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Glial Hedgehog signalling and lipid metabolism regulate neural stem cell proliferation in *Drosophila*

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1 **Title: Glial Hedgehog signalling and lipid metabolism regulate neural stem cell**
2 **proliferation in *Drosophila***

3 **Running title: Glial Hh and lipids affect NB divisions**

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18
19 **Abstract:**

20 The final size and function of the adult central nervous system (CNS) is determined
21 by neuronal lineages generated by neural stem cells (NSCs) in the developing brain.

22 In *Drosophila*, NSCs called neuroblasts (NBs) reside within a specialised
23 microenvironment called the glial niche. Here, we explore non-autonomous glial
24 regulation of NB proliferation. We show that lipid droplets (LDs) which reside within
25 the glial niche are closely associated with the signalling molecule Hedgehog (Hh).

26 Under physiological conditions, cortex glial Hh is autonomously required to sustain
27 niche chamber formation. Upon FGF-mediated cortex glial overgrowth, glial Hh non-
28 autonomously activates Hh signalling in the NBs; which in turn disrupts NB cell cycle
29 progression and its ability to produce neurons. Glial Hh's ability to signal to NB is
30 further modulated by lipid storage regulator Lipid storage droplet-2 (Lsd-2) and de
31 novo lipogenesis gene Fatty acid synthase 1 (Fasn1). Together, our data suggests that

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32 glial-derived Hh modified by lipid metabolism mechanisms can affect the
33 neighbouring NB's ability to proliferate and produce neurons.

34

35 **Keywords:** *Drosophila*, glial niche, neuroblast, Hedgehog, lipid metabolism

36

37 **Introduction:**

38 Most stem cells reside within specialized groups of cells, collectively referred to as a
39 niche, that provide the trophic, structural and nutritional microenvironment to sustain
40 and protect the stem cells during development (Scadden, 2014). The niche relays
41 developmental and physiological states of the animal to the stem cells and influences
42 the stem cells' ability to divide in accordance with the environmental state of the
43 organism. Asymmetrically dividing and multipotent neural stem cells in both
44 mammals and invertebrates are responsible for generating the adult nervous system
45 (Homem & Knoblich, 2012).

46

47 In *Drosophila*, the vast majority of NBs are specified during embryogenesis,
48 proliferate throughout larval development, and terminate divisions during pupal
49 stages. Type I NBs located within the ventral nerve cord (VNC) and the central brain
50 (CB), are the predominant type of NBs, while type II NBs are eight NB lineages
51 located on the dorsal surface of the CB (Homem & Knoblich, 2012). During each
52 type I NB cell division, NB self-renews, and produces a smaller ganglion mother cell
53 (GMC) that creates a limited number of neurons or glia (Fig 1A). The ability of NBs
54 to divide and generate appropriate progeny number and cell diversity is determined by
55 their ability to maintain asymmetric division, regulate the speed of their cell cycles,
56 and timely enter/exit the cell cycle at the beginning and end of neurogenesis (Homem
57 & Knoblich, 2012). Cell intrinsic mechanisms such as the temporal regulation of NB
58 identity via transcription factors that are expressed throughout the life time of the
59 NBs, impact on both the numbers and the types of neurons generated by the NB (Doe,
60 2017). However, more recently, attention has shifted towards understanding how cell
61 extrinsic signals are interpreted by the NBs to alter their behavior (Ramon-Canellas *et*
62 *al*, 2019).

63

64 Larval NBs and their progeny are surrounded by a scaffold of glial cell processes,
65 which form the stem cell niche in the CNS (Fig 1A). Glial cells fall into three classes:

66 (i) surface (perineural and subperineural) glia that enwrap the CNS to form the blood
67 brain barrier (BBB); (ii) cortex glia that encapsulate neuronal soma and NBs; and (iii)
68 neuropil glia that are located at the cortex–neuropil interface and form a sheath
69 around the neuropil compartments (Freeman, 2015).

70

71 The intimate relationship between glial cells and NBs has been extensively studied in
72 the context of NB entry into the cell cycle at the beginning of postembryonic
73 neurogenesis shortly after larval hatch (Ding *et al*, 2020). Feeding has been shown to
74 trigger insulin production by surface glial cells, which in turn activates the
75 insulin/insulin-like growth factor pathway in neighboring NBs and stimulates their
76 growth and proliferation via activation of the Phosphoinositide 3-kinase (PI3K)
77 signaling pathway (Chell & Brand, 2010; Sousa-Nunes *et al*, 2011). Once NBs enter
78 into the cell cycle, glial cells continue to play active roles in promoting NB
79 proliferation. These reactivated NBs are found in close association with cortex glia
80 (Hoyle, 1986; Hoyle *et al*, 1986; Peraanu *et al*, 2005), and this contact is maintained
81 through adhesion via E-cadherin. Disruption of NB-cortex glia contact affects the
82 NB's ability to undergo mitosis (Doyle *et al*, 2017; Dumstrei *et al*, 2003), and the
83 failure to expand the glial membrane also affects both neuronal survival as well as NB
84 cell cycle progression (Speder & Brand, 2018; Yuan *et al*, 2020). Diffusible
85 molecules that pass from glial cells to influence NB behavior include Dally-like (Dlp)
86 in the perineural glia, (Kanai *et al*, 2018) and Jellybelly (Jeb) in the cortex glia
87 (Cheng *et al*, 2011). Furthermore, organelles such as lipid droplets (LDs) in the glial
88 niche have been shown to buffer NBs proliferation from peroxidation chain reactions
89 induced by oxidative stress (Bailey *et al*, 2015), suggesting that glial niche and the
90 signaling molecules produced by these cells are important mediators of non-
91 autonomous regulation of NB behavior during development and environmental stress.

92

93 In this study, we investigate how the stem cell niche and its dysfunction influence
94 stem cell behavior and the consequences on the brain as a whole. We found that the
95 signaling molecule Hedgehog (Hh), involved in numerous developmental processes,
96 resides within the cortex glial membrane that surrounds NBs. Hh is autonomously
97 required to promote glial niche growth as well as acts non-cell autonomously to
98 activate the Hh signaling in the NB, triggering its delay in S-phase progression.
99 Maintaining cortex glial size is important, as overgrowth induced by FGF activation,

100 a mutation implicated in glioblastoma (Dienstmann *et al*, 2014; Jimenez-Pascual &
101 Siebzehnrbuhl, 2019; Morrison *et al*, 1994; Yamada *et al*, 1999), phenocopied the
102 effects of glial Hh activation on NBs. Indeed, inhibiting Hh, rescued NB proliferation
103 defects. Furthermore, we demonstrated that downstream of glial FGF signaling, Hh
104 activity and its ability to signal to NBs is modulated by two lipid storage regulators
105 Lsd-2 and Fasn1. Together, our data show that a dysfunctional niche can non-
106 autonomously affect NB's ability to produce the correct number of neurons that make
107 up the adult CNS. This process mechanistically involves the Hh signaling pathway
108 and its modulation by lipid metabolism.

109

110 **Results:**

111 **Hh is localised to the LDs within cortex glial cells**

112 To identify potential morphogens that facilitate glia-NB communication in the CB,
113 we assayed for secreted molecules which are known to act in a paracrine fashion. We
114 found that Hh is expressed at high levels at 96 hours After Larval Hatching (96ALH)
115 in glial cells labelled using *Repo-GAL4 > GFP* (Fig 1B-D'). Hh is a morphogen that
116 was first identified to regulate embryo segmentation and wing imaginal disc
117 development (Heemskerk & DiNardo, 1994; Nusslein-Volhard & Wieschaus, 1980).
118 In the wing disc, it is expressed in the posterior compartment and is distributed in a
119 gradient to regulate target gene expression in the anterior compartment (Fig 1E-F)
120 (Chen *et al*, 2017). In the glial niche, however, we found Hh staining was mostly
121 distributed in ring-like structures in the glial cytoplasm (yellow arrows, Fig 1C-D').
122 We then assessed if Hh is associated with specific organelles. LDs are round-shaped
123 organelles, consisting of a hydrophobic core for the storage of neutral lipids and a
124 phospholipid monolayer containing LD surface proteins. As LDs have previously
125 been reported to be enriched in the glial niche (Bailey *et al*, 2015; Kis *et al*, 2015),
126 we therefore tested if Hh ligands are associated with LDs. Using a neutral lipid stain,
127 lipidTOX to visualise LDs and either an antibody, or a BAC encoded Hh:GFP to
128 detect Hh (Chen *et al*, 2017), we found that Hh is localised to the surface of the LDs
129 in the glial niche (yellow arrows, Fig 1C-D'', I and Fig EV1A-B''), but not in the
130 wing disc (white arrows, Fig 1F-G'', J and Fig EV1C-D'') nor at earlier stages of
131 development in the CB (48 ALH, Fig EV2 A-B''). Together, our data suggests that
132 Hh is localised to the surface of LDs in the glial niche.

133

134 We next explored whether Hh-LD associations are specifically localised to a glial
135 subtype. Hh-LD associations were largely absent from both surface glial cells that
136 forms the blood brain barrier (BBB) of the CNS (white arrows, Fig EV2E-F'''), as
137 well as optic lobe glial cells (white arrows, Fig EV2C-D'''). In fact, Hh-LD
138 associations were enriched in the cortex glial cells, underneath the sheath-like surface
139 glial clone generated via *repo*-MARCM (yellow arrows, Fig EV2G). Using a cortex
140 specific driver, *NP2222-GAL4* (Awasaki *et al*, 2008; Hayashi *et al*, 2002), we
141 confirmed that the Hh-LD associations were localised to the cortex glia (Fig 1K-L'').

142

143 **Hh autonomously regulates cortex gliogenesis and non-autonomously regulates** 144 **NB proliferation**

145 In the mouse brain, Hh has been shown to promote astrocyte proliferation (Chandra *et*
146 *al*, 2015; Takezaki *et al*, 2011; Ugbo *et al*, 2017). In glioblastoma (GBM), the
147 Hh/Gli1 signaling pathway acts to accelerate cell proliferation (Chandra *et al.*, 2015).
148 To investigate the role of Hh in *Drosophila* cortex glial cells, where Hh is most
149 abundant, we used pan glial driver *repo-GAL4* to express *hh* RNAi together with
150 *UAS-Dcr2* (*Dicer-2*) which sufficiently depleted Hh expression and reduced overall
151 CNS size (Fig EV3A-D). The reduction in CNS volume was accounted for by a
152 significant decrease in Repo⁺ glial cell number and membrane size (labelled using
153 *NP2222-GAL4>GFP*, Fig 2A-D). Using a pan-glial driver *repo-GAL4*, we found
154 cortex glial chambers were significantly disrupted upon Hh knockdown (Speder &
155 Brand, 2018; Yuan *et al.*, 2020), leading to clustering of NBs (compare Fig 2G to F).
156 However, overexpression of Hh did not affect glial cell number nor membrane size
157 (Fig EV3L-O). This suggests Hh is necessary but not sufficient for glial expansion
158 during CNS development.

159

160 Given the role of Hh as a secreted ligand that can act over short range within the NB
161 lineage (Chai *et al*, 2013) and that it is highly expressed in the cortex glial niche
162 surrounding NBs, it is plausible that glial Hh non-autonomously affect NB
163 proliferation. We next explored the potential impact of glial Hh on the activities of
164 Type I NBs. As Hh is required to maintain the glial niche (Fig 2A-D, F-G) and niche
165 impairment has been shown to induce NB elimination (Read, 2018), we first assessed
166 for alterations in NB number. We found that pan-glial Hh knockdown (*repo-GAL4*)
167 did not significantly alter NB number (Fig EV3G), suggesting that NB survival is

168 unaffected. We then investigated the effects of glial Hh knockdown on NB
169 proliferation. Glial Hh knockdown using *repo-GAL4*, induced a small increase in the
170 percentage of NBs in M phase (pH3 index; Fig 2E). To assess NB S phase
171 progression, we examined EdU (5-ethynyl-2'-deoxyuridine) incorporation during a
172 15-minute time window (EdU index, Fig EV3E, yellow arrows). Here, we found EdU
173 incorporation was significantly reduced upon glial Hh knockdown, suggesting that
174 fewer NBs entered into the S phase of the cell cycle (Fig EV3F). Interestingly, a
175 similar alteration of NB pH3 and EdU index was observed upon glial niche
176 impairment caused by PI3K signalling inhibition (Speder & Brand, 2018). Therefore,
177 it is likely that glial Hh depletion indirectly causes NBs to stall at M phase via
178 inhibition of cortex glial chamber formation. To investigate the effect of glial Hh
179 depletion on NB progeny production, we conducted EdU pulse-chase assay, where
180 larvae were fed food supplemented with EdU for three hours and chased for three
181 hours in EdU-free food (Fig EV3H). Using *pros:GFP* which marks individual NB
182 lineages, we found the number of GFP⁺ and EdU⁺ cells per lineage was significantly
183 reduced upon cortex glial Hh knockdown (Fig EV3I-K). Together, these results
184 suggest that NB proliferation is inhibited upon cortex glial niche impairment caused
185 by Hh knockdown.

186
187 The sub-perineural glial Dlp and the cortex glial Jeb promote NB proliferation during
188 development; however, overexpression of these signaling molecules in the glial niche
189 was not sufficient to increase NB cell cycle rate (Cheng *et al.*, 2011; Kanai *et al.*,
190 2018). We next assessed the effect of glial Hh overexpression on NB behaviour. Pan-
191 glial induction of Hh, did not significantly alter NB number (Fig EV3G). However,
192 pan-glial and cortex glial specific Hh overexpression caused a reduction in NB EdU
193 incorporation (Fig 2H-K). Using reporter lines of Hh activity *ci-lacZ* (Schwartz *et al.*,
194 1995) and *Ptc:mCherry* (Chen *et al.*, 2017; Varjosalo & Taipale, 2008), we found that
195 Hh signaling is highly active in the NBs (Fig 2L-M'', Fig EV3P-Q''), consistent with
196 a previous report by (Chai *et al.*, 2013). Furthermore, activation of Hh transcriptional
197 activator *cubitus interruptus* (*ci^{Nc5m5m}* or *ci^{ACT}*) with a NB specific driver *dnab-GAL4*
198 (Maurange *et al.*, 2008) significantly reduced NB EdU index (Fig 2N-P),
199 phenocopying the effects of glial-Hh overexpression. Together, our data suggests that
200 high levels of glial Hh expression restricts NB cell cycle progression.
201

202 **Hh activity is modulated by Lsd-2**

203 Lipid storage droplet-2 (Lsd-2) is the *Drosophila* orthologue of the mammalian
204 perilipin2, and a widely used marker of LDs (Teixeira *et al.*, 2003) (Fig 3A-B’’).
205 Using a GFP reporter against Lsd-2 together with Hh antibody, we observed that Lsd-
206 2 and Hh colocalize to the surface of LDs in the cortex glia (yellow arrows, Fig 3C-
207 D’’). To explore whether Lsd-2 affects Hh activity, we knocked down Lsd-2 while
208 overexpressing Hh in cortex glial cells (*NP2222-GAL4>*). Lsd-2 is known to block
209 the access of lipases, thus promoting lipid storage (Teixeira *et al.*, 2003). As expected,
210 knockdown of Lsd-2 caused a significant reduction in LDs (Fig 3E-G). Furthermore,
211 this also significantly rescued the NB EdU incorporation defects caused by *hh*
212 overexpression (Fig 3H-K), suggesting Lsd-2 modulates Hh’s ability to signal to NBs.

215 **FGFR, but not EGFR or InR, activation induces cortex glial overgrowth**

216 Given that we found Hh activity is modulated by a lipid modulator Lsd-2, it is
217 possible that Hh might act in disease contexts such as glioblastoma, where glial cells
218 undergo increased proliferation and lipid metabolism alterations (Geng and Guo,
219 2017). As Hh/LD associations mostly localised to the cortex glia that enwraps NBs,
220 we explored the role for cortex glial overgrowth, using previously characterized
221 glioma models, involving the activation of FGF, InR and EGFR (Avet-Rochex *et al.*,
222 2012; Read *et al.*, 2009; Reddy & Irvine, 2011; Witte *et al.*, 2009). Firstly, we
223 characterised the effect of overexpression of a wild type form of InR, constitutively
224 activated form of the FGF receptor Heartless (Htl) and EGFR. Consistent with the
225 observation of (Avet-Rochex *et al.*, 2012), we found *htl^{ACT}* but not *InR^{wt}*
226 overexpression caused an expansion of the cortex glial niche which enwraps NBs (Fig
227 4A-C, A’-C’). In contrast, *Egfr^{ACT}* overexpression, which acts through Ras signaling,
228 didn’t affect cortex glial niche size (Fig 4A-A’, D-D’). Using cortex glial specific
229 *NP2222-GAL4* and *wrapper-GAL4* (Coutinho-Budd *et al.*, 2017; Richier *et al.*, 2017),
230 we found that cortex glial overexpression of *htl^{ACT}*, but not *InR^{wt}* or *Egfr^{ACT}* led to an
231 increase in total glial cell numbers (Fig 4E-G, I-K). Finally, *htl^{ACT}* overexpression
232 significantly increased the size of the cortex glial membrane (Fig 4E-F, H). Together,
233 our data suggest the activation of FGF, but not InR or EGFR, induced cortex glial
234 niche overgrowth.

235

236 We next explored the impact of glial niche overgrowth on NBs. Using the pan-glial
237 driver *repo-GAL4*, we measured NB cell number and its impact on the cell cycle
238 assessed with pH3 index. We found that *InR^{wt}*, but not *htl^{ACT}* and *Egfr^{ACT}*
239 overexpression, caused a reduction in NB number, reminiscent of the NB elimination
240 phenotype reported for glial *Pvr^{ACT}* overexpression (Read, 2018) (Fig 4L). In
241 addition, only overexpression of *htl^{ACT}*, which induced cortex glial overgrowth,
242 caused a reduction of NB pH3 index; while *InR^{wt}* and *Egfr^{ACT}* overexpression did not
243 significantly alter NB cell cycle progression (Fig 4M). Together, these data indicate
244 that cortex glial niche is the key glial subset that mediates glia-NB crosstalk, and its
245 expansion driven by FGF activation affected NB cell cycle progression.

246

247 **Cortex glial overgrowth mediated by FGF activation slows down NB cell cycle** 248 **progression**

249 To further investigate the effect of glial FGF activation on NB cell cycle, we assessed
250 S-phase progression via EdU incorporation. FGF activation using pan glial (*repo-*
251 *GAL4*) as well as cortex glial (*NP2222-GAL4*) drivers caused a significant reduction
252 in NB EdU index (Fig 5A-E). Consistent with the reduction in pH3 index (Fig 4M),
253 these experiments confirm that the cortex glia is responsible for NB cell cycle delay
254 induced by FGF activation.

255

256 We next examined the cell cycle length of NBs using live-cell imaging on explanted
257 brains, where three brains containing multiple NBs labelled with Dpn::GFP and
258 His::RFP (Fig 5G-H) were imaged. We found FGF activation with *repo-GAL4*
259 induced a severe slowing down of the cell cycle at 96ALH, such that we could not
260 capture any entire NB cell cycles within an 8-hour time window. Previously it was
261 reported that NBs cycle faster during earlier developmental stages (Chai *et al.*, 2013;
262 Maurange *et al.*, 2008), and therefore we imaged NB divisions at 72 ALH. We
263 observed that glial FGF activation lengthened NB cell cycle from 100.2±6.0 mins to
264 250.0±28.4 mins (Fig 5F, Movie EV1 and EV2).

265

266 To decipher which phase of the cell cycle was affected, we generated flies expressing
267 *htl^{ACT}* downstream of a *LexA* operator (*LexAop*) (Lai & Lee, 2006). Overexpressing
268 *htl^{ACT}* using *repo-LexA/LexAop* system enabled us to concurrently induce NB-specific

269 expression of Fly-Fucci using *dnab-GAL4* (as depicted in Fig 5I schematic). With
270 Fly-Fucci, cell cycle phases can be identified using combinations of two fluorescent
271 fusion proteins (Zielke *et al*, 2014). Surprisingly, we found glial FGF activation
272 significantly increased the percentage of NBs in G2/M and G1-S transition at the
273 expense of cells in G1 phase (Fig 5J-L). As the percentage of NBs in M phase
274 (reflected by pH3 index) was reduced upon glial FGF activation, we conclude that
275 these NBs are potentially stalled at G2/ G2-M and G1-S transitions of the cell cycle.
276 As a result, NBs cannot efficiently enter S phase or M phase, as indicated by reduced
277 EdU and pH3 indices (Fig 5A-E, Fig 4M), and therefore, undergo a dramatically
278 lengthened cell cycle (Fig 5F).

279
280 We expect that slowing down of the NB cell cycle upon glial FGF activation would
281 consequentially affect the number of neurons generated by NBs. To test this
282 hypothesis, we conducted EdU pulse-chase assay (3-hour feeding followed by a 4-
283 hour chase) and counted the number of EdU⁺ neurons per lineage, which are marked
284 by *dnab-GAL4::UAS-GFP*. We found that overexpression of *htl^{ACT}* with *repo-*
285 *LexA/LexAop* significantly reduced the number of EdU⁺ neurons generated per NB
286 (Fig 5M-O). Together, our data indicate that FGF activation in cortex glial cells
287 prevents NB cell cycle progression and its ability to produce the correct number of
288 neurons.

289 290 **Glial FGF activation affects NB asymmetric division, size and cell cycle exit**

291 Given that cell polarity that contributes to NB asymmetric division is established in
292 the G2/M phase, we then assessed whether NB asymmetric division is affected upon
293 glial FGF activation. In the wildtype, Inscuteable, an adaptor protein that connects the
294 aPKC/Par3/Par6 complex to the PINS/MUD/Gai complex, forms a crescent at the
295 apical side during NB mitosis (Doe, 1996); these apical complexes further direct the
296 localization of cell fate determinants (Brat/Pros/Numb) and their adaptor proteins
297 Mira (Shen *et al*, 1997) and PON to the basal cortex (as depicted in Fig EV4A). The
298 correct distribution of polarity proteins ensures the generation of a larger daughter NB
299 and a smaller GMC upon asymmetric division (Fig EV4H). We found mitotic NBs
300 displayed cytoplasmic or cortical localization of Mira and Insc upon glial FGF
301 activation (Fig EV4B-G). Furthermore, telophase NBs and GMCs were also found to
302 be more similar in size (evaluated as NB /GMC size ratio in Fig EV4I-K).

303 Intriguingly, both NB and its daughter cells were larger than their wildtype
304 counterparts, consistent with this, the average size of mitotic NBs was also
305 significantly larger upon glial *htl^{ACT}* overexpression (Fig EV4L, from $12.44 \pm 0.16 \mu\text{m}$
306 to $15.55 \pm 0.34 \mu\text{m}$). This increase in cell size was coupled with elevated cellular
307 growth, as indicated by an increase in nucleoli size (Fig EV4M-O, from $1.28 \pm 0.05 \mu\text{m}$
308 to $1.61 \pm 0.05 \mu\text{m}$). It is therefore likely that the delay between consecutive cell cycles
309 allowed these NBs to grow larger.

310

311 To investigate when glial FGF starts to impact on NB proliferation, we assessed NB
312 EdU index at 26ALH, a time point when most NBs reactivate from quiescence, and
313 commence post-embryonic neurogenesis (Fig EV4P). We found glial *htl^{ACT}*
314 overexpression did not significantly affect NB EdU incorporation during a 1-hour
315 EdU pulse (Fig EV4Q-S), suggesting that glial FGF does not affect NB reactivation at
316 the beginning of neurogenesis. We then examined whether NB termination at 24
317 hours after pupal formation (24APF) is altered (Fig EV4T). Glial *htl^{ACT}*
318 overexpression resulted in the presence of increased number of NBs at 24APF (Fig
319 EV4U-W), suggesting NB cell cycle exit is possibly delayed. However, it is not clear
320 whether NBs that persist are capable of dividing. Together we conclude glial-FGF
321 mostly exerts its effects on reducing NB cell proliferation during late larval
322 neurogenesis, coinciding with the time when Hh-LD associations become highly
323 enriched in the cortex glia. Taken together, our results revealed that NB activities
324 including its proliferation, asymmetric division and termination are affected by cortex
325 glial niche overgrowth driven by FGF activation.

326

327 **Hh mediates the effects of glial FGF signalling on NB proliferation**

328 Given that glial Hh overexpression or activation of the Hh signalling cascade in NBs
329 similarly inhibits NB proliferation and induces polarity protein delocalisation (Chai *et*
330 *al.*, 2013); it is therefore plausible that Hh lies downstream of glial FGF to regulate
331 NB activity. To test this hypothesis, we first assessed whether niche derived signals
332 including Hh and its modulators such as LD synthesis enzymes were altered. By RT-
333 qPCR, we found *hh* mRNA was upregulated by 8-fold (normalized to *rpl32*) upon
334 glial FGF activation (*repo-GAL4*, Fig 6A). Amongst genes that promote LD storage
335 (Fig 6B), *fatty acid synthase-1 (fasn1)* (Smith *et al*, 2003), and *lsd2* (an antagonist of

336 lipases which we have shown to regulate Hh activity, Fig 3), were also significantly
337 upregulated (Fig 6A). Furthermore, Hh staining was found to be more widely
338 distributed in the glial cytoplasm, rather than restricted to the ring-like structures
339 surrounding LDs upon FGF activation (Fig 6C-G, evaluated by the ratio of glial
340 cytoplasm that contains Hh staining).

341

342 We next assessed the role of Hh signalling downstream of glial FGF activation. Hh
343 knockdown (RNAi efficiency tested in (Tian *et al*, 2015)) rescued NB S-phase delay
344 (Fig 6H) without affecting glial niche size (Fig EV5A-B). In addition, we used a
345 *LexA/LexAop* binary expression system in conjunction with the *GAL4/UAS* system, to
346 simultaneously activate glial FGF and inhibit NB Hh signaling. Induction of RNAi
347 against *ci* in the NB caused a significant increase in EdU incorporation (Fig 6I, *dnab-*
348 *GAL4 > ci RNAi*). Together with glial-FGF activation, Ci knockdown also
349 significantly rescued NB EdU index (Fig 6I). Together, our results suggest that Hh
350 mediates glia-NB crosstalk downstream of glial FGF activation.

351

352 **Lipid metabolism genes lie downstream of glial FGF-NB crosstalk**

353 As we previously showed that Lsd-2 modulates Hh activity (Fig 3), we hypothesise
354 that lipid metabolism enzymes function upstream of Hh to regulate NB behaviour.
355 Consistent with this, we found induction of *hh RNAi* upon cortex glial FGF activation did
356 not alter the number of LDs (Fig EV5C-D). We next explored whether lipid
357 metabolism genes mediate the effects of glial FGF activation on NB proliferation.
358 Glial expression of RNAis targeting lipogenesis genes *fasn1* and *lsd2* efficiently
359 reduced LDs (Fig EV5E-G), but were not required for wildtype NB S phase
360 progression (Fig EV5H). However, knockdown of Fasn1 and Lsd2 significantly
361 rescued NB S-phase progression defects caused by *hhl^{ΔCT}* overexpression (Fig 6J, Fig
362 EV5I), suggesting lipid metabolism genes function downstream of glial FGF-NB
363 crosstalk.

364

365 **Fasn1 and Rasp affect glial Hh palmitoylation to regulate NB cell cycle**

366 Hh is known to be synthesized as a precursor protein (HhN) which undergoes a series
367 of post-translational modifications within the secretory pathway (Fig 7A) (Mann &
368 Beachy, 2004). The N- and C-termini of Hh proteins are covalently modified with
369 palmitate and cholesterol, respectively (Pepinsky *et al*, 1998; Porter *et al*, 1996).

370 Palmitate is a 16-carbon saturated fatty acid, which can either be diet-derived or
371 synthesized via de novo fatty acid synthesis, mediated by enzymes such as Fasn and
372 ACC (Innis, 2016; Schiller & Bensch, 1971; Slakey *et al*, 1979). Given Fasn1, an
373 enzyme involved in de novo synthesis of palmitate, is also involved in glial-FGF-NB
374 crosstalk, we assessed whether Hh palmitoylation is required downstream of glial
375 FGF signaling. Palmitoylation of the N-terminal of HhN is mediated by a dedicated
376 acyltransferase in the ER called Rasp in *Drosophila* (Fig 7A). In the embryo, Rasp
377 has been shown to be required for Hh diffusion (Chamoun *et al*, 2001; Lee &
378 Treisman, 2001; Micchelli *et al*, 2002). We found cortex glial knockdown of Rasp
379 while not required for NB S phase progression (Fig 7B), was sufficient to rescue NB
380 EdU incorporation defects caused by FGF activation (Fig 7B, *NP2222-GAL4 > htl^{ACT}*).

381
382 To test if Hh palmitoylation is required for its ability to signal to NBs, we expressed a
383 GFP tagged and non-cholesterol modifiable form of HhN in cortex glial cells
384 (Hartman *et al*, 2013; Wendler *et al*, 2006). HhN-GFP were mostly localised to
385 puncta in the glial cytoplasm, and some of these puncta made contact with NBs (Fig
386 7C-C', *NP2222-GAL4*). The number of puncta making contact with NBs was
387 significantly reduced upon knockdown of Rasp (Fig 7E-E' compared to D-D', G),
388 suggesting that palmitoylation is required for HhN transport from glia to NBs. Similar
389 to full length Hh (Fig 2J-K), glial HhN expression caused a reduction in NB EdU
390 incorporation (Fig 7H), suggesting that palmitoylated Hh is sufficient for glia-NB
391 crosstalk.

392
393 Similar to Rasp knockdown, Fasn1 knockdown caused a reduction in the number of
394 HhN-GFP puncta that made contact with NBs (Fig 7D-G). To test if Fasn1 is
395 functionally required to mediate HhN's ability to cause NB S-phase progression
396 defects, we knocked down Fasn1 while overexpressed HhN. In this setting, Fasn1
397 knockdown rescued EdU incorporation defects caused by HhN expression (Fig 7I).
398 Together, our data suggests that palmitoylation via Rasp and Fasn1, in addition to
399 regulation by Lsd-2, are required for Hh function in the context of glia-NB crosstalk.

400

401 **Glial ROS acts in parallel to lipid-Hh signaling to regulate NB proliferation in**
402 **the FGF-driven glioma model**

403 Lipid metabolism alteration in glia has previously been linked to reactive oxygen
404 species (ROS), where excessive ROS production causes LD accumulation, which in
405 turn triggers neurodegeneration (Bailey *et al.*, 2015; Liu *et al.*, 2015). Furthermore,
406 ROS upregulation is known to promote glioblastoma progression (Schieber &
407 Chandel, 2014). We therefore tested whether lipid metabolism changes in the FGF-
408 driven glioma model is correlated with ROS levels. Using ROS-inducible *gstD*
409 promoter-*GFP* reporter (Sykiotis & Bohmann, 2008), we detected a 5-fold increase in
410 *gstD-GFP* upon FGF-activation in glial cells (Appendix Fig S1A-C). To decipher the
411 effect of ROS manipulation on lipid metabolism, we overexpressed a *Catalase*
412 (Anderson *et al.*, 2005) and a *superoxide dismutase 1* (Martin *et al.*, 2009) in cortex
413 glial cells where FGF was activated to suppress ROS. And we found the number of
414 LDs, an indicator of lipid metabolism, was not altered upon ROS reduction
415 (Appendix Fig S1D', E', F', H). These manipulations were however effective in
416 partially rescuing NB EdU incorporation defects caused by glial FGF activation
417 (Appendix Fig S1I) without affecting glial overgrowth (Appendix Fig S1D, E, F, G).
418 We conclude that the production of ROS upon FGF activation in the glial niche
419 potentially acts in parallel with lipid-Hh signaling to inhibit NB proliferation.

420
421 Collectively, our findings indicate that the expression of Hh in the cortex glia is
422 required for the formation of glial chambers. Moreover, we demonstrate that
423 expression of Hh in the cortex glial niche must be restrained to prevent ectopic Hh
424 signalling in the NBs. We showed that the ability of glial Hh to signal to NBs is
425 modulated by lipid metabolism enzymes *Fasn1* and *Lsd-2*. Upon overgrowth of the
426 cortex glial niche induced by FGF activation, Hh, lipid metabolism regulators as well
427 as ROS are upregulated in the niche, which in turn slow down the NB cell cycle, and
428 affect its ability to generate a full repertoire of neurons (Fig 7J).

429

430 **Discussion:**

431 In the mammalian CNS, neuron and glia interactions are complex. Astrocytes that are
432 structurally related to the cortex glia in *Drosophila*, enwrap multiple neurons and
433 NSCs, and are known to modulate adult neurogenesis through soluble signals such as
434 morphogens and extracellular matrix proteins (Spampinato *et al.*, 2019). Similarly, the
435 *Drosophila* cortex glial niche, which form chambers that encapsulate NB and its
436 progeny, creates microenvironments that are required for NB maintenance and

437 neuronal maturation (Coutinho-Budd *et al.*, 2017; Dumstrei *et al.*, 2003; Peraanu *et*
438 *al.*, 2005; Speder & Brand, 2018; Yuan *et al.*, 2020). Furthermore, glial overgrowth
439 observed in the context of glioblastoma has recently been shown to affect NBs and
440 neuronal survival (Portela *et al.*, 2019; Read, 2018). In this study, we have identified a
441 new mechanism of glia-NB crosstalk via Hh modulated by lipid regulators, that affect
442 NB cell cycle progression and lineage size.

443 During development, NB lineages have been shown to produce its own pool of Hh
444 (Chai *et al.*, 2013), which likely act redundantly with the glial Hh. However, glial Hh
445 levels must be tightly regulated to ensure NB can progress through the cell cycle, and
446 produce the appropriate number and types of neurons. Similar to pro-proliferative
447 roles of Hh in the astrocytes of the mammalian brain (Ugbode *et al.*, 2017),
448 *Drosophila* glial- Hh is autonomously required for the growth of cortex glial cells and
449 the formation of the glia chambers which surround NBs. Hh knockdown caused NB
450 clustering which in turn induced moderate NB cell cycle delay, and reduced neuron
451 production, reminiscent of phenotypes reported for glial PI3K inhibition which also
452 caused disrupted glial chambers (Speder and Brand, 2018). This contrasts with Hh
453 and Smo mutant NB phenotype, previously reported by (Chai *et al.*, 2013), where
454 they found the loss of NB Hh signaling caused a significant increase in NB clone size.
455 Therefore, our data supports the model that glial-Hh is required for niche elaboration,
456 which in turn affects NB proliferation.

457 While moderate level of Hh is permissive for NB cell cycle progression, excess glial
458 Hh through either its overexpression or via FGF activation also affects NB cell cycle
459 progression, and its ability to produce neurons. In this case, excess glial Hh non-
460 autonomously activated Hh signaling in the NBs, which is sensitive to this ligand.
461 Different to other niche-derived secreted molecules, such as Dlp (Kanai *et al.*, 2018)
462 and Jeb (Cheng *et al.*, 2011), which are necessary but not sufficient to induce changes
463 in the NB, Hh plays a physiologically relevant role in disease models of glial
464 overgrowth caused by FGFR activation. TCGA data shows that FGFR1–4 are
465 expressed in different combinations in glioblastoma patient samples, and FGFR1 is an
466 important contributor to poor outcome in glioblastoma (Jimenez-Pascual &
467 Siebzehnrubl, 2019).

468

469 In the *Drosophila* brain, cortex glia appears to be the key glial subtype that is responsible
470 for glia-NB crosstalk. Only FGF but not glial InR and EGFR activation caused an
471 expansion of cortex glia (Avet-Rochex *et al.*, 2012; Avet-Rochex *et al.*, 2014), which
472 in turn increased Hh levels, and reduced NB proliferation. In cortex glial cells, Hh
473 closely associates with LDs which are analogous to lipoproteins, that have been
474 shown to transport Hh for long-range signalling (Palm *et al.*, 2013; Panakova *et al.*,
475 2005). Different from lipoproteins, LDs mainly act as lipid storage organelles, and are
476 less mobile. Consistent with this, upon FGF activation, genes that promote lipid
477 synthesis and storage were upregulated. So how does lipid metabolism regulate Hh
478 function? We showed two mechanisms. The first mechanism involves the interaction
479 between lipid storage regulator Lsd-2 and Hh that colocalize on the surface of LDs
480 and that Lsd-2 modulates Hh's ability to signal to NBs. It is not yet clear how Lsd-2
481 directly regulates Hh activity, but we think it is possible that Lsd-2 might physically
482 interact with Hh and affect its secretion; or alternatively, Lsd-2 competes with Hh for
483 positions on the surface of LDs, and pushes Hh into the cytoplasm. The second
484 mechanism involves Fasn and Rasp which regulates Hh-post translational
485 modification. Given that lipid synthesis and LDs have recently been reported to be
486 upregulated in glioma, and are emerging as important biomarkers and metabolic
487 targets (Geng & Guo, 2017; Guo *et al.*, 2013), it would be interesting to further
488 explore how lipid metabolism regulators affect signaling in this context. In addition
489 to the Hh-LD axis, ROS which is implicated in glioblastoma (Schieber & Chandel,
490 2014) also lies downstream of FGF mediated glia-NB crosstalk. Knockdown of both
491 Hh and ROS-axes significantly rescued NB cell proliferation defects caused by FGF
492 activation in the glial cells. The phenomenon we reported here together with other
493 reports that glioblastoma affects the survival or proliferation rate of its neighbours
494 (Portela *et al.*, 2019; Read, 2018) poses an interesting but yet unexplored prospect
495 that glioma outcompetes NBs within the CNS for limited energy and nutrient
496 resources.

498 **Materials and Methods**

499 **Fly husbandry and strains**

500 Fly stocks were reared on standard *Drosophila* media at 25 °C. Crosses for
501 overexpression and knockdown experiment were set up at 25 °C, and after a day, the
502 progenies were moved to 29 °C, unless otherwise stated.

503 The fly strains used were: *repo-GAL4* (BDSC7415), *NP2222-GAL4*
504 (KYOTO112830), *wrapper-GAL4* (Coutinho-Budd *et al.*, 2017; Richier *et al.*, 2017),
505 *dnab-GAL4* (From Alex Gould lab), *repo-LexA::GAD* (BDSC67096), *w¹¹¹⁸*, *UAS-*
506 *htl^{ACT}* (BDSC5367), *UAS-Egfr^{ACT}* (BDSC59843), *UAS-InR^{w^t}* (BDSC8262), *LexAop-*
507 *htl^{ACT}* (generated in this paper), *UAS-GFP*, *UAS-mGFP*, *UAS-dcr2*, *UAS-FUCCI*
508 (BDSC55110), *UAS-lacZ*, *UAS-luc* (BDSC64774), *UAS-hh* (from Thomas B.
509 Kornberg lab), *UAS-hh.N.GFP* (BDSC81023), *UAS-lsd2* (from Alex Gould lab),
510 *UAS-ci^{Nc5m5m(ACT)}* (from Yu Cai lab), *UAS-cat.A* (BDSC24621), *UAS-Sod1.A*
511 (BDSC24754), *UAS-mcherryRNAi* (BDSC35785), *UAS-GFPRNAi* (BDSC9331),
512 *UAS-fasn1RNAis* (NIG3523R-2, NIG3523R-6), *UAS-lsd2RNAis* (VDRC102269,
513 BDSC34617 and BDSC32846), *UAS-hhRNAis* (VDRC43255, BDSC25794), *UAS-*
514 *ciRNAis* (NIGHMJ23860, NIG2125R-2), *UAS-raspRNAis* (NIG11495R-1,
515 NIG11495R-2), *dpnGFP* (BDSC59755), *his2AV-mRFP* (Kieran Harvey lab), Hh:GFP
516 BAC, Ptc:mcherry BAC, *ci-lacZ* (all three lines are generated by Thomas B.
517 Kornberg lab), *lsd2YFP* (KYOTO115301), *gstd-GFP* (From Tatsushi Igaki lab),
518 *pros-GFP* (VDRC318418). The *repo*-MARCM stock was: *UAS-RedStinger*; *repo-flp*,
519 *repo-GAL4*, *UAS-actinGFP*; *FRT82B*, *tub-gal80* (from Joseph M. Bateman lab).
520 *w*; *FRT82B* was used to generate surface glial clones.

521 Immunostaining

522 Larval and pupal brains were dissected in PBS, fixed for 25 minutes in 4%
523 formaldehyde (Sigma-Aldrich, #F8775) in PBS and rinsed in 0.5% PBST (PBS +
524 0.5% TritonX-100 (Sigma-Aldrich, #T8787)). For immunostaining, brains were
525 incubated with primary antibodies overnight at 4 °C, followed by an overnight
526 secondary antibody incubation at 4 °C. Samples were mounted in 80% glycerol in
527 PBS for image acquisition. The primary antibodies used were mouse anti-Mira (1:50;
528 gift of Alex Gould), rat anti-Mira (1:100, Abcam, #ab197788), rabbit anti-Mira
529 (1:200, gift of Rita Sousa-Nunes), rat anti-pH3 (1:500; Abcam, #ab10543), chick anti-
530 GFP (1:2000; Abcam, #ab13970), rabbit anti-RFP (1:100, Abcam, #ab62341), mouse
531 anti-Fibrillarlin (1:200, Abcam, #ab4566), rabbit anti-Hh (1:500, gift of Isabel
532 Guerrero), rabbit anti-Insc (1:20, gift of William Chia). Secondary donkey antibodies

533 conjugated to Alexa 555 and Alexa 647 and goat antibody conjugated to Alexa 405,
534 488, 555 and 647 (Molecular Probes) were used at 1:500. DAPI (Molecular Probes)
535 was used at 1: 10,000.

536 **EdU labelling and pulse chase**

537 EdU in vitro labelling was used to identify actively dividing NBs, larval brains were
538 incubated in 10 μ M EdU /PBS for 10-15 mins (for 96ALH brains) or 1 hr (Fig EV4P-
539 S), followed by fixation, and development using Click-iT Plus EdU Cell Proliferation
540 Kit for imaging, Alexa Fluor 647 dye (Invitrogen, #C10640), following the
541 manufacturer's instruction. The brain tissues were then stained with Mira to label
542 NBs. Control and experimental brains were processed in the same tube to ensure they
543 were exposed to the same incubation conditions.

544 EdU pulse-chase was used to identify the progeny of dividing NBs. 3rd instar larvae
545 were fed with instant fly food supplemented with 100 μ g/mL EdU (Lee *et al*, 2006)
546 for 3 hours. They were then transferred to standard *Drosophila* media for a 3 or 4
547 hour-EdU chase. Wandering stage larvae were collected for brain dissection, followed
548 by fixation, development and immunostaining as described above.

549 **LD staining**

550 For LD staining, larval brains were dissected in PBS, fixed, and rinsed in PBS before
551 incubation in HCS LipidTOX Red Neutral Lipid Stain (Invitrogen, #H34476, 1:1000
552 in PBS) for 1 hour. These samples were then rinsed and mounted in PBS and imaged
553 directly to preserve LD morphology. For experiments that require LD staining
554 together with immuno-staining, the tissues were rinsed three times in PBS after
555 immunostaining to remove all PBST, and incubated with LD dyes as described above.
556 The tissues were mounted in 80% glycerol in PBS for imaging.

557 **Imaging and image processing**

558 Images were collected on a Leica SP5 or Olympus FV3000 confocal microscopes and
559 analysed using Fiji (<https://imagej.net/Fiji>). Z stacks of CBs were imaged and the
560 ventral side of the CB was shown as the representative image unless otherwise stated.

561 **Live cell imaging**

562 Dissected brains (72ALH) were cultured in Schneider's culture medium
563 supplemented with 10% inactivated FBS, 2% Penicillin-Streptomycin solution

564 (Sigma-Aldrich, #P4458), 20 μ M glutamine and Schneider's culture medium (Gibco,
565 #21720024) and dissected fat body from the same animals. The brains were imaged in
566 a μ -Slide 8 well (Ibidi, #80806) on an Olympus FV3000 microscope using resonance
567 scanner in 16Bit mode, with a 40x/0.95 lens and 2x zoom. Z stacks with 2 μ m
568 intervals were captured every 2 minutes over a period of 3-8 hours. Laser intensity
569 were kept low to avoid cytotoxicity. AVI movies were generated using Fiji.

570 All the images were processed using Adobe Photoshop and compiled using Adobe
571 Illustrator. For the purpose of better presentation, image brightness adjustments were
572 applied equally to controls and experiments.

573 **Quantitative reverse transcription PCR**

574 For gene expression analysis, 20 dissected late 3rd instar larval brains were lysed in
575 300 μ l TRI Reagent (ZYMO Research, #R2061) to form one biological replica. Three
576 biological replicates were prepared for each genotype: *repo-GAL4*> *w¹¹¹⁸* and *repo-*
577 *GAL4*> *htl^{ACT}*. Total RNA was extracted using a Direct-zol RNA Microprep Kit
578 (ZYMO Research, #R2061) and 1 μ g of total RNA from each sample was reverse
579 transcribed into cDNA using ProtoScript II First Strand cDNA Synthesis Kit (NEB,
580 #E6560S) according to the manufacturer's instructions. The qPCR was performed
581 using the stepOnePlus real-time PCR system (Applied Biosystems) using Fast SYBR
582 Green master mix reagent (Applied Biosystems, #4385612). Gene expression levels
583 were normalized to *rpl32* (in Fig 6A, one representative data set of *rpl32* was plotted)
584 and calculated using the $2^{-\Delta\Delta C_t}$ method. The primers were either designed using
585 Primer-BLAST (<https://www.ncbi.nlm.nih.gov/tools/primer-blast/>), or obtained from
586 FlyPrimerBank (<https://www.flyrnai.org/flyprimerbank>).
587 listed in Reagents Table.

588 **Molecular cloning**

589 A constitutively active form of *htl*, comprising the dimerization domain of the
590 bacteriophage λ repressor (Michelson *et al*, 1998), was amplified from the genomic
591 DNA of the fly stock: *UAS-htl^{ACT}* (BDSC5367), using a forward primer, 5'-
592 CAACTGCAACTACTGAAATCTGCC-3', and a reverse primer 5'-
593 CCCCCTCTAGATTAATAATTACACCACTTCTGC-3'. The resulting PCR product
594 was digested with *NotI* (Promega, #R6431) and *XbaI* (Promega, #R6181), which cut
595 at the restriction sites as indicated in the reverse primer (underlined above). The

596 plasmid, *pJFRC19-13XLexAop2-IVS-myr::GFP* (Addgene, #26224), was identically
597 digested to remove *myr::GFP*. The restriction fragment, *NotI-htl^{ACT}-XbaI*, was
598 subsequently cloned into the digested LexAop vector (Pfeiffer *et al.*, 2010). The
599 reconstructed plasmid was sequenced and injected into flies carrying an attP2 docking
600 site (BDSC25710).

601 **Quantification and analysis**

602 **Hh and LipidTOX intensity profile**

603 A flat line was drawn across each LDs as described in Fig 1H, and single pixel values
604 of Hh and LipidTOX staining were generated along line using “*Analyze-Plot Profile*”
605 tool of Fiji. The intensity profiles of Hh and LipidTOX for each LDs were compiled
606 using Prism-GraphPad, with Y axis reflecting pixel values and X axis reflecting relative
607 position on LDs. The relative LD position was generated by dividing the line position
608 values to each line length. This normalization was conducted because line length is
609 different in each sample due to the variations in LD size. The Hh intensity at the position
610 X_0.9 (Fig 1H, at the surface of LDs) was compared with that of the position X_1.0
611 (Fig 1H, just outside of LDs) to illustrate the Hh-LD association described in Fig 1C-
612 G”.

613 **Cortex glial membrane size, LD volume and glial number measurement**

614 Cortex glial membrane volume (*NP2222-GAL4>UAS-mGFP*) and LD volume were
615 measured from three-dimensional reconstruction of confocal Z stacks (2- μ m step-
616 size) with Volocity software (Improvision). Glial cell number were automatically
617 counted with a Fiji plug-in “DeadEasy Larval Glia” (Forero *et al.*, 2012).

618 **Cell cycle speed**

619 **The number of Type I NBs** in each brain lobe was manually counted using Fiji. Type
620 I NBs were distinguished from other *Mira*⁺ cells by size and morphology: Newly
621 generated GMCs are smaller than Type I NBs as shown in Fig EV3E (white arrows).
622 Type II NBs are associated with more *Mira*⁺ progeny cells.

623 **EdU index:** For 26ALH larval brains, EdU voxels of the whole brain were measured
624 with Volocity software (Improvision) to indicate NB re-entry into cell cycle. Glial EdU
625 voxels represents only a small amount of the total EdU voxels (Sousa-Nunes *et al.*,
626 2011). Normalized EdU voxels were calculated by dividing EdU voxels to the mean
627 voxel counts of the control. For 96ALH larval brains, the number of EdU⁺ Type I NBs

628 was counted in the CB of each brain lobe. EdU index is calculated as number of EdU⁺
629 NBs normalised to control EdU⁺ NBs. The number of EdU⁺ NBs reflects the NBs that
630 progress through S phase in a 10 or 15-min time window relative to control. We have
631 utilised GAL4 driver > *w¹¹¹⁸*, *UAS-mcherryRNAi* or *UAS-luc* interchangeably as
632 controls in our experiments, as we found EdU incorporation did not significantly alter
633 between these controls (Appendix FigS2). The total number of Type I NBs is not altered
634 in experiments where EdU index was used to determine NB cell cycle speed. For EdU
635 pulse-chase quantification, the number of progeny cells that inherit EdU from each
636 dividing NBs was counted to indicate the speed by which NBs generate their progeny.

637 **pH3 index:** represented as the % of Type I NBs in M phase (pH3⁺) / the total number
638 of Type I NBs.

639 **Cell cycle lengths in NBs** were measured as the time between two consecutive cell
640 divisions. The cell cycle lengths of NBs from at least three different ex vivo brains
641 were plotted for each genotype.

642 **CNS and Cell size measurement**

643 Brain lobes, NB, GMC and nucleoli diameters were estimated by averaging orthogonal
644 measurements of diameter, with a single confocal section on Fiji. The volume of each
645 brain lobe (CNS) was calculated by the formula: $V=4/3\pi r^3$.

646 **Localization of asymmetric determinants** was assessed in M phase NBs that display
647 a condensed metaphase plate marked by pH3. Clear crescent localisation was counted
648 as correct localisation, and cytoplasmic or cortical localisation was counted as
649 mislocalisation.

650 **The distribution of Hh in glia** was measured on Fiji with the formula: The area of
651 glia that contains Hh / Total glial area. The detailed procedure: Total glial area:
652 Adjust Threshold (Default) > Analysis > measure (area); The area of glia that
653 contains Hh: Create selection of glial channel by Adjusting Threshold (Default) >
654 Restore selection in the Hh channel > clear outside > measure area.

655 **Total intensity**

656 The sum projection confocal image was used for the intensity measurement on Fiji
657 using the formula: CTCF (corrected total cell fluorescence) = Integrated density –
658 (Area of selected cell x mean fluorescence of background readings) (McCloy *et al*,
659 2014).

660 **Statistical analysis**

661 *P*-values were calculated by two-tailed, unpaired Student's *t*-test, with equal sample
662 variance; The Welch's correction was applied in case of unequal variances.
663 Kolmogorov-Smirnov test was used to test data normality. Mann-Whitney test was
664 used when data deviated from a normal distribution. Data are presented as mean ±
665 SEM in the main text.

666

667 **Data Availability Section**

668 **This study includes no data deposited in external repositories.**

669

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686

687 **Author contributions**

688 Q.D, M.Z, F.F, T.L, S.G conducted the experiments; Q.D and L.Y.C designed the
689 experiments and wrote the paper.

690

691 **Conflict of interest**

692 The authors declare no competing interests.

693

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897

898

899 **Figure legends**

900 **Figure 1: Hh is localized to the LDs within cortex glial cells**

901 Images in this and following figures are of larval central brains (CBs) at 96ALH.

902 A) Schematic showing NBs that undergo asymmetric division to self-renew and
903 produce GMCs, which terminally differentiate to generate post-mitotic neurons or
904 glial cells (left). Each NB is surrounded by a microenvironment, composed of glial
905 cells (right).

906 B-D'') Representative images showing that Hh accumulates on the surface of LDs in
907 glial cells of the CB (yellow arrows), quantified in (I) (n=20 LDs). Glial cells are
908 marked by *repo-GAL4>GFP* and CB is circled in (B).

909 E-G'') In the posterior compartment of the developing wing disc (WD) pouch region
910 where Hh is expressed, LDs and Hh are not tightly associated (white arrows),
911 quantified in (J) (n = 17 LDs).

912 H-J) Hh-LD association is quantified by plotting the pixel intensities of both Hh
913 (cyan) and LDs (red) along a line across LDs. Y axis represents gray intensity values,
914 and X axis represents relative LD position.

915 K-L'') Hh-LD associations are observed in the cortex glia (yellow arrows, *NP2222-*
916 *GAL4>mGFP*).

917 Data information: Hh is detected with a Hh antibody and LDs are visualized with
918 LipidTOX unless otherwise stated. (D-D''), (G-G''), (L-L'') are zoomed in images of
919 (C, F, K), respectively. Scale bar=50 μ m in (C and F), scale bar=20 μ m in (D-D'', K-

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920 L''), scale bar=10 μm in (G-G''). Error bar represents SEM. In (I): Welch's t test,
921 (****) $p < 0.0001$. In (J): unpaired t test, (ns) $p = 0.7113$.

922

923

924 **Figure 2: Hh autonomously regulates cortex gliogenesis and non-autonomously**
925 **regulates NB proliferation**

926 A-D) Representative images showing that upon knockdown of Hh in cortex glial cells
927 (*NP2222-GAL4>mGFP* with *UAS-dcr2*), cortex glial membrane and overall Repo+
928 glial cell number are significantly reduced, quantified in (C) (n=7, 8 brain lobes) and
929 (D) (n=7, 8 brain lobes), respectively.

930 E-G) Hh knockdown in glia (*repo-GAL4>GFP*) results in niche disruption and
931 clustering of NBs (circled with yellow dashed line), as well as an increase in the
932 percentage of NBs in M phase (pH3⁺), quantified in (E) (n=12, 10 brain lobes).

933 H-K) Representative images showing that Hh overexpression using pan-glial (*repo-*
934 *GAL4*) and cortex-glial (*NP2222-GAL4*) drivers both result in a decrease in NB EdU
935 index, quantified in (J) (n=16, 20 brain lobes) and (K) (n=11, 14 brain lobes),
936 respectively.

937 L-M'') Representative images showing *ci-lacZ* is expressed in NBs (yellow arrows).
938 (M-M'') are zoomed in images of (L).

939 N-P) Overexpression of *ci^{ACT}* in NBs (*dnab-GAL4*) reduces EdU index, quantified in
940 (P) (n=12, 10 brain lobes).

941 Data information: NBs are marked with Mira and EdU⁺ NBs are circled by yellow
942 dashed line. Scale bar=50 μm . Error bar represents SEM. In (C): Mann-Whitney test,
943 (**) $p = 0.0059$. In (D): unpaired t test, (*) $p = 0.0176$. In (E): unpaired t test, (*)
944 $p = 0.0215$. In (J): unpaired t test, (***) $p = 0.0002$. In (K): unpaired t test, (***)
945 $p = 0.0001$. In (P): unpaired t test, (****) $p < 0.0001$.

946

947 **Figure 3: Hh activity is modulated by Lsd-2**

948 A-B'') Representative images showing that Lsd2YFP is localised to the surface of
949 LDs (yellow arrows).

950 C-D'') Lsd2YFP co-localises with Hh antibody staining in the cortex glia (yellow
951 arrows) but not surface glial cells (white arrow).

952 E-G) Lsd-2 knockdown in cortex glial cells (*NP2222-GAL4*) where *hh* is
953 overexpressed effectively reduces LD number in CB (outlined in yellow dashed
954 lines).

955 H-K) Representative images showing that NB EdU index is rescued upon Lsd-2
956 knockdown in cortex glial cells (*NP2222-GAL4*) where *hh* is overexpressed,
957 quantified in (K) (n=11, 14; 16, 14; 6, 14 brain lobes). The *NP2222-GAL4>w¹¹¹⁸* vs
958 *hh^{OE}* columns depict the same data as Fig 2K. EdU⁺ NBs are circled with yellow,
959 dashed lines.

960 Data information: (B-B'', D-D'') are zoomed in images of (A and C). Scale bar=50
961 μ m in (A, C, E-J). Scale bar=10 μ m in (B-B'' and D-D''). Error bar represents SEM.
962 In (K): unpaired t test, (***) $p=0.0001$; unpaired t test, (**) $p=0.0053$; unpaired t test,
963 (*) $p=0.0349$.

964

965 **Figure 4: Activation of *hhl^{ACT}* but not *InR^{wt}* and *Egfr^{ACT}* induces cortex glial** 966 **overgrowth**

967 A-D') Representative images showing that pan-glial overexpression of *hhl^{ACT}*, but not
968 *InR^{wt}* or *Egfr^{ACT}* causes an expansion of cortex glia that enwraps NBs. Glial cells are
969 marked with *repo-GAL4>GFP*, and NBs are marked with Mira. (A', B', C' and D')
970 are zoomed in images of (A, B, C and D), respectively.

971 E-K) Representative images showing that cortex glial overexpression of *hhl^{ACT}* but not
972 *InR^{wt}* or *Egfr^{ACT}* causes an increase in glial cell (Repo⁺) numbers and cortex glial
973 membrane size, quantified in (K) (n=10, 10, 10; 7, 12 brain lobes) and (H) (n=10, 10
974 brain lobes), respectively. *NP2222-GAL4>mGFP* is used to mark cortex glial
975 membrane in (E-G) and *wrapper-GAL4>* is used in (I, J).

976 L) Glial (*repo-GAL4>*) overexpression of *InR^{wt}* but not *hhl^{ACT}* or *Egfr^{ACT}* significantly
977 reduces the number of CB NBs (n= 10, 10, 8, 11 brain lobes).

978 M) Glial (*repo-GAL4>*) overexpression of *hhl^{ACT}* but not *InR^{wt}* or *Egfr^{ACT}* significantly
979 reduces the pH3 index of CB NBs (marked with Mira and pH3 in (A-D')) (n= 10, 10,
980 8, 11 brain lobes).

981 Data information: Scale bar=50 μ m. Error bar represents SEM. In (H): Welch's t test,
982 (****) $p<0.0001$. In (K): Welch's t test, (****) $p<0.0001$; Welch's t test, (ns)
983 $p=0.7905$; unpaired t test, (ns) $p=0.0941$. In (L): unpaired t test, (ns) $p=0.9140$;

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984 Welch's t test, (*) $p=0.0170$; unpaired t test, (ns) $p=0.9825$. In (M): unpaired t test,
985 (***) $p=0.0038$; unpaired t test, (ns) $p=0.4172$; unpaired t test, (ns) $p=0.5523$.

986

987 **Figure 5 Cortex glial overgrowth mediated by *htl^{ACT}* overexpression triggers NB**
988 **cell cycle delay**

989 A-E) Representative images showing that both pan-glial (*repo-GAL4*>) and cortex
990 glial (*NP2222-GAL4*>) *htl^{ACT}* overexpression significantly reduce NB EdU index,
991 quantified in (E) (n=15, 14; 10, 16 brain lobes). (A, B, C, D) are single sections with
992 Mira and EdU staining, and (A', B', C', D') are Z-projection of the EdU staining.

993 F-H) Representative still images from ex vivo CNS live imaging at 72ALH showing
994 that pan-glial (*repo-GAL4*) *htl^{ACT}* overexpression lengthens NB cell cycle, quantified
995 in (F) (n= 25, 6 NBs imaged from 3 brains per genotype). The cell cycle length is
996 measured as the length between consecutive divisions. NBs (Dpn::GFP, red; Histone
997 RFP, grey) are circled with blue dashed lines.

998 I) Schematic depicting concurrent glial FGF activation (*repo-LexA* > *LexAop-htl^{ACT}*),
999 and NB overexpression of fly-FUCCI (*dnab-GAL4* > *UAS-GFP::E2F1*, *UAS-*
1000 *RFP::CycB*). The fly-FUCCI system utilizes the fusion protein GFP::E2F1 (a marker
1001 for cells in G2, M and G1 phase) and RFP::CycB (a marker for cells in S, G2, M
1002 phase) to monitor cell cycle progression. Cells in G1 phase are GFP⁺ RFP⁻ (green),
1003 cells in G2/ M phase are GFP⁺ RFP⁺ (yellow), and cells in S phase are GFP⁻ RFP⁺
1004 (red), whereas, cells in G1-S transition are weakly labelled by both GFP and RFP
1005 (grey).

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1006 J-L) Representative images showing that the percentage of NBs in G1-S transition
1007 and G2/M phase are both significantly increased with significantly less cells
1008 remaining in G1 phase, quantified in (L) (n= 9, 10 brain lobes). NBs in G1 phase
1009 (Mira⁺, GFP⁺) are circled by green dashed lines; NBs in G1-S transition (Mira⁺, GFP⁻
1010 RFP⁻) are circled by grey dashed lines; NBs in S phase (Mira⁺, RFP⁺) are circled by
1011 red dashed lines; and NBs in G2/M phase (Mira⁺, GFP⁺ RFP⁺) are circled by yellow
1012 dashed lines.

1013 M-O) Representative images showing that the number of EdU⁺ neurons generated per
1014 NB is significantly reduced upon pan-glial overexpression of FGF (*repo-LexA* >
1015 *LexAop-htl^{ACT}*; yellow arrows), quantified in (O) (Box plot, the boxes extend from the
1016 25th to 75th percentiles; the median is marked by a central band inside the box; and the
1017 whiskers go down to the minimum value and up to the maximum value. n= 94, 127

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1018 NB lineages imaged from 5 and 7 brain lobes, respectively). NB lineages are marked
1019 with *dnab-gal4 > GFP*.
1020 Data information: Scale bar=50 μm in (A-D', J-K'); Scale bar=10 μm in (G, H); Scale
1021 bar=20 μm in (M, N). Error bar represents SEM unless otherwise stated. In (E):
1022 unpaired t test, (****) $p < 0.0001$; unpaired t test, (****) $p < 0.0001$. In (F): Mann-
1023 Whitney test, (****) $p < 0.0001$. In (L): G1: Mann-Whitney test, (****) $p < 0.0001$;
1024 G1-S: unpaired t test, (**) $p = 0.0064$; S: unpaired t test, (ns) $p = 0.1757$; G2/M:
1025 unpaired t test, (*) $p = 0.0171$. In (O): Mann-Whitney test, (****) $p < 0.0001$.

1026

1027 **Figure 6 Hh and lipid metabolism regulators mediate the effects of glial *htl^{ACT}* on**
1028 **NB proliferation**

1029 A) Pan-glial (*repo-GAL4*) *htl^{ACT}* overexpression causes upregulation of *hh*, *fasn1* and
1030 *lsd2* transcripts (n = 3 biological replicates pooled from 20 brains for each genotype;
1031 for each biological replicate, we ran 3 technical replicates for each PCR reaction). The
1032 lipogenesis genes (*acc*, *lipin*), and lipolysis gene *bmm* transcripts are not significantly
1033 altered. We utilised *rpl32* as a reference gene in these experiments, as it is not altered
1034 by *htl^{ACT}* overexpression. The data are represented by log2 fold change relative to the
1035 control (*repo-GAL4 > w¹¹⁸*).

1036 B) Schematic depicting lipogenesis and lipolysis. Lipogenesis begins with *de novo*
1037 synthesis of fatty acids by carboxylation of cytosolic acetyl-CoA via acetyl-CoA
1038 carboxylase (ACC) and elongation of fatty chain via fatty acid synthase (Fasn, red).
1039 Dietary-derived and *de novo*-generated fatty acids are converted into fatty acylCoA,
1040 which re-localizes to ER and participates in triglyceride (TAG) synthesis with
1041 glycerol-3 phosphate (G3P). This process is mediated by a series of enzymes:
1042 glycerol-3-phosphate acyltransferase (GPAT), acylCoA acylglycerol-3-phosphate
1043 acyltransferases (AGPAT), Lipin (a phosphatidate phosphatase), diacylglycerol
1044 acyltransferase (DGAT, red). TAG is translocated from the ER to the core of the
1045 intracellular organelles called LDs. On the surface of LDs, a triglyceride lipase called
1046 Brummer (Bmm), and its inhibitor Lsd-2, antagonistically control TAG storage.

1047 C-G) Representative images showing that Hh staining normally localised to a ring
1048 like structure (yellow arrows), becomes delocalises to the glial cytoplasm upon *htl^{ACT}*
1049 overexpression, quantified in (E) (n=4, 6 brain lobes). Glial cells are marked with
1050 *repo-GAL4 > GFP*. (E-F') are zoomed in images of (C-D').

1051 H) Cortex glial (*NP2222-GAL4>*) overexpression of two independent *hh* RNAis
 1052 significantly rescue EdU incorporation defects caused by *htl^{ACT}* overexpression (n=10,
 1053 16; 7, 12; 14, 11 brain lobes). The *NP2222-GAL4> w¹¹¹⁸* vs *htl^{ACT}* columns depict the
 1054 same data as those in Fig 5E.

1055 I) Knockdown of NB Hh signalling pathway (*dnab-GAL4> UAS-ciRNAi*) rescues NB
 1056 EdU incorporation defects induced by glial *htl^{ACT}* overexpression (*repo-LexA>*
 1057 *LexAop-htl^{ACT}*). Induction of *ciRNAi* in NBs alone increases NB EdU incorporation
 1058 (n=8, 13; 10, 10; 30, 26; 12, 7 brain lobes).

1059 J) The NB EdU incorporation defects due to cortex glial (*NP2222-GAL4>*)
 1060 overexpression of *htl^{ACT}*, is significantly rescued by overexpression of RNAis against
 1061 *fasn1* and *lsd2*, compared to corresponding control RNAis (n=10, 16; 25, 17; 14, 8
 1062 brain lobes). The *NP2222-GAL4> w¹¹¹⁸* vs *htl^{ACT}* columns depict the same data as
 1063 those in Fig 5E. The control column for *htl^{ACT}; lsd2 Ri^{KK102269}* depicts the same data as
 1064 the control column for *htl^{ACT}; hh Ri^{BL25794}* in Fig 6H.

1065 Data information: Scale bar=50 μ m. Error bar represents SEM. In (A), t test was
 1066 conducted to compare log transformed fold change: *rpl32*: Welch's t test, (ns)
 1067 $p=0.9355$; *hh*: unpaired t test, (***) $p=0.0061$; *fasn1*: unpaired t test, (*) $p=0.0242$;
 1068 *acc*: unpaired t test, (ns) $p=0.7570$; *lipin*: unpaired t test, (ns) $p=0.1222$; *lsd2*:
 1069 unpaired t test, (***) $p=0.0074$; *bmm*: unpaired t test, (ns) $p=0.9356$. In (G): Man-
 1070 Whitney test, (***) $p=0.0095$. In (H): unpaired t test, (****) $p<0.0001$; Mann-Whitney
 1071 test, (**) $p=0.0031$; unpaired t test, (**) $p=0.0012$. In (I): unpaired t test, (**) $p=0.0019$;
 1072 unpaired t test, (****) $p<0.0001$; Mann-Whitney test, (***) $p=0.0006$;
 1073 unpaired t test, (****) $p=0.0003$. In (J): unpaired t test, (****) $p<0.0001$; unpaired t
 1074 test, (****) $p<0.0001$; Welch's t test, (***) $p=0.0004$.

1076 **Figure 7 Fasn1 affects glial Hh palmitoylation to regulate its signalling to NBs**

1077 A) Schematic depicting Hh auto-processing, which starts with the cleavage of the
 1078 protein into a C-terminal part (Hh.C, yellow) and a N-terminal part (Hh.N, blue), with
 1079 simultaneous covalent addition of cholesterol. Palmitate, from either diet or *de novo*
 1080 lipogenesis (via ACC and Fasn), is added onto Hh-N, in a reaction catalysed by an
 1081 acyltransferase, encoded by *rasp*.

1082 B) Inhibition of palmitoylation (via two independent *rasp RNAis*) rescues NB EdU
 1083 incorporation defects induced by cortex glial (*NP2222-GAL4>*) *htl^{ACT}* overexpression,

Commented [HC9]: Panel legends must be in alphabetical order, i.e. (G) cannot come after (H) & (I). Please find suggested changes to the panel labels in the text.

1084 while knockdown of Rasp in cortex glial cells alone does not alter NB EdU index
1085 (n=10, 15; 10, 16; 18, 21; 26, 29 brain lobes). The *NP2222-GAL4> w¹¹¹⁸ vs htl^{ACT}*
1086 columns depict the same data as those in Fig 5E.
1087 C-C') Representative images showing that Hh.N.EGFP (which cannot undergo
1088 cholesterol modification) are found as puncta on the surface of NBs (yellow arrows)
1089 when overexpressed in neighbouring cortex glial cells (*NP2222-GAL4>*).
1090 D-G) Representative images showing that knockdown of Fasn1 in glial cells (*repo-*
1091 *GAL4>*), where Hh.N.EGFP is overexpressed, significantly reduces the number of
1092 Hh.N.EGFP puncta on the surface of NBs (yellow arrows), phenocopying the effect
1093 of Rasp knockdown, quantified in (G) (n=38, 38, 57 NBs from 4, 4, 8 brain lobes,
1094 respectively).
1095 H) Cortex glial (*NP2222-GAL4>*) overexpression of Hh.N.EGFP significantly
1096 reduces NB EdU incorporation (n= 13,12 brain lobes).
1097 I) Knockdown of Fasn1 in glial cells rescues NB EdU incorporation defects, caused
1098 by glial Hh.N.EGFP overexpression, quantified in (I) (n=15, 13 brain lobes).
1099 J) Schematic depicting our working model. During development, Hh tethered to LDs
1100 are localised to cortex glial cells, to activate gliogenesis. Non-autonomously,
1101 excessive glial Hh inhibits NB cell cycle progression. Upon cortex glial specific FGF
1102 activation, increased Hh modified by Fasn1 and Lsd-2 together with increased ROS
1103 prevent NB proliferation.
1104 Data information: (C', D', E' and F') are zoomed in images of (C, D, E and F),
1105 respectively; NBs are marked with Mira. Scale bar=50 μ m in (C, D, E, F). Scale
1106 bar=20 μ m in (C', D', E' and F'). Error bar represents SEM. In (B): unpaired t test,
1107 (ns) $p=0.9403$; unpaired t test, (****) $p<0.0001$; Mann-Whitney test, (***)
1108 $p=0.0004$; Mann-Whitney test, (**) $p=0.0047$. In (G): Welch's t test, (****)
1109 $p<0.0001$; Welch's t test, (****) $p<0.0001$. In (H): unpaired t test, (****) $p<0.0001$.
1110 In (I): unpaired t test, (****) $p<0.0001$.

Commented [HC10]: Please also change the label in the figure to G.

Commented [HC11]: Please check.

1111
1112 **Expanded View Figure legends**
1113 **Figure EV1: Hh forms complexes with LDs in the CB but not the wing discs**
1114 **(related Fig 1)**
1115 A-B'') HhGFP and LDs are associated in the CB glial cells (outlined with yellow
1116 dashed lines).

1117 (C-D'') HhGFP is not associated with LDs in the posterior wing disc (white arrows,
1118 the posterior compartment is separated from the anterior with yellow dashed lines).
1119 Data information: (B-B''), D-D'') are zoomed in images of (A and C), respectively.
1120 Scale bar=50 μm in (A and C). Scale bar=10 μm in (B-B'' and D-D'').

Commented [HC12]: Please define the yellow dashed line.

1121

1122 **Figure EV2: Hh-LD associations are specifically observed in cortex glial cells in**
1123 **the CB during late larval stages (related to Fig 1)**

1124 (A-B'') Hh and LDs are present at low levels at 48 ALH, and do not form specific
1125 association (white arrows, brain lobes are circled with purple dashed lines).

Commented [HC13]: Please define the purple dashed circle.

1126 (C-D'') Hh and LDs are not associated in the optic lobe glial cells (white arrows).

1127 (E-F'') Hh and LDs associate only in the cortex glial cells (yellow arrows) but not the
1128 surface glial cells (white arrows, surface glia are separated from cortex with magenta
1129 dashed lines).

Commented [HC14]: Please define the red dashed lines in F-F''.

1130 (G) Left and middle panel, representative image showing a surface glial clone (circled
1131 with white dashed lines, *repo*-MARCM, glial nucleus marked by Stinger in pink). Hh
1132 is localised to the cortex glial cells (yellow arrows) underneath the clone marked in
1133 green.

Commented [HC15]: Please define the white dashed line.

1134 Right panel, a schematic depicting XZ cross-section of CB glial cells and their
1135 relative position.

1136 Data information: Glial cells are visualised with *repo-GAL4>GFP* in (C-F''). (B-B'',
1137 D-D'', F-F'') are zoomed in images of (A, C, E), respectively. (C) is the same
1138 image as Fig 1C, with the optic lobe region highlighted in a yellow dashed square.

1139 Scale bar=50 μm in (C, E) and XY section in (G). Scale bar=20 μm in (A-B'', D-
1140 D'', F-F''), Scale bar=10 μm for XZ section in (G).

1141

1142 **Figure EV3: Effects of Hh overexpression and knockdown on Hh level, NB**
1143 **number and EdU index (related to Fig 2)**

1144 (A-D) Representative images showing pan-glial Hh knockdown (*repo-GAL4>GFP*
1145 with *UAS-dcr2*) efficiently reduces Hh staining in the CB (outlined with yellow
1146 dashed lines) and brain lobe size, quantified in (C) (n=6, 6 brain lobes) and (D) (n=15,
1147 16 brain lobes), respectively.

1148 (E) Representative image from EdU incorporation assays used throughout the
1149 manuscript. During a 15 min EdU pulse, type I NB (yellow arrow) and its GMC

1150 (white arrow) both incorporate EdU. EdU index quantifications include only EdU⁺
1151 type I NBs.
1152 F) Hh knockdown (*repo-GAL4>GFP* with *UAS-dcr2*) significantly reduces NB EdU
1153 index (n=12, 9 brain lobes).
1154 G) Hh knockdown or overexpression in glial cells (*repo-Gal4>*) does not
1155 significantly alter the number of CB NBs (n=12, 10; 12, 12 brain lobes).
1156 H) Schematic depicting EdU pulse-chase experiment. Larvae are fed with EdU-
1157 containing food for 3 hours and then chased with EdU-free food for 3 hours before
1158 CNS dissection at wandering stages. NBs and newly generated GMCs are marked
1159 with Mira; GMCs and newly generated neurons are marked with ProsGFP.
1160 I-K) Representative images showing that Hh knockdown in cortex glial cells
1161 (*NP2222-GAL4*) significantly reduced the number of EdU⁺ cells that are marked by
1162 ProsGFP⁺, quantified in (K) (Box plot, the boxes extend from the 25th to 75th
1163 percentiles; the median is marked by a central band inside the box; and the whiskers
1164 go down to the minimum value and up to the maximum value. n=94, 104 NB lineages
1165 imaged from 8, 8 brain lobes, respectively).
1166 L-O) Representative images showing that Hh overexpression in cortex glial cells
1167 (*NP2222-GAL4>mGFP*) does not alter cortex glial membrane size and total Repo⁺
1168 glial cell numbers, quantified in (N) (n=15, 14 brain lobes) and (O) (n=15, 14 brain
1169 lobes), respectively.
1170 P-Q'') Representative images showing Ptc:mcherry is expressed in NBs (yellow
1171 arrows). (Q-Q'') are zoomed in images of (P).
1172 Data information: Scale bar=50 μm in (A, B, E, L, M). Scale bar=20 μm in (I, J, P, Q-
1173 Q''). Error bar represents SEM. In (C): unpaired t test, (****) $p < 0.0001$. In (D):
1174 Mann-Whitney test, (**) $p = 0.0017$. In (F): Welch's t test, (*) $p = 0.0101$. In (G):
1175 unpaired t test, (ns) $p = 0.3645$; Mann-Whitney test, (ns) $p = 0.7621$. In (K): Welch's t
1176 test, (****) $p < 0.0001$. In (N): unpaired t test, (ns) $p = 0.5151$. In (O): unpaired test,
1177 (ns) $p = 0.9690$.

1179 **Figure EV4 Glial *hhl^{ACT}* overexpression affects NB asymmetric division, size and**
1180 **cell cycle exit (related to Fig 5)**

1181 A) Schematic depicting the distribution of polarity proteins in M phase NBs. Apical
1182 polarity proteins include the Par complex (aPKC/Par3/Par6), the PINS/MUD/Gai

Commented [HC16]: Please define the central band, boxes and whiskers of the boxplot.

1183 complex and the adaptor protein, Inscuteable (magenta); Basal polarity complex
1184 comprises the cell fate determinants Brat/ Pros/ Numb and their adaptor proteins Mira
1185 (red) and PON.

1186 B-G) Representative images showing that in pH3⁺ NBs, Insc and Mira mislocalize to
1187 the cytoplasm or cortex upon FGF activation in cortex glia (*NP2222-GAL4> htl^{ACT}*).

Commented [HC17]: 'NP2222-GAL4>' is labelled in the panel. Please check and adjust accordingly.

1188 H) Schematic depicting a NB undergoing telophase.

1189 I-K) Representative images showing that NBs in telophase (Mira⁺; pH3⁺) give rise to
1190 more size-symmetric daughter cells upon cortex glial (*NP2222-GAL4> htl^{ACT}*
1191 overexpression, quantified in (K) (n=43, 63 NBs from 10, 9 brain lobes, respectively).

1192 L) Glial (*repo-GAL4> htl^{ACT}* overexpression causes an increase in M phase NB
1193 diameter (n=70, 53 NBs from 12, 7 brain lobes, respectively).

1194 M-O) Representative images showing that NB nucleoli are significantly enlarged
1195 upon glial (*repo-GAL4> htl^{ACT}* overexpression, quantified in (O) (n=33, 23 NBs
1196 from 9, 7 brain lobes). NBs are marked by Histone (red), surrounded by glial cells
1197 (grey, *repo-GAL4>GFP*), nucleoli are marked by Fib (Cyan).

1198 P-S) Representative images showing that the timing of NB cell cycle entry (visualised
1199 by EdU incorporation at 26ALH) is not significantly altered by pan-glial (*repo-*
1200 *GAL4> htl^{ACT}* overexpression, quantified in (S), where EdU voxels are normalized to
1201 control (n=19, 25 brains). The region of interest is outlined in yellow.

1202 T-W) Representative images showing that the number of CB NBs (Mira⁺) at 24APF is
1203 significantly increased with pan-glial (*repo-GAL4> htl^{ACT}* overexpression, quantified
1204 in (W) (n=8, 8 brains). The region of interest is outlined by yellow dashed lines and
1205 NBs are marked with yellow arrows.

1206 Data information: NBs are outlined with yellow dashed lines in (B-G, I and J). Scale
1207 bar=50 μm in (Q, R, U, V). Scale bar=10 μm in (B-G); Scale bar=20 μm in (I, J, M

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1208 and N). Error bar represents SEM. In (K): Man-Whitney test, (***) $p=0.0002$. In (L):
1209 Welch's t test, (***) $p<0.0001$. In (O): unpaired t test, (***) $p<0.0001$. In (S):

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1210 Welch's t test, (ns) $p=0.8152$. In (W): unpaired t test, (*) $p=0.0134$.

1211

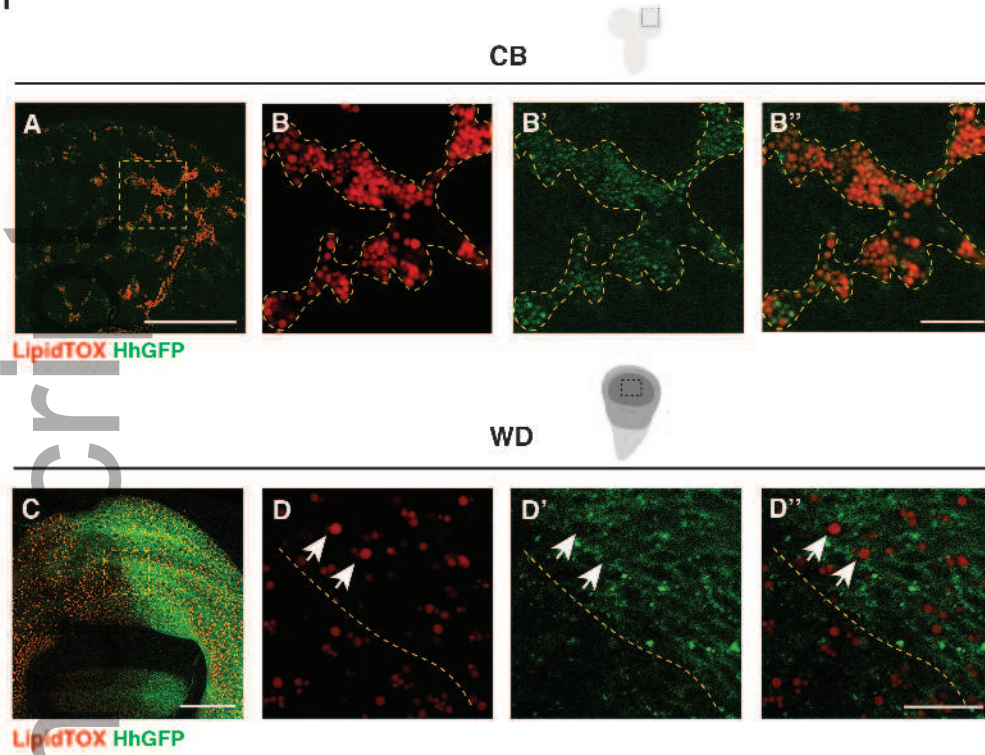
1212 **Figure EV5 Characterisation of the effects of glial *hh*, *fasn1* and *lsd2* RNAis on**
1213 **glial size, LDs and NB proliferation (related to Fig 6)**

1214 A-D) Representative images showing that induction of *hh RNAi* in cortex glial cells
1215 with *htl^{ACT}* overexpression does not alter the size of cortex glial membrane (*NP2222-*
1216 *GAL4>mGFP*) nor the number of LDs in CB (outlined by yellow dashed lines).
1217 E-G) Representative images showing that glial (*repo-GAL4>*) induction of RNAis
1218 against *fasn1* and *lsd2* efficiently reduce the number of LDs in CB (outlined by
1219 yellow dashed lines).
1220 H) knockdown of lipogenesis genes *fasn1* and *lsd2* or overexpression of *lsd2* using a
1221 cortex glial driver (*NP2222-GAL4>*) does not significantly affect NB EdU index
1222 (n=10, 10; 15, 10; 10, 10 brain lobes).
1223 I) The NB EdU incorporation defects due to cortex glial (*NP2222-GAL4*)
1224 overexpression of *htl^{ACT}* is rescued by overexpression of additional RNAi lines
1225 against *fasn1* and *lsd2* (related to Fig 6J; n=10, 16; 14, 14; 8, 10 brain lobes). The
1226 *NP2222-GAL4> w¹¹¹⁸* vs *htl^{ACT}* columns depict the same data as those in Fig 5E.
1227 Data information: Scale bar=50 μ m in (A-G). Error bar represents SEM. In (H):
1228 Mann-Whitney test, (ns) $p=0.9555$; unpaired t test, (ns) $p=0.1799$; unpaired t test,
1229 (ns) $p=0.9574$. In (I): unpaired t test, (****) $p<0.0001$; unpaired t test, (***)
1230 $p=0.0008$; unpaired t test, (**) $p=0.0032$.

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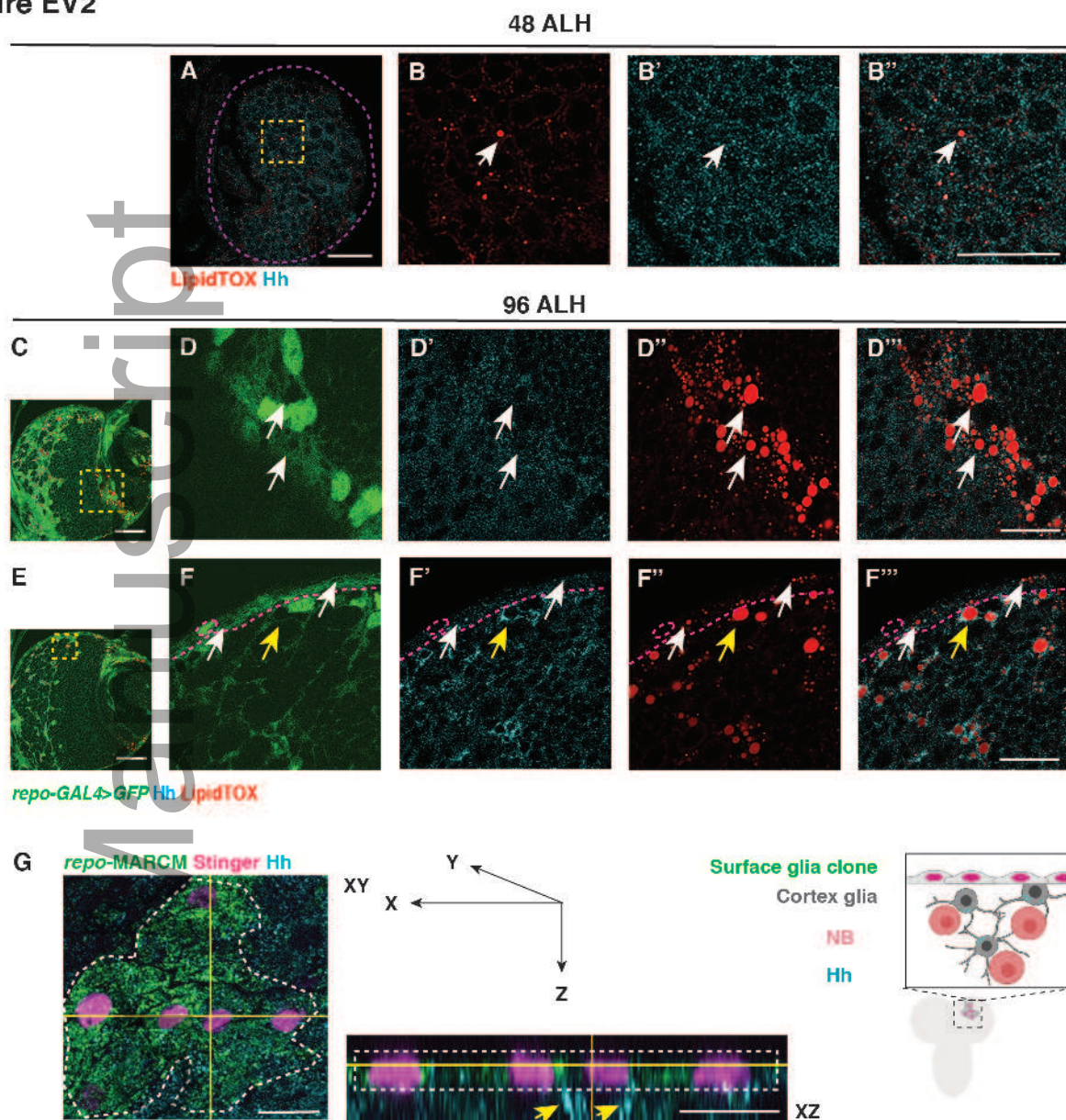
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Figure EV1



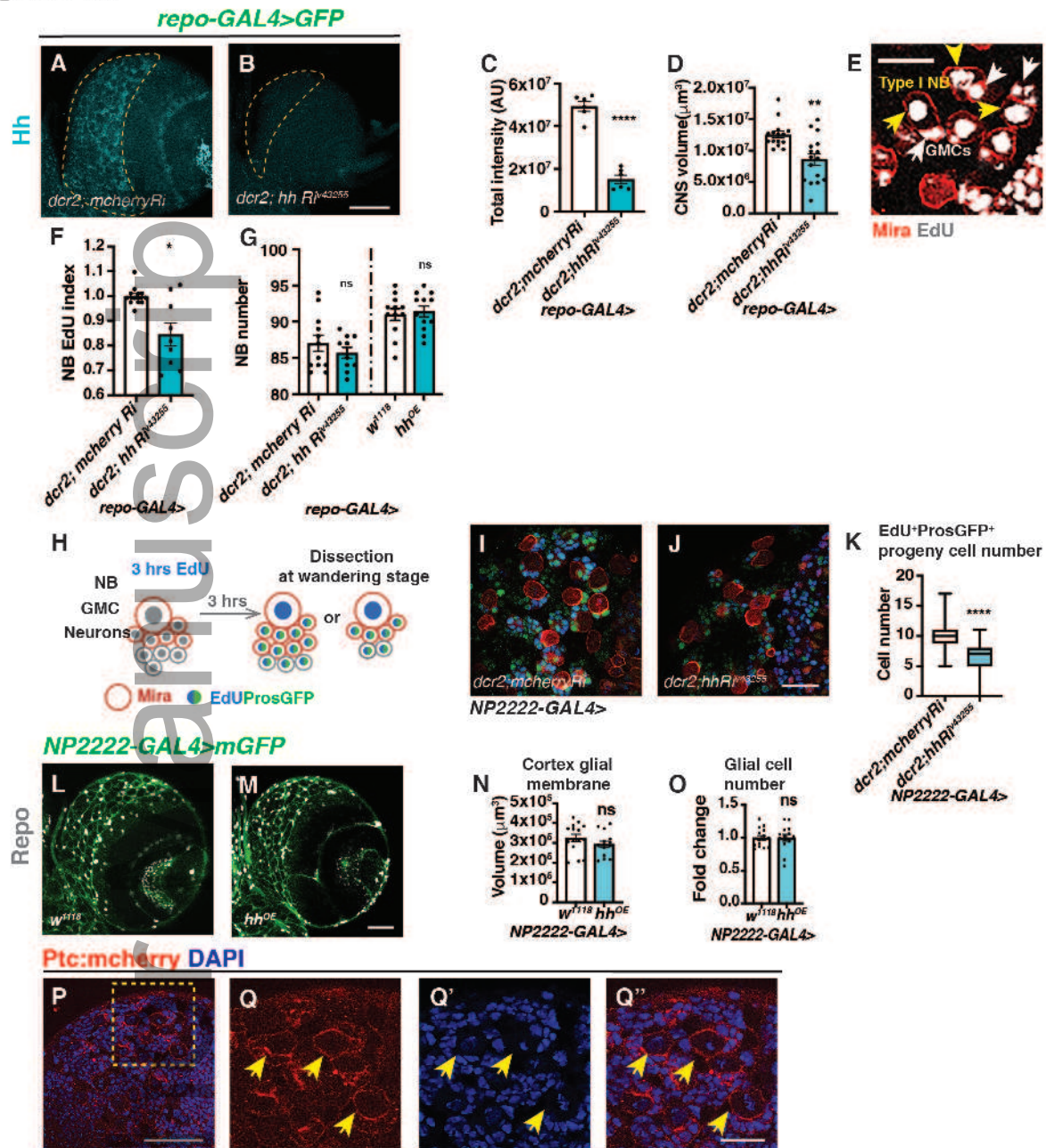
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Figure EV2



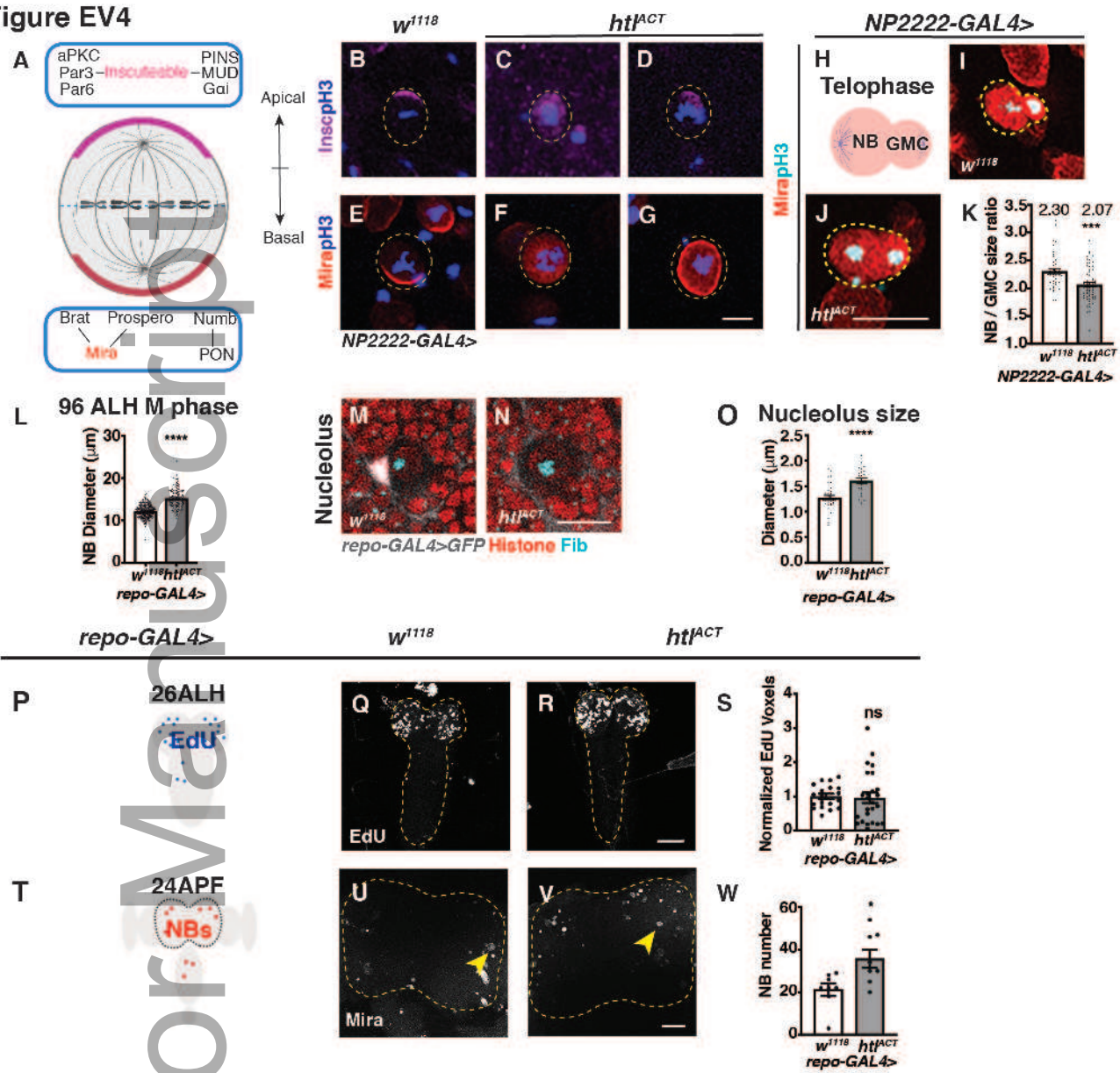
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Figure EV3



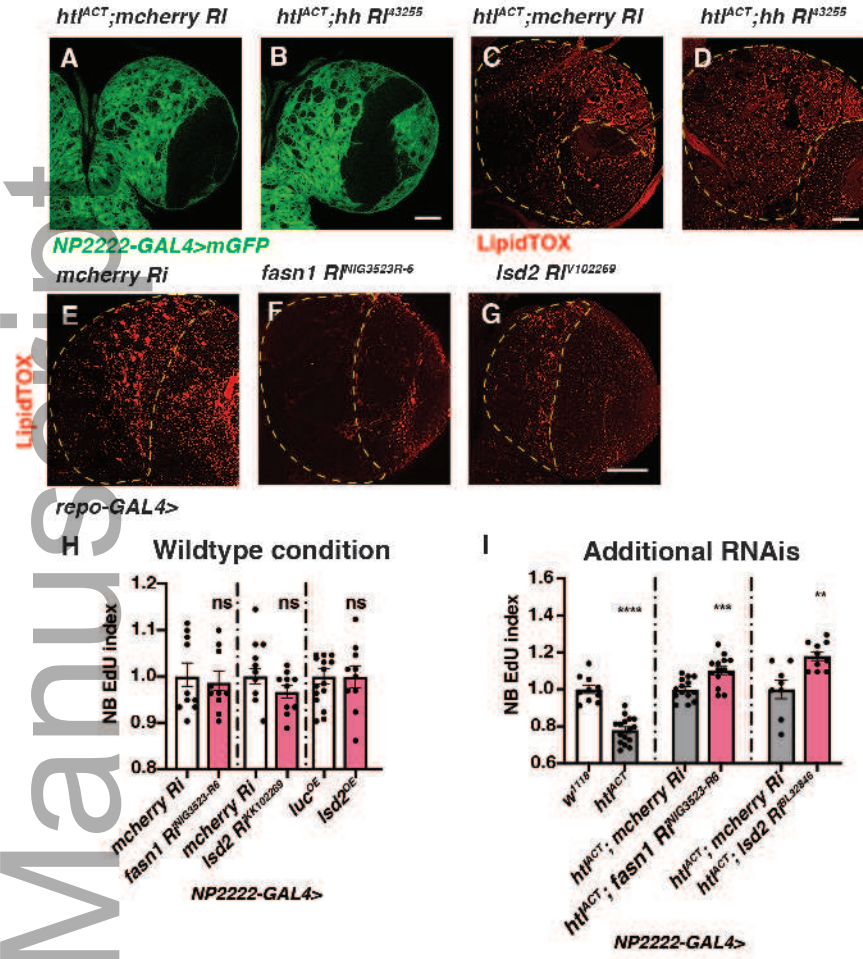
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Figure EV4



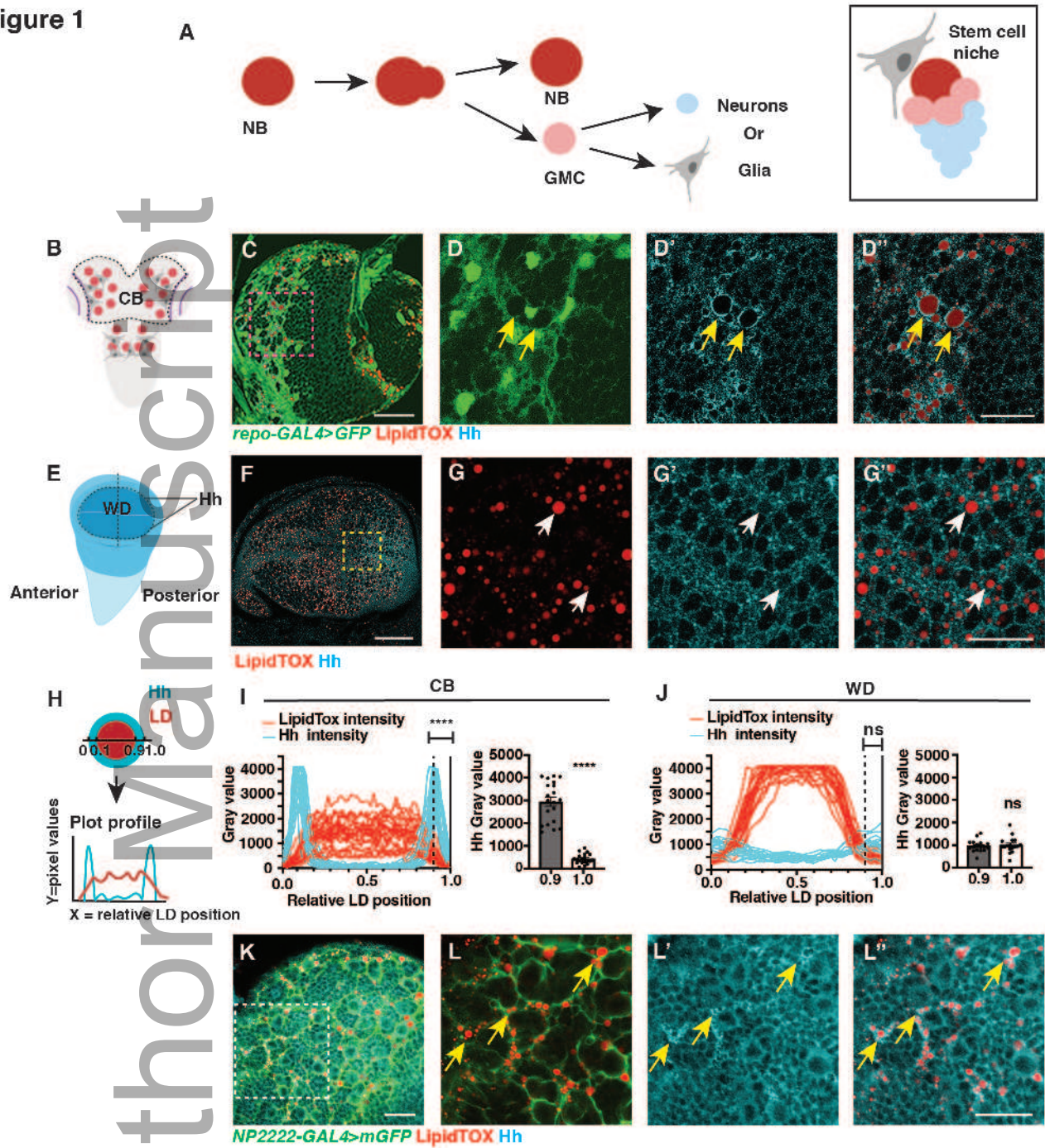
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Figure EV5



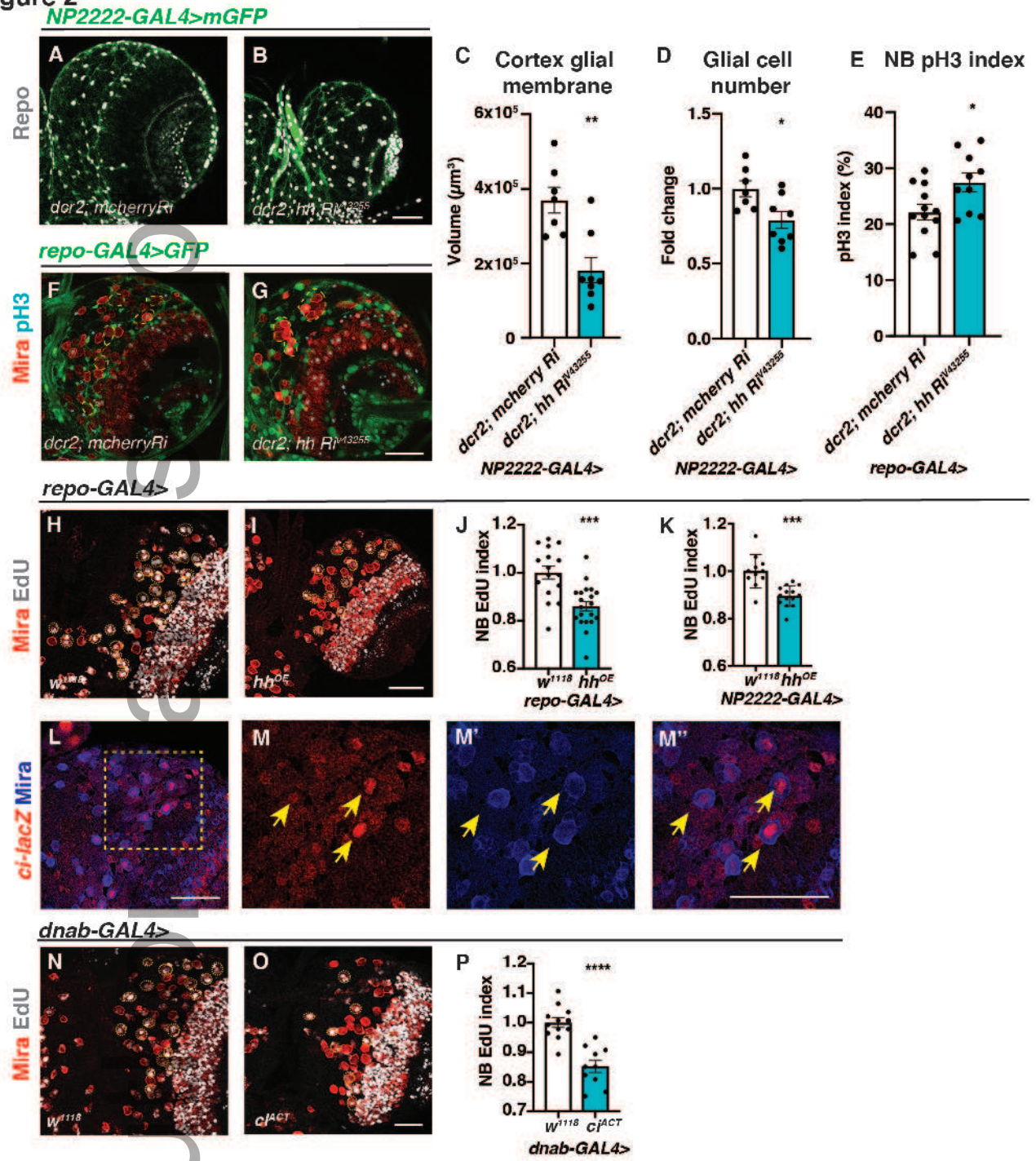
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Figure 1



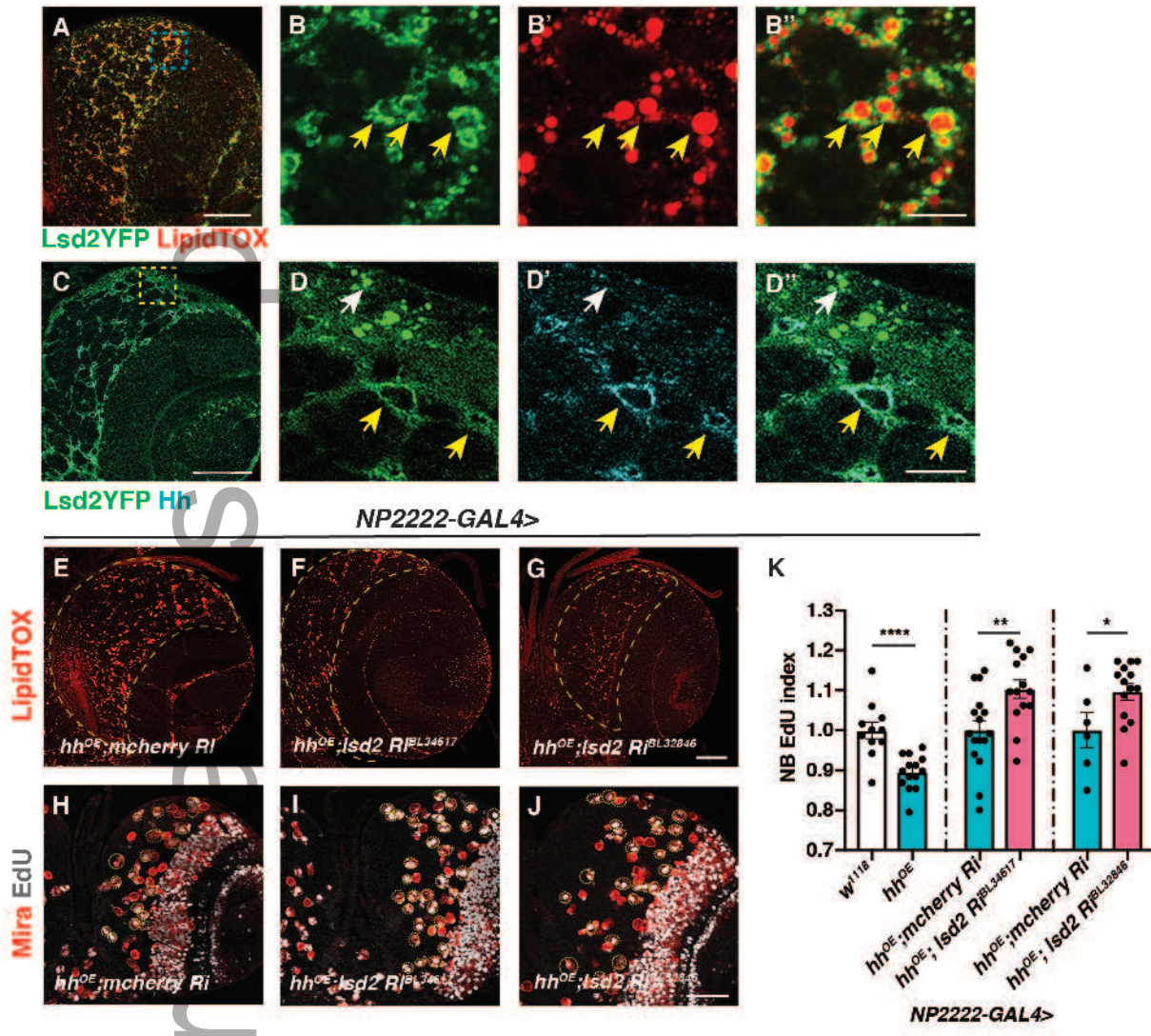
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Figure 2



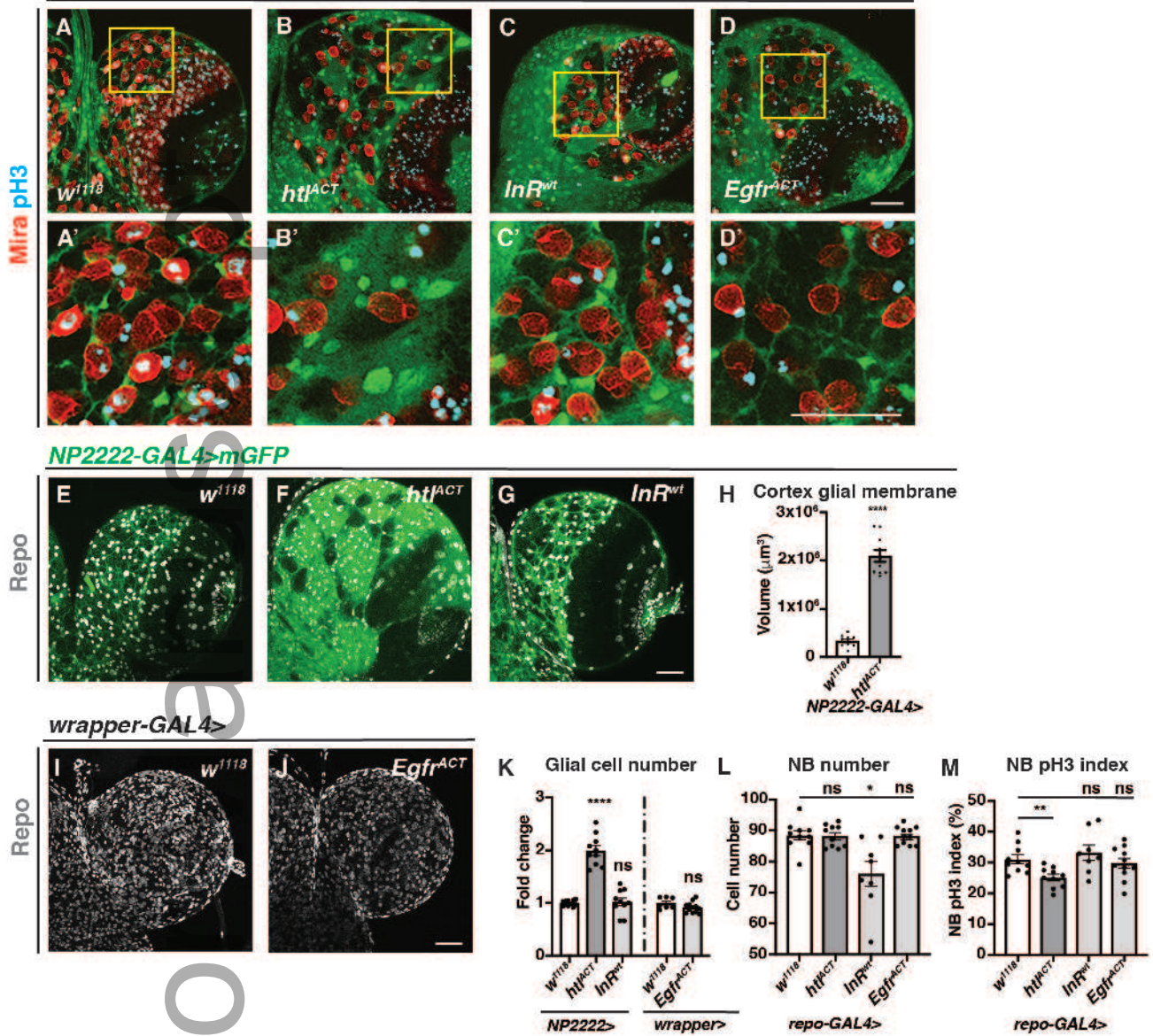
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Figure 3



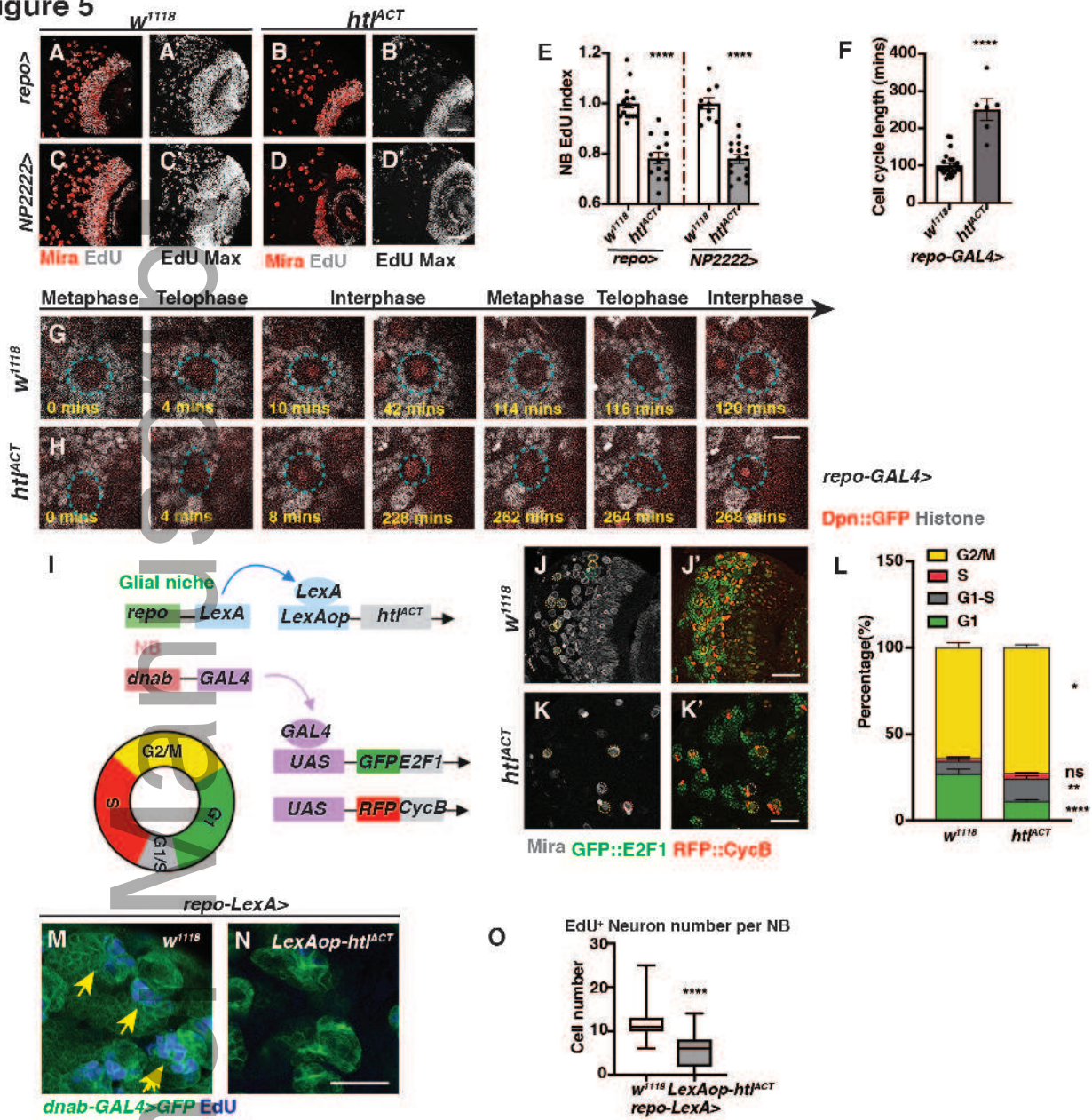
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Figure 4
repo-GAL4>GFP



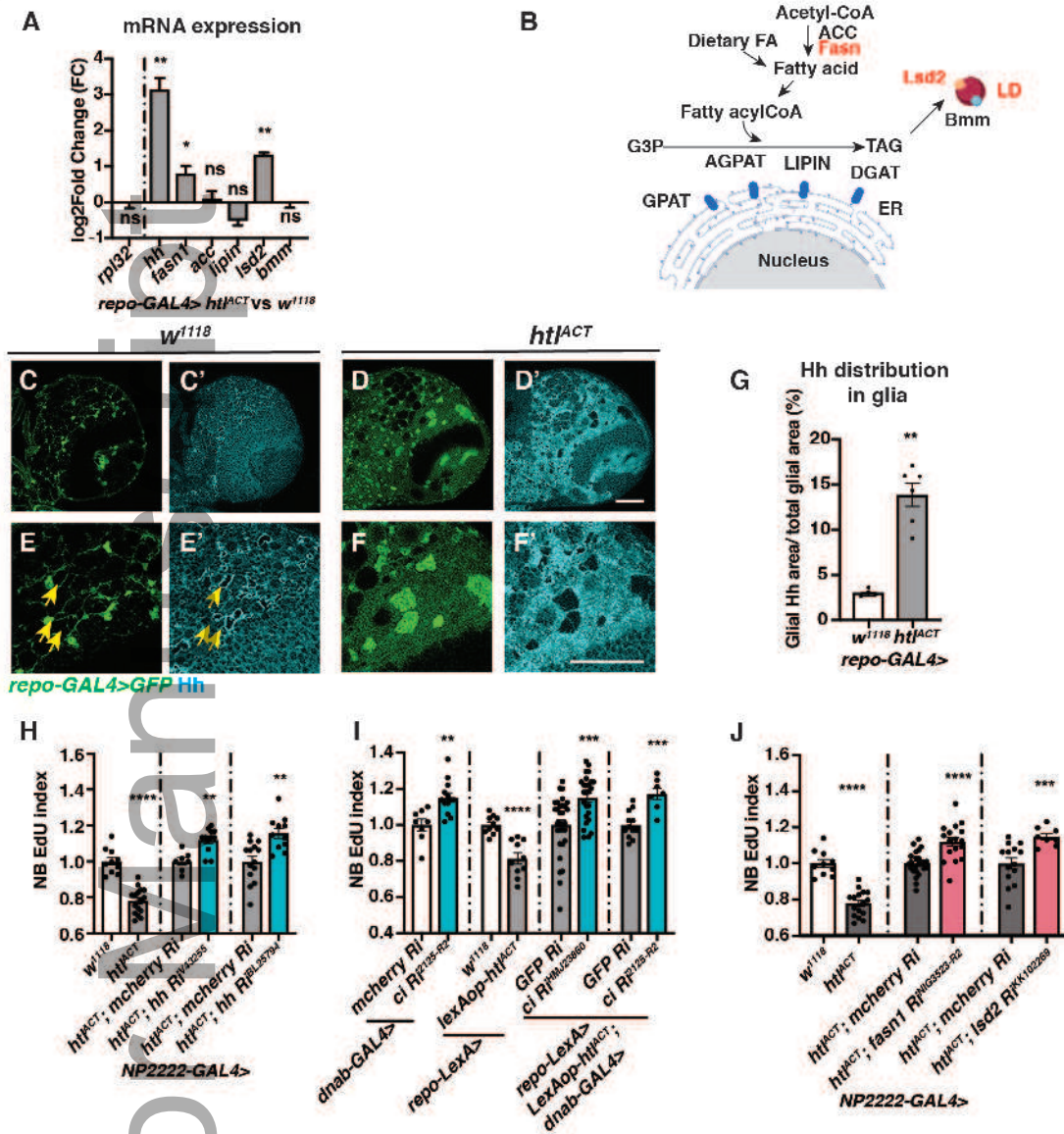
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Figure 5



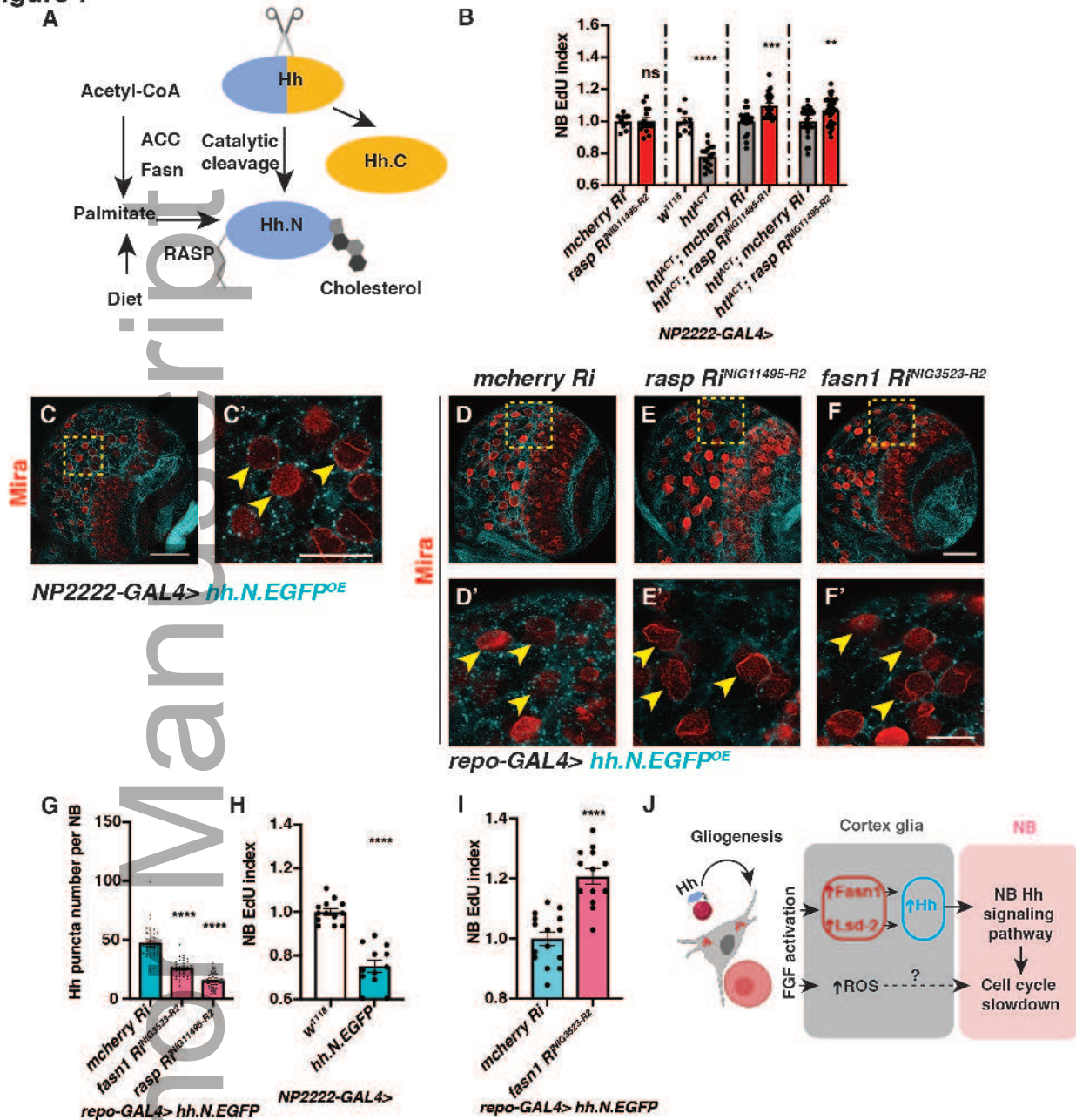
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Figure 6



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Figure 7



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