A hippocampus-accumbens code guides goal-directed appetitive behavior

Oliver Barnstedt^{1,2 \boxtimes}, Petra Mocellin^{1,3}, and Stefan Remy^{1,2,4,5 \boxtimes}

¹Department of Cellular Neuroscience, Leibniz Institute for Neurobiology, 39118 Magdeburg, Germany
 ²German Center for Neurodegenerative Diseases (DZNE), 53127 Bonn & 39120 Magdeburg, Germany
 ³International Max Planck Research School for Brain & Behavior (IMPRS), 53175 Bonn, Germany
 ⁴Center for Behavioral Brain Sciences (CBBS), 39118 Magdeburg, Germany
 ⁵German Center for Mental Health (DZGP), Magdeburg site, 39106 Magdeburg, Germany

Neurons in dorsal hippocampus (dHPC) encode a rich repertoire of task-relevant environmental features, while downstream regions such as the nucleus accumbens (NAc) translate this information into task-adaptive behaviors. Yet, the contents of this information stream and their acute role in behavior remain largely unknown. Here, we used in vivo dual-color two-photon imaging to simultaneously record from both large numbers of hippocampal neurons and an identified subpopulation of NAcprojecting neurons during goal-directed navigation towards a hidden reward zone. We found that NAc-projecting neurons contain enriched spatial information and enhanced representations of non-spatial task-relevant behaviors such as deceleration and appetitive licking, both of which could be elicited by optogenetic activation of dHPC terminals in NAc. A generalized linear model revealed enhanced conjunctive coding in NAc-projecting dHPC neurons, improving the identification of the reward zone. We propose that dHPC routes specific reward-related spatial and behavioral state information to guide NAc action selection.

19 hippocampus | nucleus accumbens | spatial memory | place cell | reward
 20 learning | conjunctive coding | appetitive behavior | calcium imaging

21 Correspondence:
22 oliver.barnstedt@lin-magdeburg.de or stefan.remy@lin-magdeburg.de

3 Introduction

Memories allow an organism to use past experience to optimize current and future behaviors (1). The hippocampus (HPC) is widely recognized as one of the main sites of memory-related plasticity (2–4). Yet, while our understanding of memory processing within the HPC has greatly advanced, surprisingly little is known about the translation of this information into behavioral action: which hippocampal output pathways send which types of information to guide memory-driven behavior? This lack of knowledge corresponds with the scarcity of data available on the activity profile of large numbers of identified hippocampal projection neurons while animals perform memory-dependent behaviors.

The HPC is a major part of the brain's limbic system, processing various kinds of memory by receiving highly processed sensory information from the entorhinal cortex, and sending behaviorally relevant outputs to diverse brain regions via hippocampal Cornu Ammonis field 1 (CA1) and subiculum (5). Its elevated role for spatial memory is highlighted by the presence of spatially tuned "place cells" that form a "cognitive map" to support goal-directed navigation (6, 7).

This cognitive map is further supported by neuronal coding of navigationally relevant features such as borders, speed, reward/goal locations (8–11), but also non-spatial information such as that about future decisions or behavioral tasks (12, 13). The conjunction of this variety of encoded features is suggested to provide a "scaffold" that supports the formation of episodic memories (14).

Previous models of hippocampal memory processing assumed largely homogeneous cell populations (15, 16). However, hippocampal principal neurons are increasingly recognized as structurally and functionally diverse in terms of their morphology, electrophysiology, transcriptomic cell types, anatomical differences across its three axes, and projection patterns (17, 18). Such heterogeneity may provide the hippocampus with the intrinsic flexibility to meet the various demands of diverse environments (17), and to route task-relevant information to specific output targets (19, 20).

One target receiving strong projections from both dorsal and ventral CA1 and subiculum is the nucleus accumbens (NAc) (21, 22), a basal ganglia brain structure crucial for valuebased action selection (23, 24). Described as the "interface between limbic and motor circuitry" (25), it has been suggested as the main site of transformation from a hippocampal spatial code into a motivation-driven motor code (26). In line with this, NAc neurons are spatially tuned, required for spatial memory acquisition and consolidation, and display task-dependent synchrony with dHPC neurons (27–30). Indeed, disabling HPC

NAc projections diminished conditioned place preference (CPP) (31, 32), and optogenetic stimulation was sufficient to artificially induce CPP (33, 34). While these findings cement the NAc's role as an indispensable hippocampus output node, surprisingly little is known about which spatial, contextual and behavioral patterns of hippocampal information the NAc receives. Here, we set out to understand the specific contents of this information stream during goal-directed navigation and its acute role in behavior. We hypothesized that the HPC may selectively route both spatial as well as other task-relevant information to aid the NAc in action selection.

To test this, we employed dual-color two-photon imaging, capturing *in vivo* calcium signals of large populations of neurons in dHPC, while using a projection-specific red fluorophore to allow identification of NAc-projecting neurons (dHPC $^{\to NAc}$). Directly comparing dHPC $^{\to NAc}$ activity with

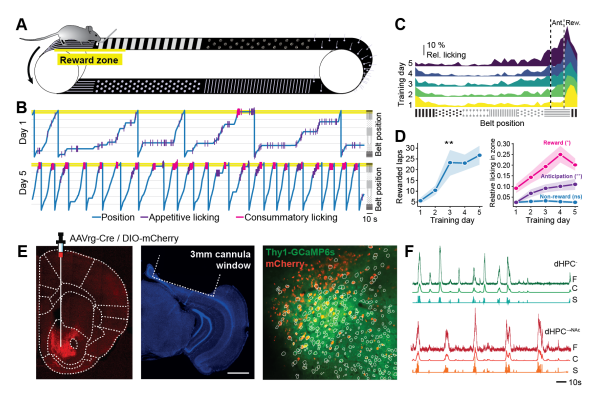


Figure 1. Dual-color two-photon calcium imaging of dHPC allows projection-specific activity monitoring during goal-directed navigation. (A-D) Training mice on a spatial reward learning task. (A) Schematic of behavioral task. Mice moved on a cue-enriched self-propelled treadmill belt of 360 cm length and obtained a liquid reward when licking at a spout in a hidden reward zone of 30 cm length (yellow). (B) Representative traces of one mouse's behavior on training day 1 and day 5. (C) Licking in anticipation (Ant.) and reward (Rew.) zone increases over the course of training. (D) Number of rewarded laps per training session (F(4) = 6.344, GG-correction), anticipatory (F(4) = 3.803) and reward licking (F(4) = 4.276). GG-correction) significantly increased over the course of five training days (n = 18) mice, repeated-measures ANOVA). See also Figure S1. (E) Schematic of dual-color projection neuron imaging method. Thy1-GCaMP6s mice were injected with AAVrg-Cre in the medial NAc and DIO-mCherry in dHPC. Representative coronal brain slice showing axonal mCherry expression in NAc (AP -1.3; left). Representative coronal brain slice stained with DAPI (blue) of dHPC, showing the outlines of the 3 mm cannula window used for imaging; scale bar represents 1 mm (second left). Field of view of one sample experiment showing Thy1-GCaMP6s expression in green and mCherry expression of putative NAc-projecting neurons in red; outlines show detected components used for analysis (right). (F) Two representative neurons' raw ("F"), denoised and deconvolved ("C") and event ("S") traces; red traces indicate mCherry co-expression (dHPC Nac). See also Figure S2. All data are presented as mean \pm SEM. *p < 0.05, **p < 0.01.

the rest of the dHPC population (dHPC⁻), we found enhanced spatial tuning in dHPC - NAc neurons, with strong modulation by local cue boundaries and a reward zone. We also found elevated coding of low velocities and appetitive licking, both of which could be elicited by optogenetic stimulation. Lastly, we show stronger conjunctive coding of space, velocity, and appetitive licking, which improves classification of the reward zone with a linear decoder. We thus propose that dHPC routes reward context-enriched spatial and behavioral state information to bias NAc action selection.

Results

Dual-color calcium imaging during goal-directed navigation. To investigate coding of spatial information and goal-directed behavior in dHPC -NAc neurons, we trained food-restricted mice on a head-fixed spatial reward learning task. Mice had to run on a self-propelled treadmill, traversing a 360 cm long textile belt with six differently textured zones, including one otherwise unmarked 30 cm long fixed reward zone (35, 36). Licking a spout in this zone causes a liquid reward to be dispensed once per lap (Figure 1A and Figure S1A-E). Mice underwent five days of training in which they learned to obtain more rewards by progressively increasing

their licking in both reward zone and a 30 cm "anticipation 111 zone" preceding the reward zone (Figure 1B-D and Figure 112 \$1F-N).

113

122

To understand the neural coding properties of large numbers 114 of dHPC neurons in behaving animals, we turned to 115 dual-color two-photon imaging in mice pan-neuronally expressing the calcium indicator GCaMP6s, as well as the static 117 red marker mCherry in defined NAc-projecting neurons. For 118 this, we injected AAVrg-Cre in NAc and DIO-mCherry in 119 dHPC (CA1/subiculum border region) of Thy1-GCaMP6s mice (37). This approach allowed us to obtain dynamic calcium signals both in a large majority of mCherry-negative hippocampal neurons (dHPC⁻) and specifically mCherry coexpressing NAc-projecting neurons (dHPC^{→NAc}), simultaneously within the same field of view using the same calcium 125 indicator (Figure 1E and Figure S2). It allowed us to overcome constraints of electrophysiological studies such as relatively low sample sizes (19, 20) or indirect connectivity measurements (38, 39). Optical access to dorsal CA1 and prosubiculum (also known as proximal subiculum (40, 41)) was established by implanting a chronic hippocampal window after virus injections (Figure S1C) (42, 43). Imaging data was 132 acquired after 5 days of behavioral training, was motioncorrected using NormCorre, and spatio-temporal components

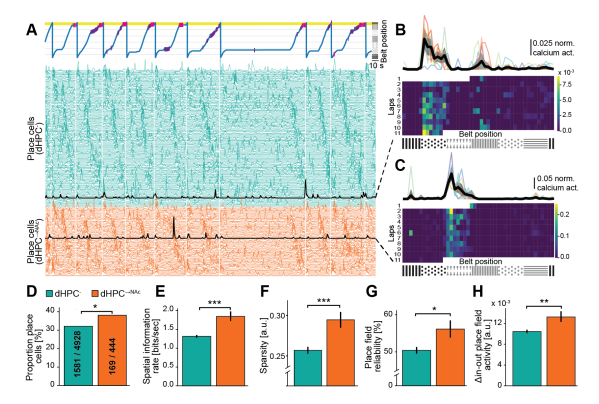


Figure 2. dHPC - Nac neurons carry enhanced and more reliable spatial information. (A) Excerpt from one representative recording session showing behavioral activity (top) and denoised neural activity of identified place cells (bottom) over the course of several minutes from one mouse. White vertical lines mark new laps. Traces are ordered according to whether neurons co-expressed mCherry (red; dHPC⁻)Ac) or not (green; dHPC⁻) and their respective place fields. Black traces are example neurons shown in (B and C). (B and C) Spatial activity of one neuron without (B) and with (C) mCherry co-expression over one session. Calcium events are binned by position on the belt (x axis) and laps (y axis) and show consistent activity at one position on the belt. (D-H) Comparisons of spatial tuning characteristics between dHPC⁻ (green) and dHPC⁻ (red) neurons. NAc-projecting neurons contain a higher proportion of place cells (D, $\chi^2(1, 5372) = 6.364$); these place cells contain more spatial information per second (E, Welch's t(186.55) = 4.770), show increased sparsity (F. Welch's t(218.79) = 3.657), higher reliability (G, probability of firing maximally within their place field per lap; Welch's t(203.46) = 2.479), and higher in-place field activity (H, Welch's t(190.32) = 2.884). n = 6 mice, 19 imaging sessions, 5,372 (inc. 444 mCherry-coexpressing) neurons, 1,750 (inc. 169 mCherry-coexpressing) place cells. *p < 0.05, **p < 0.01, ***p < 0.001. See also Figure S3.

vere extracted using constrained non-negative matrix factorization (CNMF) (44). We thus obtained calcium signals for further analysis from a total of 5,372 GCaMP-expressing neurons including 444 putative dHPC→NAc neurons in 6 mice across 19 imaging sessions (Figure 1F and Figures S2 and S3). These numbers approximate previously established proortions of NAc-projecting neurons in dorsal prosubiculum and distal CA1 (20, 45, 46).

136

138

140

144

146

148

150

Enhanced spatial coding by dHPC→NAc neurons. Given ne dHPC's well-established role in representing spatial information and the role of dHPC^{→NAc} projections in spatial memory tasks (31, 32), we wondered if and how spatial information may be encoded by dHPC^{→NAc} neurons compared to dHPC⁻ neurons. For this, we extracted calcium events for each neuron and compared this activity across the linear spatial environment of the treadmill belt as mice traversed it lap by lap. Indeed, we found large numbers of both dHPC⁻ and dHPC^{→NAc} neurons with repeatedly elevated calcium levels on the same positions of the belt (Figure 2A-C and Figure S3). Comparing each neuron's spatial information content (47) with that of a randomly shuffled distribution, we identified a significantly higher proportion of such place cells in dHPC→NAc neurons (169/444 neurons; 38 %) compared to dHPC⁻ neurons (1,581/4,928 neurons; 32 %; χ^2 , p = 0.012; Figure 2D). Within this population of place cells, we further 159 analyzed how specifically space is encoded, and found that dHPC→NAc place cells had a higher spatial information rate (47) (p < 0.001, Welch's t-test; Figure 2E). They also had significantly higher levels of sparsity (p < 0.001, Welch's ttest; Figure 2F), a measure for how diffuse a neuron is firing in the spatial domain (48). Furthermore, both the relative calcium activity (activity inside the place field – activity outside the place field; p = 0.0044, Welch's t-test) and reliability of in-place field activity per lap (p = 0.014, Welch's t-test) were significantly higher for dHPC^{NAc} place cells compared to dHPC⁻ neurons (Figure 2G-H). These results suggest that the dHPC routes enhanced and more reliable spatial information to NAc compared to the general dHPC population, in line with previous results pointing towards the necessity of dHPC^{→NAc} projections for spatial memory expression (31, 32).

Place fields are modulated by local cue boundaries and are overrepresented near the reward zone. Previous studies found that place fields are often not homogenously distributed across the environment but can be modulated by salient environmental features such as textures, borders, or reward/goal zones (11, 49, 50). We hypothesized that information about such spatial features may be preferen-

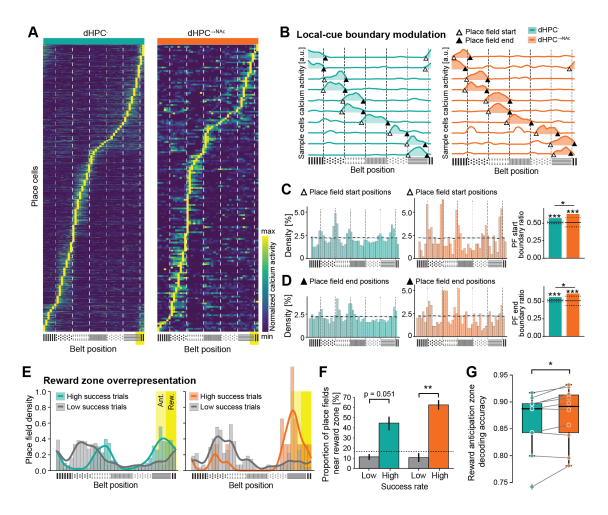


Figure 3. dHPC^{→NAc} place fields are modulated by local cues and reward zone. (A) Heat maps of dHPC⁻ (left) and dHPC^{-→NAc} (right) place cells' normalized positionbinned average calcium events, ordered by place field location. Texture boundaries are indicated as white dashed lines. Yellow rectangle represents reward zone. (B) Example place fields with edges near belt texture boundaries. Primary place fields are indicated by color fill. Triangles mark start (no fill) and end (fill) points. Dashed black lines represent texture boundaries. (C and D) Place field edges accumulate near texture boundary areas. Histograms of dHPC⁻ (green) and dHPC^{-→NAc} (red) neurons' place field start (C) and end (D). Dotted line and shade represent average and 95th CI of 1000x randomly shuffled place fields. Both dHPC⁻ and dHPC⁻ nlace field start and end positions are significantly overrepresented at the 99.9th percentile (dotted black lines) compared to a randomly shuffled distribution. Start ($\chi^2(1,5134)=5.735$) and end positions (χ²(1, 5217) = 4.397) of dHPC → NAc place fields are furthermore significantly overrepresented compared to the dHPC population. (E and F) Place cells are overrepresented near reward zone in high success trials. (E) Histograms (bars) and kernel density estimations (KDEs; lines) of place field centers for dHPC⁻ (left) and dHPC →NAc (right) neurons, split into high success trials (green/red) and low success trials (gray). Reward zone (Rew.; yellow) and anticipation zone (Ant.; bright yellow) are indicated as rectangles. (F) Proportion of place fields in reward and vicinity zone is significantly higher in high-success trials (colored bars) compared to low-success trials (gray bars) in NAc-projecting neurons (red) but not in dHPC⁻ neurons (green). 2-way ANOVA, $F_{\text{success}}(1,1) = 54.918$, p < 0.001, $F_{\text{projection}}(1,1) = 0.958$, p = 0.338, $F_{\text{interaction}}(1,1) = 2.969, p = 0.098.$ Post-hoc Welch's t-tests with Bonferroni correction: t_{dHPC} -(3.561) = 3.698; t_{dHPC} -Nac (3.479) = 8.671; $t_{\text{low success}}(13.629) = 0.093, p = 0.098$ 1; thigh success (3.952) = 2.075, p = 0.215. Dashed line represents an even distribution of reward and anticipation zone. (G) A linear classifier shows significantly increased decoding accuracy of reward anticipation zone based on dHPC $^{\rightarrow NAc}$ neural activity compared to that of sample size-matched dHPC $^{-}$ neurons. Wilcoxon's t-test, W(9) = 5.0, n = 10 imaging sessions. All data are presented as mean \pm SEM. *p < 0.05, **p < 0.01, ***p < 0.001. See also Figure S4.

tially routed to NAc, given the relevance of NAc for spatial memory (29, 30) and particularly for reward-related behaviors (51).

Upon inspection of place cells' spatially binned calcium profiles (Figure 3A-B and Figures S3, S4A), we noticed an overabundance of place fields that seemed to cover entire texture zones, starting at the beginning of a texture zone and ending before the next texture zone began. To quantify this observation, we plotted the densities of place field start and end positions across the belt and compared the observed densities with those of randomly shuffled place fields. We found a significant overrepresentation of place field start and end positions (but not centers) near texture boundaries for both dHPC⁻ and dHPC^{→NAc} populations (ratios >99.9th percentile, permutation test). This effect was more pronounced 197 in dHPC $^{\rightarrow NAc}$ neurons (PF start: p = 0.033, PF end: p =0.036, χ^2 ; Figure 3C-D and Figure S4A-C). This suggests that dHPC → NAc neurons are more strongly modulated by local cue boundaries.

Mice were trained to lick for reward in a hidden reward 202 zone. Previous studies have shown that such zones and their 203 preceding vicinity are often over-represented by place cells 204 (35, 52, 53) – we hypothesized that this effect might involve 205 dHPC→NAc neurons, given the NAc's role in reward-related 206 behaviors (51). When pooled across all sessions, we found 207 little evidence of such an overrepresentation (Figure \$4D-E). Mouse behavior generally shows great variability though, 2009 and previous work demonstrated the dependence of such an 210

188

190

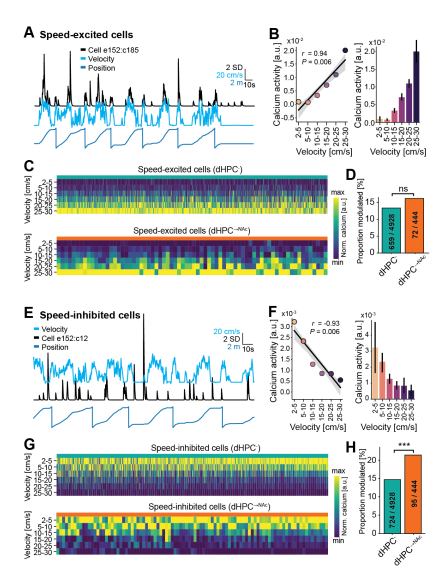


Figure 4. Speed-inhibited cells are overrepresented in $dHPC^{\rightarrow NAc}$ neurons. (A-D) Speed-excited dHPC neurons. (A and B) Representative example of one speed-excited neuron. (A) Sample traces of velocity, position and one neuron's denoised calcium activity. Note the increased calcium activity in times of high velocity, irrespective of position. (B) Linear regression on average calcium events per velocity bin shows a significant positive relationship (slope = 7.43×10⁻⁴, intercept = -2.097×10^{-3} , r = 0.937, p = 0.0058). (C) Heatmaps of speed-binned normalized calcium activity of all significantly positively speed-modulated dHPC⁻ (top) and dHPC^{→NAc} (bottom) neurons. (D) Proportions of speed-excited neurons are comparable between dHPC⁻ and dHPC^{→NAc} populations $(\chi^2(1, 5372) = 2.565, p = 0.109)$. (E-H) Speed-inhibited dHPC neurons. (E and F) Representative example of one speedinhibited neuron. (E) Sample traces of velocity, position and one neuron's denoised calcium activity. Note the increased calcium activity in times of low velocity, irrespective of position. (F) Linear regression on average calcium events per velocity bin shows a significant negative relationship (slope = -1.0437×10^{-4} , intercept = 2.814×10^{-3} , r = -0.933, p =0.0065). (G) Heatmaps of speed-binned normalized calcium activity of all significantly negatively speed-modulated dHPC (top) and $dHPC^{\rightarrow NAc}$ (bottom) neurons. (H) Negatively tuned neurons are overrepresented in the NAc-projecting population $(\chi^2(1, 5372) = 13.66, p = 0.00022)$. Speed-modulated neurons were classified as showing a significant linear regression at p < 0.05 after Benjamini/Hochberg FDR correction. All data are presented as mean ± SEM. ns: not significant, ***p < 0.001.

overrepresentation on individual task success (35, 52). We thus divided sessions into high- and low-success based on lick performance (lick precision and reward dispensation, see Methods) and found significantly more dHPC $^{\rightarrow \text{NAc}}$ place cells near the reward zone (reward and anticipation zones) in high-success sessions compared to low-success sessions (p = 0.0036, Welch's t-test). In contrast, dHPC $^{-}$ neurons only showed a trend towards significance (p = 0.051, Welch's t-test; Figure 3E-F). In line with previous studies, both neuronal populations also showed a strong correlation between the success rate (percentage of rewarded laps) and the proportion of place fields near the reward zone across sessions (r = 0.62; Figure S4F).

If reward zone information is preferentially encoded in dHPC^{→NAc} populations and NAc neurons play an integral part in reward-related behaviors, we would expect linear decoders to identify reward or anticipation zones more accurately based on dHPC^{→NAc} activity compared to that of dHPC⁻. To test this, we trained a linear classifier based on a support vector machine (SVM) with calcium activity and reward zone information on odd/even laps and tested decod-

ing accuracy on even/odd laps, using either dHPC $^-$ or sample size-matched dHPC $^{\to NAc}$ populations (see Methods). We found that, within individual sessions, decoding accuracy of the reward anticipation zone was significantly enhanced for dHPC $^{\to NAc}$ populations (p=0.0195, Wilcoxon's test; Figure 3G). These findings demonstrate significant modulation of dHPC $^{\to NAc}$ neurons by local cue boundaries and enhanced reward zone coding.

Enhanced coding of low velocities in dHPC→NAc neurons. As correct performance in the spatial reward learning task goes hand in hand with a reduction in velocity and an increase of licking near the reward zone (see Figure S1K-243 N), we wondered if such non-spatial task-relevant behavioral features were encoded by dHPC neurons and its projections to NAc (9, 54, 55). We hypothesized that NAc may have privileged access to information on low velocities as mice generally slow down near the reward zone, presumably to allow for better discrimination and to engage in anticipatory licking. In line with previous analyses of speed coding in hippocampal and parahippocampal regions (8, 56), we averaged each neuron's calcium activity per velocity bin from

213

215

217

219

221

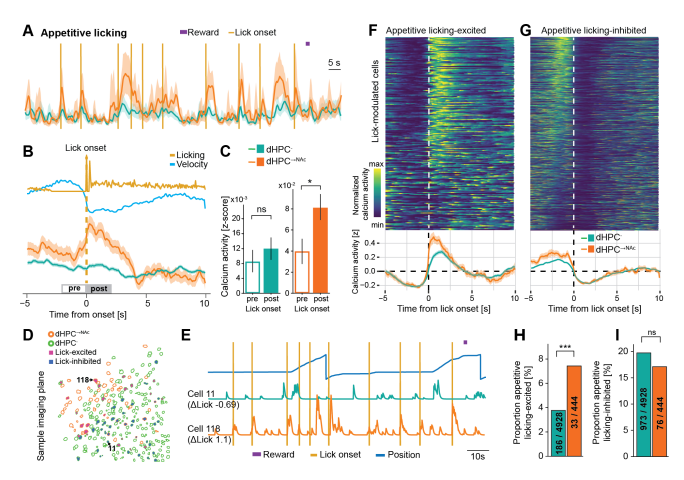


Figure 5. dHPC $^{\rightarrow$ NAc} neurons are excited by appetitive licking and are over-represented in lick-excited neurons. (A-C) Appetitive licking is accompanied by increased neural activity in dHPC $^{\rightarrow}$ NAc neurons but not dHPC $^{-}$ neurons. (A) Representative example showing reward dispensation (purple bar) and appetitive licking onsets (golden vertical lines) and population calcium activity from dHPC $^{-}$ (green) and dHPC $^{\rightarrow}$ NAc (red) neurons. Note the robust increase of neural activity around lick onsets in the dHPC $^{\rightarrow}$ NAc population. (B) Event-triggered average traces around appetitive licking onset, including licking and speed as well as population average calcium activity of dHPC $^{-}$ (green) and dHPC $^{\rightarrow}$ NAc (red) neurons. Gray rectangles indicate time windows for comparing calcium activity shown in (C). (C) Calcium activity is differentially modulated by appetitive licking onset only in dHPC $^{\rightarrow}$ NAc neurons; two-way mixed ANOVA; $F_{\text{licktiming}}(1, 5370) = 2.843$, p = 0.0918, $F_{\text{projection}}(1, 5370) = 43.779$, p < 0.001, $F_{\text{interaction}}(1, 5370) = 7.073$, p = 0.0079. Post-hoc t-tests with Bonferroni correction: $t_{\text{dHPC}}(4927) = 0.871$, p = 0.768; $t_{\text{dHPC}}\rightarrow$ NAc (443) = 2.470, p = 0.0277. (D and E) Example imaging session and traces showing lick-excited and lick-inhibited neurons. (D) Field of view showing spatial profiles of dHPC $^{-}$ (green outlines) and dHPC $^{\rightarrow}$ NAc (red outlines), some of which are classified as lick-excited (violet fill) or lick-inhibited (dark blue fill); neurons #11 and #118 are highlighted. (E) Behavioral traces and calcium activity of sample neurons #11 (lick-inhibited) and #118 (lick-excited). (F and G) Heatmaps (top) and event-triggered calcium activity averages (bottom) of neurons classified as appetitive licking-excited (F) and licking-inhibited neurons is not different between populations; $\chi^2(1, 5372) = 1.626$, p = 0.202. All data are presented as mean ± SEM. ns: not significant, *p < 0.005, ****p < 0.001. See also Figure S5.

2 to 30 cm/s and regressed this activity against velocity. Neurons with a significant regression model (after correcting for false discovery rate) and positive slope were classified as speed-excited (Figure 4A-B). We found around 15 % of neurons were positively speed-modulated, with comparable proportions between dHPC⁻ and dHPC⁻NAc neurons (13 % vs. 16 %, p = 0.109, χ^2 ; Figure 4C-D). We also identified neurons with a significant velocity regression but a negative slope (Figure 4E-F). We found approximately 15 % of such speed-inhibited neurons, with a significantly larger proportion among the dHPC⁻NAc population (15 % vs. 21 %, p = 0.00022, χ^2 ; Figure 4G-H). These results suggest widespread modulation of dHPC neuronal activity by non-spatial features such as velocity, with dHPC⁻NAc neurons specifically over-representing low speeds.

Overrepresentation of appetitive licking-excited dHPC → NAc neurons. Besides a decrease in velocity when

approaching the reward zone, mice also increasingly engaged 270 in licking behavior (see Figure 1B-D). Given the NAc's dual 271 role in appetitive and consummatory behaviors (57), we 272 wondered if licking behaviors might be reflected in the neural 273 activity of dHPC→NAc neurons. For this, we distinguished 274 between consummatory licking which occurs after a reward 275 is dispensed and allows the mouse to consume the reward 276 provided, and appetitive licking which is an operant behavior 277 that will lead to reward dispensation when performed at 278 the correct location on the belt. We found a significant 279 decrease of calcium activity during reward consumption in 280 both dHPC⁻ and dHPC $^{\rightarrow NAc}$ populations (Figure S5A-C). Appetitive licking, on the other hand, had no apparent effect on neural activity in dHPC- neurons but coincided with 283 a significant increase in calcium activity in the dHPC^{→NAc} population that began about one second before lick onset 285 and then fell below baseline 4 seconds after licking (Figure 5A-C). We investigated if this population-averaged data 287

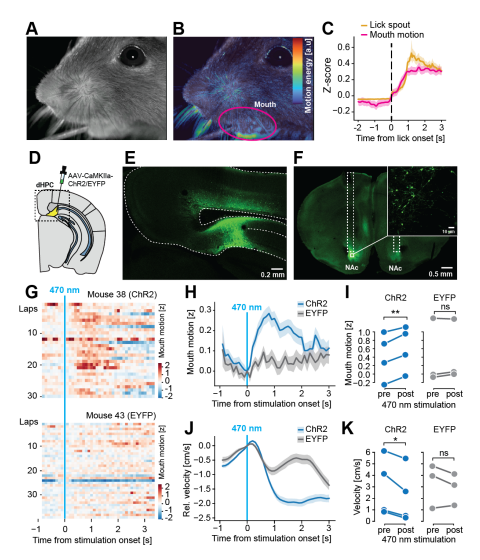


Figure 6. Optogenetic stimulation of dHPC axons in NAc induces mouth movement and deceleration. (A-C) Tracking of orofacial movements via infrared camera recordings. (A) Example still image from near-infrared camera, sampled at 75 Hz. (B) False-color coded motion energy (pixel-by-pixel intensity difference to previous image) overlaid over sample image from (A). Automatically segmented mouth region is indicated by a purple line. (C) Average motion energy in mouth region increases with licking. Golden line shows analog lick spout signal. (D-F) Experimental approach for optogenetic $dHPC\overset{\cdot}{\rightarrow}{}^{NAc}$ stimulation. (D) Injection schematic. (E) Somatic expression of ChR2-EYFP in dorsal prosubiculum. (F) Axonal expression of ChR2-EYFF in the NAc, where light fibers are placed (tracts indicated by white dotted lines). (G) Representative examples of mouth motion around onset of optogenetic stimulation in an animal expressing ChR2 (top) or EYFP control (bottom). (H) Trial-averaged mouth motion activity around time of optogenetic stimulation in ChR2 (blue) and EYFP (gray)-expressing mice. (I) Mouth motion is significantly increased with optogenetic stimulation in ChR2 animals (t(3) = 7.485) p = 0.00494) but not EFYP animals (t(2) = 1.353; p = 0.309). Paired t-tests; n = 4 mice (ChR2), n = 4= 3 mice (EYFP). (J) Trial-averaged relative velocity around time of optogenetic stimulation in ChR2 (blue) and EYFP (gray)-expressing mice. (K) Velocity is significantly decreased with optogenetic stimulation in ChR2 animals (t(3) = -3.551; p = 0.0381) but not EFYP animals t(2) = -1.263; p = 0.334. Paired ttests; n = 4 mice (ChR2), n = 3 mice (EYFP). All data are presented as mean ± SEM. ns: not significant, *p < 0.05. **p < 0.01.

is reflected on the single-cell level and if there are individual ells that are reliably modulated by appetitive licking (Figure 5D-E and Figure S5D-E). Comparing the pre- and post-lick neural activity for each neuron for each appetitive lick event, we identified a total of 1,268 neurons (24 %) that were significantly (negatively or positively) modulated by appetitive licking (Figure 5F-I). Interestingly, while the proportion of lick-inhibited neurons was comparable between dHPC⁻ and dHPC $^{\rightarrow \text{NAc}}$ populations (19.7 % vs 17.1 %, p = 0.20, χ^2 ; Figure 5I), we found a significantly larger proportion of lickexcited neurons in dHPC→NAc populations (3.8 % vs 7.4 %, $< 0.001, \chi^2$; Figure 5H). These findings suggest that dHPC does not route reward information per se to NAc, but rather information on appetitive behaviors required to obtain such

Optogenetic activation of dHPC terminals in NAc induces mouth movement and deceleration. Given the NAc's hypothesized role as an interface between limbic and motor systems (25), we investigated if the observed increase of calcium activity in dHPC → NAc neurons may be a corollary signal of motor action or indeed serve a causal role in appetitive behaviors (58). To enable high-resolution behavioral

tracking, we monitored mouse orofacial movements using a 310 high-speed near-infrared camera (Figure 6A-C). To test for a 311 causal role of excitatory dHPC→NAc projections in appetitive 312 behaviors, we injected animals with either CaMKIIa-driven ChR2 or EYFP into dHPC and implanted light fibers in the NAc (n = 4/3 mice; Figure 6D-F). After habituating mice to 315 run on the treadmill and receive rewards upon licking on the lick spout, mice were given 5 mW of 473 nm 20 Hz (5 ms duration) pulsed laser light for up to 10 seconds upon entry into a hidden light stimulation zone.

We found that, shortly after stimulation onset, ChR2- 320 expressing mice reliably showed increased mouth movement 321 for up to two seconds after stimulation, while we observed 322 no effects in mice expressing EYFP ($p_{ChR2} = 0.0099, p_{EYFP} = 323$ 0.617, paired *t*-tests; Figure 6G-H). In line with this, we also 324 found a significant deceleration of running on the treadmill 325 upon light delivery in ChR2 animals but not EYFP animals 326 $(p_{ChR2} = 0.0381, p_{EYFP} = 0.334, paired t-tests; Figure 6J-K)$. 327 These findings support the idea that dHPC→NAc projections may not simply represent task-related non-spatial behavioral features but may in fact represent a driving force in the generation of task-relevant appetitive behavior.

289

291

293

294

295

296

297

300

301

302

303

305

307

319

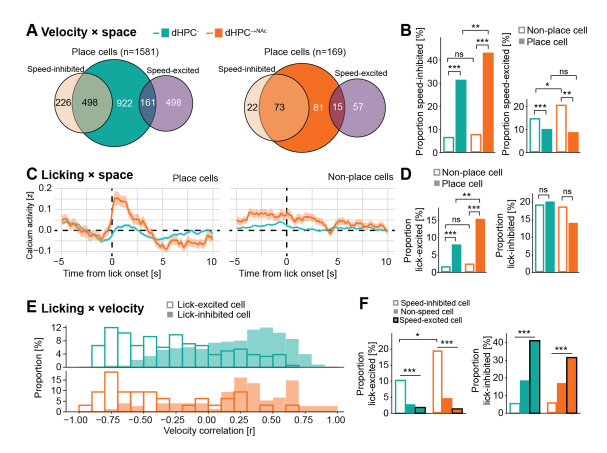


Figure 7. Enhanced conjunctive coding of space, velocity and licking in NAc-projecting neurons. (A and B) Conjunctive coding of space and velocity. (A) Venn diagrams of dHPC" (left) and dHPC "NAc" (right) place cells and their overlaps with negatively (ocher) and positively (purple) tuned speed cells. Numbers shown refer to absolute numbers of neurons classified. (B) Proportions of dHPC⁻ (green) and dHPC^{-NAc} (red) place cells (fill) and non-place cells (no fill) that are also significantly speedmodulated. A larger proportion of place cells is speed-inhibited compared to non-place cells (dHPC⁻: χ²(1, 4928) = 522.71; dHPC^{-NAc}: χ²(1, 444) = 75.02). dHPC^{-NAc} place cells also have a higher proportion of speed-inhibited cells than dHPC⁻ place cells ($\chi^2(1, 1750) = 8.977$). Speed-excited neurons are overrepresented in non-place cells compared to place cells (dHPC⁻: $\chi^2(1, 4928) = 20.034$; dHPC^{-NAc}: $\chi^2(1, 444) = 9.966$). dHPC^{-NAc} non-place cells also have a higher proportion of speed-excited cells than dHPC⁻ non-place cells ($\chi^2(1, 3622) = 6.255$). (C and D) Conjunctive coding of space and licking. (C) Event-triggered average calcium traces for dHPC⁻ (green) and dHPC $^{-NAc}$ (red) place cells (left) and non-place cells (right). (D) Proportions of lick-excited neurons are significantly enriched in dHPC $^{-}$ ($\chi^2(1, 4928) = 114.515$) and dHPC^{→NAc} (χ²(1, 444) = 23.248) place cells compared to non-place cells. The proportion of dHPC^{→NAc} lick-excited place cells is also higher than the proportion of dHPC lick-excited place cells ($\chi^2(1, 1750) = 9.442$); there is no difference between non-place cells ($\chi^2(1, 3622) = 0.488$, p = 0.485). There is no difference in the proportions of lick-inhibited place and non-place dHPC $^-$ and dHPC $^-$ and dHPC $^-$ neurons (χ^2 s, all $\rho > 0.05$). (E and F) Conjunctive coding of velocity and licking. (E) Histogram of lick-excited (no fill) and lick-inhibited (fill) cells' velocity correlations (green: dHPC⁻; red: dHPC⁻→NAc). Note the negative velocity correlations of lick-excited cells and positive velocity correlations for lick-inhibited cells. (F) Proportions of lick-excited (left) and lick-inhibited (right) cells among speed-inhibited (no fill), non-speed-modulated (fill) and speed-excited (fill, black stroke) dHPC⁻ (green) and dHPC⁻NAc (red) cells. Proportions of lick-excited cells are overrepresented in speed-inhibited cells (dHPC⁻: \(\chi^2(2, 4928) = 100.484; \) dHPC^{-NAc}: $\chi^2(2, 444) = 27.608$). Lick-excited neurons are further enriched in speed-inhibited dHPC $^{-NAc}$ neurons compared to dHPC $^{-}$ neurons ($\chi^2(1, 830) = 6.564$). Proportions of lick-inhibited cells are overrepresented in speed-excited cells (dHPC⁻: $\chi^2(2, 4928) = 290.832$; dHPC^{->NAc}: $\chi^2(2, 444) = 18.825$). All data are presented as mean ± SEM. ns: not significant, *p < 0.05, **p < 0.01, ***p < 0.001.

Enhanced conjunctive coding of space, velocity, and appetitive behaviors in dHPC - NAc neurons. We identified cells modulated by space, velocity, and appetitive licking. Previous studies suggested that individual hippocampal neurons do not necessarily exclusively code for one single feature but are instead able to conjunctively encode various environmental properties (9, 55, 59, 60). Such conjunctive coding may be particularly relevant for downstream linear decoders to select task-appropriate actions (61, 62). We thus investigated speed and lick modulation of projection-specific place cells and interactions between velocity and lick modulation.

We first analyzed speed coding in dHPC^{→NAc} place cells and compared it to the dHPC⁻ population (Figure 7A-B). We found about one third of hippocampal place cells were also speed-inhibited, in contrast to only about 7 % of non-place cells. This effect was particularly pronounced in dHPC→NAc

place cells (43 % vs. 31 %, p = 0.0027, χ^2). Conversely, 349 place cells were significantly less likely to be speed-excited 350 than non-place cells (10 % vs. 15 %, p < 0.001, χ^2), an effect that was again more pronounced in dHPC^{→NAc} neurons. 352 This shows that dHPC place cells, and in particular those projecting to NAc, are more likely to be speed-inhibited, and less 354 likely to be speed-excited.

We next analyzed lick modulation of place cells and, surprisingly, found that the previously observed lick-related increase in calcium activity (Figure 5B) was largely carried by 358 place cells and not by non-place cells (Figure 7C). This effect seems to be mostly carried by lick-excited neurons that 360 are significantly overrepresented in place cells compared to 361 non-place cells (8 % vs. 2 %, p < 0.001, χ^2), particularly 362 in dHPC $^{\to NAc}$ neurons (15 % vs. 3 %, p < 0.001, χ^2 ; Fig. 363 ure 7D). Lick-inhibited neurons, on the other hand, were 364 distributed equally between place and non-place dHPC⁻ and 365

336

337

338

340

342

343

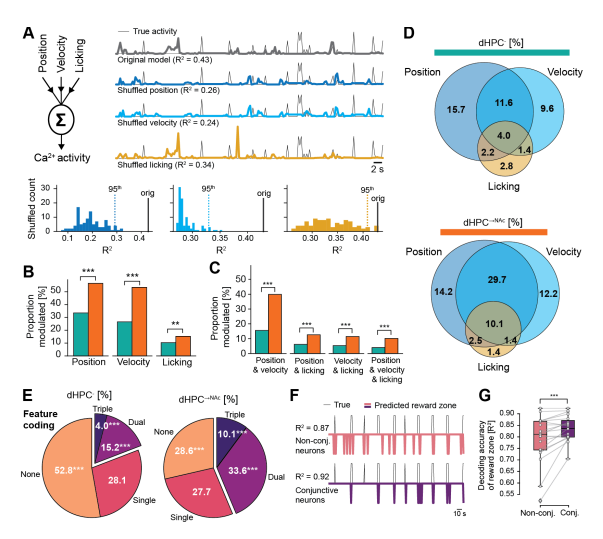


Figure 8. A generalized linear model confirms enhanced conjunctive coding in NAc-projecting neurons. (A) Schematic of the generalized linear model (GLM) using position, velocity, and appetitive licking data as predictors for each neuron's calcium activity (left). Example modelling approach for a triple conjunctive neuron (right). Upper traces show true calcium activity (downsampled and normalized) of test dataset (thin black line) as well as predictions of original model (thick gray) and one example each for each shuffled feature. Note the reductions in prediction quality when features are shuffled. Bottom histograms show R^2 distributions for 100 shuffled models for each feature. Dotted line represents value of 95th percentile, thick black line represents original model's R^2 . (B) Increased proportions of dHPC $^{-NAc}$ neurons modulated by position ($\chi^2(1, 5372) = 93.634$), velocity ($\chi^2(1, 5372) = 141.86$), and licking ($\chi^2(1, 5372) = 10.050$). (C) Increased proportions of conjunctive coding in dHPC $^{-NAc}$ neurons for position & velocity ($\chi^2(1, 5372) = 163.97$), position & licking ($\chi^2(1, 5372) = 26.029$), velocity & licking ($\chi^2(1, 5372) = 27.145$), and position & velocity & licking ($\chi^2(1, 5372) = 34.993$). (D) Venn diagrams showing overlap of neurons GLM-classified by position, velocity and licking in dHPC $^{-NAc}$ (bottom) neurons. (E) Proportions of n-feature coding neurons in dHPC $^{-}$ (left) and dHPC $^{-NAc}$ (right) populations. Non-coding neurons are overrepresented in dHPC $^{-}$ neurons ($\chi^2(1, 5372) = 0.0057$); dual-coding ($\chi^2(1, 5372) = 0.361$) and triple-coding ($\chi^2(1, 5372) = 30.447$) neurons are overrepresented in dHPC $^{-NAc}$ neurons. (F and G) A linear classifier to decode presence of reward zone. (F) Example true presence of reward zone (thin line) and decoder predictions based on non-conjunctive neurons (pink; top) and conjunctive-coding neurons (Wilcoxon's W(18) = 14.0). **p < 0.001. **p < 0.001. See also Figure S6.

dHPC $^{\to NAc}$ neurons. These findings show that a large majority of lick-excited neurons, especially in the dHPC $^{\to NAc}$ population, also code for spatial information. Finally, we investigated interactions between lick and velocity modulation. Comparing velocity correlations of lick-excited and lick-inhibited neurons, we observed a clear skew of lick-excited cells to have more negative velocity correlations and lick-inhibited cells to have more positive velocity correlations, visible in both dHPC $^-$ and dHPC $^{\to NAc}$ populations (Figure 7E). This results in significantly more speed-inhibited cells to be lick-excited compared to speed-excited cells (11 % vs. 2 %, p < 0.001, χ^2), an effect that was even more pronounced in dHPC $^{\to NAc}$ neurons (20 % vs. 1 %, p < 0.001, χ^2 ; Figure 7F). Conversely, speed-excited neurons were much more likely to be lick-inhibited than speed-inhibited neurons (40

% vs. 6 %, p < 0.001, χ^2). These results demonstrate that there is a strong inverse relationship between lick and velocity modulation. 381

One caveat of such conjunctive coding analyses is that in our behavioral task, trained mice often show highly stereotypical behavior, such that mice would mostly lick at one location where they would also slow down (see Figure 1C and Figure S1K-N). In light of this, conjunctive coding could be an epiphenomenon of collinear behavioral features. To account for this collinearity, we modelled the influence of three key behavioral features (space, velocity, and appetitive licking) on the activity of each neuron by building a generalized linear model for each neuron (GLM; Figure 8A). On average, we found that our models could explain close to 40% of the variance observed in our test datasets (Figure S6A),

367

369

370

371

372

374

with dHPC^{NAc} neurons showing increased feature importance for position and licking (Figure S6B). To determine significant contributions of the three behavioral features, we next built 3×100 models in which one of the behavioral features was randomly shuffled against time, and compared the variance explained to the original model (Figure 8A; see also ref (63)). Thus, if neural activity spuriously coincided with the activity of one behavioral feature that could be similarly explained by a largely collinear behavioral feature, the model's predictive performance should be mostly unaffected by this shuffling. The average drop in variance explained to the full model was significantly stronger in dHPC^{→NAc} neurons compared to dHPC- for position and velocity but not licking (Figure S6C). We classified significant modulation as behavioral features whose shuffling led to a reduction in variance explained in more than 95% of shuffled models. We found that cells thus encoding space, velocity, and licking were overrepresented in dHPC→NAc compared to dHPCneurons (space: p < 0.001; velocity: p < 0.001; licking: p= 0.0015; χ^2 ; Figure 8B). Importantly, we also found significantly increased proportions of conjunctive coding for all three feature combinations as well as triple-conjunctive neurons in dHPC→NAc neurons compared to dHPC- (all combinations p < 0.001, χ^2 ; Figure 8C-D). This results in a significantly higher proportion of conjunctive coding neurons in $dHPC^{\rightarrow NAc}$ neurons compared to $dHPC^{-}$ (44 % vs. 19 %, p $< 0.001, \chi^2$; Figure 8E). As conjunctive coding has been suggested to aid downstream linear decoders to select task-appropriate actions (55, 61), we wondered if this increased conjunctive code might allow linear decoders to identify the presence of the reward zone more correctly in our task. We thus trained an SVMbased linear classifier on each trial's odd/even laps' reward zones and tested the decoding accuracy on even/odd laps, based on conjunctive or randomly sample size-matched nonconjunctive coding neurons (Figure 8F). We found enhanced reward zone decoding accuracy for conjunctive compared to non-conjunctive coding neurons (p < 0.001, Wilcoxon's test; Figure 8G). Thus, our data suggest that the enhanced conjunctive coding observed in dHPC→NAc neurons allows NAc neurons to better identify the reward zone and guide taskappropriate appetitive behavior.

Discussion

While our understanding of the principal mechanisms by which internal and environmental features are processed by hippocampal circuits has greatly advanced (5, 6), the information content of its various output streams and their specific contributions to behavior remain largely unresolved. The NAc, with its proposed role as an integrator between limbic and motor systems (25) seems an ideal candidate to transform hippocampal mnemonic and contextual information into task-relevant behaviors (26), but evidence for such a role has remained sparse. One main finding was the identification of a larger proportion of place cells among the dHPC → NAc population that also en-

coded more spatial information and showed both enhanced

trial-to-trial reliability and in-place-field activity. Our data 452 confirm previous suggestions that the NAc receives spatial information from dHPC (26) and likely explains previously described spatial tuning of NAc neurons (29). The dHPC $^{\rightarrow NAc}$ population's observed enhanced spatial tuning and stronger 456 modulation of local cues parallels previously described differences between deep and superficial dHPC neurons (18). Indeed, vHPC→NAc projection neurons have been shown to accumulate mostly in deep layers (64), raising the possibility that previously observed anatomical-functional differences could at least partly be explained by differences in projection targets, thus adding another important layer of complexity to pyramidal cell heterogeneity (17).

463

484

Place cells in CA1 and subiculum have long been appreciated to overrepresent reward/goal zones (53, 65, 66) in a way 466 that correlates with behavioral performance (35, 52). This effect likely depends on the interaction of inputs from entorhinal cortex layer 3, dopaminergic inputs from VTA, and 469 hippocampal NMDA receptors (36, 52, 67). While previous studies suggested a role for a largely undefined dedicated 471 population of dHPC cells in this overrepresentation (65) and 472 differences have been observed along the radial dCA1 axis 473 (35), evidence for projection-specific reward zone representations has remained sparse. Although synchronous activity has been observed between dHPC and NAc during spatial reward learning tasks (38, 68, 69), in line with a central role of 477 NAc in reward learning (reviewed in ref (70)), our findings present direct evidence of increased reward zone overrepresentation by dHPC→NAc neurons. Interestingly, a previous 480 study (19) showed projection-specific ambiguity of reward 481 zone activity depending on vCA1→NAc collaterals. Future studies should investigate potential functional differences in 483 collaterals of dHPC projections (41).

Successful behavioral performance in our spatial reward 485 learning task depended on a decrease in velocity and increased appetitive lick activity as mice approached the hidden 487 reward zone. In line with previous work (8, 9, 12, 20), we 488 found a considerable proportion of speed-modulated dHPC neurons. Interestingly, NAc-projecting neurons were more likely to be speed-inhibited, suggesting an elevated role in 491 reward approach behaviors (see also ref (12)). Similarly, 492 we observed significant modulation of hippocampal neurons 493 during appetitive and consummatory licking (see also refs (12, 54)). This modulation was dichotomous: consummatory licking resulted in widespread suppression of dHPC activity, while appetitive licking led to enhanced activity in dHPC→NAc neurons. Activity of dHPC neurons, and especially those projecting to NAc, thus seems to mirror the activity within NAc: Inhibition of NAc D1R medium spiny neurons (MSNs) has been observed during lick behavior (71–73) 501 and was shown to be both necessary (71, 74) and sufficient 502 (71, 75) for consummatory licking. We found that the origin 503 of these signals may be localized upstream, similar to recent 504 findings of an inhibitory permissive drive for licking in vHPC projections to the NAc (72, 76) and that inhibiting these projections facilitates licking (72). This suggests a similar involvement of dHPC⁻ and vHPC-accumbens projections for 508

consummatory licking.

511

512

514

516

518

527

531

533

534

554

557

Contrasting this consumption-related inhibition, we found appetitive lick-related dHPC→NAc excitation and optogenetic induction of lick-related behaviors. This dichotomy likely reflects previously described differences in the NAc's role for consummatory and appetitive behaviors (57): vHPC→NAc activity was shown to increase during active food investigation (76) and around the time of lever pressing to obtain a reward (Iyer E, Muir J, Namuhoranye B, Bagot, R. Glutamatergic afferents to the nucleus accumbens integrate outcomes in reward-learning. Poster S01-181, FENS Forum 2022), while optogenetic stimulation of this projection facilitated nose poke and lever pressing behaviors (33, 77). Given that optogenetic stimulation of D2R MSNs in the NAc shell was also found to produce repetitive jaw movements (78), it is likely that D1R and D2R MSNs, both of which are targeted by HPC axons (34, 79, 80), might play dichotomous roles in appetitive and consummatory behaviors (see also refs (34, 81)). Future studies should aim to resolve the behavioral roles of specific NAc cell types targeted by HPC neurons.

In our analysis of dHPC→NAc coding properties, we uncovered more neurons individually coding for two or three aspects of space, velocity, and lick activity specifically in identified projection neurons. Such conjunctive coding, or mixed selectivity, has been previously found in hippocampal neurons to combine coding of space with direction, velocity, appetitive behaviors, behavioral tasks, future paths ("splitter cells"), context, or objects (9, 12, 13, 55, 66, 82). These conjunctions are hypothesized to combine "where" and "what" information to disambiguate different experiences occurring in the same location, thus building a "scaffold" for episodic memories that allows unambiguous later retrieval (14, 83, 84). Relatedly, theoretical and experimental studies demonstrated that such high-dimensional coding facilitates action selection by putative downstream linear decoders tasked with action selection and generating behavior (61, 62, 85), and was found to scale with task demands and performance (55, 62, 85–87). Indeed, our data show that conjunctive coding improved reward zone detection by a linear decoder, similar to a recent study in retrosplenial cortex that demonstrated enhanced decoding near a reward zone by conjunctive coding neurons (87). In line with previous findings of increased conjunctive coding in prefrontal cortex

NAc projection neurons (88) and elevated proportions of splitter cells in dSub \rightarrow NAc neurons (20), we thus propose that dHPC routes strongly conjunctive task-relevant information to facilitate NAc action selection.

The NAc has been described as a key node transforming motivational information from the limbic system into motor behaviors (25), but the information flow of specific projection neurons has remained elusive. Our data demonstrate that dHPC routes enhanced spatial information to the NAc that is conjunctively enriched by further spatial and non-spatial task-relevant features. We show that this conjunctive code improves linear decoding to guide downstream action selection, and that dHPC can drive the execution of appetitive motor behaviors via the basal ganglia, as early studies suggested

(89). Thus, our findings identify a direct role for the hippocampus in the generation of motor behaviors and build an important bridge in our understanding of how sensory and mnemonic processes guide behavioral action.

570

583

601

602

Materials and Methods

Animals. Experiments were performed in adult male and female mice. C57Bl/6 (n = 16) and Thy1-GCaMP6s (n = 6) mice (GP4.3; The Jackson Laboratory, Bar Harbor, USA) were bred under specific pathogen-free conditions. Heterozygous mice were group-housed with 12 hours reversed dark light cycle at 21°C and ad libitum food/water access until mice had recovered from surgery. Experiments were performed during the dark phase. All experiments were performed according to the Directive of the European Communities Parliament and Council on the protection of animals used for scientific purposes (2010/63/EU) and were approved by the animal care committee of North Rhine-Westphalia, Germany.

Viral vectors. To co-express the static red fluorophore 584 mCherry in Thy1-GcaMP6s animals, we injected retrograde (rg) serotype (90) adeno-associated virus (AAV) under control of the polyglycokinase (pgk) promoter (AAVrg-pgk-Cre, Catalog #24593-AAVrg, titer: 1×10^{13} viral genomes (vg) ml⁻¹, Addgene, Watertown, USA) in the NAc to enter axons and induce Cre expression in NAc-projecting neurons. Then, we used double-floxed inverse open reading frame (DIO) Cre-dependent mCherry under the human synapsin promoter in dHPC (AAV5-hSyn-DIO-mCherry, titer: 1.1×10^{13} vg/ml, Catalog #50459-AAV5, Addgene, Watertown, USA). For optogenetic experiments, we used unfloxed ChR2 (91) or control EYFP in excitatory neurons under the CaMKII promoter in the dHPC of C57Bl/6 mice (AAV2-CaMKIIhChR2(H134R)-EYFP-WPRE, titer: 4×10¹² transducing units (TU), UNC #AV4381E, Vector Core at the University of North Carolina, Chapel Hill, USA; rAAV2-CaMKII-EYFP, titer: 4.3×10¹² TU, UNC #AV6650, Vector Core at the University of North Carolina, Chapel Hill, USA).

Stereotactic virus injections. For stereotactic injection of 603 AAVs, mice were anesthetized with an intraperitoneal (i.p.) injection of a mixture of ketamine (0.13 mg/g) and xylazine (0.01 mg/g). Mice were head-fixed using a head holder (MA-6N, Narishige, Tokyo, Japan), placed into a motorized stereotactic frame (LuigsNeumann, Ratingen, Germany) and warmed by a self-regulating heat pad (Fine Science Tools, Heidelberg, Germany). After skin incision (5 mm) and removal of the periosteum, placement of the injection was determined in relation to bregma. A 0.5 mm wide hole was drilled through the skull (Ideal micro drill, World Precision Instruments, Berlin, Germany). Stereotactic coordinates were taken from Franklin and Paxinos, 2008 (The Mouse Brain in Stereotaxic Coordinates, Third Edition, Academic Press). To induce retrograde labelling of NAc-projecting neurons in dHPC, 2× 500 nl of AAVrg-pgk-Cre were injected into the ipsilateral (right) nucleus accumbens (-1.3 mm 619 anterior-posterior, -1.0 mm lateral, 5.0 and 4.3 mm ventral, relative to Bregma) at 100 nl/min, using a UltraMicroPump, 34G cannula and Hamilton syringe (World Precision Instruments, Berlin, Germany). To label projection neurons in red, 200 nl of AAV5-hSyn-DIO-mCherry were injected into the ipsilateral dorsal hippocampus (3.38 mm anterior-posterior, -2.5 mm lateral, 1.8 mm ventral, relative to Bregma, at a 10° angle). For optogenetic experiments, 200 nl of either AAV2-CaMKII-hChR2(H134R)-EYFP-WPRE or rAAV2-CaMKII-EYFP were injected each bilaterally into dorsal hippocampus (3.38 mm anterior-posterior, \pm 2.5 mm lateral, 1.8 mm ventral, relative to Bregma, at a 10° angle). After surgery, buprenorphine (0.05 mg/kg) was administered thrice daily for 3 consecutive days. Implant surgery followed two weeks after AAV injection.

Cranial window surgery. For awake hippocampal Ca2+ imaging, a window was surgically implanted over the right dorsal hippocampus, one week after virus injections. The hippocampal window was assembled from a 1.7 mm long stainless-steel cannula (3 mm outer diameter) and a round cover slip (3 mm diameter). The coverslip was glued to the end of the cannula using UV curable adhesive (NOA81, Thorlabs, Dachau/Munich, Germany). Mice were anesthetized and prepared for surgery as described above. The skin over the parietal skull and the periosteum were removed and wound edges sealed with Vetbond tissue adhesive (3M Animal Care Products, St Paul, USA). Skull surface was roughened by briefly applying gel etchant phosphoric acid (37.5 %; Kerr Dental, Scafati, Italy), carefully washing the skull surface and applying two-component dental adhesive (OptiBond FL, Kerr Dental, Scafati, Italy). A circular piece of the skull (3 mm in diameter) centered over the injection was carefully cut out using a sharp drill (Ideal micro drill, World Precision Instruments, Berlin, Germany). The dura was removed with forceps and mild vacuum suction was used to slowly remove the cortex within the craniotomy. Blood and aspirated tissue were washed out using a constant flow of artificial cerebrospinal fluid (ACSF). Intracranial aspiration was continued until the external capsule was exposed. The external capsule remained intact. The hippocampal window was manually inserted. During continuous perfusion with ACSF, the hippocampal window was lowered until sealing onto the external capsule. The hippocampal window was fixated, and the craniotomy sealed using UV curable dental cement (Gradia Direct Flo, GC Corporation, Tokyo, Japan). An angular metal bar (Luigs & Neumann, Ratingen, Germany) for head fixation was placed paramedian on the skull. After surgery, buprenorphine (0.05 mg/kg) was administered thrice daily for 3 days.

Behavioral task. 3-4 days after recovery from surgery, mice were provided with spinning wheels in their cages and were placed under a reverse light/dark cycle. One week after surgery, food restriction and two-week habituation schedules were initiated. Mice were food-restricted by providing about 80 % of their measured daily food pellets every 24 h. In the course of this, mice lost about 10-20 % of their origi-

nal weight before the start of training. Habituation consisted 676 of progressive exposure of mice to manual handling by the 677 experimenter, obtaining milk rewards through a metal cannula, gentle manual head fixation, the treadmill apparatus, and, finally, head-fixed running on an unmarked treadmill belt with random rewards provided through a metal cannula 681 lick spout after licking on it. The self-propelled treadmill (Luigs & Neumann) consisted of three rotating cylinders covered by a 7 cm wide and 360 cm long textile belt (Luigs & 684 Neumann) including six differently textured zones: horizontal and vertical glue stripes, glue dots, Velcro dots, vertical 686 tape stripes and upright nylon spikes. The reward zone for imaging experiments was 30 cm long and was placed between the end of the horizontal glue stripes and the beginning of the vertical tape stripes. The position of the mouse was recorded via an optical sensor (Luigs & Neumann) measuring the rotation of the treadmill cylinder underneath the 692 mouse. Lick signals were measured by an analog piezo sensor; one full belt rotation was measured by an optical infrared sensor. All signals were collected at 10 kHz by an I/O board 695 (USB-6212 BNC, National Instruments, Austin, USA) and 696 recorded using custom-written Python software. After two 697 weeks of habituation, mice were placed daily for 15 minutes on a cued (see above) treadmill belt on which they needed to lick on the metal spout in this hidden reward zone to obtain 700 a liquid reward in the form of condensed milk. Milk was 701 released by a miniature peristaltic pump (RP-Q1-S-P45A-DC3V, Takasago Fluidic Systems, Nagoya, Japan) that was 703 triggered by custom-written Python software via an I/O board (USB-6212 BNC, National Instruments, Austin, USA). After five days of training, calcium activity was recorded while mice performed the learned task.

Infrared camera behavioral tracking. Headfixed mouse 708 behavior was continuously monitored by simultaneously using two monochrome CCD cameras (Basler acA 780-75gm) 710 positioned at approximately 15 cm from the mouse. To capture face dynamics, we used a high-resolution zoom lens (50 712 mm FL, Thorlabs MVL50TM23); for body dynamics, we 713 used a wide-angle lens (12 mm FL, Edmund Optics #33-303). Infrared illumination was provided via two 850 nm 715 LED arrays (Thorlabs LIU850A), and cameras were outfitted 716 with 850/40 nm bandpass filters (Thorlabs FB850-40). Both 717 cameras' positions were aligned for each mouse before the 718 start of recordings. Camera images were acquired at 25 or 719 75 Hz with 782×582 pixels using pylon Camera Software 720 Suite (Basler), each frame triggered by TTL pulses from the recording software. Files were saved in compressed MP4 format before further processing.

Two-photon calcium imaging. Two-photon imaging was 724 performed using an upright LaVision BioTec (Bielefeld, Germany) TrimScope II resonant scanning microscope, equipped 726 with a Ti:sapphire excitation laser (Chameleon Ultra II, 727 Coherent, Santa Clara, USA) operated at 920 nm for GCaMP6s fluorescence excitation, a second 1045 nm fixedwavelength laser (Spectra Physics HighQ-2-IR, Newport 730 Corp., Irvine, USA) for mCherry fluorescence excitation, 731

706

722

and a 16x objective (N16XLWD-PF, Nikon, Düsseldorf, Germany). GCaMP6s fluorescence emission was isolated using a band-pass filter (525/40, Semrock, Rochester, USA) and detected using a GaAsP PMT (H7422-40, Hamamatsu, Herrsching am Ammersee, Germany). mCherry fluorescence emission was isolated using a band-pass filter (590/40, Semrock, Rochester, USA) and detected using a GaAsP PMT (H7422-40, Hamamatsu, Herrsching am Ammersee, Germany). Both lasers were aligned such that they excited the same focal plane. Imspector software (LaVision BioTec) was used for microscope control and image acquisition. Image series (2 channels, 1024×1024 pixels or 512×512 pixels, 350– 850 µm square field of view) were acquired at 15.2 Hz or 30.5 Hz, respectively. Frame capture signals were recorded by an I/O board (USB-6212 BNC, National Instruments, Austin, USA) that allowed subsequent data synchronization.

734

735

736

737

739

751

758

760

762

764

766

770

773

777

781

Optogenetic stimulation. For optogenetic experiments, directly following virus injections, the skin over the parietal skull and the periosteum were removed and wound edges sealed with Vetbond tissue adhesive (3M Animal Care Products, St Paul, USA). Skull surface was roughened by briefly applying gel etchant phosphoric acid (37.5 %; Kerr Dental, Scafati, Italy), carefully washing the skull surface and applying two-component dental adhesive (OptiBond FL, Kerr Dental, Scafati, Italy). Two small 0.5 mm wide holes were drilled through the skull (Ideal micro drill, World Precision Instruments, Berlin, Germany) and a two-ferrules fiber-optic cannula (TFC_200/230-0.37_5.5mm_TS2.0_FLT, Doric Lenses, Quebec, Canada) was implanted bilaterally on top of NAc (coordinates: +1.3 mm anterior-posterior, +/- 1 mm lateral, -4.9 mm ventral from brain surface, relative to bregma). The craniotomy was then sealed using UV curable dental cement (Gradia Direct Flo, GC Corporation, Tokyo, Japan). An angular metal bar (Luigs & Neumann, Ratingen, Germany) for head fixation was placed paramedian on the skull. After surgery, buprenorphine (0.05 mg/kg) was administered thrice daily for 3 days. After one week of recovery, the food restriction and a two-week habituation schedules were started. After successful habituation, headfixed mice received light stimulation via a fiber-coupled 473 nm diode laser (LuxX 473-80, Omicron-Laserage) through branching fiberoptic patchcord (BFP(2)_200/220/900-.37_2m_FCM*-2xZF1.25; Doric Lenses, Quebec, Canada), at 5 mW light intensity, measured at the fiber output. Customwritten Python scripts were used to send a trigger signal to an analog isolated pulse generator (Model 2100, A-M Systems, Sequim, USA) once animals entered a hidden optogenetic stimulation zone. The pulse generator produced 5 ms long pulses at 20 Hz for a maximum duration of 10 seconds or until the mouse passed beyond the stimulation zone.

Behavioral analysis. Analog signals for position, licking, reward pump, full rotation of belt, and digital signals from infrared camera triggers, two-photon scanning triggers, and optogenetic stimulation triggers were collected at 10 kHz and saved as TDMS files. Data processing and analysis were performed in Python. Position signals in cm were reconstructed

using the optical sensor detecting a full rotation of the treadmill belt. Velocity was calculated using a Kalman filter applied to the position signal. Behavioral data were then downsampled to either match the sampling rate of camera tracking (25/75 Hz) for training data or to match the sampling rate of two-photon imaging (15/30 Hz), by using each time window's arithmetic mean (velocity), median (position, lap number, camera/scanner trigger), maximum (reward pump, optogenetic trigger), or standard deviation (licking). Discrete lick events were detected using Scipy's find_peaks function with a minimum temporal distance of 0.33 s and a dynamic minimum height threshold that was individually determined by inspecting the synchronized infrared camera video. Lick 800 bouts were classified as lick events that were <2 seconds apart from one another. Appetitive lick onsets were defined as lick bout onsets that were preceded by at least 3 seconds absence of lick events. Consummatory licking onset was defined as the first lick event between 0.5 seconds up to 5 seconds after reward pump trigger. Rewarded laps were defined as laps in 806 which the reward pump was triggered by the animal's licking. Relative licking refers to the cumulative analog lick signal per position bin or reward zone. Successful laps were defined as laps in which mice showed at least 50 % of relative licking in reward and anticipation zones and received a reward. High success trials were defined as trials consisting of at least 50 % of successful laps.

Calcium signal processing. Two-photon imaging data was 814 processed using custom-written software in Python, largely based on CaImAn (v1.6.2; (44)). Green and red channel 16-bit TIFF stack files (512×512 or 1024×1024 pixels times 9,000 or 18,000 frames) were first resampled to 1 px/µm before motion-correcting the red static channel using NormCorre piecewise rigid (parameters: max shifts = 40, num_frames_split = 2000, overlaps = 46, splits_els = 4, strides = 255). The motion-corrected red channel image was then averaged over t and used for later identification of mCherry-positive components. Motion correction vectors were then applied on the dynamic green channel before using constrained non-negative matrix factorization (CNMF) for cell segmentation. Resulting traces were detrended and 827 deconvolved before filtering spatiotemporal components using quality criteria followed by identity- and behavior-blind manual curation based on visual inspection of spatial and temporal footprints and the quality of deconvolution. To identify mCherry-positive components, resulting spatial footprints were overlaid over the red channel average and a dynamic threshold applied to visually match the optimal signal discovery.

Place field analysis. Continuous belt positions were binned 836 into 45 bins of 8 cm length. For calcium signals, deconvolved events were used to avoid differential effects of GCaMP6s calcium signal tails at different animal velocities across space. Only time points with a velocity >2 cm/s were considered for spatial tuning analysis. Spatial information 841

813

(SI) was calculated as follows (47):

$$SI = \sum_{i} o_i a_i \log_2(a_i/\bar{a}) \tag{1}$$

where i denotes the i-th spatial bin, o_i is the animal's occupancy at spatial bin i, a_i is the mean of deconvolved events at spatial bin i, and \bar{a} is the overall mean calcium activity. Place cells were defined as cells whose SI was higher than the 95th percentile of 1000x randomly position-shuffled SI values (see ref (65)). Each shuffle consisted of randomly circularly shifting the activity time course by at least 500 frames, segmenting the activity in n blocks that were randomly permuted. n was automatically chosen for each session to match approximately twice the number of laps run to account for running differences between animals. The resulting shuffled trace was then used to calculate SI as above. Spatial information rate in bits per second was calculated by multiplying SI with the average firing rate during times of movement (>2 cm/s). Firing rate was defined as the number of deconvolved calcium events >0 per second. Sparsity was calculated according to ref (48), using the same denotation as above:

Sparsity =
$$\frac{(\sum o_i a_i)^2}{(\sum o_i a_i^2)}$$
 (2)

Place field reliability was calculated as the fraction of laps in which the maximum spatially binned deconvolved calcium activity occurred within the cell's place field. Δ in-out place field activity was defined as the average deconvolved calcium event activity within a cell's place field subtracted by that activity outside the place field. Place fields were obtained by first averaging deconvolved calcium event activity at velocities >2 cm/s per 45 spatial bins, replicating the resulting trace by a factor of two to account for circularity of the belt, and applying a Savitzky-Golay filter (using Scipy's savgol function with a window of 5 frames at the second order polynomial) to account for skewed place field activity. The resulting filtered spatial calcium activity was then searched for peaks using Scipy's find peaks function (width: 1 frame; prominence: 1.5 standard deviations; relative height: 0.8), with the most prominent peak used as primary place field and left/right interpolated intersection points as place field beginning/end. Place fields that would stretch into the replicated positions were translated back to the original 45 spatial bins. Place field boundary ratio was calculated as segmenting position bins into texture transition or middle zones, depending on their proximity to a texture transition, and calculating the ratio of place field beginnings/ends in a transition zone over those in a middle zone. To compare this ratio with a shuffled distribution, all place fields underwent 1000 iterations of randomly circularly shifting both beginnings and ends simultaneously and saving the resulting ratio both cell populations after each iteration. Place field centers were defined as the center of mass (COM) of significantly spatially modulated neurons (see above). For this, calcium activity at velocities >2 cm/s per 45 spatial bins was transformed to polar coordinates with q as position bin and r as average deconvolved calcium event activity at that position (see also ref (65)). The

two-dimensional centroid was calculated, and the resulting 893 angle transformed back to belt position to yield the place field center. "Place fields near reward zone" refers to place cells with COMs in either the anticipation zone (starting 30 cm 896 before reward zone) or reward zone.

897

936

937

Decoding of reward and anticipation zones. For decoding of reward and anticipation zones, all non-zero deconvolved calcium event activity was normalized into quantiles. Position and calcium activity were only considered from time 901 points with velocity >2 cm/s. Data were downsampled to 902 1 Hz by accumulating calcium activity and averaging position data. Positions in the 30 cm reward zone or the preceding 30 cm anticipation zone were binarized (1 inside zone, 0 outside zone). Models were trained using calcium data of either (non-)projecting or (non-)conjunctive neurons and 100 randomly sample size-matched neurons of the respectively larger population, ultimately using the average R^2 of the random sampling procedure. Neurons with fewer than 10 910 time bins of non-zero calcium activity in train/test datasets 911 were excluded from training/testing of the model. A linear support vector machine (SVM) classifier (Sklearn's linear_model.SGDClassifier function; loss: hinge, penalty: L2) was cross-validated by training on even laps and testing on 915 odd laps, and vice versa, using the average of the resulting 916 R^2 . For models comparing (non-)projection neurons only imaging sessions with >10 projection neurons were included. 918

Velocity and lick coding. Velocity modulation was assessed by linear regression of each neuron's average deconvolved calcium event activity against 1 cm/s velocity bins from 2 up to 30 cm/s using Scipy's stats.linregress function (see also ref (56)). Cells were considered to be significantly velocity modulated at P_{adjusted} < 0.05 after correcting for false discovery rate by using Benjamini/Hochberg correction. Significantly velocity modulated cells with positive slopes were termed "speed-excited" and those with negative slopes "speed-inhibited". To assess population-level appetitive/consummatory lick modulation, average deconvolved calcium activity before and after each lick event (see above for definition) were compared within the respectively described time windows. For single-cell level appetitive lick modulation analysis, each neuron's average pre/post deconvolved calcium activity was compared across events using Wilcoxon's non-parametric paired test. Cells were classified as "licking-excited" if they were significantly modulated and showed a positive difference, or "licking-inhibited" if they showed a significant negative difference.

Generalized linear model. To determine cellular coding of space, velocity, and licking, we used a generalized linear 940 model to predict each neuron's calcium activity. We normalized calcium activity by dividing each neuron's decon- 942 volved calcium events by their signal-to-noise ratio and the 943 resulting standard deviation. Normalized calcium activity was then smoothed using a Gaussian window of 0.5 seconds (SD = 2) and temporally downsampled by averaging 946 to 3 Hz. Finally, downsampled non-zero calcium data were

normalized into quintiles. The feature matrix consisted of normalized position, velocity, and lick data. Position data were binned into 45 equally sized spatial bins (8 cm each) and median-averaged to 3 Hz, used as factors. Velocity data were normalized by dividing data by its mean and averaging to 3 Hz. Lick bouts were calculated as described above. Appetitive lick bouts were defined as all lick bouts occurring up to 5 seconds before and at least 5 seconds after reward dispensation. Appetitive lick bouts were downsampled by median-averaging to 3 Hz. Both downsampled velocity and downsampled licking (appetitive lick bouts) were replicated with time shifts spanning 3 seconds before to 3 seconds after original timing at 3 Hz to account for anticipatory or delayed calcium activity. Generalized linear Poisson models were trained, validated and tested using Python's H2O library (H2OGeneralizedLinearEstimator; lambda: 0) with train/validation/test ratio of 0.8/0.1/0.1. The resulting R^2 score based on the test dataset was saved and compared to that of randomly shuffled feature models. For each predictor group (position, velocity, licking), 100 shuffled models were generated by circularly shifting the respective group's data randomly between 200 and 700 time points, saving each model's R^2 score. Significant predictors for each neuron's calcium activity were defined as those whose random shuffling procedure resulted in reductions in \mathbb{R}^2 for more than 95 % of cases. This means that at least 96 of the 100 models with one randomly shuffled predictor group had to have decreased R^2 values compared to the full model, in order for this predictor to be considered significant. Conjunctive-coding neurons were defined as those with at least two significant predictor

950

951

952

953

954

957

959

961

963

965

967

971

978

980

982

984

993

997

Infrared camera recordings analysis. Markerless pose estimation (Deep Graph Pose (92) and DeepLabCut (93)) was used to detect facial and body movements for both types of videos. For this, a deep neural network was trained to automatically discriminate 15 markers for videos of the body (paw, tail and head segments) and 13 markers for videos of the face (6 for pupil, eye, nose, mouth, etc). The network was trained on a large variety of lighting conditions and angles until it reached satisfactory performance. Mouth facial regions of interest (ROIs) were automatically segmented using rideo-averaged marker points of nose tip, the eye's tear duct and the mouth as stable landmarks. Mouth motion energy was calculated as the ROI's average pixel-by-pixel intensity differences from t_{-1} to t_0 , and z-scoring the resulting activ-

Statistical analysis. Statistical analysis was performed using Python's pingouin library. Statistical tests are indicated in the figure legends and text. To evaluate statistical significance, data from Figures 1D and 3F was subjected to oneor two-way ANOVAs. Data in Figures 2D, 4D, 4H, 5H-I, 7B, 7D, 7F, 8B-C, and 8E was subjected to chi-square tests. Data in Figures 2E-H was subjected to Welch's t-tests. Data in Figures 3C-D was subjected to bootstrapping analysis (see above) and chi-square tests. Data in Figure 3G and 8G was subjected to Wilcoxon's test for nonparametric data. Data in Figure 5C was subjected to a repeated-measures ANOVA fol- 1004 lowed by post-hoc t-tests with Bonferroni correction. Data 1005 in Figures 6I and 6K was subjected Student's paired t-test. 1006 For all analyses data are presented as mean ± SEM, and the 1007 threshold for significance was at p < 0.05.

ACKNOWLEDGEMENTS

The authors would like to thank Janelle Pakan, Hugo Spiers, Magdalena Sauvage, Marlene Bartos, and Thomas Mrsic-Flögel as well as the rest of the Cellular Neuroscience Department for their insightful comments and discussions while preparing this manuscript. In addition, the authors would like to thank Falko Fuhrmann, Pavol Bauer, and Rüdiger Geis for their help with the treadmill setup, cranial surgery, and two-photon imaging. The resources for all experiments were provided by the German Center for Neurodegenerative Diseases (DZNE) in Bonn and the Leibniz Institute for Neurobiology (LIN) in Magdeburg. The authors received funding from the German Research Foundation (DFG) SFB grant 1089 (P.M., S.R.) and ERC Consolidator grant Sub-D-Code (O.B., S.R.).

1009

1011

1012

1015

1016

1017

1018

1019

1020

1021

1024

1025

1026

1027

1029

1030

1032

1033

1036

1038

1048

1050

1052

1053

1054

1055

1056

1057

1058

1059

1060

1061

1062

1063

1064

1065

1066

1067

1068

1069

1070

1071

1072

1073

1074

1075

1076

1079

AUTHOR CONTRIBUTIONS

O.B. and S.R. conceived and designed experiments. O.B. performed and analyzed imaging experiments. O.B. and P.M performed optogenetic experiments, O.B. analyzed these. O.B. wrote the manuscript with help from S.R. and P.M.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

CODE AND DATA AVAILABILITY

Code and data will be made available upon final publication

Bibliography

- Natalie Biderman, Akram Bakkour, and Daphna Shohamy. What Are Memories For? The Hippocampus Bridges Past Experience with Future Decisions. Trends in Cognitive Sciences, 24(7):542-556, 2020. ISSN 1364-6613. doi: 10.1016/j.tics.2020.04.004.
- T. V. P. Bliss and T. Lømo. Long-lasting potentiation of synaptic transmission in the dentate area of the anaesthetized rabbit following stimulation of the perforant path. The Journal of Physiology, 232(2):331-356, 1973. ISSN 0022-3751. doi: 10.1113/jphysiol.1973.sp010273.
- William Beecher Scoville and Brenda Milner. LOSS OF RECENT MEMORY AFTER BILAT-ERAL HIPPOCAMPAL LESIONS. Journal of Neurology, Neurosurgery & Psychiatry, 20(1): 11, 1957. ISSN 0022-3050. doi: 10.1136/jnnp.20.1.11
- L.R. Squire and E.R. Kandel. Memory: From Mind to Molecules. Scientific American Library series. Roberts & Company, 2009. ISBN 9780981519418.
- Per Andersen, Richard Morris, David Amaral, Tim Bliss, and John O'Keefe. The hippocam pus book. Oxford university press, 2006
- J O'Keefe and L Nadel. The Hippocampus as a Cognitive Map, 1978
- PE Sharp and C Green. Spatial correlates of firing patterns of single cells in the subiculum of the freely moving rat. Journal of Neuroscience, 14(4):2339-2356, 1994. ISSN 0270-6474 doi: 10.1523/jneurosci.14-04-02339.1994
- Motosada Iwase, Takuma Kitanishi, and Kenji Mizuseki. Cell type, sub-region, and layerspecific speed representation in the hippocampal-entorhinal circuit. Scientific Reports, 10 (1):1407, 2020. doi: 10.1038/s41598-020-58194-1.
- B. L. McNaughton, C. A. Barnes, and J. O'Keefe. The contributions of position, direction, and velocity to single unit activity in the hippocampus of freely-moving rats. Experimental Brain Research, 52(1):41-49, 1983, ISSN 0014-4819, doi: 10.1007/bf00237147
- Edvard I. Moser, Emilio Kropff, and May-Britt Moser. Place Cells, Grid Cells, and the Brain's Spatial Representation System. Neuroscience, 31(1):69-89, 2008. ISSN 0147-006x. doi: 10.1146/annurev.neuro.31.061307.090723
- 11. Nils Nyberg, Éléonore Duvelle, Caswell Barry, and Hugo J. Spiers. Spatial goal coding in the hippocampal formation. Neuron, 2022. ISSN 0896-6273. doi: 10.1016/j.neuron.2021.12.012.
- SI Wiener, CA Paul, and H Eichenbaum. Spatial and behavioral correlates of hippocampal neuronal activity. The Journal of Neuroscience, 9(8):2737-2763, 1989. ISSN 0270-6474 doi: 10.1523/jneurosci.09-08-02737.1989.
- Emma R. Wood, Paul A. Dudchenko, R.Jonathan Robitsek, and Howard Eichenbaum, Hippocampal Neurons Encode Information about Different Types of Memory Episodes Occurring in the Same Location. Neuron, 27(3):623-633, 2000. ISSN 0896-6273. 10.1016/s0896-6273(00)00071-4.
- Timothy E.J. Behrens, Timothy H. Muller, James C.R. Whittington, Shirley Mark, Alon B. Baram, Kimberly L. Stachenfeld, and Zeb Kurth-Nelson. What Is a Cognitive Map? Organizing Knowledge for Flexible Behavior. Neuron, 100(2):490-509, 2018. ISSN 0896-6273. doi: 10.1016/i.neuron.2018.10.002.
- Kenneth D. Harris, Jozsef Csicsvari, Hajime Hirase, George Dragoi, and György Buzsáki. Organization of cell assemblies in the hippocampus. Nature, 424(6948):552-556, 2003. ISSN 0028-0836. doi: 10.1038/nature01834.
- Alessandro Treves and Edmund T. Rolls. Computational analysis of the role of the hippocampus in memory. Hippocampus, 4(3):374-391, 1994. ISSN 1098-1063. doi: 10.1002/hipo.450040319.
- Mark S. Cembrowski and Nelson Spruston. Heterogeneity within classical cell types is the rule: lessons from hippocampal pyramidal neurons. Nature Reviews Neuroscience, 20(4): 193-204 2019 ISSN 1471-003X doi: 10.1038/s41583-019-0125-5
- Ivan Soltesz and Attila Losonczy. CA1 pyramidal cell diversity enabling parallel information 1077 processing in the hippocampus. Nature Neuroscience, 21(4):484-493, 2018. ISSN 1097-1078 6256. doi: 10.1038/s41593-018-0118-0

- S. Ciocchi, J. Passecker, H. Malagon-Vina, N. Mikus, and T. Klausberger, Selective informa-1080 19. tion routing by ventral hippocampal CA1 projection neurons. Science, 348(6234):560-563, 1081 1082 2015. ISSN 0036-8075. doi: 10.1126/science.aaa3245.
- 1083 20. Takuma Kitanishi, Ryoko Umaba, and Kenji Mizuseki. Robust information routing by dorsal 1084 subiculum neurons. Science Advances, 7(11):eabf1913, 2021. ISSN 2375-2548. doi: 1085 10.1126/sciadv.abf1913.
- 1086 H.J. Groenewegen, E. Vermeulen-Van der Zee, A. te Kortschot, and M.P. Witter. Organiza-1087 tion of the projections from the subjculum to the ventral striatum in the rat. A study using anterograde transport of Phaseolus vulgaris leucoagglutinin. Neuroscience, 23(1):103-120, 1088 1089 1987. ISSN 0306-4522. doi: 10.1016/0306-4522(87)90275-2.
- 1090 A.E. Kelley and V.B. Domesick. The distribution of the projection from the hippocampal 1091 formation to the nucleus accumbens in the rat: An anterograde and retrograde-horseradish peroxidase study. Neuroscience, 7(10):2321-2335, 1982. ISSN 0306-4522. doi: 10.1016/ 1092 1093 0306-4522(82)90198-1.
- Stan B. Floresco. The Nucleus Accumbens: An Interface Between Cognition, Emotion, 1094 and Action. Annual Review of Psychology, 66(1):25-52, 2015. ISSN 0066-4308. doi: 1095 10.1146/annurev-psych-010213-115159. 1096
- 1097 Francesco Mannella, Kevin Gurney, and Gianluca Baldassarre. The nucleus accumbens as a nexus between values and goals in goal-directed behavior: a review and a new 1098 1099 hypothesis. Frontiers in Behavioral Neuroscience, 7:135, 2013. ISSN 1662-5153. doi: 1100 10.3389/fnbeh.2013.00135
- Gordon J. Mogenson, Douglas L. Jones, and Chi Yiu Yim. From motivation to action: Func-1101 1102 tional interface between the limbic system and the motor system. Progress in Neurobiology, 14(2-3):69-97, 1980. ISSN 0301-0082. doi: 10.1016/0301-0082(80)90018-0 1103
- 1104 C.M.A. Pennartz, R. Ito, P.F.M.J. Verschure, F.P. Battaglia, and T.W. Robbins. The hippocam-1105 pal-striatal axis in learning, prediction and goal-directed behavior. Trends in Neurosciences, 1106 34(10):548-559, 2011. ISSN 0166-2236. doi: 10.1016/j.tins.2011.08.001.
- 1107 Yukiori Goto and Patricio O'Donnell. Synchronous Activity in the Hippocampus and Nucleus 1108 Accumbens In Vivo. The Journal of Neuroscience, 21(4):RC131-RC131, 2001. ISSN 0270-1109 6474. doi: 10.1523/jneurosci.21-04-j0003.2001.
- Carien S. Lansink, Pieter M. Goltstein, Jan V. Lankelma, Bruce L. McNaughton, and Cyriel 1110 M. A. Pennartz. Hippocampus Leads Ventral Striatum in Replay of Place-Reward Infor-1111 mation. PLoS Biology, 7(8):e1000173, 2009. ISSN 1544-9173. doi: 10.1371/journal.pbio. 1112 1113
- A.M. Lavoie and S.J.Y. Mizumori. Spatial, movement- and reward-sensitive discharge by 1114 29. 1115 medial ventral striatum neurons of rats. Brain Research, 638(1-2):157-168, 1994. ISSN 0006-8993. doi: 10.1016/0006-8993(94)90645-9. 1116
- Geke E. Ploeger, Berry M. Spruijt, and Alexander R. Cools. Spatial Localization in the 1117 30. Morris Water Maze in Rats:: Acquisition Is Affected by Intra-Accumbens Injections of the 1118 Dopaminergic Antagonist Haloperidol. Behavioral Neuroscience, 108(5):927-934, 1994. 1119 ISSN 0735-7044. doi: 10.1037/0735-7044.108.5.927. 1120
- 1121 31. R Ito, T W Robbins, C M Pennartz, and B J Everitt. Functional Interaction between the 1122 Hippocampus and Nucleus Accumbens Shell Is Necessary for the Acquisition of Appetitive Spatial Context Conditioning. Journal of Neuroscience, 28(27):6950-6959, 2008. ISSN 1123 0270-6474. doi: 10.1523/jneurosci.1615-08.2008. 1124
- 32. Stéphanie Trouche, Vadim Koren, Natalie M. Doig, Tommas J. Ellender, Mohamady El-1125 Gaby, Vítor Lopes-dos Santos, Hayley M. Reeve, Pavel V. Perestenko, Farid N. Garas, 1126 Peter J. Magill, Andrew Sharott, and David Dupret. A Hippocampus-Accumbens Tripartite 1127 Neuronal Motif Guides Appetitive Memory in Space. Cell, 176(6):1393-1406.e16, 2019. 1128 1129 ISSN 0092-8674. doi: 10.1016/j.cell.2018.12.037.
- 1130 33. Jonathan P. Britt, Faiza Benaliouad, Ross A. McDevitt, Garret D. Stuber, Roy A. Wise, and 1131 Antonello Bonci. Synaptic and Behavioral Profile of Multiple Glutamatergic Inputs to the Nucleus Accumbens, Neuron, 76(4):790-803, 2012, ISSN 0896-6273, doi: 10.1016/i. 1132 1133 neuron, 2012, 09, 040.
- 1134 34. Tara A. LeGates, Mark D. Kvarta, Jessica R. Tooley, T. Chase Francis, Mary Kay Lobo, 1135 Meaghan C. Creed, and Scott M. Thompson. Reward behaviour is regulated by the strength 1136 of hippocampus-nucleus accumbens synapses. Nature, 564(7735):258-262, 2018. ISSN 1137 0028-0836. doi: 10.1038/s41586-018-0740-8.
- 1138 35 Nathan B. Danielson, Jeffrey D. Zaremba, Patrick Kaifosh, John Bowler, Max Ladow, and 1139 Attila Losonczy. Sublayer-Specific Coding Dynamics during Spatial Navigation and Learning 1140 in Hippocampal Area CA1. Neuron, 91(3):652-665, 2016. ISSN 0896-6273. doi: 10.1016/ i.neuron.2016.06.020. 1141
- 1142 Christine Grienberger and Jeffrey C. Magee. Entorhinal cortex directs learning-related 36. 1143 changes in CA1 representations. Nature, 611(7936):554-562, 2022. ISSN 0028-0836. 1144 doi: 10.1038/s41586-022-05378-6.
- Hod Dana, Tsai-Wen Chen, Amy Hu, Brenda C. Shields, Caiying Guo, Loren L. Looger, 1145 1146 Douglas S. Kim, and Karel Svoboda. Thy1-GCaMP6 Transgenic Mice for Neuronal Popula-1147 tion Imaging In Vivo. PLoS ONE, 9(9):e108697, 2014. doi: 10.1371/journal.pone.0108697.
- 1148 Lucas Sjulson, Adrien Peyrache, Andrea Cumpelik, Daniela Cassataro, and György Buzsáki. Cocaine Place Conditioning Strengthens Location-Specific Hippocampal Cou-1149 pling to the Nucleus Accumbens. Neuron, 98(5):926-934.e5, 2018. ISSN 0896-6273. doi: 1150 1151 10.1016/j.neuron.2018.04.015.
- E.T. Tabuchi, A.B. Mulder, and S.I. Wiener. Position and behavioral modulation of syn-1152 1153 chronization of hippocampal and accumbens neuronal discharges in freely moving rats Hippocampus, 10(6):717-728, 2000. ISSN 1050-9631. doi: 10.1002/1098-1063(2000)10: 1154 5<717::aid-hipo1009>3.0.co;2-3.
- 1156 Michael S. Bienkowski, Ian Bowman, Monica Y. Song, Lin Gou, Tyler Ard, Kaelan Cotter, 1157 Muye Zhu, Nora L. Benavidez, Seita Yamashita, Jaspar Abu-Jaber, Sana Azam, Darrick 1158 Lo, Nicholas N. Foster, Houri Hintiryan, and Hong-Wei Dong. Integration of gene expres-1159 sion and brain-wide connectivity reveals the multiscale organization of mouse hippocam-1160 pal networks. Nature Neuroscience, 21(11):1628-1643, 2018. ISSN 1097-6256. doi: 1161
- 1162 Mark S. Cembrowski, Matthew G. Phillips, Salvatore F. DiLisio, Brenda C. Shields, Johan 1163 Winnubst, Jayaram Chandrashekar, Erhan Bas, and Nelson Spruston. Dissociable Struc-1164 tural and Functional Hippocampal Outputs via Distinct Subiculum Cell Clas (5):1280-1292.e18, 2018. ISSN 0092-8674. doi: 10.1016/j.cell.2018.03.031. 1165

42. Daniel A Dombeck, Christopher D Harvey, Lin Tian, Loren L Looger, and David W Tank. Functional imaging of hippocampal place cells at cellular resolution during virtual navigation. 1167 Nature Neuroscience, 13(11):1433-1440, 2010, ISSN 1097-6256, doi: 10.1038/nn.2648

1166

1168

1170

1171

1172

1173

1174

1177

1178

1179

1181

1182

1183

1185

1186

1187

1189

1190

1191

1192

1193

1195

1197

1198

1199

1200

1201

1203

1204

1205

1206

1207

1208

1209

1211

1213

1215

1216

1217

1219

1220

1221

1222

1223

1224

1225

1226

1227

1228

1229

1230

1231

1232

1233

1234

1235

1236

1237

1238

1240

1241

1242

1244

1245

1246

- Falko Fuhrmann, Daniel Justus, Liudmila Sosulina, Hiroshi Kaneko, Tatjana Beutel, Detlef 1169 Friedrichs, Susanne Schoch, Martin Karl Schwarz, Martin Fuhrmann, and Stefan Remy. Locomotion, Theta Oscillations, and the Speed-Correlated Firing of Hippocampal Neurons Are Controlled by a Medial Septal Glutamatergic Circuit. Neuron, 86(5):1253-1264, 2015. ISSN 0896-6273. doi: 10.1016/j.neuron.2015.05.001.
- Andrea Giovannucci, Johannes Friedrich, Pat Gunn, Jérémie Kalfon, Brandon L Brown, Sue Ann Koay, Jiannis Taxidis, Farzaneh Najafi, Jeffrey L Gauthier, Pengcheng Zhou, 1175 Baljit S Khakh, David W Tank, Dmitri B Chklovskii, and Eftychios A Pnevmatikakis. CalmAn 1176 an open source tool for scalable calcium imaging data analysis. eLife, 8:e38173, 2019. doi: 10.7554/elife.38173
- P.A. Naber and M.P. Witter. Subicular efferents are organized mostly as parallel projections: A double-labeling, retrograde-tracing study in the rat. Journal of Comparative Neurol-1180 ogy, 393(3):284–297, 1998. ISSN 1096-9861. doi: 10.1002/(sici)1096-9861(19980413)393: 3<284::aid-cne2>3.0.co;2-y
- Johan Winnubst, Erhan Bas, Tiago A. Ferreira, Zhuhao Wu, Michael N. Economo, Patrick Edson, Ben J. Arthur, Christopher Bruns, Konrad Rokicki, David Schauder, Donald J. Ol- 1184 bris, Sean D. Murphy, David G. Ackerman, Cameron Arshadi, Perry Baldwin, Regina Blake, Ahmad Elsayed, Mashtura Hasan, Daniel Ramirez, Bruno Dos Santos, Monet Weldon, Amina Zafar, Joshua T. Dudman, Charles R. Gerfen, Adam W. Hantman, Wyatt Korff, Scott M. Sternson, Nelson Spruston, Karel Svoboda, and Jayaram Chandrashekar. Reconstruction of 1,000 Projection Neurons Reveals New Cell Types and Organization of Long-Range Connectivity in the Mouse Brain. Cell, 179(1):268-281.e13, 2019. ISSN 0092-8674. doi: 10.1016/j.cell.2019.07.042.
- William Skaggs, Bruce McNaughton, and Katalin Gothard. An Information-Theoretic Approach to Deciphering the Hippocampal Code. In Advances in Neural Information Processing Systems, volume 5. Morgan-Kaufmann, 1993.
- MW Jung, SI Wiener, and BL McNaughton. Comparison of spatial firing characteristics of units in dorsal and ventral hippocampus of the rat. The Journal of Neuroscience, 14(12): 1196 7347-7356, 1994, ISSN 0270-6474, doi: 10.1523/ineurosci,14-12-07347,1994
- Steven Poulter, Sang Ah Lee, James Dachtler, Thomas J. Wills, and Colin Lever. Vector trace cells in the subiculum of the hippocampal formation. Nature Neuroscience, pages 1-10, 2020. ISSN 1097-6256. doi: 10.1038/s41593-020-00761-w.
- Chia-Hsuan Wang, Joseph D. Monaco, and James J. Knierim. Hippocampal Place Cells Encode Local Surface-Texture Boundaries. Current Biology, 30(8):1397-1409.e7, 2020. 1202 ISSN 0960-9822. doi: 10.1016/j.cub.2020.01.083.
- Regina M. Carelli. The Nucleus Accumbens and Reward: Neurophysiological Investigations in Behaving Animals. Behavioral and Cognitive Neuroscience Reviews, 1(4):281-296, 2002. ISSN 1534-5823. doi: 10.1177/1534582302238338.
- David Dupret, Joseph O'Neill, Barty Pleydell-Bouverie, and Jozsef Csicsvari. The reorganization and reactivation of hippocampal maps predict spatial memory performance. Nature Neuroscience, 13(8):995-1002, 2010. ISSN 1097-6256. doi: 10.1038/nn.2599
- 53. Stig A Hollup, Sturla Molden, James G Donnett, May-Britt Moser, and Edvard I Moser. 1210 Accumulation of Hippocampal Place Fields at the Goal Location in an Annular Watermaze Task. The Journal of Neuroscience, 21(5):1635-1644, 2001. ISSN 0270-6474. doi: 10. 1212 1523/ineurosci,21-05-01635,2001.
- 54. Anh Son Ho, Etsuro Hori, Phuong Hong Thi Nguyen, Susumu Urakawa, Takashi Kondoh, 1214 Kunio Torii, Taketoshi Ono, and Hisao Nishijo. Hippocampal neuronal responses during signaled licking of gustatory stimuli in different contexts. Hippocampus, 21(5):502-519, 2011. ISSN 1050-9631. doi: 10.1002/hipo.20766
- Debora Ledergerber, Claudia Battistin, Jan Sigurd Blackstad, Richard J. Gardner, Menno P. 1218 Witter, May-Britt Moser, Yasser Roudi, and Edvard I. Moser, Task-dependent mixed selectivity in the subiculum. Cell Reports, 35(8):109175, 2021. ISSN 2211-1247. doi: 10.1016/i.celrep.2021.109175.
- Emilio Kropff, James E. Carmichael, May-Britt Moser, and Edvard I. Moser. Speed cells in the medial entorhinal cortex. Nature, 523(7561):419-424, 2015. ISSN 0028-0836. doi: 10 1038/nature14622
- Trevor W Robbins and Barry J Everitt. Neurobehavioural mechanisms of reward and motivation. Current Opinion in Neurobiology, 6(2):228-236, 1996. ISSN 0959-4388. doi: 10 1016/s0959-4388(96)80077-8
- Nick T.M. Robinson, Lucie A.L. Descamps, Lloyd E. Russell, Moritz O. Buchholz, Brendan A. Bicknell, Georgy K. Antonov, Joanna Y.N. Lau, Rebecca Nutbrown, Christoph Schmidt-Hieber, and Michael Häusser. Targeted Activation of Hippocampal Place Cells Drives Memory-Guided Spatial Behavior. Cell, 183(6):1586-1599.e10, 2020. ISSN 0092-8674. doi: 10.1016/j.cell.2020.09.061. first paper showing a causal role of place cells in behaviour.
- R W Komorowski, J R Manns, and H Eichenbaum. Robust Conjunctive Item-Place Coding by Hippocampal Neurons Parallels Learning What Happens Where. Journal of Neuroscience, 29(31):9918-9929, 2009. ISSN 0270-6474. doi: 10.1523/jneurosci.1378-09.2009.
- Xinyu Zhao, Ching-Lung Hsu, and Nelson Spruston. Rapid synaptic plasticity contributes to a learned conjunctive code of position and choice-related information in the hippocampus. Neuron, 110(1):96-108.e4, 2022. ISSN 0896-6273. doi: 10.1016/j.neuron.2021.10.003.
- Stefano Fusi, Earl K Miller, and Mattia Rigotti. Why neurons mix: high dimensionality for higher cognition. Current Opinion in Neurobiology, 37:66-74, 2016. ISSN 0959-4388. doi: 10.1016/j.conb.2016.01.010.
- Mattia Rigotti, Omri Barak, Melissa R. Warden, Xiao-Jing Wang, Nathaniel D. Daw, Earl K. Miller, and Stefano Fusi. The importance of mixed selectivity in complex cognitive tasks. Nature, 497(7451):585-590, 2013. ISSN 0028-0836. doi: 10.1038/nature12160
- Simon Musall, Matthew T. Kaufman, Ashley L. Juavinett, Steven Gluf, and Anne K. Churchland. Single-trial neural dynamics are dominated by richly varied movements. Nature Neu roscience, 22(10):1677-1686, 2019. ISSN 1097-6256. doi: 10.1038/s41593-019-0502-4.
- Teruhiro Okuyama, Takashi Kitamura, Dheeraj S. Roy, Shigeyoshi Itohara, and Susumu Tonegawa. Ventral CA1 neurons store social memory. Science, 353(6307):1536-1541, 1249 2016. ISSN 0036-8075. doi: 10.1126/science.aaf7003
- Jeffrey L. Gauthier and David W. Tank. A Dedicated Population for Reward Coding in the 1251

- Hippocampus, Neuron, 99(1):179-193,e7, 2018, ISSN 0896-6273, doi: 10.1016/j.neuron. 1252 2018.06.008. 1253
- 1254 66. EJ Markus, YL Qin, B Leonard, WE Skaggs, BL McNaughton, and CA Barnes, Interactions 1255 between location and task affect the spatial and directional firing of hippocampal neurons. 1256 The Journal of Neuroscience, 15(11):7079-7094, 1995. ISSN 0270-6474. doi: 10.1523/ 1257 ineurosci.15-11-07079.1995.
- 1258 Colin G McNamara, Álvaro Tejero-Cantero, Stéphanie Trouche, Natalia Campo-Urriza, and 1259 David Dupret, Dopaminergic neurons promote hippocampal reactivation and spatial mem-1260 ory persistence. Nature Neuroscience, 17(12):1658-1660, 2014. ISSN 1097-6256. doi: 1261 10 1038/nn 3843
- 1262 William E. DeCoteau, Catherine Thorn, Daniel J. Gibson, Richard Courtemanche, Partha 1263 Mitra, Yasuo Kubota, and Ann M. Graybiel. Learning-related coordination of striatal and hippocampal theta rhythms during acquisition of a procedural maze task. Proceedings of 1264 1265 the National Academy of Sciences, 104(13):5644-5649, 2007. ISSN 0027-8424. doi: 10. 1073/pnas.0700818104 1266
- 1267 Marielena Sosa, Hannah R. Joo, and Loren M. Frank. Dorsal and Ventral Hippocampal Sharp-Wave Ripples Activate Distinct Nucleus Accumbens Networks. Neuron, 105(4):725-1268 741.e8, 2020. ISSN 0896-6273. doi: 10.1016/j.neuron.2019.11.022. 1269
- Jeremy J. Day and Regina M. Carelli. The Nucleus Accumbens and Pavlovian Reward 1270 127 Learning. The Neuroscientist, 13(2):148-159, 2007. ISSN 1073-8584. doi: 10.1177/ 1272 1073858406295854
- Eoin C. O'Connor, Yves Kremer, Sandrine Lefort, Masaya Harada, Vincent Pascoli, Clément 1273 1274 Rohner, and Christian Lüscher. Accumbal D1R Neurons Projecting to Lateral Hypothalamus Authorize Feeding. Neuron, 88(3):553-564, 2015. ISSN 0896-6273. doi: 10.1016/j.neuron. 1275 1276
- Sean J. Reed, Christopher K. Lafferty, Jesse A. Mendoza, Angela K. Yang, Thomas J. 1277 1278 Davidson, Logan Grosenick, Karl Deisseroth, and Jonathan P. Britt. Coordinated Reductions in Excitatory Input to the Nucleus Accumbens Underlie Food Consumption. Neuron, 1279 1280 99(6):1260-1273.e4, 2018. ISSN 0896-6273. doi: 10.1016/j.neuron.2018.07.051
- 1281 S A Taha and H L Fields. Encoding of Palatability and Appetitive Behaviors by Distinct Neuronal Populations in the Nucleus Accumbens. Journal of Neuroscience, 25(5):1193-1282 1202, 2005, ISSN 0270-6474, doi: 10.1523/ineurosci.3975-04.2005 1283
- 74. Luis Prado, Jorge Luis-Islas, Osvaldo I. Sandoval, Liliana Puron, Moreno M. Gil, Alvaro 1284 1285 Luna, Mario A. Arias-García, Elvira Galarraga, Sidney A. Simon, and Ranier Gutierrez. Activation of Glutamatergic Fibers in the Anterior NAc Shell Modulates Reward Activity in 1286 1287 the aNAcSh, the Lateral Hypothalamus, and Medial Prefrontal Cortex and Transiently Stops 1288 Feeding. The Journal of Neuroscience, 36(50):12511-12529, 2016. ISSN 0270-6474. doi: 10.1523/ineurosci.1605-16.2016. 1289
- CS Maldonado-Irizarry, CJ Swanson, and AE Kelley. Glutamate receptors in the nucleus 1290 accumbens shell control feeding behavior via the lateral hypothalamus. The Journal of Neu-1291 roscience, 15(10):6779-6788, 1995. ISSN 0270-6474. doi: 10.1523/jneurosci.15-10-06779. 1292 1293
- Ryan W. S. Wee, Karyna Mishchanchuk, Rawan AlSubaie, and Andrew F. MacAskill. Hip-1294 76. pocampal ghrelin signalling informs the decision to eat. bioRxiv, page 2021.11.05.467326, 1295 2021. doi: 10.1101/2021.11.05.467326.1296
- David Lindenbach, Giada Vacca, Sovon Ahn, Jeremy K, Seamans, and Anthony G. 1297 Phillips. Optogenetic modulation of glutamatergic afferents from the ventral subiculum 1298 to the nucleus accumbens: Effects on dopamine function, response vigor and locomo-1299 tor activity. Behavioural Brain Research, 434:114028, 2022. ISSN 0166-4328. doi: 1300 10.1016/j.bbr.2022.114028. 1301
- 1302 Noriaki Koshikawa, Yasuhiro Miwa, Kazunori Adachi, Masafumi Kobayashi, and Alexan-1303 der R. Cools. Behavioural effects of 7-OH-DPAT are solely due to stimulation of dopamine D2 receptors in the shell of the nucleus accumbens; jaw movements. European Journa. 1304 of Pharmacology, 308(3):227-234, 1996. ISSN 0014-2999. doi: 10.1016/0014-2999(96) 1305 1306 00301-9
- Emma E Boxer, JungMin Kim, Brett Dunn, and Jason Aoto. Ventral subjculum inputs to 1307 nucleus accumbens medial shell preferentially innervate D2R medium spiny neurons and 1308 1309 contain calcium permeable AMPARs. bioRxiv, page 2022.10.12.511974, 2022. doi: 10. 1310 1101/2022 10 12 511974
- 1311 Zhao Li, Zhilong Chen, Guoqing Fan, Anan Li, Jing Yuan, and Tonghui Xu. Cell-Type-1312 Specific Afferent Innervation of the Nucleus Accumbens Core and Shell. Frontiers in Neuroanatomy, 12:84, 2018, ISSN 1662-5129, doi: 10.3389/fnana,2018,00084. 1313
- 1314 Mary Kay Lobo, Herbert E. Covington III, Dipesh Chaudhury, Allyson K. Friedman, 1315 HaoSheng Sun, Diane Damez-Werno, David M. Dietz, Samir Zaman, Ja Wook Koo, 1316 Pamela J. Kennedy, Ezekiell Mouzon, Murtaza Mogri, Rachael L. Neve, Karl Deisseroth, Ming-Hu Han, and Eric J. Nestler. Cell Type-Specific Loss of BDNF Signaling Mimics Opto-1317 1318 genetic Control of Cocaine Reward. Science, 330(6002):385-390, 2010. ISSN 0036-8075. 1319 doi: 10.1126/science.1188472
- 1320 Roberto A. Gulli, Lyndon R. Duong, Benjamin W. Corrigan, Guillaume Doucet, Sylvain Williams, Stefano Fusi, and Julio C. Martinez-Trujillo. Context-dependent representations 1321 of objects and space in the primate hippocampus during virtual navigation. Nature Neuro-1322 1323 science, 23(1):103-112, 2020. ISSN 1097-6256. doi: 10.1038/s41593-019-0548-3
- Hisham E. Atallah, Michael J. Frank, and Randall C. O'Reilly. Hippocampus, cortex, and 1324 basal ganglia: Insights from computational models of complementary learning systems. 1325 Neurobiology of Learning and Memory, 82(3):253-267, 2004. ISSN 1074-7427. doi: 10. 1326 1327 1016/j.nlm.2004.06.004.
- 1328 Randall C. O'Reilly and James L. McClelland. Hippocampal conjunctive encoding, storage, 1329 and recall: Avoiding a trade-off. Hippocampus, 4(6):661-682, 1994. ISSN 1098-1063. doi: 1330 10.1002/hipo.450040605
- Atsushi Kikumoto and Ulrich Mayr. Conjunctive representations that integrate stimuli, re-1331 1332 sponses, and rules are critical for action selection. Proceedings of the National Academy of Sciences, 117(19):10603-10608, 2020. ISSN 0027-8424. doi: 10.1073/pnas.1922166117. 1333
- 1334 Wenhao Dang, Russell J Jaffe, Xue-Lian Qi, and Christos Constantinidis. Emergence of 1335 Non-Linear Mixed Selectivity in Prefrontal Cortex after Training. The Journal of Neuro-1336 science, 41(35):JN-RM-2814-20, 2021. ISSN 0270-6474. doi: 10.1523/jneurosci.2814-20. 1337

87. Weilun Sun, Ilseob Choi, Stovan Stovanov, Oleg Senkov, Evgeni Ponimaskin, York Winter. 1338 Janelle M. P. Pakan, and Alexander Ditvatey. Context value updating and multidimensional neuronal encoding in the retrosplenial cortex. Nature Communications, 12(1):6045, 2021. doi: 10.1038/s41467-021-26301-z.

1339

1340

1341

1342

1344

1345

1346

1347

1348

1349

1350

1353

1354

- Malavika Murugan, Hee Jae Jang, Michelle Park, Ellia M. Miller, Julia Cox, Joshua P. Taliaferro, Nathan F. Parker, Varun Bhave, Hong Hur, Yupu Liang, Alexander R. Nectow, 1343 Jonathan W. Pillow, and Ilana B. Witten. Combined Social and Spatial Coding in a Descending Projection from the Prefrontal Cortex. Cell. 171(7):1663-1677.e16. 2017. ISSN 0092-8674. doi: 10.1016/j.cell.2017.11.002.
- Gordon J. Mogenson and Mark Nielsen. A study of the contribution of hippocampal—accumbens—subpallidal projections to locomotor activity. Behavioral and Neural Biology, 42(1):38-51, 1984. ISSN 0163-1047. doi: 10.1016/s0163-1047(84)90412-6.
- D. Gowanlock R. Tervo, Bum-Yeol Hwang, Sarada Viswanathan, Thomas Gaj, Maria Lavzin, Kimberly D. Ritola, Sarah Lindo, Susan Michael, Elena Kuleshova, David Ojala, Cheng-Chiu 1351 Huang, Charles R. Gerfen, Jackie Schiller, Joshua T. Dudman, Adam W. Hantman, Loren L. Looger, David V. Schaffer, and Alla Y. Karpova. A Designer AAV Variant Permits Efficient Retrograde Access to Projection Neurons. Neuron, 92(2):372–382, 2016. ISSN 0896-6273. doi: 10.1016/j.neuron.2016.09.021.
- Georg Nagel, Tanjef Szellas, Wolfram Huhn, Suneel Kateriya, Nona Adeishvili, Peter 1356 Berthold, Doris Ollig, Peter Hegemann, and Ernst Bamberg. Channelrhodopsin-2, a directly light-gated cation-selective membrane channel. Proceedings of the National Academy of 1358 Sciences, 100(24):13940-13945, 2003. ISSN 0027-8424. doi: 10.1073/pnas.1936192100.
- Anqi Wu, E Kelly Buchanan, Matthew R Whiteway, Michael Schartner, Guido Meijer, Jean-Paul Noel, Erica Rodriguez, Claire Everett, Amy Norovich, Evan Schaffer, Neeli Mishra, C Daniel Salzman, Dora Angelaki, Andrés Bendesky, The International Brain Laboratory, John Cunningham, and Liam Paninski. Deep Graph Pose: a semi-supervised deep graphical model for improved animal pose tracking. bioRxiv, page 2020.08.20.259705, 2020. doi: 10.1101/2020.08.20.259705.
- Alexander Mathis, Pranav Mamidanna, Kevin M. Cury, Taiga Abe, Venkatesh N. Murthy, Mackenzie Weygandt Mathis, and Matthias Bethge. DeepLabCut: markerless pose estimation of user-defined body parts with deep learning. Nature Neuroscience, 21(9):1281-1289, 1368 2018. ISSN 1097-6256. doi: 10.1038/s41593-018-0209-y.

Supplementary videos

Supplemental Video 1. In vivo dual-color two-photon 1371 calcium imaging of a mouse during goal-directed nav-1372 igation. Representative example of dual-color two-photon calcium imaging combined with goal-directed navigation and 1374 lick behavior. Infrared camera images show face (top left) and body (top right) tracking. Calcium activity is shown 1376 by overlaying denoised GCaMP6s activity (green) onto av-1377 erage mCherry signal (red; bottom left); traces represent de-1378 noised GCaMP6s activity of mCherry-negative (green) and 1379 mCherry-positive (red; NAc-projecting) hippocampal neu-1380 rons (bottom right). Playback speed is $2 \times$ original speed. 1381

Supplemental Video 2. Optogenetic stimulation induces mouth movement. Five representative optogenetic 1383 stimulation trials from one experimental mouse expressing 1384 ChR2 and one control mouse expressing EYFP only. A light 1385 fiber in NAc stimulated excitatory hippocampal projections by shining 473 nm laser light at times indicated by blue 1387 1388 squares (top) and blue traces (bottom). Optogenetic stimulation repeatedly induces lick behavior and a decrease in ve-1389 locity in ChR2-expressing animals but not those expressing EYFP. Playback speed equals original speed.

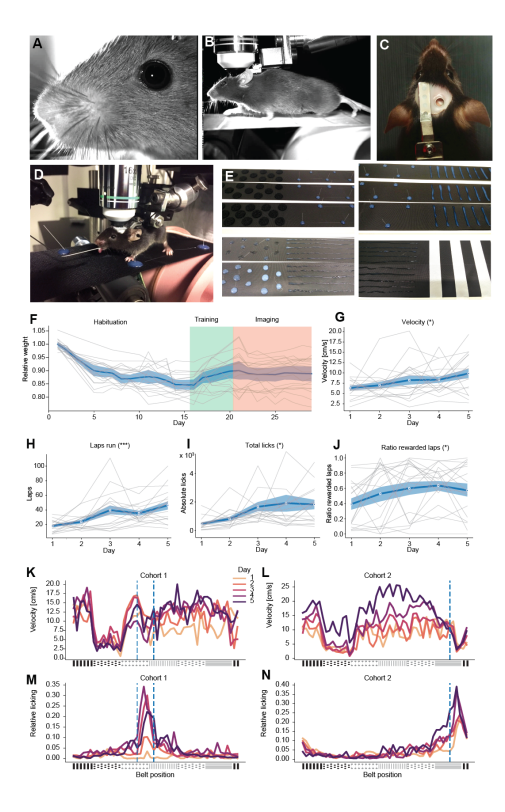


Figure S1. Spatial reward learning task. (A and B) Sample infrared images of face (A) and body (B) that were continuously captured at 25/75 Hz. (C) Image of one head-fixed experimental mouse from top, illustrating craniotomy and head-fixation. (D) Sample image of two-photon imaging of a head-fixed mouse on treadmill belt, licking at the lick spout. (E) Sample images of the six different belt texture zones, including Velcro dots, nylon spikes, vertical glue stripes, glue dots, horizontal glue stripes, white tape stripes (from left top to right bot-(F) Weight changes after introduction of food restriction and during training and imaging. (G-J) Average velocity (G, F(4) =2.631, p = 0.0430), number oflaps run (H, F(4) = 8.771, p <0.001, GG-corrected), number of licks (I, F(4) = 3.883, p = 0.0326,GG-corrected), and ratio of rewarded laps (J, F(4) = 3.331, p =0.0157) increase over the course of five training days (all repeatedmeasures ANOVAs). Gray lines indicate data points of individual animals, blue shade represents SEM, blue line represents mean. (K-N) Average velocity (K and L) and average licking (M and N) changes across belt position and days of mice tested with reward zone in "center" of belt (K and M) and "end" of belt (L and N). Blue dashed line represents presence of reward zone. n = 9 (cohort 1), n = 9 (cohort 2). All data are presented as mean \pm SEM. *p < 0.05, ***p <0.001.

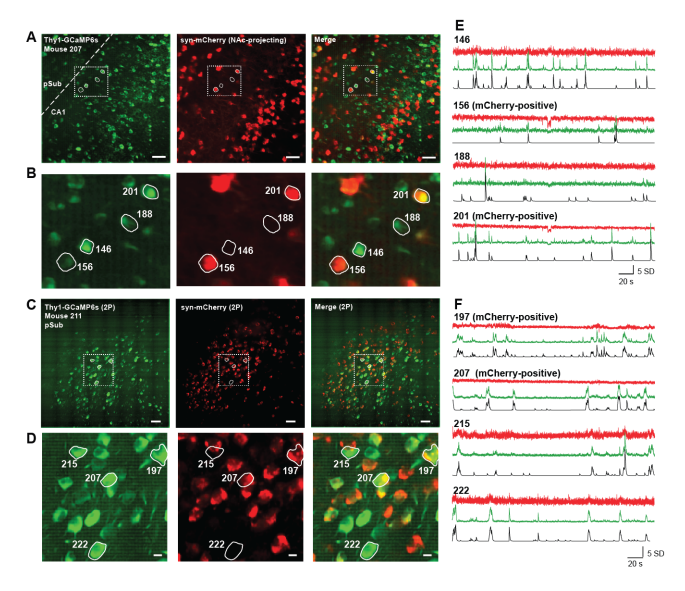


Figure S2. In vivo dual-color two-photon imaging of dynamic GCaMP and static mCherry signals. (A) Sample field of view (FOV) of one imaged region crossing putative CA1 and prosubiculum (pSub). Green channel shows GCaMP6s local correlations image, red channel shows average motion-corrected mCherry signal. (B) Detail of region in (A) clearly outlining spectrally separate channels. (C and D) Sample FOV of another region in putative prosubiculum (C) with detailed view (D). Scale bars represent 50 μ m (A and C) or 10 μ m (B and D). (E and F) Z-scored fluorescent temporal sample traces of regions of interest (ROIs) shown in (A-D). Red traces show raw fluorescence dynamics from imaging red channel, green traces show raw fluorescence dynamic from green channel, and black traces show denoised calcium signal.

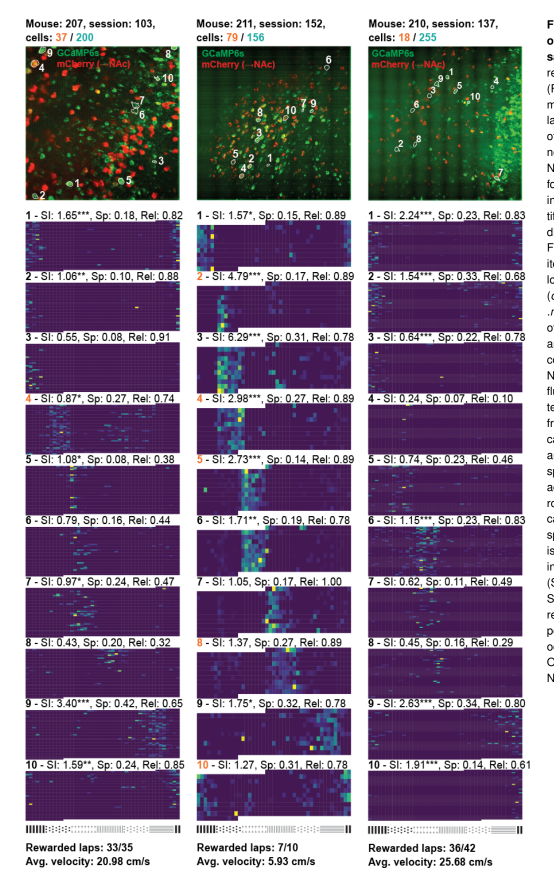


Figure S3. Spatial tuning of different neurons in sample FOVs. Three representative fields of view (FOVs) from three different mice are shown, with singlelap spatial calcium activity of each 10 representative neurons, some putatively FOV in-NAc-projecting. formation is shown on top, including numbers of identified dHPC (green) and $dHPC^{\rightarrow NAc}$ (red) neurons. FOVs shown are composites of CalmAn maximum local correlation images (caiman.summary images .max correlation image) of GCaMP channel (green) and the averaged motioncorrected red channel with NAc-projecting mCherry fluorescence. Contours of ten representative neurons from each FOV are indicated with white outlines and numbers that refer to spatially averaged calcium activity below. Each neuron's normalized average calcium activity across 45 spatial bins per lap (y axis) is shown with key spatial information values above (SI: spatial information (47), Sp: Sparsity (48), Rel: reliability of each neuron's per-lap maximum activity to occur within the place field). Orange numbers refer to NAc-projecting neurons.

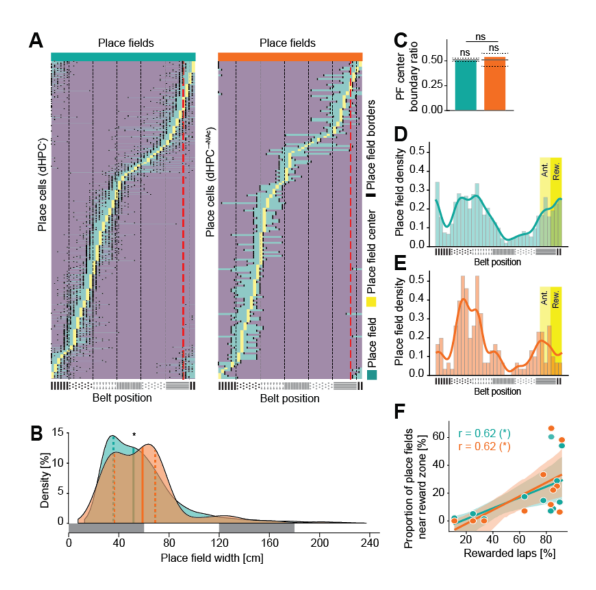


Figure S4. Place field distribution across the belt and behavioral performance. (A) Extent of place fields of dHPC[−] (left) and dHPC^{→NAC} (right) place cells, sorted by each place cell's center of mass (COM). Turquoise represents place field, yellow represents COM, black lines represent left and right borders of place fields. Red dashed line represents beginning of reward zone, black dashed lines represent belt texture borders. (B) Width of place fields is unevenly distributed between dHPC[−] (green; quartiles 35.6 cm / 51.7 cm / 69.0 cm) and dHPC^{→NAC} (red; quartiles 36.5 cm / 58.9 cm / 69.0 cm) place cells (Kolmogorov-Smirnov test, D = 0.116, p = 0.0299). Gray bars represent belt texture zones (60 cm). (C) Place field centers are not biased with respect to texture boundaries (percentile < 95th, permutation test) and the ratio is not different between neuronal populations (χ^2 (1, 5372) = 1.646, p = 0.1995). (D and E) Place field density between reward and anticipation zones compared to the rest of the belt for dHPC[−] (D) and dHPC^{→NAC} (E) populations. (F) Reward and anticipation zone overrepresentation correlates with behavioral success (percentage of rewarded laps per session) for both dHPC[−] (green) and dHPC^{→NAC} (red) populations (Pearson correlation; r_{dHPC} -(15) = 0.622, p = 0.0107; r_{dHPC} -NAC (10) = 0.619, p = 0.0421). ns: not significant, *p < 0.05.

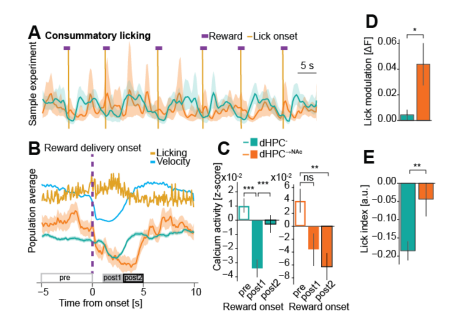


Figure S5. Lick modulation during consumption and general licking. (A-C) Consummatory licking leads to a depression in neural activity. (A) Sample experiment showing reward dispensation (purple bars) and consummatory licking onsets (golden vertical lines) and population calcium activity from dHPC⁻ (green) and dHPC⁻NAc (red) neurons. Note the robust depression of neural activity during times of reward consumption. (B) Event-triggered average traces around reward delivery onset, including licking and speed as well as population average calcium activity of dHPC⁻ (green) and dHPC⁻NAc (red) neurons. Gray rectangles indicate time windows for comparing calcium activity shown in (C). (C) Calcium activity is differentially modulated by reward onset (two-way mixed ANOVA; $F_{\text{reward_timing}}(2, 10740) = 19.380$, p(GG-corrected) < 0.001, $F_{\text{projection}}(1, 5370) = 0.798$, p = 0.372, $F_{\text{interaction}}(2, 10740) = 5.559$, p = 0.0039). Post-hoc pairwise t-tests with Bonferroni correction were performed for each interaction and the results indicated with asterisks. (D) Lick modulation (average of each trial's pre-post difference) is significantly increased in dHPC^{-NAc} neurons (Welch's t(496.11) = 2.379, p = 0.0177). (E) Lick index (average calcium activity difference during licking) is significantly increased in dHPC^{-NAc} neurons (Welch's t(454.19) = 2.729, p = 0.0066). All data are presented as mean \pm SEM. *p < 0.05, **p < 0.01, ***p < 0.001.

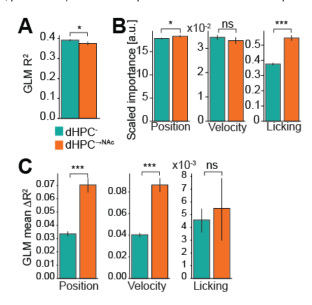


Figure S6. Generalized linear model reveals coding differences for dHPC $^{\rightarrow$ NAc} neurons. (A) GLMs' explained variance R^2 is significantly lower for dHPC $^{\rightarrow$ NAc} neurons (Welch's t(615.15) = 2.147, p = 0.032). (B) Scaled feature importance (standardized GLM coefficients retrieved by H2O's varimp() method) is higher in dHPC $^{\rightarrow$ NAc} neurons for position (Welch's t(2508.09) = 2.162, p = 0.031) and licking (Welch's t(1879.75) = 9.328, p < 0.001), but not for velocity (Welch's t(2609.15) = 0.881, p = 0.378). (C) Mean R^2 differences between full and feature-shuffled models are significantly greater in dHPC $^{\rightarrow$ NAc} neurons for position (Welch's t(523.23) = 5.984, p < 0.001), velocity (Welch's t(497.69) = 7.704, p < 0.001), but not for licking (Welch's t(552.63) = 0.3626, p = 0.717). All data are presented as mean \pm SEM. ns: not significant, *p < 0.05, ***p < 0.001.