

1 **Choosing memory retrieval strategies:**
2 **a critical role for inhibition in the dentate gyrus**

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1 **Abstract**

2

3 Remembering the location of food is essential for survival. Rodents and humans employ
4 mainly hippocampus-dependent spatial strategies, but when being stressed they shift to
5 striatum-mediated stimulus-based strategies. To investigate underlying brain circuits, we
6 tested mice with a heightened stress susceptibility due to a lack of the GABA-synthetizing
7 enzyme GAD65 (GAD65^{-/-} mice) in a dual solution task. Here, GAD65^{-/-} mice preferred to
8 locate a food reward in an open field via a proximal cue, while their wildtype littermates
9 preferred a spatial strategy. The analysis of cFos co-activation across brain regions and of
10 stress-induced mRNA expression changes of GAD65 pointed towards the hippocampal dorsal
11 dentate gyrus (dDG) as a central structure for mediating stress effects on strategy choices via
12 GAD65. Reducing the GAD65 expression locally in the dDG by a shRNA mediated knock down
13 was sufficient to replicate the phenotype of the global GAD65 knock out and to increase dDG
14 excitability. Using DREADD vectors to specifically interfere with dDG circuit activity during
15 dual solution retrieval but not learning confirmed that the dDG modulates strategy choices
16 and that a balanced excitability of this structure is necessary to establish spatial strategy
17 preference. These data highlight the dDG as a critical hub for choosing between spatial and
18 non-spatial foraging strategies.

19

20 **Keywords:** spatial learning, stimulus-based learning, cognitive flexibility, dentate gyrus,
21 GAD65, inhibition

22

23 **Abbreviations:** ACC: anterior cingulate cortex; BLA: basolateral amygdala; CA: Cornu
24 ammonis; CeA: central amygdala; CNO: clozapine-N-oxide; CORT: corticosterone; dDG:
25 dorsal dentate gyrus; DREADD: designer receptors exclusively activated by designer drugs;
26 fEPSP: field postsynaptic potentials; GAD65: glutamic acid decarboxylating enzyme, 65 kDa
27 isoform; GR: glucocorticoid receptors; LA: lateral amygdala; mPFC: medial prefrontal cortex;
28 MR: mineralocorticoid receptors; PL/IL: Pre-/ Infralimbic cortex

1 **1. Introduction**

2

3 Remembering the location of a food source is indispensable for survival. Navigating to such a
4 location can be achieved via different neuronal systems dependent on the underlying
5 information processing and operations performed. Rodents and humans deploy two major
6 strategies: They can form an internal map by learning spatial relationships between
7 landmarks or they use a stimulus response strategy, where a sequence of motor events
8 needs to be performed upon specific cues, e.g. turn right at the gas station and then left at
9 the bakery (Packard and Goodman, 2012). While the spatial strategy involves the
10 hippocampus and medial temporal lobe structures, the cue response strategy activates the
11 dorsal striatum (Goodman, 2021; Packard and Goodman, 2013). Moreover, studies in human
12 and rodents demonstrate that prefrontal cortical areas are involved in processing stimulus-
13 reward associations and in choosing specific actions upon these stimuli. Together with
14 cingulate cortical areas, they also process cue- and spatial-related information such as the
15 position of landmarks or of the own position in space. By forming a network with the
16 hippocampus and striatum, prefrontal and cingulate cortical areas mediate choosing spatial
17 or cue-based response strategies when navigating to a reward (Cowen et al., 2012;
18 Kennerley et al., 2006; Rich and Shapiro, 2009; Rudebeck and Izquierdo, 2022).

19 While a spatial strategy allows for creating shortcuts and is more flexible, individuals under
20 stress favor a cue response learning strategy. To investigate this shift in strategy preferences
21 Schwabe et al. (Schwabe et al., 2008) developed a paradigm where mice had to learn the
22 location of an escape hole on a hole board by either using distal spatial cues for navigation
23 or respond to a bottle located next to the escape hole. While control mice relied on a spatial
24 reward-location strategy, chronically stressed mice preferred responding to the proximal
25 cue. A similar dual solution setup in humans using a computer-based memory game
26 demonstrated that participants with a higher self-reported stress score shifted towards a
27 stimulus-response strategy as well. In a follow-up study, the authors could demonstrate that
28 acute stress, the administration of the stress hormone corticosterone (CORT) and the
29 activation of mineralocorticoid receptors (MR) induce a similar switch of memory systems
30 (Schwabe et al., 2010a, 2010b). However, stress and stress hormones have detrimental
31 effects on hippocampus-dependent learning and long-term potentiation (Roosendaal et al.,
32 2003; Sandi and Pinelo-Nava, 2007) and may already affect spatial strategy preference by

1 impairing spatial learning but not strategy choice during retrieval. Moreover, detailed
2 knowledge about possible molecular mechanisms of stress-induced shifts and their location
3 within the strategy choice neuronal network remains elusive to date.

4 In our current study we investigated the involved neuronal networks and their potential
5 molecular regulation, focusing on the role of GABAergic interneurons and the key enzyme in
6 GABA synthesis, glutamic acid decarboxylase (GAD). Within the cortex, hippocampus and
7 striatum neuronal activity is shaped by local inhibitory interneurons that use GABA as a
8 neurotransmitter. The GABA-synthetizing enzyme GAD65 critically determines the activity-
9 dependent GABA pools and expression levels of this enzyme are modulated by stress
10 exposure as well as corticosterone (Albrecht et al., 2021b; Bergado-Acosta et al., 2008;
11 Bowers et al., 1998). Mice with a total knock out of GAD65 (GAD65^{-/-}) show increased
12 anxiety, avoidance and hyperarousal as well as a generalized fear memory and deficits in
13 fear extinction (Müller et al., 2015). They provide an interesting model to further elucidate
14 molecular mechanisms and a potential contribution of the inhibitory system to strategy
15 choice in a dual solution paradigm. We therefore compared strategy preferences of GAD65-
16 ^{-/-} mice and their wildtype littermates in a dual solution task based on locating a food reward
17 in the open field. GAD65^{-/-} mice did not show a spatial strategy preference, even though
18 spatial learning was intact in these animals. Analysis of neuronal co-activation of brain
19 regions via cFos and the analysis of stress-induced expression changes of GAD65 pointed
20 towards the dorsal dentate gyrus (dDG) as a potential hub for mediating strategy choices via
21 GAD65. Accordingly, the local knock down of GAD65 in the dDG reproduced a phenotype
22 reminiscent of the total knock out in the dual solution task and induced an increased
23 excitability of the DG circuit. Using chemogenetic manipulation of the dDG network to
24 dissect effects on learning vs. retrieval confirmed the importance of DG excitation/ inhibition
25 balance for strategy choices during retrieval.

26

27

28 **2. Material and methods**

29

30 **2.1 Animals**

31 All mice were bred and raised in the animal facility at the Institute of Biology, Otto-von-
32 Guericke University Magdeburg. The animals were kept in groups of 2–6 on an inverse 12 h

1 light/dark cycle (lights on at 7 pm with a 30 min dawn phase) with food and water *ad*
2 *libitum*. Animal housing and animal experiments were in accordance with European
3 regulations for animal experiments and approved by the Landesverwaltungsamt Saxony-
4 Anhalt (Permission Nr. 42502-2-1177 and -1517). GAD65 knock out mice (GAD65^{-/-}) and
5 their wildtype littermates (GAD65^{+/+}) were obtained from a heterozygous breeding and
6 genotyped with allele-specific polymerase chain reaction as described previously (Stork et
7 al., 2000). Further mouse lines engaged were wild type C57BL/6BomTac (M&B Taconic,
8 Germany), SST-Cre (B6(Cg)- Ssttm2.1(cre)Zjh) and PV-Cre (B6;129P2-Pvalbtm1(cre)Arbr/J)
9 mice. Founders of Cre lines were obtained from Jackson laboratories and maintained in a
10 homozygous breeding for SST-Cre and a heterozygous breeding for PV-Cre mice with
11 genotypes being determined with allele-specific polymerase chain reaction using the
12 supplier's recommendations. From all lines young adult male mice, aged 10-16 weeks, were
13 used for experiments. All experiments were performed during the dark, active phase of the
14 animals and handling was done under red light illumination.

15

16 **2.2 Stress protocol**

17 Animals were randomly assigned to restrained stress or control group. Restrained animals
18 were placed in a plastic cylinder (3 cm diameter x 11.5 cm length) with holes for breathing at
19 the top for 10 min that prevented side movement and restricted movements forth and back.
20 Home cage controls were handled in a fresh cage in parallel. Afterwards all mice were
21 returned to their home cage for 24 h until preparation of trunk blood and brain tissue after
22 decapitation under brief inhalation anesthesia with isoflurane.

23

24 **2.3 Corticosterone ELISA**

25 Trunk blood was collected and allowed to clot at room temperature for 30 min. Clotted
26 material was removed by centrifugation at 3000 rpm for 15 min. Serum was removed
27 immediately and stored at -80°C until further use. Corticosterone serum levels were
28 measured with the IBL Corticosterone ELISA kit (IBL International, Hamburg, Germany)
29 according to the manufacturer's instructions.

30

31 **2.4 Dual solution task**

1 Five days before the training of a reward location task, animals were food restricted until
2 85% of the starting body weight was reached. Everyday each mouse received 4 choco
3 krispies (Kellogg Deutschland, Hamburg, Germany) in a plastic lab dish (8.5 cm diameter)
4 within its home cage. For habituation to the training environment, mice were placed for 20
5 min in an open field (50 cm x 50 cm, 10 lux low light illumination) equipped with patterned
6 paper cuts as distal cues at the wall. After approximately 45 min break in the homecage,
7 mice were reintroduced to the open field, this time equipped with 4 plastic lab dishes (8.5
8 cm diameter) in the center of each quadrant. One of the dishes contained a wooden toy
9 block (2 cm x 2 cm and 5 cm high) as a proximal cue and a choco krispie placed right next to
10 it. The other 3 dishes contained 1 choco krispie, but access was restricted by a lid with holes
11 for olfactory information. The position of the reward and the proximal cue was randomized
12 across batches but kept consistent over training trials. In each training trial, a mouse was
13 placed randomly in one of the 2 corners equidistant from the dish with the krispie for start
14 and were allowed to explore the arena freely for 4 min. The latency to eat the choco krispie
15 for each trial was assessed as an indicator for learning progress. Each mouse went through 6
16 training trials with intertrial intervals of approx. 20 min in the home cage.

17 Animals performed a retrieval trial 24 h after training for which the proximal cue was placed
18 diagonally opposite of the dish which contained the reward previously. Choco krispies were
19 absent in all dishes during retrieval. Again, the respective mouse started the trial in a corner
20 and the first head contact with a dish in a 4 min session was noted by an experimenter blind
21 to genotype and manipulation: a “spatial” choice was made when the mouse entered the
22 previously rewarded dish, a “cued” choice was noted when the dish with the proximal cue
23 was chosen and when any of the other 2 dishes was entered, the response was “neither”. If
24 a mouse did not visit any dish within 4 min, it was classified as “no response”.

25 Spatial learning abilities of GAD65^{-/-} mice were tested in separate groups, first by applying
26 training and testing without any proximal cue and, second, by presenting a proximal cue
27 during training but not retrieval. As a measure for spatial memory the time spent at the
28 target position during retrieval was used in these modified paradigms.

29

30 **2.5 cFos immunohistochemistry**

31 A subgroup of GAD65^{-/-} mice and their GAD65^{+/+} littermates received transcatheter
32 perfusion with phosphate buffered 4% paraformaldehyde solution at pH 6.8, as previously

1 described (Raza et al., 2017). After postfixation overnight in the same fixative and overnight
2 immersion in wt/vol 20% sucrose at 4°C, brains were cut into 30 µm thick, serial coronar
3 sections at the level of the PFC, posterior ACC/ dorsal striatum and dorsal hippocampus/
4 amygdala, stored as free-floating sections in phosphate buffered saline at 4°C and then
5 incubated with a rabbit antibody against cFos (1:1000, Cell signaling #2250, Danvers, MA,
6 USA) for 48 h. As secondary antibody biotinylated goat anti-rabbit (1:200, Jackson
7 ImmunoResearch, Ely, UK) in phosphate buffer (PB) with 0.2% Triton (PBT) was used and
8 labelled with Cy2 via Streptavidin (1:1000, Jackson ImmunoResearch, Ely, UK) in PB. Nuclei
9 were visualized with DAPI (300 nM, Thermofischer, Waltham, MA, USA). Immunostainings
10 were imaged using a DMI6000 epifluorescence microscope (Leica, Wetzlar, Germany) in
11 both hemispheres from 2 slices per animal and region. cFos-positive cells were counted
12 manually within the selected brain areas and cell density was normalized to area size
13 measured with imageJ software. The cell counts per target area were averaged for each
14 individual animal and used for statistical comparison between groups.

15

16 **2.6 qPCR**

17 GAD65^{-/-} and GAD65^{+/+} mice were sacrificed 24 h after restraint stress by decapitation
18 under deep anesthesia with isoflurane. Brains were quickly removed and snap frozen in
19 methylbutane cooled by liquid nitrogen and stored at -80°C. Brains were mounted at a
20 cryostat at -20°C and coronar sections were removed until the start of the dorsal striatum
21 (1.34 mm from Bregma). Striatal tissue was harvested with a stainless-steel sample corer
22 (diameter 0.5 mm, Fine Science Tools, Heidelberg, Germany). Then further sections were
23 removed until the dorsal pole of the hippocampus emerged (starting at -1.34 mm from
24 Bregma) and dDG tissue was harvested with a 0.35 mm diameter sample corer (Fine Science
25 Tools, Heidelberg, Germany). Lysis of tissue samples and isolation of total RNA was
26 performed with the RNeasy Micro Plus kit (Qiagen, Hilden, Germany) according to
27 manufacturer's instructions. For first-strand cDNA synthesis the Takara PrimeScript RT-PCR
28 Kit (Takara Bio, Shiga, Japan) was used with 50 µM Oligo (dT) and 50 µM random hexamer
29 first strand primers according to the manufacturer's instructions.

30 Real-time PCR was performed with a 1:5 dilution of cDNA using QuantStudio3 real-time PCR
31 apparatus (Life Technologies) and TaqMan® reagents with predesigned assays for the target
32 genes GAD65 (alias GAD2, assay ID: Mm00484623_m1), GAD67 (alias GAD1, assay ID:

1 Mm00725661_s1), MR (alias NR3C2, assay ID: Mm01241596_m1) and glucocorticoid
2 receptors (GR, alias NR3C1, assay ID: Mm00433832_m1) as well as for the housekeeping
3 gene Glycerinaldehyd-3-phosphat-Dehydrogenase (GAPDH; endogenous control; Life
4 Technologies) that was labeled with another fluorescent dye, allowing for quantitative
5 evaluation. Multiplex PCR samples were run in triplicates with 50 cycles of 15 s at 95°C and 1
6 min at 60°C, preceded by a 2 min 50°C decontamination step with Uracil-N-glycosidase and
7 initial denaturation at 95°C for 10 min. Gene expression relative to control groups (relative
8 quantification, RQ) was calculated using the ddCT method by normalizing the mean cycle
9 threshold (CT) of each triplicate assay first to the overall content of cDNA using GAPDH as an
10 internal control ($dCT; dCT[\text{target gene}] = (CT [\text{target gene}] - (CT [\text{GAPDH}]))$) and then to the
11 control group with $ddCT = dCT(\text{sample}) - \text{mean } dCT (\text{control group})$. RQ values were
12 obtained by calculating $RQ\% = (2^{-ddCT}) \times 100$.

13

14

15 **2.7 Viral injections**

16 Mice under inhalation anesthesia with isoflurane (1.5-2.5 Vol% in O₂) were placed on a
17 stereotaxic frame (World Precision Instruments, Berlin, Germany). 33G injection needles
18 attached to 10 µl NanoFil microsyringes (World Precision Instruments, Berlin, Germany)
19 were placed bilaterally in drillholes at -1.94 mm anterioposterior, ±1.3 mm mediolateral
20 from Bregma and were lowered -1.7 mm dorsoventral from the brain surface into the dorsal
21 dentate gyrus. Viral vectors were injected bilaterally at a volume of 1 µl/side at a flow rate of
22 0.1 µl/min using a digital microsyringe pump (World Precision Instruments, Berlin,
23 Germany).

24 To silence GAD65, a small hairpin RNA construct was generated against the oligonucleotide
25 sequence 5'-GCATGCTTCCTACCTCTTTCA-3' (corresponding to NM 008078, base pairs 1599-
26 1619 in mouse and NM_012563.1, base pairs 1337-1357 in rat) and for control a random
27 sequence shRNA (5'-TCGTCATGACGTGCATAGG -3') were generated and cloned for
28 production of lentiviral particles as described and validated in Tripathi et al. (2021). Lentiviral
29 particles were injected with a concentration of 10⁹ particles/µl. Virus spread was determined
30 by the expression of the marker GFP (see Suppl. Fig. 1). The knock down efficiency for
31 GAD65 in the mouse hippocampus was determined by western blot protein expression (see
32 Suppl. Fig. S2).

1
2 For chemogenetic manipulation of DG circuits designer receptors exclusively activated by
3 designer drugs (DREADDs)-based tools were employed. The following DREADD adeno-
4 associated viruses (AAV), which were gifts from Bryan Roth and obtained via Addgene
5 (Watertown, MA, USA), were used: pAAV-CaMKIIa-hM4D(Gi)-mCherry (Addgene viral prep
6 #50477-AAV8; <http://n2t.net/addgene:50477>; RRID:Addgene_50477) and pAAV-CaMKIIa-
7 GFP (Addgene viral prep #50469-AAV8; <http://n2t.net/addgene:50469>;
8 RRID:Addgene_50469) were used for silencing excitatory cells in the DG. hSyn-DIO-hM4Di-
9 mCherry (Addgene viral prep #44362; <http://n2t.net/addgene:44362>;
10 RRID:Addgene_44362), AAV hSyn-DIOhM3Dq-mCherry (Addgene viral prep #44361;
11 <http://n2t.net/addgene:44361>; RRID:Addgene_44361) and AAV hSyn-DIO-mCherry
12 (Addgene viral prep #50459; <http://n2t.net/addgene:50459> ; RRID:Addgene_50459) were
13 used for chemogenetically manipulating interneurons in the respective cre-transgenic lines
14 (Raza et al., 2017). AAVs were injected at $> 5 \times 10^{12}$ particles/ μ l.

15 For activation of chemogenetic hM4Di and hM3Dq receptors, animals received an i.p.
16 injection of 10 mg kg⁻¹ clozapine-N-oxide (Enzo Life Sciences, Germany) in physiological
17 saline 1 h before either retrieval or training of the dual solution task.

18 For histological verification of viral vector expression, animals were transcardially perfused
19 at the end of the experiment and 3-4 alternating 30 μ m sections of the dorsal hippocampus
20 (from Bregma AP: -1.58 to AP: -2.18 mm) were imaged using a DMI6000 epifluorescence
21 microscope (Leica, Wetzlar, Germany). As observed previously, marker expression is
22 expected in the dendrites and axons of dDG granule cell (Raza et al., 2017). Only mice with a
23 correct bilateral marker expression in the molecular layer and the mossy fibers of the dDG
24 were therefore considered for behavioral experiment.

25

26 **2.8 Slice electrophysiology**

27 14 to 16 days after the shRNA-mediated viral knock down of GAD65 in the dorsal DG, mice
28 were decapitated under deep anesthesia with isoflurane and their brains were quickly
29 removed and submerged into cold ($\sim 4^{\circ}\text{C}$) carbogenated (5% CO₂/95% O₂) artificial
30 cerebrospinal fluid (aCSF, composition in mM: 129 NaCl, 21 NaHCO₃, 3 KCl, 1.6 CaCl₂, 1.8
31 MgSO₄, 1.25 NaH₂PO₄, 10 glucose). Parasagittal brain slices including the transverse-like
32 dorsal hippocampal sections ($\sim 400 \mu$ m) were cut from the dorsal pole on an angled platform

1 of approx. 12°. Three to four slices per hemisphere were transferred to an interface chamber
2 perfused with aCSF at $30 \pm 0.5^\circ\text{C}$ (flow rate: 1.8 ± 0.2 mL/min, pH 7.4, osmolarity ca. 300
3 mOsmol/kg) and incubated at least for 90 min before extracellular field recordings were
4 performed.

5 For dDG electrophysiology (Fig. 3Ci), a bipolar tungsten wire stimulation electrode (with
6 exposed tips of approx. 20 μm , tip separations of approx. 75 μm , electrode resistance in
7 ACSF: approx. 100 k Ω) was placed at the perforant path (PP) presumably stimulating both
8 medial and lateral fibers. Population spike (PS) and positive-going field postsynaptic
9 potential (fEPSP) were recorded with a borosilicate glass electrode filled with aCSF (1-2 M Ω)
10 from granule cell layer of DG at 70–100 μm depth as previously described (Albrecht et al.,
11 2016). Stable PP-induced responses were verified for ~ 10 min (0.033 Hz with pulse durations
12 of 100 μs) before obtaining an input–output (I–O) curve (Stimulation intensities in μM : 10, 20,
13 30, 40, 50, 75, 100, 150, 200). Paired pulse (PP) responses (Stimulus intervals in ms: 10, 25,
14 50, 100, 250, 500) were recorded using the ~ 50 –60 % of the maximum PS amplitude.

15 Signals were sampled at a frequency of 5 kHz, pre-amplified with a custom-made amplifier,
16 low-pass filtered at 3 kHz and stored on a computer hard disc for offline analysis. Evoked
17 potentials PS and fEPSP were analyzed using MATLAB (MathWorks, Natick, MA) and Spike2-
18 based analysis tools (CED, Cambridge). For the PP-DG electrophysiology, the slope of
19 positive-going fPSP was analyzed by calculating the slope (V/s) between the 20 and 80% of
20 the fEPSP amplitudes. The DG PS amplitude was measured by calculating the average of
21 peak-to-peak amplitude (mV) of descending and ascending phase of PS. For calculating the
22 paired-pulse ratio, the magnitude of the second evoked potential was divided by the first
23 one. For methodological details on mossy fiber-to-CA3 synapse or Schaffer collateral-to-CA1
24 synapse electrophysiology see supplementary material (Suppl. Fig. 3 and 4).

25

26

27 **2.9 Statistics**

28 For all experiments, normal distribution of the data was assessed with Shapiro-Wilk's test.
29 Equality of variances was tested using Levene's test. Comparisons of two groups were made
30 with Student's t-test or with Mann-Whitney U-test, in case of non-normal distribution of the
31 data. In case of unequal variances, a Welch test was applied. For comparing measures over
32 time (e.g. during dual solution learning), stimulation intensities or intervals (during slice

1 recordings), repeated measure ANOVA was applied, if necessary with Greenhouse-Geißer
2 corrections for sphericity issues. Correlation of cFos intensities between areas were assessed
3 using Pearson's correlation coefficient. For comparing the distribution of strategy choices
4 between groups the likelihood-ratio chi-square (X^2) test, optimized for small sample sizes,
5 was used. For statistical analysis of the data GraphPad Prism software (version 9, SD,
6 California) and SPSS (version 28, IBM, New York) was used. Differences were considered
7 statistically significant if p-value (p) < 0.05.

8
9

10 **3. Results**

11

12 **3.1 GAD65^{-/-} mice preferred a cue-based strategy in the dual solution reward memory** 13 **task.**

14 To assess the stress susceptibility of GAD65^{-/-} mice a brief submission to 10 min restrained
15 stress was performed and CORT serum levels were determined 24 h in trunk blood
16 (GAD65^{+/+} control: n=7; GAD65^{+/+} restrained: n=6; GAD65^{-/-} control: n=6; GAD65^{-/-}
17 restrained: n=5). CORT levels were elevated in GAD65^{-/-} irrespective of stress (Fig. 1A; Mann-
18 Whitney-U-test for main effects in non-normal distributed data set: U=106, p=0.047). Paired
19 comparisons revealed that GAD65^{-/-} mice had higher serum levels of CORT than their
20 wildtype littermates under control conditions (control GAD65^{+/+} vs. ^{-/-}: U=38, p=0.014),
21 while 24 h after restrained stress no genotype differences were observed (restrained
22 GAD65^{+/+} vs. ^{-/-}: U=18, p=0.662).

23 Both, GAD65^{-/-} (n=18) and GAD65^{+/+} (n=16) mice learned the food location well during
24 training in the dual solution task, with a trend to lower latencies in wildtype mice (Fig. 1 B;
25 Repeated Measure ANOVA with Greenhouse-Geißer correction: trials F(2.466,
26 78.913)=56.333, p<0.001; trials x genotype F(2.466, 78.913)=0.405, p=0.711; genotype
27 F(1,32)=3.831, p=0.059). During retrieval, GAD65^{+/+} mice showed a preference for a spatial
28 response strategy, while GAD65^{-/-} showed a more diverse response pattern (Fig. 1B;
29 likelihood-ratio chi square test $X^2(2)=16.569$, p=0.001).

30 When no proximal cue was present during training and retrieval, mice of both genotypes
31 (GAD65^{+/+}: n=9; GAD65^{-/-}: n=7) spent most of the time during the retrieval in the target
32 position previously probed with the reward (Fig. 1C; Repeated Measure ANOVA with

1 Greenhouse-Geißer correction for position: $F(1.067, 14.942)=15.358$, $p=0.001$; with $p<0.01$
2 for target vs. other positions, LSD posthoc test). No main effects of genotype ($F(1,14)=2.949$,
3 $p=0.108$) nor significant interactions of position x genotype: $F(1.067, 14.942)=4.062$, $p=0.06$
4 were observed. GAD65^{-/-} mice showed no significantly increased time at the target position
5 compared to their wildtype littermates (paired comparison, Mann-Whitney-U-test: $U=44.5$,
6 $p=0.174$). When a cue was present during training but not retrieval, mice of both genotypes
7 (GAD65^{+/+}: $n=8$; GAD65^{-/-}: $n=10$) spent more time in the target position (Fig. 1D; Repeated
8 Measure ANOVA for position: $F(3, 66)=12.255$, $p<0.001$; with $p<0.01$ for target vs. other
9 positions, LSD posthoc test) and no effects of genotype ($F(1,22)=1.555$, $p=0.225$) or
10 interactions of position x genotype: $F(3,66)=1.463$, $p=0.233$) were observed. Thus, while
11 stress-susceptible GAD65^{-/-} mice lost the spatial preference observed in GAD65^{+/+} mice in a
12 dual solution task, they were able to acquire spatial information for the reward location and
13 could use this information for retrieval if no other option was available.

14

15 **3.2 Interregional interaction and GAD65 expression regulation after stress point** 16 **towards the dDG as a hub for stress-related effects**

17 After the retrieval, the number of cFos-positive neurons (per area) was counted in selected
18 brain regions of spatial-preferring GAD65^{+/+} mice ($n=5$) vs. GAD65^{-/-} with a cue-based
19 retrieval choice ($n=5$). No significant differences were observed in any of the selected areas
20 (Fig. 2A; T-/ Mann-Whitney-U-Tests for PL/IL: $T(8)=0.843$, $p=0.424$; ACC: $T(8)=-0.141$,
21 $p=0.891$; striatum: $T(8)=-0.211$, $p=0.838$; DG: $T(8)=0.583$, $p=0.576$; CA1: $T(8)=0.318$, $p=0.758$;
22 CA3: $T(8)=-0.539$, $p=0.604$; LA: $T(8)=0.912$, $p=0.388$; BLA: $T(8)=0.458$, $p=0.659$; CeA: $U=14$,
23 $p=0.814$). However, a co-activation analysis using Pearson's correlation co-efficient revealed
24 negative correlations between the prefrontal areas (PL/IL) or the ACC and the dDG, as well
25 as a positive correlation between the dDG and the central amygdala in GAD65^{+/+} mice with
26 a spatial preference. In GAD65^{-/-} mice the correlation patterns of the DG with prefrontal
27 cortical areas, the striatum and the amygdala appeared reversed between groups (see Fig.
28 2B for color-coded correlation coefficients).

29 An important role of the dDG in mediating stress-related effects on dual solution strategy
30 choice was further supported by the expression analysis 24 h after restrained stress ($n=5-6$
31 for control, $n=6$ for restrained). Such a brief stressor before training has been shown to shift
32 retrieval strategies, at least in part via CORT signaling on mineralocorticoid receptors (MR).

1 24 h after restrained stress, mRNA levels of MR were increased in the dDG of GAD65+/+
2 mice (Fig. 2Ci; $T(10)=-2.343$, $p=0.041$), while the other receptor for CORT, GR, remained
3 unchanged ($T(10)=-0.433$, $p=0.674$). Moreover, in wildtype mice restrained stress lastingly
4 reduced the expression of GAD65 in the dDG (Fig. 2Ci; Mann-Whitney-U-test, $U=3$, $p=0.03$),
5 but not GAD67 (T-test, $T(9)=0.114$, $p=0.912$). The expression of MR and GR was not altered
6 in the dDG of GAD65-/- mice 24 h after restrained stress (Fig. 2Cii; MR: T-test $T(11)=0.572$,
7 $p=0.579$; GR: T-test $T(11)=0.541$, $p=0.6$). In the dorsal striatum no changes in expression
8 levels were observed in GAD65+/+ mice (Fig. 2Di; GAD65: Welch-tests for unequal variances,
9 $T(8.212)=0.041$, $p=0.968$; GAD67: T-test $T(10)=1.361$, $p=0.203$; MR: $T(10)=-0.157$, $p=0.878$;
10 GR: $T(10)=-1.09$, $p=0.301$) and in GAD65-/- mice (Fig. 2Dii; MR: T-test $T(11)=-1.817$, $p=0.097$;
11 GR: T-test $T(11)=-0.466$, $p=0.65$).
12 Together, these data suggested a relevant role for the dDG in stress responses and GAD65
13 expression regulation in wildtype mice that prompted us to specifically target this region in
14 the dual solution task.

15

16 **3.3 Local knock down of GAD65 in the dDG is sufficient to induce a loss of spatial** 17 **strategy preference in the dual solution task**

18 Based on the co-activation analysis, we then used a recently established shRNA-mediated
19 lentiviral vector for GAD65 knock down and injected it into the dDG (Tripathi et al., 2021).
20 Two weeks after virus injection training in the dual solution task took place. Mice with the
21 local GAD65 knock down (GAD65 KD, $n=18$) acquired the task as successfully as mice injected
22 with a control vector containing a random sh sequence (CTR, $n=16$; Fig. 3B; Repeated
23 Measure ANOVA with Greenhouse-Geisser correction: trials: $F(2.023,64.734)=46.477$,
24 $p<0.001$; trials x group: $F(2.023,64.734)=0.426$, $p=0.657$; group: $F(1,32)=1.386$, $p=0.248$).
25 Different strategy choice patterns were visible during retrieval 24h later (Fig. 3B;
26 $X^2(2)=6.478$, $p=0.039$), with CTR but not GAD65 KD mice showing a spatial preference.

27 To assess the impact of GAD65 KD on excitability of the dDG, perforant-path induced fEPSP
28 and PS responses were recorded in the DG granule layer in slices prepared from a different
29 set of CTR and GAD65 KD mice (Fig. 3Ci). Analysis of I/O curves revealed an increased
30 somatic fEPSP slopes in GAD65 KD slices (Fig. 3Cii; Repeated Measure ANOVA with
31 Greenhouse-Geisser correction: stimulation intensity $F(1.397,39.12)=86.93$, $p<0.0001$;
32 stimulation intensity x group: $F(8,224)=6.352$, $p<0.0001$; group: $F(1,28)=5.965$, $p=0.0212$)

1 and PS amplitudes (Fig. 3Ciii; Repeated Measure ANOVA with Greenhouse-Geißer
2 correction: stimulation intensity $F(2.396, 67.09)=63.80$, $p<0.0001$; stimulation intensity x
3 group: $F(8,224)=3.205$, $p=0.0018$; group: $F(1,28)=2.931$, $p=0.0979$). Accordingly, the
4 maximum evoked somatic fEPSP slope (Fig. 3Civ; Mann-Whitney-U-test, $U=39$, $p=0.0027$) as
5 well as PS amplitude (Fig. 3Cv; T-test, $T(28)=2.327$, $p=0.0274$) were increased in slices from
6 GAD65 KD mice. When determining paired pulse responses as a measure of short term
7 plasticity, no significant differences were observed after GAD65 KD for the somatic fEPSP
8 slopes (Fig. 3Cvi; Repeated Measure ANOVA with Greenhouse-Geißer correction: interval
9 $F(3.361, 87.39)=17.12$, $p<0.0001$; interval x KD $F(5,130)=1.366$, $p=0.2411$; KD
10 $F(1,26)=0.01234$, $p=0.9124$) and for the PS amplitudes (Fig. 3Cvii: Repeated Measure ANOVA
11 with Greenhouse-Geißer correction: interval: $F(1.664, 46.59)=64.26$, $p<0.0001$; interval x
12 group: $F(5,140)=0.2413$, $p=0.9435$; group: $F(1,28)=0.5033$, $p=0.4839$).

13 Of note, we found no evidence for altered neurotransmission and plasticity in the MF-CA3
14 (Suppl. Fig. 3) and SC-CA1 (Suppl. Fig. 4) synapses in hippocampal slices of GAD65 KD mice,
15 indicating that GAD65 KD induced changes in the excitability of dDG remain local without
16 altering baseline synaptic transmission and plasticity in the downstream regions of the
17 hippocampal trisynaptic circuit.

18 Taken together, a local GAD65 knock down in the dDG induced a similar phenotype in the
19 dual solution task as the total knock out of GAD65 and increased the excitability of this
20 structure, thereby suggesting an important role of dDG excitability in spatial strategy
21 preference.

22

23 **3.4 Modulation of dDG excitatory activity during retrieval affects strategy choices**

24 To dissect whether inhibition by dDG interneurons modulates strategy choices during dual
25 solution task retrieval, chemogenetic activation and inhibition of interneurons before
26 retrieval were performed via DREADDs in interneuron-specific cre-driver lines (Fig. 4A). SST-
27 Cre mice were used to target somatostatin-positive (SST(+)) interneurons in the dDG and to
28 increase (hM3Dq, $n=8$) or reduce (hM4Di, $n=8$) their activity, in comparison to controls (CTR,
29 $n=9$). The groups did not differ in their learning performance, as expected (Fig. 4B; Repeated
30 Measure ANOVA with Greenhouse-Geißer correction for trial $F(2.784,61.243)=18.357$,
31 $p<0.001$; trial x group $F(5.568,61.243)=0.846$, $p=0.533$; group $F(2,22)=1.934$, $p=0.168$).
32 Modulating SST(+) interneuron activity by CNO ligand injection before retrieval induced no

1 differences and all groups showed spatial dominating response patterns (Fig. 4B, upper
2 panel; $X^2(4)=0.196$, $p=0.906$). Manipulating parvalbumin-positive (PV(+)) interneurons was
3 achieved by injecting activating (hM3Dq, $n=10$), inactivating (hM4Di, $n=10$) and control
4 constructs (CTR, $n=9$) in the dDG of PV-Cre-mice and applying CNO again 1 h before retrieval.
5 As expected, training 24 h earlier was not affected (Fig. 4B, lower panel; Repeated Measure
6 ANOVA with Greenhouse-Geißer correction for trial $F(1.788,50.063)=42.462$, $p<0.001$; trial x
7 group $F(3.567,50.063)=1.381$, $p=0.256$; group: $F(2,28)=1.793$, $p=0.185$). Activating or
8 inactivating PV(+) did not shift response patterns significantly either (Fig. 5C; $X^2(2)=3.714$,
9 $p=0.446$). However, spatial preference was less established in control animals of this strain
10 (50% in PV CTR) compared to all other control mouse cohorts used in this study, suggesting
11 strain effects.

12 For comparison, dDG excitatory cells were chemogenetically inhibited before retrieval by
13 using inhibitory DREADD constructs under the CamKII promotor (Fig. 4C; hM4Di, $n=14$; vs.
14 control construct, $n=13$). Both, hM4Di and the CTR groups learned the location of the reward
15 well during training (Fig. 4B; Repeated Measure ANOVA with Greenhouse-Geißer correction
16 for trial $F(1.476,36.9)=28.779$, $p<0.001$; trial x group interaction $F(1.476,36.9)=2.642$,
17 $p=0.099$; group $F(1,25)=3.919$, $p=0.059$). After inactivating excitatory DG cells during
18 retrieval, strategy choices were shifted from spatial pattern in CTR to cue-based strategy in
19 the hM4Di groups (Fig. 4D; $X^2(2)=6.924$, $p=0.031$). Noteworthy, inhibiting DG activity before
20 training did not induce a shift in strategy choice (Suppl. Fig. 5). Together, a direct
21 contribution of either SST- or PV-positive DG interneuron populations to strategy choices
22 during dual solution retrieval could not be confirmed, despite their potential role in
23 modulating DG excitability. Targeting, excitatory cells of the dDG during retrieval, however,
24 induced a strategy shift, suggesting a specific role of this region in strategy choice during
25 retrieval rather than in acquiring strategies during training.

26

27

28 **4. Discussion**

29 The adaptation to stress involves various neurophysiological and behavioral responses,
30 including a switch from previously acquired spatial to cue-based search strategies (Schwabe
31 et al., 2010a, 2010b, 2008). Our findings suggest that these alterations involve a regulation

1 of inhibitory functions in the dDG, via regulation of the GABA synthesizing enzyme GAD65,
2 and excitability of the DG granule cells.

3 As one of the two isoenzymes synthesizing GABA, GAD65 expression is regulated in an
4 activity-dependent manner and is modulated by stress exposure (Albrecht et al., 2021b;
5 Bergado-Acosta et al., 2008; Bowers et al., 1998). GAD65^{-/-} mice display a behavioral
6 phenotype associated with a maladaptation to challenging environments, including
7 increased anxiety, increased fear memory as well as a heightened generalization of
8 conditioned fear and reduced fear extinction (see (Müller et al., 2015), for review). In the
9 current study increased plasma levels of corticosterone were observed in GAD65^{-/-} mice
10 handling controls, already without previous exposure to restrained stress. While basal
11 differences in corticosterone levels had not been observed in previous studies (Stork et al.,
12 2003), likely due to differences in rearing and analysis conditions including circadian rhythm,
13 they further support the notion of an altered stress response system in these mice.

14 Stress and elevated CORT levels have been shown to induce a shift in strategy preference in
15 mice favoring stimulus-response based strategies (Schwabe et al., 2010a). We therefore
16 tested unstressed, naïve GAD65^{-/-} mice in a dual solution foraging task, based on locating a
17 food reward in an open field. Like in other protocols based on aversive and appetitive stimuli
18 (see (Goodman, 2021), for review), mice have the option to build either a spatial map and
19 navigate in relation to distal cues like signs at the walls or to respond to cues placed
20 proximal to the goal, in our case the food reward. After establishing both strategies during
21 the memory acquisition phase, the preferred solution strategy can be assessed during
22 retrieval by moving the proximal cue to a different position. Mice will then either direct their
23 search to the cue or to the trained location (spatial solution). GAD65^{+/+} mice preferred a
24 spatial strategy in that case, as seen in wildtype mice in other dual solution paradigms e.g.
25 using a holeboard (Schwabe et al., 2008). GAD65^{-/-} mice, however, showed a cued response
26 in 50% of the individuals, as observed previously in chronically or acutely pre-stressed mice
27 (Schwabe et al., 2008, 2009). Importantly, GAD65^{-/-} mice showed in our protocol no
28 aversion against novel food nor any alterations in activity or anxiety-like behavior in the
29 open field (see Suppl. Fig. 6, 20-min habituation before the dual solution training), thereby
30 rendering confounding effects of heightened anxiety on learning or retrieval unlikely.

31 Across different dual solution paradigms, stress hormones such as CORT and activation of
32 the basolateral amygdala have been demonstrated to induce stress-related shifts of strategy

1 choices (Packard, 2009; Packard and Goodman, 2012). As these factors may impair spatial
2 memory acquisition (Roosendaal et al., 2003; Sandi and Pinelo-Nava, 2007), we tested
3 whether GAD65^{-/-} mice have a spatial learning deficit. However, these mice were able to
4 acquire spatial memory for a food reward when only a spatial solution was available or when
5 the proximal cue was absent during the retrieval. This suggests that GAD65^{-/-} mice have no
6 spatial learning or memory deficit per se, in line with undisturbed context processing in
7 aversive fear conditioning paradigms (Bergado-Acosta et al., 2008; Sangha et al., 2009).
8 Together, unstressed GAD65^{-/-} mice prefer a cue-based response strategy just like stressed
9 mice in previous studies, but neither disturbed emotional processing nor spatial learning
10 deficits appear accountable for this effect.

11 Previous studies suggest that both systems, hippocampus and dorsal striatum, are activated
12 during dual solution training and their cellular activity relates to strategy choices (Kathirvelu
13 and Colombo, 2013; Yagi et al., 2017). Studies in humans and rodents show that decision
14 making processes such as choosing an appropriate strategy also depend on the mPFC and
15 ACC (Rich and Shapiro, 2009; Rudebeck and Izquierdo, 2022). Using the neuronal activity
16 marker cFos, we mapped neuronal activation in these brain regions and found no
17 differences in neuronal activation patterns of spatial-preferring GAD65^{+/+} vs. cued-
18 preferring GAD65^{-/-} mice, further supporting that hippocampus and striatum-based retrieval
19 systems are intact in GAD65^{-/-} mice. However, when correlating activities between brain
20 areas, we noted a shift in the co-activation specifically of the dDG with prefrontal cortical
21 areas and the central amygdala areas between genotypes, indicative of a change in
22 functional connectivity.

23 The dDG is perfectly suited to integrate cognitive processes and emotional arousal and
24 plasticity and is modulated by stress and amygdala activation (Albrecht et al., 2021a; Fa et
25 al., 2014). Information processing in the DG is achieved by strong inhibition via GABAergic
26 interneurons that form local networks with the excitatory granule cells (Acsády and Káli,
27 2007) and may mediate stress effects and the impact of GAD65 deficiency on strategy
28 preferences. We therefore assessed whether a brief restrained stress exposure, a procedure
29 that induced a shift in strategy choice in previous studies (Schwabe et al., 2010b), alters the
30 expression of GAD65 in the dDG of wildtype mice. Indeed, GAD65 mRNA expression was
31 reduced in the dDG while remaining unaltered in the striatum of GAD65^{+/+} mice. Moreover,
32 an increased expression of MR was observed in the dDG of GAD65^{+/+} mice, but not in their

1 striatum. In GAD65^{-/-} mice MR levels remained unchanged in the dDG and only a slight, but
2 insignificant increase in MR expression was visible in the striatum, further supporting an
3 altered regulation of the stress axis in GAD65^{-/-} mice. The activation of MR has previously
4 been linked to the stress-induced strategy shift in dual solution paradigms. An orally
5 administered MR antagonist could prevent stress-induced shifts in strategy choices
6 (Schwabe et al., 2010a), while mice overexpressing MR in excitatory neurons showed a
7 stronger switch to cued response strategies after stress (ter Horst et al., 2012). Thus,
8 increased MR levels in the dDG of wildtype mice after stress underline a possible central role
9 of the dDG.

10 Accordingly, we then commenced by mimicking a molecular aspect of stress exposure by
11 reducing the expression of GAD65 only in the dDG using lentiviral vectors. This treatment
12 clearly replicated the lack of spatial strategy preference observed in the total GAD65^{-/-} mice.
13 Field recordings in DG slices from such animals demonstrated that a GAD65 knock down
14 furthermore increased excitability of dDG circuits without altering paired pulse responses.
15 Previous studies in GAD65^{-/-} mice found a reduced paired pulse inhibition, but only in the
16 CA1 area (Tian et al., 1999). The current results suggest a heightened excitability of the DG,
17 in accordance with a reduced inhibitory activity due to GAD65 knock down.

18 Two major populations of GABAergic interneurons shape the local circuits of the DG. We
19 next investigated therefore their potential impact on strategy choices in the dual solution
20 task by chemogenetic intervention. This approach also allowed for a targeted manipulation
21 of the training vs. retrieval phase. First, we manipulated somatostatin interneurons, which
22 are predominantly located in the hilus of the dDG and mediate inhibition of distal dendrites
23 of excitatory granule cells (Tallent, 2007). These cells control the size of engrams for
24 contextual representations (Raza et al., 2017; Stefanelli et al., 2016) and play a role in spatial
25 memory formation (Morales et al., 2021). However, neither activating nor inactivating
26 somatostatin-positive interneurons before retrieval affected strategy choices, leaving spatial
27 preference undisturbed in the dual solution paradigm. We then tested parvalbumin-positive
28 interneurons, which represent mostly basket and chandelier cells located close to the
29 granule cell layer that mediate inhibition near the soma and proximal dendrites of granule
30 cells (Bartholome et al., 2020). Their chemogenetic activation in the DG has been shown to
31 reduce anxiety and to enhance contextual fear memory extinction (Zou et al., 2016). In our
32 experiments, activating PV-positive interneurons resulted in about 70% of mice choosing a

1 cued response strategy. However, only 50% of the control vector-injected mice of the PV-Cre
2 mouse line showed a preference for the spatial solution, suggesting a genetic background
3 effect and different basal behavior of this mouse line in the dual solution paradigm. While it
4 is intriguing to speculate that increased dDG inhibition by activating PV-positive
5 interneurons may contribute to shifts in strategy choices, future experiments using different
6 interventional approaches are necessary to answer this question.

7 Finally, in addition to manipulating specific interneuron populations, we used a
8 chemogenetic vector that allowed for the inhibition of excitatory cells of the dDG (under
9 control of the CamKIIa promotor). The use of an activating chemogenetic construct in these
10 mice however had to be abandoned after pilot experiments, since the locomotor activity of
11 the animals was low and side effects such as seizures were frequently observed. Focusing on
12 inactivating DG excitatory cells before retrieval, we observed a shift towards a cue-
13 preference. By contrast, inactivating the excitatory dDG cells before training did not induce
14 such a shift (see Suppl. Fig. 5). Together, these findings suggest that either increasing (via
15 GAD65 knock down and potentially also by blocking PV interneurons) or decreasing the
16 excitability of the DG (via chemogenetic inhibition of excitatory cells) both induces a shift
17 towards cue-based strategies specifically in the retrieval phase of the dual solution
18 paradigm. This suggests that an optimum excitation/ inhibition balance in the dDG is
19 required for its proper functioning within a strategy choice network that may comprise the
20 striatum and frontal cortical areas. Moreover, a recent study demonstrates that mossy cells
21 of the hilus respond as well to objects explored in specific spatial configurations (GoodSmith
22 et al., 2022). Mossy cells are the second excitatory cell population of the dDG next to granule
23 cells. They are located in the hilus, receive inputs from granule cells as well as from back
24 projections of CA3 and they target other mossy cells, granule but also basket cells.
25 Depending on neuromodulatory inputs (i. e. acetylcholinergic inputs), mossy cells can
26 provide a feed forward inhibition of granule cells, but also directly activate sparse granule
27 cell populations (Schafman, 2016). They may therefore provide a switch for DG activity
28 during the dual solution task depending on arousal and comprise an interesting target to
29 study local circuits of the DG in future studies.

30

31 **5. Conclusions and future perspectives**

32

1 Setting out to investigate neuronal circuits underlying memory-based foraging strategies, we
2 identified an unexpected role of the DG in mediating strategy selection during memory
3 retrieval. Alterations in GAD65-mediated inhibition in the DG seem to provide a molecular
4 mechanism of such an adaptive response. It is well known that the dDG is embedded in a
5 functional network of decision-making processes, comprising for example the mPFC, the ACC
6 and the anterior thalamic nucleus (Méndez-Couz et al., 2015), that may support its role as a
7 hub for strategy selection. In fact, cFos-based regional activation analysis indicated altered
8 functional interaction of DG and ACC in GAD65^{-/-} mice during the dual solution task. Direct
9 hippocampal inputs to the frontal cortex stem mainly from the ventral hippocampal CA1
10 areas but not the DG (Eichenbaum, 2017), but a knock down of GAD65 in the DG resulted in
11 no alterations of excitability of CA1 or CA3 (see Suppl. Fig. 3 and 4). One promising relay
12 station to be considered is the supramammillary nucleus (SUM), which projects
13 bidirectionally to the ACC, to the PFC and specifically to the dorsal DG (Pan and
14 McNaughton, 2004) and synchronizes its activity with the dDG during spatial memory
15 retrieval (Li et al., 2020). Deconstructing this network and the contribution of such PFC-dDG-
16 relay stations to spatial and cue-based information processing and retrieval will be an
17 attractive task for future studies and may provide novel insights in networks of decision-
18 making processes.

19

20

21

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4

5 **Declaration of interest**

6 None

7

8

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37

1 **Figure Legends**

2 **Fig. 1: Strategy preference in GAD65^{-/-} mice. (A)** GAD65^{-/-} mice showed elevated
3 corticosterone serum levels already under baseline control conditions that were not further
4 elevated 24 h after a brief restraint stress episode, suggesting altered stress susceptibility.
5 **(B)** In the dual solution task (scheme on top), GAD65^{-/-} mice showed a good learning
6 performance of the reward location over 6 trials. When tested for their strategy preference
7 24 h later, they did not display a spatial preference in comparison to their wildtype
8 littermates. **(C)** When the experimental setup only allowed for a spatial solution in order to
9 learn the reward location, GAD65^{-/-} mice showed a spatial preference for the reward
10 location during retrieval. **(D)** Similarly, when a cue was present during training but not during
11 retrieval, GAD65^{-/-} mice still displayed a spatial preference similar to their wildtype
12 littermates. All values means \pm sem. * significant difference between genotypes, $p < 0.05$; ***
13 $p < 0.001$; +++ significant learning effect over trials, $p < 0.001$; \$\$ significant difference of
14 target position compared to other positions, $p < 0.01$; \$\$\$ $p < 0.001$.

15

16 **Fig. 2: Interregional activation patterns during dual solution retrieval and GAD65**
17 **expression regulation after stress point towards the dDG as a hub for stress-related**
18 **effects. (A)** GAD65^{-/-} mice and their wildtype littermates were trained in the dual solution
19 task and perfused 1 h after retrieval brains for subsequent cFos analysis. Example images are
20 shown for cFos labeling (green: cFos; cyan: DAPI nuclear staining; white arrow head:
21 exemplary cFos-positive cell; scale bar = 100 μ m) in the dorsal dentate gyrus. No difference
22 in the number of cFos positive cells was evident in any of the regions analyzed between
23 GAD65^{+/+} mice preferring a spatial solution (n=5) and GAD65^{-/-} mice using cue-based
24 strategy (n=5). **(B)** Interregional correlations of cFos activation after DS retrieval and strategy
25 choice demonstrated shifts in co-activation of the dDG together with PL/IL, and CeA in
26 between the groups. **(C)** The dDG was also susceptible to gene expression changes induced
27 by restrained stress. In (i) GAD65^{+/+} mice GAD65 mRNA levels were downregulated and
28 mineralocorticoid receptor (MR) upregulated 24h after brief restrained stress, while (ii) MR
29 expression was unaltered in GAD65^{-/-} mice (n=5-7 per group). **(D)** In comparison, no stress-
30 induced expression changes were observed in the striatum of (i) GAD65^{+/+} and (ii) GAD65^{-/-}
31 mice (n=5-7 per group). Expression of the isoform GAD67 and of the glucocorticoid receptor

1 (GR) remained unchanged in both regions. All values means \pm sem. + significant Pearson's
2 correlation, $p < 0.05$; ++ $p < 0.01$; * significant difference between genotypes, $p < 0.05$.

3

4 **Fig. 3: Local knock down of GAD65 in the dDG leads to loss of spatial preference in a dual**

5 **solution task. (A)** Vector scheme and workflow for local knock down of GAD65 in the dDG

6 before dual solution training and retrieval. **(B)** The local knock down of GAD65 in the dDG is

7 sufficient to replicate the phenotype of total GAD65^{-/-} mice. **(C)** Electrophysiological profile

8 of the dDG after local GAD65 knock down (CTR: n=18 slices from 9 animals; GAD65 KD: n=12

9 slices from 8 animals). (i) Scheme for perforant-path induced somatic field postsynaptic

10 potential (fEPSP) and population spike (PS) responses. Input-output (I/O) curves

11 demonstrate an enhanced excitability after GAD65 KD in the dDG evident by (ii) increased

12 fEPSP slope and (iii) PS amplitudes. (iv) Accordingly, the maximum evoked fEPSP slope and

13 (v) PS amplitude is increased in GAD65 KD slices. No differences in the paired pulse ratios

14 were observed for (vi) the fEPSP slopes or (vii) the PS amplitudes. All values means \pm sem.

15 +++ significant learning effect over trials, $p < 0.001$; * significant difference from CTR group,

16 $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$. &&& $p < 0.001$, significant stimulus intensity effect.

17

18 **Fig. 4. Modulation of dDG excitatory activity during retrieval affects strategy choices. (A)**

19 Scheme for chemogenetic activation/ inhibition of somatostatin (SST)- and parvalbumin

20 (PV)-positive interneurons during dual solution retrieval. **(B, upper panel)** Activation or

21 inhibition of dDG SST(+) cells during dual solution retrieval did not affect strategy choice

22 (CTR: n=9; 3Dq: n=8; 4Di: n=8). **(B, lower panel)** Activation or inhibition of PV(+)

23 interneurons during dual solution retrieval did not affect strategy choice significantly, but

24 spatial strategy preference appeared diminished in the PV-hM3Dq group (not significant;

25 CTR: n=9; 3Dq: n=10; 4Di: n=10). **(C)** For comparison, chemogenetic inhibition of excitatory

26 dDG cells was achieved using CamKII-hM4Di constructs. Expression of the marker mCherry is

27 observed in the molecular layer and mossy fibers, containing dendritic and axonal

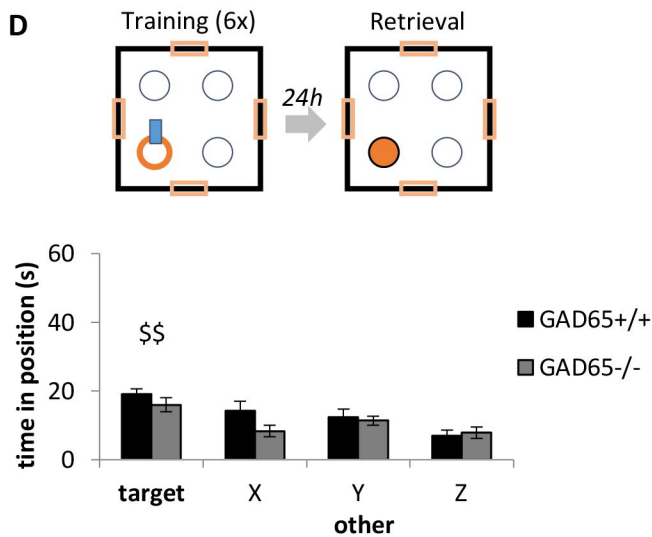
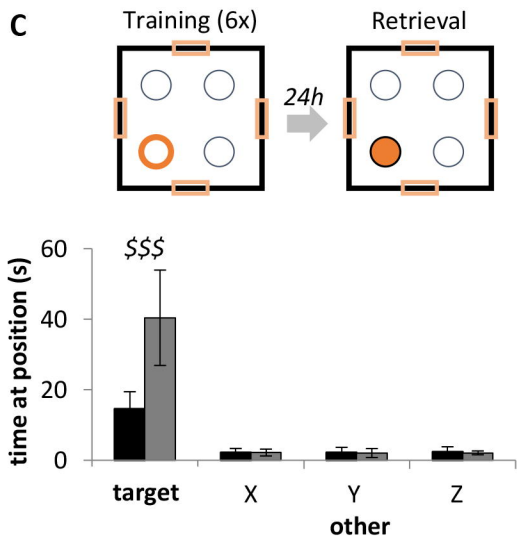
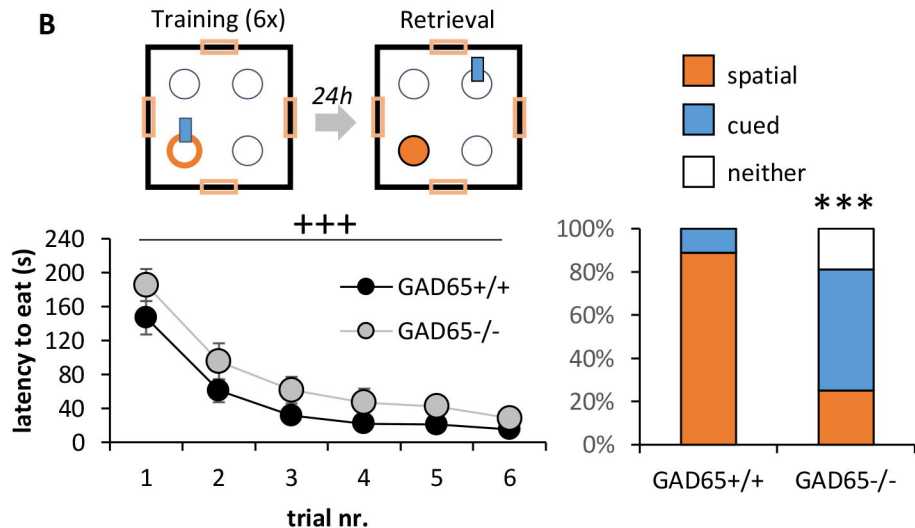
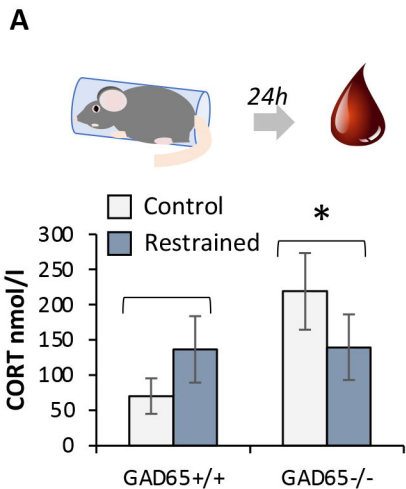
28 compartments of dDG granule cells. **(D)** Inhibiting excitatory cell activity during retrieval

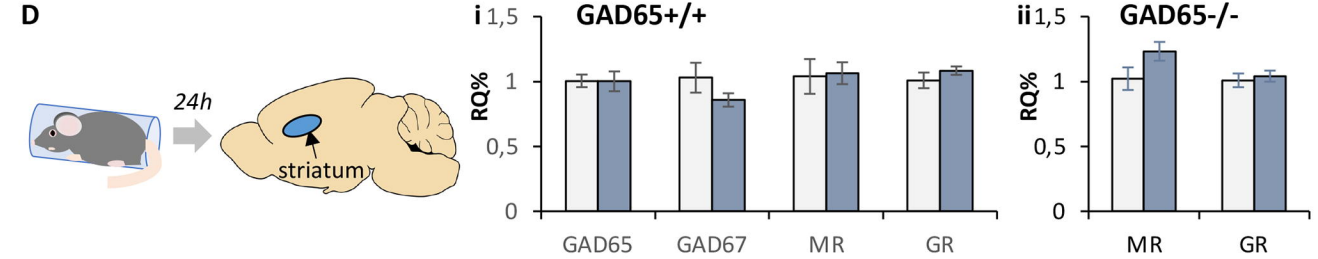
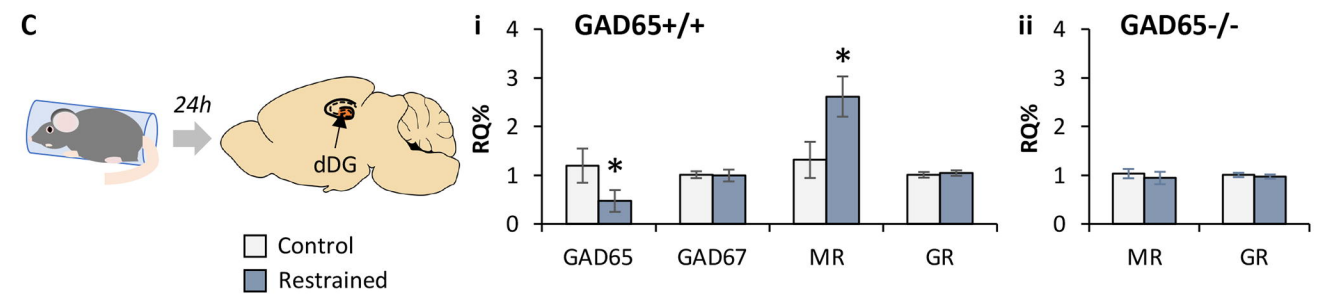
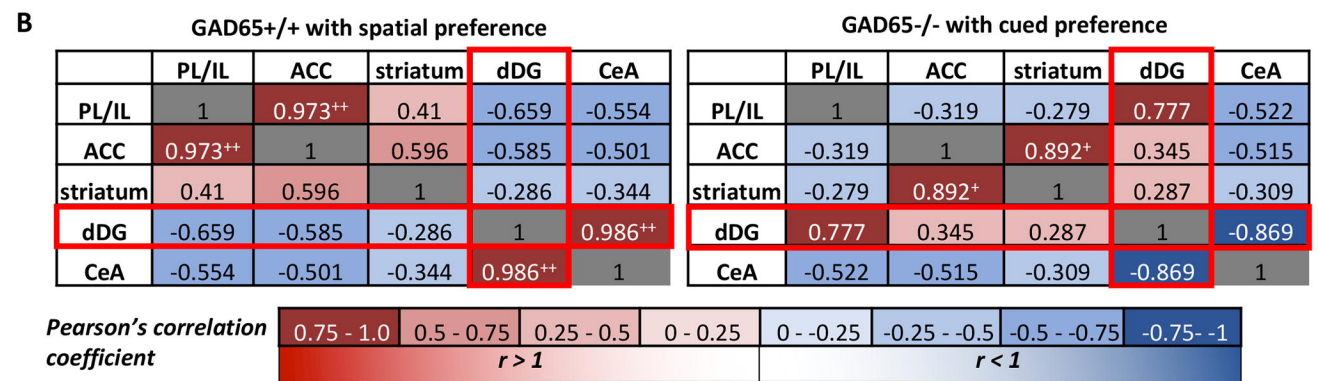
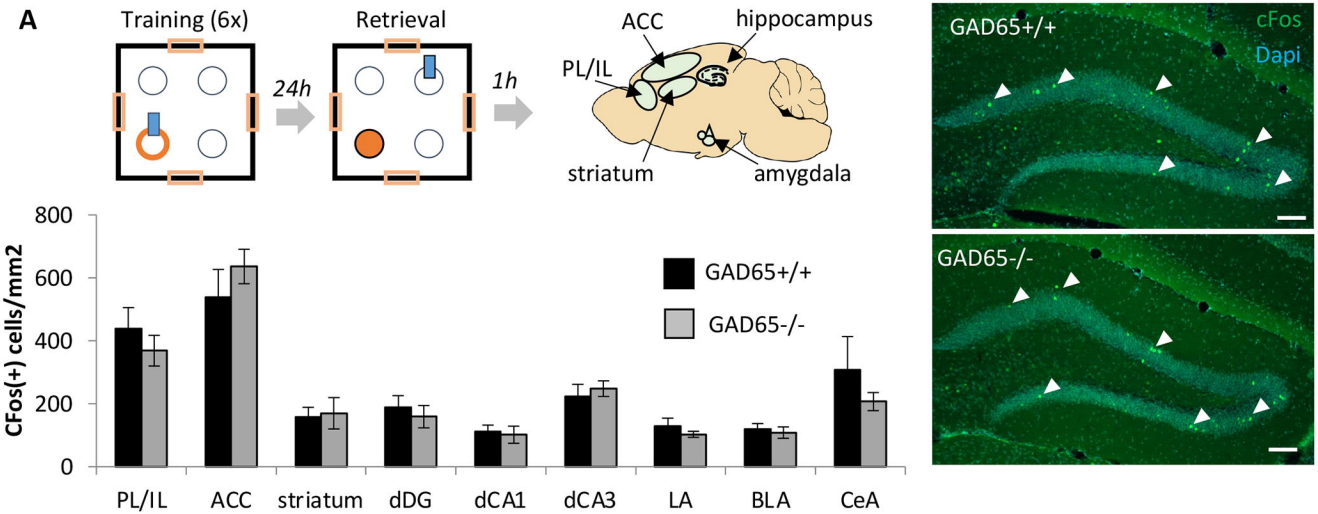
29 induced a strategy shift with a loss of spatial preference (n=14; CTR: n=13). All groups did not

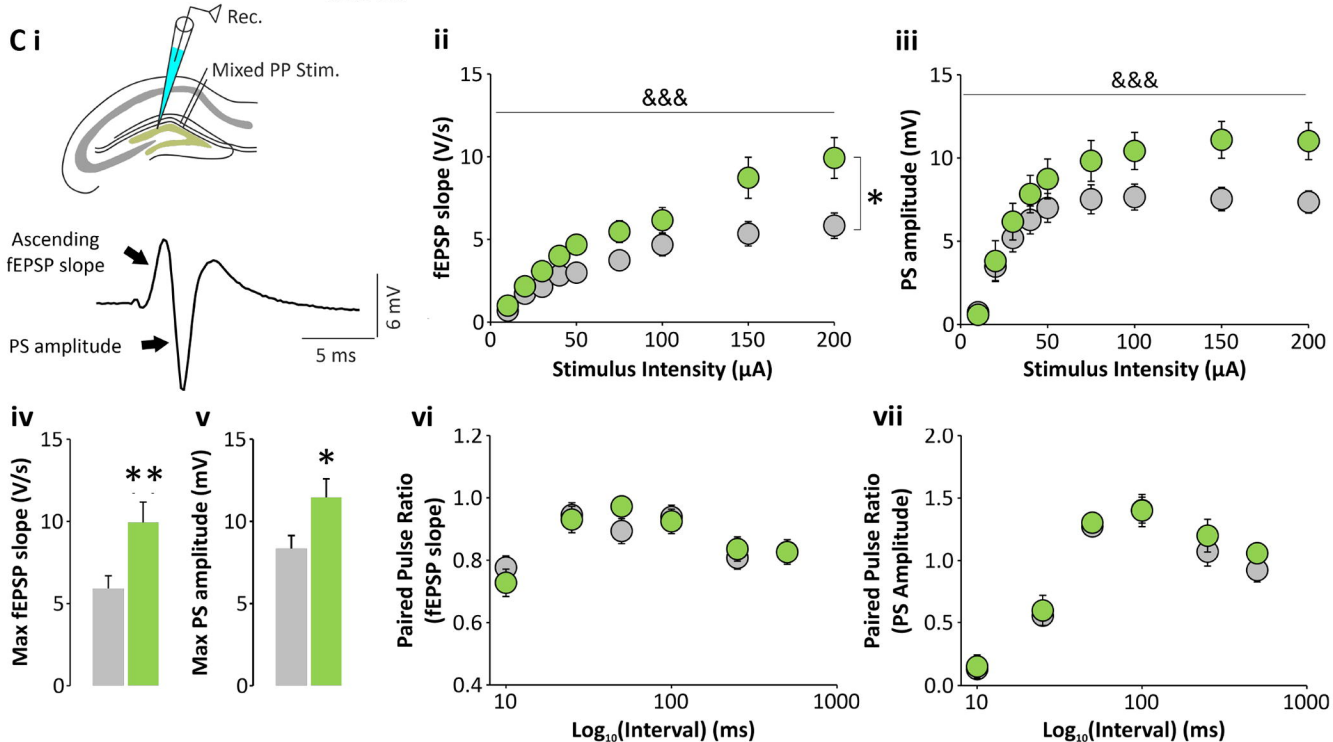
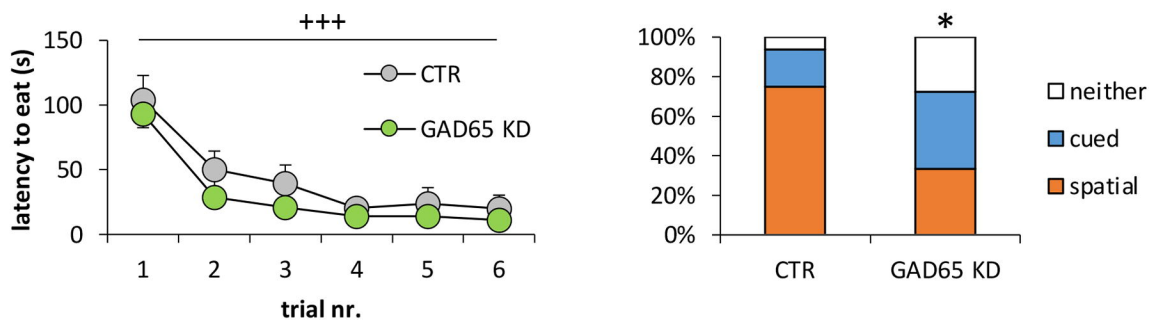
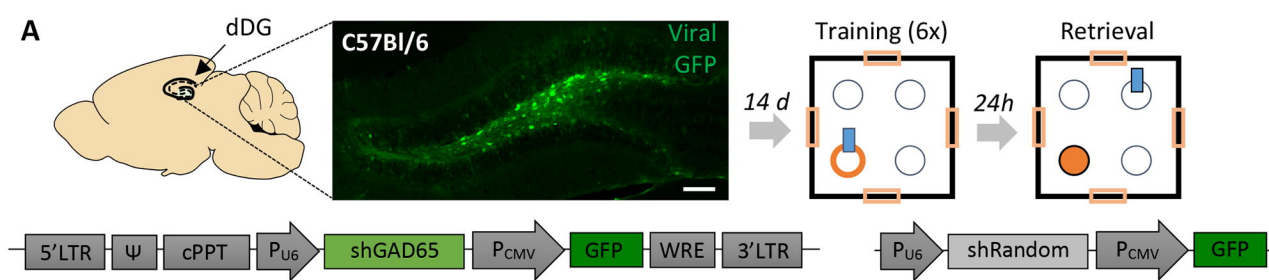
30 differ in their training performance. All values means \pm sem. +++ significant learning effect

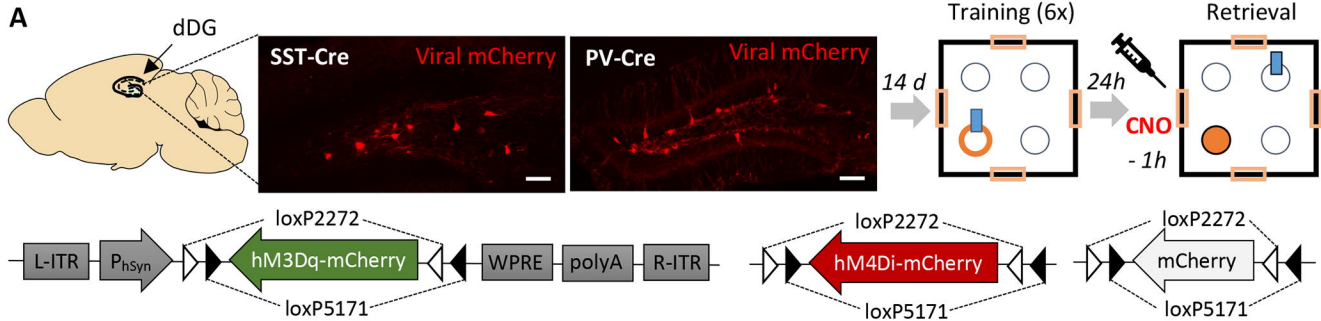
31 over trials, $p < 0.001$; * significant difference to CTR group, $p < 0.05$.

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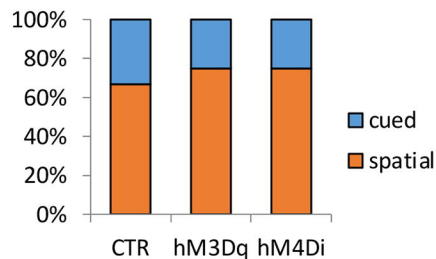
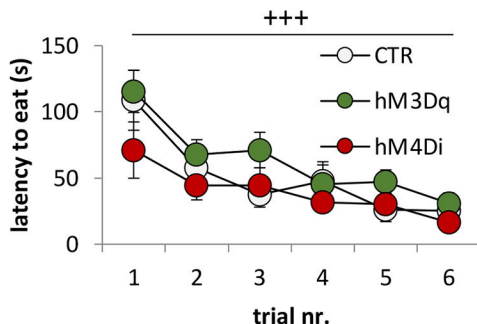
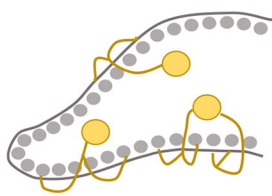




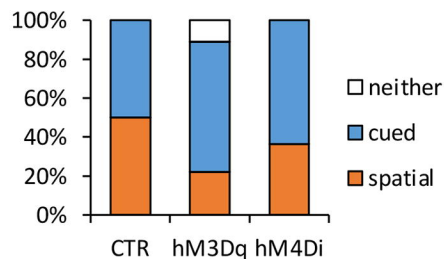
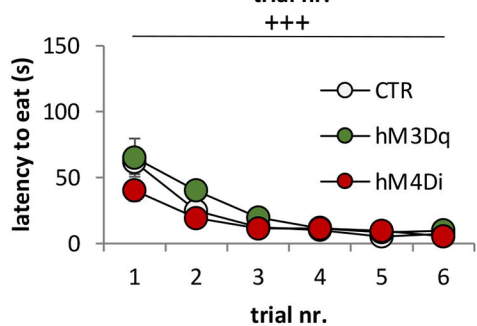
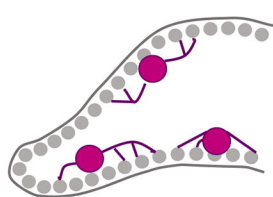




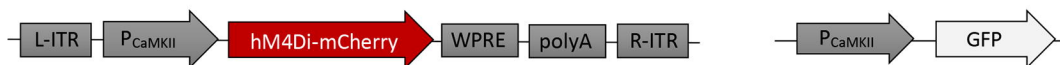
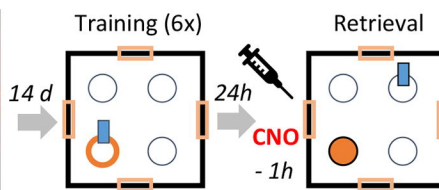
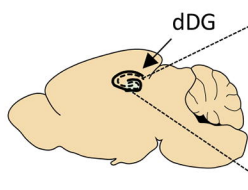
B Inhibition/ Activation of SST interneurons



Inhibition/ Activation of PV interneurons



C



D Inhibition of excitatory cells

