

Inhibitory neurons defined by a synaptogenic molecule impair memory discrimination

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SUMMARY

The CA3 region is central to hippocampal function during learning and memory because of its unique connectivity. CA3 pyramidal neurons are the targets of huge, excitatory mossy fiber synapses from DG axons and have an unusually high degree of excitatory recurrent connectivity. Thus, inhibition likely plays an outsized importance in constraining runaway excitation and shaping CA3 ensembles during learning and memory. Here, we investigate the function of a group of dendrite-targeting, hippocampal GABAergic neurons defined by expression of the synaptogenic adhesion molecule, Kirrel3. We discovered that activating Kirrel3-expressing GABAergic neurons impairs memory discrimination by inhibiting CA3 pyramidal neurons in novel contexts. Kirrel3 is required for DG-to-GABA synapse formation and variants in Kirrel3 are strong risk factors for neurodevelopmental disorders. Thus, our work suggests that Kirrel3-GABA neurons are a critical source of feed-forward inhibition from DG to CA3 during contextual memory whose activity may be specifically disrupted in some brain disorders.

KEYWORDS

interneuron, inhibition, memory discrimination, memory generalization, CA3, hippocampus, contextual fear conditioning, Kirrel3, mossy fiber synapse

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AUTHOR CONTRIBUTIONS

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DECLARATION OF INTERESTS

The authors declare no competing interests.

INTRODUCTION

The hippocampus is essential for learning and memory and, in particular, the CA3 region is critical for recalling and integrating contextual information¹⁻⁷. CA3 pyramidal neurons, the principal excitatory cells of the region, receive excitatory inputs from the DG. These DG inputs are extremely large mossy fiber boutons that synapse onto multi-headed CA3 spines. These powerful, excitatory synapses have up to 30 active zones and are known as “detonator” synapses because it is possible that release from a single bouton activates a CA3 neuron^{8,9}. In addition, CA3 pyramidal neurons are interconnected via numerous recurrent collateral synapses. This combination of mossy fiber detonator synapses and extensive recurrent connectivity means that inhibition likely has an outsized importance in shaping CA3 activity and preventing runaway excitation¹⁰⁻¹³. In fact, DG mossy fiber synapses sprout filopodia directly from the presynaptic bouton to synapse with local GABAergic (GABA) inhibitory neurons that provide feed-forward inhibition to CA3^{11,14}. Despite the importance of inhibition in CA3, most hippocampal studies have focused on inhibition in the DG and CA1; regions that are easier to access and have more traditional synaptic structures than CA3. Recent work has begun to describe CA3 GABA neuron dynamics¹⁵ and appreciate the role of feed-forward inhibition in remote memory¹⁶⁻¹⁸, but there remain large gaps in knowledge about how different types of CA3 GABA neurons modulate learning and memory.

GABA neurons are far fewer in number than excitatory neurons, but they are incredibly diverse and essential for proper brain function¹¹. Sequencing studies estimate that the hippocampus houses up to 30 transcriptionally distinct types of GABA neurons¹⁹⁻²³. It is thought that different transcriptional profiles impart distinct morphological and functional properties to GABA neurons. Despite this diversity, the genetic toolset to identify and manipulate different subpopulations of GABA neurons in mouse models is limited and, consequently, only a few cardinal GABA subpopulations are well studied. In contrast, we investigated the function of a previously unstudied subpopulation of GABA neurons identified by the synaptogenic cell adhesion molecule Kirrel3. We previously discovered that Kirrel3 is necessary to properly form DG-to-GABA mossy fiber filopodia synapses that provide feed-forward inhibition to CA3^{24,25} and, consequently, loss of Kirrel3 results in increased CA3 neuron activity²⁴. Together, these observations suggest that Kirrel3 regulates a critical source of feed-forward inhibition between DG and CA3, and that this inhibition is mediated by Kirrel3-expressing GABA (K3-GABA) neurons. Interestingly, K3-GABA neurons are not identified by any one cardinal GABA neuron marker²⁴. Thus, we generated a new driver line that expresses Flp recombinase in Kirrel3-expressing cells and used intersectional genetics to specifically study K3-GABA neurons.

Here, we demonstrate that activating K3-GABA neurons, but not parvalbumin GABA neurons, reduces memory discrimination in a contextual fear conditioning task. We further show that K3-GABA neurons in the CA3 and hilus are a transcriptionally heterogeneous population of dendrite-targeting GABA neurons that selectively inhibit CA3 pyramidal activity in novel contexts. Thus, our work supports an alternative classification of GABA neurons based on synapse-specific molecules, which have potential to drive a shared connectivity pattern among otherwise transcriptionally heterogeneous groups of neurons²⁶. Moreover, because Kirrel3 variants are associated with neurodevelopmental disorders²⁷⁻⁴¹, our findings and new tools lay the foundation for studying the specific circuits and behaviors that may be disrupted by Kirrel3 variants.

RESULTS

Kirrel3-expression marks a subpopulation of GABA neurons. To study K3-GABA neurons, we generated a mouse line that expresses Flp recombinase under control of the endogenous Kirrel3 gene. We inserted the Flp sequence just after the start codon of Kirrel3 (Figure 1A) along with a viral 2A sequence to allow for normal expression of Kirrel3 protein (Figure 1B). To test the specificity of our new mouse line and identify K3-GABA but not DG neurons, which also express Kirrel3, we crossed the Kirrel3-Flp line with a Gad2-Cre line expressing Cre in all GABA neurons⁴² and to a reporter line known

as RC:FLTG⁴³. This reporter labels cells co-expressing both Flp and Cre with GFP (K3-GABA neurons) and cells expressing Flp alone with tdTomato (DG neurons) (Figure 1C). There is no suitable antibody to label Kirrel3 in tissue, so we used fluorescent in situ hybridization (FISH) with RNA probes against Kirrel3, GAD1, and GFP to test if the GFP positive cells in the triple transgenic line are GABA neurons that express Kirrel3 (Figure 1D). We imaged cells in area CA3 and found that the transgenic line functions as expected. 94% \pm 14 of Kirrel3 cells in CA3 are GABA neurons (mean \pm sd, Figure 1E) and 86% \pm 17 of GFP cells express Kirrel3 (mean \pm sd, Figure 1E). This experiment also indicates that 34% \pm 14 of all GABA neurons express Kirrel3 mRNA (mean \pm sd, Figure 1E). This is slightly more than we previously reported (~19%) using immunohistochemistry on Kirrel3 knockout mice, which express GFP instead of Kirrel3²⁴ and is likely due to different sensitivities of each method.

We previously tested if K3-GABA neurons could be described by a common GABA subtype marker but did not find a match²⁴. We revisited this idea using our new reporter mouse. We stained K3-Flp;Gad2-Cre;RC:FLTG mice for GFP to mark K3-GABA neurons and a selection of GABA markers, including parvalbumin as shown in Figure 1F. We again found that K3-GABA neurons are not identified by any one GABA marker (Figure 1G). Instead of examining individual marker proteins one at a time, we also used publicly available single cell sequencing data from mouse hippocampus to determine if K3-GABA neurons fall into a single transcriptomic GABA class. Data from the DropViz database uses unsupervised clustering to group GABA neurons into 27 distinct transcriptomic classes²². We determined the percent of cells in each group that had one or more reads of a Kirrel3 transcript. All classes except one, had some cells that express Kirrel3, no class had more than 30% of its cells express Kirrel3, and most classes had between 5-15% of their cells with at least one Kirrel3 transcript (Figure 1H, Table S1). Similar results were obtained using data from the Allen Institute²³ (not shown). Taken together, we conclude that Kirrel3 mRNA is expressed by up to 1/3 of all hippocampal GABA neurons and they are a heterogenous group that is not categorized by a single transcriptomic class.

Specific activation of K3-GABA neurons using intersectional viral vectors. To study the function of K3-GABA neurons in area CA3, we needed a tool to selectively activate them in vivo. We used a newly developed ConVERGD intersectional expression vector that is Cre and Flp-dependent⁴⁴. It contains a LoxP-flanked ribozyme, which needs to be removed by Cre to prevent mRNA degradation, and an inverted, Frt-flanked payload driven by the human synapsin promoter⁴⁴. We tested the functionality of an AAV-ConVERGED vector with the activating DREADD payload, hM3D-mCherry (Figure 2A). We first confirmed that hM3D-mCherry expression depended on both Flp and Cre in vitro (Figure S2A). Next, we injected the hM3D-mCherry ConVERGD virus into hippocampal area CA3 of K3-Flp; Gad2-Cre double heterozygous mice to specifically express hM3D-mCherry in K3-GABA neurons. We observed cell labeling in a pattern consistent with the known location of K3-GABA neurons with no off-target expression in K3-Flp or wildtype mice (Figure 2B and S2B). We used a dual FISH/immunostaining protocol to confirm that the hM3D-mCherry protein is selectively expressed in cells that express Kirrel3 and Gad1 mRNA (K3-GABA neurons) (Figure 2C). Taken together, our data indicate that we achieved specific targeting of K3-GABA neurons.

We next tested if the ConVERGD-hM3D-mCherry activates K3-GABA neurons after injection with the agonist, deschloroclozapine (DCZ). K3-Flp; Gad2-Cre mice were injected with the virus. Two weeks later, mice were intraperitoneally injected with saline or DCZ 10 minutes before placing them in a behavior box and subjecting them to 4-foot shocks as used for fear conditioning. 30 minutes later, mice were perfused, and sections were stained for the immediate early gene cFos, a common measure of recent neuronal activity. We found that DCZ administration significantly increased the percent of K3-GABA neurons expressing cFos compared to saline (Figure 2D, E), indicating that the hM3D tool activates K3-GABA neurons as expected.

Activating K3-GABA neurons impairs memory discrimination. We next sought to determine if and how activating K3-GABA neurons alters mouse behavior in a hippocampal-dependent learning and memory task. To do this, we established an expanded contextual fear conditioning paradigm (Figures

3A, B). We conditioned mice in context A where they received 4 unsigned foot shocks. 24 hours later, we tested their fear memory by measuring their freezing behavior when placing them back in context A without foot shock and in a similar but slightly different context B. The next day, we again tested them in context A and in a very distinct context C. The time spent freezing in each context is quantified and the difference in freezing relative to context A is quantified as the discrimination ratio ($A/(A+B)$) between the two contexts on each day. Wildtype mice are generally poor at distinguishing between similar contexts A and B but reliably discriminate between distinct contexts A and C⁴⁵⁻⁴⁷ (Figure S3A and S3B). This behavioral paradigm allows us to evaluate several aspects of learning and memory upon manipulation of K3-GABA neuron activity including, if the mice can learn to associate context A with the shock at all, if memory discrimination between contexts A and B improves, or if discrimination between contexts A and C worsens.

Next, we tested if activating K3-GABA neurons alters the learning and memory of mice using this paradigm. Male and female K3-Flp; Gad2-Cre heterozygous mice were bilaterally injected in the CA3 region with ConVERGD-hM3D-mCherry AAVs and randomly divided into a control group that received saline injections or an experimental group that received DCZ injections. Because the function of K3-GABA neurons was completely unknown, we initially opted to give saline or DCZ injections prior to all contexts (Figure 3A). All mice conditioned normally to foot shock on training day 0 (Figure S3C, D, H), and showed normal fear conditioning in context A and B on day 1 (Figure 3C, D). However, on day 2 we observed a significant increase in the fear response when K3-GABA neurons were activated by DCZ in the neutral context C (Figure 3C). This resulted in a significant decrease in the A/C discrimination ratio compared to saline controls (Figure 3D) and no difference in the A/B and A/C discrimination ratios within the DCZ condition (Figure 3D). Importantly, mice that did not express the DREADD but were injected with DCZ had normal behavior (Figure S3A and S3B). Thus, our data suggests that K3-GABA activation impairs memory discrimination by increasing fear generalization in a neutral context.

To test if this phenotype is specific to K3-GABA neurons or if it would result from activating any similar sized subpopulation of GABA neurons, we expressed hM3D-mCherry in parvalbumin (PV) neurons (Figure 3E). PV-Cre mice were bilaterally injected with a Cre-dependent hM3D-mCherry and tested in the same contextual fear conditioning paradigm. We found that the DREADD appropriately activated PV-GABA neurons as measured by cFos expression (Figure S3G) but that activating PV-GABA neurons did not alter learning or memory discrimination on either day compared to saline controls (Figure 3F,G). We confirmed proper hM3D-mCherry expression and targeting in all mice used for experiments (Figure S3I and S3K). Together, our data indicate that K3-GABA neurons have a specific function in memory discrimination compared to PV neurons.

Activating K3-GABA neurons decreases CA3 activity in novel contexts. To begin to understand how activating K3-GABA neurons might result in impaired memory discrimination at the circuit level, we sought to determine if activating K3-GABA neurons affects CA3 pyramidal neuron activity. We first imaged K3-GABA projections to determine generally where they contact CA3 neurons by sparsely labeling K3-GABA neurons with a fluorescent reporter. We could clearly see axonal labeling in the CA3 SR and SO layers (Figure 4A, B), suggesting K3-GABA neurons target CA3 pyramidal cell dendrites.

Next, we tested if activating K3-GABA neurons with DREADDs inhibits CA3 neuron activity using cFos staining. One week after mice were run through the full contextual fear conditioning paradigm (Figure 3A), we injected mice with either saline or DCZ and then returned them to the familiar context A with foot shock. Following cFos immunostaining, we found that DCZ injection activates K3-GABA neurons compared to saline as reported in Figure 2 (data again shown in Figure 4C), but surprisingly, we observed no corresponding change in CA3 neuron activity as measured by the density of cFos positive cells in the pyramidal cell layer in the same mice (Figure 4D). We then reasoned that the DCZ injected mice froze normally in every exposure to context A in the behavioral tests (Figure 3A) so we analyzed cFos activity 30 minutes after mice were exposed to a novel and neutral enriched environment, which is

similar to the scenario in which they displayed abnormal behavior. In this condition, we again observed an increase in cFos-positive K3-GABA neurons (Figure 4C, E), indicating that our DREADD is working, and we also observed a significant decrease in the number of cFos-positive CA3 neurons (Figure 4F). This suggests that activation of K3-GABA neurons selectively decreases net CA3 neuron activity in response to novel, but not familiar, contexts. Interestingly, DCZ alone was not enough to activate K3-GABA neurons in the home cage condition (Figure 4C), suggesting that K3-GABA neurons can only be activated under specific environmental conditions. Together, this strongly suggests that K3-GABA neuron activity specifically inhibits CA3 pyramidal neurons in novel environments.

DISCUSSION

In sum, we demonstrate that activating K3-GABA neurons decreases the number of active CA3 neurons and causes mice to generalize fear to a novel, neutral environment. These results are consistent with what is broadly known about memory and ensemble size, which is that increasing ensemble size improves memory discrimination, while decreasing ensemble size worsens memory discrimination^{48–52}. Consistent with these other studies in different brain regions, activating K3-GABA neurons ultimately decreased CA3 ensemble size and worsened memory discrimination.

More specifically, we observed that K3-GABA neuron activation caused mice to generalize fear to a novel, neutral context. Why might this happen at the cellular level? We speculate that is because at any given time active CA3 neurons come from two pools. Some CA3 neurons are activated directly from the DG by the huge mossy fiber detonator synapses, while others are activated by subsequent recurrent connectivity and entorhinal inputs. It is unlikely that dendrite-targeting K3-GABA neurons would prevent powerful mossy fiber synapse transmission. However, K3-GABA neuron activity may constrain CA3 activity by limiting the amount of downstream recurrent CA3 activation and entorhinal inputs. If the strong DG inputs encode core features of the memory, such as being taken out of their home cage and put in a behavior box, while recurrent CA3 activity contributes to encoding or retrieving contextual details, then it would be predicted that activating K3-GABA neurons would retain the core fear memory but lose contextual detail as we observed. There are certainly other ways that K3-GABA neurons could affect CA3 activity including feed-back inhibition to DG and by modulating other types of GABA neurons or entorhinal inputs^{7,46,53–56}. Future work will need to map the input and output connectivity of K3-GABA neurons in more detail, but our work clearly implicates K3-GABA neurons as an important modulator of CA3 activity.

Our results presented here also reveal several intriguing features about the specificity of K3-GABA neuron activity. First, we showed that activation of K3-GABA neurons, and not PV-GABA neurons, decreases memory discrimination. This indicates that K3-GABA neurons not only have a distinct function from PV neurons, but that the behavioral changes we observed here are specific to K3-GABA neurons and not due to simply modulating any similar sized population of GABA neurons. Second, we observed that activating K3-GABA neurons decreased CA3 activity in novel contexts but not in a familiar context. This suggests that CA3 neuron activity is tightly controlled by multiple different kinds of neurons, and they are uniquely vulnerable to K3-GABA neuron activity only under certain conditions. Third, we observed that K3-GABA neurons expressing the activating DREADD are not responsive to DCZ in the home cage condition, but they are upon contextual stimulation. This suggests that either K3-GABA neurons are strongly inhibited in a familiar environment or that their baseline activity level is not in the working range of the DREADD receptor. Our experiments using cFos as a readout for K3-GABA and CA3 activity cannot reveal subtle changes in activity dynamics but is an important foundation for understanding their net activity in vivo before and after behavior. Moreover, the highly specific activity of K3-GABA neurons could relate to the synapse-specific nature of the Kirrel3 protein. DG neurons make synapses with many types of GABA neurons via two types of synapses; typical en passant synapses along the entire length of the axon and mossy fiber filopodia synapses that are only found protruding from large mossy fiber boutons^{14,25,57}. We previously found that loss of Kirrel3 protein specifically reduces the density of DG-to-GABA mossy fiber filopodia synapses but not DG-to-GABA en passant

synapses²⁵. Thus K3-GABA neurons in area CA3 may be highly specialized to receive mossy fiber filopodia input and could be an important avenue to investigate this poorly understood aspect of the hippocampal circuit.

Finally, our work is one of the first to study a group of GABA neurons based on the expression of a molecule that is a strong driver of synaptic connectivity. Thus, K3-GABA neurons, though transcriptionally heterogenous, may be functionally linked in the circuit by sharing common input and output neurons driven, at least in part, by Kirrel3. Cell type-specific study of GABA neurons using cardinal markers like PV, SST, and VIP enabled groundbreaking discoveries in circuit motifs, but it is becoming clear that even these tools label heterogenous populations of GABA neurons with diverse circuit functions^{58,59}. The generation of new tools to study distinct cell types, combined with intersectional approaches like those used here, should propel our understanding of circuit function in the future. Moreover, missense variants and copy number changes in Kirrel3 are strongly associated with increased risk for autism spectrum disorders, intellectual disabilities, and other neurodevelopmental disorders²⁷⁻⁴¹. Thus, understanding how changing K3-GABA neuron activity affects brain circuits could have significant clinical relevance and, taken together, bridges molecular, cellular, and systems neuroscience.

FIGURE LEGENDS

Figure 1. Kirrel3 expression marks a heterogenous group of GABA neurons. A) Schematic of the FlpO-2A insertion site in the mouse Kirrel3 gene. B) Western blot of brain lysates from 2 months old mice show that the Flp transgenic line expresses normal levels of Kirrel3 protein. Lysates from Kirrel3 knockout (KO) and wildtype (WT) are used as controls. Coomassie stained membrane indicates equal protein loading. C) A hippocampal section from an adult triple transgenic mouse (heterozygous for all transgenes) stained with antibodies against TdTomato (red), GFP (green), and vGlut1 (blue). D) A magnified image of a hippocampal section labeled by fluorescent in situ hybridization (FISH). Arrows indicate K3-GABA neurons co-expressing Kirrel3, GFP, and GAD1 mRNA. E) Quantification of FISH as indicated. Each column represents a mouse and each dot represents one section. Average percentages are shown as a dotted line. n = 7 adult mice (3 male and 4 female) Error bars represent SEM. F) An adult triple transgenic mouse stained for GFP to label K3-GABA neurons (green) and parvalbumin (magenta). Note the lack of overlap. G) Table showing the percent of K3-GABA neurons that express the indicated marker as assessed by immunostaining triple transgenic mice similar to F. n = 3 female adult mice. Total number of neurons counted for each marker is indicated. H) Graph showing the percent of cells with at least one Kirrel3 RNA read in each transcriptomic class of GABA neuron from the DropViz data set. Classes that express parvalbumin (PV) and somatostatin (SST) are indicated. The total number of cells in each class is indicated. No Kirrel3 transcripts were found in any cells in group 4.

Figure 2. Selectively targeting and activating K3-GABA neurons. A) Schematic showing the ConVERGD AAV construct with an hM3D-mCherry transgene. B) Representative hippocampal image from a K3-Flp;Gad2-Cre mouse with ConVERGD-hM3D-mCherry targeted to area CA3. C) Magnified image from a hippocampal section processed for dual FISH/immunohistochemistry from a K3-Flp;Gad2-Cre mouse infected with ConVERGD-hM3D-mCherry. D) Representative images showing cFos (green) in K3-GABA neurons expressing hM3D-mCherry (magenta) specifically after DCZ injection (right). Arrowheads indicate K3-GABA neurons that express cFos (white). E) Quantification of % of infected K3-GABA neurons that express cFos for saline and DCZ treated mice after foot shock. n = 4 males each. Each point represents one mouse, up to 3 sections and 6 hippocampi counted per mouse. Error bars represent SEM, nested T-test, p=0.0058.

Figure 3. K3-GABA activation impairs memory discrimination. A) Schematic of behavior paradigm. B) Description of contextual cues between conditioning context A, similar context B, and neutral context C. C) Time spent freezing (%) when placed in indicated contexts after saline (gray) or DCZ (blue)

injection. All mice were adult K3-Flp; Gad2-Cre heterozygotes injected with ConVERGD hM3D-mCherry. n = 14 saline, 14 DCZ. (D) Data from (C) plotted as a discrimination ratio relative to context A, where 0.5 marks no discrimination and is calculated by the percent freezing time for A/A+X, where X is either context B or C depending on the day, ** indicates p=0.0021, *** indicates p=0.0001, ns indicates p=0.6263. E) Representative image of a hippocampal section from a PV-Cre mouse infected with a Cre-dependent hM3D-mCherry AAV (magenta) and stained with anti-PV antibodies (green). Overlap appears white. F) Time spent freezing (%) when placed in indicated contexts after saline (gray) or DCZ (red) injection. All mice were adult PV-Cre heterozygous mice injected with Cre-dependent hM3D-mCherry. F) Discrimination ratios for PV-Cre mice showed normal discrimination for saline and DCZ injected mice, n = 19 saline, 16 DCZ, ** indicates p=0.0034, *** indicates p=0.0003, and n.s. indicates p=>0.9999. Error bars represent SEM, ANOVA multiple comparisons. Males and females are indicated.

Figure 4. K3-GABA neurons inhibit CA3 in novel contexts. A, B) K3-Flp;Gad2-Cre heterozygous mouse infected with an AAV that expresses GFP in a Flp and Cre-dependent manner to label K3-GABA neurons. B shows a magnified image of CA3 from a section sparsely labeled. s.o.; stratum oriens, s.p.; stratum pyramidale, s.l.; stratum lucidum, s.r.; stratum radiatum. C) % of hM3D-mCherry infected K3-GABA neurons that are cFos positive after saline (grey) and DCZ injection (blue) in indicated environments. HC = homecage, FS = foot shock, EE = enriched environment. Nested ANOVA with post-tests, ** indicates p=0.0058, **** indicates p=<0.0001. ns for FS indicates p=0.5174, ns for HC indicates p=.1583. D) Number of cFos-positive CA3 neurons per mm² for saline (gray) and DCZ treated (blue) mice in indicated contexts n = 4-6 per group. Each dot represents a hippocampus, each column represents a mouse, nested ANOVA with post-tests, * indicates p=0.0113. Error bars represent SEM. E) Representative images of hippocampal CA3 sections immunostained for cFos in hM3D-mCherry K3-Flp;Gad2-Cre mice treated with saline or DCZ.

MATERIALS AND METHODS

Mouse lines: The Kirrel3-Flp line FlpO-P2A was inserted in frame and immediately upstream of the Kirrel3 start codon so that Kirrel3 protein expression is unperturbed. We inserted FlpO at the start because the Kirrel3 gene undergoes alternative splicing at its C-terminus. Transgenic mice were generated at the University of Nebraska School of Medicine Transgenic core using the Easi-CRISPR method of homologous recombination⁶⁰. Briefly, single cell C57Bl6 mouse zygotes were injected with preassembled Cas9 ribonucleotide complexes containing Cas9 protein, a Kirrel3 crRNA (AGGAATGAGACCTTTCCAGC), and tracrRNA along with a 1,536 bp single strand DNA (ssDNA) repair template containing the FlpO-P2A sequence and ~100bp homology arms. A founder mouse was fully sequenced to ensure there were no mutations at insertion sites and the mouse was bred to a wildtype C57Bl6 mouse to propagate the strain. The donor ssDNA sequence is as follows:
GCCGCGCTTGAAGAGAACTAACTGCACACCCAAGTTGCCGCGGCTGCCGCGCGCTGAGGAATGAGACCTTTCCGCT
AGCATGGCTCCTAAGAAGAAGAGGAAGGTGATGAGCCAGTTCGACATCCTGTGCAAGACCCCCCAAGGTGCTGGTGC
GGCAGTTCGTGGAGAGATTCCGAGAGGCCAGCGGCGAGAAGATCGCCAGCTGTGCCGCCGAGCTGACCTACCTGTGCT
GGATGATCACCCACAACGGCACCGCCATCAAGAGGGCCACCTTCATGAGCTACAACACCATCATCAGCAACAGCCTGAGC
TTCGACATCGTGAACAAGAGCCTGCAGTTCAAGTACAAGACCCAGAAGGCCACCATCCTGGAGGCCAGCCTGAAGAAGCT
GATCCCCGCCTGGGAGTTCACCATCATCCCTTACAACGGCCAGAAGCACCAGAGCGACATCACCGACATCGTGTCCAGCC
TGCAGCTGCAGTTCGAGAGCAGCGAGGAGGCCGACAAGGGCAACAGCCACAGCAAGAAGATGCTGAAGGCCCTGCTGT
CCGAGGGCGAGAGCATCTGGGAGATCACCGAGAAGATCCTGAACAGCTTCGAGTACACCAGCAGGTTCCACCAAGACCAA
GACCCTGTACCAGTTCCTGTTCCCTGGCCACATTCATCAACTGCGGCAGGTTCCAGCGACATCAAGAACGTGGACCCCAAGA
GCTTCAAGCTGGTGCAGAACAAGTACCTGGGCGTGATCATTAGTGCCTGGTGACCGAGACCAAGACAAGCGTGTCCAG
GCACATCTACTTTTTTCAGCGCCAGAGGCGAGGATCGACCCCTGGTGTACCTGGACGAGTTCCTGAGGAACAGCGAGCCC
GTGCTGAAGAGAGTGAACAGGACCGGCAACAGCAGCAGCAACAAGCAGGAGTACCAGCTGCTGAAGGACAACCTGGTG
CGCAGCTACAACAAGGCCCTGAAGAAGAACGCCCTACCCCTCTTCGCTATCAAGAACGGCCCTAAGAGCCACATCGG
CAGGCACCTGATGACCAGCTTTCTGAGCATGAAGGGCCTGACCGAGCTGACAAACGTGGTGGGCAACTGGAGCGACAAG
AGGGCCTCCGCCGTGGCCAGGACCACCTACACCCACCAGATCACCGCCATCCCCGACCACTACTTCGCCCTGGTGTCCA
GGTACTACGCCTACGACCCCATCAGCAAGGAGATGATCGCCCTGAAGGACGAGACCAACCCCATCGAGGAGTGGCAGCA
CATCGAGCAGCTGAAGGGCAGCGCCGAGGGCAGCATCAGATACCCCGCCTGGAACGGCATCATCAGCCAGGAGGTGCT
GGACTACCTGAGCAGCTACATCAACAGGCGGATCGGAAGCGGAGCTACTAACTTCAGCCTGCTGAAGCAGGCTGGAGAC
GTGGAGGAGAACCCTGGACCTATGAGACCTTTCCAGCTGGATTTGCTCTTCTCTGCTTCTTCTCTCAGTCAAGGtaggaa
cccgcgccggaggtggagcaaatgctgg.

Other mice used are: Gad2-IRES-Cre (JAX stock #010802)⁴², RC::FLTG (JAX stock #026932)⁴³, and PV-Cre (JAX stock #017320), and C57BL/6J wildtype mice (JAX stock #000664). All mice were maintained and conducted in accordance with NIH guidelines on the care and use of animals approved by the University of Utah IACUC committee.

Immunohistochemistry: Mice were transcardially perfused with 1X PBS followed by 20mL of 4% PFA prior to brain extraction. Brains were then stored in 4% PFA overnight and cut in 50µm sections around the hippocampal formation. Sections were incubated in blocking solution (PBS with 3% bovine serum albumin and 0.2% Triton-X100) for 1 hour. Sections were then left overnight with gentle shaking in primary antibody solution, which was prepared with the antibody diluted in PBS with 3% BSA. After 3 washes with PBS, sections were then incubated with secondary antibody solution for 2 hours at room temperature. A final 3 washes were performed with the third wash substituting PBS for a 1:10,000 Hoechst solution prior to mounting.

Antibodies: Primary antibodies were used as follows: rabbit and goat anti-GFP 1:5000 (Abcam), rabbit anti-cFos 1:500 (Santa Cruz Biotech), rabbit anti-Npas4 1:500 (Activity Signaling), rabbit anti-GABA 1:5000 (Sigma), rabbit anti-calretinin 1:2000 (Swant), mouse anti-parvalbumin 1:5000 (Swant), rat anti-somatostatin 1:500 (Chemicon), rabbit anti-calbindin d28k 1:2000 (Swant), rabbit anti-VIP 1:500 (Immunostar), rabbit anti-PV 1:250 (Swant), rabbit anti-RFP 1:1000 (Fisher NBP267102). All secondary antibodies were made in donkey and are from Jackson ImmunoResearch Laboratories Inc. and used at 1:1000.

Stereotaxic surgeries: Adult mice over the age of 8-weeks were injected with Buprenorphine (0.1mg/kg) intraperitoneally an hour prior to operation. Mice were anesthetized in an airtight container ventilated

with oxygen and isoflurane and then readied in stereotaxic surgery area. An injection of lidocaine (2mg/kg) is administered subcutaneously near the intended incision area which is shaved and sterilized. Small holes were drilled in the skull to expose the ROI for viral intracranial injection. The viral payload of 0.5ul was delivered via picospritzer to the desired region with a 0.03 ms pulse at 60 psi. Upon completion, incisions were sealed with VetBond (Amazon) and mice were injected subcutaneously with Rimadyl (5mg/kg Carprofen). Mice were allowed to recover on heating pad and observed for signs of distress before returning to their home cage. Follow-ups continued for three days and consisted of weighing mice and administering Rimadyl to ensure healthy recovery.

Viruses: Adeno-associated viruses (AAVs) were made using standard iodixanol gradient purification and the pHP.eb capsid⁶¹. Viral titers for all viruses used were 10^{11} to 10^{13} . Viruses used are as follows: ConVERGD construct (pHP.eb)hSyn.hM3Dm.Cherry and (AAV8)hSyn.hM3D.HA⁴⁴ provided by Lindsay Schwarz lab; (AAV9)hSyn.DIO.hM3D.mCherry (Addgene 44361); INTR SCT construct (pHP.eb)hSyn.Cre-on/Flp-on.YFP (Addgene 55650)⁶².

Enriched environment: Mice are removed from their home cage and injected with either saline or 1.0mg/kg DCZ then immediately placed into a 40cm x 40cm plastic container with 5 novel objects evenly spaced apart. Mice are allowed to explore freely for 40-minutes prior to perfusion.

Contextual Fear Conditioning: Mice underwent a 3-day contextual fear conditioning protocol. Mice were habituated up for 3 days prior to conditioning day (Day 0) for 1 hour. On conditioning day 0 and after habituation, mice are then injected i.p. with either saline or 1.0mg/kg DCZ 10-minutes prior to being placed into behavior boxes (MedAssociates NIR). Behavior boxes are furnished with distinguishing features such as floor grating, light, scent, roofing, and sound that they will associate with conditioned "Context A". Mice 4 shocks (1mA, 2-second duration, 1-minute isi) are delivered after an initial 3-minute acclimation period. Mice then rest for an additional minute, for a total 8-minute duration conditioning protocol. On Test Day 1, mice are habituated and injected as described earlier. Half of conditioned mice are exposed to conditioned Context A for 8-minutes or "intermediate" Context B (see Figure 3B). Mice are returned to their homecage and mouse room for a minimum of 5 hours before their next context exposure. Mice initially exposed to Context A are now exposed to Context B, and vice versa. This process is repeated for Test Day 1, with "neutral" Context C replacing intermediate Context B

cFos analysis: Confocal images (Zeiss 980) were taken at 10 and 20X objective as Z-stacks and tiles, exported as OME-TIFF files, and analyzed in FIJI blind to condition or genotype. For image analysis in FIJI, all images were z-projected for maximum intensity, despeckled, and split channels. ROIs are drawn for the CA3 region in the virus mCherry channel, and only the pyramidal region in the cFos channel to measure area. The cFos channel background is subtracted by a rolling ball radius value of 15, then thresholded to three times the inflection point of the signal histogram values for a consistent signal-to-noise ratio. Particles are analyzed with a circularity range of 0-1.0 and size of 30-infinity. The counts are then reviewed and saved as ROI to overlap with the virus mCherry channel and count co-expressing cells. The percent signal area within and outside of the ROI of the mCherry channel is recorded, while cell bodies are individually counted by hand-drawn ROIs. Then the number of overlapping ROIs for mCherry cell bodies and cFos counts are recorded. Counts are verified by merging channels and examining merged fluorescence.

Fluorescent in situ hybridization (FISH): Fluorescent in situ hybridization chain reaction (HCR) HCR was performed as previously described⁶³. Mouse brains were cryo-sectioned to 30- μ m slices, mounted on slides, fixed [4% paraformaldehyde (PFA)] and washed in PBS. Before processing samples according to protocol HCR v3.0 (Invitrogen), slices were treated with 1mg/ml proteinase K-treated (TE buffer) and equilibrated in SSC buffer. Custom HCR probes were designed and generated by Molecular Instruments for Gad1, Kirrel3, and GFP. After nuclear staining with Hoechst in PBS, coverslips were mounted in Fluoromount-G (Southern Biotech catalog #0100-01) and imaged (Zeiss LSM 710). For protein amplification, samples are incubated in freshly prepared 4% PFA made with DEPC water and PBS for 10 minutes, then washed in 1X DEPC treated PBS (pH 7.4) for 5 minutes at room

temperature three times. Samples are then blocked for 1 hour at room temperature (100uL/slide applied over top of sections) inside humidified chamber (does not need to be RNase free) and 75μL primary antibody solution (anti-mCherry 1:1000) is added per slide and covered with an RNase free coverslip. Samples are incubated for 1 hour at room temperature and then overnight at 4°C in humidified chamber. The next day, samples are washed in 1X DEPC treated PBST (pH 7.4) for 5 minutes at room temperature for a total of three washes before adding the 2° antibody at 1:1000 then incubating for 1 hour at room temperature inside humidified chamber. Samples are washed in 1X DEPC treated PBST (pH 7.4) for 5 minutes at room temperature another three times and mounted by adding 250μL antifade mounting reagent on top of sample, then cover-slipped for microscopy.

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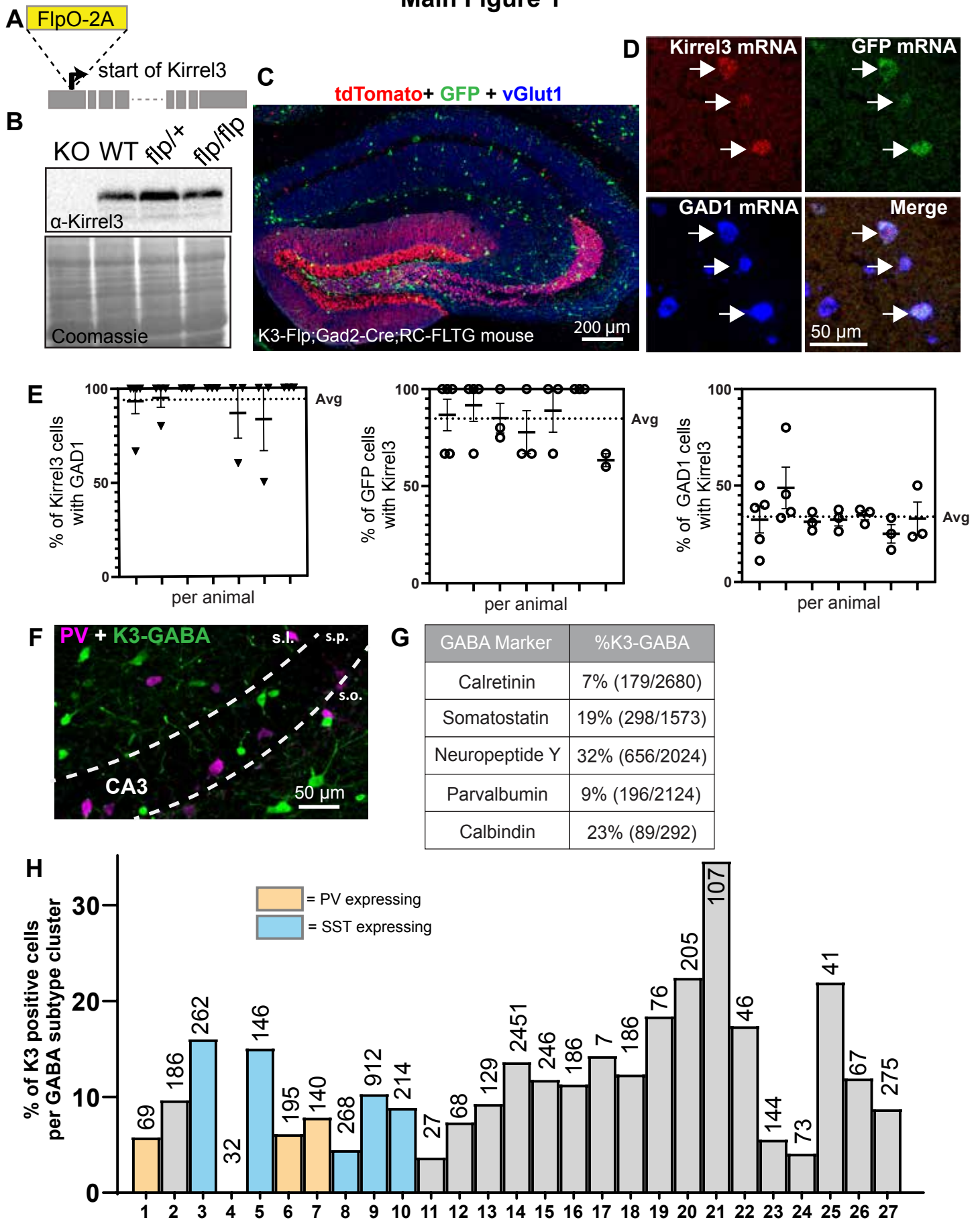
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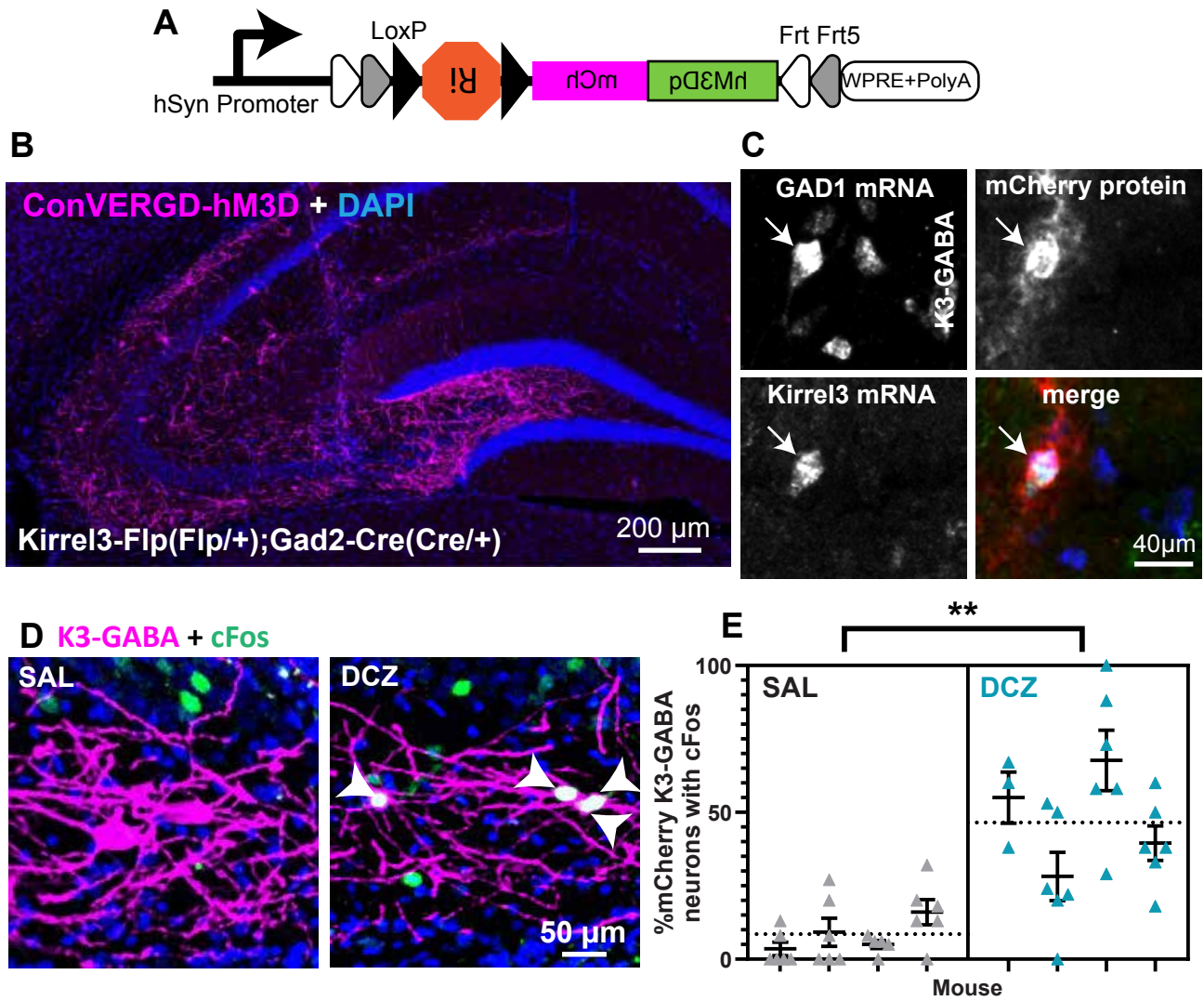
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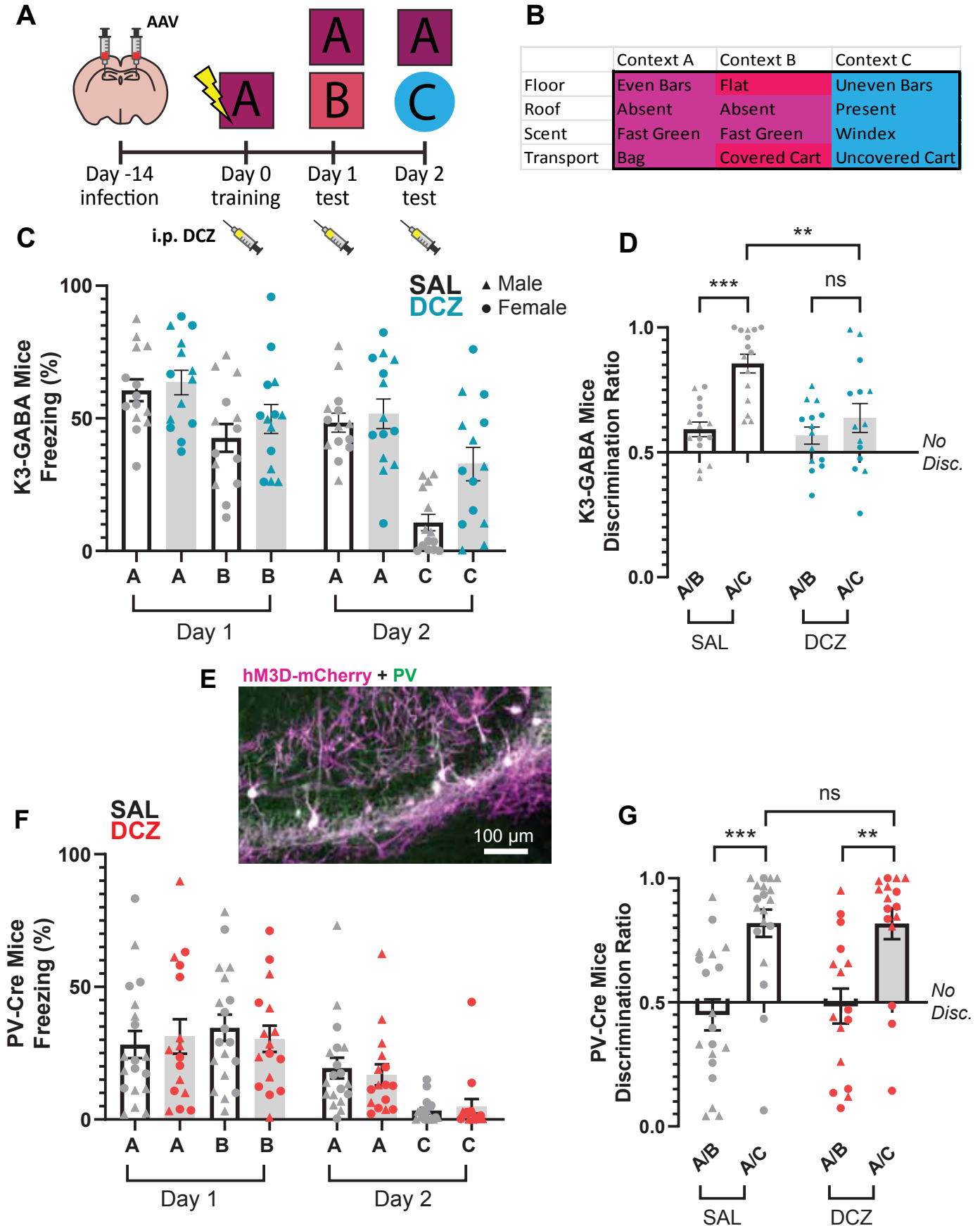
Main Figure 1



Main Figure 2

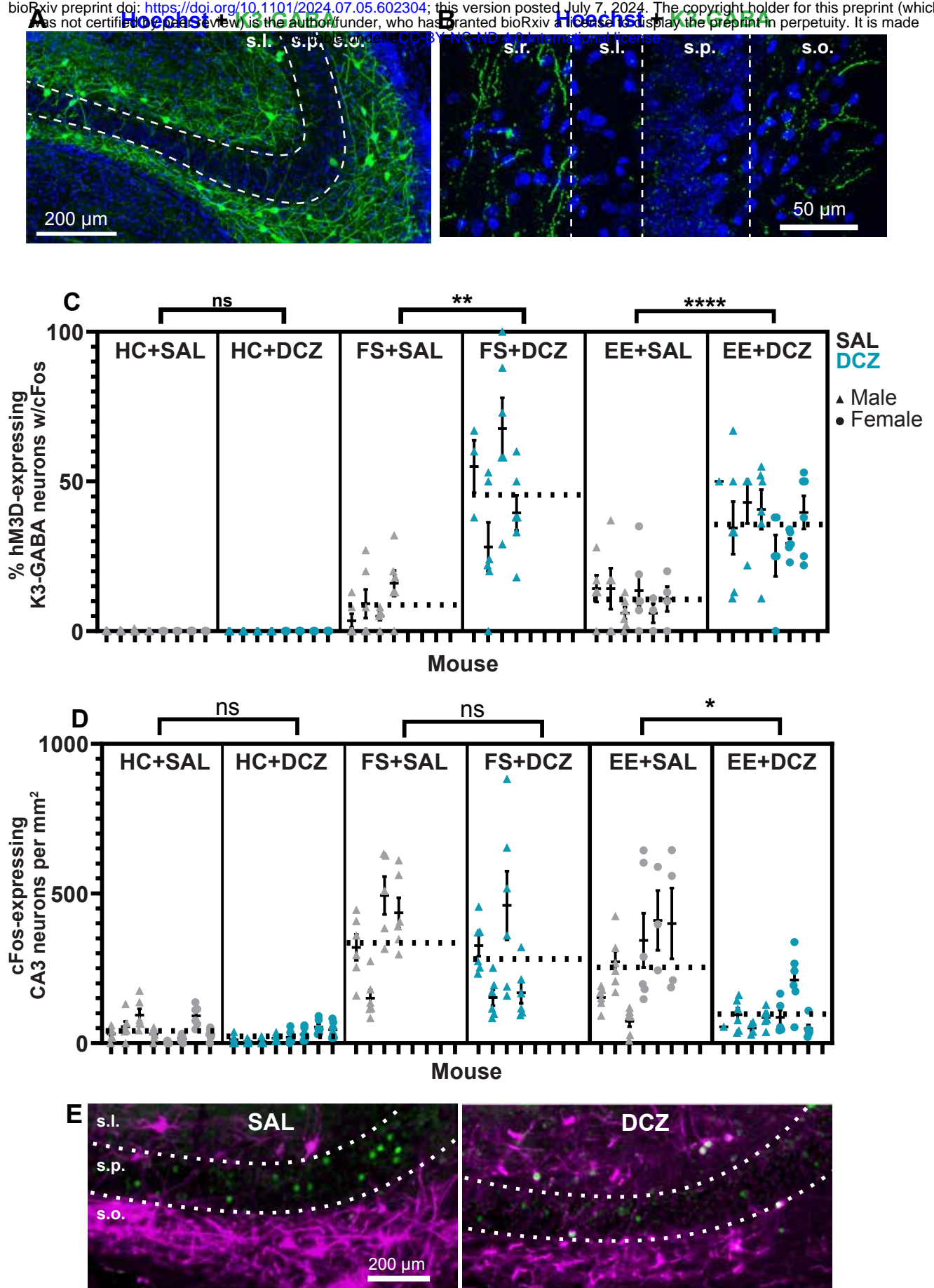


Main Figure 3



Main Figure 4

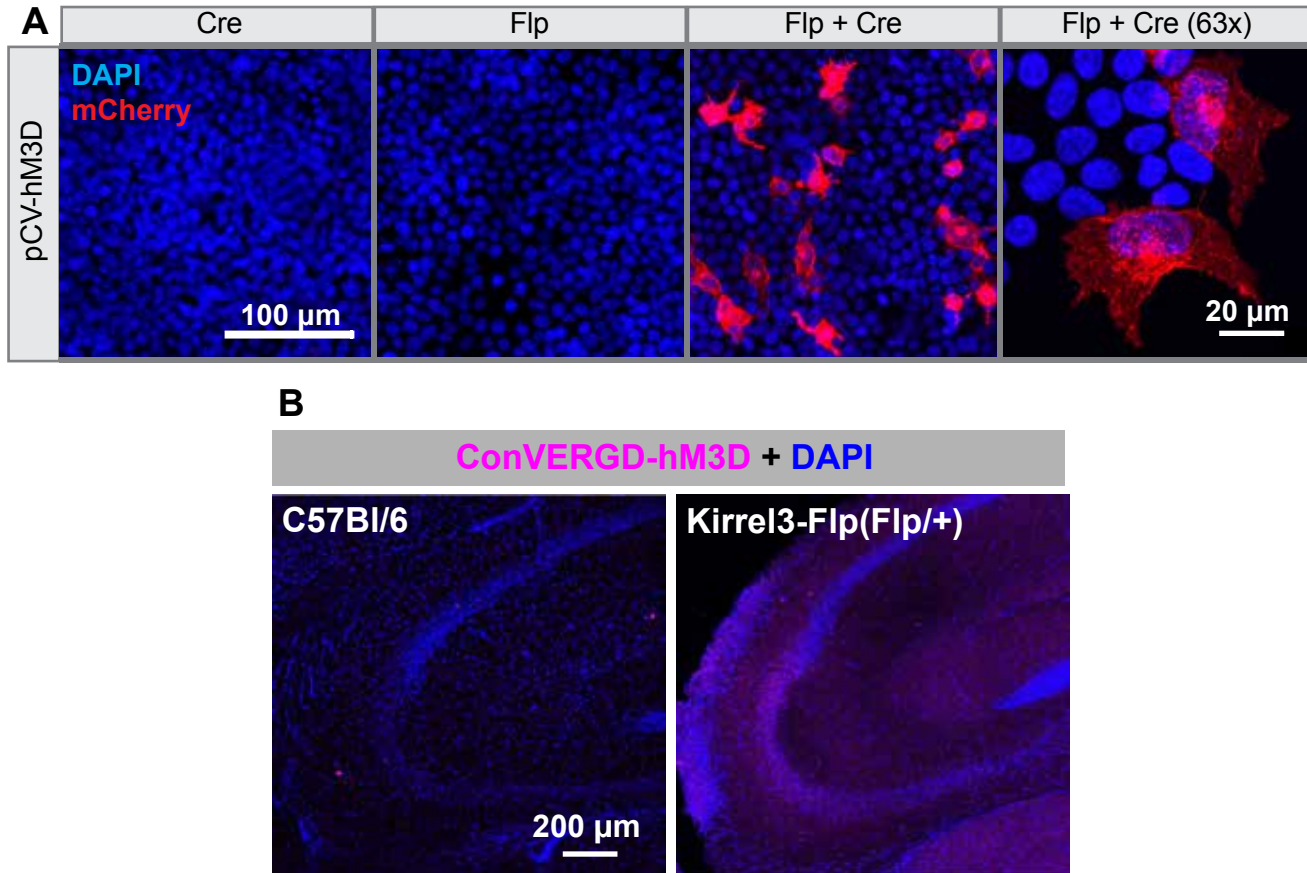
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ID	NAME
1	Pvalb.C1ql1
2	Id2.Sfrp2
3	Sst.Atp2b4
4	Slc17a8_Sst.Ncam2
5	Sst.Spon1
6	Pvalb.Tac1
7	Pvalb.Gfra1
8	Sst.Rbp4
9	Sst.Grm1
10	Sst.Pcdh11x
11	Cplx3.Rxfp1
12	Cplx3.Tox
13	Id2.Tac1
14	Id2.Bcl11b
15	Id2.Prir
16	Htr3a.Nnat
17	Chat_Htr3a.Chat
18	Htr3a.Phlda1
19	Htr3a.Efna5
20	Htr3a.Cpne5
21	Htr3a.Ecel1
22	Htr3a.Ibsp
23	Htr3a.Krt73
24	Slc17a8_Htr3a.Kctd12
25	Htr3a.Chrm2
26	Htr3a.Htr1b
27	Htr3a.Sema3c

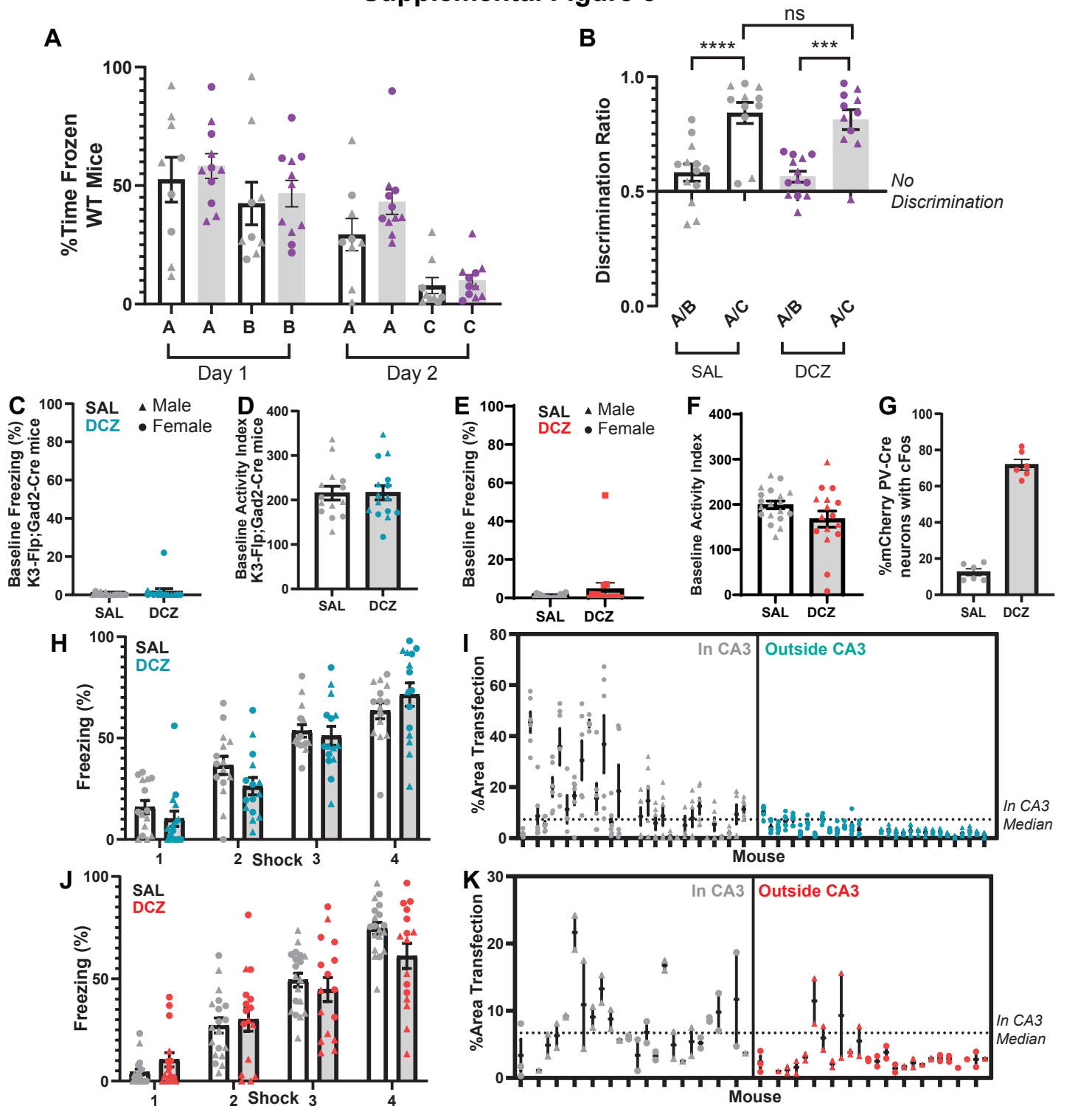
Supplemental Table 1: Key for neuron groups in Figure 1H

Supplemental Figure 2



Supplemental Figure 2. A) Representative images of HEK293 cells co-transfected with hSyn-ConVERGD-hM3D-mCherry (pCV-hM3D) and either Cre, Flp, or Flp and Cre plasmids. Far right column: representative image at 63x, all other images taken at 20x. B) Representative hippocampal CA3 images of hSyn-ConVERGD-hM3D-mCherry AAV infection in wildtype C57Bl/6J and Kirrel3-Flp mice show no expression of the DREADD as expected.

Supplemental Figure 3



Supplemental Figure 3. A) Mouse behavior plotted as percent time spent freezing for wild-type (WT) mice when placed in indicated contexts after saline (gray) or DCZ (purple) injection. B) Time spent freezing (%) plotted as a discrimination ratio relative to context A for WT mice. 0.5 equals no discrimination. No differences between saline or DCZ, $n = 13$ saline, 11 DCZ, ANOVA with multiple comparisons. C, D) Baseline freezing (C) and activity index (D) prior to foot shock for K3-Flp;Gad2-Cre mice expressing hM3D-mCherry. E, F) Baseline freezing (E) and activity index (F) prior to foot shock for PV-Cre mice expressing hM3D-mCherry. G) % of hM3D-mCherry infected PV neurons that express cFos after foot shock, $n = 6$ per group. H) % time spent freezing for one minute after indicated shock for K3-Flp;Gad2-Cre mice expressing hM3D-mCherry. I) Plot indicating the percent area of hM3D-mCherry signal in CA3 versus outside CA3 to measure transfection efficacy for K3-Flp;Gad2-Cre mice. Each dot represents a brain section. The median value for "in CA3" is indicated by the dotted line. J) % time spent freezing for one minute after indicated shock for PV-Cre mice expressing hM3D-mCherry. K) Same as I but for PV-Cre mice. Error bars represent SEM.