1	Engram reactivation during memory retrieval predicts long-term memory performance
2	in aged mice
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29 Abstract

30 Age-related cognitive decline preferentially targets long-lasting episodic memories that 31 require intact hippocampal function. Memory traces (or engrams) are believed to be encoded 32 within the neurons activated during learning (neuronal ensembles), and recalled by 33 reactivation of the same population. However, whether engram reactivation dictates memory 34 performance late in life is not known. Here, we labelled neuronal ensembles formed during 35 object location recognition learning in the dentate gyrus, and analyzed the reactivation of this 36 population by long-term memory recall in young adult, cognitively impaired- and unimpaired-37 aged mice. We found that reactivation of memory-encoding neuronal ensembles at long-term 38 memory recall was disrupted in impaired but not unimpaired-aged mice. Furthermore, we 39 showed that the memory performance in the aged population correlated with the degree of 40 engram reactivation at long-term memory recall. Overall, our data implicates recall-induced 41 engram reactivation as a prediction factor of memory performance in aging. Moreover, our 42 findings suggest impairments in neuronal ensemble stabilization and/or reactivation as an 43 underlying mechanism in age-dependent cognitive decline. 44

Keywords: aging, dentate gyrus, hippocampus, memory trace, neuronal ensembles, object
location memory.

48 **1. Introduction**

49 Age-related cognitive decline refers to the gradual decrease in cognitive performance 50 throughout the aging process, mostly affecting long-term storage of episodic and spatial 51 memories that depend on hippocampal function (Burke and Barnes, 2006). The dentate 52 gyrus (DG) is the hippocampal subregion most sensitive to the effects of advanced age 53 (Small et al., 2011, 2004). The DG undergoes anatomical and physiological changes as well 54 as alterations in the transcriptomic profile that are thought to underlie the aging-associated 55 dysfunction (Burke and Barnes, 2010; lanov et al., 2016). It has recently been shown that 56 subsets of DG neurons activated during learning (i.e., neuronal ensembles) belong to the 57 memory engram, and that the reactivation of this neuronal population at memory recall is 58 necessary and sufficient to evoke memory retrieval in young adult mice (Josselyn et al., 59 2015; Tonegawa et al., 2015). Moreover, dysfunctions in neuronal ensemble reactivation and 60 memory retrieval mechanisms have been proposed as an underlying cause of dementia in 61 pathological conditions, such as Alzheimer's disease (Perusini et al., 2017; Poll et al., 2020; 62 Roy et al., 2016), in which aging presents a major risk factor. However, whether memory 63 impairments observed late in life are associated with disruptions in neuronal ensemble 64 reactivation is not known. 65 Here, we hypothesized that differences in DG neuronal ensemble dynamics underlie the 66 inter-individual variability in long-term memory (LTM) performance in the aged population. 67 We used tagging tools to label and characterize the formation and reactivation of neuronal 68 ensembles associated with long-term spatial recognition memory, a form of memory 69 particularly susceptible to decline with age. We showed that impairments in long-term spatial 70 memory during aging are associated with impaired reactivation of memory-encoding 71 neuronal ensembles at LTM recall. Moreover, we found that the degree of engram 72 reactivation in aged mice correlated with long-term spatial memory performance. Overall, our 73 findings identified novel cellular correlates of age-related memory decline. 74

75 2. Materials and Methods

76 2.1. Animals. 8-10 weeks or 18-20 months old male C57BL/6JRj mice (Janvier Labs, 77 Germany) were individually housed on a 12 h light/dark cycle and ad libitum access to food 78 and water (after stereotaxic surgery regular food was replaced by doxycycline-containing diet 79 (40mg/kg, BioServ, Flemington, NJ, USA)). Experiments were carried out during the light 80 phase. All of the animal procedures were approved and performed according to the 81 European Community Council Directive 86/609/EEC. 82 2.2. Recombinant adeno associated viruses (rAAVs). rAAVs were produced and purified as 83 described previously (Gulmez Karaca et al., 2020; Zhang et al., 2007). The RAM-HA-GFP 84 viral construct was generated by insertion of a HA-tagged GFP expression cassette into the 85 multiple cloning site (MCS) of rAAV-RAM (rAAV-RAM-d2TTA::TRE-MCS-WPRE-pA) that 86 was a kind gift from Yingxi Lin (Addgene plasmid # 63931; http://n2t.net/addgene:6393; 87 RRID:Addgene 63931)) (Gulmez Karaca et al., 2020; Sørensen et al., 2016). 88 2.3. Stereotaxic surgery. 250 nl of rAAV-RAM viral solution was injected into the dentate 89 gyrus (DG) of the hippocampus at the following coordinates relative to Bregma: - 2.0 mm 90 anteroposterior, ± 1.3 mm medio-lateral, – 2.4 mm dorsoventral at the rate of 80-100 nl / min. 91 The needle was left in place for 10 min before and after each injection to allow DG-specific 92 diffusion of the fluid. Only the mice with viral spread throughout the DG of at least 60 µm 93 range in the AP axis were considered for behavior and further analysis. Throughout the 94 study, a total of 10 mice were rejected from the analysis due to insufficient viral spread (5 95 mice) and/or lack of exploration during behavioral training or testing (5 mice). 96 2.4. Behavior paradigms. Three weeks after stereotaxic surgery and following 5 days of 97 handling, the spatial object recognition (SOR) test was performed. SOR training involved four 98 sessions; each lasted 6 min and was separated by 3 min intervals in the home-cage. In the 99 first session, mice explored an empty black, square open field (50 cm × 50 cm × 50 cm) with 100 a visual cue placed on one wall. This session was also used for the open field test as 101 described previously (Brito et al., 2020b, 2020a; Gulmez Karaca et al., 2018). The Smart 102 Video Tracking Software (Panlab, Harvard Apparatus) was used to score the time spent in 103 the central zone (32% of the arena), number of center entries and total distance travelled in

104 the open field test. In the next three sessions of the SOR training, mice freely explored two 105 diagonally located distinct objects (a glass bottle and a metal tower) in the arena. After the 106 training, mice were placed in the home-cage and were undisturbed until the memory test. 107 During the memory test (24 h post-training), one of the two objects was displaced in the 108 arena (while the other one remained in the original position), which the mice explored freely 109 for 6 min. Object exploration was defined as the animal sniffing the object or pointing its nose 110 to the object within 1-cm distance. When the animal leaned on the object but did not direct 111 the nose towards it, object exploration was not considered. SOR memory performance was 112 assessed by the formula ($T_{displaced}$ /($T_{nondisplaced}$ + $T_{displaced}$) × 100) where T represents time 113 exploring an object. Assigning the aged mice into cognitively impaired and unimpaired 114 groups was performed based on the mean SOR memory performance of the young group. 115 Aged mice displaying memory performance lower than one standard deviation from the 116 mean of young mice were designated as cognitively impaired. 117 2.5. Immunohistochemistry. 2 h after the start of the SOR test, mice were perfused 118 intracardially with 4% paraformaldehyde (PFA) (Sigma, Munich, Germany) and free-floating 119 brain slices (at a thickness of 20 µm) were immunostained as previously described (Gulmez 120 Karaca et al., 2018; Oliveira et al., 2012). Primary antibodies were used at the following 121 concentrations: anti-HA tag (Covance MMS-101R (1:1000)), anti-Fos (Cell Signaling #2250, 122 1:1000)). 123 2.6. Image acquisition and analysis. Z-stacks of DG images (3 frames with 2 µm interval at 124 20x magnification) were acquired with Nikon A1R confocal microscope (at Nikon Imaging 125 Center, BioQuant, Heidelberg) using NIS-Elements software or with Leica SP8 confocal 126 microscope using LAS X software. Maximum projection files of each stack were imported in 127 Fiji (Schindelin et al., 2012), and Fos⁺ or HA-GFP⁺ neurons were manually marked after 128 background subtraction and application of a signal threshold. Reactivation rate 129 ((GFP⁺Fos⁺)/(GFP⁺)×100), and similarity index ((GFP⁺Fos⁺)/((GFP⁺)+(Fos⁺)-130 (GFP⁺Fos⁺))×100 was calculated as previously described (Cowansage et al., 2014; Gulmez

131 Karaca et al., 2020; Milczarek et al., 2018). Data was normalized to the mean of the aged

impaired group to avoid artifacts that may be caused from different viral expressions among

133 experimental batches.

134 2.7. Statistical analysis. Blinding to experimental conditions was applied to image and 135 behavioral analysis. Each data was subjected to a normality test (Shapiro-Wilk normality test, 136 alpha=0.05) before further analysis. For normally distributed data ordinary one-way ANOVA 137 followed by Tukey's multiple comparisons test, whereas for non-normally distributed data, 138 Kruskal-Wallis test followed by Dunn's multiple comparisons test was used to compare three 139 groups. When comparing two dependent data sets, paired t-test was performed if the data 140 set showed a normal distribution, if not, Wilcoxon matched-pairs signed rank test was 141 applied. For correlation analysis, Pearson correlation test was applied when the data 142 followed a normal distribution. Nonparametric Spearman correlation was computed in case of 143 a non-normally distributed dataset. Statistical analysis was performed using GraphPad prism 144 for Mac OS X, version 8.

145

146 3. Results

147 3.1. Neuronal ensemble tagging in the DG of young, cognitively-impaired or -unimpaired 148 aged mice. We aimed at characterizing DG neuronal ensembles that hold long-term 149 representations of object location memory in order to identify possible correlates of cognitive 150 dysfunction in the aged population. To label the neuronal ensemble activated during learning, 151 we used the robust activity marking (RAM) system (Sørensen et al., 2016) that allowed 152 learning-dependent expression of HA-tagged GFP (Sørensen et al., 2016) (Figure 1A). We 153 have previously confirmed that this tool reliably tags the DG neuronal ensembles associated 154 with object location memory (Gulmez Karaca et al., 2020). First, we stereotaxically delivered 155 recombinant adeno associated viruses (rAAVs) containing the RAM-HA-GFP viral construct 156 into the DG of young-adult (2 months old) or aged (18-20 months old) mice. Through the 157 removal of doxycycline diet, we tagged the neuronal population activated during object 158 location training (Figure 1A). When tested 24 h after training all young adult mice displayed 159 higher than chance (50%) preference for the displaced object (Figure 1B), indicating intact

160 long-term object location memory. In contrast, the performance of the aged group was 161 heterogenous; some mice performed similarly to young mice, whereas others showed 162 preferences closer to chance level (Figure 1B). To be able to characterize the aged 163 population according to individual differences in cognitive performance, we sorted the aged 164 group into aged impaired (AI) and aged unimpaired (AU) based on the mice's long-term 165 spatial memory scores (Figure 1B). Aged mice displaying preferences for the displaced 166 object lower than one standard deviation below the mean of young mice (64.3%, represented 167 with a dashed line in Figure 1B) were considered impaired. Importantly, we confirmed that all 168 groups displayed comparable object exploration times during the training session (Figure 169 1C) and demonstrated similar locomotion and anxiety-like behavior in open-field test (Figure 170 **1D-F**). This indicates that poor LTM performance in AI mice was not due to lower exploratory 171 behavior or motor or anxiety impairments. 172 3.2. Cognitively impaired aged mice exhibit impaired recall-induced neuronal ensemble 173 reactivation in the upper blade of the DG. Next, we investigated the reactivation of spatial 174 memory-associated neuronal ensembles in response to LTM recall. To this end, the 175 expression of HA-GFP and Fos in the DG, that represents active neurons during learning or 176 LTM recall respectively, was analyzed 2 h after object location test (Figure 2A). The 177 neuronal population coexpressing HA-GFP and Fos represented the population active in both 178 episodes, i.e., reactivated neuronal ensemble (Figure 2B). Given the previously described 179 differences in responsiveness of the upper and lower blades of the DG to environmental 180 exposure (Chawla et al., 2013; Gulmez Karaca et al., 2020; Marrone et al., 2012; Ramirez-181 Amaya et al., 2013), we analyzed the two subregions separately. We found that AI mice 182 exhibited reduced recall-driven reactivation of the neuronal population activated by learning 183 in the DG compared to young or AU mice (Figure 2C), whereas the reactivation rates 184 observed in the DG of young and AU mice were similar (Figure 2C). Interestingly, the 185 impaired reactivation rate was specific for the neuronal ensembles located in the upper, but 186 not the lower, blade of the DG in Al mice (Figure 2C). Although Al mice were able to 187 reactivate the learning-activated neuronal population at a rate significantly higher than what

188 would occur by chance, the observed overlap above chance level was higher both in the 189 young and AU groups compared to the AI group (Figure 2D). The analysis of the two DG 190 blades independently, showed that, similarly to what we observed for the reactivation rate 191 (Figure 2C), the effect originated from the upper blade (Figure 2D). There was no significant 192 difference in the reactivation above chance levels of young and AU mice (Figure 2D). 193 Furthermore, to evaluate whether the learning-activated neuronal ensemble pattern was 194 reinstated equally in all of the three groups at memory recall, we applied a second formula, 195 the similarity index (Milczarek et al., 2018). This formula complements reactivation rate by 196 also including the number of the neurons activated solely by recall, and thereby allows to 197 measure the degree of similarity in the neuronal activity pattern at learning and recall. We 198 observed that AI mice exhibited the same selective impairment in the DG upper blade 199 (Figure 2E). We then confirmed that the identified differences between the AI and AU were 200 not due to differences in the number of neurons activated by learning (GFP⁺) (Figure 2F) or 201 by recall (Fos⁺) (**Figure 2G**) in the aged population. Although learning triggered the activation 202 of fewer neurons in the AU mice than the young (Figure 2F), this did not correlate with 203 reactivation rate in the DG upper blade (p=0.2286, r=-0.3073, Spearman correlation, data not 204 shown) or with memory performance (p=0.1566, r=-0.3594, Pearson correlation, data not 205 shown) in the aged. Altogether, this set of experiments revealed that despite being able to 206 activate a similar proportion of neurons upon learning or memory recall, cognitively impaired 207 mice exhibit disrupted neuronal ensemble reactivation at LTM recall in the upper blade of the 208 DG.

3.3. LTM performance of aged mice correlates with the reactivation rate in the upper blade of the DG. Finally, we tested whether the neuronal ensemble reactivation rate in the upper DG blade correlates with memory performance in aged mice. For this, we performed Pearson correlation analysis between LTM performance and reactivation rate in the aged population (i.e., pooled data of AI and AU mice). Remarkably, we found a significant positive correlation between the rate of neuronal ensemble reactivation in the total DG and the memory score of the mice (p=0.0272, r=0.5396, Spearman correlation, data not shown) which primarily

216	originated from the upper blade of the DG (Figure 2H), but was not present in the lower
217	blade of the DG (p=0.3668, r=0.2496, Spearman correlation, data not shown). Similar to the
218	reactivation rate, we observed a correlation between the degree of the similarity index in the
219	total DG and long-term memory performance in aged mice (p=0.0064, r=0.6437, Spearman
220	correlation, data not shown) that was selectively present in the upper blade of the DG
221	(Figure 2I), but not in the lower blade of the DG (p=0.6874, r=0.1128, Spearman correlation,
222	data not shown). Overall, these results demonstrated that cognitive performance during
223	aging positively correlates with neuronal ensemble reactivation during the recall of LTM.

224

225 4. Discussion

226 In this study, we tagged the neuronal ensemble formed during a spatial recognition task in 227 the DG of young, aged-impaired and aged-unimpaired mice and identified a cellular correlate 228 of cognitive performance in aged mice. We showed that cognitively impaired aged mice 229 exhibit impairments in recruiting the original memory engram at long-term spatial memory 230 recall compared to young or cognitively unimpaired aged mice. We further showed that the 231 degree of engram reactivation correlates with the strength of LTM in the aged population. 232 We focused on long-term object location memory which has previously been shown to be 233 impaired by aging (Oliveira et al., 2012; Wimmer et al., 2012). Increasing evidence suggests 234 that each memory is encoded by a subset of neurons (neuronal ensembles) that are 235 synchronously activated upon learning and reactivated by the retrieval of the memory 236 (Josselyn et al., 2015; Tonegawa et al., 2015). A previous study by Penner and colleagues 237 showed that neuronal reactivation upon short-term re-exposure to a previously visited 238 context is reduced in the DG, but not CA1, of aged compared to young adult mice (Penner et 239 al., 2011). These findings suggest that impairments in neuronal reactivation may be 240 associated with cognitive deficits in aged individuals (Penner et al., 2011). Here, we 241 characterized DG neuronal ensemble during the formation and retrieval of long-term 242 recognition memory and correlated with the performance in the same task. We found that 243 aging-related impairments in long-term object location memory are not associated with the

244 size of the DG neuronal population activated by learning or recall, but rather with the fidelity 245 of reactivation of the encoding population during memory retrieval. Aged impaired and 246 unimpaired mice showed similar number of neurons activated by learning or by recall. In 247 contrast, cognitively impaired mice had significantly disrupted reactivation rates compared to 248 unimpaired aged, or young adult mice. Intriguingly, the differences were found in the 249 neuronal population located in the upper, but not lower, blade of the DG. This is in line with 250 previous studies showing that behaviorally-induced expression of activity regulated genes 251 occurs primarily in the upper blade of the DG (Chawla et al., 2013, 2005; Erwin et al., 2020; 252 Gulmez Karaca et al., 2020; Marrone et al., 2012; Ramirez-Amaya et al., 2013). These 253 reports indicate that the sparse population of DG upper blade neurons form the spatial 254 memory neuronal ensemble. Therefore, it may be expected that alterations in ensemble 255 dynamics are specific to this subregion. Finally, our findings showed a positive correlation 256 between the neuronal ensemble reactivation rate in the upper blade of the DG and LTM 257 performance of aged mice. These findings suggest a novel mechanism that may underlie 258 long-term spatial memory integrity in the aged population. 259 The underlying molecular and physiological causes of disrupted neuronal ensemble 260 reactivation in aged cognitively-impaired aged mice are not understood. It is well established 261 that aging is accompanied by anatomical and physiological changes in the DG. Namely, 262 fewer synaptic contacts and impaired synaptic plasticity in aged versus young adults have 263 been reported (Burke and Barnes, 2010). Interestingly, age-related impairments in LTP 264 maintenance correlated with memory performance (Rosenzweig and Barnes, 2003). Thus, 265 deficits in the reactivation of the original encoding neuronal population may result from 266 impaired neuronal ensemble stabilization during memory consolidation, which could emerge 267 from the inability to strengthen the connectivity between ensemble neurons.

268 Recently, a few studies showed that LTM impairments in a mouse model of Alzheimer's

disease (AD) are linked to deficits in neuronal ensemble reactivation (Perusini et al., 2017;

270 Poll et al., 2020; Roy et al., 2016). Our findings now show that alterations in neuronal

271 ensemble properties are a common mechanism in aging and aging-associated pathological

- 272 conditions. This underscores the need for therapeutic approaches targeted at facilitating
- 273 engram reactivation to restore age-associated memory impairments.
- 274

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- 281

282 Disclosure statement

- 283 The authors declare no competing financial interests.
- 284

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369

370 Figure legends

371

Figure 1. Tagging neuronal ensembles formed during spatial object-location learning in the DG of

373 young, cognitively-impaired or -unimpaired aged mice. A) Schematic representation of the RAM-based

- 374 viral construct used for neuronal ensemble tagging and of the experimental design. B) Percentage of
- 375 preference for displaced object during the object location test session. Dashed line (64.3%) represents
- the threshold applied to categorize AI (n=9) and AU (n=8) mice (Y: mean=76.07, SD=11.80, n=16). C)
- 377 Total object exploration time during the training session of object-location test (p =0.9237, Kruskal-
- Wallis test with Dunn's multiple comparisons: Y (n=13) vs AU (n=8): p>0.9999; Y vs AI (n=9):
- 379 p>0.9999; AU vs AI: p>0.9999). D) Representative trajectories during open-field test. E) Total
- distance travelled in open-field test (F(2, 27)=0.0072, p =0.9303, One-way ANOVA with Tukey's

381 multiple comparisons test: Y (n=13) vs AU (n=8): p=0.9694; Y vs AI (n=9): p=0.9287; AU vs AI: 382 p=0.9936). F) Percentage of time in the center (p =0.8604, Kruskal-Wallis test with Dunn's multiple 383 comparisons: Y (n=13) vs AU (n=8): p>0.9999; Y vs AI (n=9): p>0.9999; AU vs AI: p>0.9999) and 384 number of center entries (F(2,27)=0.05143, p=0.9500, one-way ANOVA with Tukey's multiple 385 comparisons: Y (n=13) vs AU (n=8): p=0.9826; Y vs AI (n=9): p=0.9468; AU vs AI: p=0.9926) in open-386 field test. ns: not significant. ITR: inverted terminal repeat, DG: Dentate gyrus of the hippocampus, 387 Dox: doxycycline, AU: aged-unimpaired, AI: aged-impaired, Y: young, rAAV: recombinant adeno-388 associated viruses, TRE: tetracycline responsive element, pRAM: robust activity marking promoter. 389 Error bars represent s.e.m.

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391 Figure 2. Age-related cognitive decline is associated with disrupted DG neuronal ensemble 392 reactivation at LTM recall. A) Schematic representation of the experimental design. B) Representative 393 images showing the immunohistochemical analysis of HA-GFP and Fos expression in young, AU and 394 Al groups. Scale bars represent 50 µm. Dashed lines define DG granule cell layer. White arrows 395 indicate cells with overlapping HA-GFP and Fos signal, whereas cyan and magenta arrows indicate 396 individual HA-GFP and Fos signals, respectively, C) Normalized reactivation rates of learning-397 activated neuronal ensembles at memory recall (Total DG: p=0.0690, Kruskal-Wallis test with Dunn's 398 multiple comparisons: Y vs AU: p>0.9999; Y vs AI: p=0.1484; AU vs AI: p=0.1146; Upper blade: 399 p=0.0003, Kruskal-Wallis test with Dunn's multiple comparisons: Y vs AU: p>0.9999; Y vs AI: 400 p=0.0009; AU vs AI: p=0.0016; Lower blade: p=0.8702, Kruskal-Wallis test with Dunn's multiple 401 comparisons: Y vs AU: p>0.9999; Y vs AI: p>0.9999; AU vs AI: p>0.9999). D) Observed- over chance-402 overlap between HA-GFP and Fos expressions in young, AU and AI groups (Observed- vs chance-403 overlap: Total Y: W = 91, p = 0.0002 by Wilcoxon test; Total AU: W = 36, p = 0.0078 by Wilcoxon test; 404 Total AI: W = 43, p = 0.0078 by Wilcoxon test; Upper blade Y: W = 91, p = 0.0002 by Wilcoxon test; 405 Upper blade AU: W = 36, p = 0.0078 by Wilcoxon test; Upper blade AI: W = 43, p = 0.0078 by 406 Wilcoxon test: Lower blade Y: W = 89, p = 0.0005 by Wilcoxon test; Lower blade AU: W = 34, p =407 0.0156 by Wilcoxon test; Lower blade AI: W = 33, p = 0.0547 by Wilcoxon test) (Observed/chance: 408 Total DG: F(2,27)=7.201, p=0.0031, one-way ANOVA with Tukey's multiple comparisons: Y vs AU: 409 p=0.6181; Y vs AI: p=0.0152; AU vs AI: p=0.0039; Upper blade: p=0.0010, Kruskal-Wallis test with 410 Dunn's multiple comparisons: Y vs AU: p>0.9999; Y vs AI: p=0.0062; AU vs AI: p=0.0022; Lower 411 blade: F(2,27)=1.336, p=0.2797, one-way ANOVA with Tukey's multiple comparisons: Y vs AU:

412 p=0.5108; Y vs AI: p=0.7924; AU vs AI: p=0.2557). E) Normalized similarity indices of neuronal 413 ensemble populations activated at learning or at LTM recall (Total DG: p=0.0299, Kruskal-Wallis test 414 with Dunn's multiple comparisons: Y vs AU: p>0.9999; Y vs AI: p=0.0493; AU vs AI: p=0.0789; Upper 415 blade: p=0.0002, Kruskal-Wallis test with Dunn's multiple comparisons: Y vs AU: p>0.9999; Y vs AI: 416 p=0.0003; AU vs AI: p=0.0050; Lower blade: p=0.6309, Kruskal-Wallis test with Dunn's multiple 417 comparisons: Y vs AU: p>0.9999; Y vs AI: p>0.9999; AU vs AI: p>0.9999). F) Percentage of HA-GFP⁺ 418 neurons (Total DG: p=0.0275, Kruskal-Wallis test with Dunn's multiple comparisons: Y vs AU: 419 p=0.0395; Y vs AI: p=0.1667; AU vs AI: p>0.9999; Upper blade: p=0.0123, Kruskal-Wallis test with 420 Dunn's multiple comparisons: Y vs AU: p=0.0160; Y vs AI: p=0.1255; AU vs AI: p>0.9999; Lower 421 blade: p=0.0842, Kruskal-Wallis test with Dunn's multiple comparisons: Y vs AU: p=0.1364; Y vs AI: 422 p=0.2767; AU vs AI: p>0.9999). G) Percentage of Fos⁺ neurons (Total DG: p=0.1560, Kruskal-Wallis 423 test with Dunn's multiple comparisons: Y vs AU: p=0.4933; Y vs AI: p=0.2275; AU vs AI: p>0.9999; 424 Upper blade: p=0.1724, Kruskal-Wallis test with Dunn's multiple comparisons: Y vs AU: p=0.4823; Y 425 vs AI: p=0.2690; AU vs AI: p>0.9999; Lower blade: F(2,27)=3.460, p=0.0459, one-way ANOVA with 426 Tukey's multiple comparisons: Y vs AU: p=0.0665; Y vs AI: p=0.1263; AU vs AI: p=0.9298). H) 427 Spearman correlation between the normalized neuronal ensemble reactivation in the upper blade of 428 the DG and memory performance of aged mice at long-term object-location test (p<0.0001, r=0.8396, 429 n=17, Spearman correlation). I) Spearman correlation between the normalized similarity index in the 430 upper blade of the DG and memory performance of aged mice at long-term object-location test 431 (p<0.0001, r=0.8236, n=17, Spearman correlation). In all graphs: Y (n=13), AI (n=9), AU (n=8), *p<0.05: **p<0.01; ***p<0.001; [#]p<0.05; ^{##}p<0.01; ^{###}p<0.001; ns: not significant. DG: Dentate gyrus of 432 433 the hippocampus, Dox: doxycycline, AU: aged-unimpaired, AI: aged-impaired, rAAV: recombinant 434 adeno-associated viruses, pRAM: robust activity marking promoter. Error bars represent s.e.m.

Figure 1

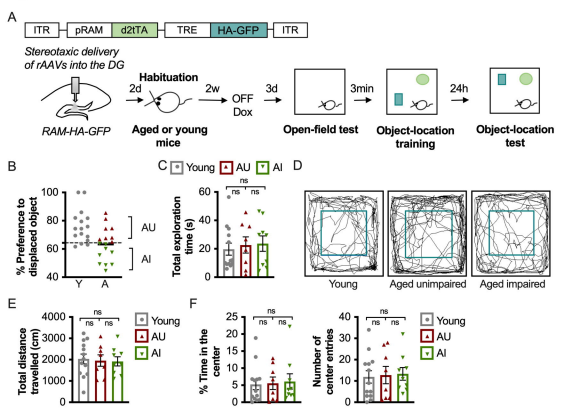


Figure 2

