

# 1        **Ankyrin2 is required for neuronal morphogenesis and long-** 2        **term memory and interacts genetically with HDAC4**

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11        **Keywords: Ankyrin repeat, Ankyrin2, ANK3, histone deacetylase, HDAC4, memory,**  
12        **neuron, *Drosophila*, mushroom body**

## 13        **Abstract**

14        Dysregulation of *HDAC4* expression and/or subcellular distribution results in impaired  
15        neuronal morphogenesis and long-term memory in *Drosophila melanogaster*. A  
16        recent genetic screen for genes that interact in the same molecular pathway as *HDAC4*  
17        identified the cytoskeletal adapter *Ankyrin2* (*Ank2*). Here we sought to investigate the  
18        role of *Ank2* in neuronal morphogenesis, learning and memory, and to examine the  
19        nature of interaction with *HDAC4*. We found that *Ank2* is expressed widely throughout  
20        the *Drosophila* brain where it localizes predominantly to axon tracts. Pan-neuronal  
21        knockdown of *Ank2* in the mushroom body, a region critical for memory formation,  
22        resulted in defects in axon morphogenesis, and similarly reduction of *Ank2* in lobular  
23        plate tangential neurons of the optic lobe disrupted dendritic branching and  
24        arborization. Conditional knockdown of *Ank2* in the mushroom body of adult  
25        *Drosophila* significantly impaired long-term courtship memory, and this requirement  
26        for *Ank2* was isolated to gamma ( $\gamma$ ) neurons of the mushroom body. As overexpression  
27        of *HDAC4* in  $\gamma$  neurons also impairs the formation of long-term courtship memory, this  
28        suggests that any functional relationship between these proteins during LTM likely  
29        occurs in  $\gamma$  neurons. We determined that the genetic interaction requires the presence  
30        of nuclear *HDAC4* and is not dependent on a conserved putative ankyrin-binding motif  
31        present in *HDAC4*. In summary, we provide the first characterization of the expression  
32        pattern of *Ank2* in the adult *Drosophila* brain and demonstrate that *Ank2* is critical for  
33        morphogenesis of the mushroom body and for the molecular processes required in  
34        the adult brain for formation of long-term memories.

## 35 36        **Introduction**

37        Histone deacetylase 4 (*HDAC4*) is a member of the Class IIa family of histone  
38        deacetylases, which are categorized by the presence of an extended N-terminal  
39        regulatory region and their ability to undergo nucleocytoplasmic shuttling  
40        (GROZINGER AND SCHREIBER 2000; MCKINSEY *et al.* 2000; WANG *et al.* 2000; WANG  
41        AND YANG 2001; CHAWLA *et al.* 2003; SCHLUMM *et al.* 2013). *HDAC4* is expressed  
42        widely throughout the brain and dysregulation of *HDAC4* expression and/or

43 subcellular distribution has been implicated in several neurodevelopmental and  
44 neurodegenerative disorders (WILLIAMS *et al.* 2010b; LI *et al.* 2012; SANDO *et al.*  
45 2012; HERRUP *et al.* 2013; MIELCAREK *et al.* 2013; SHEN *et al.* 2016; TRAZZI *et al.*  
46 2016). Loss of *HDAC4* is associated with Chromosome 2q37 deletion syndrome [MIM  
47 600430], the clinical features of which include developmental delay, autistic features  
48 and intellectual disability (WILLIAMS *et al.* 2010b; WILLIAMS *et al.* 2010a; MORRIS *et al.*  
49 2012a; MORRIS *et al.* 2012b; VILLAVICENCIO-LORINI *et al.* 2013). Developmental delay is  
50 the most common feature, and while haploinsufficiency of *HDAC4* displays variable  
51 penetrance with respect to intellectual disability (WHEELER *et al.* 2014), the  
52 predominant genetic cause is considered to be loss of function of *HDAC4* (DAG *et al.*  
53 2019). *HDAC4* has been shown to regulate synaptic plasticity and memory formation  
54 in several animal models (WANG *et al.* 2011; KIM *et al.* 2012a; SANDO *et al.* 2012;  
55 FITZSIMONS *et al.* 2013; SCHWARTZ *et al.* 2016; MAIN 2021); In mice, brain-specific  
56 conditional knockout of *HDAC4* results in impaired spatial memory (KIM *et al.* 2012b;  
57 KIM *et al.* 2012a) and in *Drosophila*, RNAi knockdown of *HDAC4* in the brain also  
58 impairs long-term memory (LTM) formation (FITZSIMONS *et al.* 2013). Disruption of  
59 normal subcellular distribution is also detrimental to neurodevelopment and  
60 cognition; a recent study identified seven individuals presenting with developmental  
61 delay, intellectual disability and other neurodevelopmental deficits, all of whom had  
62 *de novo* mutations in the 14-3-3 binding site within *HDAC4*. This motif is required for  
63 phosphorylation-dependent nucleocytoplasmic shuttling of *HDAC4*, and thus the  
64 disruption to 14-3-3 binding is presumed to disrupt normal *HDAC4* shuttling, leading  
65 to nuclear accumulation (WAKELING *et al.* 2021). In mice, a truncated mutant of *HDAC4*  
66 that accumulates in the nucleus causes deficits in spatial memory. This mutant lacks  
67 the deacetylase domain (SANDO *et al.* 2012), however vertebrate *HDAC4* is catalytically  
68 inactive (LAHM *et al.* 2007; BOTTOMLEY *et al.* 2008), reviewed by (FITZSIMONS 2015).  
69 Similarly, overexpression of *HDAC4* in the adult mushroom body, a brain region  
70 essential for memory formation in *Drosophila* (HEISENBERG *et al.* 1985; MCBRIDE *et al.*  
71 1999), prevented LTM formation, as did a catalytically inactive mutant (FITZSIMONS *et al.*  
72 2013). This memory phenotype was recapitulated with a mutant of *HDAC4* that is  
73 restricted to the nucleus, and when expressed during development it also impaired  
74 morphogenesis of mushroom body axons (MAIN 2021).

75

76 Increased nuclear *HDAC4* has also been observed in hippocampal pyramidal neurons  
77 in mouse models of Alzheimer's disease (SHEN *et al.* 2016). Moreover, *HDAC4* was  
78 found to accumulate in pyramidal cells and layer III of the frontal cortex in post-  
79 mortem brains from individuals with Alzheimer's disease, with the abundance of  
80 nuclear *HDAC4* correlating with the clinical dementia scores (HERRUP *et al.* 2013; SHEN  
81 *et al.* 2016). These data together indicate that nuclear *HDAC4* impairs cognitive  
82 function and that dysregulation of nucleocytoplasmic shuttling contributes to disease  
83 progression.

84

85 To gain further knowledge of the molecular pathway through which *HDAC4* acts, we  
86 previously carried out a genetic enhancer screen in *Drosophila* photoreceptors for  
87 genes that interact genetically with *HDAC4* (SCHWARTZ *et al.* 2016). Expression of  
88 *HDAC4* in the eye results in a mild "rough" eye phenotype, as characterized by  
89 disruption of the regular ommatidial patterning. We identified a group of cytoskeletal

90 regulators including *Ankyrin2* (*Ank2*), that when knocked down in combination with  
91 *HDAC4* overexpression resulted in an enhanced rough eye phenotype, indicative of a  
92 genetic interaction. We focused our attention on further investigating *Ank2*, since the  
93 altered expression of the human homologue *ANK3* has been associated with a variety  
94 of neurodevelopmental disorders including intellectual disability, epilepsy, attention  
95 deficit hyperactivity disorder (IQBAL *et al.* 2013), bipolar disorder (TESLI *et al.* 2011) and  
96 autism spectrum disorder (BI *et al.* 2012). Moreover, *ANK3* single nucleotide  
97 polymorphisms have been associated with schizophrenia (ATHANASIU *et al.* 2010; YUAN  
98 *et al.* 2012; NIE *et al.* 2015a; GUO *et al.* 2016; HUGHES *et al.* 2018) and Alzheimer's  
99 disease (MORGAN *et al.* 2008).

100

101 Ankyrins are adapter proteins that link the underlying spectrin-actin cytoskeleton to  
102 integral membrane proteins such as ion channels, anion exchangers, signaling proteins  
103 and cell adhesion molecules (BENNETT 1978; BENNETT AND BAINES 2001; MOHLER *et al.*  
104 2002). Canonical ankyrins are comprised of an N-terminal membrane binding domain,  
105 which contains 24 ANK repeat motifs organized as two anti-parallel  $\alpha$ -helices that  
106 mediate protein-protein interactions. They also contain a central spectrin-binding  
107 domain, a death domain and a less conserved C-terminal regulatory domain (SEDGWICK  
108 AND SMERDON 1999; CUNHA AND MOHLER 2009). *Drosophila Ank2* displays high similarity  
109 to human *ANK3* (IQBAL *et al.* 2013), sharing 57% amino acid identity over the whole  
110 protein and 71.2% identity across the ankyrin repeat region. Similarly to *ANK3*,  
111 expression of *Ank2* is restricted to neurons and a number of transcript isoforms are  
112 also expressed. In the embryonic nervous system, shorter isoforms of *Ank2* (*Ank2-S*)  
113 localize to the cell bodies of neurons, whereas medium *Ank2-M* isoforms localize to  
114 axons and are essential for viability (HORTSCH *et al.* 2002). A longer *Ank2-L* isoform  
115 containing an additional C-terminal domain localizes to axons and synaptic boutons at  
116 the neuromuscular junction. Mutants with disrupted *Ank2-L* expression display a loss  
117 of synapse stability as well as a reduction in the terminal bouton size, disassembly of  
118 presynaptic active zone and retraction of the synaptic microtubule cytoskeleton (KOCH  
119 *et al.* 2008; PIELAGE *et al.* 2008), thus *Ank2-L* is required for synapse stability and normal  
120 morphology at the neuromuscular junction in *Drosophila* larvae.

121

122 Despite comprehensive investigation of the roles that *Ank2* plays in neuronal  
123 development, the role of *Ank2* in the adult brain and functional consequences of  
124 reduced *Ank2* has received little attention. Here we aimed to investigate the role of  
125 *Ank2* in learning, memory and development of the *Drosophila* brain as well as to  
126 examine the nature of the interaction between *HDAC4* and *Ank2* and to determine  
127 whether this is an important mechanism through which *HDAC4* regulates neuronal  
128 development and memory.

129

## 130 **METHODS**

### 131 **Fly strains**

132 All flies were raised on standard medium on a 12-hour light/dark cycle and maintained  
133 at a temperature of 25°C unless otherwise indicated. *w*[\*];  
134 *P*{*w*[+*mW.hs*]=*GawB*}*OK107 ey*[*OK107*]/*In*(4)*ci*[*D*], *ci*[*D*] *pan*[*ciD*] *sv*[*spa-pol*] (*OK107-*  
135 *GAL4*), *w*[\*]; *P*{*w*[+*mC*]=*GAL4-ninaE.GMR*}12 (*GMR-GAL4*),  
136 *P*{*w*[+*mW.hs*]=*GawB*}*elav*[*c155*] (*elav-GAL4*), *P*{*w*+*mW.hs*=*GawB*}*c739* (*c739-GAL4*),

137 *w<sup>1118</sup>;P{w+mW.hs=GawB}c305a (c305a-GAL4), w<sup>1118</sup>;P{w+mW.hs=GawB}1471 (1471-*  
138 *GAL4), w[1118]; P{y[+t7.7] w[+mC]=GMR16A06-GAL4}attP2 (R16A06-GAL4) and*  
139 *P{w[+mW.hs]=GawB}3A (3A-GAL4) were obtained from the Bloomington *Drosophila**  
140 *Stock Center. w[\*]; P{w{+mW.hs}=GawB}NP1131 (NP1131-GAL4) and w[1118];*  
141 *PBac{EGFP-IV}ank2[KM0104] (Ank2::EGFP) were obtained from the Kyoto Stock*  
142 *Center. P{w+mC=tubP-GAL80<sup>ts</sup>}10 (tubP-GAL80<sup>ts</sup>), p{MEF2-GAL4.247}(MB247-GAL4)*  
143 *and w(CS10) strains were kindly provided by R. Davis (The Scripps Research Institute,*  
144 *Jupiter, FL). P{KK106729}VIE-260B (UAS-Ank2 RNAi, VDRC ID 107369) and*  
145 *w<sup>1118</sup>;P{GD12247}v40638 (UAS-Ank2 RNAi, VDRC ID 40638) were obtained from the*  
146 *Vienna *Drosophila* Resource Center. All strains were outcrossed for a minimum of five*  
147 *generations to w(CS10) flies. A homozygous line harbouring w(CS10); P{w+mC=tubP-*  
148 *GAL80<sup>ts</sup>}10 and P{w+mW.hs=GawB}OK107 (tubP-GAL80<sup>ts</sup>; OK107-GAL4) was*  
149 *generated by standard genetic crosses, as was (elav-GAL4; tubP-GAL80<sup>ts</sup>), (c739-GAL4;*  
150 *tubP-GAL80<sup>ts</sup>), (tubP-GAL80<sup>ts</sup>, 1471-GAL4), (tubP-GAL80<sup>ts</sup>; MB247-GAL4), (tubP-*  
151 *GAL80<sup>ts</sup>, c305a-GAL4) (tubP-GAL80<sup>ts</sup>; R16A06-GAL4) and (tubP-GAL80<sup>ts</sup>, NP1131-*  
152 *GAL4). UAS-DmHDAC4 and UAS-DmHDAC4 3SA have been described previously (MAIN*  
153 *2021). UAS-DmHDAC4 ΔAnk was generated by site directed mutagenesis of UAS-*  
154 *DmHDAC4 with the following amino acid substitutions: P48A L50A P51A and I53A.*  
155 *The UAS-Ank2<sub>190-946</sub>-HA construct consists of a 2268 bp N-terminal region of Ank2*  
156 *containing the ankyrin repeat region (nucleotides 1123 - 3393 of Ank2, NCBI reference*  
157 *NM\_001274607, which corresponds to amino acids 190 - 946). This construct with a*  
158 *C-terminal 3x HA epitope tag was generated and subcloned into pUASTattB by*  
159 *Genscript (NJ, USA). Transgenic flies were generated by Genetivision (Houston, TX,*  
160 *USA) using the VK22 docking site at 2R(57F5).*

161

## 162 **Immunohistochemistry**

163 Whole flies were fixed in PFAT/DMSO (4% paraformaldehyde in 1X PBS +0.1% Triton  
164 X-100 +5% DMSO) for one hour then brains were microdissected in 1X PBS. Brains  
165 were post-fixed in PFAT/DMSO for 20 mins and blocked in immunobuffer (5% normal  
166 goat serum in 1X PBS +0.5% Triton X-100) for three hours prior to incubation with  
167 primary antibody of rabbit anti-Ank2-L (1:1000, gift from H. Aberle) (Koch *et al.* 2008),  
168 rabbit anti-GFP (Abcam, ab290 1:10,000), mouse anti-nc82 (1:100), mouse anti-Futsch  
169 (1:20), mouse anti-Repo (1:20) and mouse anti-Fasciclin II (Fas II, 1:200). Brains were  
170 then incubated with secondary antibody (goat anti-mouse Alexa 488 or 555, or goat  
171 anti-rabbit Alexa 488 or 555, Sigma Aldrich, 1:500) and mounted with Antifade. The  
172 monoclonal antibodies anti-Brp (nc82, developed by E. Buchner), anti-futsch (22C10,  
173 developed by S. Benzer and N. Colley), anti-Repo (8D12, developed by C. Goodman),  
174 and anti-FasII (1D4, developed by C. Goodman) were obtained from the  
175 Developmental Studies Hybridoma Bank developed under the auspices of the NICHD  
176 and maintained by The University of Iowa, Department of Biology, Iowa City, IA 52242.  
177 For confocal microscopy, images were captured with a Leica TCS SP5 DM6000B  
178 Confocal Microscope and images were processed with Leica Application Suite  
179 Advanced Fluorescence (LAS AF) software and Image J (NIH). For quantification of  
180 dendrite branch length, the total shaft and major branch length inclusive of all six  
181 neuronal shafts and their major visible branches were traced using the SNT program  
182 in the ImageJ NeuroAnatomy plugin, which allows branching trace plots to be  
183 reproduced from the dendritic arborizations (Avery *et al.*, 2017). Total shaft and

184 branch lengths were traced and these measurements were then added together to  
185 produce a total sum branch length. Statistical analysis was carried out with the  
186 student's *t*-test with significance set at  $p < 0.05$ .

187

### 188 **Western Blotting**

189 Whole cell extracts were prepared from 100 snap-frozen heads by homogenizing in  
190 RIPA buffer, followed by centrifugation at 13,000 g for 2 minutes at 4°C. Lysates (30  
191 µg) were resolved on 4-20% SDS-PAGE gels (Biorad) and transferred onto  
192 nitrocellulose membranes. Membranes were blocked for >1 hour at room  
193 temperature in 5% skim milk powder in TBST (50 mM Tris, 150 mM NaCl, 0.05%  
194 Tween-20, pH 7.6) then incubated overnight at 4°C in primary antibody, washed 3 x 5  
195 mins in TBST then incubated one hour in secondary anti-mouse, anti-rat or anti-rabbit  
196 HRP-conjugated antibodies (GE Life Sciences) as appropriate. Following 3 x 5 min  
197 washes in TBST, proteins were detected with Amersham ECL Prime (GE Life Sciences).  
198 The following antibodies were used: rabbit anti-GFP (Abcam ab290, 1:10,000); rabbit  
199 anti-Myc (Roche, 1:1,000); rat anti-HA (Roche, 1:1,000) and mouse anti- $\alpha$ -tubulin  
200 (12G10 clone, DSHB, 1:500).

201

### 202 **Co-immunoprecipitation**

203 Whole cell extracts were prepared as per the western blotting method above.  
204 Immunoprecipitation (IP) was performed with the Pierce Classic IP Kit (Thermo  
205 Scientific) according to the manufacturer's instructions. Anti-Myc or anti-HA antibody  
206 (1 µL) was incubated overnight with 1 mg of lysate. Following elution in 2x sample  
207 buffer, IP samples were processed for SDS-PAGE and western blotting with anti-HA or  
208 anti-Myc alongside 30 µg input samples. Anti- $\alpha$ -tubulin (1:500) was used as a loading  
209 control.

210

### 211 **RT-qPCR**

212 *elav-GAL4* females were crossed to *UAS-Ank2* RNAi males to generate progeny in  
213 which *Ank2* was knocked down in all neurons; and progeny of *elav-GAL4* crossed to  
214 *w(CS10)* served as the control. To confirm knockdown, total RNA was extracted from  
215 *Drosophila* heads from three independent crosses with the RNeasy Mini kit (Qiagen)  
216 according to the manufacturer's instructions. cDNA was synthesized from 1 µg of  
217 total RNA with Transcriptor (Roche) as per the manufacturer's instructions. RT-qPCR  
218 was conducted using SsoFast-EvaGreen (BioRad) reaction master on a Lightcycler II  
219 480 instrument (Roche), following manufacturer's instructions. The following primers  
220 were used: *Ank2*for 5'-GGCCGATATGGCACAAAACC-3', *Ank2*rev  
221 5'TTCTTTCGACGGTGGTACGG-3', *EF1 $\alpha$ 48D*for 5'-ACTTTGTTCGAATCCGTCGC-3',  
222 *EF1 $\alpha$ 48D*rev 5'-TACGCTTGTCGATACCACCG-3'. A 5-fold dilution of cDNA from control  
223 flies was used as template to prepare a standard curve to confirm efficiency of the  
224 PCR reactions. Relative quantification was conducted using  $2^{-\Delta\Delta Ct}$  method, normalizing  
225 to the housekeeping gene *Ef1 $\alpha$ 48D* (LIVAK AND SCHMITTGEN 2001). *Ank2* expression was  
226 reduced to  $0.42 \pm 0.12$  (mean  $\pm$  standard error) of that of the control, student's *t*-test  
227  $t_{(12)}=4.74$ ,  $p < 0.001$ .

228

### 229 **Courtship Suppression Assay**

230 The repeat training courtship suppression assay was used to assess 24-hour long-term  
231 courtship memory. This is an experience-dependent assay in which wild-type male  
232 flies that have been previously rejected by a mated female will reduce their courtship  
233 behavior towards a new mated female. During mating, the male pheromone cVA is  
234 transferred to the female, and the presence of this pheromone on the female causes  
235 the male to reduce his courtship towards her. Males that have previously experienced  
236 rejection will suppress courtship towards another mated female due to an enhanced  
237 response to cVA (EJIMA *et al.* 2005; KELEMAN *et al.* 2007) and this form of courtship  
238 memory is termed cVA-retrievable memory (RAUN *et al.* 2021). The detailed  
239 methodology has been described previously (FITZSIMONS AND SCOTT 2011; FITZSIMONS *et*  
240 *al.* 2013; FREYMUTH AND FITZSIMONS 2017). For training, single virgin males (3-5 days post  
241 eclosion) of each genotype were placed into individual training chambers. A freshly  
242 mated wild-type female was placed with each male to be trained, whereas sham  
243 control males were housed alone. Over the seven-hour training period, multiple bouts  
244 of courtship were observed in the trained group. The female fly was then aspirated  
245 from the training chamber and the males were left in their chambers for the 24 hours,  
246 prior to testing. Each trained or sham male fly was then placed into a testing chamber  
247 containing a mated wild-type female and was scored for the time spent performing  
248 stereotypic courtship behaviors over the ten-minute period. A courtship index (CI) was  
249 calculated as the proportion of the ten-minute period spent courting. A mean CI for  
250 each group was determined, and from this a memory index (MI) was calculated by the  
251 following equation:  $MI = 1 - (CI \text{ of each trained fly} / \text{mean CI of sham group})$   
252 ( $n \geq 16/\text{group}$ ). The MI was measured on a scale of 0 to 1, a score of 0 indicating  
253 memory was no different than untrained sham controls. In all experiments, the scorer  
254 was blind to the genotype of the flies. For assessment of immediate short-term  
255 memory, the training session was reduced to one hour and flies were tested  
256 immediately after training. For assessment of learning, the male was placed with a  
257 mated female for an hour and the first ten minutes and last ten minutes were scored  
258 for courtship behavior. The learning index was calculated as  $1 - (CI \text{ last 10 mins} / CI \text{ first}$   
259  $10 \text{ mins})$ . For statistical analyses, data was arcsine transformed to approximate a  
260 normal distribution and one-way ANOVA with post-hoc Tukey's HSD test was used to  
261 assess significance ( $p < 0.05$ ).

262

### 263 **Scanning electron microscopy (SEM)**

264 Flies were anaesthetised using FlyNap (Carolina Biologicals) and fixed in primary  
265 modified Karnovsky's fixative (3% glutaraldehyde, 2% formaldehyde in 0.1 M  
266 phosphate buffer, pH 7.2) with Triton X-100 by vacuum infiltration. They were then  
267 placed in fresh fixative and incubated at room temperature > 8 hours, followed by 3 x  
268 10 min washes in phosphate buffer (0.1 M, pH 7.2). Dehydration was carried out via a  
269 graded ethanol series for ten to fifteen minutes at 25%, 50%, 75%, 95%, and 100%  
270 ethanol, followed by a final one-hour incubation in 100% ethanol. The flies were then  
271 critical point dried using CO<sub>2</sub> and 100% ethanol (Polaron E3000 series II drying  
272 apparatus). Heads were removed, mounted onto aluminium stubs and sputter coated  
273 with gold (Baltex SCD 050 sputter coater). Imaging was performed with a FEI Quanta  
274 200 Environmental Scanning Electron Microscope at an accelerating voltage of 20 kV.  
275 To determine the sizes of each eye for comparison between genotypes, ImageJ  
276 software was used to draw a line surrounding the eye and calculate the area in

277 arbitrary units. Statistical analysis was performed with one-way ANOVA with post-hoc  
278 Tukey's HSD test with significance set at  $p < 0.05$ .

279

280 To provide a semi-quantitative analysis of SEM images, a scoring system was  
281 developed based on observations of phenotypes resulting from overexpression of one  
282 or two copies of *UAS-HDAC4* in a previous study (SCHWARTZ *et al.* 2016). No defects:  
283 the eye appears wild-type - ommatidia are organised in a regular array with no fusion  
284 and mechanosensory bristles are correctly positioned between each ommatidium.  
285 Mild: Presence of one of the following phenotypes; between 5 - 10 instances of an  
286 abnormal number of interommatidial bristles, mild ommatidial disorganisation or  
287 fusion of ommatidia in up to two areas. Moderate: all mild phenotypes were  
288 collectively observed or one of the following phenotypes was observed; between 10  
289 - 20 instances of an abnormal number of interommatidial bristles, moderate  
290 disorganisation or fusion of ommatidia in up to five areas. Major: all moderate  
291 phenotypes were collectively observed or one of the following phenotypes was  
292 observed; more than 20 instances of an abnormal number of interommatidial bristles,  
293 major disorganisation of the ommatidial array, fusion of ommatidia in up to 10 areas  
294 with few large areas of fusion or up to 50 collapsed ommatidia. Severe: all major  
295 phenotypes were collectively observed or one of the following phenotypes was  
296 observed; severe disorganisation, fusion in more than 10 areas or multiple large  
297 patches, more than 50 collapsed ommatidia or severe collapsing of ommatidia  
298 resulting in central hole-like cavities. Statistical analysis was assessed with the Fisher's  
299 Exact test with significance set at  $p < 0.05$ .

300

## 301 **RESULTS**

### 302 **Characterisation of Ank2 expression in the brain**

303 To date, the expression and localisation pattern of Ank2 has been described in the  
304 *Drosophila* neuromuscular junction (KOCH *et al.* 2008; PIELAGE *et al.* 2008) however the  
305 expression pattern in the adult brain has not been characterized.  
306 Immunohistochemistry on whole mount brains with an antibody that detects Ank2-L  
307 (KOCH *et al.* 2008) indicated a broad expression profile with high expression in the optic  
308 lobes, antennal lobes, mushroom body and axon tracts throughout the brain (Fig  
309 1A,B). Colocalization with the axonal marker Futsch (HUMMEL *et al.* 2000) confirmed  
310 that Ank2-L localizes to axon tracts across the adult brain (Fig 1C). Ank2-L did not co-  
311 distribute with the glial marker Reversed Polarity (Repo) (ALFONSO AND JONES 2002),  
312 confirming its specific neuronal expression pattern (Fig 1D). Since the mushroom body  
313 is a critical structure for memory (HEISENBERG *et al.* 1985; MCBRIDE *et al.* 1999), we  
314 examined the expression and subcellular distribution of Ank2 in this brain area in more  
315 detail. The intrinsic neurons of the mushroom body are the Kenyon cells, which  
316 receive input from the olfactory system (TURNER *et al.* 2008). The cell bodies of the  
317 approximately 2,500 Kenyon cells are clustered in the posterior dorsal region of the  
318 brain and extend their dendrites anteriorly into a globular region known as the calyx,  
319 which is organized into an array of microglomeruli, each comprising the large synaptic  
320 bouton of projection neurons from the antennal lobe surrounded by Kenyon cell  
321 dendrites (LEISS *et al.* 2009). Their axons form a bundled fiber termed the pedunculus  
322 and project towards the anterior portion of the brain, forming five distinct lobes; the  
323 vertical  $\alpha$  and  $\alpha'$  lobes and the medial  $\beta$ ,  $\beta'$  and  $\gamma$  lobes (CRITTENDEN *et al.* 1998; LEE AND

324 Luo 1999) (see Fig. 3F). We examined the colocalization of Ank2-L with the neuronal  
325 cell adhesion molecule Neuroglian (Nrg), which has been shown to interact with Ank2  
326 (ENNEKING *et al.* 2013; SIEGENTHALER *et al.* 2015). Nrg is the sole *Drosophila* orthologue  
327 of the L1-CAM family of proteins (BIEBER *et al.* 1989), which enables axon guidance  
328 through the mushroom body. We confirmed that Ank2-L and Nrg codistributed in  
329 multiple axon tracts including the axons of the mushroom body, where both were  
330 observed in the  $\alpha$ ,  $\beta$  and  $\gamma$  lobes (Fig 1E). Ank2-L was also concentrated in axon tracts  
331 surrounding the calyx of the mushroom body (Fig 1F).

332

### 333 **Ank2 is essential for axon and dendrite morphogenesis**

334 We previously found that overexpression of *HDAC4* impairs morphogenesis of  
335 mushroom body axons, with deficits in axon branching and guidance observed as  
336 missing  $\alpha$  and/or  $\beta$  lobes, misdirected axons as well as the appearance of fused  $\beta$   
337 lobes, resulting from defects in axon termination across the midline (MAIN 2021). To  
338 investigate whether Ank2 is also required for axon morphogenesis, the morphology of  
339 the mushroom body in brains with reduced Ank2 was analyzed via detection of  
340 Fasciclin II. This cell adhesion molecule is highly expressed in the  $\alpha$ ,  $\beta$  and  $\gamma$  lobes of  
341 the mushroom body (CRITTENDEN *et al.* 1998) and is a commonly used marker to  
342 visualise mushroom body lobe architecture (Fig 2A). Pan-neuronal knockdown of *Ank2*  
343 with an inverted repeat hairpin that targets all long isoforms of *Ank2* mRNA for  
344 degradation resulted in a variety of phenotypic defects of the mushroom body,  
345 including thin lobes, missing lobes and guidance abnormalities (Fig 2B-F). GAL4 activity  
346 increases at higher temperatures and accordingly we observed more severe defects  
347 when the temperature was raised during larval development (Table 1).

348

349 During *Drosophila* embryonic and larval stages, *Ank2* mutants exhibit reduced  
350 dendritic branching, and in *Drosophila* dopaminergic neurons, knockdown of *Ank2*  
351 results in decreased dendritic branching points, leading to a reduced total branch  
352 length and a lack of branching complexity (AVERY *et al.* 2017). To that end, we  
353 investigated whether Ank2 is required for dendrite morphogenesis in the adult  
354 *Drosophila* brain. As branching and elongation of Kenyon cell dendrites is difficult to  
355 visualize, we instead examined lobular plate tangential cells (LPTCs) of the visual  
356 system. The LPTCs are a group of six interneurons in the optic lobe that provide an  
357 ideal model system for investigating dendrite growth and branching as they display  
358 stereotypical dendritic arborization (LEISS *et al.* 2009). Individual dendrites are easily  
359 visualised via expression of Lifeact, a GFP-fused F-actin binding peptide (RIEDL *et al.*  
360 2008) with the *3A-GAL4* driver (SCOTT *et al.* 2002) (Figure 2G), and branch length can  
361 be traced and quantified (Figure 2H, I). The characteristic arborization pattern of the  
362 six neurons was disrupted by expression of *Ank2* RNAi with severely reduced dendritic  
363 projections (Figure 2J, K) leading to reduced total branch length (Figure 2L). These data  
364 suggest that wild-type levels of Ank2 are required for both axon branching, guidance  
365 and elongation as well as normal dendritic branching and arborization.

366

### 367 **LTM requires Ank2 expression in the $\gamma$ lobe of the mushroom body**

368 We next assessed whether Ank2 was required for memory formation with the repeat  
369 training courtship suppression assay. This test evaluates the memory of a male  
370 following exposure to an unreceptive mated female. Following this failure of mating,



371 a male suppresses his courtship activity towards mated females to which he is  
372 subsequently presented. After seven hours of training, males form a stable long-term  
373 memory that lasts for at least 24 hours (KELEMAN *et al.* 2007; FITZSIMONS AND SCOTT 2011;  
374 FITZSIMONS *et al.* 2013). After this time, each male is placed with a new freshly mated  
375 (unreceptive) female and a courtship index is calculated by dividing the amount of  
376 time each male spends courting by the total duration of the observation period. A  
377 memory index is calculated by comparing the time a trained male spends courting to  
378 that of a sham male. A score of zero indicates that memory is impaired and no  
379 different from untrained sham controls, whereas a higher memory index indicates  
380 intact memory. This form of courtship memory has been recently described as “cVA-  
381 retrievable memory” to differentiate from the associative memory formed when  
382 virgin females are used for testing, which uses different circuitry for memory retrieval  
383 (RAUN *et al.* 2021). Learning and immediate short-term memory were unaffected by  
384 pan-neuronal knockdown of *Ank2* (Fig 3A,B). Long-term courtship memory is  
385 dependent on an intact mushroom body, therefore unsurprisingly, pan-neuronal  
386 knockdown of *Ank2* during development resulted in a significant and severe loss of  
387 LTM formation compared to control genotypes (Fig 3C). This was not due to an effect  
388 on courtship behavior as sham males of each genotype all spent approximately the  
389 same percentage of time courting (87 to 89%, Figure 3D).

390  
391 To avoid the developmental deficits resulting from decreased *Ank2* expression and  
392 allow assessment of the role of *Ank2* specifically in adult memory processes,  
393 knockdown of *Ank2* was restricted to the mature brain with GAL80ts, a temperature  
394 sensitive inhibitor of GAL4 activity (MCGUIRE *et al.* 2004) (Fig 3E). Flies were raised at  
395 the permissive temperature of 19°C at which GAL80ts is active. Seventy-two hours  
396 after eclosion, male flies from the F1 progeny were collected individually and  
397 transferred to 30°C to inactivate GAL80ts and thus induce pan-neuronal RNAi  
398 expression. After three days, males were tested in the courtship suppression assay.  
399 Adult-specific knockdown of *Ank2* in all neurons resulted in impairment of LTM  
400 formation (Fig 3G) and when knockdown of *Ank2* was restricted to the adult  
401 mushroom body (Fig 3F) with *OK107-GAL4*, this impairment remained (Fig 3H). The  
402 three Kenyon cell subtypes are structurally distinct with individually identifiable  
403 transcriptomes (CROSET *et al.* 2018) and distinct roles in learning and memory (JOINER  
404 AND GRIFFITH 1999; KELEMAN *et al.* 2007), thus we next investigated whether there was  
405 a differential requirement for *Ank2* in specific mushroom body subtypes by restricting  
406 expression of *Ank2* RNAi to each subtype individually (Fig 3F). Expression with the  $\alpha/\beta$   
407 and  $\gamma$  neuron driver *MB247-GAL4* abolished LTM formation (Fig 3I). Knockdown in the  
408  $\alpha/\beta$  neurons or  $\alpha'/\beta'$  neurons did not significantly alter LTM (Fig 3J,K) whereas  
409 knockdown in  $\gamma$  neurons with *NP1131-GAL4* prevented LTM formation (Fig 3L). The  $\gamma$   
410 neuron driver *1471-GAL4* did not quite impair memory to significant levels (Fig 3M),  
411 however this is a much weaker than *NP1131-GAL4* (Aso *et al.* 2009). We therefore  
412 tested an additional  $\gamma$  neuron driver *R16A06-GAL4* which drives very strong expression  
413 in the gamma lobe, but minimal expression elsewhere in the brain (JENETT *et al.* 2012),  
414 and employed a second independent *Ank2* RNAi line that also targets all long forms of  
415 *Ank2* mRNA, which together resulted in a significant reduction in LTM (Fig 3N).

416

417 Taken together these data show that Ank2-L is required for normal mushroom body  
418 development and in the adult brain, wild-type levels of Ank2-L are required in the  $\gamma$   
419 lobes for normal LTM formation. This is strikingly similar to the phenotypes resulting  
420 from manipulation of HDAC4 expression in that it is also required in the  $\gamma$  lobe for LTM  
421 but not STM (FITZSIMONS *et al.* 2013).

422

#### 423 **HDAC4 does not physically interact with Ank2 in *Drosophila* neurons**

424 The N-terminal region of HDAC4 contains an ankyrin repeat binding domain consisting  
425 of a PxLPxI/L motif (Fig 4A), which in mammalian cells binds the ankyrin repeat region  
426 of ANKRA2 and RFXANK (WANG *et al.* 2005; XU *et al.* 2012; NIE *et al.* 2015b). This motif  
427 is conserved in *Drosophila* HDAC4, and the ankyrin repeat of RFXANK and ANKRA2  
428 both share 52% amino acid similarity with that of Ank2, thus it is possible that Ank2  
429 and HDAC4 could interact physically through the ankyrin repeat of Ank2. To test for a  
430 physical interaction, we generated flies that co-express the ankyrin repeat-containing  
431 domain of Ank2 with a C-terminal HA tag (UAS-Ank2<sub>190-946</sub>-HA) and Myc-tagged  
432 *Drosophila* HDAC4 in the brain. Coimmunoprecipitation on head lysates revealed no  
433 detectable interaction of HDAC4 with Ank2 via pulldown with either anti-HA or anti-  
434 Myc (Fig 4B,C), therefore while *Ank2* and *HDAC4* interact genetically, they do not  
435 appear to physically interact at a detectable level in the brain. An obvious explanation  
436 for the genetic interaction between the two would be that HDAC4 regulates  
437 expression of *Ank2*, however previous RNA-seq data in which we expressed DmHDAC4  
438 or a nuclear-restricted mutant of human HDAC4 did not show a significant change in  
439 transcription of *Ank2* (SCHWARTZ *et al.* 2016; MAIN 2021). We also confirmed that  
440 increased HDAC4 expression does not alter the level of Ank2 protein (Fig 4D,E).

441

#### 442 **Nuclear HDAC4 mediates the genetic interaction with Ank2**

443 We previously showed that knockdown of *Ank2* in the eye resulted in a mild  
444 developmental impairment that was significantly enhanced when combined with  
445 *HDAC4* overexpression, indicating the two genes interact in the same molecular  
446 pathway (SCHWARTZ *et al.* 2016). The HDAC4-induced impairments in eye development  
447 are largely a consequence of nuclear accumulation of HDAC4, as expression of a  
448 mutant variant of human HDAC4 that is sequestered in the nucleus resulted in a more  
449 severe phenotype than overexpression of wild-type human *HDAC4*, whereas the  
450 phenotype resulting from expression of a cytoplasm-restricted mutant was mild (MAIN  
451 2021). To that end, we next investigated whether the genetic interaction between  
452 *Ank2* and *HDAC4* in the eye is also dependent on the nuclear presence of *Drosophila*  
453 HDAC4. In addition, we sought to determine whether the genetic interaction is  
454 dependent on the putative ankyrin repeat-binding motif region of HDAC4. If not, this  
455 would provide further confirmation that the genetic interaction between *HDAC4* and  
456 *Ank2* is not through direct physical binding.

457

458 We first confirmed our previous observations by co-expressing *DmHDAC4* and *Ank2*  
459 RNAi under the control of *GMR-GAL4*, which drives expression in post-mitotic cells  
460 posterior to the morphogenetic furrow (FREEMAN 1996), and examining the phenotype  
461 of adult eyes. We also raised the temperature to 27°C to increase GAL4 activity and  
462 thus increase the degree of knockdown in order to examine the resulting phenotypic  
463 defects in eye development in more detail. Control flies (*GMR-GAL4* crossed to the

464 background *w[CS10]* strain) showed predominantly normal ommatidial alignment and  
465 no evidence of fusion (Fig 5A), whereas eyes with reduced *Ank2* expression had fused,  
466 collapsed and misaligned ommatidia lacking some interommatidial bristles (Fig 5B).  
467 Overexpression of *DmHDAC4* also resulted in missing/disorganised bristles and  
468 misaligned and fused ommatidia (Fig 5C). Combined expression of *DmHDAC4* and  
469 *Ank2* RNAi resulted in a more severe rough eye phenotype consisting of major areas  
470 of ommatidial fusion, severe misalignment with hole-like cavities within the  
471 ommatidia, and missing/disorganised bristles (Fig 5D). The severity of phenotypes was  
472 scored (Table S1) and the percentage of eyes displaying the most severe phenotype  
473 was significantly higher when *DmHDAC4* and *Ank2* RNAi were combined (Fig 5I). In  
474 addition to the severe eye phenotypes observed above, the eyes were also  
475 significantly physically smaller than those expressing either *Ank2* RNAi or *DmHDAC4*  
476 individually (Fig 5J). Together these synergistic phenotypes are consistent with our  
477 previous findings (SCHWARTZ *et al.* 2016) and provide further confirmation that *Ank2*  
478 and *HDAC4* genetically interact together to influence eye development.

479  
480 To further investigate the mechanism of this interaction, we examined whether it was  
481 dependent on the nuclear activity of HDAC4. A nuclear-restricted mutant of *HDAC4*  
482 (*DmHDAC4 3SA*) resulted in a more severe phenotype than wild-type HDAC4, with  
483 increased disorganisation and fusion of ommatidia and a reduced number of bristles  
484 (Fig 5E), which we have previously observed (MAIN 2021). *DmHDAC4 3SA* interacted  
485 synergistically with *Ank2* (Fig 5F), confirming that nuclear activity of HDAC4 is required  
486 for the genetic interaction. To address whether the putative ankyrin binding motif in  
487 *Drosophila* HDAC4 is required we substituted residues that have been shown to be  
488 important for this interaction in mammals; P48 L50 P51 and I53 of the PxLPxI/L motif  
489 (Fig 4A) to alanines to create the *DmHDAC4 ΔAnk mutant*. Expression of this mutant  
490 also resulted in a moderate rough eye phenotype consisting of ommatidial fusion and  
491 misalignment (Figure 5G), indicating that the presence of this motif is not required for  
492 the *HDAC4* overexpression-induced eye defects. This mutant still interacted  
493 genetically with *Ank2*, resulting in a significantly more severe rough eye phenotype  
494 and reduced eye size when expressed in combination with *Ank2* (Fig 5H), therefore  
495 this interaction does not depend on the presence of the putative ankyrin-binding  
496 motif of HDAC4, which is consistent with the lack of physical interaction with *Ank2*.

## 497 498 **DISCUSSION**

499 Here, we provide the first characterization of the expression pattern of *Ank2* in the  
500 adult *Drosophila* brain and demonstrate that *Ank2* is critical for normal development  
501 of the mushroom body and for formation of long-term memories.

502  
503 *Ank2* expression was observed throughout the brain and localized predominantly to  
504 axon tracts, including the lobes of the mushroom body where it colocalized with *Nrg*.  
505 As the mushroom body is a critical structure for memory formation (HEISENBERG *et al.*  
506 1985; MCBRIDE *et al.* 1999), and we previously showed that HDAC4 is required in the  
507 mushroom body for normal memory formation, and that overexpression of HDAC4  
508 impairs mushroom body development (FITZSIMONS *et al.* 2013; MAIN 2021), we focused  
509 our attention on examining the role of *Ank2* in both developmental and post-  
510 developmental processes in this brain region. Pan-neuronal knockdown of *Ank2* in the

511 developing mushroom body resulted in deficits in axon guidance and elongation. *Ank2*  
512 has been demonstrated to interact with *Nrg* through a FIGQY motif within the  
513 intracellular domain of *Nrg* (ENNEKING *et al.* 2013). *Nrg* is required for normal  
514 mushroom body lobe development, whereby the extracellular domain of *Nrg*  
515 mediates cellular adhesion between axons of different mushroom body subtypes for  
516 guidance into the pedunculus and lobes. This interaction relies on the presence of  
517 *Ank2* in either the ingrowing or substrate axon neurons where it is proposed to  
518 stabilise the transaxonal *Nrg* complex (SIEGENTHALER *et al.* 2015). A hypomorphic *Nrg*  
519 mutant displayed deficits in axon growth and guidance in the mushroom body  
520 (SIEGENTHALER *et al.* 2015), which are very similar to those we observed on knockdown  
521 of *Ank2*. In addition, *Nrg* also interacts with a second cytoskeletal adapter protein  
522 Moesin (*Moe*) through a FERM domain in the intracellular domain of L1 CAMs (DICKSON  
523 *et al.* 2002), creating a ternary complex between *Ank2*, *Nrg* and *Moe* (SIEGENTHALER *et al.*  
524 *et al.* 2015). *Moe* is highly expressed in the mushroom body and distributes to the lobes  
525 on activation by phosphorylation (FREYMUTH AND FITZSIMONS 2017), and likely acts to link  
526 the *Ank2*-*Nrg* complex to the actin cytoskeleton in mushroom body axons.  
527 Interestingly we also previously showed that *HDAC4* interacts genetically with *Moe*  
528 (SCHWARTZ *et al.* 2016), and moreover, reduction of *Moe* also shows similar disruption  
529 to mushroom body development as with *Ank2* and *Nrg*, with defects in axon  
530 elongation and guidance (SIEGENTHALER *et al.* 2015; FREYMUTH AND FITZSIMONS 2017).  
531 Together these data support the evidence for a functional relationship between *Ank2*,  
532 *Nrg* and *Moe* in mushroom body development.

533  
534 Pan-neuronal knockdown of *Ank2* also severely impaired 24-hour LTM without  
535 affecting courtship behavior, learning, or immediate courtship memory. It was  
536 previously found that decreased expression of *Ank2* in the mushroom body with  
537 *OK107-GAL4* did not result in learning deficits but caused significant impairment to  
538 one-hour STM in the same assay (IQBAL *et al.* 2013). Similarly, pan-neuronal  
539 knockdown of *Ank2* with *elav-GAL4* impaired one hour olfactory memory (HIGHAM *et al.*  
540 *et al.* 2019) and mushroom body-specific knockdown with *nSyb-GAL4* impaired three-  
541 hour memory when tested in the olfactory conditioning assay (WALKINSHAW *et al.*  
542 2015). However, in these studies the drivers are expressed in larvae at which time the  
543 mushroom body is developing (NICOLAI *et al.* 2003; OGIENKO *et al.* 2020; KOBLER *et al.*  
544 2021), and an intact mushroom body is required for normal olfactory STM (HEISENBERG  
545 *et al.* 1985) and associative STM lasting longer than 30 mins (MCBRIDE *et al.* 1999).  
546 Furthermore, output from  $\gamma$  neurons is required for short-term courtship memory  
547 (KELEMAN *et al.* 2012) and knockdown of *Ank2* in  $\gamma$  neurons during mushroom body  
548 development was recently found to result in a shortened axon initial segment in third  
549 instar larval brains (SPURRIER *et al.* 2019). In light of the severe mushroom body defects  
550 we observed on knockdown of *Ank2*, and that *Ank2* has been implicated in synapse  
551 stability (HORTSCH *et al.* 2002; KOCH *et al.* 2008), it is unsurprising that memory would  
552 be impaired, and from these data it cannot be determined whether *Ank2* plays a  
553 specific role in memory or whether the defects are a result of impaired morphogenesis  
554 of the mushroom body. To that end, in order to dissociate developmental effects from  
555 the molecular processes required for LTM in an adult brain, knockdown of *Ank2* was  
556 restricted to mature neurons with *GAL80*. A specific deficit in 24 hour LTM was  
557 observed when *Ank2* was knocked down in the adult mushroom body, and subsequent

558 testing of GAL4 drivers that restrict expression to specific mushroom body subtypes  
559 revealed that knockdown in just the  $\gamma$  neurons was sufficient to impair memory. We  
560 previously showed that overexpression of *HDAC4* in  $\gamma$  neurons also impaired the  
561 formation of long-term courtship memory (SCHWARTZ *et al.* 2016) and strikingly, in  
562 accordance with the similarity in mushroom body defects, knockdown of *Moe* resulted  
563 in the same phenotype as *Ank2* knockdown and *HDAC4* overexpression (FITZSIMONS *et*  
564 *al.* 2013; FREYMUTH AND FITZSIMONS 2017), suggesting that a functional relationship  
565 between these proteins also occurs during LTM formation.

566  
567 The identification that *Ank2* is required in the  $\gamma$  neurons of the mushroom body for  
568 normal cVA-retrievable memory is consistent with current models of the circuitry that  
569 facilitates this memory (KELEMAN *et al.* 2012; RAUN *et al.* 2021). cVA-retrievable  
570 memory involves activation of aSP13 dopaminergic neurons which innervate the  $\gamma$ 5  
571 compartment at tip of the  $\gamma$  lobe. This results in increased synaptic transmission from  
572  $\gamma$  neurons to glutamatergic M6 output neurons, which themselves feedback to  
573 innervate aSP13 neurons to form a recurrent activation loop (KELEMAN *et al.* 2012; ZHAO  
574 *et al.* 2018). Long-term memory requires a later reactivation of aSP13 neurons, which  
575 is dependent on sleep (DAG *et al.* 2019). Expression of epigenetic regulators required  
576 for cVA-retrievable memory, including *Rpd3*, *HDAC4* and *G9a* has also been pinpointed  
577 to a requirement in only  $\gamma$  neurons. Similarly, the cytoplasmic polyadenylation  
578 element-binding protein *Orb2* is required for translation of synaptic mRNA during LTM  
579 consolidation in the  $\gamma$  lobe (KRUTTNER *et al.* 2015).

580  
581 Although *Ank2* and *HDAC4* both localize to axons of the mushroom body, and  
582 overexpression of *HDAC4* in the mushroom body also results in impaired axon  
583 elongation and guidance defects as well as deficits in LTM formation (FITZSIMONS *et al.*  
584 2013; FREYMUTH AND FITZSIMONS 2017; MAIN 2021), we found no evidence of a physical  
585 interaction in *Drosophila* neurons. We further investigated the interaction between  
586 *HDAC4* and *Ank2* in the eye and observed that a nuclear-restricted mutant of *HDAC4*  
587 interacted genetically with *Ank2*, however this was not dependent on the presence of  
588 a conserved putative ankyrin-binding motif in *HDAC4*. Given the striking similarity of  
589 the *Ank2* and *Moe* knockdown phenotypes with *HDAC4* overexpression, it is possible  
590 that when increased in abundance in Kenyon cell nuclei, *HDAC4* may act indirectly to  
591 disrupt the normal role of the *Ank2*, *Nrg*, and *Moe* complex. It is not yet known how  
592 an *HDAC4*-mediated signal from the nucleus modulates these or other proteins  
593 involved in neuronal morphogenesis and LTM. We previously found that *HDAC4* also  
594 interacts genetically with several components of the SUMOylation machinery,  
595 including the SUMO E2-conjugating enzyme *Ubc9*, which is also required for long-term  
596 courtship memory and interacts genetically with *HDAC4* during this process (SCHWARTZ  
597 *et al.* 2016). *HDAC4* has been proposed to act as an E3 ligase to enhance SUMOylation  
598 of target proteins (GREGOIRE AND YANG 2005; ZHAO *et al.* 2005). Given that SUMOylation  
599 regulates neuronal protein activity (HENLEY *et al.* 2014) and regulates memory  
600 formation (YANG *et al.* 2012; CHEN *et al.* 2014; LEE *et al.* 2014; YU *et al.* 2020), it would  
601 be worthwhile to further investigate whether any of the *HDAC4*-interacting genes are  
602 SUMOylated and whether this is impacted by altering the expression level or  
603 subcellular distribution of *HDAC4*.

604

605 In our screen for genes that enhanced the *HDAC4*-induced rough eye phenotype  
606 (SCHWARTZ *et al.* 2016), the largest group of genes we identified composed of  
607 regulators of the actin/spectrin cytoskeleton such as *Trio*, *NetrinB*, *Derailed*, *Ankyrin*,  
608 *Ankyrin2*, and *Moesin*, suggesting that *HDAC4* might regulate memory through  
609 interaction with genes involved in remodelling of the actin/spectrin cytoskeleton. This  
610 is a phenomenon which is believed to regulate the structural changes including  
611 polymerization/depolymerization or alterations of the underlying actin cytoskeleton  
612 that underpin learning and memory (ENGERT AND BONHOEFFER 1999; KRUCKER *et al.* 2000;  
613 LAMPRECHT AND LEDOUX 2004; LEISS *et al.* 2009; OJELADE *et al.* 2013; LAMPRECHT 2014). We  
614 previously showed that expression of a constitutively active mutant of *Moe* increases  
615 the density of F-actin in spine-like structures in LPTC neurons (FREYMUTH AND FITZSIMONS  
616 2017). In mammals, AnkG is present at synapses and is required for formation of  
617 nanodomain structures in the perisynaptic spine and spine neck of dendrites, the  
618 contents of which include adhesion molecules, F-actin and CaMKII, thus facilitating  
619 interaction with the actin cytoskeleton for F-actin rearrangements that underlie  
620 dynamic alterations in spine morphology. The geometry of the spine in this region  
621 increases on induction of long-term potentiation in primary cortical neurons, and  
622 reduction of AnkG prevents spine enlargement (SMITH *et al.* 2014). It is also notable  
623 that inhibition of actin polymerization within the mushroom body of the honeybee  
624 enhanced associative olfactory memory (GANESHINA *et al.* 2012). As reduced *HDAC4*  
625 also impairs LTM, indicating an essential role in memory formation, the impact of both  
626 increased and reduced *HDAC4* on cytoskeletal rearrangement warrants further  
627 investigation.

628

### 629 **Acknowledgements**

630 We thank Hermann Aberle (Heinrich-Heine-Universität, Düsseldorf, Germany) for the  
631 Ank2-L antibody and Max Scott for constructive comments on the manuscript. We also  
632 thank the Manawatu Microscopy and Imaging Centre, Massey University for  
633 assistance with confocal and SEM.

634

### 635 **Funding**

636 This work was supported by the Royal Society of New Zealand (Marsden grant  
637 MAU1702 to HLF) and the Massey University Research Fund.

638

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911

912

913 **Table 1. Frequency of mushroom body phenotypes resulting from knock down of**

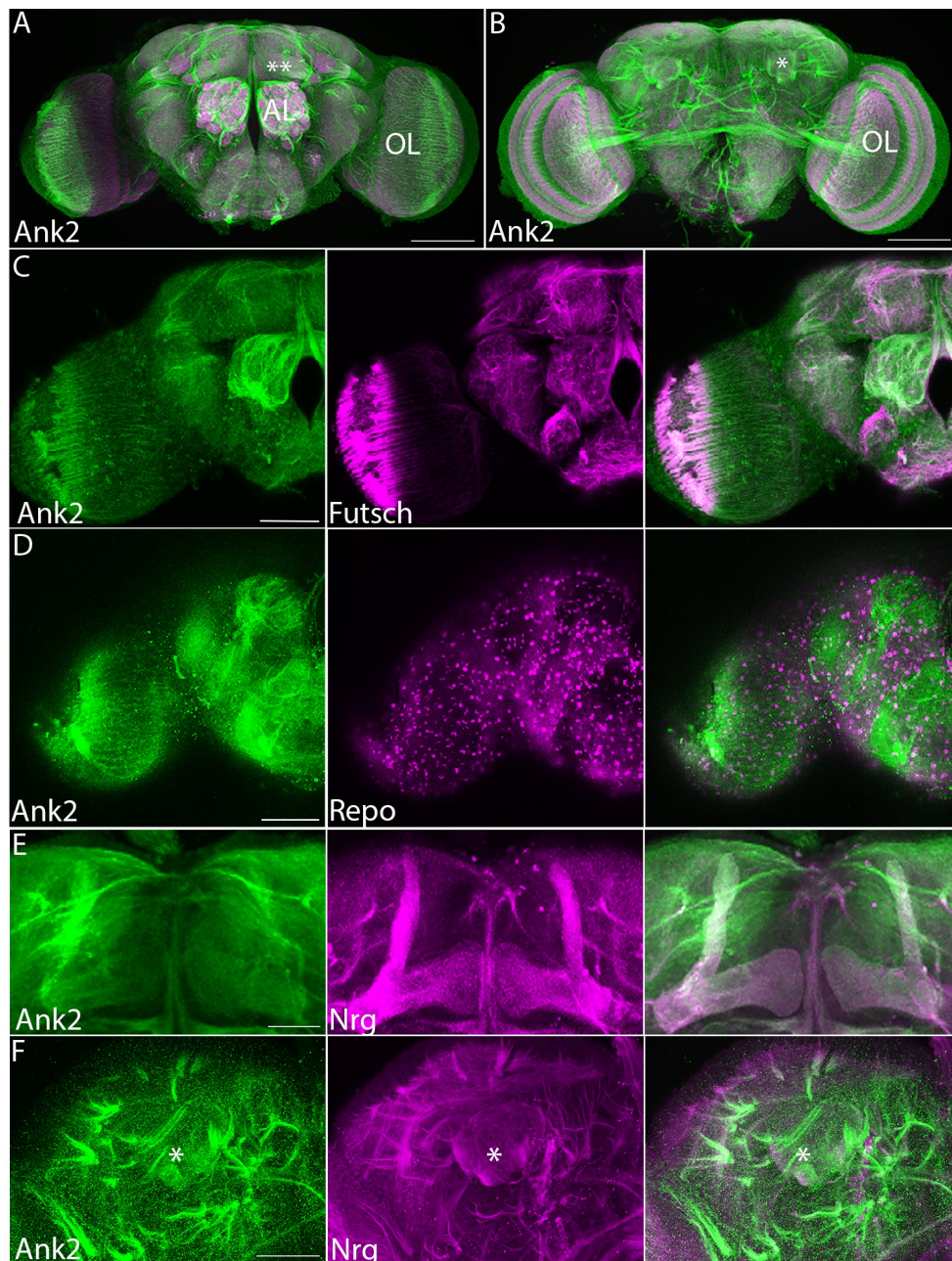
914 **Ank2**. The percentage of brains displaying each phenotype was calculated from the  
 915 total number of brains analyzed for each genotype (n) at 22, 25 and 30°C. Statistical  
 916 analysis was performed with Fisher's Exact Test. Knockdown of *Ank2* resulted in  
 917 significantly more brains with thin lobes (p=0.0032), elongation and guidance deficits  
 918 (p=0.0032), or absent  $\alpha$  and/or  $\beta$  lobes (p=0.0014), than controls at 25°C. The more  
 919 severe phenotypes of completely absent  $\alpha$  and/or  $\beta$  lobes were more pronounced at  
 920 higher temperatures (27°C compared to 22°C, p=0.0015).

921

Genotype	<i>elav&gt;w(CS10)</i>	<i>elav&gt; Ank2 RNAi</i>	<i>elav&gt; Ank2 RNAi</i>	<i>elav&gt; Ank2 RNAi</i>
Temperature	25°C	22°C	25°C	27°C
<i>n</i>	20	30	30	16
Thin $\alpha/\beta$ or $\gamma$ lobes	0%	37%	23%	62%
Fused $\beta$ lobes	0%	13%	10%	0%
Outgrowth or guidance defect	0%	37%	23%	50%
Absent $\alpha/\beta$ or $\gamma$ lobes	0%	6%	40%	50%

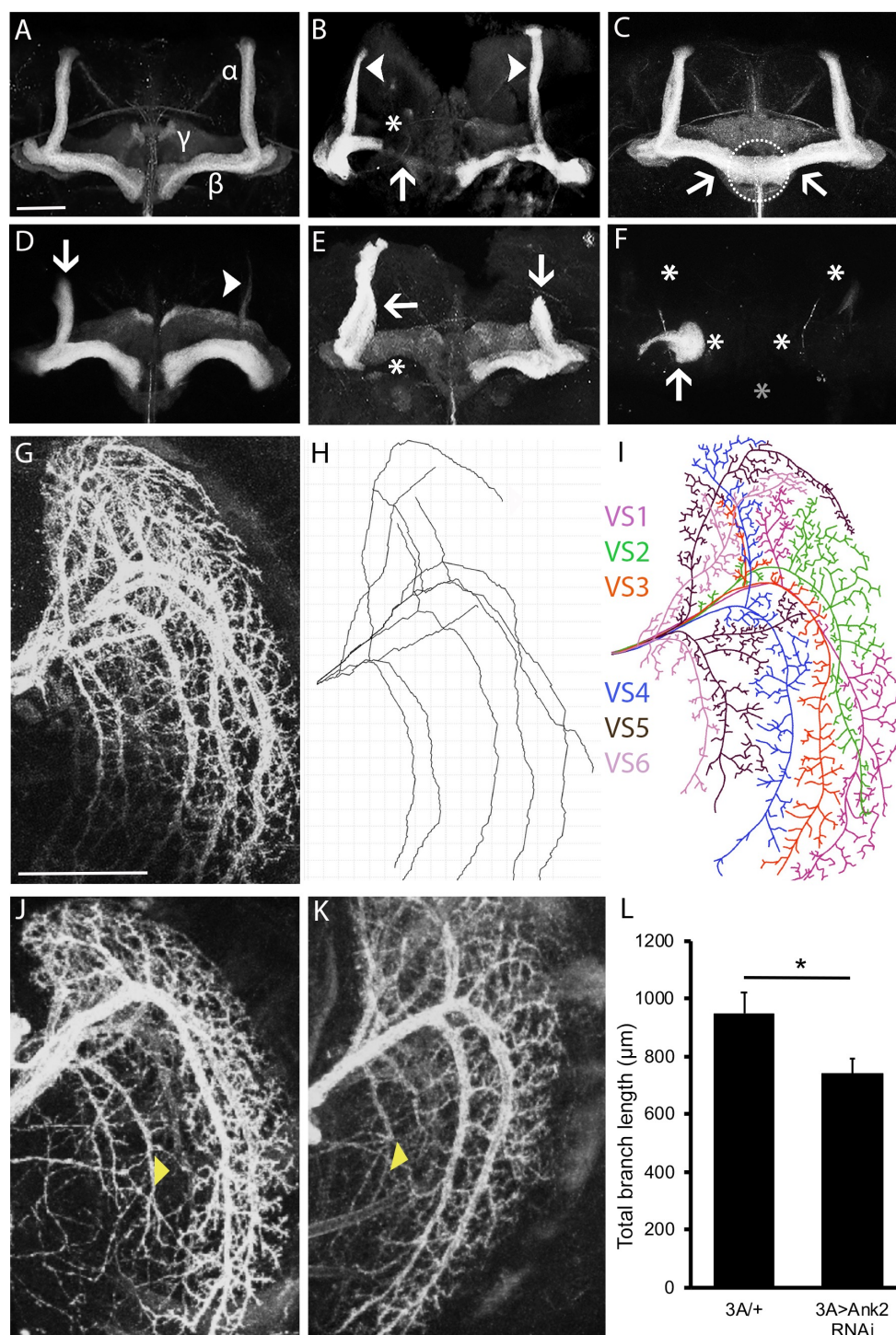
922

## Figure 1



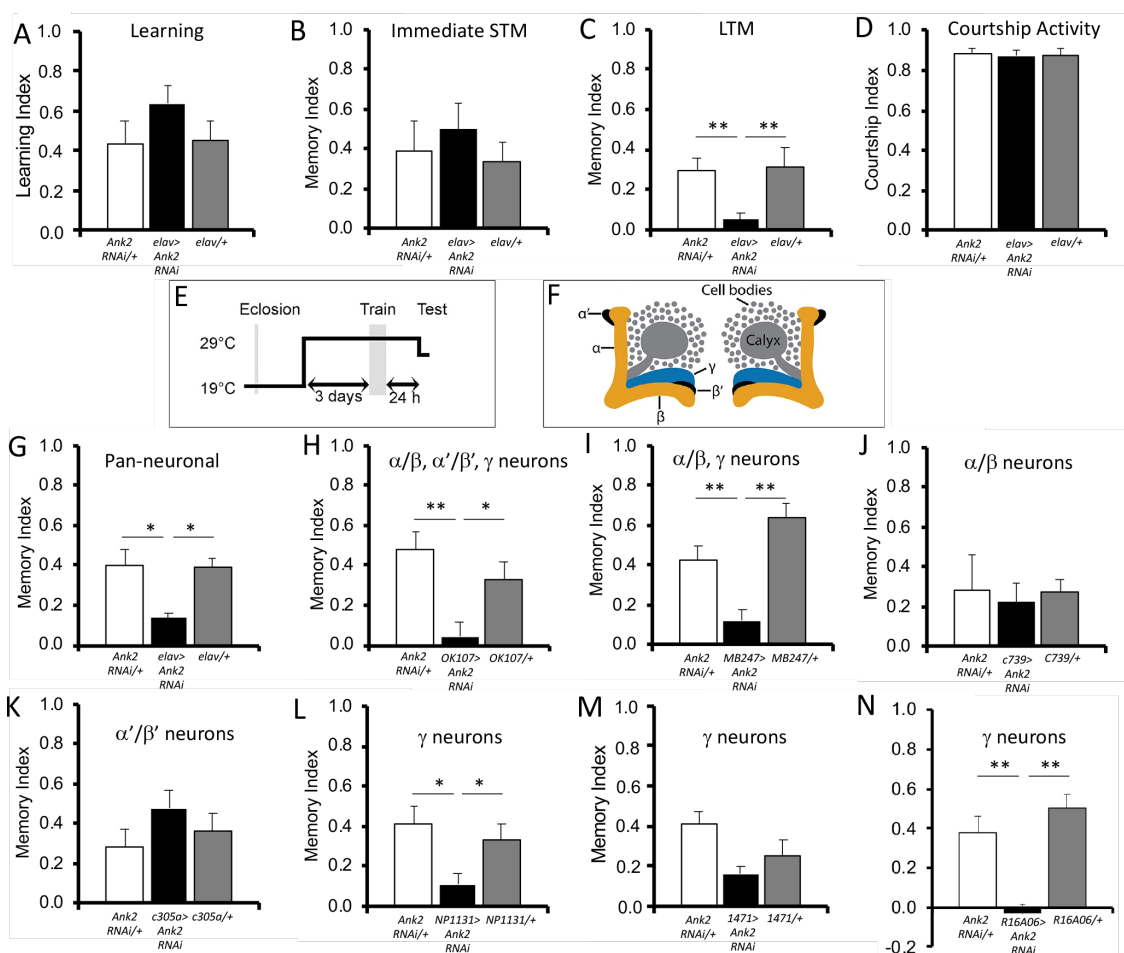
**Figure 1. Expression of Ank2 in the adult brain.** Immunohistochemistry on whole mount wild-type brains indicates widespread distribution of Ank2-L. A,B Confocal projection of brains labelled with Ank2-L (green) and nc82 (Bruchpilot, magenta) antibodies, to highlight the synaptic neuropil. Images are Z-stacks of 1  $\mu\text{m}$  optical sections. A, Anterior confocal projection. AL, antennal lobe, OL optic lobe, \*\*indicates the  $\gamma$  lobe of the mushroom body. Scale bar = 100  $\mu\text{m}$  B-F. Co-labelled proteins are shown in magenta and labelled in the middle panel. B. Posterior confocal projection, \*indicates the calyx of the mushroom body. Scale bar = 100  $\mu\text{m}$ . C. Immunohistochemistry with Ank2-L and 22C10 (Futsch) antibodies showing codistribution in neurons, with widespread localization to axon tracts. Scale bar = 50  $\mu\text{m}$ . D. Ank2-L does not codistribute with pan-glial marker Repo. Scale bar = 50  $\mu\text{m}$ . E. Ank2-L colocalizes with Nrg in the mushroom body lobes. Scale bar = 50  $\mu\text{m}$ . F. Ank2-L also codistributes with Nrg in axon tracts surrounding the calyx (asterisk). Scale bar = 50  $\mu\text{m}$ .

## Figure 2



**Figure 2. Reduced Ank2 expression disrupts neuronal development.** A-F. Immunohistochemistry with anti-FasII on whole mount brains reveals morphological defects of the mushroom body resulting from pan-neuronal expression of *Ank2* RNAi driven by *elav-GAL4*. Knockdown was confirmed by RT-qPCR (as described in the methods). All images are frontal confocal projections through the mushroom body. Scale bar = 50  $\mu$ m. A. Wild-type mushroom body stained with anti-Fas II to highlight the  $\alpha$ ,  $\beta$  and  $\gamma$  lobes. B. Thin  $\alpha$  lobes (arrowheads), a prematurely terminated  $\beta$  lobe (arrow) and a missing  $\gamma$  lobe (asterisk). C.  $\beta$  lobes (arrows) have crossed the midline and appear fused (circle). D. A thin (arrowhead) and a prematurely terminated  $\alpha$  lobe (arrow). E. Misoriented  $\beta$  lobe (arrow) which projects anteriorly rather than its usual medial orientation (asterisk). A prematurely terminated  $\alpha$  lobe is also present (arrow). F. None of the lobes have elongated (asterisks) and axon stalling is observed whereby the axons grow in a ball-like structure (arrow). Grey asterisk indicates the midline. G-J. Immunohistochemistry on whole mount brains with anti-GFP driven by *3A-GAL4* to detect Lifect in LPTCs whole mount brains. All images are confocal projections through the optic lobe of the brain. G. The dendritic arbor of the six neurons comprising the vertical system of LPTCs in a wild-type brain is visualised with anti-GFP. Scale bar = 50  $\mu$ m. H. Dendritic trace generated using SNT Tracer (Image J). I. Cartoon trace of the confocal micrograph showing the dendritic branching of each of the vertical system neurons. J,K. Knockdown of *Ank2* results in defects in dendritic branching (arrow heads). L. Total dendrite branch length was reduced by knockdown of *Ank2* (student's *t*-test  $t_{(36)}=2.27$ ,  $p<0.05$ ).

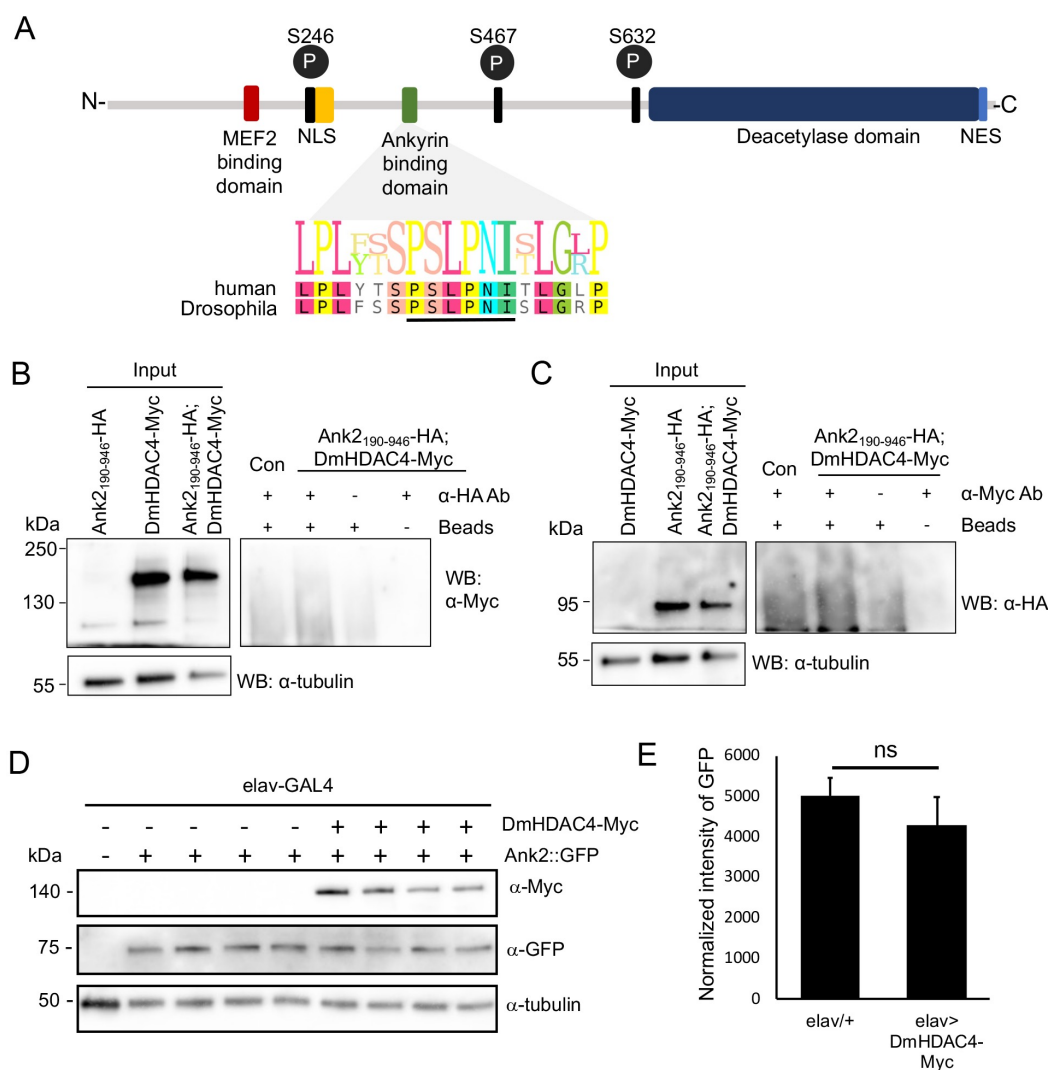
Figure 3





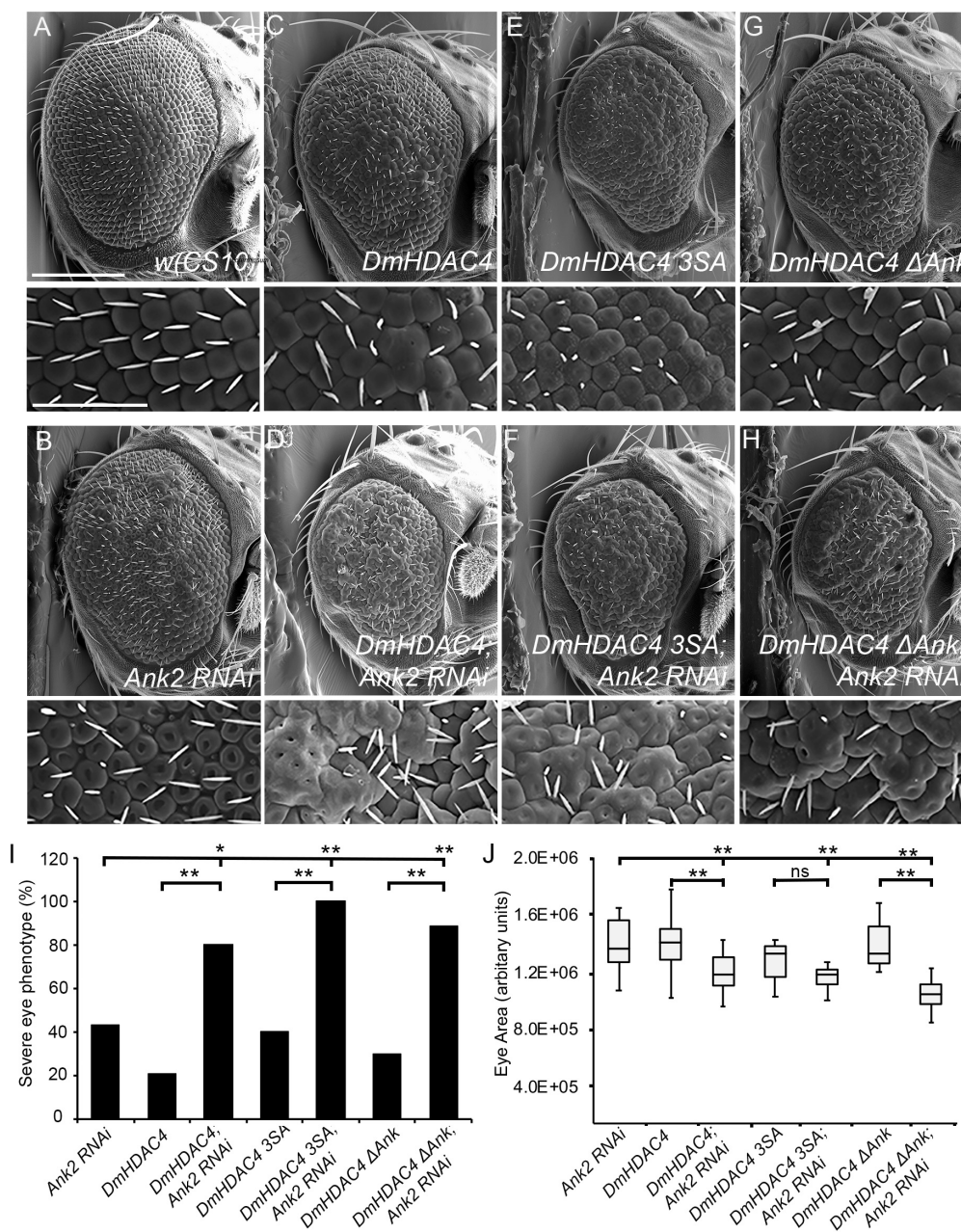
**Figure 3. Ank2 is required for long-term memory.** Learning and memory were assessed with the courtship suppression assay. The controls included in each assay are the GAL4 driver (plus tubP-GAL80ts where indicated) crossed to CS, and UAS-*Ank2* crossed to CS, such that the progeny are heterozygous for either the driver or the RNAi. A-D. *elav-GAL4* and UAS-*Ank2* RNAi flies were crossed to achieve pan-neuronal knockdown of *Ank2* in progeny. A. Learning was unaffected knockdown of *Ank2* (ANOVA,  $F_{(2,47)}=0.002$ ,  $p=0.252$ ). B. Immediate memory was also unaffected (ANOVA,  $F_{(2,45)}=0.044$ ,  $p=0.819$ ). C. *Ank2* knockdown impaired long-term memory (ANOVA,  $F_{(2,60)}=7.31$ ,  $p<0.001$ ; *post-hoc* Tukey's HSD,  $**p<0.01$ ). D. Courtship activity was not impaired by pan-neuronal knockdown of *Ank2* (ANOVA,  $F_{(2,51)}=0.14$ ,  $p=0.870$ ). E-K. Knockdown of *Ank2* in the adult mushroom body impairs LTM. *Ank2* was knocked down in specific regions of the brain by crossing *Ank2* RNAi to the indicated driver line and *tubP-Gal80ts*. E. Schematic diagram depicting the induction of expression in the adult mushroom body. Expression was restricted to the adult brain by raising flies at raised at 19°C, at which temperature GAL80 represses GAL4. After eclosion, when flies were 3-5 days old, the temperature was raised to 30°C for 72 hours, after which training commenced. At this temperature GAL80 is inactivated, allowing GAL4 to induce transgene expression. Twenty-four hours after training, the flies were equilibrated to 25°C for one hour prior to testing. F. Schematic diagram labelling the lobes of the mushroom body in which *Ank2* was knocked down. G. Pan-neuronal knockdown of *Ank2* in the adult brain impairs long term memory (ANOVA,  $F_{(2,54)}=0.317$ ,  $p<0.01$ ; *post-hoc* Tukey's HSD,  $*p<0.05$ ). H. Similarly, memory is also impaired when knockdown of *Ank2* is restricted to the mushroom body (ANOVA,  $F_{(2,52)}=0.922$ ,  $p<0.001$ ; *post-hoc* Tukey's HSD,  $**p<0.01$ ,  $*p<0.05$ ). I. When knockdown is restricted to the  $\alpha/\beta$  and  $\gamma$  neurons of the mushroom body, long-term memory is still disrupted (ANOVA,  $F_{(2,51)}=0.923$ ,  $p<0.0001$ ; *post-hoc* Tukey's HSD,  $**p<0.01$ ). J. Reduction of *Ank2* in just the  $\alpha/\beta$  neurons has no significant effect on long-term memory (ANOVA,  $F_{(2,41)}=0.025$ ,  $p=0.819$ ). K. There is also no impairment when *Ank2* is reduced in the  $\alpha'/\beta'$  neurons (ANOVA,  $F_{(2,51)}=0.122$   $p=0.372$ ). L. *Ank2* is required in the  $\gamma$  lobes, as knockdown with *NP1131-GAL4* impairs LTM (ANOVA,  $F_{(2,46)}=0.312$ ,  $p<0.01$ ; *post-hoc* Tukey's HSD,  $*p<0.05$ ). M. The weaker  $\gamma$  lobe driver *1471-GAL4* reduced LTM, however this was not quite significant (ANOVA,  $F_{(2,59)}=0.210$ ,  $p=0.056$ ). Knockdown of *Ank2* with the stronger  $\gamma$  lobe driver *R16A06-GAL4* did impair LTM significantly (ANOVA,  $F_{(2,33)}=18.57$ ,  $p<0.0001$ , *post-hoc* Tukey's HSD,  $**p<0.01$ ).

## Figure 4



**Figure 4. Ank2 does not bind HDAC4 and nor is its expression regulated by HDAC4.** A. Domain structure of human HDAC4 showing binding sites conserved between *Drosophila* and human HDAC4. The amino acid sequence of the region containing the RFXANK/ANKRA2 binding site in human HDAC4 is shown, with the corresponding amino acid sequence in *Drosophila* HDAC4. NLS, nuclear localisation sequence, NES, nuclear export sequence. Ps circled in black are serine residues that when phosphorylated provide binding sites for 14-3-3 mediated nuclear export. B,C. Co-immunoprecipitation of Ank2<sub>190-946</sub>-HA and DmHDAC4-Myc from whole cell lysates of fly heads with either anti-Myc or anti-HA. Both blots were probed with anti-tubulin as a loading control. Input samples = 30 μg. B. Following immunoprecipitation with anti-HA, DmHDAC4-Myc was not detected upon probing with anti-Myc. Inputs include Ank2<sub>190-946</sub>-HA, which is not detected by Myc, DmHDAC4-Myc, and Ank2<sub>190-946</sub>-HA; DmHDAC4-Myc to confirm expression. C. In the reciprocal experiment, flies expressing Ank2<sub>190-946</sub>-HA; DmHDAC4-Myc were subjected to IP with anti-Myc, however Ank2<sub>190-946</sub>-HA was not detected upon probing with anti-HA. D. *elav-GAL4; Ank2::GFP* flies were crossed to *w(CS)10* and *UAS-DmHDAC4-Myc* and whole head lysates of progeny were generated for western blotting. Samples were processed from four independent crosses. Blots were probed with anti-Myc to verify expression of *DmHDAC4-Myc* and anti-GFP to determine whether the amount of Ank2::GFP normalized to tubulin is altered in the presence of *DmHDAC4-Myc*. E. There was no significant change in the level of Ank2::GFP on expression of *DmHDAC4-Myc*.

Figure 5



**Figure 5. *Ank2* interacts genetically with *HDAC4* in the eye.** A-H. Scanning electron micrographs of *Drosophila* eyes expressing *Ank2* RNAi and/or *HDAC4* variants. The genotypes indicated in each panel were generated by crossing *GMR-GAL4* females to males carrying each *UAS-HDAC4* construct or *Ank2* RNAi to the *w(CS10)* control. Top panel: Scale bar = 200  $\mu$ m. Bottom panel: Scale bar = 100  $\mu$ m. I. The percentage of eyes displaying severe phenotypes are shown for each genotype. Severe phenotypes were scored as severe disorganisation with fused ommatidia in more than ten areas, and/or more than 50 collapsed ommatidia with crevices or cavities. \* $p < 0.05$ , \*\* $p < 0.01$  following one-tailed Fisher's exact test. p-values: *Ank2* RNAi:*DmHDAC4*; *Ank2* RNAi = 0.0247, *Ank2* RNAi:*DmHDAC4 3SA*; *Ank2* RNAi = 0.0006, *Ank2* RNAi:*DmHDAC4  $\Delta$ Ank*; *Ank2* RNAi = 0.0028, *DmHDAC4*:*DmHDAC4*; *Ank2* RNAi = 0.0004, *DmHDAC4 3SA*:*DmHDAC4 3SA*; *Ank2* RNAi = 0.0005, *DmHDAC4  $\Delta$ Ank*:*DmHDAC4  $\Delta$ Ank*; *Ank2* RNAi = 0.0002. J. Eye sizes were quantified by tracing a line around each eye and calculating the area in arbitrary units using ImageJ software and plotting on Box and whisker plots to show the variation in eye size for each genotype. \*\* $p < 0.01$  following one-way ANOVA and post-hoc Tukey's HSD test for significance. p-values: *w(cs10)*:*Ank2* RNAi = 0.007, *w(cs10)*:*DmHDAC4* = 0.009, *w(cs10)*:*DmHDAC4*; *Ank2* RNAi = 0.001, *Ank2* RNAi:*DmHDAC4*; *Ank2* RNAi = 0.001, *DmHDAC4*:*DmHDAC4*; *Ank2* = 0.001, *w(cs10)*:*DmHDAC4 3A* = 0.001, *w(cs10)*:*DmHDAC4 3A*; *Ank2* RNAi = 0.001, *DmHDAC4 3A*:*DmHDAC4 3A*; *Ank2* RNAi = 0.285, *w(cs10)*:*DmHDAC4  $\Delta$ Ank* = 0.001, *w(cs10)*:*DmHDAC4  $\Delta$ Ank*; *Ank2* RNAi = 0.001, *DmHDAC4  $\Delta$ Ank*:*DmHDAC4  $\Delta$ Ank*; *Ank2* RNAi = 0.001. n = number of eyes per sample. The number of eyes analyzed per genotype is indicated in Table S1.