (BLA), the 36 37 basomedial amygdala (BMA), the medial amygdala (MEA) and cortical amygdala, each of which sends projections to vH (McDonald and Mott, 2016; Petrovich et al., 2001; Strange et al., 2014). These nuclei, and 38 their projections to vH are thought to be crucial for the learning of reward and threat associated cues, and for 39 the generation of appropriate goal-directed and anxiety-like behaviour (Beyeler et al., 2018, 2016; Felix-Ortiz et 40 al., 2013; Felix-Ortiz and Tye, 2014; Hitchcott and Phillips, 1997; McHugh et al., 2004; Pi et al., 2020; Richardson 41

et al., 2004; Selden et al., 1991; Sheth et al., 2008; Yang and Wang, 2017). Thus, it is commonly assumed that
powerful and specific synaptic connectivity between these two structures is crucial for the maintenance of such
behaviours. However, there is limited information describing the organisation of such functional connectivity
between amygdala input and neurons in vH (Bazelot et al., 2015; Felix-Ortiz et al., 2013; Pi et al., 2020).

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This lack of understanding is compounded by the fact that the ventral hippocampus, in particular its output 6 structure the ventral CA1 and subiculum – where the majority of BA input is found – is organised as a parallel 7 circuit, such that the majority of neurons project to only one downstream area (Gergues et al., 2020; Naber and 8 Witter, 1998; Wee and MacAskill, 2020). Thus, while vH has powerful connections to the nucleus accumbens 9 10 (NAc), the prefrontal cortex (PFC) and back to the BA, each of these projections arises from a distinct population of neurons. Importantly each of these projection populations is increasingly shown to underlie unique 11 behavioural functions (Adhikari et al., 2010; Jimenez et al., 2018; LeGates et al., 2018; Sanchez-Bellot and 12 MacAskill, 2020). For example, vH^{NAc} neuron activity is high during motivated behaviour and around rewarded 13 locations (Ciocchi et al., 2015; Okuvama et al., 2016; Reed et al., 2018), is necessary for place-value 14 associations (LeGates et al., 2018; Trouche et al., 2019) and can promote spatial and instrumental reinforcement 15 (Britt et al., 2012; LeGates et al., 2018). In contrast, vHPFC activity is proposed to support the resolution of 16 approach avoidance conflict, and contribute to spatial working memory (Padilla-Coreano et al., 2016; Sanchez-17 Bellot and MacAskill, 2020; Spellman et al., 2015), while vH^{BA} activity is proposed to support contextual learning 18 (Jimenez et al., 2018). However, it remains unclear how the activity of these distinct populations in vH is 19 differentially controlled to promote these functions. We reasoned that a means for this control would be 20 projection-specific innervation from BA. 21

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The circuit organisation of the nuclei in the BA is similar to classic cortical circuitry – with the majority of neurons classed as either excitatory pyramidal neurons, or local inhibitory interneurons (McDonald and Mott, 2016). However, there is also evidence suggesting the presence of long-range inhibitory projection neurons throughout BA (Dedic et al., 2018; McDonald et al., 2012; McDonald and Zaric, 2015; Seo et al., 2016). Similar inhibitory projections from cortex are hypothesised to have a crucial regulatory role in modulating hippocampal circuit function (Basu et al., 2016; Melzer et al., 2012), but the connectivity and function of BA long range inhibitory input in vH has never been directly investigated.

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In this study, we used a combination of retrograde tracing, electrophysiology and channelrhodopsin-2-assisted 31 circuit mapping to show that BA provides both excitatory and direct inhibitory input to distinct projection 32 populations within vH. We show that excitatory projections uniquely target vH neurons that project to NAc and 33 34 back to the BA, and do not connect with neurons that project to PFC. In contrast, long range inhibitory input preferentially targets BA projecting vH neurons. Next, using a simple network model constrained by our 35 36 electrophysiology recordings, we predicted that the ability of BA input to drive motivation- and value-promoting vH projections to NAc was dependent on the coactivation of inhibitory input from BA. Finally, we confirmed these 37 predictions using in vivo optogenetics and genetically targeted pharmacology to show that long range inhibition 38 is required for the generation of spatial place preference. Together our results outline a novel inhibitory projection 39

- 1 from amygdala to ventral hippocampus that defines the activity of vH output neurons, and is able to control
- 2 hippocampal output to promote the formation of spatial place preference.
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1 RESULTS

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BA input into ventral hippocampus is both excitatory and inhibitory.

While the majority of investigation of BA-vH connectivity 5 is focussed on projections specifically from the BLA, it is 6 known that multiple BA nuclei project to vH (McDonald 7 and Mott, 2016). Therefore, we first determined the 8 spatial distribution of neurons in BA that send input into 9 vH by injecting a fluorescently conjugated cholera toxin 10 beta subunit (CTX β) into the ventral part of the 11 hippocampus (Figure 1A). $CTX\beta$ is taken up by 12 13 presynaptic terminals at the injection site, and retrogradely transported to label the soma of afferent 14 neurons. After two weeks, we serially sectioned labelled 15 brains and mapped labelled cell locations to the Allen 16 Brain atlas (Fürth et al., 2018; Wee and MacAskill, 2020). 17 We found that neurons sending input to vH were widely 18 dispersed throughout the entire BA, including in BLA, 19 BMA, and MEA, as well as in more cortical amygdala 20 areas (Figure 1B-D, McDonald and Mott, 2016; Strange 21 et al., 2014). Overall, this experiment confirmed that 22 23 there is large input from disperse BA nuclei to vH, focussed around the posterior BMA and BLA. 24

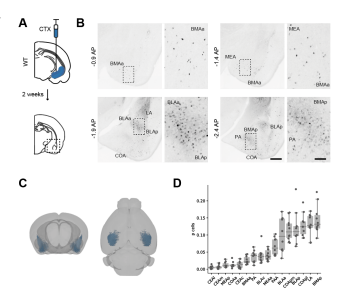


Figure 1 | Distribution of BA input to vH.

A. Schematic of experiment. $CTX\beta$ was injected into vH, 2 weeks later coronal slices of BA were examined for retrogradely labelled neurons.

B. Example slices showing widespread labelling throughout numerous BA nuclei. Scale bar = 500 um, 100 um (zoom).

C. Whole brain distribution of labelled BA neurons.

D. Summary showing proportion of labelled BA cells in each nuclei.

Full size figures are reproduced at the end of the manuscript.

We next tested whether BA input to vH may be both excitatory and inhibitory (McDonald and Mott, 2016). We 26 repeated our experiment using a vGAT::cre::dtomato reporter mouse. In this experiment, CTX β labelled neurons 27 in BA could be distinguished as either GABAergic (vGAT+), or putatively excitatory (vGAT-) based on 28 fluorescence colocalization. Using this approach, we found that a small but consistent proportion of BA neurons 29 that projected to vH were GABAergic (Figure 2A, Figure S1A, B). Using whole-brain registration as before, we 30 found inhibitory projection neurons were intermingled with excitatory projection neurons, such that there was no 31 obvious anatomical separation between inhibitory and classic excitatory projections. Supporting this, both were 32 found in consistent proportions (~5% of labelled neurons) throughout each nucleus in BA, and across all three 33 anatomical axes (Figure S1A-C). Thus, in addition to the classically described excitatory projection from BA to 34 vH, there is a parallel inhibitory projection arising from GABAergic neurons from across the BA. 35

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We next investigated if these projections made functional connections onto vH pyramidal neurons. To recruit both excitatory and inhibitory projections, we expressed ChR2 under a pan-neuronal synapsin promoter (hsyn-ChR2) in the BA using an injection of adenoassociated virus (AAV) centred on posterior BMA and BLA. Two weeks later we prepared acute slices of vH from these animals and performed whole-cell recordings from

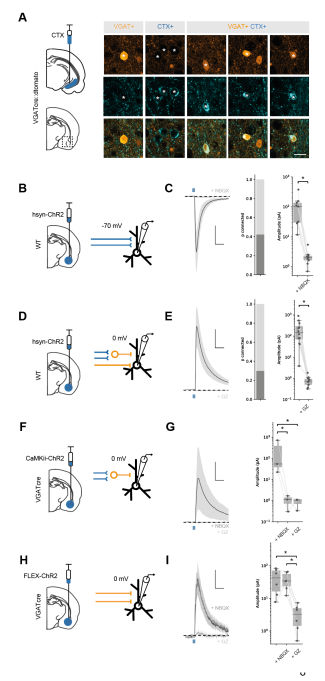


Figure 2 | BA input to vH is both excitatory and inhibitory.

A. CTX β injection in vH in a vGAT::cre::dtomato mouse line reveals inhibitory neurons (vGAT+), putative excitatory neurons that project to vH (CTX+) and inhibitory neurons that project to vH (vGAT+ CTX+). Scale bar = 20 um.

B. Schematic showing experimental setup. ChR2 was expressed using the pan-neuronal synapsin promoter using an AAV injection in BA. After allowing for expression, whole cell recordings were performed in voltage clamp at -70 mV in vH.

C. Brief pulses of blue light evoke excitatory currents that are blocked by the AMPA receptor antagonist NBQX. *Left,* Average current trace pre and post NBQX. *Middle,* proportion of recorded cells connected (with time-locked response to light). *Right,* Amplitude before and after NBQX. Note log scale. NBQX blocks excitatory currents evoked by BA input. Scale bar = 50 pA, 10 ms.

D, **E**. As **B**,**C** but for voltage clamp at 0 mV before and after the GABA receptor antagonist gabazine. Gabazine blocks inhibitory currents evoked by BA input. Scale bar = 50 pA, 10 ms.

F. Feedforward inhibition isolated using ChR2 expression under the CaMKii promoter.

G. Brief pulses of blue light evoked inhibitory currents at 0 mV that are blocked by the AMPA receptor antagonist NBQX. *Left,* Average current trace pre and post NBQX and GZ. *Right,* Amplitude before and after NBQX and GZ. Note log scale. NBQX blocks inhibitory currents evoked by CaMKii BA input, indicating it is solely feedforward. Scale bar = 50 pA, 10 ms.

H, **I**. As for **F**,**G** but direct inhibitory input isolated using ChR2 expression only in vGAT+ BA neurons. NBQX has no effect on direct inhibitory connection, while it is blocked by GZ, indicating it is a direct, long range inhibitory connection. Scale bar = 15 pA, 10 ms.

pyramidal neurons in the axon-rich CA1/ proximal subiculum border (Figure 2B). Recording in voltage clamp at -70 mV, we could isolate excitatory currents in response to blue light in ~40% of recorded neurons that were blocked by bath application of the AMPA receptor antagonist NBQX (Figure 2C). In the same neurons, we could also record inhibitory currents at 0 mV in ~30% of cells that were blocked by the GABA receptor antagonist gabazine (Figure 2E). Thus, BA

9 input makes excitatory and inhibitory connections with vH pyramidal neurons via AMPA and GABA receptors.

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Our retrograde tracing experiments (Figure 2A) suggested that in addition to classic feed-forward inhibition 11 (where excitatory axons make connections with local interneurons to disynaptically inhibit pyramidal neurons), 12 BA input also contained axons originating from inhibitory neurons, that would putatively make direct inhibitory 13 connections. To confirm this possibility, we first used a pharmacological approach (Figure S1D-F). Using mice 14 injected with hsyn-ChR2 in BA as above, we recorded inhibitory currents in vH pyramidal neurons at 0 mV. We 15 first removed feedforward inhibition with bath application of the AMPA receptor antagonist NBQX. Interestingly, 16 17 while inhibition was completely blocked in a subset of neurons (8/12), in the remaining population inhibitory currents persisted. This finding suggests that - consistent with our retrograde anatomy - a proportion of this 18 inhibitory input was due to a direct long range inhibitory projection from the BA. Consistent with this prediction, 19

the remaining current was blocked by bath application of gabazine, indicating that it was a GABA receptor
 mediated current.

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To test this more explicitly, we again used vGAT::cre mice where cre is expressed only in GABAergic neurons, 4 and expressed ChR2 in BA using either a CaMKii promoter - to confine expression to only putative excitatory 5 pyramidal neurons (Felix-Ortiz et al., 2013; Pi et al., 2020), or using a cre-dependent cassette to restrict ChR2 6 only to putative GABAergic neurons (Seo et al., 2016). Consistent with the presence of both excitatory and 7 inhibitory projections, CaMKii+ BA input evoked strong inhibitory currents at 0 mV (Figure 2G), but these currents 8 were blocked by bath application of NBQX, showing that the inhibitory currents were a result of solely 9 feedforward inhibition. In contrast, vGAT+ BA input also showed robust input at 0 mV (Figure 2I), but this 10 inhibitory current was insensitive to NBQX application, but blocked by gabazine, suggesting a direct inhibitory 11 connection. 12

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Together, these experiments define a novel, direct inhibitory projection from BA to vH. Thus, contrary to previous
 assumptions, BA provides two parallel projections to pyramidal neurons in vH, one excitatory, and one inhibitory.

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18 BA excitatory and inhibitory input selectively connects with unique vH output populations

The relatively sparse connectivity in our results above suggest that both excitatory and inhibitory BA input may connect with only a proportion of pyramidal neurons in vH. The CA1/proximal subiculum border of vH is composed of multiple populations of neurons organised as parallel projections (Figure S2, Gergues et al., 2020; Naber and Witter, 1998; Wee and MacAskill, 2020). Therefore, we hypothesised that this low connectivity may be an indication that BA input connects differentially with neurons that project to either NAc, PFC or back to BA.

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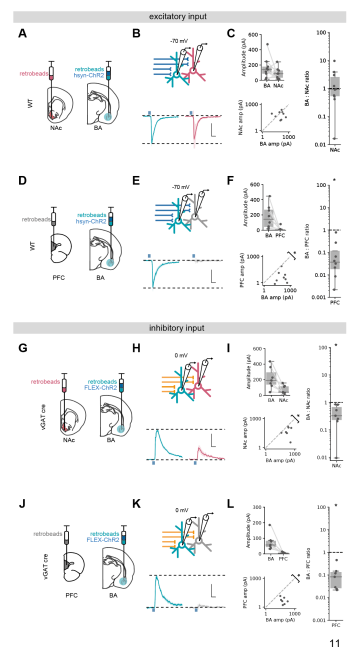
To investigate this possibility, we again injected ChR2 into BA, but also retrograde tracers into either BA and NAc, or BA and PFC. This allowed us, two weeks later, to prepare acute slices and obtain whole cell recordings from pairs of fluorescently-identified neurons in vH projecting to each downstream target. Paired recordings of neurons in the same slice and field of view allowed for a comparison of synaptic input while controlling for confounds such as injection volume, and the exact location in CA1/subiculum.

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We first compared excitatory input in voltage clamp at -70mV, as before with pan-neuronal expression of ChR2 using the synapsin promoter. Sequential paired recordings of vH^{BA} and vH^{NAc} neurons showed that light-evoked excitatory BA input was on average equivalent onto both populations (Figure 3A-C). In contrast, paired recordings of vH^{BA} and vH^{PFC} neurons revealed an almost complete lack of excitatory input onto vH^{PFC} neurons (Figure 3D-F).

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We next investigated long range inhibitory input using vGAT::cre mice and expressing cre dependent ChR2 in BA. Paired recordings of vH^{BA} and vH^{NAc} neurons showed a marked bias of inhibitory input to vH^{BA} neurons, with consistently smaller input onto neighbouring vH^{NAc} neurons (Figure 3G-I). Similarly to excitatory input, pairs of vH^{BA} and vH^{PFC} projecting neurons showed essentially no connectivity from BA to vH^{PFC} neurons (Figure 3J-L).



12 well placed to define their differential activity.

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14 BA excitatory and inhibitory input interact with local inhibitory circuitry in vH

We next wanted to understand how BA input may interact with the local vH circuit to define activity of the different 15 output populations. vH output populations have been shown to be strongly connected with local interneurons 16 to form both feedforward and feedback inhibitory circuitry, and this connectivity can vary on a cell-type specific 17 basis (S.-H. Lee et al., 2014; Soltesz and Losonczy, 2018). Thus we next wanted to ask 3 questions about the 18 layout of the vH circuit, and how it is influenced by BA input: (1) Does excitatory and inhibitory BA input connect 19 directly with local interneurons in vH? (2) Do pyramidal neurons from each projection population connect with 20 local interneurons to provide feedback inhibition? (3) Are there differences in how local interneurons connect 21 with pyramidal neurons from different projection populations? 22

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Figure 3 | Excitatory and inhibitory BA input differentially targets vH output populations.

A. Schematic of experiment vH^{NAc} and vH^{BA} neurons were labelled with retrobead injections, and ChR2 was expressed pan neuronally in BA.

B. Paired, fluorescently targeted recordings from neurons in each pathway and recording of light evoked currents. *Top*, recording setup. *Bottom*, average light evoked currents in vH^{BA} (*green*) and vH^{NAc} (*red*) neurons. Scale bar = 0.5 vH^{BA} response, 10 ms.

C. Summary of amplitude of light evoked BA input in pairs of vH^{NAc} and vH^{BA} neurons (*top*). When displayed as a scatter plot (*bottom*), or as the ratio of vH^{NAc}: vH^{BA} (*right*), the amplitudes cluster on the line of unity, indicating these population shave equal input. Note log axis.

D-F. As **A-C** but for pairs of vH^{BA} and vH^{PFC} neurons. Note when displayed as a scatter and a ratio of vH^{PFC}:vH^{BA} amplitudes are below the line of unity, indicating input preferentially innervates vH^{BA} neurons.

G-L. As **A-F** but for inhibitory input from BA isolated by expressing FLEX ChR2 in a vGAT::Cre line. Note when displayed as a scatter and a ratio, both vH^{PFC} and vH^{NAc} amplitudes are below the line of unity, indicating inhibitory input preferentially innervates vH^{BA} neurons in both cases. Scale bar = 0.5 vH^{BA} response, 10 ms.

Overall, these experiments suggest that excitatory input from BA equally targets vH neurons projecting to either NAc or BA, but not with those projecting to PFC. In contrast, inhibitory input from BA preferentially targets vH neurons projecting to BA, has a weak connection to those that project to NAc, and again avoids those projecting to PFC. Together, this shows that both excitatory and inhibitory BA input to vH have unique and distinct connectivity patterns with vH output circuitry, and suggests it is

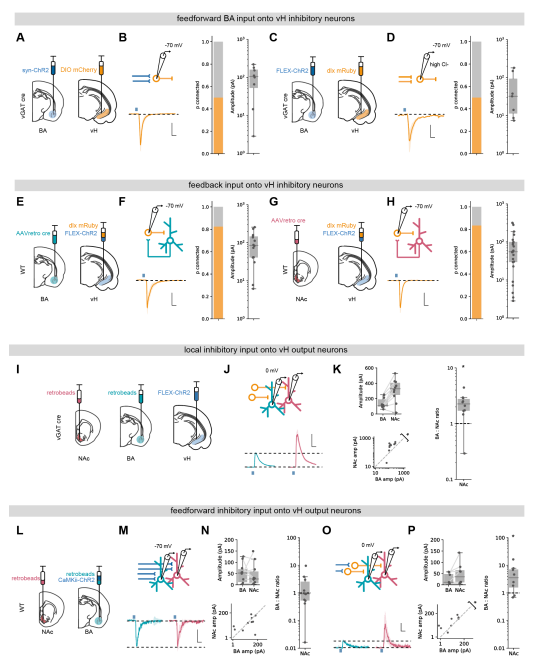


Figure 4 | BA input interacts with local inhibitory circuitry that is biased towards vH^{NAc} neurons.

A. schematic of experiment. ChR2 was expressed in BA, and DIO mCherry was expressed in vH in vGAT:cre mice to label local interneurons.

B. *Left*, Recording configuration to record excitatory connectivity at – 70 mV (top). Average light evoked current in interneurons in vH. Scale bar = 50 pA, 10 ms. *Right*, Summary of probability of connection (left) and amplitude of connected currents (right).

C,D. As **A**, **B** but for inhibitory input isolated using FLEX ChR2 expression in vGAT:cre mice as before. Note recordings were performed in high Cl-, so inward currents were measured at -70 mV.

E. Experimental setup for investigating feedback connectivity from vH^{BA} neurons. AAVretro was injected into BA, and FLEX ChR2 and dlx-mRuby into vH to allow recordings from dlx+ interneurons, and measurement of light evoked currents from vH^{BA} activation.

F. Left, Recording configuration to record excitatory connectivity at – 70 mV (top). Average light evoked current in dlx+ interneurons in vH. Right, Summary of probability of connection (left) and amplitude of connected currents (right).

G,H. As **E,F** but for feedback input from vH^{NAc} neurons.

I. Schematic of experiment, vH^{NAc} and vH^{BA} cells were labelled with injections of retrobeads, while ChR2 was expressed in vH interneurons using FLEX ChR2 in a vGAT::cre mouse.

J. Paired, fluorescently targeted recordings from neurons in each pathway at 0 mV and recording of light evoked currents. *Top*, recording setup. *Bottom,* average light evoked currents in vH^{BA} (*green*) and vH^{NAc} (*red*) neurons. Scale bar = 1 vH-BA response, 10 ms.

K. Summary of amplitude of light evoked BA input in pairs of vH^{NAc} and vH^{BA} neurons (*top*). When displayed as a scatter plot (*bottom*), or as the ratio of vH^{NAc}: vH^{BA} (*right*), the amplitudes cluster above the line of unity, indicating that local inhibition preferentially innervates vH^{NAc} neurons. Note log axis.

L-N. as I, J but for CaMKii input recorded at -70 mV. Note as in Figure 3 there is equal input onto both populations. Scale bar = 0.5 vH^{BA} response, 10 ms.

O,P. as in **M**, **N** but recording at 0 mV to isolate feedforward inhibition. Note that the amplitudes cluster above the line of unity, indicating that feedforward inhibition preferentially innervates vH^{NAc} neurons. Scale bar = 1 vH-BA response, 10 ms.

- 1 We first asked whether BA excitatory and inhibitory input targeted interneurons in vH. To do this we combined
- 2 ChR2 input mapping with an AAV injection in vH to express interneuron-specific fluorescent markers (Cho et al.,

2015; Dimidschstein et al., 2016). This allowed us to record from fluorescently identified interneurons in vH, and record light-evoked excitatory or inhibitory input from BA (Figure 4A-D). We found similar levels of both excitatory and inhibitory connectivity to input from BA onto local interneurons as we found with pyramidal neurons (in both cases ~50 % of recorded neurons were connected). Thus, both inhibitory and excitatory input from BA connect with local interneurons as well as pyramidal projection neurons in vH.

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We next wanted to investigate if vH^{BA} and vH^{NAc} neurons connected to local interneurons to form the basis of a 7 feedback inhibitory circuit (S.-H. Lee et al., 2014). To do this we injected a retrogradely transported AAV 8 (AAVretro) in either NAc and BA to express cre recombinase in NAc or BA projecting vH neurons respectively. 9 In the same surgery we injected a combination of cre-dependent ChR2 and the fluorescent reporter dlx-mRuby 10 into vH. This allowed us to obtain whole cell recordings from fluorescently identified vH interneurons, while 11 activating neighbouring projection neurons. Voltage clamp recordings at -70 mV showed robust responses from 12 both vH^{NAc} and vH^{BA} neurons onto local interneurons (~80 % of recorded neurons were connected in each 13 condition, Figure 4E-H), confirming previous studies suggesting strong feedback inhibition in vH (S.-H. Lee et 14 al., 2014). 15

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Finally, we asked if local interneurons differentially innervate vH^{BA} and vH^{NAc} neurons. We expressed ChR2 in 17 vGAT+ interneurons in vH using a vGAT::cre mouse line, and injected different coloured retrobeads into NAc 18 and BA. Two weeks later we then obtained paired, whole cell recordings from vH^{BA} and vH^{NAc} neurons, and 19 investigated light-evoked inhibitory synaptic input at 0mV. We found that local inhibitory connectivity was 20 markedly biased towards vH^{NAc} neurons (Figure 4I-K), where inhibitory connections onto vH^{NAc} neurons were 21 on average twice the strength of those onto neighbouring vH^{BA} neurons. Thus, activation of local interneurons 22 in vH, either via direct input from BA or via feedback from local pyramidal neurons, results in biased inhibition of 23 vH^{NAc} neurons, and has a much smaller effect of neighbouring vH^{BA} neurons. 24

This marked asymmetry of local inhibitory connectivity led us to predict that feedforward inhibition activated by excitatory BA input may also differentially impact the two output populations. We tested this using ChR2 expressed in BA under the control of the CaMKii promoter to limit expression to excitatory projections. As before, excitatory input in this experiment was equivalent in both vH^{BA} and vH^{NAc} neurons (Figure 4L-N). In contrast, and as predicted, feedforward inhibition recorded at 0 mV was markedly biased towards vH^{NAc} neurons (Figure 4O, P).

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Together these experiments show that local interneurons in vH make biased connections onto vH^{NAc} neurons. This biased innervation of interneurons towards vH^{NAc} neurons suggests greater influence of both feedforward inhibition from BA, but also feedback inhibition resulting from activation of local pyramidal neurons.

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38 A circuit model predicts a role for long range inhibition in promotion of vH^{NAc} activity

Our results so far suggest that the connectivity of both excitatory and inhibitory BA input into vH is very specific, and interacts with a number of interconnected elements in the local vH circuit. In order to investigate the overall

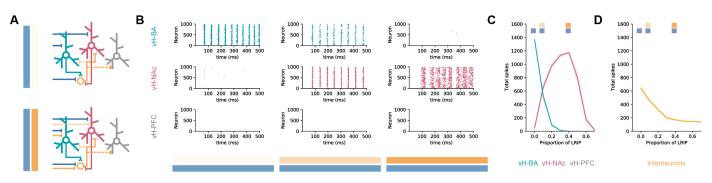


Figure 5 | Co-activation of inhibitory and excitatory input switches vH activity from vH^{BA} to vH^{NAc}

A. Schematic of integrate and fire model. Three populations of projection neurons (vH^{NAc}, *red*; vH^{BA}, *green*; vH^{PFC}, *grey*) and local interneurons (*orange*) are innervated by excitatory (blue, *top*) as well as inhibitory (orange, *bottom*) BA input. Connectivity is defined from results in previous figures.

B. Increasing the proportion of inhibitory relative to excitatory BA input has opposite effects on vH^{BA} and vH^{NAc} spiking. Each graph shows a raster of spiking for each neuron across a 500 ms period. Note high vH^{BA} spiking with no inhibitory input, and high vH^{NAc} spiking with high inhibitory input. vH^{PFC} neurons never fire as they are not innervated by BA, and only receive background input.

C. Summary of pyramidal neuron activity. With increasing inhibitory input, activity shifted from vH^{BA} to vH^{NAc} neurons. Markers indicate proportions plotted in **B.**

D. Long range inhibition reduces local interneuron firing, removing preferential feedback inhibition onto vH^{NAc} neurons, allowing them to fire.

influence of BA input in a more holistic way, we built a simple integrate-and-fire network (Stimberg et al., 2019),
 containing three separate projection populations in vH (to BA, NAc and PFC), local interneurons, excitatory and
 inhibitory input from BA, and background synaptic input from other structures. We then constrained the
 connectivity between these groups of neurons using the results of our circuit analysis (Figure 5A).

- We first looked at excitatory BA input alone, and found that this robustly activated vH^{BA} neurons in our model and had no effect on vH^{PFC} activity – consistent with the lack of connectivity to this population (see Figure 3). However, there was also a marked lack of vH^{NAc} activity, despite these neurons receiving equivalent excitatory synaptic input from BA. This was due to asymmetrical targeting by local inhibition (see Figure 4), and thus a combination of feedback and feedforward inhibition effectively silencing vH^{NAc} neurons, despite them receiving excitatory drive.
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We next incrementally added increasing proportions of long-range inhibitory input from BA to the model, such 13 that there was co-activation of both long-range inhibitory AND excitatory input. We found that increasing 14 inhibitory input resulted in a switch in the activity of the different populations (Figure 5B, C). While vHPFC neurons 15 remained silent, vH^{NAc} neuron activity increased as direct inhibition increased, and vH^{BA} neuron activity 16 decreased. This difference peaked around 40% long range inhibition, where vH^{BA} neurons were effectively silent, 17 and vH^{NAc} neurons were firing robustly. This was due to long range inhibition efficiently removing feedforward 18 and feedback inhibition onto vH^{NAc} neurons (Figure 5D) – both by direct inhibition of local interneuron activity, 19 but also by inhibiting vH^{BA} neurons that provide the bulk of feedback inhibitory drive. This effect was robust 20 across a wide range of feedforward and feedback connectivity (Figure S3). 21

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This circuit analysis suggests that specific connectivity of excitatory BA input into vH may not be the major determinant of vH^{BA} and vH^{NAc} neuron activity. In fact, it is the presence of direct inhibitory input from BA that

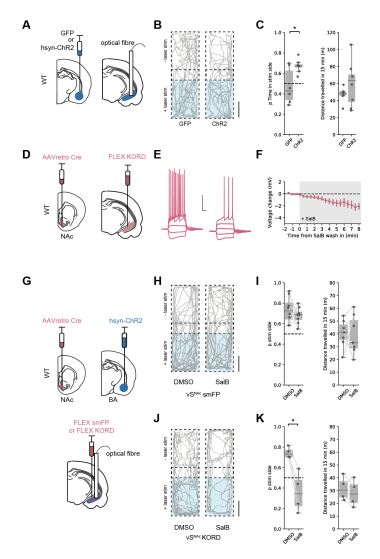


Figure 6 | BA input supports real time place preference dependent on vH^{NAc} neurons.

A. Schematic of experiment. GFP or pan-neuronal ChR2 were expressed in BA, and an optic fibre implanted in vH.

B. Real time place preference (RTPP) assay. One side of a chamber was paired with 20 Hz blue light stimulation. Example trajectories of GFP (left) and ChR2 (right) expressing animals over the 15 min RTPP session. Note increased occupancy of light-paired (stim) side in ChR2 animals. Scale bar = 15 cm.

C. Summary of RTPP. Left, Proportion of time spent on stim side (*left*) and total distance travelled (*right*) in GFP and ChR2 animals. Note consistent preference for stim side in ChR2 animals.

D. Strategy to express KORD in vH^{NAC} neurons.

E,F. Bath application of SalB (100 nm) hyperpolarises KORD expressing vH^{NAc} neurons, and reduces AP firing. See Figure S4 for full quantification. Scale bar = 30 mV, 100 ms.

G. Schematic of strategy to inhibit vH^{NAc} neurons during BA input driven RTPP.

H,I. As **B,C** but comparing the effect of either DMSO (vehicle) or SalB (KORD agonist) injections 15 mins before testing in control mice. Note consistent RTPP in both conditions indicating no effect of SalB in control mice.

 ${\bf J,K}.$ As ${\bf H,I},$ but in mice expressing KORD in ${\bf vH}^{\rm NAc}$ neurons. Note loss of RTPP in SalB injected mice compared to controls.

defines which projection population is active. With

no inhibition present, activity is confined to a reciprocal projection back to BA, however, when inhibition is present
 there is a switch to increased activity to NAc.

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6 **BA** input to vH can support RTPP via activation of vH^{NAc} neurons

A hallmark of activation of vH^{NAc} activation is the ability to promote real-time place preference (RTPP, Britt et al., 2012; LeGates et al., 2018). The results of our circuit modelling suggested that co-activation of BA inhibitory
and excitatory input to vH results in vH^{NAc} activation. We reasoned that BA input to vH may also support RTPP
via activation of vH^{NAc} neurons *in vivo*, and that this would depend on the coactivation of inhibitory as well as
excitatory BA projections.

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We tested if activation of both excitatory and inhibitory BA input supported RTPP by unilaterally injecting either GFP, or ChR2 under the pan-neuronal synapsin promoter into BA, and implanting optical fibres in vH (Figure 6A). We then carried out a RTPP test where one side of a rectangular arena was paired with 20 Hz blue light stimulation of BA terminals in vH. Consistent with our circuit analysis showing BA input activating vH^{NAc} neurons,

this stimulus supported RTPP in ChR2 expressing animals compared to GFP controls, with no change in the
 total distance moved during the session (Figure 6B,C).

3

From our circuit model, we predicted that this RTPP should be abolished by a reduction in the activity of vH^{NAc} 4 neurons. We next directly tested this using a combination of optogenetic RTPP to activate BA input, and the 5 Kappa Opioid Receptor Designer receptor exclusively activated by designer drugs (KORD) to reversibly inhibit 6 vH^{NAc} neurons (Vardy et al., 2015). We first tested the efficacy of KORDs expressed in vH^{NAc} neurons, and 7 confirmed that the KORD agonist salvinorin B (SalB) hyperpolarised vH^{NAc} neurons, and resulted in a decrease 8 in current-induced action potential firing (Figure 6D-F, Figure S4). We next combined this KORD mediated 9 inhibition with the optogenetic RTPP assay. We expressed pan-neuronal ChR2 in BA, KORDs in vH^{NAc} neurons, 10 and implanted an optical fibre unilaterally in vH (Figure 6G). We then carried out the RTPP assay 15 mins after 11 a subcutaneous injection of either SalB, or vehicle control (DMSO, Figure 6H-K). We found that after DMSO 12 injection, there was still robust RTPP in both control and KORD expressing mice. After SalB, control animals 13 again still had robust RTPP. However, after injection of SalB in KORD expressing animals, RTPP was abolished. 14 Together these experiments support our circuit model, where coactivation of both excitatory and inhibitory BA 15 input to vH supports real time place preference through the activation of vH^{NAC} neurons. 16

17 18

19 Excitatory BA input to vH supports RTPP only when vH^{BA} activity is inhibited.

In contrast to activation of both excitatory and inhibitory BA input into vH, another prediction from our circuit modelling is that excitatory BA input alone would not activate vH^{NAc} neurons, and thus would not support RTPP. We tested this prediction using ChR2 expressed under the CaMKii promoter to target only excitatory BA input to vH (see Figure 2). We injected either GFP or ChR2 under the CaMKii promoter in BA and implanted an optical fibre in vH, before carrying out an RTPP assay as before (Figure 7A). Consistent with the predictions from our circuit analysis, this assay showed that the light stimulus was unable to support RTPP in either GFP or ChR2 expressing animals (Figure 7B,C).

27

Our reasoning for this lack of RTPP was that excitatory BA input results in vH^{BA} neuron activity, and this recruits strong local feedback inhibition that preferentially reduces the activity of vH^{NAc} neurons (Figure 5) that are required to support RTPP (Figure 6). We therefore hypothesised that reducing vH^{BA} neuron activity (in effect mimicking the effect of the direct BA inhibitory projection) may increase vH^{NAc} activity and support RTPP from only excitatory BA input. This reasoning was supported our circuit model, where removing vH^{BA} activity increased the activity of vH^{NAc} neurons when no BA inhibitory input was present (Figure S5).

34

To test this hypothesis, we first ensured KORD-expressing vH^{BA} neuron excitability was inhibited by bath application of SalB (Figure 7D-F, Figure S4). Next, we injected ChR2 under the CaMKii promoter in BA to target only excitatory input into vH. In the same surgery we combined this with an injection of AAVretro cre in BA and cre-dependent KORD in vH to target vH^{BA} neurons, and implanted an optical fibre unilaterally in vH (Figure 7J). After allowing for expression, we then performed the RTPP assay 15 mins after injection of either SalB, or vehicle

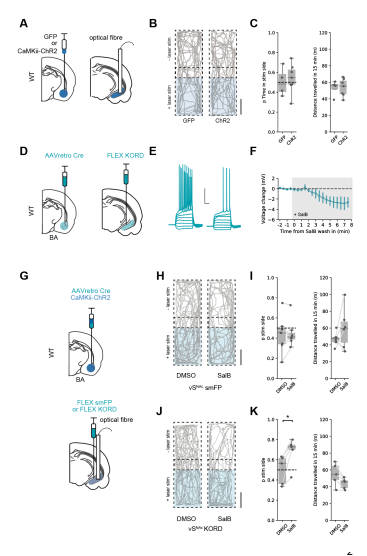


Figure 7 | Excitatory BA input supports real time place preference only after inhibition of vH^{BA} neurons.

A. Schematic of experiment. GFP or excitation-specific CaMKii ChR2 were expressed in BA, and an optic fibre implanted in vH.

B. Real time place preference (RTPP) assay. One side of a chamber was paired with 20 Hz blue light stimulation. Example trajectories of GFP (left) and ChR2 (right) expressing animals over the 15 min RTPP session. Note lack of preference for light-paired (stim) side in either group. Scale bar = 15 cm

C. Summary of RTPP. Left, Proportion of time spent on stim side (*left*) and total distance travelled (*right*) in GFP and ChR2 animals. Note lack of preference for stim side in either condition.

D. Strategy to express KORD in vH^{BA} neurons.

E,F. Bath application of SalB (100 nm) hyperpolarises KORD expressing vH^{BA} neurons, and reduces AP firing. See Figure S4 for full quantification. Scale bar = 30 mV, 100 ms.

G. Schematic of strategy to inhibit vH^BA neurons during BA input driven RTPP.

H,I. As **B,C** but comparing the effect of either DMSO (vehicle) or SalB (KORD agonist) injections 15 mins before testing in control mice. Note lack of RTPP in both conditions indicating no effect of SalB in control mice.

J,K. As **H,I**, but in mice expressing KORD in vH^{BA} neurons. Note induction of RTPP in SalB injected mice compared to controls.

control as before (Figure 7H-K). Consistent with our previous results, there was no RTPP in either

group after DMSO injections, or in control animals after SalB injection. However, after SalB injections in KORD
 expressing animals, light stimulation now supported RTPP.

5

This experiment supports our hypothesis that vH^{NAc} activity and hence RTPP is crucially dependent on the activity of both excitatory and inhibitory input from BA. Excitatory BA input to vH can only support RTPP if accompanied by inhibition of BA-projecting vH neurons, in effect mimicking the effect of BA inhibitory input on the circuit. Our model predicts that this reduction in vH^{BA} activity removes local feedback inhibition (Figure S5), and allows excitatory BA input to drive vH^{NAc} activity, which can support place preference.

1 DISCUSSION

2

In this study, we have defined a novel long-range inhibitory projection from BA to ventral hippocampus. We show that this novel projection exists in concert with a parallel excitatory projection, and that the presence of its inhibitory influence can dramatically shift vH output in response to BA activity. While excitation alone preferentially drives a reciprocal projection back to BA, coactivation of both excitatory and inhibitory input preferentially drives a separate projection to NAc, that can support place-value associations.

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We found that in addition to classically described excitatory input from BA to vH, there was also direct inhibitory 9 10 projection (Figures 1, 2). Excitatory input from BA to vH has been widely studied, and is distributed across a large range of subnuclei, ranging from the MEA to the BLA, and well as cortical amygdala (McDonald and Mott, 11 2016). Each of the distinct nuclei of the amygdala are thought to control various aspects of cue-dependent 12 learning and carry out unique roles during behaviour. Increasingly, function has been assigned to BA based on 13 anatomical location. For example, anterior basolateral, basomedial and central amygdala have unique 14 contributions to fear learning and extinction (Adhikari et al., 2015; Ciocchi et al., 2010; Kim et al., 2016; LeDoux, 15 2000), while more posterior and medial regions of BA are increasingly associated with reward-learning, value 16 calculations and prosocial behaviours (Chen et al., 2019; Kim et al., 2016; Lutas et al., 2019; Malvaez et al., 17 2019; Pi et al., 2020; Shemesh et al., 2016). However, the role within each of these nuclei is also diverse - with 18 interspersed neurons involved in encoding behaviour across a wide range of different situations (Beyeler et al., 19 2016; Felix-Ortiz et al., 2013; Felix-Ortiz and Tye, 2014; Gründemann et al., 2019; Kim et al., 2016; Namburi et 20 al., 2015). We found that the BA inhibitory projection arose from GABAergic neurons interspersed between 21 excitatory projection neurons throughout the entire extent of the BA (Figure S2). Thus, in addition to 22 systematically investigating the synaptic targeting and behavioural contribution of the input from different nuclei 23 separately, it will be important to assess the differential contribution of excitatory and inhibitory drive, most likely 24 through the use of intersectional genetic and anatomical approaches (Fenno et al., 2014; Kim et al., 2016). 25

Our results suggest that inhibitory input from BA to vH may be important for motivated behaviour, in particular 27 we show that coactivation of both excitatory and inhibitory projections from BA, and not excitation alone, is 28 29 essential for promoting place-value associations (Figure 6, 7). Long range inhibitory projections from classical excitatory projection areas have been increasingly identified as having a key role in shaping circuit output and 30 for defining motivated behaviour. For example, functional inhibitory projections from PFC to NAc (A. T. Lee et 31 al., 2014), and BA to PFC (Seo et al., 2016) have both been shown to modulate value-based and reward 32 behaviour, including the support of real time place preference and aversion. The hippocampus also receives 33 34 long range inhibitory input from numerous regions including entorhinal cortex (Basu et al., 2016; Melzer et al., 2012), septum (Schlesiger et al., 2021) and PFC (Malik et al., 2021). While these studies focussed on dorsal 35 36 hippocampal circuitry and a role for these projections in memory and navigation, due to the known dichotomy between dorsal and ventral hippocampal function (Fanselow and Dong, 2010; Strange et al., 2014), it would be 37 interesting to investigate the presence and function of such long range inhibitory projections into vH. In particular 38 whether a role in motivated behaviour and place preference was specific to BA input, or due to the dorso-ventral 39 location of this input in hippocampus. Interestingly, long range inhibition from entorhinal cortex, septum and PFC 40

all preferentially target interneurons and avoid pyramidal neurons (Basu et al., 2016; Malik et al., 2021; Melzer et al., 2012; Schlesiger et al., 2021). In contrast our data shows that BA long range inhibition connects with both interneurons and pyramidal neurons (Figure 3), similar to that seen in long range inhibitory projections from BA to PFC (Seo et al., 2016). This suggests that there may at least in part be interesting input specific connectivity across the different long range inhibitory inputs into hippocampus.

6

We investigated the synaptic and circuit basis by which BA input could promote such motivated behaviour. The 7 ventral hippocampus is increasingly viewed as being composed as a series of parallel output streams, where 8 pyramidal neurons in the CA1 / subiculum border are composed of multiple populations each projecting to a 9 distinct downstream region including the NAc, the PFC and the BA. Each projection population in vH underlies 10 a unique role during behaviour. In particular, vH^{NAc} neurons have been shown to be key for motivated behaviour, 11 and the association of reward with a particular place or time (Britt et al., 2012; Ciocchi et al., 2015; LeGates et 12 al., 2018; Okuyama et al., 2016; Trouche et al., 2019). We found that both excitatory and inhibitory input from 13 BA made specific connections onto each of these projection populations (Figure 3, 4, 5), such that the balance 14 of excitation and inhibition from BA into vH is well placed to determine their relative activity. Excitatory input 15 alone preferentially activated vH^{BA} neurons, while excitatory and inhibitory input together preferentially activated 16 vH^{NAc} neurons (Figure 5). Thus, BA input is well placed to define the activity of specific vH output pathways in 17 response to a particular environment, state or task. More specifically, the level of inhibitory input form BA can 18 control real time place preference by defining the activity of vH^{NAc} neurons (Figure 6,7). 19

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In addition to the role of BA and vH in value based and motivated behaviour, multiple studies have examined 21 the role of excitatory BLA input into vH in the generation of anxiety-like behaviour (Felix-Ortiz et al., 2013; Pi et 22 al., 2020). The vH has a key role in the generation of appropriate behaviour in anxiogenic environments (Gray 23 and McNaughton, 2003; Kjelstrup et al., 2002; McHugh et al., 2004). This is thought to be achieved both by 24 resolving approach avoidance conflict during decision making via vH^{PFC} projection neurons (Padilla-Coreano et 25 al., 2016; Sanchez-Bellot and MacAskill, 2020), but recently also via generation of a specific anxiogenic state 26 defined via projections to the lateral hypothalamus (LH, Jimenez et al., 2018). Neither excitatory or inhibitory BA 27 input connect with vHPFC neurons (Figure 3), suggesting innervation from other local or long-range afferent 28 regions may be key for this behavioural role (Sanchez-Bellot and MacAskill, 2020). However, BA input does 29 innervate vH^{LH} neurons (Gergues et al., 2020; Wee and MacAskill, 2020), and thus it is interesting to note the 30 possibility that the anxiogenic influence of excitatory, anterior BLA input (Felix-Ortiz et al., 2013; Pi et al., 2020) 31 may be via this distinct circuit. vH^{LH} neurons are present in more distal areas of ventral subiculum, with only a 32 minority present in the CA1/proximal subiculum border region considered in this study (Wee and MacAskill, 33 34 2020). However, how BA input interacts with distal subicular circuits that project to distinct downstream regions including hypothalamus and retrosplenial cortex (Cembrowski et al., 2018; Kim and Spruston, 2012); and how 35 36 inhibitory and excitatory input interact with this circuit is an interesting future direction.

Finally, our study focussed on the postsynaptic influence of BA inhibitory projections, and the molecular identity of these projection neurons remains unknown. Anatomical studies have suggested that BA inhibitory projections are preferentially observed in somatostatin (SOM)- and neuropeptide Y (NPY)-expressing neurons (McDonald et al., 2012; McDonald and Zaric, 2015), and almost completely absent in parvalbumin (PV)- and vasoactive

intestinal peptide (VIP)-expressing neurons. Thus, there is the potential for inhibitory input to be from both 1 specific nuclei in BA (Figure 1, Figure S2), but also different genetically defined populations of inhibitory neurons, 2 as is seen for excitatory amygdala projections (Kim et al., 2016). Similarly, in our study we did not differentiate 3 BA input onto different types of inhibitory interneuron in vH. There is enormous diversity of interneuron types 4 throughout the hippocampus (Group et al., 2008), each of which involved in distinct parts of the circuit calculation 5 - such as dendritic targeting SOM- and VIP-expressing neurons, perisomatic PV-expressing interneurons, and 6 cholecystokinin- (CCK) expressing interneurons. Inhibitory input from entorhinal cortex preferentially innervates 7 CCK interneurons (Basu et al., 2016), while input from PFC specifically innervates VIP interneurons (Malik et 8 al., 2021). Thus, how BA input differentially innervates these populations is an important and interesting future 9 question. 10

11

Overall we have defined a novel circuit that allows BA input to define the activity of parallel output pathways from vH to control motivated behaviour. The anatomical and functional specificity of this circuit provides an ideal substrate upon which to control reward and value-based learning and decision making, and helps to explain the multiple and varied roles attributed to this circuit.

1 METHODS

2

3 Animals

6 - 10 week old (adult) male and female C57 / bl6J mice provided by Charles River were used except where
noted. To target inhibitory neurons we used the Slc32a1(VGAT)-IRES-Cre (#016962) knock-in line obtained
from Jackson laboratories and bred in-house. Mice were housed in cages of 2 - 4 and kept in a humidity- and
temperature-controlled environment under a 12 h light/dark cycle (lights on 7 am to 7 pm) with ad-libitum access
to food and water. All experiments were approved by the U.K. Home Office as defined by the Animals (Scientific
Procedures) Act, and University College London ethical guidelines.

10

11 Stereotaxic surgery

12

13 Retrograde tracers:

- 14 Red and Green fluorescent retrobeads (Lumafluor, Inc.) for electrophysiological recordings.
- 15 Cholera toxin subunit B ($CTX\beta$) tagged with Alexa 555, 488 or 647 (Molecular Probes) for histology experiments.
- 16

18

17 Viruses:

19	AAV2/1-CaMKII-GFP	(a gift from Edward Boyden; Addgene #64545)
20	AAV2retro-CAG-Cre	(UNC vector core)
21	AAV2/1-EF1a-FLEX-hChR2(H134R)-EYFP	(a gift from Karl Deisseroth; Addgene #20298-AAV1)
22	AAV2/1-hSyn-hChR2(H134R)-EYFP	(a gift from Karl Deisseroth; Addgene #26973-AAV1)
23	AAV2/1-CaMKII-hChR2(H134R)-EYFP	(a gift from Karl Deisseroth; Addgene #26969-AAV1)
24	pAAV2/8-hSyn-dF-HA-KORD-IRES-mCitrine	(a gift from Bryan Roth; Addgene #6541-AAV8)
25	AAV2/1.CAG.FLEX.Ruby2sm-Flag.WPRE	(a gift from Loren Looger; Addgene #98928-AAV1)
26	AAV2/9-mDlx-NLS-mRuby2	(a gift from Viviana Gradinaru; Addgene #99130-AAV1)
27	pAAV2/1-Ef1a-fDIO mCherry	(a gift from Karl Deisseroth ; Addgene 114471-AAV1)

28

29

30 Surgery:

Stereotaxic injections were performed on 7 - 10 week old mice anaesthetized with isofluorane (4 % induction, 1 31 - 2 % maintenance) and injections carried out as previously described (Sanchez-Bellot and MacAskill, 2020; 32 Wee and MacAskill, 2020). Briefly, the skull was exposed with a single incision, and small holes drilled in the 33 skull directly above the injection site. Injections are carried out using long-shaft borosilicate glass pipettes with 34 a tip diameter of ~ 10 - 50 µm. Pipettes were back-filled with mineral oil and front-filled with ~ 0.8 µl of the 35 substance to be injected. A total volume of 250 - 300 nl of each virus was injected at each location in ~ 14 or 36 28 nl increments every 30 s. If two or more substances were injected in the same region they were mixed prior 37 to injection. The pipette was left in place for an additional 10 - 15 min to minimize diffusion and then slowly 38 removed. If optic fibres were also implanted, these were inserted immediately after virus injection, secured with 39

1 - 2 skull screws and cemented in place with C&B superbond. Injection coordinates were as follows (mm relative to bregma):

3

4	infralimbic PFC:	ML: ± 0.4,	RC: + 2.3,	and DV: - 2.4
5	nucleus accumbens:	ML: ± 0.9,	RC: + 1.1,	and DV: - 4.6
6	basal amygdala:	ML: ± 3.4,	RC: - 1.7,	and DV: - 5.8
7	ventral hippocampus:	ML: ± 3.2,	RC: - 3.7,	and DV: - 4.7

8

After injection, the wound was sutured and sealed, and mice recovered for ~30 min on a heat pad before they were returned to their home cage. Animals received carprofen in their drinking water (0.05 mg / ml) for 48 hrs post-surgery as well as subcutaneously during surgery (0.5 mg / kg). Expression occurred in the injected brain region for ~2 weeks for WT animals and ~4 weeks for vGAT animals until behavioural testing, preparation of acute slices for physiology experiments, or fixation for histology. The locations of injection sites were verified for each experiment.

15

16 Anatomy

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18 Histology:

Mice were perfused with 4% PFA (wt / vol) in PBS, pH 7.4, and the brains dissected and postfixed overnight at 19 4°C as previously described (MacAskill et al., 2014; Sanchez-Bellot and MacAskill, 2020; Wee and MacAskill, 20 2020). 70 µm thick slices were cut using a vibratome (Campden Instruments) in either the transverse, coronal 21 or sagittal planes as described in the figure legends. Slices were mounted on Superfrost glass slides with 22 ProLong Gold or ProLong Glass (for visualization of GFP) antifade mounting medium (Molecular Probes). 23 NucBlue was included to label gross anatomy. Imaging was carried out with a Zeiss Axio Scan Z1, using 24 standard filter sets for excitation/emission at 365-445/50 nm, 470/40-525/50 nm, 545/25-605/70 nm and 640/30-25 690/50 nm. Raw images were analyzed with FIJI. 26

27

28 Whole Brain Registration:

29 Cell counting of cholera-labelled inputs was conducted using WholeBrain (Fürth et al., 2018; Wee and MacAskill, 2020). After acquiring the imaged sections and exporting them as 16-bit depth image files, images were 30 manually assigned a bregma coordinate (AP -6.0 to 0.0 mm) and processed using WholeBrain (Fürth et al., 31 2018) and custom cell counting routines written in R (Wee and MacAskill, 2020). The workflow comprised of (1) 32 segmentation of cells and brain section. (2) registration of the cells to the Allen Brain Atlas (ABA) and (3) analysis 33 of anatomically registered cells. As tissue section damage impairs the automatic registration implemented on 34 the WholeBrain platform, sections with poor registration were manually registered to the atlas plate using 35 36 corresponding points to clear anatomical landmarks. Once all cells had been registered, the cell counts were further manually filtered from the dataset to remove false-positive cells (e.g. debris). 37

38

Each cell registered to a brain region was classified as belonging to an anatomically defined region as defined
 by the ABA brain structure ontology. Information on the ABA hierarchical ontology was scraped from the ABA

API (link: http://api.brain-map.org/api/v2/structure graph download/1.json) using custom Python routines. For 1 guantification of input fractions, cells residing in different layers within the same structure, e.g. COAa1, COAa2 2 etc, were agglomerated across layers and subdivisions and counted as residing in one single region (e.g. 3 COAa). Structures included as part of BA were: 'BLAa', 'BLAv', 'BLAp', 'BMAa', 'BMAp', 'LA', 'COAa', 'COApl', 4 'COApm', 'MEAa', 'MEAav', 'MEApd', 'MEApv', 'CEAc', 'CEAm', 'CEAI', 'PAA', 'PA'. For colocalization of VGAT+ 5 and CTX_β labelled neurons, images acquired as above were manually annotated with single and dual labelled 6 neurons using Napari (napari contributors, 2019, doi:10.5281/zenodo.3555620). Whole brain distributions were 7 visualised using the Brainrender package for python (Claudi et al., 2020). 8

9

10 Electrophysiology

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12 Slice preparation:

Hippocampal recordings were studied in acute transverse slices. Mice were anaesthetized with a lethal dose of 13 ketamine and xylazine, and perfused intracardially with ice-cold external solution containing (in mM): 190 14 sucrose, 25 glucose, 10 NaCl, 25 NaHCO₃, 1.2 NaH₂PO₄, 2.5 KCl, 1 Na⁺ ascorbate, 2 Na⁺ pyruvate, 7 MgCl₂ 15 and 0.5 CaCl₂, bubbled with 95% O₂ and 5% CO₂. Slices (400 µm thick) were cut in this solution and then 16 17 transferred to artificial cerebrospinal fluid (aCSF) containing (in mM): 125 NaCl, 22.5 glucose, 25 NaHCO₃, 1.25 NaH₂PO₄, 2.5 KCl, 1 Na⁺ ascorbate, 3 Na⁺ pyruvate, 1 MgCl₂ and 2 CaCl₂, bubbled with 95% O₂ and 5% CO₂. 18 After 30 min at 35 °C, slices were stored for 30 min at 24 °C. All experiments were conducted at room 19 temperature (22-24 °C). All chemicals were from Sigma, Hello Bio or Tocris. 20

21

22 Whole-cell electrophysiology:

Whole-cell recordings were made from hippocampal pyramidal neurons retrogradely labelled with retrobeads 23 which were identified by their fluorescent cell bodies and targeted with Dodt contrast microscopy, as previously 24 described (MacAskill et al., 2014; Sanchez-Bellot and MacAskill, 2020; Wee and MacAskill, 2020). For 25 sequential paired recordings, neurons were identified within a single field of view at the same depth into the 26 slice. The recording order was counterbalanced to avoid any potential complications that could be associated 27 with rundown. For current clamp recordings, borosilicate recording pipettes $(4 - 6 M\Omega)$ were filled with (in mM): 28 135 K-gluconate, 10 HEPES, 7 KCl, 10 Na-phosphocreatine, 10 EGTA, 4 MgATP, 0.4 NaGTP. For voltage 29 clamp experiments, three internals were used, First, in Figure 2, 3 and 4I-P, a Cs-gluconate based internal was 30 used containing (in mM): 135 Gluconic acid, 10 HEPES, 7 KCI, 10 Na-phosphocreatine, 4 MgATP, 0.4 NaGTP, 31 10 TEA and 2 QX-314. Excitatory and inhibitory currents were electrically isolated by setting the holding potential 32 at -70 mV (excitation) and 0 mV (inhibition) and recording in the presence of APV. Experiments in Fig 4A,B,E-H 33 were carried out using current clamp internal in APV in order to carry out post stimulation analysis of intrinsic 34 properties of recorded interneurons. Finally, to record inhibitory currents at -70 mV in Fig.4C,D we used a high 35 chloride internal (in mM): 135 CsCl, 10 HEPES, 7 KCl, 10 Na-phosphocreatine, 10 EGTA, 4 MgATP, 0.3 NaGTP, 36 10 TEA and 2 QX-314. Recordings were made using a Multiclamp 700B amplifier, with electrical signals filtered 37 at 4 kHz and sampled at 10 kHz. 38

Presynaptic glutamate release was triggered by illuminating ChR2 in the presynaptic terminals of long-range
 inputs into the slice, as previously described (Sanchez-Bellot and MacAskill, 2020; Wee and MacAskill, 2020).
 Wide-field illumination was achieved via a 40 x objective with brief pulses of blue light from an LED centered at
 470 nm (CoolLED pE-4000 / Thorlabs M470L4-C1, with appropriate excitation-emission filters). Light intensity
 was measured as 4 –7 mW at the back aperture of the objective and was constant between all cell pairs.

6

7 Electrophysiology data acquisition and analysis:

Electrophysiology data were acquired using National Instruments boards and WinWCP (University of Strathclyde). Optical stimulation was via wide field irradiance with 473 nm LED light (CoolLED) as described above. Data was analysed using custom routines written in Python 3.6, imported using the neo package in python (Garcia et al., 2014). For connectivity analysis, a cell was considered connected if the average of lightinduced response was greater than 2 standard deviations above baseline. Amplitudes of responses were calculated as the average of a 2ms window around the peak of the response. Current step data (Figure S2) was analysed using routines based around the eFEL package in python (Blue Brain Project).

15

16 Integrate and fire model

An integrate and fire model was constructed using the Brian2 package in python (Stimberg et al., 2019). 1000 17 vH-BA, vH-NAc and vH-PFC neurons were modelled interspersed with 80 interneurons (S.-H. Lee et al., 2014). 18 Neurons were set to have a leak conductance, resting potential, spike threshold and membrane capacitance 19 based on the literature and our current clamp recordings (Figure S2): leak conductance 5.5 nS; resting potential 20 -70 mV, spiking threshold -35 mV, membrane capacitance 200 pF. Connectivity of the local vH circuit was based 21 on our electrophysiology recordings. AMPA receptor connections were 1 nS and were modelled with a tau of 5 22 ms. GABA receptor mediated connections were 3 nS and modelled with a tau of 10 ms. Feedback connectivity 23 from each pyramidal neuron population was connected at a probability of 0.1. The probability of connection of 24 local interneurons to pyramidal neurons was based on Figure 4, and was 0.8 for vH-NAc neurons, and 0.4 for 25 vH-BA and vH-PFC neurons, each with a 3 nS GABA conductance. To simulate excitatory BA input, neurons 26 were supplied with 50,000 BA inputs timed as a Poisson distribution with an average rate of 10 Hz. Each neuron 27 was connected to this input with a probability of 0.1, where the strength of the synaptic connection was randomly 28 29 drawn from a normal distribution defined by our electrophysiology experiments in Figure 3 (vH-BA 0.3 nS +/-0.2, vH-NAc 0.3 nS +/- 0.2, vH-PFC 0.03 nS +/- 0.2, interneurons 0.3 nS +/- 0.2). To simulate BA inhibitory input, 30 neurons were again supplied with 50,000 BA inputs timed as a Poisson distribution with an average rate of 10 31 Hz, but the connection probability was calculated as a proportion of excitatory input, and varied across runs. As 32 before the strength of each synaptic connection was randomly drawn from a normal distribution defined by our 33 34 electrophysiology experiments in Figure 4 (vH-BA 0.3 nS +/- 0.2, vH-NAc 0.08 nS +/- 0.2, vH-PFC 0.03 nS +/-0.2, interneurons 0.3 nS +/- 0.2). Each simulation was run 5 times at each level of inhibitory connection strength, 35 36 with the length of simulation 500 ms for each run. To investigate the influence of feedforward and feedback connection probability (Figure S4) we systematically altered these parameters for each run. Model output was 37 analysed as total spikes produced by each neuronal population over the course of 500 ms. 38

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- 40

1 Behaviour

After sufficient time for surgical recovery and viral expression (>4 weeks), mice underwent multiple rounds of habituation. Mice were habituated to the behavioural testing area in their home cage for 30 min prior to testing each day. Mice were habituated to handling for at least 3 days, followed by 1-2 days of habituation to the optical tether in their home cage for 10 min.

6

7 Real time place preference:

Axon terminals were labelled as described above, and a 200 µm optical fibre was implanted unilaterally 100 um 8 above the stimulation area (vH). After habituation (above), behaviour was assessed using a real time place 9 preference (RTPP) task. On Day 1 mice were exposed to the 3 chamber arena (24 cm x 16 cm x 30 cm) for 15 10 min without stimulation to allow habituation, and also to ensure no large side bias was present. The testing 11 chamber was made out of black acrylic, was symmetrical and had no odour, visual or tactile cues to distinguish 12 either side of the arena. The arena was thoroughly wiped down with 70% ethanol between each trial. Mice were 13 excluded if they spend more than 80% of their time in one side of the chamber during this habitation session. 14 On day 2, 20 Hz light stimulation was delivered via a 473 nm laser, coupled to a patch cord (7-10 mW at the 15 end of the patch cord) to activate ChR2 positive terminals. Real-time light delivery was based on the location of 16 the mouse in the RTPP apparatus, where light stimulation occurred only when the mouse was in the light-paired 17 side of the arena. The paired side was chosen randomly for each mouse and each session, thus in combination 18 with the lack of explicit cues in the chamber, this assay represents acute place preference and not learned 19 preference over sessions. Time spent in the light paired, and control side of the arena over the course of the 15 20 minute session was scored for each mouse using automated tracking analysis (Bonsai). For experiments 21 involving pharmacogenetics (Fig 6, 7), mice first underwent habituation and laser only trials as before, and data 22 from control animals were used to replicate the original RTPP cohort (Figure 6A-C and 7A-C). Next, mice were 23 given 1 - 2 daily s.c. injections of DMSO for habituation, before undergoing two further days of testing - first with 24 DMSO as a control and with 10 mg/kg SalB the next day to avoid any spill over effects of the SalB injection. All 25 injections were given 15 minutes prior to RTPP session. Control mice for optogenetics expressed GFP in BA. 26 Control mice for KORD experiments consisted of a mixture of mice expressing smFP in vH^{NAc} neurons and mice 27 lacking expression in vH, all of which received an injection of both DMSO and SalB. No differences were seen 28 29 across the two conditions and so data were pooled.

30

31 Statistics

Summary data are reported throughout the figures either as boxplots, which show the median, 75th and 95th 32 percentile as bar, box and whiskers respectively, or as line plots showing mean +/- s.e.m. Example physiology 33 34 and imaging traces are represented as the median +/- s.e.m across experiments. Data were assessed using statistical tests described in the supplementary statistics summary, utilising the Pingouin statistical package for 35 python (Vallat, 2018). Significance was defined as P < 0.05, all tests were two sided. No statistical test was run 36 to determine sample size a priori. The sample sizes we chose are similar to those used in previous publications. 37 Animals were randomly assigned to a virus cohort (e.g. ChR2 versus GFP), and where possible the 38 experimenter was blinded to each mouse's virus assignment when the experiment was performed. This was 39 sometimes not possible due to e.g. the presence of the injection site in the recorded slice. 40

1 ACKNOWLEDGEMENTS

We thank members of the MacAskill laboratory for helpful comments on the manuscript. A.F.M. was supported by a Sir Henry Dale Fellowship jointly funded by the Wellcome Trust and the Royal Society (grant number 109360/Z/15/Z) and by a UCL Excellence Fellowship. R.A. was supported by a King Fahad Medical City Studentship. R.W.S.W. was supported by a UCL Graduate Research Scholarship and a UCL Overseas

6 Research Scholarship. K.M. was supported by the Wellcome Trust 4-year PhD in Neuroscience at UCL

- 7 (grant number 215165/Z/18/Z).
- 8 9

10 AUTHOR CONTRIBUTIONS

Conceptualization, R.A. and A.F.M.; Methodology, R.A., R.W.S.W and A.F.M.; Investigation, R.A.,
R.W.S.W., K.M., A.R., D.R. and A.F.M.; Formal Analysis, R.A., R.W.S.W., A.R., and A.F.M.; Writing –
Original Draft, A.F.M.; Writing – Review & Editing, R.A. and A.F.M.; Funding Acquisition, R.A. and A.F.M.;
Supervision, A.F.M.

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17 DECLARATION OF INTERESTS

- 18 The authors declare no competing interests.
- 19

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21 DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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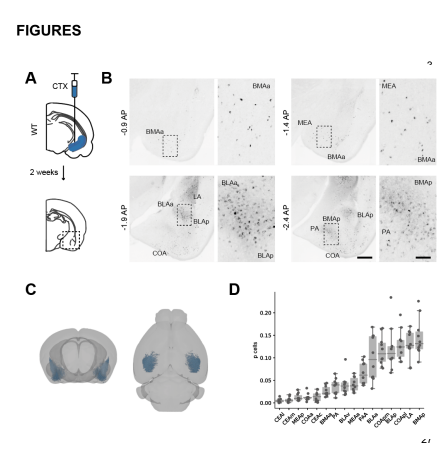


Figure 1 | Distribution of BA input to vH.

A. Schematic of experiment. $CTX\beta$ was injected into vH, 2 weeks later coronal slices of BA were examined for retrogradely labelled neurons.

B. Example slices showing widespread labelling throughout numerous BA nuclei. Scale bar = 500 um, 100 um (zoom).

C. Whole brain distribution of labelled BA neurons.

D. Summary showing proportion of labelled BA cells in each nuclei.

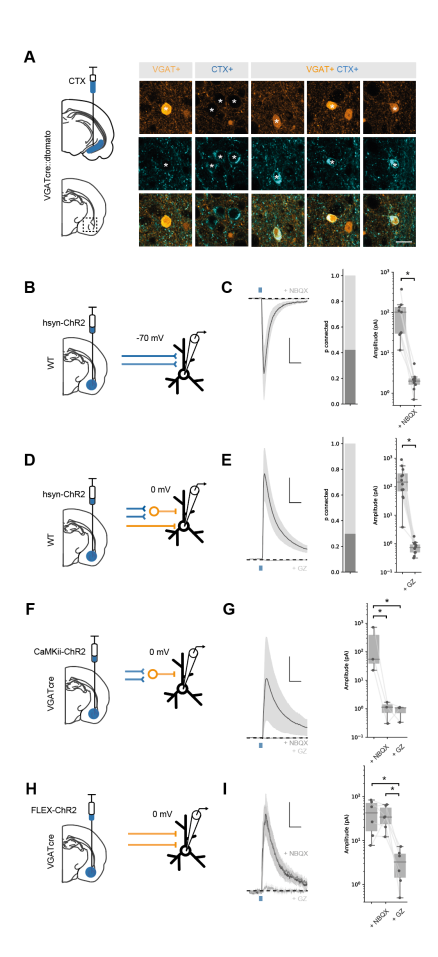


Figure 2 | BA input to vH is both excitatory and inhibitory.

A. CTX β injection in vH in a vGAT::cre::dtomato mouse line reveals inhibitory neurons (vGAT+), putative excitatory neurons that project to vH (CTX+) and inhibitory neurons that project to vH (vGAT+ CTX+). Scale bar = 20 um.

B. Schematic showing experimental setup. ChR2 was expressed using the pan-neuronal synapsin promoter using an AAV injection in BA. After allowing for expression, whole cell recordings were performed in voltage clamp at – 70 mV in vH.

C. Brief pulses of blue light evoke excitatory currents that are blocked by the AMPA receptor antagonist NBQX. *Left*, Average current trace pre and post NBQX. *Middle*, proportion of recorded cells connected (with time-locked response to light). *Right*, Amplitude before and after NBQX. Note log scale. NBQX blocks excitatory currents evoked by BA input. Scale bar = 50 pA, 10 ms.

D, **E**. As **B**,**C** but for voltage clamp at 0 mV before and after the GABA receptor antagonist gabazine. Gabazine blocks inhibitory currents evoked by BA input. Scale bar = 50 pA, 10 ms.

F. Feedforward inhibition isolated using ChR2 expression under the CaMKii promoter.

G. Brief pulses of blue light evoked inhibitory currents at 0 mV that are blocked by the AMPA receptor antagonist NBQX. *Left,* Average current trace pre and post NBQX and GZ. *Right,* Amplitude before and after NBQX and GZ. Note log scale. NBQX blocks inhibitory currents evoked by CaMKii BA input, indicating it is solely feedforward. Scale bar = 50 pA, 10 ms.

H, I. As for F,G but direct inhibitory input isolated using ChR2 expression only in vGAT+ BA neurons. NBQX has no effect on direct inhibitory connection, while it is blocked by GZ, indicating it is a direct, long range inhibitory connection. Scale bar = 15 pA, 10 ms.

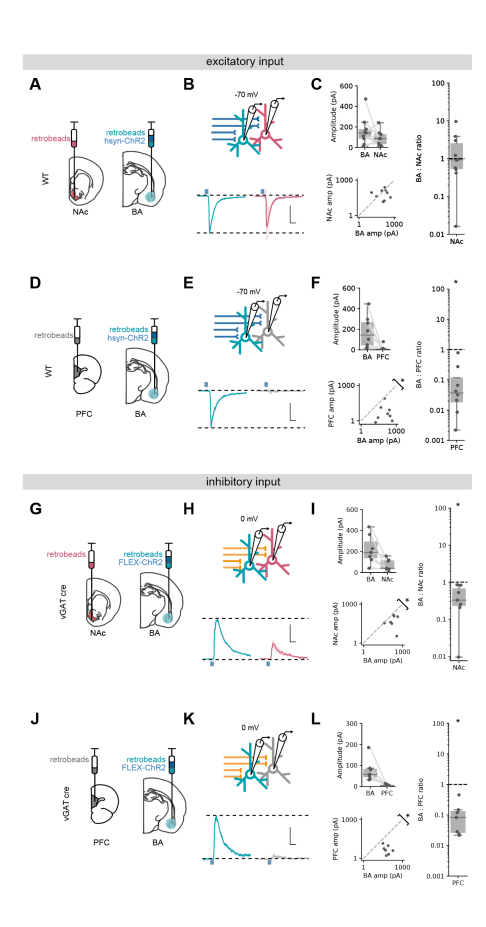


Figure 3 | Excitatory and inhibitory BA input differentially targets vH output populations.

A. Schematic of experiment vH^{NAc} and vH^{BA} neurons were labelled with retrobead injections, and ChR2 was expressed pan neuronally in BA.

B. Paired, fluorescently targeted recordings from neurons in each pathway and recording of light evoked currents. *Top*, recording setup. *Bottom,* average light evoked currents in vH^{BA} (*green*) and vH^{NAc} (*red*) neurons. Scale bar = 0.5 vH^{BA} response, 10 ms.

C. Summary of amplitude of light evoked BA input in pairs of vH^{NAc} and vH^{BA} neurons (*top*). When displayed as a scatter plot (*bottom*), or as the ratio of vH^{NAc}: vH^{BA} (*right*), the amplitudes cluster on the line of unity, indicating these population shave equal input. Note log axis.

D-F. As **A-C** but for pairs of vH^{BA} and vH^{PFC} neurons. Note when displayed as a scatter and a ratio of vH^{PFC}:vH^{BA} amplitudes are below the line of unity, indicating input preferentially innervates vH^{BA} neurons.

G-L. As **A-F** but for inhibitory input from BA isolated by expressing FLEX ChR2 in a vGAT::Cre line. Note when displayed as a scatter and a ratio, both vH^{PFC} and vH^{NAc} amplitudes are below the line of unity, indicating inhibitory input preferentially innervates vH^{BA} neurons in both cases. Scale bar = 0.5 vH^{BA} response, 10 ms.

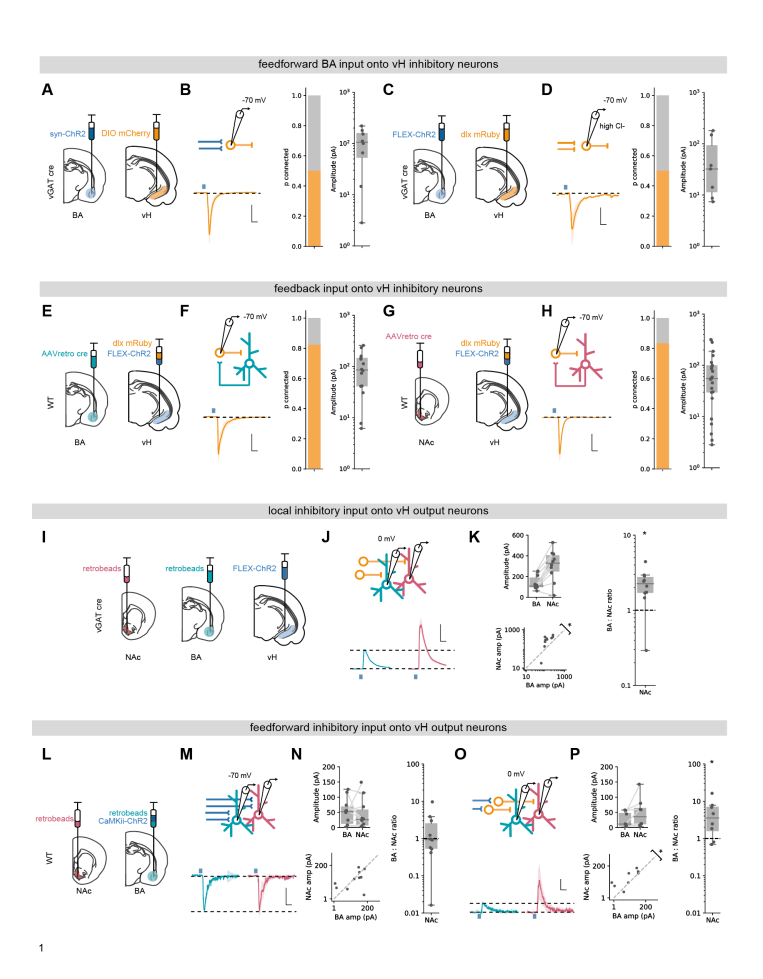


Figure 4 | BA input interacts with local inhibitory circuitry that is biased towards vH^{NAc} neurons.

A. schematic of experiment. ChR2 was expressed in BA, and DIO mCherry was expressed in vH in vGAT:cre mice to label local interneurons.

B. *Left*, Recording configuration to record excitatory connectivity at – 70 mV (top). Average light evoked current in interneurons in vH. Scale bar = 50 pA, 10 ms. *Right*, Summary of probability of connection (left) and amplitude of connected currents (right).

C,D. As **A**, **B** but for inhibitory input isolated using FLEX ChR2 expression in vGAT:cre mice as before. Note recordings were performed in high Cl-, so inward currents were measured at -70 mV.

E. Experimental setup for investigating feedback connectivity from vH^{BA} neurons. AAVretro was injected into BA, and FLEX ChR2 and dlx-mRuby into vH to allow recordings from dlx+ interneurons, and measurement of light evoked currents from vH^{BA} activation.

F. Left, Recording configuration to record excitatory connectivity at – 70 mV (*top*). Average light evoked current in dlx+ interneurons in vH. Right, Summary of probability of connection (*left*) and amplitude of connected currents (*right*).

G,H. As **E,F** but for feedback input from vH^{NAc} neurons.

I. Schematic of experiment, vH^{NAc} and vH^{BA} cells were labelled with injections of retrobeads, while ChR2 was expressed in vH interneurons using FLEX ChR2 in a vGAT::cre mouse.

J. Paired, fluorescently targeted recordings from neurons in each pathway at 0 mV and recording of light evoked currents. *Top*, recording setup. *Bottom*, average light evoked currents in vH^{BA} (*green*) and vH^{NAc} (*red*) neurons. Scale bar = 1 vH-BA response, 10 ms.

K. Summary of amplitude of light evoked BA input in pairs of vH^{NAc} and vH^{BA} neurons (*top*). When displayed as a scatter plot (*bottom*), or as the ratio of vH^{NAc}: vH^{BA} (*right*), the amplitudes cluster above the line of unity, indicating that local inhibition preferentially innervates vH^{NAc} neurons. Note log axis.

L-N. as I, J but for CaMKii input recorded at -70 mV. Note as in Figure 3 there is equal input onto both populations. Scale bar = 0.5 vH^{BA} response, 10 ms.

O,P. as in **M**, **N** but recording at 0 mV to isolate feedforward inhibition. Note that the amplitudes cluster above the line of unity, indicating that feedforward inhibition preferentially innervates vH^{NAc} neurons. Scale bar = 1 vH-BA response, 10 ms.

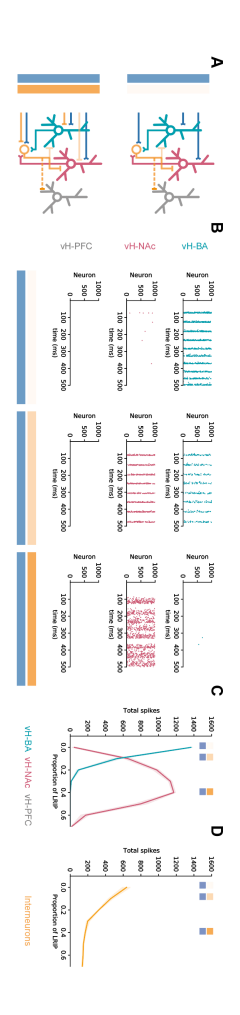


Figure 5 | Co-activation of inhibitory and excitatory input switches vH activity from vH^{BA} to vH^{NAC}

A. Schematic of integrate and fire model. Three populations of projection neurons (vH^{NAc}, *red*; vH^{BA}, *green*; vH^{PFC}, *grey*) and local interneurons (*orange*) are innervated by excitatory (blue, *top*) as well as inhibitory (orange, *bottom*) BA input. Connectivity is defined from results in previous figures.

B. Increasing the proportion of inhibitory relative to excitatory BA input has opposite effects on vH^{BA} and vH^{NAc} spiking. Each graph shows a raster of spiking for each neuron across a 500 ms period. Note high vH^{BA} spiking with no inhibitory input, and high vH^{NAc} spiking with high inhibitory input. vH^{PFC} neurons never fire as they are not innervated by BA, and only receive background input.

C. Summary of pyramidal neuron activity. With increasing inhibitory input, activity shifted from vH^{BA} to vH^{NAC} neurons. Markers indicate proportions plotted in **B**.

D. Long range inhibition reduces local interneuron firing, removing preferential feedback inhibition onto vH^{NAc} neurons, allowing them to fire.

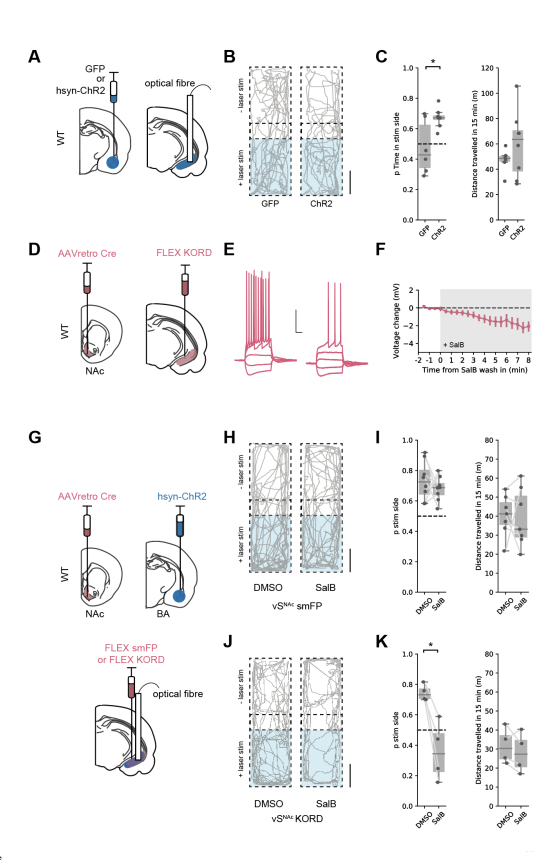


Figure 6 | BA input supports real time place preference dependent on vH^{NAc} neurons.

A. Schematic of experiment. GFP or pan-neuronal ChR2 were expressed in BA, and an optic fibre implanted in vH.

B. Real time place preference (RTPP) assay. One side of a chamber was paired with 20 Hz blue light stimulation. Example trajectories of GFP (left) and ChR2 (right) expressing animals over the 15 min RTPP session. Note increased occupancy of light-paired (stim) side in ChR2 animals. Scale bar = 15 cm.

C. Summary of RTPP. Left, Proportion of time spent on stim side (*left*) and total distance travelled (*right*) in GFP and ChR2 animals. Note consistent preference for stim side in ChR2 animals.

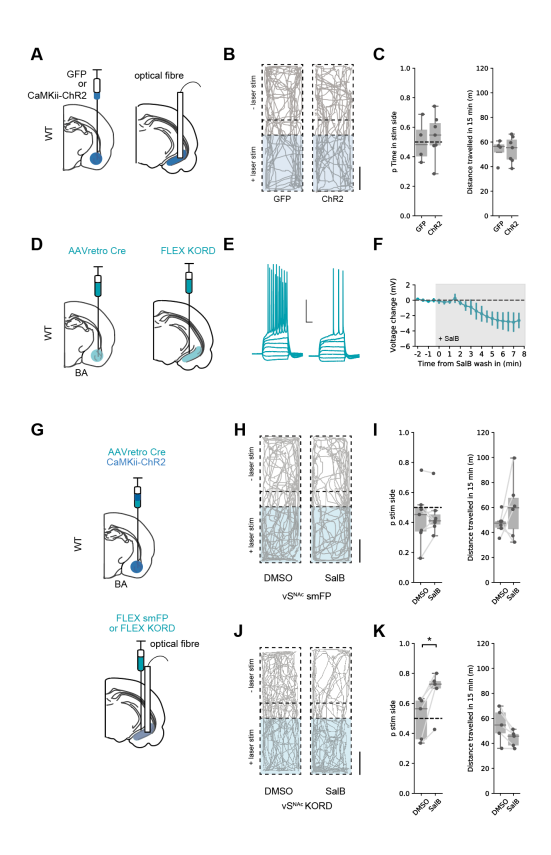
D. Strategy to express KORD in vH^{NAc} neurons.

E,F. Bath application of SalB (100 nm) hyperpolarises KORD expressing vH^{NAc} neurons, and reduces AP firing. See Figure S4 for full quantification. Scale bar = 30 mV, 100 ms.

G. Schematic of strategy to inhibit vH^{NAc} neurons during BA input driven RTPP.

H,I. As **B,C** but comparing the effect of either DMSO (vehicle) or SalB (KORD agonist) injections 15 mins before testing in control mice. Note consistent RTPP in both conditions indicating no effect of SalB in control mice.

J,K. As H,I, but in mice expressing KORD in vH^{NAc} neurons. Note loss of RTPP in SalB injected mice compared to controls.



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Figure 7 | Excitatory BA input supports real time place preference only after inhibition of vH^{BA} neurons.

A. Schematic of experiment. GFP or excitation-specific CaMKii ChR2 were expressed in BA, and an optic fibre implanted in vH.

B. Real time place preference (RTPP) assay. One side of a chamber was paired with 20 Hz blue light stimulation. Example trajectories of GFP (left) and ChR2 (right) expressing animals over the 15 min RTPP session. Note lack of preference for light-paired (stim) side in either group. Scale bar = 15 cm

C. Summary of RTPP. Left, Proportion of time spent on stim side (*left*) and total distance travelled (*right*) in GFP and ChR2 animals. Note lack of preference for stim side in either condition.

D. Strategy to express KORD in vH^{BA} neurons.

E,F. Bath application of SalB (100 nm) hyperpolarises KORD expressing vH^{BA} neurons, and reduces AP firing. See Figure S4 for full quantification. Scale bar = 30 mV, 100 ms.

G. Schematic of strategy to inhibit vH^{BA} neurons during BA input driven RTPP.

H,I. As B,C but comparing the effect of either DMSO (vehicle) or SalB (KORD agonist) injections 15 mins before testing in control mice. Note lack of RTPP in both conditions indicating no effect of SalB in control mice.

J,K. As H,I, but in mice expressing KORD in vH^{BA} neurons. Note induction of RTPP in SalB injected mice compared to controls.

1 SUPPLEMENTARY FIGURES



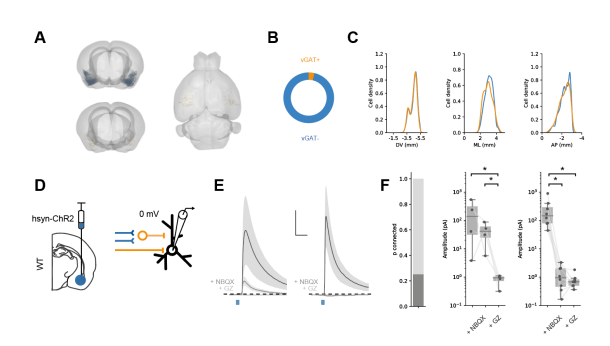


Figure S1 | Feedforward and direct inhibitory input from BA to vH

A. Whole brain distribution of labelled excitatory (blue) and inhibitory (orange) BA neurons after injection of CTX β into vH.

B. Proportion of CTX β labelled neurons that are vGAT+ (orange) and vGAT- (blue).

C. Distribution of CTX β neurons that were vGAT+ (orange) and vGAT- (blue) across three anatomical axes. Note similar overall distribution.

D. Schematic of experiment. Pan neuronal synapsin promoter expresses ChR2 in both inhibitory and excitatory neurons, allowing assessment of feedforward and direct BA input to vH.

E. Brief pulses of blue light evoked inhibitory currents at 0 mV. Average current traces pre and post NBQX and GZ. In a proportion of neurons (left) currents not completely blocked by the AMPA receptor antagonist NBQX, but removed by GABA receptor antagonist gabazine indicating presence of direct inhibition. In other recorded neurons, currents were completely blocked by NBQX (right), indicating only feedforward inhibition. Scale bar = 50 pA, 10 ms.

F. *Left*, proportion of neurons with inhibitory currents that received direct inhibition. *Right*, Amplitude before and after NBQX and GZ in two groups of neurons. Note log scale.

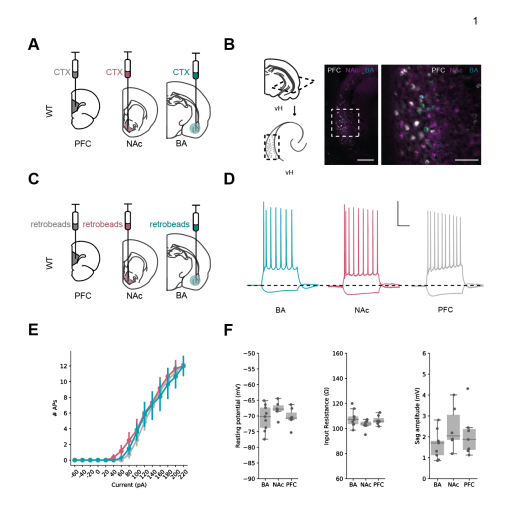


Figure S2 | Parallel output populations in vH.

A. Schematic of experiment, three differently tagged CTX β tracers were injected into PFC, NAc and BA.

B. horizontal section of CA1 / subiculum in vH showing interspersed but non overlapping labelling.

C. Strategy for electrophysiology recordings – projection populations were fluorescently labelled with retrobead injections into downstream projection areas.

D. Examples of positive (+160 pA) and negative (-40 pA) current steps in fluorescently targeted neurons from each population. Scale bar = 30 mV, 100 ms.

E,F. No large differences in input / output curve, resting potential, input resistance or sag amplitude across the three populations.

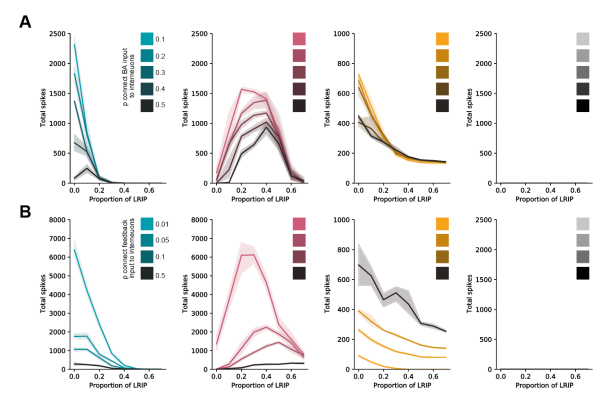


Figure S3 | The switch in vH^{BA} and vH^{NAc} activity is robust over a wide range of feedforward and feedback connectivity.

A. Total spiking of model in vH^{BA} neurons (green), vH^{NAc} neurons (red), vH^{PFC} neurons (grey) and interneurons (orange) with increasing levels of BA inhibition, with different levels in feedforward inhibition (darker colours). Note that although absolute firing changes, the switch from vH^{BA} to vH^{NAc} activity with increasing inhibitory input is maintained.

B. As in (**A**) but for different levels of feedback connectivity. Note that again, the switch from vH^{BA} to vH^{NAc} with increasing inhibitory input is maintained

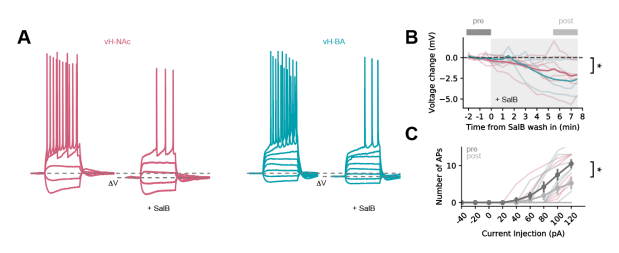


Figure S4 | SalB wash in reduces activity of KORD expressing neurons.

A. example traces before and after SalB wash in in vH^{NAc} neuron (red) and vH^{BA} neuron. Note lower resting potential and reduced spiking after SalB wash in.

B. Summary of effect of SalB wash in on resting potential for vH^{NAc} neurons (red) and vH^{BA} neurons. Thin lines are individual experiments, old line is group average.

C. Summary of effect of SalB wash in on current injection-induced action potential firing. Thin lines show individual vH^{NAc} neurons (red) and vH-BA neurons, grey lines show group average.

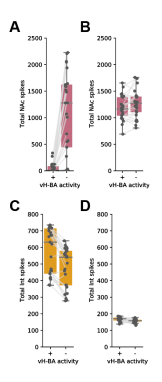


Figure S5 | Removing vH^{BA} activity from the integrate of fire model increases vH^{NAc} activity in response to excitatory but not excitatory and inhibitory BA input.

A. Total vH^{NAc} activity with and without vH^{BA} activity. Individual data points for all configurations of model presented in Figure S3. Note large increase in vH^{NAc} activity on vH^{BA} silencing across a wide range of feedback and feedforward connectivity.

B. As in (**A**) but with both excitatory and inhibitory input (inhibitory input scaled to 40 % of excitatory input (as in example in Figure 5). vH^{BA} silencing has no effect, as these neurons are already effectively silenced by BA inhibitory input.

C,D. As in (A,B) but for interneuron firing. vH-BA silencing reduces interneuron firing, contributing to the increase in vH-NAc activity.

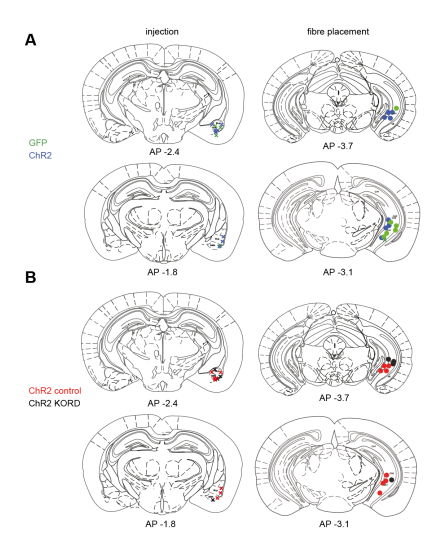


Figure S6 | Histology for behavioural experiments in Figure 6.A. Histology for experiments in Figure 6A-C

B. Histology for experiments in Figure 6G-K

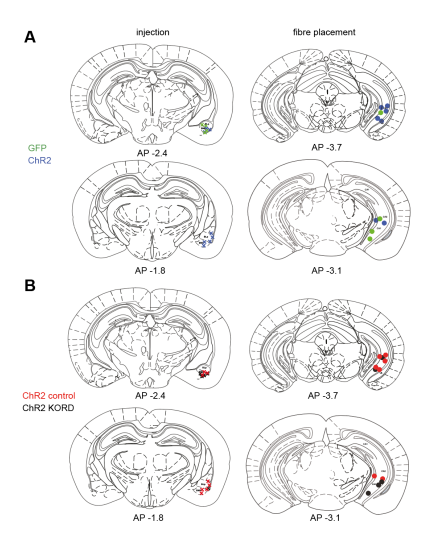


Figure S7| Histology for behavioural experiments in Figure 7.

A. Histology for experiments in Figure 7A-C

B. Histology for experiments in Figure 7G-K

1 STATISTICAL SUMMARY

Figure	Descriptors	n	Test used	Statistic	p-value
2c	Synapsin -70 mV	9	Paired t-test	t ₍₈₎ = 10.04	0.00008
	Amplitude (pA)		(log transformed data)		
	+/- NBQX				
2e	Synapsin 0 mV	12	Paired t-test	t ₍₁₁₎ = 10.15	6.35 x10 ⁻⁷
	Amplitude (pA)		(log transformed data)		
	+/- GZ				
2g	CaMKii 0 mV	3	Repeated-measures ANOVA	F _(2,4) = 23.4	0.006
	Amplitude (pA)		(log transformed data)		
	Baseline				
	+NBQX		Tukey post hoc test		
	+GZ		Baseline vs NBQX	t ₍₂₎ = 4.73	0.001
			Baseline vs GZ	t ₍₂₎ = 4.84	0.001
			NBQX vs GZ	t ₍₂₎ = 0.12	0.9
2i	vGAT 0 mV	6	Repeated-measures ANOVA	F _(2,10) = 16.7	0.0006
	Amplitude (pA)		(log transformed data)		
	Baseline				
	+NBQX		Tukey post hoc test		
	+GZ		Baseline vs NBQX	t ₍₂₎ = 0.06	0.9
			Baseline vs GZ	t ₍₂₎ = 4.92	0.001
			NBQX vs GZ	t ₍₂₎ = 4.98	0.001
3c	BA:NAc -70 mV	9	Wilcoxon Rank Sum	W = 15	0.43
	Amplitude (pA)				
3f	BA:PFC -70 mV	8	Wilcoxon Rank Sum	W = 0	0.0018
	Amplitude (pA)				
3i	BA:NAc vGAT 0 mV	7	Wilcoxon Rank Sum	W = 0	0.016
	Amplitude (pA)				
31	BA:PFC vGAT 0 mV	7	Wilcoxon Rank Sum	W = 0	0.016
01	Amplitude (pA)				
4k	BA:NAc local vGAT 0 mV	10	Wilcoxon Rank Sum	W = 2	0.006
	Amplitude (pA)				
4n	BA:NAc CaMKii -70 mV	10	Wilcoxon Rank Sum	W = 22	0.625
	Amplitude (pA)				-
4p	BA:NAc CaMKii 0 mV	10	Wilcoxon Rank Sum	W = 3	0.04
.6	Amplitude (pA)				
	F				
6c	GFP vs ChR2	GFP = 6	t-test	t _(5.9) = 2.61	0.041
	p stimulated side	ChR2 = 8		-(0.0)	
6c 6f 6i-k	GFP vs ChR2	GFP = 6	t-test	t _(9.2) = 1.27	0.23
	distance travelled	ChR2 = 8		(J.Z) ······	5.20
	Voltage change in	7	Repeated measures ANOVA		
	SalB (mV)		Effect of time	F _(24,144) = 5.94	2.64 x10 ⁻¹²
	DMSO vs SalB	Cont = 7	Mixed ANOVA	· (24,144) - J.J4	2.04 × 10
		Cont = 7 KRD = 4		F 0.00	0.010
	Control vs KORD	<u>κκυ</u> = 4	Effect of group	$F_{(1,10)} = 9.99$	
			Effect of drug	$F_{(1,10)} = 10.6$	0.009
			Interaction	F _(1,10) = 9.75	0.010

6i	DMSO vs SalB	8	Paired t-test	$t_{(7)} = 0.84$	0.43
	p stimulated side				
6i	DMSO vs SalB	8	Paired t-test	t ₍₇₎ = 0.99	0.35
	distance travelled				
6k	DMSO vs SalB	4	Paired t-test	t ₍₃₎ = 4.72	0.018
	p stimulated side				
6k	DMSO vs SalB	4	Paired t-test	t ₍₃₎ = 1.12	0.34
	distance travelled				
7c	GFP vs ChR2	GFP = 4	t-test	t _(6.4) = 0.40	0.70
10	p stimulated side	ChR2 = 7		10.4) 0.10	
7c	GFP vs ChR2	GFP = 4	t-test	t _(6.9) = 0.08	0.94
	distance travelled	ChR2 = 8			
7f	Voltage change in	3	Repeated measures ANOVA		
	SalB (mV)		Effect of time	F _(19,38) = 2.95	0.002
7i-k	DMSO vs SalB	Cont = 7	Mixed ANOVA		
	Control vs KORD	KRD = 5	Effect of group	$F_{(1,10)} = 3.05$	0.11
			Effect of drug	F _(1,10) = 8.96	0.013
			Interaction	F _(1,10) = 9.20	0.013
7i	DMSO vs SalB	7	Paired t-test	t ₍₆₎ = 0.33	0.75
	p stimulated side				
7i	DMSO vs SalB	7	Paired t-test	t ₍₆₎ = 0.70	0.51
	distance travelled				
7k	DMSO vs SalB	5	Paired t-test	t ₍₄₎ = 4.13	0.014
	p stimulated side				
7k	DMSO vs SalB	5	Paired t-test	t ₍₄₎ = 2.77	0.05
	distance travelled				

1 SUPPLEMENTARY STATSTICS SUMMARY

Figure	Descriptors	n	Test used	Statistic	p-value
S1f	With direct inhibition	4	Repeated-measures ANOVA	F _(2,6) = 7.9	0.02
	Amplitude (pA)		(log transformed data)		
	Baseline				
	+NBQX		Tukey post hoc test		
	+GZ		Baseline vs NBQX	t ₍₃₎ = 0.7	0.7
			Baseline vs GZ	t ₍₃₎ = 4.7	0.001
			NBQX vs GZ	t ₍₃₎ = 3.5	0.002
S1f	No direct inhibition	8	Repeated-measures ANOVA	F _(2,14) = 84.7	1.5 x10 ⁻⁸
	Amplitude (pA)		(log transformed data)		
	Baseline				
	+NBQX		Tukey post hoc test		
	+GZ		Baseline vs NBQX	t ₍₇₎ = 12.1	0.001
			Baseline vs GZ	t ₍₇₎ = 12.7	0.001
			NBQX vs GZ	t ₍₇₎ = 0.53	0.84
S2f	Resting Potential (mV)	BA = 9	One-Way ANOVA	F _(2,19) = 1.38	0.27
	BA, NAc, PFC	NAc = 7			
		PFC = 7			
S2f	Inpit Resistance (Ohms)	BA = 9	One-Way ANOVA	F _(2,19) = 1.77	0.20
	BA, NAc, PFC	NAc = 7			
		PFC = 7			
S2f	Sag Amplitude (mV)	BA = 9	One-Way ANOVA	F _(2,19) = 1.40	0.27
	BA, NAc, PFC	NAc = 7			
		PFC = 7			
S4b	Pre vs post SalB	vH-BA = 3	Mixed ANOVA		
		vH-NAc = 7	Effect of group	$F_{(1,8)} = 0.29$	0.6
			Effect of drug	F _(1,8) = 13.8	0.007
			Interaction	$F_{(1,8)} = 0.35$	0.57
S4c	Pre vs post SalB	vH-BA = 2	Mixed ANOVA		
		vH-NAc = 6	Effect of group	$F_{(1,6)} = 0.21$	0.66
			Effect of drug	F _(1,6) = 31.3	0.001
			Interaction	$F_{(1,6)} = 0.75$	0.42

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