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1 2 3	Homeostatic and tumourigenic activity of SOX2+ pituitary stem cells is controlled by the LATS/YAP/TAZ cascade				
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# 29 ABSTRACT

31	SOX2 positive pituitary stem cells (PSCs) are specified embryonically and persist
32	throughout life, giving rise to all pituitary endocrine lineages. We have previously
33	shown the activation of the STK/LATS/YAP/TAZ signalling cascade in the
34	developing and postnatal mammalian pituitary. Here, we investigate the function of
35	this pathway during pituitary development and in the regulation of the SOX2 cell
36	compartment. Through loss- and gain-of-function genetic approaches, we reveal that
37	restricting YAP/TAZ activation during development is essential for normal organ size
38	and specification from SOX2+ PSCs. Postnatal deletion of LATS kinases and
39	subsequent upregulation of YAP/TAZ leads to uncontrolled clonal expansion of the
40	SOX2+ PSCs and disruption of their differentiation, causing the formation of non-
41	secreting, aggressive pituitary tumours. In contrast, sustained expression of YAP
42	alone results in expansion of SOX2+ PSCs capable of differentiation and devoid of
43	tumourigenic potential. Our findings identify the LATS/YAP/TAZ signalling cascade
44	as an essential component of PSC regulation in normal pituitary physiology and
45	tumourigenesis.
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51	Key words: pituitary stem cell, SOX2, Hippo, YAP, pituitary tumour
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# **INTRODUCTION**

SOX2 is crucial transcription factor involved in the specification and maintenance of multiple stem cell populations in mammals. Pituitary stem cells express SOX2 and contribute to the generation of new endocrine cells during embryonic development and throughout postnatal life<sup>1, 2</sup>. The pituitary gland is composed of three parts, the anterior, intermediate and posterior lobes (AL, IL and PL, respectively). The AL and IL contain hormone-secreting cells, which are derived from an evagination of the oral ectoderm expressing SOX2, termed Rathke's pouch (RP). SOX2+ cells, both in the embryonic and adult pituitary, can differentiate into three endocrine cell lineages, which are marked by transcription factors PIT1 (POU1F1)<sup>3</sup>, TPIT (TBX19)<sup>4</sup> and SF1 (NR5A1)<sup>5</sup>, and differentiate into hormone-secreting cells (somatotrophs, lactotrophs, thyrotrophs, corticotrophs, melanotrophs and gonadotrophs, which express growth hormone, prolactin, thyrotropin, adrenocorticotropin, melanotropin and gonadotropin, respectively). SOX2+ PSCs are highly proliferative during the first 2-3 weeks of life, in concordance with major organ growth, after which they reach a steady low proliferative capacity that contributes to maintain normal homeostasis and physiological adaptation of the pituitary gland<sup>6,7</sup>. 

Contrary to other organs, where somatic stem cells are shown to be able to become 78 79 transformed into cancer stem cells, the roles of SOX2+ PSCs in tumourigenesis 80 remain poorly understood, possibly due to the patchy knowledge of the pathways 81 regulating SOX2+ PSC fate and proliferation. Pituitary tumours are common in the population, representing 10-15% of all intracranial neoplasms<sup>8,9</sup>. Adenomas are the 82 83 most common adult pituitary tumours, classified into functioning, when they secrete 84 one or more of the pituitary hormones, or non-functioning if they do not secrete 85 hormones. In children, adamantinomatous craniopharyngioma (ACP) is the most 86 common pituitary tumour. Targeting oncogenic beta-catenin in SOX2+ PSCs in the 87 mouse generates clusters of senescent SOX2+ cells that induce tumours resembling 88 ACP in a paracrine manner, i.e. the tumours do not derive from the targeted SOX2+ PSCs<sup>1, 10</sup>. Up to 15% of adenomas and 50% of ACP display aggressive behaviour 89 90 with invasion of nearby structures including the hypothalamus and visual tracts, associated with significant morbidity and mortality<sup>11</sup>. Pituitary carcinomas exhibiting 91 metastasis are rare but can develop from benign tumours<sup>12, 13, 14</sup>. Whether SOX2+ 92 cells can cell autonomously contribute to pituitary neoplasia has not been hitherto 93 94 demonstrated.

95

The Hippo pathway controls stem cell proliferation and tumourigenesis in several
organs such as in the liver<sup>15, 16</sup>, intestines<sup>17</sup> and lung<sup>18, 19</sup>. In the core phosphorylation
cascade, STK3/4 kinases phosphorylate and activate LATS1/2 serine/threonineprotein kinases, which in turn phosphorylate co-activators Yes-associated protein
(YAP1, a.k.a. YAP) and WW domain-containing transcription regulator protein 1
(WWTR1, a.k.a. TAZ) that are subsequently inactivated through degradation and
cytoplasmic retention<sup>20</sup>. Active YAP/TAZ associate with TEAD transcription factors,

103	promoting the transcription of target genes such as <i>Cyr61</i> and <i>Ctgf</i> <sup>21, 22, 23</sup> . YAP/TAZ
104	have been shown to promote proliferation and the stem cell state in several organs,
105	and can also lead to transformation and tumour initiation when overexpressed <sup>24, 25, 26</sup> .
106	The involvement of YAP/TAZ in the function of tissue-specific SOX2+ stem cells
107	during development and homeostasis has not been shown. We previously reported
108	strong nuclear localisation of YAP and TAZ exclusively in SOX2+ stem cells of
109	developing Rathke's pouch and the postnatal anterior pituitary of mice and humans,
110	and enhanced expression in human pituitary tumours composed of uncommitted cells,
111	including ACPs and null-cell adenomas <sup>27, 28</sup> , which do not express any of the lineage
112	transcription factors PIT1, TPIT or SF1. In these populations we detected
113	phosphorylation of YAP at serine 127 (S127) indicating LATS kinase activity.
114	Together these point to a possible function for LATS/YAP/TAZ in normal pituitary
115	stem cells and during tumourigenesis. Here, we have combined genetic and molecular
116	approaches to reveal that deregulation of the pathway can promote and maintain the
117	SOX2+ PSC fate under physiological conditions and that major disruption of this axis
118	transforms SOX2+ PSCs into cancer-initiating cells giving rise to aggressive tumours.
119	
120	RESULTS
121	
122	Sustained conditional expression of YAP during development promotes SOX2+
123	PSC fate
124	To determine if YAP and TAZ function during embryonic development of the
125	pituitary, we used genetic approaches to perform gain- and loss-of-function
126	experiments. We first expressed a constitutive active form of YAP(S127A) using the
127	Hesx1-Cre driver, which drives Cre expression in Rathke's pouch (RP) and the

128	hypothalamic primordium from 9.5dpc, regulated by administration of doxycycline
129	through the reverse tetracycline-dependent transactivator (rtTA) system ( $R26^{rtTA/+}$ ; see
130	Methods for details, Scheme Fig1A). Analyses were restricted to embryonic time
131	points. As expected, we confirmed accumulation of total YAP protein but not of TAZ
132	or pYAP(S127), throughout the developing pituitary and hypothalamus of
133	<i>Hesx1</i> <sup>Cre/+</sup> ; <i>R26</i> <sup>rtTA/+</sup> ; <i>Col1a1</i> <sup>tetO-Yap/+</sup> (hereafter YAP-TetO) embryos at 15.5dpc, but
134	not of Cre-negative controls (Fig1B, Figure 1 – figure supplement 1A). Likewise, the
135	YAP downstream target Cyr61 (Fig1B) was also upregulated. Morphologically, YAP-
136	TetO mutants displayed a dysplastic anterior pituitary, which was more medially
137	compacted and lacked a central lumen, making it difficult to distinguish between the
138	developing anterior and intermediate lobes (Fig 1C). Immunofluorescence staining
139	against SOX2 at 15.5dpc demonstrated loss of SOX2 in the most lateral regions of
140	control pituitaries (arrows in Fig. 1C), where cells are undergoing commitment; yet
141	mutant pituitaries had abundant SOX2 positive cells in the most lateral regions
142	(arrowheads in Fig1C). Immunostaining for LHX3, which is expressed in the
143	developing anterior pituitary <sup>29</sup> , was used to demarcate AL and IL tissue. Staining
144	using antibodies against lineage markers PIT1, TPIT and SF1 revealed a concomitant
145	reduction in committed cell lineages throughout the gland (Fig1D; PIT1 0.35% in
146	mutants compared with 30.21% in controls (Student's t-test $P < 0.0001$ , n=3 for each
147	genotype), TPIT 1.03% in mutants compared with 9.81% in controls (Student's t-test
148	P=0.0012, n=3 for each genotype), SF1 0.34% in mutants compared with 4.14% in
149	controls (Student's t-test $P=0.0021$ , n=3 for each genotype)). We therefore conclude
150	that sustained activation of YAP prevents lineage commitment and is sufficient to
151	maintain the progenitor state during embryonic development.

153	We did not obtain any live $Hesx1^{Cre/+}$ ; $R26^{rtTA/+}$ ; $Col1a1^{tetO-Yap/+}$ pups at birth when
154	treated with doxycycline from 5.5dpc (n=5 litters). To bypass the embryonic lethality
155	of these early inductions, we commenced doxycycline treatment from 9.5dpc, the
156	onset of RP formation (Figure 1 – figure supplement 1B).
157	$Hesx1^{Cre/+}$ ; $R26^{rtTA/+}$ ; $Colla1^{tetO-Yap/+}$ pups were viable and were maintained on
158	doxycycline until P24, at which point the experimental end point was reached due
159	to excessive weight loss and animals had to be culled following UK Home Office
160	Regulations. Histological analyses of pituitaries revealed multiple anterior lobe cysts
161	per gland, localising predominantly in the ventral AL (n=4) (Figure 1 – figure
162	supplement 1C). These structures developed in YAP-accumulating regions and were
163	lined by SOX2+ cells (Figure 1 – figure supplement 1D). The proportion of SOX2+
164	cells throughout the AL was increased, as was the percentage of SF1+ cells, whereas
165	PIT1+ cell numbers were significantly decreased and the TPIT lineage, identified by
166	ACTH antibody staining, was unaffected (Figure 1 – figure supplement 1E). The total
167	number of cycling Ki-67+ cells showed a trend towards a decrease
168	in $Hesx1^{Cre/+}$ ; $R26^{rtTA/+}$ ; $Colla1^{tetO-Yap/+}$ mutants relative to controls, which did not
169	reach significance (Figure 1 – figure supplement 1F). The cystic structures observed
170	in Hesx1 <sup>Cre/+</sup> ;R26 <sup>rtTA/+</sup> ;Col1a1 <sup>tetO-Yap/+</sup> mutants were reminiscent of Rathke's cleft
171	cyst (RCC), which is a benign developmental anomaly of the pituitary characterised
172	by the presence of ciliated and secretory cells, expression cytokeratins and frequent
173	expression of p63. Immunostaining revealed that cysts were lined by cytokeratin+
174	cells using the AE1/AE3 pan-cytokeratin cocktail and basal cells were positive for
175	nuclear p63 in $Hesx1^{Cre/+}$ ; $R26^{rtTA/+}$ ; $Col1a1^{tetO-Yap/+}$ mutant pituitaries (Figure 1 –
176	figure supplement 1G). Staining using antibodies against ARL13B and Acetylated $\alpha$ -
177	Tubulin (Lys40) marking cilia, revealed multi-ciliated cells along the cyst lining

(Figure 1 – figure supplement 1H). Combined staining using Alcian Blue and the
Periodic Acid-Schiff technique (AB/PAS) to recognise mucins, detected royal bluestained mucous cells lining the cysts (Figure 1 – figure supplement 1H). Taken
together, we conclude that sustained activation of YAP during embryonic and
postnatal pituitary development, promotes maintenance and abnormal expansion of
SOX2+ epithelia during development, resulting in the formation of cysts that
resemble RCC.

185

186 Next, we generated embryos null for TAZ and conditionally lacking YAP in the

187 *Hess1* expression domain (Figure 1 – figure supplement 2A-E).

188  $Hesx1^{Cre/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{-l-}$  double mutants were obtained at expected ratios during 189 embryonic stages until 15.5dpc, however the majority of  $Taz^{-/-}$  mutants with or

190 without compound *Yap* deletions showed lethality at later embryonic and early

191 postnatal stages<sup>30</sup> (Supplementary File 1). The developing pituitary gland of

192  $Hesx1^{Cre/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{-l-}$  double mutants appeared largely normal at 13.5 dpc by

193 histology (Figure 1 – figure supplement 2A). Immunostaining against SOX2 to mark

194 embryonic progenitors and postnatal stem cells did not reveal differences in the

195 spatial distribution of SOX2+ cells between double mutants compared to controls

196 ( $Hesx1^{+/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{+/+}$  and  $Hesx1^{+/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{+/-}$ ) at 13.5 dpc, 16.0 dpc (Figure 1 –

197 figure supplement 2B) or P28, even in regions devoid of both TAZ and active YAP

198 (Figure 1 – figure supplement 2C,D). This suggests that YAP/TAZ are not required

199 for SOX2+ cell specification or survival. Likewise, analysis of commitment markers

- 200 PIT1and SF1 as well as ACTH to identify the TPIT lineage, did not show any
- 201 differences between genotypes (Figure 1 figure supplement 2E). Together, these
- 202 data suggest there is no critical requirement for YAP and TAZ during development

for the specification of SOX2+ cells or lineage commitment, but that YAP functions
to promote the SOX2 cell identity.

205

# LATS, but not STK, kinases are required for normal pituitary development and differentiation

208 Since sustained activation of YAP led to an embryonic phenotype, we reasoned that

209 YAP/TAZ need to be regulated during embryonic development. To determine if STK

210 and LATS kinases are important in YAP/TAZ regulation we carried out genetic

211 deletions in the pituitary.

212

213 Conditional deletion of *Stk3* and *Stk4* (also called *Mst2* and *Mst1*) in

214  $Hesx1^{Cre/+}$ ; Stk3<sup>fl/fl</sup>; Stk4<sup>fl/fl</sup> embryos did not lead to a pituitary phenotype (Figure 2 –

215 figure supplement 1). A reduction of over 75% in total STK3/4 proteins in mutants

216 was confirmed by western blot on total lysates from  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl}$  controls and

217  $Hesx1^{Cre/+}$ ; Stk3<sup>fl/fl</sup>; Stk4<sup>fl/fl</sup> mutants (Figure 2 – figure supplement 1B). Mutant

218 pituitaries were macroscopically normal at birth (Figure 2 – figure supplement 1A),

and showed comparable expression patterns of TAZ, YAP, pYAP to controls lacking

220 *Cre*, without distinct accumulation of YAP or TAZ (Figure 2 – figure supplement

221 1C). The distribution of SOX2+ cells was comparable between mutants and controls

222 (Figure 2 – figure supplement 1C). Normal lineage commitment was evident by

immunofluorescence staining for PIT1, TPIT and SF1 at P10 (Figure 2 – figure

supplement 1D). Mutant animals remained healthy and fertile until P70, at which

225 point pituitaries appeared histologically normal (Figure 2 – figure supplement 1E).

226 Since deletion of *Stk3/4* at embryonic stages does not affect embryonic or postnatal

227 pituitary development, we conclude these kinases are not critical for YAP/TAZ

regulation in the pituitary.

229

230	We next focused on perturbing LATS kinase function, as we have previously shown			
231	strong expression of Lats1 in the developing pituitary and postnatal kinase activity in			
232	SOX2+ stem cells <sup>27</sup> . However, <i>Hesx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup></i> embryos showed unaffected			
233	pituitary development and normal localisation and levels of YAP and TAZ as			
234	assessed by immunofluorescence (Figure 2 – figure supplement 2A,B) when			
235	compared with controls. mRNA in situ hybridisation against Lats2 at P2, revealed			
236	abundant Lats2 transcripts upon conditional deletion of Lats1, suggesting a			
237	compensatory upregulation of Lats2 in the absence of LATS1 (Figure 2 – figure			
238	supplement 2C), similar to previous reports of elevated YAP/TAZ signalling inducing			
239	Lats2 expression <sup>31</sup> .			
240				
241	To overcome potential functional redundancy, we deleted both Lats1 and Lats2 in RP.			
	To overcome potential functional redundancy, we deleted both <i>Lats1</i> and <i>Lats2</i> in RP. Deletion of <i>Lats2</i> alone ( $Hesx1^{Cre/+}$ ; Lats2 <sup>fl/fl</sup> ), did not reveal any developmental			
241				
241 242	Deletion of <i>Lats2</i> alone ( <i>Hess1</i> <sup><i>Cre/+</i></sup> ; <i>Lats2</i> <sup><i>fl/fl</i></sup> ), did not reveal any developmental			
241 242 243	Deletion of <i>Lats2</i> alone ( <i>Hess1<sup>Cre/+</sup>;Lats2<sup>fl/fl</sup></i> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified			
<ul><li>241</li><li>242</li><li>243</li><li>244</li></ul>	Deletion of <i>Lats2</i> alone ( <i>Hesx1<sup>Cre/+</sup>;Lats2<sup>fl/fl</sup></i> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified at normal Mendelian proportions (Supplementary File 2). Similarly, deletion of any			
<ul> <li>241</li> <li>242</li> <li>243</li> <li>244</li> <li>245</li> </ul>	Deletion of <i>Lats2</i> alone ( <i>Hess1<sup>Cre/+</sup>;Lats2<sup>fl/fl</sup></i> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified at normal Mendelian proportions (Supplementary File 2). Similarly, deletion of any three out of four <i>Lats</i> alleles did not affect pituitary development and were identified			
<ul> <li>241</li> <li>242</li> <li>243</li> <li>244</li> <li>245</li> <li>246</li> </ul>	Deletion of <i>Lats2</i> alone ( <i>Hesx1<sup>Cre/+</sup>;Lats2<sup>fl/fl</sup></i> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified at normal Mendelian proportions (Supplementary File 2). Similarly, deletion of any three out of four <i>Lats</i> alleles did not affect pituitary development and were identified at normal ratios, similar to other tissues <sup>32</sup> . Homozygous <i>Hesx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup>;Lats2<sup>fl/fl</sup></i>			
<ul> <li>241</li> <li>242</li> <li>243</li> <li>244</li> <li>245</li> <li>246</li> <li>247</li> </ul>	Deletion of <i>Lats2</i> alone ( $Hesx1^{Cre/+}$ ; <i>Lats2</i> <sup>fl/fl</sup> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified at normal Mendelian proportions (Supplementary File 2). Similarly, deletion of any three out of four <i>Lats</i> alleles did not affect pituitary development and were identified at normal ratios, similar to other tissues <sup>32</sup> . Homozygous $Hesx1^{Cre/+}$ ; <i>Lats1</i> <sup>fl/fl</sup> ; <i>Lats2</i> <sup>fl/fl</sup> mutants were identified at embryonic stages at reduced Mendelian ratios and were			
<ul> <li>241</li> <li>242</li> <li>243</li> <li>244</li> <li>245</li> <li>246</li> <li>247</li> <li>248</li> </ul>	Deletion of <i>Lats2</i> alone ( $Hesx1^{Cre/+}$ ; <i>Lats2</i> <sup>fl/fl</sup> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified at normal Mendelian proportions (Supplementary File 2). Similarly, deletion of any three out of four <i>Lats</i> alleles did not affect pituitary development and were identified at normal ratios, similar to other tissues <sup>32</sup> . Homozygous $Hesx1^{Cre/+}$ ; <i>Lats1</i> <sup>fl/fl</sup> ; <i>Lats2</i> <sup>fl/fl</sup> mutants were identified at embryonic stages at reduced Mendelian ratios and were			
<ul> <li>241</li> <li>242</li> <li>243</li> <li>244</li> <li>245</li> <li>246</li> <li>247</li> <li>248</li> <li>249</li> </ul>	Deletion of <i>Lats2</i> alone ( <i>Hesx1</i> <sup><i>Crel+</i></sup> ; <i>Lats2</i> <sup><i>fllfl</i></sup> ), did not reveal any developmental morphological anomalies (Figure 2 – figure supplement 2D) and pups were identified at normal Mendelian proportions (Supplementary File 2). Similarly, deletion of any three out of four <i>Lats</i> alleles did not affect pituitary development and were identified at normal ratios, similar to other tissues <sup>32</sup> . Homozygous <i>Hesx1</i> <sup><i>Crel+</i></sup> ; <i>Lats1</i> <sup><i>fllfl</i></sup> ; <i>Lats2</i> <sup><i>fllfl</i></sup> mutants were identified at embryonic stages at reduced Mendelian ratios and were absent at P0-P2, suggesting embryonic and perinatal lethality (Supplementary File 2).			

to controls lacking *Cre* (Fig2A, n=4). Total TAZ and YAP proteins accumulated

253	throughout the developing gland in double mutants (arrowheads) but only in the
254	SOX2+ periluminal epithelium of controls (arrows). The same regions showed a
255	marked reduction in pYAP-S127 staining, which is observed in SOX2+ cells of the
256	control (Fig2A). These findings are in line with LATS1/2 normally regulating YAP
257	and TAZ in the pituitary and demonstrate successful deletion in RP. The mutant
258	pituitary was highly proliferative (Fig2B, Figure 2 – figure supplement 2F; Ki-67
259	index average 47.42% $\pm$ 1.73 SEM in control versus 76.04% $\pm$ 9.11 SEM in the double
260	mutant, P=0.0067, Student's t-test) and the majority of cells expressed SOX2
261	(Fig2A,C) but not SOX9 (Fig2B, Figure 2 – figure supplement 2F).
262	
263	By 15.5dpc the pituitary was grossly enlarged and exerting a mass effect on the brain,
264	had cysts and displayed areas of necrosis (asterisks Fig2, Figure 2 - figure
265	supplement 2E, n=5). Staining for Endomucin to mark blood vessels revealed poor
266	vascularisation in $Hesx1^{Cre/+}$ ; Lats $1^{f1/f1}$ ; Lats $2^{f1/f1}$ mutants compared to the ample
267	capillaries seen in the control (Fig2C), which may account for the necrosis. This could
268	be due to a direct inhibition of vascularisation or a consequence of the rapid growth of
269	this embryonic tumour. We frequently observed ectopic residual pituitary tissue at
270	more caudal levels, reaching the oral epithelium and likely interfering with
271	appropriate fusion of the sphenoid, similar to other phenotypes involving pituitary
272	enlargement (arrows Fig2C) <sup>33, 34, 35</sup> . Immunofluorescence to detect active (non-
273	phosphorylated) YAP revealed abundant staining throughout the pituitary at 15.5dpc,
274	compared to the control where active YAP localises in the SOX2 epithelium (Fig2C).
275	Immunofluorescence using specific antibodies against lineage commitment markers
276	PIT1, TPIT and SF1 at 15.5dpc revealed very few cells expressing PIT1, TPIT and
277	SF1 in the double mutant (Fig2D; PIT1 9.14% in mutants compared with 51.4% in

278controls (Student's *t*-test P < 0.0001); TPIT 4.0% in mutants compared with 11.4% in279controls (Student's *t*-test P < 0.007); SF1 2.1% in mutants compared with 6.5% in280controls (Student's *t*-test P > 0.05) n=3 mutants and 5 controls), suggesting failure to281commit into the three lineages. These data suggest that the LATS/YAP/TAZ axis is282required for normal embryonic development of the anterior pituitary and that283LATS1/2 kinases control proliferation of SOX2+ progenitors and their progression284into the three committed lineages.

285

#### 286 Loss of LATS kinases results in carcinoma-like murine tumours

Postnatal analysis of *Hesx1<sup>Cre/+</sup>*;*Lats1<sup>fl/fl</sup>* pituitaries revealed that by P56, despite 287 288 developing normally during the embryonic period, all glands examined exhibited 289 lesions of abnormal morphology consisting of overgrowths, densely packed nuclei 290 and loss of normal acinar architecture (n=15). To minimise the likely redundancy by 291 LATS2 seen at embryonic stages, we generated *Lats1* mutants additionally haploinsufficient for Lats2 (Hesx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup>;Lats2<sup>fl/+</sup>). These pituitaries also 292 developed identifiable lesions accumulating YAP and TAZ (Figure 3 – figure 293 294 supplement 1A), which were observed at earlier time points (P21 n=4), the earliest 295 being 10 days, indicating increased severity. The number of lesions observed per 296 animal was similar between the two models at P56 (3-8 per animal). Deletion of Lats2 alone (*Hess1<sup>Cre/+</sup>*;*Lats2<sup>fl/fl</sup>*), which is barely expressed in the wild type pituitary, did 297 298 not result in any defects (Figure 3 – figure supplement 1B). We focused on the  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$  double mutants for further analyses. 299

300

301 Histological examination of  $Hesx1^{Cre/+}$ ;  $Lats1^{fl/fl}$ ;  $Lats2^{fl/+}$  pituitaries confirmed the 302 abnormal lesions were tumours, characterised by frequent mitoses, focal necrosis, and

303	a focal squamous differentiation, as well as the occasional presence of cysts (Fig3A).
304	These lesions were identical to those in <i>Hesx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup></i> pituitaries (not shown).
305	These tumours accumulated YAP/TAZ and upregulated expression of targets Cyr61
306	and Ctgf (Fig3B), confirming the validity of the genetic manipulation (Fig3B).
307	Tumours were also frequently observed in the anterior and intermediate lobe (Figure
308	3 – figure supplement 1C). Analysis of proliferation by Ki-67 immunostaining
309	revealed an elevated mitotic index of 7-28% in tumours (mean 15.46, SEM $\pm 2.74$ ),
310	compared to 2.97% (SEM $\pm 1.2$ ) mean in control pituitaries not carrying the Lats1
311	deletion (Fig3C).
312	In keeping with the morphological evidence of epithelial differentiation (Fig3A), the
313	tumours were positive for cytokeratins using AE1/AE3 (multiple keratin cocktail)
314	(Figure 3 – figure supplement 1E). Furthermore, the tumours showed focal
315	morphological evidence of squamous differentiation and showed positive nuclear p63
316	staining, frequently expressed in squamous carcinomas (Figure 3 – figure supplement
317	1E). In contrast, the tumours did not show immunohistochemical evidence of
318	adenomas i.e. negative for neuroendocrine markers, which all types of adenomas are
319	typically positive for: the neuroendocrine marker synaptophysin and neuron-specific
320	enolase (Figure 3 – figure supplement 1F). The lesions were also negative for
321	chromogranin A, a neuroendocrine granule marker often expressed in clinically non-
322	functioning pituitary adenomas. Tumours were also negative for vimentin, expressed
323	by spindle cell oncocytoma, an uncommitted posterior pituitary tumour (Figure 3 –
324	figure supplement 1F). Moreover, immunostaining against PIT1, TPIT and SF1
325	showed only sparse positive cells within the lesions, suggesting lack of commitment
326	into endocrine precursors and supporting the undifferentiated nature of the tumour
327	cells (Fig3D). Consistent with a tumourigenic phenotype, and role for LATS1

328 genomic stabilisation<sup>36</sup>, staining for gamma-H2A.X detected elevated DNA damage

329 in cells of the mutant pituitaries compared with controls (Figure 3 – figure supplement

1D). The absence of adenoma or oncocytoma markers together with the histological

appearance, observation of focal necrosis and a high mitotic index support the

- 332 features of squamous carcinoma.
- 333

#### **SOX2** +ve cells are the cell of origin of the tumours

335 Tumour regions were mostly composed of SOX2 positive cells, a sub-population of

which also expressed SOX9 (Fig3E, Figure 3 - figure supplement 1A; 85-97% of

337 cells, 7 tumours across 4 pituitaries). Close examination of the marginal zone

epithelium, a major SOX2+ stem cell niche of the pituitary, revealed a frequent

339 'ruffling' resembling crypts, likely generated through over-proliferation of the

340 epithelial stem cell compartment (Fig3F). To determine if the cell of origin of the

341 tumourigenic lesions is a deregulated SOX2+ stem cell, we carried our specific

deletion of LATS1/2 in postnatal SOX2+ cells using the tamoxifen-inducible Sox2-

343 *CreERT2* driver, combined with conditional expression of membrane-GFP in targeted

344 cells ( $Sox2^{CreERT2/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$ ;  $R26^{mTmG/+}$ ).

345

Tamoxifen induction at P5 or P21, led to abnormal lesions in the anterior pituitary within three months in all cases. We focused our analyses on inductions performed at P5, from which time point all animals developed lesions by P35 (Fig4A). Similar to observations in  $Hesx1^{Cre/+}$ ; Lats1<sup>fl/fl</sup>; Lats2<sup>fl/+</sup> animals, these areas strongly accumulated YAP and TAZ (Fig4B), activated expression of targets *Cyr61* and *Ctgf*, displayed ruffling of the AL epithelium (Fig4C, Figure 4 – figure supplement 1E) and lacked lineage commitment markers (Fig4D, Figure 4 – figure supplement 1A). These

353 lesions showed a similar marker profile to *Hesx1-Cre*-targeted tumours, with positive

354 p63 and AE1/AE3 staining (Figure 4 – figure supplement 1B). Lineage tracing

355 confirmed expression of membrane GFP in tumourigenic lesions, characterised by the

accumulation of YAP and expansion of SOX2+ cells, suggesting they were solely

derived from SOX2+ cells (Fig4E, Figure 4 – figure supplement 1C). Taken together,

358 our data support that LATS kinase activity is required to regulate the pituitary stem

359 cell compartment. Loss of LATS1 is sufficient to drive deregulation of SOX2+

360 pituitary stem cells, generating highly proliferative non-functioning tumours with

Conditional deletion of LATS1/2 kinases in the pituitary has revealed how these

361 features of carcinomas.

362

364

# 363 YAP expression is sufficient to activate pituitary stem cells.

365 promote an expansion of SOX2+ve stem cells in the embryonic and postnatal gland at the expense of differentiation. To establish if this effect was mediated through YAP 366 alone, we used the tetracycline-controlled conditional YAP-TetO system to promote 367 YAP (S127A) protein levels in postnatal pituitaries of Hesx1<sup>Cre/+</sup>;R26<sup>rtTA/+</sup>;Colla1<sup>tetO-</sup> 368 <sup>Yap/+</sup> mice. We treated YAP-TetO animals with doxycycline from P21 to P105 (12 369 370 week treatment, Fig5A). We did not observe the formation of tumours at any stage 371 analysed (n=12, Figure 5 – figure supplement 1A). Similarly, we did not observe the 372 formation of lesions when treating from P5. This is in contrast with the unequivocal tumour formation observed in Sox2<sup>CreERT2/+</sup>;Lats1<sup>fl/fl</sup>;Lats2<sup>fl/+</sup> mice. Elevation of YAP 373 374 protein levels was confirmed following three weeks of doxycycline treatment (P42), 375 displaying patchy accumulation, likely a result of genetic recombination efficiencies 376 (Fig5B). Consistent with pathway activation, there was robust elevation in the 377 expression of transcriptional targets Cyr61 and Ctgf following treatment (Figure 5 –

378 figure supplement 1B), however at significantly lower levels compared to

379  $Sox2^{CreERT2/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$  deletions (Figure 5 – figure supplement 1E), and there 380 was no elevation in phosphorylated inactive YAP (Fig5B).

381

382 Immunofluorescence against SOX2 demonstrated a significant increase in the number 383 of SOX2+ cells as a proportion of the anterior pituitary (Fig5B,F; 18.0% compared to 384 12.1% in controls, P=0.0014), a finding recapitulated by SOX9 that marks a subset of 385 the SOX2 population (Fig5B). This increase in the percentage of SOX2+ cells was 386 maintained at all stages analysed (Fig5F) and did not affect the overall morphology of 387 the pituitary. At P42 we observed a significant increase in proliferation among the 388 SOX2+ pituitary stem cells from 3% in controls to 15% in mutants (P=0.027). 389 SOX2+ cells make up 10% of all cycling cells (Ki-67%) in normal pituitaries, 390 however in mutants this increased to 25%, suggesting a preferential expansion of the 391 SOX2+ population, rather than an overall increase in proliferation (Fig5C). No 392 additional marked differences were observed in samples analysed at P63 (6 weeks of 393 treatment, n=3), however longer treatment (P21 to P105) resulted in sporadic regions 394 of expanded SOX2+ cells (Figure 5 – figure supplement 1C). These regions did not 395 express the commitment marker PIT1 and were identifiable by haematoxylin/eosin 396 staining. In contrast to tumour lesions generated following loss of LATS kinases, 397 these were not proliferative, were positive for pYAP and did not accumulate high 398 levels of YAP/TAZ (n=6 lesions). Together these results suggest that the sustained 399 expression of constitutive active YAP can activate the proliferation of SOX2 stem 400 cells, but in contrast to deletion of LATS1, this alone is not oncogenic. 401

402 To establish if the expansion of pituitary stem cells following forced expression of

403 YAP is reversible, we administered doxycycline to YAP-TetO animals for three

404 weeks (P21 to P42) by which point there is a robust response, followed by

- 405 doxycycline withdrawal for three weeks (until P63) to allow sufficient time for YAP
- 406 levels to return to normal (scheme Fig5D). Immunofluorescence against total YAP
- 407 protein confirmed restoration of the normal YAP expression pattern and levels after
- 408 recovery (Fig5E), and mRNA in situ hybridisation detected a reduction in expression
- 409 of YAP/TAZ targets *Cyr61* and *Ctgf* (Figure 5 figure supplement 1D). Following
- 410 recovery from high levels of YAP, the number of SOX2+ cells reduced to comparable

411 levels as in controls (around 10% of the total anterior pituitary) (Fig5E,F). This

412 suggests that the effects of YAP overexpression on the stem cell population are

413 transient following three weeks of treatment (Fig5F).

414

415 Finally, to determine if SOX2+ cells could differentiate into hormone-producing cells

416 after the reduction in YAP levels, we expressed constitutive active YAP only in

417 SOX2+ cells whilst lineage tracing this population

418  $(Sox2^{CreERT2/+}; R26^{rtTA/mTmG}; Collal^{tetO-Yap/+})$ . We induced SOX2+ cells by low-dose

419 tamoxifen administration at P21 and treated with doxycycline for three weeks,

420 followed by doxycycline withdrawal for a further three weeks (Fig5G). Larger clones

421 of SOX2 derivatives were observed at P63 in  $Sox2^{CreERT2/+}$ ;  $R26^{rtTA/mTmG}$ ;  $Colla1^{tetO-}$ 

422 *Yap/+* animals compared to controls, and these still contained SOX2+ cells (Fig5H).

423 Following withdrawal, we were able to detect GFP+ derivatives of SOX2+ cells,

- 424 which had differentiated into the three lineages (PIT1, SF1 and ACTH, marking
- 425 corticotrophs of the TPIT lineage) (Fig5I). Taken together, these findings confirm that
- 426 sustained expression of YAP is sufficient to maintain the SOX2+ state and promote
- 427 activation of normal SOX2+ pituitary stem cells in vivo, driving expansion of this

428 population.

# **DISCUSSION**

433	Here we establish that regulation of LATS/YAP/TAZ signaling is essential during
434	anterior pituitary development and can influence the activity of the stem/progenitor
435	cell pool. LATS kinases, mediated by YAP and TAZ, are responsible for controlling
436	organ growth, promoting an undifferentiated state and repressing lineage
437	commitment. Loss of both Lats1 and Lats2, encoding potent tumour suppressors,
438	leads to dramatic tissue overgrowth during gestation, revealing a function for these
439	enzymes in restricting growth during pituitary development. The involvement of
440	YAP/TAZ and dysfunction of the kinase cascade is emerging in multiple paediatric
441	cancers, which are often developmental disorders <sup>37</sup> .
442	
443	Loss of <i>LATS1</i> heterozygosity has been reported in a range of human tumours <sup>38, 39, 40,</sup>
443 444	Loss of <i>LATS1</i> heterozygosity has been reported in a range of human tumours <sup>38, 39, 40,</sup> <sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i>
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444 445	<sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i> in mice resulted in a variety of soft tissue sarcomas and stromal cell tumours <sup>42</sup> . The
444 445 446	<sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i> in mice resulted in a variety of soft tissue sarcomas and stromal cell tumours <sup>42</sup> . The anterior lobe of these animals appeared hyperplastic with poor endocrine cell
444 445 446 447	<sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i> in mice resulted in a variety of soft tissue sarcomas and stromal cell tumours <sup>42</sup> . The anterior lobe of these animals appeared hyperplastic with poor endocrine cell differentiation leading to combined hormone deficiencies, but the presence of tumours
444 445 446 447 448	<sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i> in mice resulted in a variety of soft tissue sarcomas and stromal cell tumours <sup>42</sup> . The anterior lobe of these animals appeared hyperplastic with poor endocrine cell differentiation leading to combined hormone deficiencies, but the presence of tumours was not noted. We report that loss of <i>Lats1</i> alone is sufficient to drive anterior and
444 445 446 447 448 449	<ul> <li><sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i> in mice resulted in a variety of soft tissue sarcomas and stromal cell tumours<sup>42</sup>. The anterior lobe of these animals appeared hyperplastic with poor endocrine cell differentiation leading to combined hormone deficiencies, but the presence of tumours was not noted. We report that loss of <i>Lats1</i> alone is sufficient to drive anterior and intermediate lobe tumour formation. This phenotype is accelerated following</li> </ul>
444 445 446 447 448 449 450	<sup>41</sup> leading to an increase in YAP/TAZ protein levels. Previous global deletion of <i>Lats1</i> in mice resulted in a variety of soft tissue sarcomas and stromal cell tumours <sup>42</sup> . The anterior lobe of these animals appeared hyperplastic with poor endocrine cell differentiation leading to combined hormone deficiencies, but the presence of tumours was not noted. We report that loss of <i>Lats1</i> alone is sufficient to drive anterior and intermediate lobe tumour formation. This phenotype is accelerated following additional deletion of one copy of <i>Lats2</i> . Phenotypically identical tumour lesions were

453	LATS activation in other tissues <sup>43</sup> , did not lead to any pituitary defects despite
454	reduction in STK3/4 levels. These data suggest that perhaps the residual activity of
455	STK3/4 is sufficient for LATS1/2 activation. Alternatively, regulation of LATS1/2 by
456	kinases other than STK3/4 is possible in the pituitary, meaning deletion of $Stk3/4$
457	alone is insufficient to result in significant LATS function impairment. Similar
458	situations have been reported in other organs where LATS are functioning <sup>43</sup> . The
459	resulting non-secreting tumours in our mouse models are composed predominantly of
460	SOX2+ stem cells and display signs of squamous differentiation. Rare cases of
461	squamous cell carcinoma have been reported as primary pituitary tumours <sup>44</sup> , but more
462	frequently, arising within cysts that are normally non-neoplastic epithelial
463	malformations <sup>45, 46</sup> . In the embryonic YAP-TetO model, where constitutive active
464	YAP (S127A) was expressed during pituitary development, cysts phenocopying
465	Rathke's cleft cyst, develop by postnatal stages. Target elevation is not as high in
466	YAP-TetO pituitaries, as following the deletion of LATS1/2, indicating that signaling
467	levels are likely to be critical for progression between these phenotypes.
468	Although human pituitary carcinomas are only diagnosed as such after metastasis, the
469	tumours generated in our LATS1/2 mouse models fit their histopathological profile.
470	Genetic lineage tracing identified SOX2+ cells as the cell of origin of the tumours;
471	this observation could have ramifications regarding involvement of the
472	LATS/YAP/TAZ pathway in the establishment or progression of human pituitary
473	tumours composed of uncommitted cells. In cancer stem cells of osteosarcoma and
474	glioblastoma, SOX2 antagonises upstream Hippo activators, leading to enhanced
475	YAP function <sup>47</sup> . We recently reported enhanced expression of YAP/TAZ in a range of
476	non-functioning human pituitary tumours, compared to functioning adenomas, and
477	that Lats1 knock-down in GH3 pituitary mammosomatotropinoma cells results in

478 repression of the *Gh* and *Prl* promoters<sup>28</sup>. Therefore, YAP/TAZ, perhaps in a positive
479 feedback loop with SOX2, are likely to function both to promote the maintenance of
480 an active pituitary stem cell state as well as to inhibit differentiation.

481

482 By dissecting the downstream requirement for YAP in pituitary regulation by the 483 LATS/YAP/TAZ axis, we found that expression of constitutively active YAP 484 (S127A) is sufficient to push SOX2+ pituitary stem cells into an activated state, 485 leading to expansion of the stem cell cohort (see Model, Fig6). YAP has previously 486 been indicated to promote the stem cell state in other tissues, e.g. pancreas, neurons, mammary glands<sup>48</sup>. However, this does not fully recapitulate the LATS deletion 487 488 phenotypes, as it did not lead to the formation of tumours during the time course of 489 YAP activation (12 weeks). Interestingly, since the levels of target activation are 490 significantly greater in Lats1/2 deletions that in YAP-TetO activation, initiation of 491 tumourigenesis may be associated with levels of signalling rising above a threshold. 492 However, the temporal control of expressing the mutation is critical, as seen in other tumour models<sup>49</sup>. Instead, the findings identify an isolated role for YAP in promoting 493 494 the expansion of the SOX2+ stem cell pool and restoring their proliferative potential 495 to levels akin to the most active state during postnatal pituitary growth. Activity of 496 YAP/TAZ is reduced in dense tissues, resulting in a decrease in stemness. One 497 mechanism through which this is achieved is by crosstalk with other signaling pathways regulating stem cell fate<sup>50, 51</sup>. For example, a decrease in YAP/TAZ activity 498 499 removes inhibition on Notch signalling, resulting in higher levels of differentiation and a drop in stem cell potential<sup>52</sup>. In the pituitary, Notch plays a role in the 500 501 maintenance of the SOX2 stem cell compartment and is involved in regulating differentiation<sup>53, 54, 55, 56</sup>. The downstream mechanisms of YAP action on SOX2+ 502

pituitary stem cells, as well as the likely crosstalk with other signalling pathwaysremain to be explored.

505

506	In summary, our findings highlight roles for LATS/YAP/TAZ in the regulation of
507	pituitary stem cells, where fine-tuning of their expression can make the difference
508	between physiological stem cell re-activation and tumourigenesis, of relevance to
509	other organs. We reveal this axis is involved in the control of cell fate commitment,
510	regulation of regenerative potential and promotion of tumourigenesis. These findings
511	can aid in the design of treatments against pituitary tumours and in regenerative
512	medicine approaches targeting the regulation of endogenous stem cells.

513

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527

# 529 MATERIALS & METHODS

# 530 Key Resources Table

Reagent type	Designation	Source or	Identifiers	Additional
(species) or		reference		Information
resource				
Genetic reagent	Hesx1 <sup>Cre/+</sup>	PMID:	RRID:MGI:5314529	
(M. musculus)		17360769		
Genetic reagent	Sox2 <sup>CreERT2/+</sup>	PMID:	MGI:5512893	
(M. musculus)		24094324		
Genetic reagent	Lats 1 <sup>fl/fl</sup>	Jackson	Stock #: 024941,	
(M. musculus)		Laboratory	RRID: MGI:5568576	
Genetic reagent	Lats2 <sup>fl/fl</sup>	Jackson	Stock #: 025428,	
(M. musculus)		Laboratory	RRID: MGI:5568577	
Genetic reagent	Stk4 <sup>fl/fl</sup> ;Stk3 <sup>fl/fl</sup>	Jackson	Stock #: 017635,	PMID:
(M. musculus)		Laboratory	RRID: MGI:5301573	20080689
Genetic reagent	R26 <sup>rtTA/+</sup>	Jackson	Stock: #: 016999	PMID:
(M. musculus)		Laboratory	RRID: MGI:5292520	15941831
Genetic reagent	Colla1 <sup>tetO-Yap/+</sup>	PMID:	MGI:5430522	
(M. musculus)		22363786		
Genetic reagent	$R26^{mTmG/+}$	Jackson	Stock #: 007576	PMID:
(M. musculus)		Laboratory	RRID:MGI:3722405	17868096
Genetic reagent	Taz <sup>-/-</sup>	Jackson	Stock #: 011120,	PMID:

(M. musculus)		Laboratory	RRID: MGI:4420900	17636028
Genetic reagent	Yap <sup>f1/f1</sup>	PMID:	MGI:5316446	
(M. musculus)		21376238		
Antibody	Rabbit	Atlas	Cat# HPA007415	IF: 1:1000
	polyclonal anti-	Antibodies	RRID:AB_1080602	
	TAZ			
Antibody	Rabbit	Cell Signaling	Cat# 4912S	IF: 1:1000
	polyclonal anti-	Technology	RRID:AB_2218911	
	YAP			
Antibody	Rabbit	Cell Signaling	Cat# 4911S	IF: 1:1000
	polyclonal anti-	Technology	RRID:AB_2218913	
	pYAP			
Antibody	Rabbit	Abcam	Cat# ab97959	IF: 1:2000
	polyclonal anti-		RRID:AB_2341193	
	SOX2			
Antibody	Rat monoclonal	Abcam	Cat# ab106100	IF: 1:1000
	anti-EMCN		RRID:AB_10859306	
	(V.7C7.1)			
Antibody	Chicken	Abcam	Cat# ab13970	IF: 1:300
	polyclonal anti-		RRID:AB_300798	
	GFP			
Antibody	Goat polyclonal	Immune	Cat# GT15098	IF: 1:250
	anti-SOX2	Systems	RRID:AB_2732043	
		Limited		
Antibody	Rabbit	Abcam	Cat# ab16667	IF: 1:300
	1	1	1	1

	monoclonal		RRID:AB_302459	
	anti-Ki-67			
Antibody	Rabbit	Proteintech	Cat# 17711-1-AP,	IF: 1:100
	polyclonal anti-	Group	RRID:AB_2060867	
	ARL13B			
Antibody	Mouse	Sigma-Aldrich	Cat# MABT868	IF: 1:200
	monoclonal			
	anti-Acetylated-			
	αΤUΒ			
Antibody	Rabbit	Abcam	Cat# ab185230	IF: 1:300
	monoclonal		RRID:AB_2715497	
	anti-SOX9			
Antibody	Rabbit	Abcam	Cat# ab205270	IF: 1:300
	monoclonal			
	anti-Active YAP			
	EPR19812			
Antibody	Rabbit	Prof. S.		IF: 1:1000
	polyclonal anti-	Rhodes		
	PIT1	(Indiana		
		University		
Antibody	Rabbit	Prof. J. Drouin		IF: 1:1000
	polyclonal anti-	(Montreal		
	TPIT	IRCM)		
Antibody	Mouse	Life	Cat# N1665	IF: 1:200
	monoclonal	Technologies	RRID:AB_2532209	

	anti-SF1	(Thermo		
		Fisher		
		Scientific)		
Antibody	Rabbit	Abcam	Cat# ab2893	IF: 1:1000
	polyclonal anti-		RRID:AB_303388	
	gamma H2A.X			
	(phospho S139)			
Antibody	Rabbit	Bethyl	Cat# A300-466A	WB:
	polyclonal anti-	Laboratories	RRID:AB_2148394	1:5000
	STK3/4			
Antibody	Mouse	R&D Systems	Cat# MAB5410	WB:
	monoclonal		RRID:AB_2169416	1:1000
	anti-Cyclophilin			
	В			
	(Clone#			
	549205)			
Antibody	Rabbit	Cell Signaling	Cat# 5741	IF: 1:300
	monoclonal	Technology	RRID:AB_10695459	
	anti-Vimentin			
	(D21H3)			
Antibody	Biotinylated	Abcam	Cat# ab6720	IF: 1:350
	Goat polyclonal		RRID:AB_954902	
	anti-rabbit			
Antibody	Goat polyclonal	Life	Cat# A11039	IF: 1:300
	anti-chicken	Technologies	RRID:AB_2534096	

	Alexa Fluor 488	(Thermo		
		Fisher		
		Scientific)		
Antibody	Goat polyclonal	Life	Cat# A21434	IF: 1:300
	anti-rat Alexa	Technologies	RRID:AB_2535855	
	Fluor 555	(Thermo		
		Fisher		
		Scientific)		
Antibody	Biotinylated	Abcam	Cat# ab6788	IF: 1:350
	Goat polyclonal		RRID:AB_954885	
	anti-mouse			
Antibody	Donkey	Abcam	Cat# ab150133	IF: 1:300
	polyclonal anti-			
	goat Alexa Fluor			
	488			
Antibody	Streptavidin	Life	Cat# S21381	IF: 1:500
	Alexa Fluor 555	Technologies	RRID:AB_2307336	
Antibody	Goat HRP-	Cell Signaling	Cat# 7074	WB:
	linked anti-	Technology	RRID:AB_2099233	1:2000
	rabbit			
Antibody	Goat HRP-	Cell Signaling	Cat# 7076	WB:
	linked anti-	Technology	RRID:AB_330924	1:2000
	mouse			
Antibody	Mouse	Dako	Cat# M351529	IHC: 1:100
	monoclonal			

	anti-AE1/AE3			
Antibody	Mouse	Dako	Cat# M086901	IHC: 1:400
	monoclonal			
	anti-			
	Chromogranin			
Antibody	Mouse	Novocastra	Cat# NCL-L-CD56-	IHC 1:15
	monoclonal		504	
	anti-NCAM			
Antibody	Mouse	Dako	Cat# M087329	IHC 1:1000
	monoclonal			
	anti-NSE			
Antibody	Mouse	A. Menarini	Cat# MP163	IHC 1:100
	monoclonal	Diagnostics		
	anti-p63			
Antibody	Mouse	Dako	Cat# M731529	IHC 1:2
	monoclonal		RRID:AB_2687942	
	anti-			
	Synaptophysin			
Commercial	TSA kit	Perkin Elmer	Cat# NEL753001KT	
assay or kit				
Commercial	TSA Blocking	Perkin Elmer	Cat# FP1020	
assay or kit	Reagent			
Commercial	ABC kit	Vector	Cat# Vector PK-6100	
assay or kit		Laboratories	RRID:AB_2336819	
Commercial	BCA assay	Thermo Fisher	Cat# 23227	

assay or kit				
Commercial	UltraView	Ventana	Cat# 760-500	
assay or kit	Universal DAB	Medical		
	Detection Kit	Systems		
Commercial	VectaFluor	Vector	Cat# DK-2488	
assay or kit	Excel R.T.U.	Laboratories	RRID:AB_2336775	
	Antibody Kit,			
	DyLight 488			
	Anti-Mouse			
Chemical	Doxycycline	Alfa Aesar	Cat# J60579	2mg/ml
compound, drug	hyclate			
Chemical	Sucrose	Sigma-Aldrich	Cat# S0389	10mg/ml
compound, drug				
Chemical	Tamoxifen	Sigma-Aldrich	Cat# T5648	0.15mg/g
compound, drug				
Chemical	Hoechst 33342	Life	Cat# H3570	1:10000
compound, drug		Technologies		
Chemical	Laemmli buffer	Bio-Rad	Cat# 1704156	
compound, drug				
Chemical	Clarity Western	Bio-Rad	Cat# 170-5060	
compound, drug	ECL Substrate			
Chemical	Alcian Blue	Alfa Aeser	Cat# J60122	1%
compound, drug				
Chemical	Acetic acid	VWR	Cat# 20103	3%
compound, drug				

Chemical	Periodic acid	VWR	Cat# 29460	1%
compound, drug				
Chemical	Schiff's reagent	Thermo Fisher	Cat# 88017	
compound, drug		Scientific		
Software,	GraphPad Prism	GraphPad	RRID:SCR_015807	
algorithm		Software		
		(www.graphpa		
		<u>d.com</u> )		
Software,	Fiji	Schindelin et	RRID:SCR_002285	
algorithm		al., 2012		
		(Fiji.sc)		
Software,	ImageLab	BioRad		
algorithm				
Other	Probe: Ctgf	ACDBio	Cat# 314541	
Other	Probe: Cyr61	ACDBio	Cat# 429001	
Other	Probe: Lats2	ACDBio	Cat# 420271	
Other	Probe: Nr5a1	ACDBio	Cat# 445731	
Other	Probe: Tbx19	ACDBio	Cat# 484741	
Other	Probe: Poulfl	ACDBio	Cat# 486441	

532

533

# 534 Animals

535 Animal husbandry was carried out under compliance of the Animals (Scientific

536 Procedures) Act 1986, Home Office license and KCL ethical review approval.

- 537 The  $Hesx1^{Cre/+57}$ ,  $Sox2^{CreERT2/+1}$ ,  $Yap^{fl/fl 25}$ ,  $Taz^{--30}$ (JAX:011120),  $R26^{mTmG/+1}$
- <sup>58</sup>(JAX:007576), *ROSA26<sup>rtTA/+ 59</sup>* (JAX:016999), *Collal<sup>tetO-Yap/+ 60</sup>*(MGI:5430522),

539  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl \, 61}$  (JAX:017635), and  $Lats1^{fl/fl \, 51}$  (JAX:024941) and  $Lats2^{fl/fl}$ 

- $540 \quad {}^{51}$ (JAX:025428) have been previously described.
- 541 Tamoxifen (Sigma, T5648) was administered to experimental mice by intraperitoneal
- 542 injection at a single dose of 0.15mg/g body weight, or two equal doses on sequential
- 543 days, depending on the experiment. Mice for growth studies were weighed every
- 544 week. For embryonic studies, timed matings were set up where noon of the day of
- 545 vaginal plug was designated as 0.5dpc.
- 546 For YAP-TetO experiments, crosses between  $Hesx1^{Cre/+}$ ;  $R26^{+/+}$ ;  $Colla1^{+/+}$  and
- 547  $Hesx1^{+/+}$ ;  $R26^{rtTA/rtTA}$ ;  $Collal^{tetO-Yap/tetO-Yap}$  animals were set up to generate
- 548 *Hesx1<sup>Cre/+</sup>;R26<sup>rtTA/+</sup>;Col1a1<sup>tetO-Yap/+</sup>* offspring (hereby YAP-TetO) and control
- 549 littermates, or crosses between  $Sox2^{CreERT2/+}$ ;  $R26^{mTmG/mTmG}$ ;  $Collal^{+/+}$  and  $Sox2^{+/+}$ ;
- 550  $R26^{rtTA/rtTA}$ ; Collal<sup>tetO-Yap/tetO-Yap</sup> animals were set up to generate
- 551  $Sox2^{CreERT2/+}$ ;  $R26^{rtTA/mTmG}$ ;  $Collal^{tetO-Yap/+}$  offspring. Whilst treated with the
- 552 tetracycline analogue doxycycline, YAP-TetO expressed rtTA from the *ROSA26*
- 553 locus in *Cre*-derived cells, enabling YAP S127A expression from the *Collal* locus.
- 554 For embryonic studies between 5.5dpc and 15.5dpc (scheme, Fig1A), doxycycline
- 555 (Alfa Aesar, J60579) was administered to pregnant dams in the drinking water at
- 556 2mg/ml, supplemented with 10% sucrose. For postnatal analyses animals were treated
- 557 with doxycycline or vehicle (DMSO) as described, from the ages specified for
- 558 individual experiments on the *Hesx1<sup>Cre/+</sup>* driver, or directly following tamoxifen
- administration for animals on the  $Sox2^{CreERT2/+}$  driver. Both male and female mice and
- 560 embryos where included in the studies.
- 561

#### 562 **Tissue preparation**

563 Embryos and adult pituitaries were fixed in 10% neutral buffered formalin (Sigma)

564 overnight at room temperature. The next day, tissue was washed then dehydrated

through graded ethanol series and paraffin-embedded. Embryos up to 13.5dpc were

- sectioned sagittal and all older embryo and postnatal samples were sectioned frontal,
- 567 at a thickness of 7µm for immunofluorescence staining, or 4µm for RNAscope
- 568 mRNA *in situ* hybridisation.

569

# 570 RNAscope mRNA in situ hybridisation

571 Sections were selected for the appropriate axial level, to include Rathke's pouch or

572 pituitary, as described previously <sup>27</sup>. The RNAscope 2.5 HD Reagent Kit-RED assay

573 (Advanced Cell Diagnostics) was used with specific probes: *Ctgf*, *Cyr61*, *Lats2* (all
574 ACDBio).

575

# 576 H&E staining

577 Sections were dewaxed in histoclear and rehydrated through graded ethanol series

from 100% to 25% ethanol, then washed in distilled  $H_2O$ . Sections were stained with

579 Haematoxylin QS (Vector #H3404) for 1 minute, and then washed in water. Slides

580 were then stained in eosin in 70% ethanol for 2 minutes and washed in water. Slides

- 581 were dried and coverslips were mounted with VectaMount permanent mounting
- 582 medium (Vector Laboratories H5000).

583

## 584 Immunofluorescence and immunohistochemistry

585 Slides were deparaffinised in histoclear and rehydrated through a descending graded

586 ethanol series. Antigen retrieval was performed in citrate retrieval buffer pH6.0, using

587 a Decloaking Chamber NXGEN (Menarini Diagnostics) at 110°C for 3mins.

588 Tyramide Signal Amplification (TSA) was used for staining using antibodies against

589 YAP (1:1000, Cell Signaling #4912S), pYAP (1:1000, Cell Signaling #4911S), TAZ

- 590 (1:1000, Atlas Antibodies #HPA007415) and SOX2 (1:2000, Abcam ab97959) with
- 591 EMCN (1:1000, Abcam ab106100) staining as follows: sections were blocked in TNB
- 592 (0.1M Tris-HCl, pH7.5, 0.15M NaCl, 0.5% Blocking Reagent (Perkin Elmer
- 593 FP1020)) for 1 hour at room temperature, followed by incubation with primary
- antibody at 4C overnight, made up in TNB. Slides were washed three times in TNT
- 595 (0.1MTris-HCl pH7.5, 0.15M NaCl, 0.05% Tween-20) then incubated with secondary
- antibodies (biotinylated anti-rabbit (1:350 Abcam ab6720) and anti-Rat Alexa Fluor

597 555 (1:300, Life Technologies A21434) for 1 hour at room temperature and Hoechst

598 (1:10000, Life Technologies H3570). Slides were washed again then incubated in

- 599 ABC reagent (ABC kit, Vector Laboratories PK-6100) for 30 mins, followed by
- 600 incubation with TSA conjugated fluorophore (Perkin Elmer NEL753001KT) for ten
- 601 minutes. Slides were washed and mounted with VectaMount (Vector Laboratories
- 602 H1000).
- 603 For regular immunofluorescence sections were blocked in blocking buffer (0.15%
- 604 glycine, 2mg/ml BSA, 0.1% Triton-X in PBS), with 10% sheep serum (donkey serum
- for goat SOX2 antibody) for 1 hour at room temperature, followed by incubation with
- 606 primary antibody at 4C overnight, made up in blocking buffer with 1% serum.
- 607 Primary antibodies used were against SOX2 (1:250, Immune Systems Ltd GT15098),
- 608 active YAP (1:300, Abcam ab205270), GFP (1:300, Abcam ab13970), Ki-67 (1:300,
- 609 Abcam ab16667), SOX9 (1:300, Abcam ab185230), PIT1 (1:1000, Gift from S.
- 610 Rhodes, Indiana University), TPIT (1:1000, Gift from J. Drouin, Montreal), SF1
- 611 (1:200, Life Technologies N1665), Gamma H2A.X (1:1000, Abcam ab2893),

612	Vimentin (1:300, Cell Signaling #5741), Caspase (1:300, Cell Signaling #9661S).
613	Slides were washed in PBST then incubated with secondary antibodies for 1 hour at
614	room temperature. Appropriate secondary antibodies were incubated in blocking
615	buffer for 1 hr at room temperature (biotinylated anti-rabbit (1:350, Abcam ab6720),
616	biotinylated anti-mouse (1:350, Abcam ab6788), anti-chicken 488 (1:300, Life
617	Technologies A11039), anti-goat 488 (1:300, Abcam ab150133). Slides were washed
618	again using PBST and incubated with fluorophore-conjugated Streptavidin (1:500,
619	Life Technologies S21381 or S11223) for 1 hour at room temperature, together with
620	Hoechst (1:10000, Life Technologies H3570). Slides were washed in PBST and
621	mounted with VectaMount (Vector Laboratories, H1000).
622	
623	Immunohistochemistry for the remaining antigens were undertaken on a Ventana
624	Benchmark Autostainer (Ventana Medical Systems) using the following primary
625	antibodies and antigen retrieval: AE1/AE3 (1:100, Dako M351529), CC1 (36
626	minutes, Ventana Medical Systems 950-124); Chromogranin (1:400, Dako
627	M086901), CC1 (36 minutes, Ventana Medical Systems 950-124); NCAM (1:15,
628	Novocastra NCL-L-CD56-504), CC1 (64 minutes, Ventana Medical Systems 950-
629	124); NSE (1:1000, Dako M087329), CC1 (36 minutes, Ventana Medical Systems
630	950-124); p63 (1:100, A. Menarini Diagnostics), CC1 (64 minutes, Ventana Medical
631	Systems 950-124) and Synaptophysin (1:2, Dako M731529), CC2 (92 minutes,
632	Ventana Medical Systems 950-124). Targets were detected and viewed using the
633	ultraView Universal DAB Detection Kit (Ventana Medical Systems, 760-500)
634	according to manufacturer's instructions.
635	

# 636 Alcian Blue with Periodic Acid-Schiff staining (AB/PAS)

Following deparaffinisation and rehydration, sections were taken through distilled
water then placed in Alcian Blue solution (1% Alcian Blue (Alfa Aeser J60122) in
3% acetic acid (VWR International 20103)) for 20 minutes. Sections were then placed
in 1% periodic acid (VWR 29460) for 10 minutes, washed in distilled water and
transferred to Schiff's reagent (Thermo Fisher Scientific 88017) for 10 minutes,
followed by washing in distilled water for 5 minutes. Sections were then routinely
dried, cleared and mounted.

644

# 645 Western blotting

646 Dissected anterior pituitaries were flash frozen in liquid nitrogen and stored at -80°C.

647 Frozen pituitaries were each lysed in 30μl of lysis buffer (5mM Tris, 150mM NaCl,

648 1% protease and phosphatase inhibitor (Abcam ab201119), 5μM EDTA, 0.1% Triton-

649 X, pH7.6) and sonicated at 40% power, twice for ten cycles of: two seconds on/two

650 seconds off, using a Vibra-Cell Processor (Sonics). Protein concentration was

determined using the Pierce BCA protein assay kit (Thermo #23227) and all samples

652 were diluted to 4mg/ml in Laemmli buffer (Biorad #161-0747). Proteins were

denatured at 95°C for 5 minutes. Samples were run on a 10% Mini-PROTEAN TGX

polyacrylamide gel (BioRad #4561033), then transferred using Trans-Blot Turbo

transfer machine (BioRad) onto polyvinylidene difluoride membranes (BioRad

41704156). Membranes were blocked with 5% non-fat dairy milk (NFDM) in TBST

657 (20mM Tris, 150mM NaCl, 0.1% Tween-20, pH7.6), cut, then incubated with

primary antibodies overnight at 4°C as follows: anti-STK3/STK4 (1:5000, Bethyl

Laboratories #A300-466A) or Cyclophilin B (1:1000, R&D Systems #MAB5410) in

660 5%NFDM. The next day, membranes were washed in TBST, incubated with

secondary antibodies HRP-conjugated anti-Rabbit (1:2000, Cell Signaling #7074) or

- 662 HRP-conjugated anti-Mouse (1:2000, Cell Signaling #7076) in 5% NFDM for 1hr at
- 663 room temperature. After washing in TBST, membranes were treated with Clarity
- 664 Western ECL substrate (Biorad #170-5060) and bands visualised using the ChemiDoc
- 665 Touch Imaging System (BioRad). Protein abundance was analysed using ImageLabs
- 666 (BioRad).
- 667

## 668 Imaging

- 669 Wholemount images were taken with a MZ10 F Stereomicroscope (Leica
- 670 Microsystems), using a DFC3000 G camera (Leica Microsystems). For bright field
- 671 images, stained slides were scanned with Nanozoomer-XR Digital slide scanner
- 672 (Hamamatsu) and images processed using Nanozoomer Digital Pathology View.
- 673 Fluorescent staining was imaged with a TCS SP5 confocal microscope (Leica
- 674 Microsystems) and images processed using Fiji <sup>62</sup>.
- 675

# 676 Quantifications and Statistics

677 Cell counts were performed manually using Fiji cell counter plug-in; 5-10 fields were
678 counted per sample, totalling over 1500 nuclei, across 3-7 pituitaries. Statistical

analyses and graphs were generated in GraphPad Prism (GraphPad Software) and the

- 680 following tests were performed to determine significance: Student's *t*-tests between
- controls and mutants for Figures 1D, 2D, S1bD, S1bE (n=3 of each genotype), S4
- 682 (n=4 of each genotype) and 5C (n=4-5 of each genotype); unpaired *t*-test for Figures
- 683 S2bA (n=3 per genotype) and S2bF (n=6 sections across two samples per genotype);
- 684 two-tailed *t*-test for Figure 3C (n=3 controls, 7 mutants); two-way ANOVA with
- 685 Sidak's multiple-comparison test for Figures 5F (n=4-5 of each genotype). For
- 686 quantification of target expression by RNAscope mRNA in situ hybridisation (Figure

- 687 S5), the area of positive staining (red fluorescence) from 4 $\mu$ m sections was
- determined from images using thresholding in Fiji, and quantified as a percentage of
- total pituitary area in the same image. For statistical testing, one-way ANOVAs with
- 690 Tukey's multiple comparisons were performed (n=4 mutants per genotype). Error
- bars in graphs show  $\pm$  standard error of the mean, unless otherwise indicated.
- 692 Quantification of STK3/4 by western blot was carried out on 2 control
- 693  $(Stk3^{fl/fl}; Stk4^{fl/fl})$  and 3 mutant  $(Hesx1^{Cre/+}; Stk3^{fl/fl}; Stk4^{fl/fl})$  samples. A Student's t-test
- 694 was carried out on normalised band intensities. Chi-squared tests were used to
- 695 determine significant deviations of observed from expected genotypes presented as
- tables in Supplementary Files 1 and 2.
- 697
- 698
- 699 FIGURE LEGENDS

### Figure 1 Regulation of YAP is required for normal morphogenesis and lineage commitment during pituitary development.

- 702 A. Schematic outlining the time course of doxycycline (DOX) treatment administered
- to pregnant dams from  $Hesx1^{Cre/+} \propto R26^{rtTA/rtTA}$ ;  $Colla1^{tetO-Yap/tetO-Yap}$  crosses for the
- embryonic induction of YAP(S127A) expression in  $Hesx1^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-}$
- 705 *Yap/+* (YAP-TetO) mutant embryos as well as controls that do not express
- 706 YAP(S127A) ( $Hesx1^{+/+}$ ; $R26^{rtTA/+}$ ; $Colla1^{tetO-Yap/+}$  controls shown here). **B.**
- 707 Immunofluorescence staining against YAP and TAZ on frontal pituitary sections at
- 708 15.5dpc confirms accumulation of YAP protein in YAP-TetO compared to control
- sections, but no increase in TAZ levels. RNAscope mRNA in situ hybridisation
- against the YAP/TAZ target *Cyr61* confirms an increase in transcripts in the anterior
- 711 pituitary as well as the hypothalamus where the Cre is also active (arrows). C.
- 712 Haematoxylin and eosin staining of frontal pituitary sections from 15.5dpc control
- and YAP-TetO embryos showing pituitary dysmorphology in mutants.
- 714 Immunofluorescence staining for LHX3 to mark anterior pituitary tissue and SOX2 to
- 715 mark pituitary progenitors shows the persistence of SOX2 protein in lateral regions of

- the gland in YAP-TetO mutants (arrowheads) when they have lost SOX2 expression
- 717 in controls (arrows) (magnified boxed region in SOX2, corresponding to dashed box
- 718 in LHX3). **D.** Immunofluorescence staining for lineage-committed progenitor markers
- 719 PIT1, TPIT and SF1 reveals very few cells expressing commitment markers in YAP-
- 720 TetO compared to control. Graph showing quantification of committed cells of the
- three anterior pituitary endocrine lineages, positive for PIT1, TPIT and SF1, as a
- 722 percentage of total nuclei of  $Hesx1^{+/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-Yap/+}$  control and
- *Hesx1*<sup>Cre/+</sup>; R26<sup>rtTA/+</sup>; Colla1<sup><math>tetO-Yap/+</sup> (YAP-TetO) mutant pituitaries at 15.5dpc</sup></sup>
- 724 (Student's *t*-test; PIT1: *P*<0.0001 (\*\*\*\*), TPIT: *P*=0.0012 (\*\*), SF1: *P*=0.0021 (\*\*)).
- Scale bars 100µm, 50µm in magnified boxed regions in C. See also figure
- supplements 1 and 2.
- 727

### Figure 2 Pituitary-specific deletion of *Lats1* and *Lats2* during development leads to pituitary overgrowth and defects in lineage commitment.

- 730 **A.** Haematoxylin and eosin staining on sagittal sections from
- 731  $Hesxl^{Cre/+};Latsl^{fl/fl};Lats2^{fl/fl}$  (mutant) and  $Hesxl^{+/+};Latsl^{fl/fl};Lats2^{fl/fl}$  (control)
- embryos at 13.5dpc reveals anterior pituitary dysmorphology and overgrowth in
- 733 mutants (dashed outline). Immunofluorescence staining for TAZ, YAP and pYAP
- reveals accumulation of TAZ and YAP in overgrown mutant tissue (arrowheads,
- normal epithelial expression indicated by arrows in control) and lack of staining for
- 736 pYAP (S127). Immunofluorescence for SOX2 shows the presence of SOX2+
- 737 progenitors throughout the abnormal tissue in mutants. **B.** Immunofluorescence
- staining for late progenitor marker SOX9 shows localisation in few cells of the
- 739 pituitary of mutants at 13.5dpc. Immunofluorescence staining for Ki-67 indicates
- 740 cycling cells throughout the mutant pituitary. C. Immunofluorescence staining for
- 741 SOX2 and Endomucin (EMCN) on frontal pituitary sections at 15.5dpc shows
- expansion of the SOX2+ progenitor compartment compared to controls and a
- reduction in vasculature marked by Endomucin. Immunofluorescence for non-
- phosphorylated (Active) YAP shows strong expression throughout the mutant gland
- compared to the control. Areas of necrosis in mutant tissue indicated by asterisks.
- 746 Ventral overgrowth extending into the oral cavity between the condensing sphenoid
- bone indicated by arrows. **D.** Immunofluorescence staining for lineage-committed
- 748 progenitor markers PIT1, TPIT and SF1 reveals only sporadic cells expressing

- 749 commitment markers in  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/fl}$  mutants compared to controls.
- 750 Boxes showing magnified regions. Dashed lines demarcate anterior pituitary tissue.
- 751 Graph showing quantification of committed cells of the three anterior pituitary
- endocrine lineages, positive for PIT1, TPIT and SF1, as a percentage of total nuclei of
- 753  $Hesxl^{+/+};Latsl^{fl/fl};Lats2^{fl/fl}$  control and  $Hesxl^{Cre/+};Latsl^{fl/fl};Lats2^{fl/fl}$  mutant pituitaries
- 754 at 15.5dpc (Student's *t*-test; PIT1: *P*<0.0001 (\*\*\*\*), TPIT: *P*=0.007 (\*\*), SF1:
- 755 P>0.05). Scale bars 100µm. See also figure supplement 2.
- 756

### 757 Figure 3 Pituitary specific loss of *Lats1* leads to tumour formation.

- 758 A. Haematoxylin and eosin staining of frontal sections from
- 759  $Hesx1^{Cre/+};Lats1^{fl/fl};Lats2^{fl/+}$  (mutant) and control pituitaries at P56 demonstrates
- 760 overgrown tumourigenic regions in mutants. These show focal necrosis, cysts and a
- 761 squamous morphology (magnified regions) not seen in controls. Asterisk indicates
- necrosis. **B.** Immunofluorescence staining for TAZ, YAP and pYAP(S127) show
- accumulation of TAZ and YAP but not pYAP in the mutant but not in the control.
- 764 RNAscope mRNA *in situ* hybridisation against YAP/TAZ targets *Ctgf* and *Cyr61*
- reveals an increase in transcripts on mutant tissue compared to control. C. Graph of
- the proliferation index in control and mutant samples at P56 shows a significant
- increase in cycling cells in the  $Hesx1^{Cre/+}$ ;  $Lats1^{fl/fl}$ ;  $Lats2^{fl/+}$  mutant pituitaries
- compared to controls (control percentage Ki-67: 2.967±1.2 SEM, n=3; mutant:
- 769 15.46 $\pm$ 2.74 n=7. *P*=0.0217 (\*), two-tailed *t*-test). Images show representative
- examples of Ki-67 immunofluorescence staining. D. Immunofluorescence staining for
- 1771 lineage-committed progenitor markers PIT1, TPIT and SF1 shows the near absence of
- committed cells in tumours. **E.** Immunofluorescence staining for pituitary stem cell
- markers SOX2 and SOX9 reveal that tumour lesions have abundant positive cells
- compared to the control, whilst Endomucin (EMCN) staining shows poor
- vascularisation. **F.** The marginal zone epithelium of  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$
- 776 mutant pituitaries develops invaginations as seen by haematoxylin and eosin staining.
- 777 Immunofluorescence staining against SOX2 shows the maintenance of a single-
- 1778 layered epithelium. Scale bars 100 μm. Boxes indicate magnified regions. See also
- figure supplement 1.
- 780

## Figure 4 SOX2+ pituitary stem cells are the cell-of-origin of tumours generated in the absence of *Lats1*.

- 783 A. Schematic outlining the experimental time line of inductions in
- 784  $Sox2^{CreERT2/+}$ ; Lats  $I^{fl/fl}$ ; Lats  $2^{fl/+}$  (mutant) and  $Sox2^{+/+}$ ; Lats  $I^{fl/fl}$ ; Lats  $2^{fl/+}$  (control)
- animals. Representative images of haematoxylin and eosin staining of frontal sections
- of control and mutant pituitaries at P35, revealing a hyperplastic anterior pituitary in
- 787 the mutant with areas of necrosis (asterisks). **B.** Immunofluorescence staining reveals
- tumourigenic lesions in  $Sox2^{CreERT2/+}$ ; Lats  $I^{fl/fl}$ ; Lats  $2^{fl/+}$  that display increased levels of
- 789 TAZ and YAP staining compared to the control. C. RNAscope mRNA in situ
- hybridisation against *Ctgf* and *Cyr61* shows elevated transcripts in tumourigenic
- resions. Insets (i) and (ii) show invaginations in the epithelium of the mutant. **D.**
- 792 Immunofluorescence staining for lineage-committed progenitor markers PIT1, TPIT
- and SF1 showing a reduction in staining in tumourigenic lesions compared to control
- 794 pituitaries. E. Lineage tracing of SOX2+ cells in
- 795  $Sox2^{CreERT2/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}R26^{mTmG/+}$  reveals that tumour regions accumulating
- 796 YAP as seen by immunofluorescence, are composed of GFP+ cells at P35. Scale bars
- 500μm in A; 100μm in B, D, E; 250μm in C. See also figure supplement 1.
- 798

## Figure 5 Postnatal expression of constitutively active YAP increases leads to an activation of SOX2+ pituitary stem cells.

- 801 **A.** Schematic outlining the time course of doxycycline (DOX) treatment administered 802 to  $Hesx1^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-Yap/+}$  (YAP-TetO) and  $Hesx1^{+/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-Yap/+}$
- 803  $Y_{ap/+}$  controls to drive expression of YAP-S127A in mutant pituitaries. **B.** At P42 (3)
- 804 weeks of treatment), immunofluorescence staining on frontal anterior pituitary
- 805 sections detects strong total YAP expression in YAP-TetO mutants compared to the
- so control and no increase in pYAP(S127). Immunofluorescence for SOX2 and SOX9
- 807 reveals an expanded population of stem cells in YAP-TetO compared to control
- 808 (quantification in F). C. Graph showing the percentage of double Ki-67+SOX2+ cells
- 809 as a proportion of the total SOX2+ (P=0.027 (\*)) or Ki-67+ (P=0.006 (\*\*))
- 810 populations at P42 (n=3 pituitaries per genotype). There is an increase in the numbers
- 811 of cycling SOX2 cells in YAP-TetO mutant compared to controls. The image shows a
- 812 representative example of double immunofluorescence staining against Ki-67 and
- 813 SOX2 in a control and YAP-TetO section. **D.** Schematic outlining the time course of
- 814 doxycycline (DOX) treatment administered to  $Hesx1^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Collal^{tetO-Yap/+}$
- 815 (YAP-TetO) and  $Hesx1^{+/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-Yap/+}$  controls to drive expression of
- 816 YAP-S127A in mutant pituitaries for three weeks, followed by a three-week recovery

- 817 period in the absence of DOX. E. Immunofluorescence staining against YAP, SOX2
- 818 and SOX9 on control and YAP-TetO pituitaries treated as in D, shows comparable
- 819 expression of YAP, SOX2 and SOX9 between genotypes. F. Graph of quantification
- 820 of SOX2+ cells as a percentage of total nuclei in control and YAP-TetO pituitaries at
- 821 P42 *P*=0.0014 (\*\*); P63 *P*=0.0044 (\*\*); P105 *P*<0.0001(\*\*\*\*) (n=3 pituitaries per
- 822 genotype). Following the Recovery treatment scheme in D, there is no significant
- 823 difference in the numbers of SOX2+ cells between genotypes. G. Schematic outlining
- the time course of tamoxifen induction and doxycycline (DOX) treatment
- 825 administered to  $Sox2^{CreERT2/+}$ ;  $R26^{rtTA/mTmG}$ ;  $Collal^{tetO-Yap/+}$  (mutant) and
- 826  $Sox2^{CreERT2/+}$ ;  $R26^{mTmG/+}$ ;  $Collal^{+/+}$  (control) animals to drive expression of YAP-
- 827 S127A in SOX2+ cells of mutants. H. Lineage tracing of SOX2+ cells and
- 828 immunofluorescence staining against SOX2 and GFP shows an expansion of GFP+
- 829 cells compared to controls at P63, where a proportion of cells are double-labelled. I.
- 830 Immunofluorescence staining against commitment markers PIT1, SF1 and terminal
- 831 differentiation marker ACTH (TPIT lineage) together with antibodies against GFP
- 832 detects double-labelled cells (arrows) across all three lineages in
- 833  $Sox2^{CreERT2/+}$ ;  $R26^{rtTA/mTmG}$ ;  $Collal^{tetO-Yap/+}$  pituitaries following the recovery period.
- 834 Graph of quantification of GFP+;PIT1+, GFP+;SF1+ and GFP+;ACTH+ cells as a
- 835 percentage of total GFP+ cells in  $Sox2^{CreERT2/+}$ ;  $R26^{rtTA/mTmG}$ ;  $Collal^{tetO-Yap/+}$  pituitaries
- at P63. Scale bars 100 $\mu$ m. Data in C. and F. represented as mean  $\pm$  SEM, analysed
- 837 with Two-Way ANOVA with Sidak's multiple comparisons. See also figure
- supplement 1.
- 839

## Figure 6 Model of stem cell activity following regulation by the LATS/YAP/TAZ cascade in the anterior pituitary.

- 842 SOX2+ pituitary stem cells express YAP and TAZ (green spheres). During normal
  843 developmental and postnatal expansion (normal regulation), pituitary stem cells are
- 844 maintained as a balanced pool while generating endocrine cells of three committed
- 845 lineages (red, blue, yellow). Expression of constitutively active YAP-S127A in
- 846 pituitary stem cells leads to an elevation in target gene expression, an expansion of
- 847 pituitary stem cell numbers and maintenance of the SOX+ state, preventing lineage
- 848 commitment. When YAP-S127A expression ceases, commitment into the endocrine
- 849 lineages takes place. Genetic deletion of LATS kinases (LATS1 as well as one or two
- 850 copies of LATS2), results in YAP and TAZ accumulation, major elevation in target

gene expression, repression of lineage commitment, continued expansion of SOX2+

cells and tumour formation.

853

854 855

### 856 FIGURE SUPPLEMENT LEGENDS

857

## 858 Figure 1 – figure supplement 1 Regulation of YAP and TAZ during pituitary 859 development.

860 **A.** Hematoxylin and eosin staining on frontal sections through the pituitary from

- 861 control and YAP-TetO heads after DOX treatment from 5.5dpc until 15.5dpc. **B.**
- 862 Schematic outlining the time course of doxycycline (DOX) treatment administered to
- 863  $Hessl^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Collal^{tetO-Yap/+}$  (YAP-TetO) and  $Hessl^{+/+}$ ;  $R26^{rtTA/+}$ ;  $Collal^{tetO-Yap/+}$
- 864 *Yap/+* controls to drive expression of YAP-S127A in mutant pituitaries during
- 865 embryonic as well as postnatal development. **C.** Hematoxylin and eosin (H&E)
- staining of control and YAP-TetO pituitaries at P24. Higher magnification images
- show the presence of cysts in the YAP-TetO mutant. White arrows indicate cells with
- 868 enlarged nuclei surrounding the cysts and yellow arrows indicate ciliated cells. **D.**
- 869 Immunofluorescence staining against total YAP on frontal sections at P24 confirms
- accumulation of YAP protein in YAP-TetO compared to control sections, especially
- 871 in the ventral anterior lobe. Immunofluorescence staining against SOX2 shows an
- 872 expansion of SOX2+ epithelia lining cysts. E. Immunofluorescence staining for
- 873 lineage-committed progenitor markers PIT1, TPIT and of ACTH marking the SF1
- 874 lineage in control and YAP-TetO sections at P24. The number of SOX2+ and lineage-
- 875 committed cells is quantified in the graph below. Note there is a significant increase
- 876 in the proportion of SOX2+ cells in YAP-TetO mutants (Student's *t*-test, *P*<0.0001
- 877 (\*\*\*\*)), decrease in PIT1+ cells (Student's *t*-test,  $P \le 0.0002$  (\*\*\*)), increase in SF1+
- 878 cells (Student's *t*-test,  $P \le 0.0066$  (\*\*)) and no significant change in ACTH+ cells. **F**.
- 879 Immunofluorescence staining against Ki-67 marking cycling cells in control and
- 880 YAP-TetO sections at P24. Graph showing the percentage of Ki-67+ cells across total
- anterior pituitary cells. There is a trend towards a reduction in the proportion of
- 882 cycling cells in YAP-TetO mutants, which is not significant (Student's *t*-test,
- 883 *P*>0.05). **G.** Immunohistochemistry using antibodies against p63 and the AE1/AE3
- 884 cytokeratin cocktail in YAP-TetO mutants at P24 revealing positive cells lining the

- 885 cysts (arrowheads). H. Immunofluorescence staining using antibodies against
- ARL13B and Acetylated α-Tubulin, staining components of cilia, reveals ciliated
- cells lining the cysts. Staining for Alcian Blue and Period Acid Schiff (AB/PAS) to
- 888 differentiate between acidic and neutral mucins reveals royal blue-stained mucous
- cells lining the cysts. Scale bars 1mm in A, 500µm in C and 100µm in magnified
- panels in C,  $100\mu m$  in D, E, F and  $50\mu m$  in G and H.
- 891

### 892 Figure 1 – figure supplement 2 Regulation of YAP and TAZ during pituitary 893 development.

- A. Hematoxylin and eosin staining on sagittal pituitary sections of 13.5dpc
- 895  $Hesx l^{Cre/+}; Yap^{fl/fl}; Taz^{-/-}$  (mutant) and  $Hesx l^{+/+}; Yap^{fl/+}; Taz^{+/-}$  (control) showing
- 896 comparable morphology. **B.** Immunofluorescence staining using antibodies against
- 897 SOX2 in  $Hesx1^{Cre/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{-/-}$  and control at 13.5dpc (sagittal) and 16.5dpc
- 898 (frontal) showing the presence of SOX2+ cells in both genotypes. C.
- 899 Immunofluorescence staining for SOX2, Endomucin (EMCN) and active YAP in P28
- 900  $Hesx1^{Cre/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{-/-}$  and control pituitaries, identifies SOX2+ cells in regions that
- 901 are negative for active YAP (mice are null for TAZ) and normal vasculature. **D.**
- 902 Graph quantifying the percentage of SOX2+ cells expressing active YAP in control
- 903 and  $Hesx1^{Cre/+}$ ;  $Yap^{fl/fl}$ ;  $Taz^{-/-}$  mutant pituitaries at P28. There is a reduction in double-
- 904 positive cells in the mutant, which did not reach significance. **E.**
- 905 Immunofluorescence staining for lineage committed progenitor markers PIT1 and
- 906 SF1, as well as ACTH marking corticotrophs (TPIT lineage), reveals the presence and
- 907 normal localisation of cells from the three lineages in a P28  $Hesx1^{Cre/+}; Yap^{fl/fl}; Taz^{-/-}$
- 908 mutant. Scale bars 100µm.
- 909

### 910 Figure 2 – figure supplement 1 Pituitary-specific loss of *Stk3* and *Stk4* does not

- 911 affect SOX2 cell specification or lineage commitment.
- 912 A. Dorsal view of wholemount  $Hesx1^{Cre/+}$ ;  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl}$  (mutant) and  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl}$
- 913 (control) pituitaries at P0 showing comparable morphology and size at birth. B.
- 914 Western blot to determine levels of STK3 and STK4 proteins in
- 915  $Hesx1^{Cre/+}$ ;  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl}$  mutant pituitaries compared to controls at P35, using an
- 916 antibody against total STK3 and STK4 proteins. Comparison of STK3/4 band
- 917 intensities confirms a significant reduction in mutants (Student's *t*-test, *P*=0.00032

- 918 (\*\*\*)). STK3/4 bands normalised to the housekeeping protein Cyclophilin B. C.
- 919 Immunofluorescence staining using antibodies against SOX2, TAZ, Endomucin
- 920 (EMCN), YAP and pYAP at P0, indicating comparable staining between control and
- 921 mutant samples. **D.** Immunofluorescence staining against lineage commitment
- 922 markers PIT1, TPIT and SF1 shows normal lineage commitment in a
- 923  $Hesx1^{Cre/+}$ ;  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl}$  mutant pituitary compared to the control at P10. E.
- 924 Hematoxylin and eosin staining through frontal sections of  $Hesx1^{Cre/+}$ ;  $Stk3^{fl/fl}$ ;  $Stk4^{fl/fl}$
- 925 and control pituitaries at P70. AL: anterior lobe, IL: intermediate lobe, PL: posterior
- 926 lobe. Scale bars 100μm.
- 927

### Figure 2 – figure supplement 2 Isolated deletions of *Lats1* or *Lats2* in the pituitary do not affect development.

930 **A.** Hematoxylin and eosin staining of a sagittal section of  $Hesxl^{Cre/+}$ ; Lats  $l^{fl/fl}$  at

- 931 13.5dpc showing normal morphology (see Figure 2A for control). Dashed lines
- 932 demarcate developing Rathke's pouch. Immunofluorescence staining for TAZ and
- 933 YAP reveals a normal expression pattern and no gross protein accumulation (compare
- by to control, Figure 2A) **B.** Dorsal view of wholemount  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$  (mutant) and
- 935 *Hesx1<sup>Cre/+</sup>* (control) pituitaries at P0 showing comparable morphology and size at
- 936 birth. C. RNAscope mRNA in situ hybridisation against Lats2 shows an increase in
- 937 transcripts in the anterior pituitary following deletion of Lats1 (Hesx1<sup>Cre/+</sup>;Lats1<sup><math>fl/fl</sup>)</sup>
- 938 compared to control ( $HesxI^{Cre/+}$ ), where *Lats2* expression is barely detectable. **D**.
- Hematoxylin and eosin staining of a sagittal section of  $Hesx1^{Cre/+}$ ;  $Lats2^{fl/fl}$  at 13.5dpc
- showing normal morphology (see Figure 2A for control). Dashed lines demarcate
- 941 developing Rathke's pouch. E. Hematoxylin and eosin staining on frontal sections
- 942 through 15.5dpc embryonic heads of  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/fl}$  (mutant) and control
- 943 ( $Hesx1^{+/+}$ ;  $Lats1^{fl/fl}$ ;  $Lats2^{fl/fl}$ ) genotypes, at the levels indicated in the cartoon. Note the
- 944 hyperplastic pituitary at both axial levels, exerting mass effect on the brain. Asterisk
- 945 indicates necrosis. Graph showing quantification of pituitary size at 15.5dpc as
- 946 measured by the area occupied by the pituitary in matched histological sections
- 947 between control and mutant embryos.  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/fl}$  mutant pituitaries
- 948 are significantly larger (average 0.7195mm<sup>2</sup>) compared to controls (average
- 949 0.1994mm<sup>2</sup>) (Student's *t*-test, *P*=0.0003 (\*\*\*)). **F**. Quantification of Ki-67+ and
- 950 SOX9+ cells across the whole Rathke's pouch of  $Hesx1^{Cre/+}$  (control) and
- 951 *Hesx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup>* (mutant) pituitaries at 13.5dpc. There is a significant increase in

- 952 cycling cells in mutants, marked by Ki-67 (Student's *t*-test, *P*=0.0067 (\*\*)). The
- 953 proportion of SOX9+ cells is comparable between genotypes. Scale bars 100μm in A-
- 954 D, 1mm in E.
- 955

## Figure 3 – figure supplement 1 Analysis of tumourigenic lesions in postnatal pituitaries following pituitary-specific deletion of *Lats1*.

- 958 A. Immunofluorescence staining for TAZ and active YAP reveal lesions of
- 959 accumulation at P21 in  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$  compared to
- 960  $Hesx1^{+/+};Lats1^{fl/fl};Lats2^{fl/+}$  control. Immunofluorescence staining using antibodies
- 961 against SOX2 and Endomucin (EMCN) show these lesions are composed of SOX+
- 962 stem cells and have reduced vascularisation. B. Hematoxylin and eosin staining of
- 963 frontal sections from  $Hesx1^{Cre/+}$ ; Lats  $2^{fl/fl}$  and  $Hesx1^{Cre/+}$  control pituitaries at P56
- showing comparable histology. **C.** Immunofluorescence staining against SOX2 and
- 965 Endomucin on an intermediate lobe lesion (asterisk) in a  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$
- 966 pituitary compared to control. **D.** Immunofluorescence staining against DNA damage
- 967 marker gamma H2A.X showing positive cells in  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$  mutants.
- 968 E. P56 Immunohistochemistry using antibodies against p63 and the AE1/AE3
- 969 cytokeratin cocktail, both positive in pituitary carcinomas, showing abundant staining
- 970 in  $Hesxl^{Cre/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$  compared to control. Note that membrane staining
- 971 detected in controls is background for both antibodies. F. Immunohistochemistry
- 972 using antibodies against synaptophysin, neural-specific enolase (NSE) and
- 973 chromogranin demonstrate tumourigenic lesions in  $Hesx1^{Cre/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$  are
- 974 negative for adenoma markers. Lesions are negative for vimentin by
- 975 immunofluorescence staining, commonly marking spindle-cell oncocytoma in the
- 976 pituitary. Scale bars 100µm in A, C-F; 500µm in B. PL: posterior lobe, IL:
- 977 intermediate lobe, AL: anterior lobe.
- 978

### 979 Figure 4 – figure supplement 1 Analysis of tumourigenic lesions in postnatal

- 980 pituitaries following SOX2-specific deletion of Lats1.
- 981 A. Graph of quantification of lineage commitment markers PIT1, TPIT and SF1, as a
- 982 percentage of all anterior pituitary cells, in  $Sox2^{+/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$  (control) and
- 983  $Sox2^{CreERT2/+}$ ; Lats  $1^{fl/fl}$ ; Lats  $2^{fl/+}$  (mutant) pituitaries. There is a significant reduction in
- 984 the percentage of committed cells of all three lineages in mutants compared to
- 985 controls (Student's *t*-test; PIT1: *P*<0.0001 (\*\*\*\*), TPIT: *P*<0.0001 (\*\*\*\*), SF1:

- 986 P=0.004 (\*\*)). B. Immunohistochemistry using specific antibodies against p63 and
- 987 cytokeratin cocktail AE1/AE3 on frontal sections of  $Sox2^{CreERT2/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$
- 988 (mutant) and  $Sox2^{+/+}$ ; Lats  $I^{fl/fl}$ ; Lats  $2^{fl/+}$  (control) pituitaries at P35, revealing positive
- 989 staining in mutants. Note that the membrane staining in controls is background for
- both antibodies. C. Double immunofluorescence staining against total YAP and GFP,
- as well as SOX2 and GFP in consecutive sections of a tumourigenic lesion from
- 992  $Sox2^{CreERT2/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$ ; R26<sup>*mTmG/+*</sup> pituitaries at P35. Lineage tracing of
- 993 SOX2+ cells, detected using GFP reveals abundant staining in the tumour lesion,
- characterised by accumulation of YAP and SOX2+ cells (yellow arrowheads). Scale
- 995 bars 100μm.

996

- Figure 5 figure supplement 1 Postnatal expression of constitutively active YAP
  increases leads to an activation of SOX2+ pituitary stem cells.
- 999 A. Schematic outlining the time course of doxycycline (DOX) treatment administered
- 1000 to  $Hesx1^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-Yap/+}$  (YAP-TetO) and  $Hesx1^{+/+}$ ;  $R26^{rtTA/+}$ ;  $Colla1^{tetO-Yap/+}$
- 1001 *Yap/+* controls to drive expression of YAP-S127A in mutant pituitaries. Hematoxylin
- and eosin staining of control and YAP-TetO pituitaries at P42 (3 weeks treatment),
- 1003 P63 (6 weeks treatment) and P105 (12 weeks treatment). **B.** RNAscope mRNA *in situ*
- 1004 hybridisation against YAP targets *Cyr61* and *Ctgf* showing increased transcripts in
- 1005 YAP-TetO sections compared to controls at P42. C. Analysis of YAP-TetO mutants
- 1006 at P105: double immunofluorescence staining against SOX2 and Ki-67 reveals
- 1007 regions of expanded SOX2+;Ki-67- cells compared to the normal expression pattern
- 1008 in the control. This region is SOX9+, does not accumulate TAZ or YAP and
- 1009 expresses pYAP as does normal anterior pituitary epithelium. Immunofluorescence
- 1010 against PIT1 shows the absence of commitment to this lineage, a pattern not seen in
- 1011 the control. Hematoxylin and eosin staining in consecutive sections identifies this
- 1012 region, which does not have neoplastic features. **D.** Schematic outlining the time
- 1013 course of doxycycline (DOX) treatment administered to
- 1014  $Hessl^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Collal^{tetO-Yap/+}$  (YAP-TetO) and  $Hessl^{+/+}$ ;  $R26^{rtTA/+}$ ;  $Collal^{tetO-Yap/+}$
- 1015 *Yap/+* controls to drive expression of YAP-S127A in mutant pituitaries for three weeks,
- 1016 followed by a three-week recovery period in the absence of DOX. Hematoxylin and
- 1017 eosin staining of control and YAP-TetO pituitaries. RNAscope mRNA in situ
- 1018 hybridisation shows comparable levels of expression of targets *Cyr61* and *Ctgf*. E.
- 1019 Graph comparing total fluorescence of *Cyr61* and *Ctgf* by Fast Red RNAscope

1020 mRNA in situ hybridisation across sections from control,

 $Hess1^{Cre/+}$ ;  $R26^{rtTA/+}$ ;  $Collal^{tetO-Yap/+}$  (YAP-TetO) and  $Sox2^{CreERT2/+}$ ;  $Lats1^{fl/fl}$ ;  $Lats2^{fl/+}$ 1021

- 1022 anterior pituitaries, normalised for total anterior pituitary area. There is a significant
- increase in the expression of both targets in  $Sox2^{CreERT2/+}$ ; Lats  $l^{fl/fl}$ ; Lats  $2^{fl/+}$  pituitaries 1023
- compared to other genotypes (one-way ANOVA with Tukey's post hoc test; Control 1024
- $v Sox2^{CreERT2/+}$ : Lats  $I^{fl/fl}$ : Lats  $2^{fl/+}$ : P<0.0001 for Cvr61 (\*\*\*\*). P=0.001 for Ctgf (\*\*\*): 1025
- YAP-TetO v Sox2<sup>CreERT2/+</sup>;Lats1<sup>fl/fl</sup>;Lats2<sup>fl/+</sup>: P<0.0001 for Cvr61 (\*\*\*\*), P=0.0049 1026
- 1027 for Ctgf(\*\*)). Scale bars 250µm in A, 100µm in B-D.
- 1028
- 1029 SUPPLEMENTARY FILE LEGENDS
- 1030

### 1031 **Supplementary File 1**

- 1032 Table showing expected and observed frequency of genotypes from
- $Hesx 1^{Cre/+}; Yap^{fl/fl}; Taz^{+/-} \ge Yap^{fl/fl}; Taz^{+/-}$  at embryonic 15.5 dpc and postnatal day 0-2. 1033
- 1034 Embryonic: P=0.3471, Chi-square test (two tailed). Postnatal: P=0.0003 (\*\*\*), Chi-
- square test (two tailed). 1035
- 1036

### 1037 **Supplementary File 2**

- Table showing expected and observed frequency of genotypes from 1038
- $Hesx1^{Cre/+}$ ;  $Lats1^{fl/+}$ ;  $Lats2^{fl/+}$  x  $Lats1^{fl/fl}$ ;  $Lats2^{fl/fl}$  and  $Hesx1^{Cre/+}$ ;  $Lats1^{fl/+}$ ;  $Lats2^{fl/+}$  x 1039
- *Lats*  $l^{fl/fl}$ ; *Lats*  $2^{fl/+}$  at embryonic 15.5 dpc and postnatal day 0-2. Embryonic: P<0.0001 1040
- (\*\*\*\*), Chi-square test (two tailed). Postnatal: P<0.0001 (\*\*\*\*), Chi-square test (two 1041 1042 tailed).
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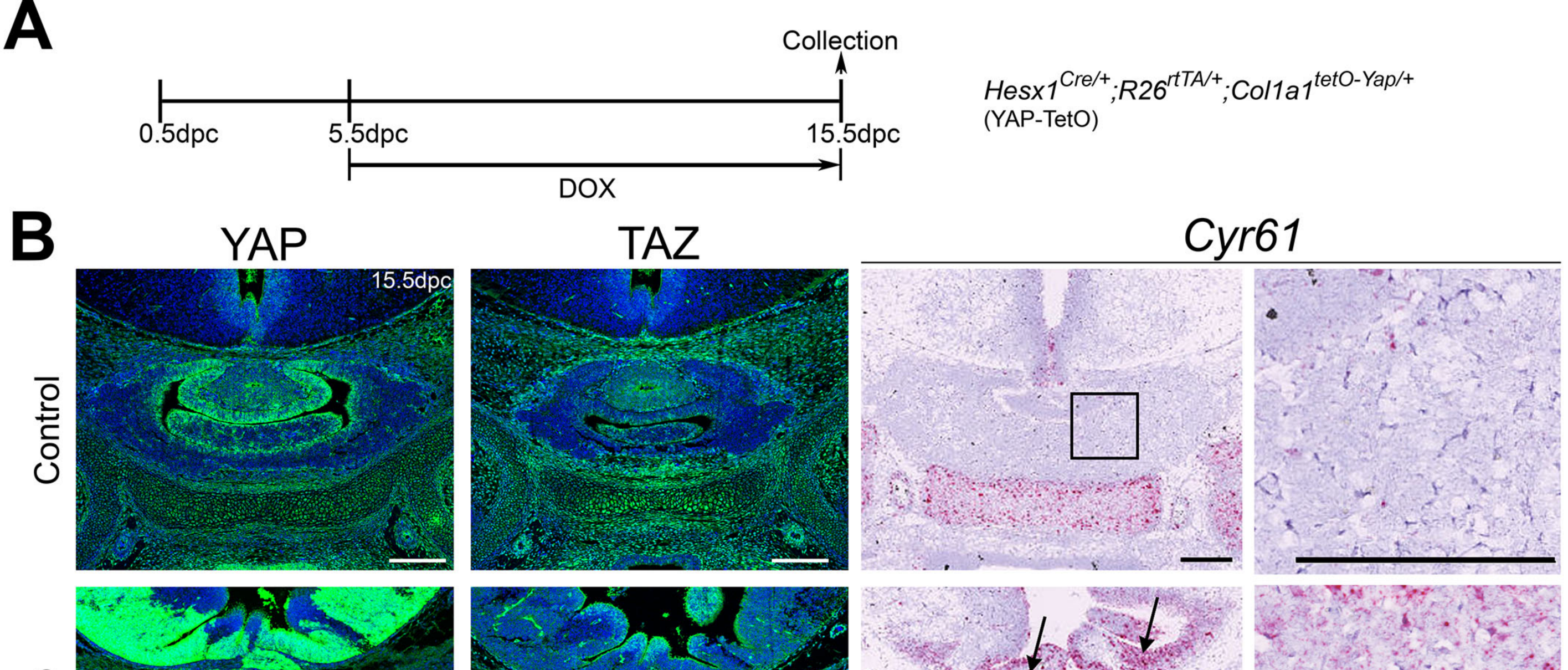
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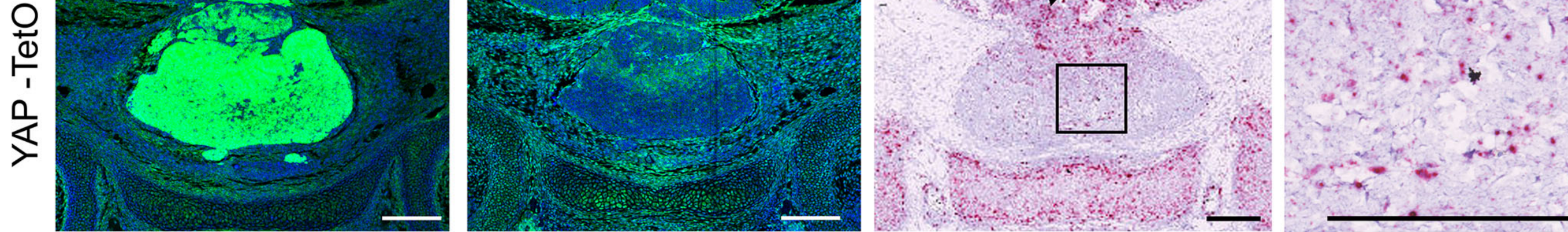
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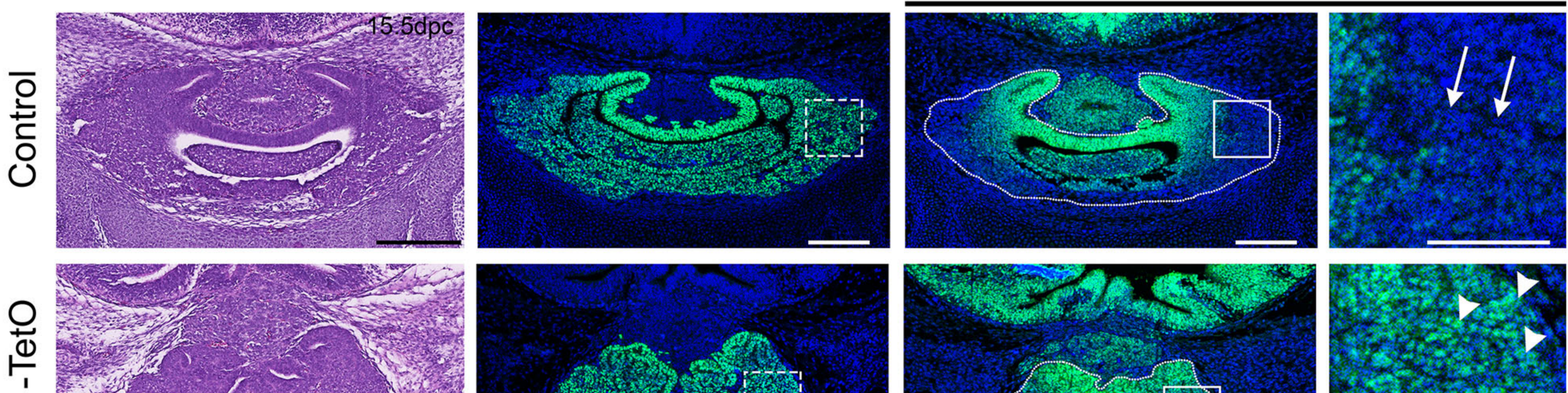


## H&E

С

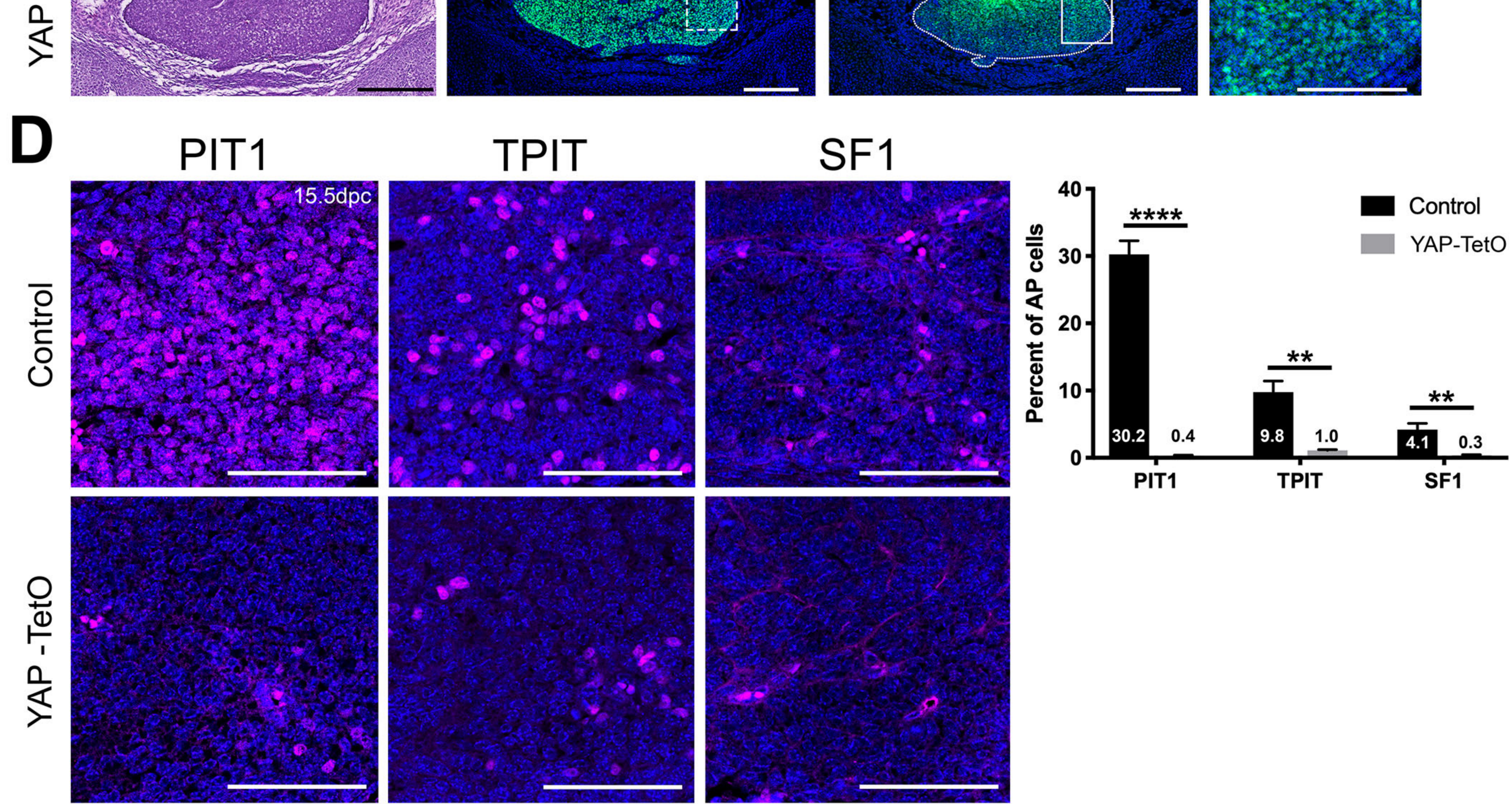
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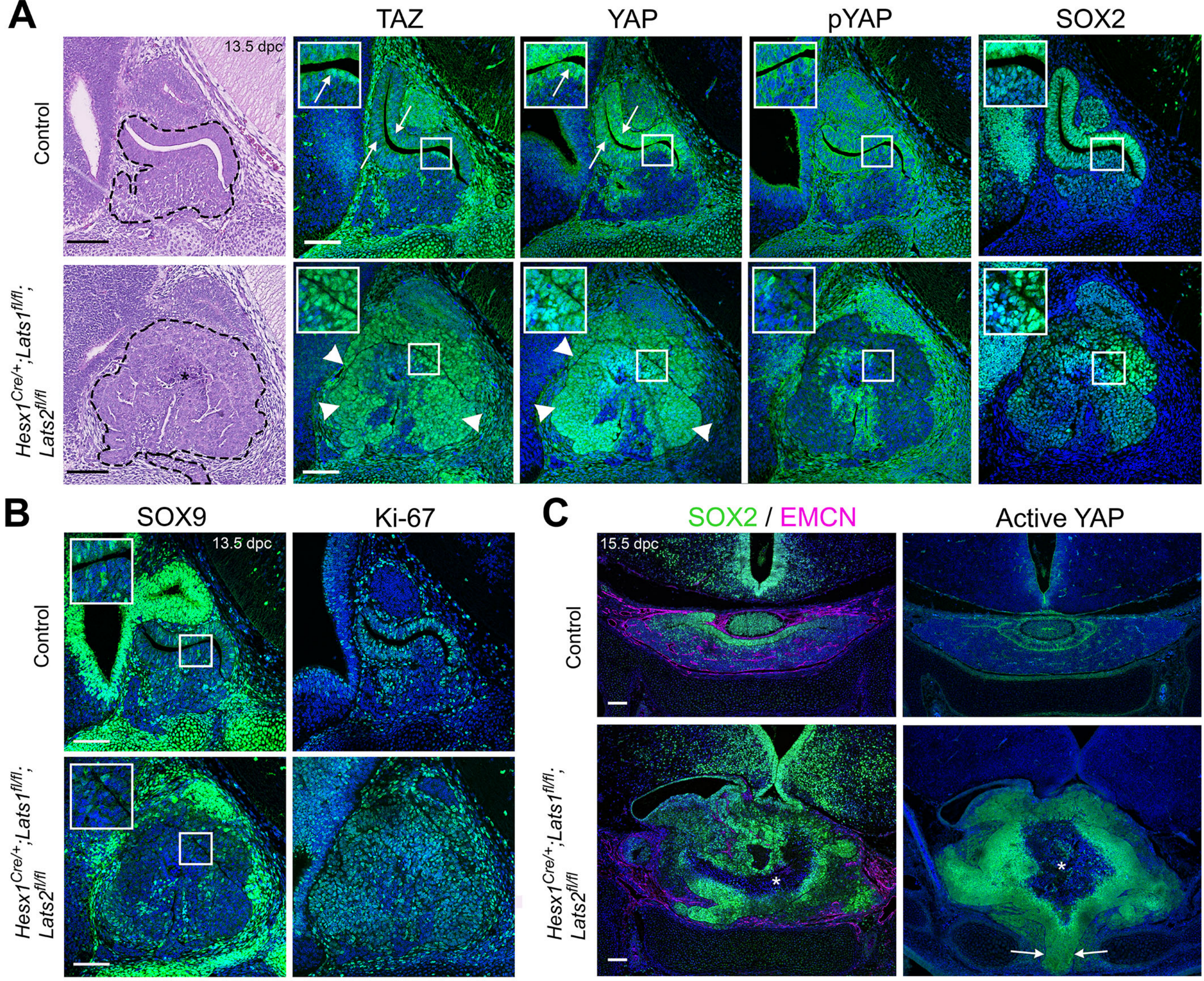






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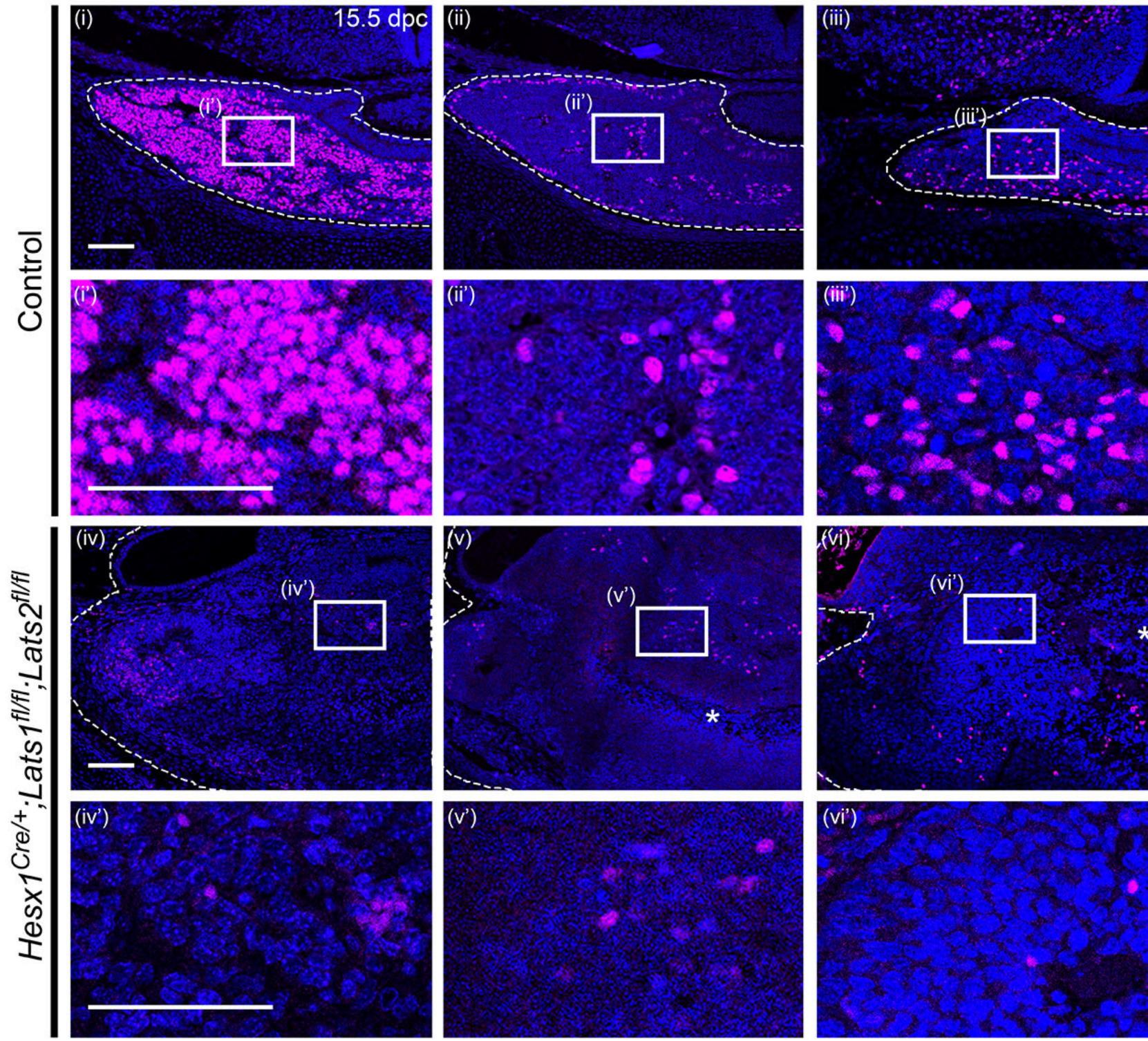




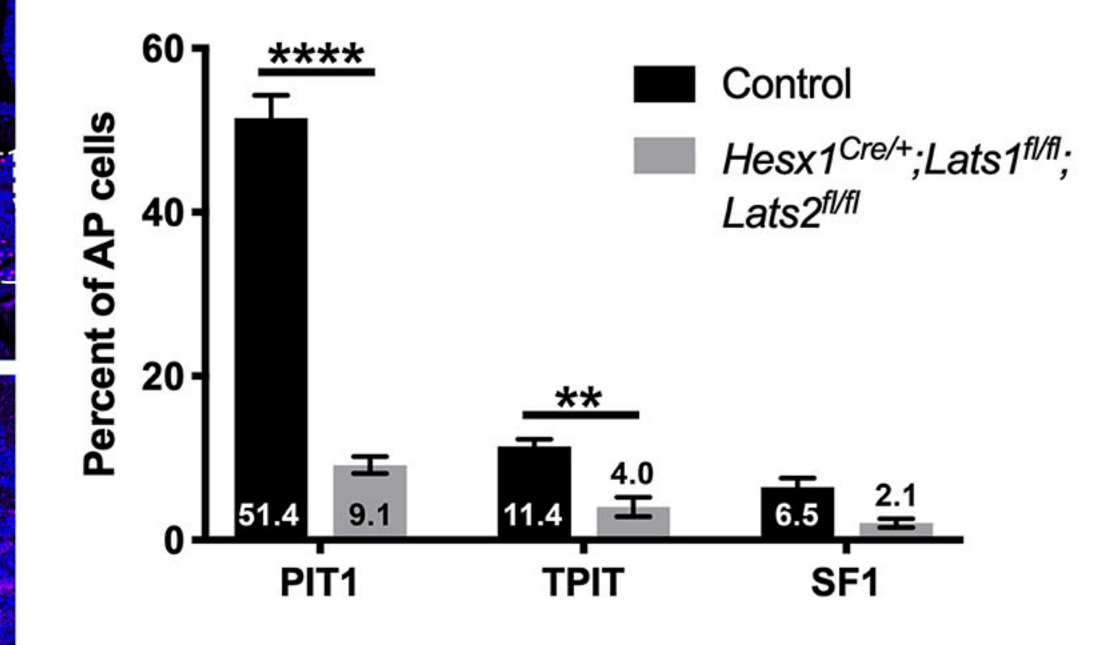
SF1

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PIT1



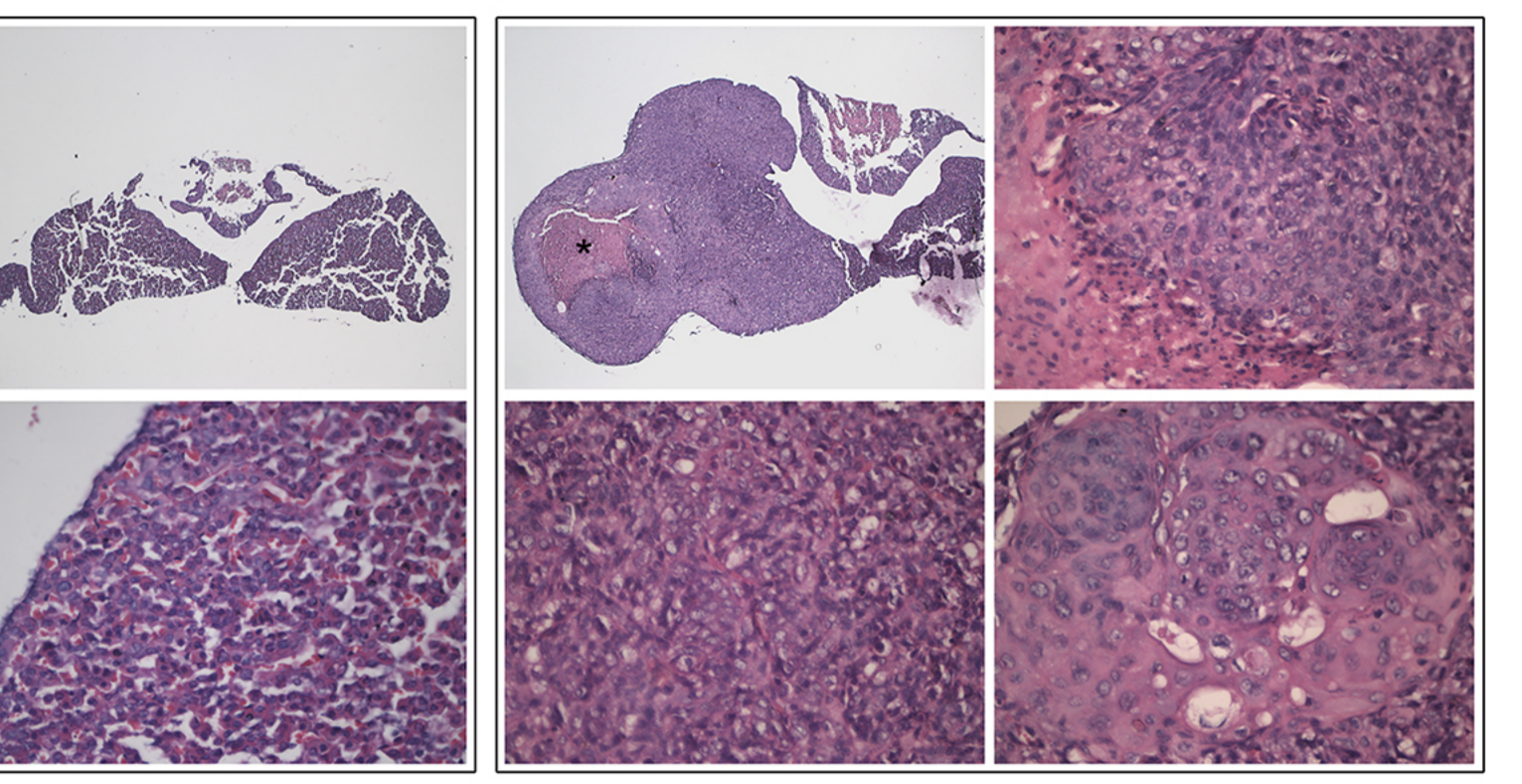
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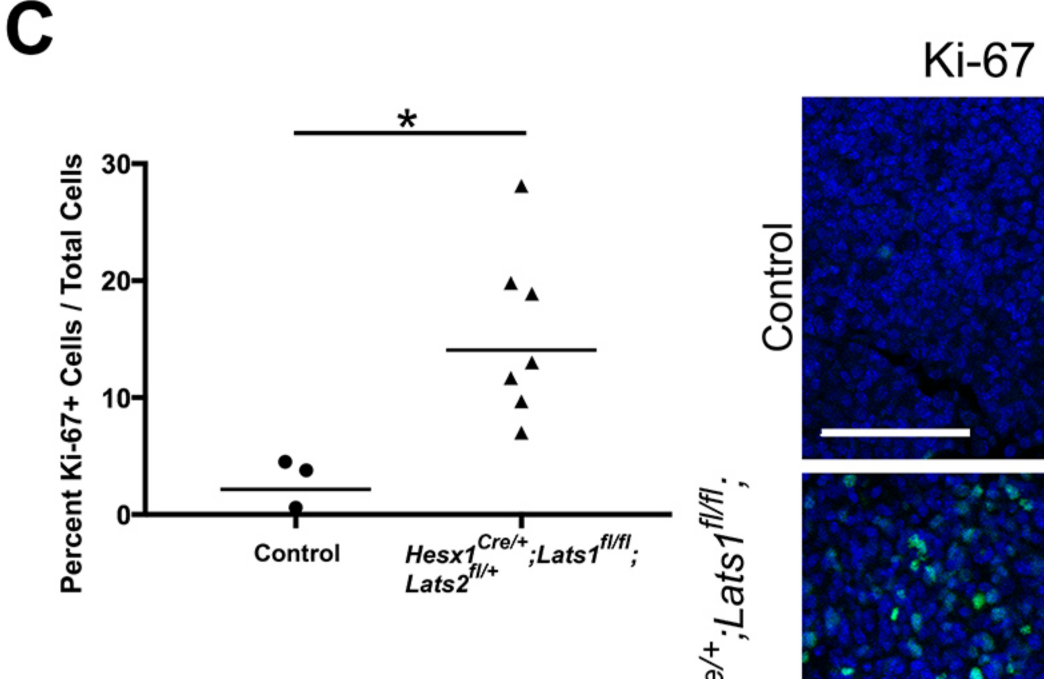




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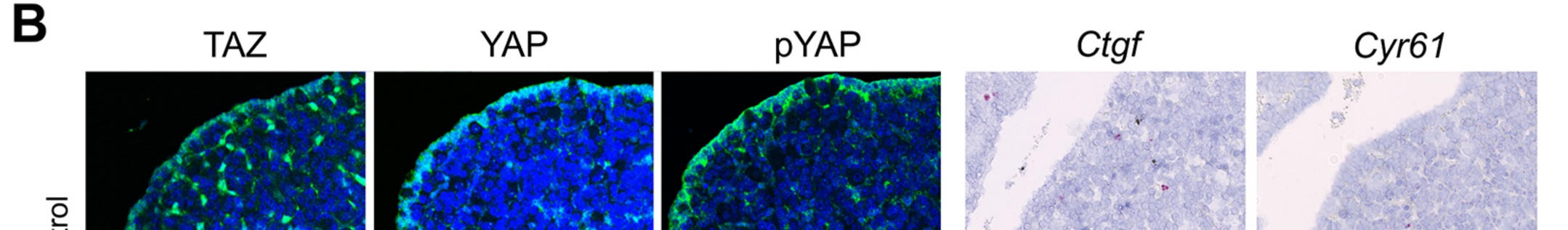


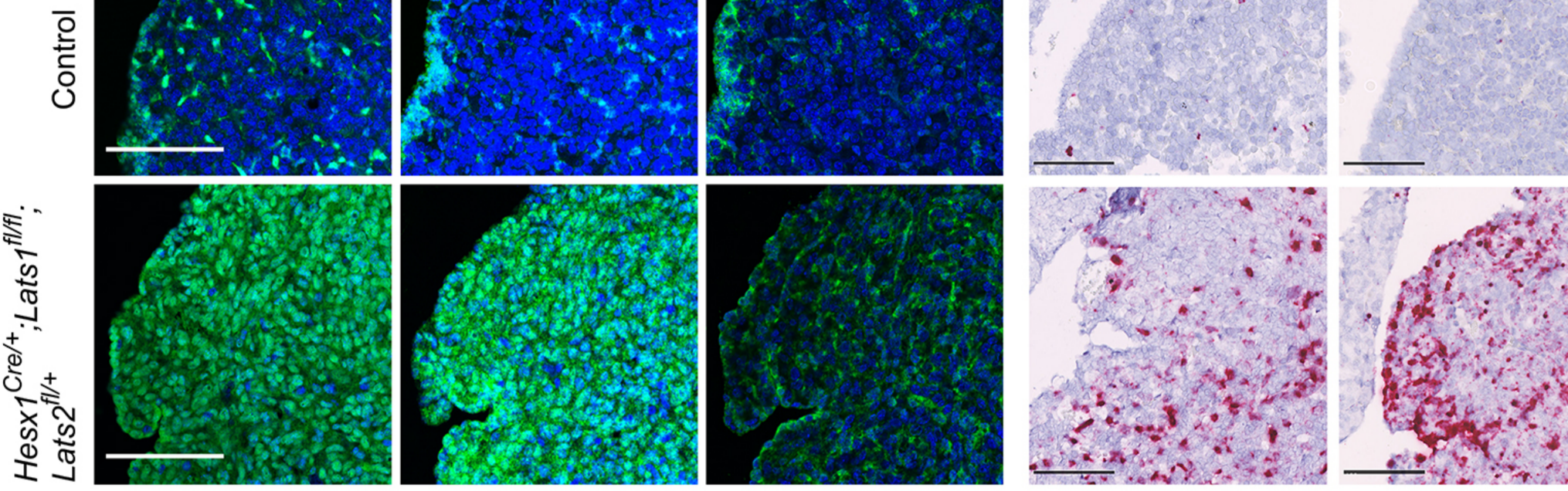


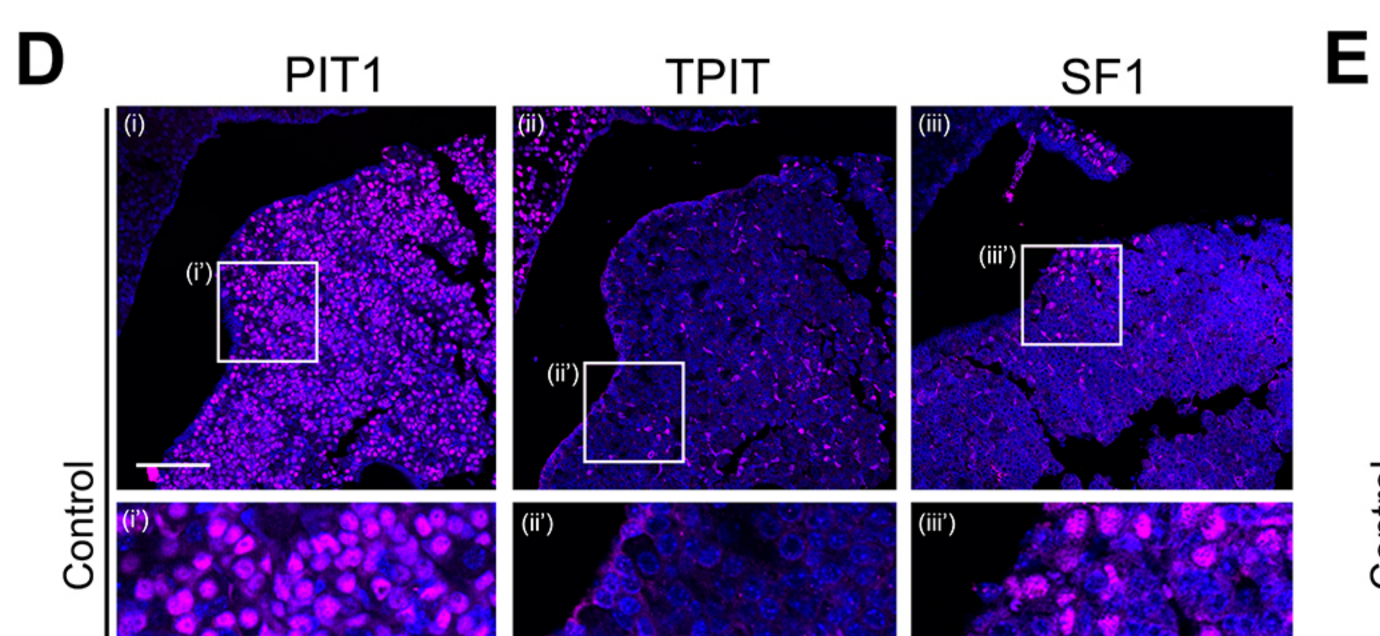
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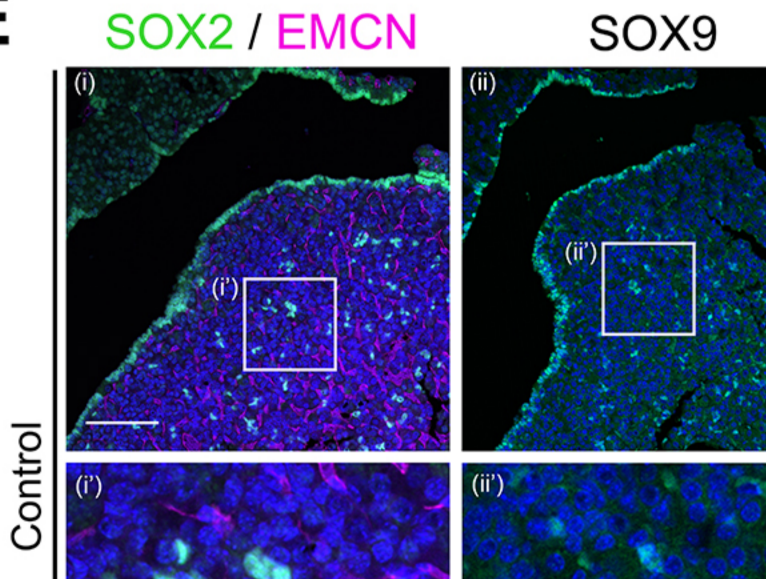
Hesx1

atsi



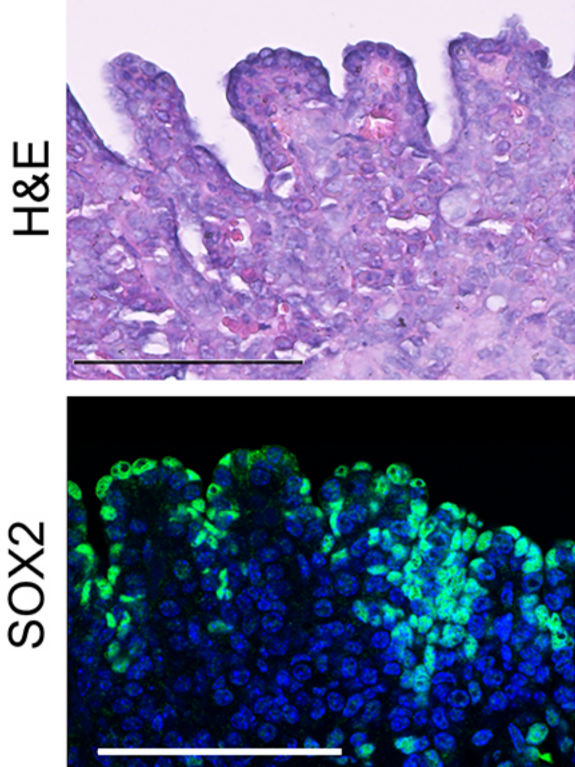




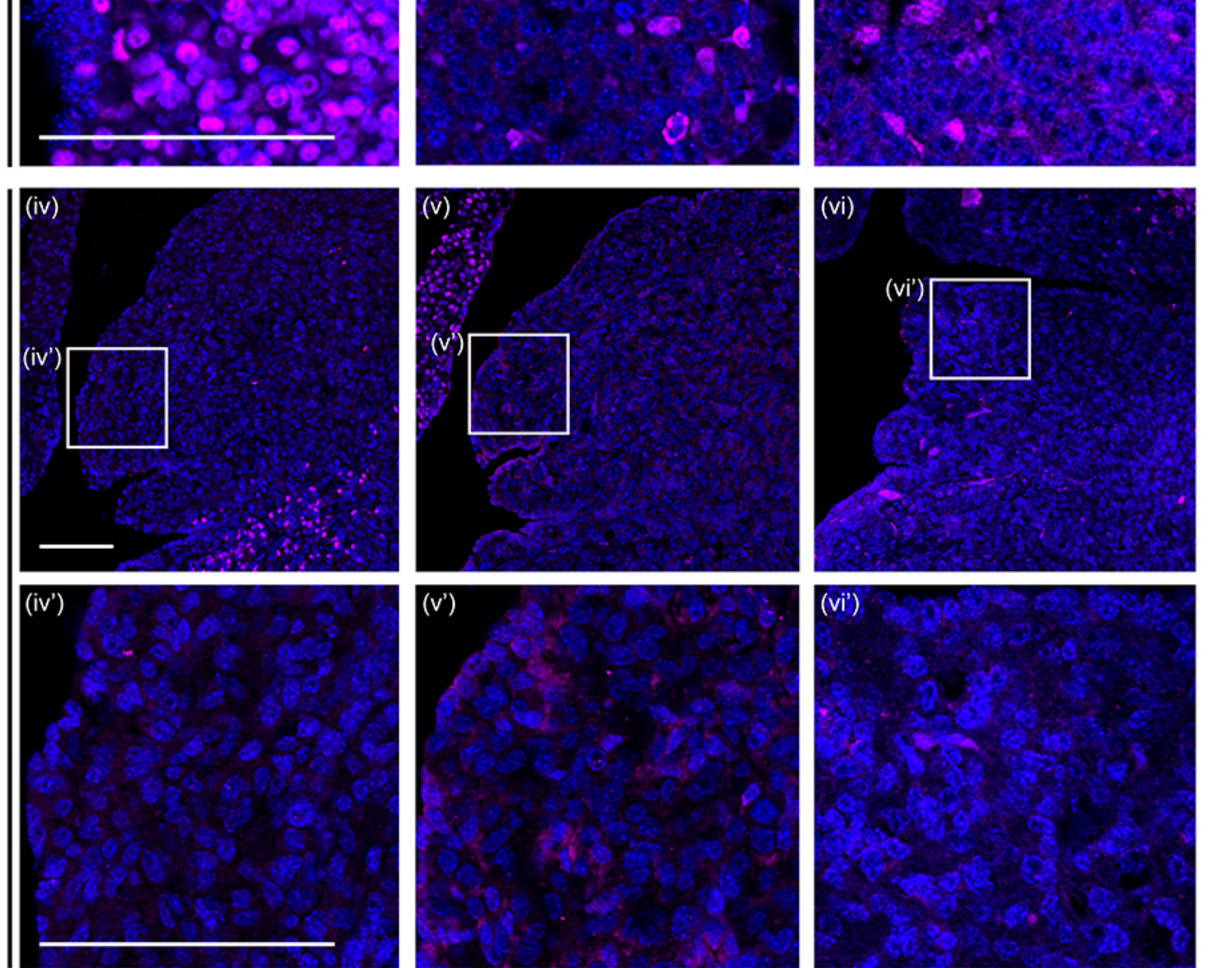


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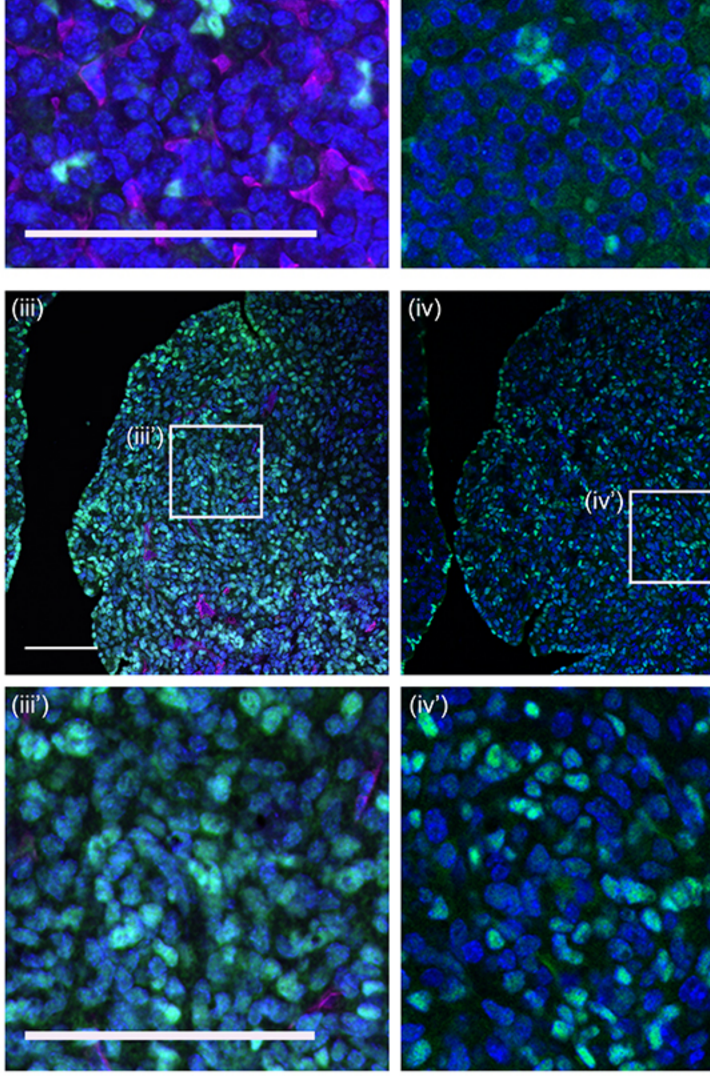
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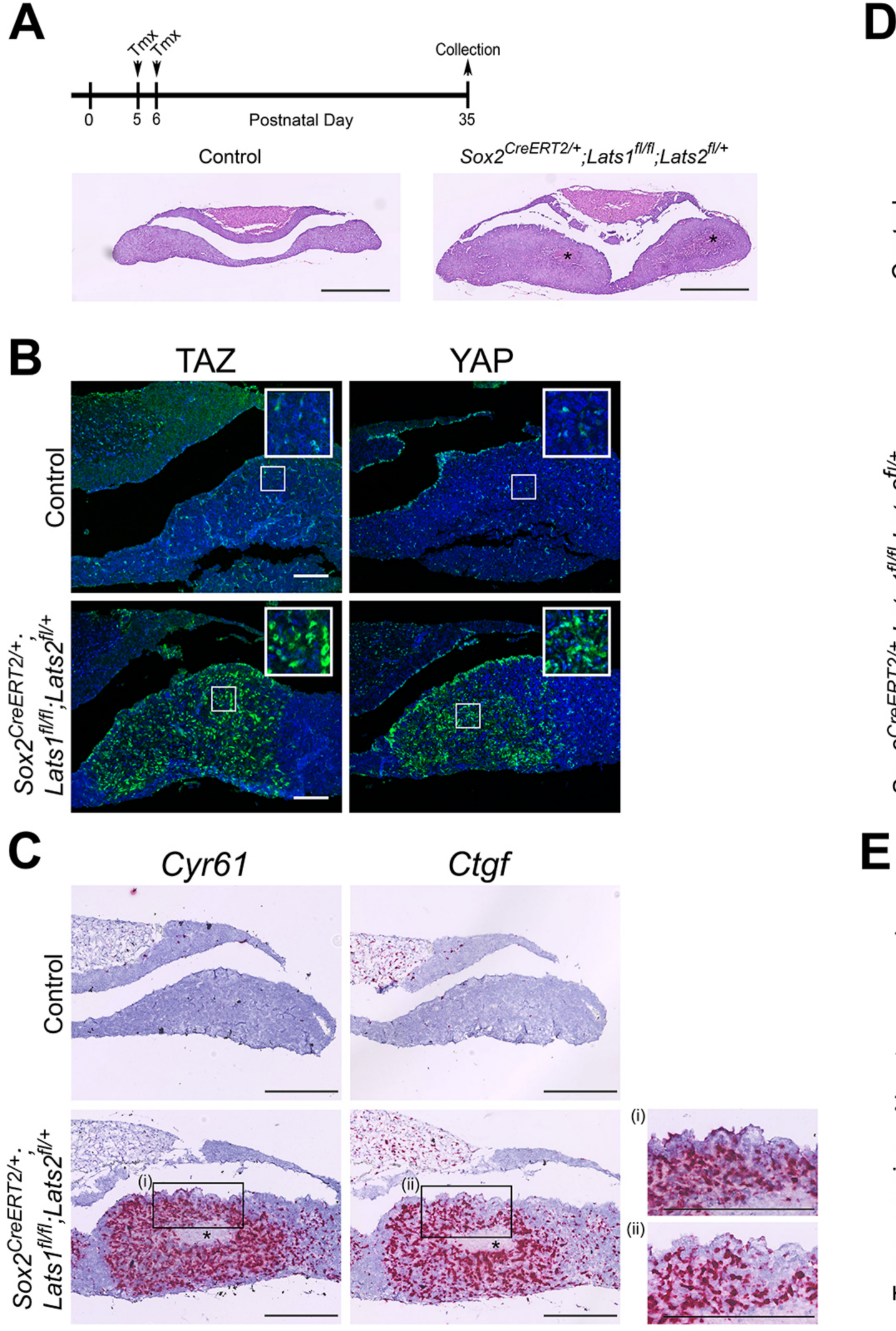
## .ats1<sup>fl/fl</sup>;Lats2<sup>fl/+</sup> Cre/ -Hesx

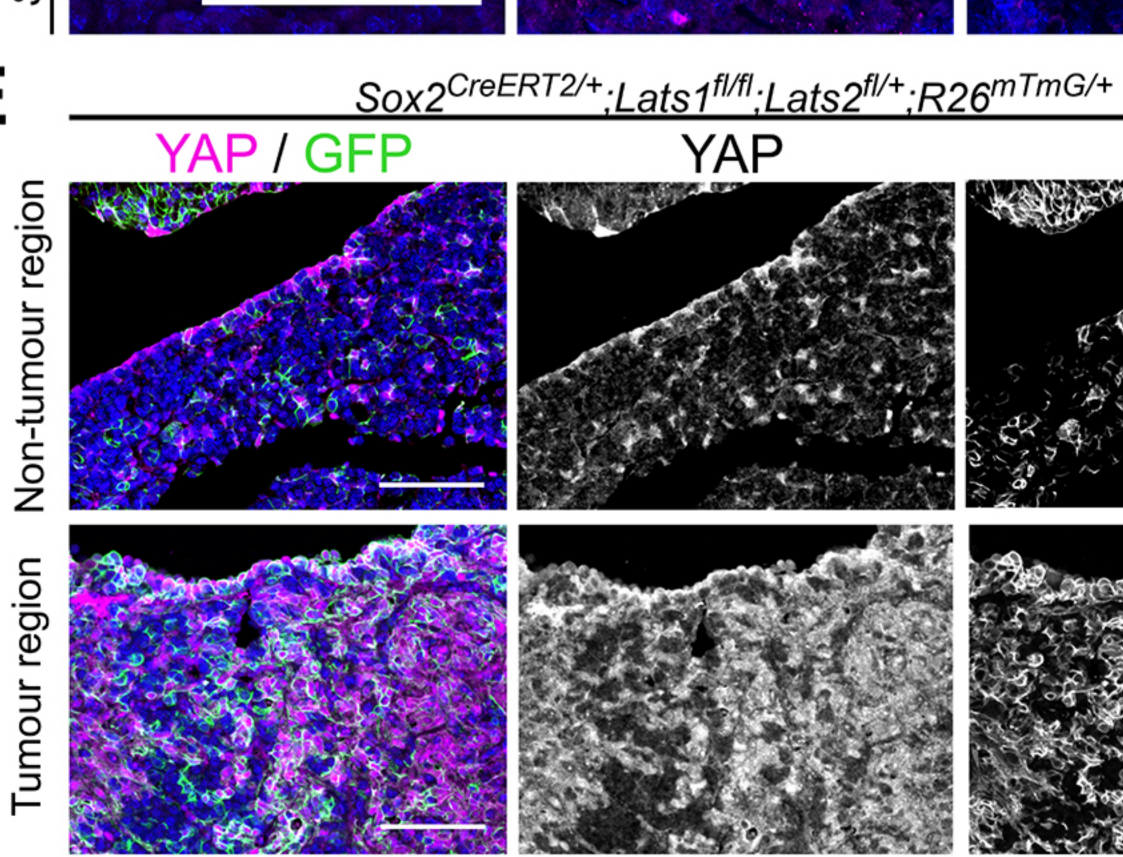


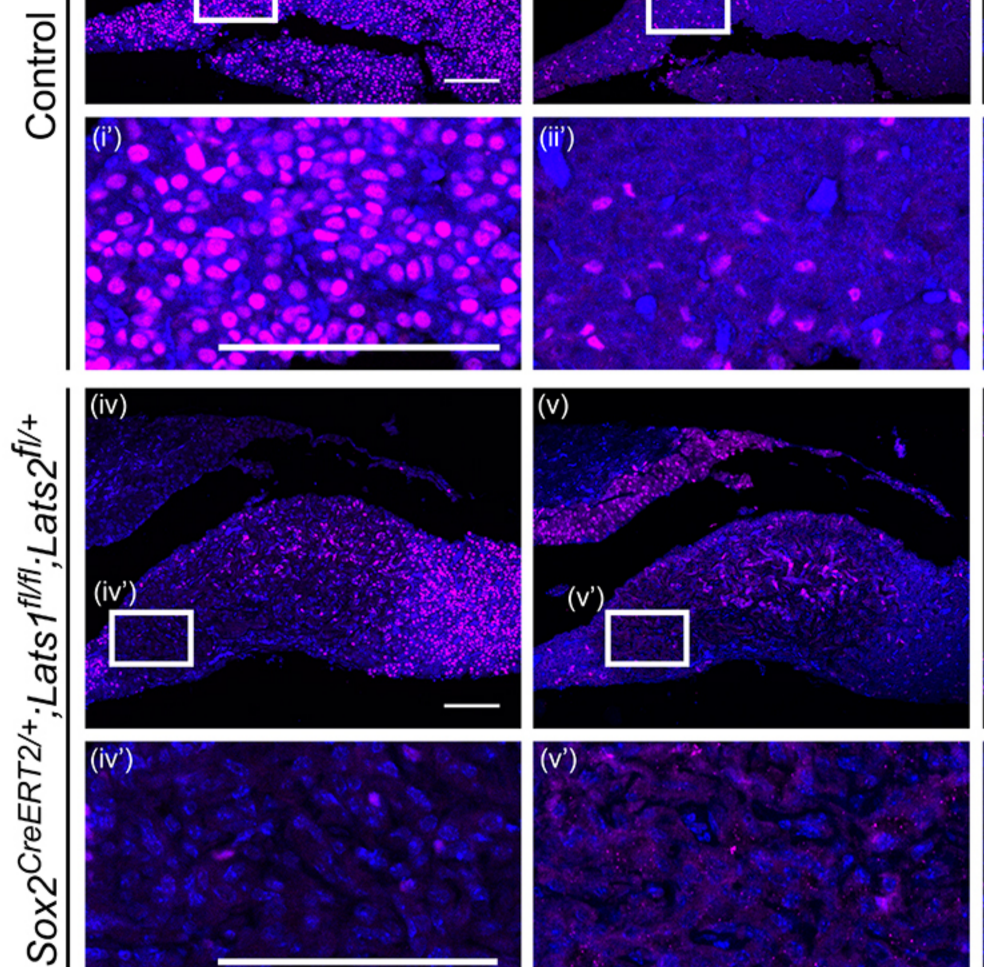
ats1<sup>fl/fl</sup>.











Lotte

(i')

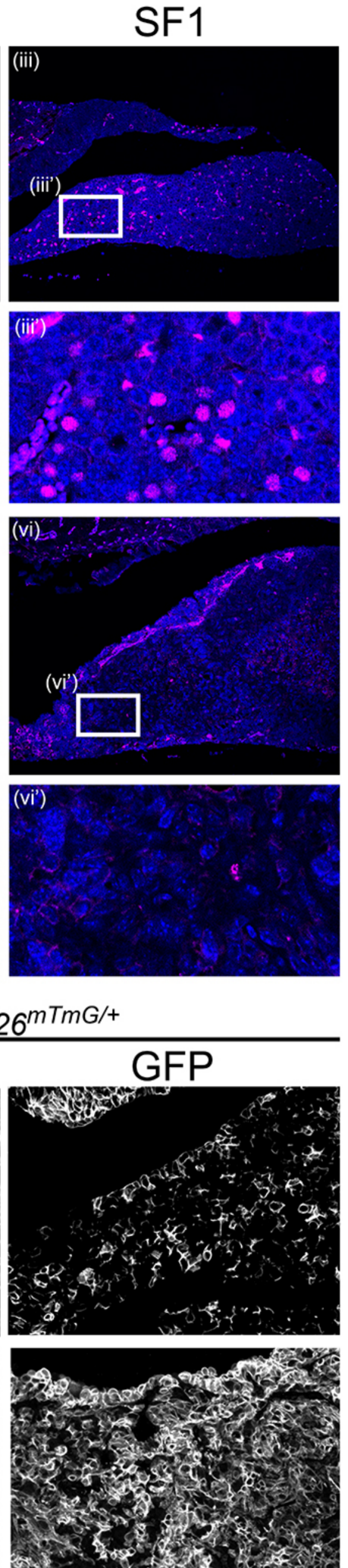
(i)

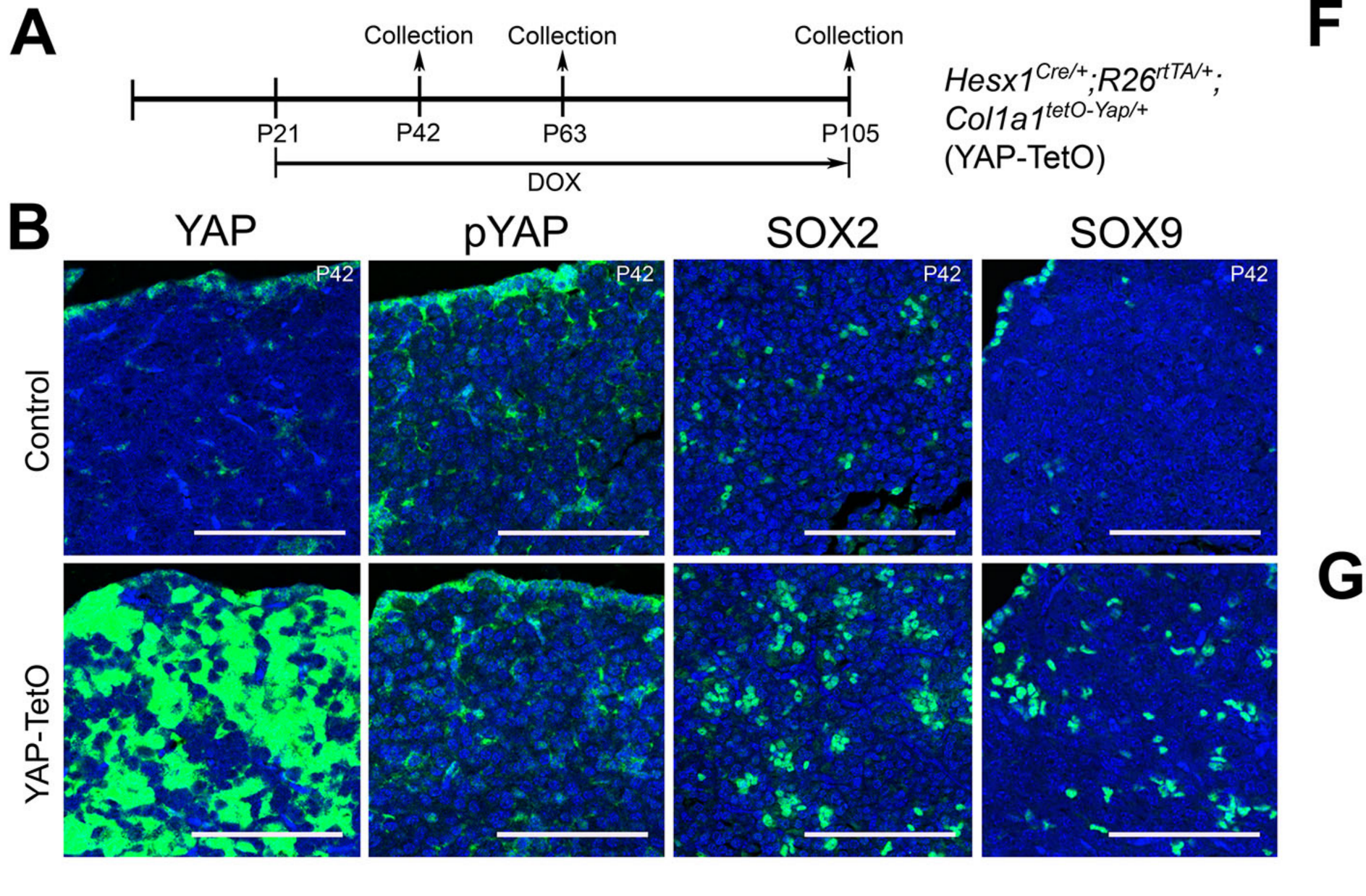
PIT1

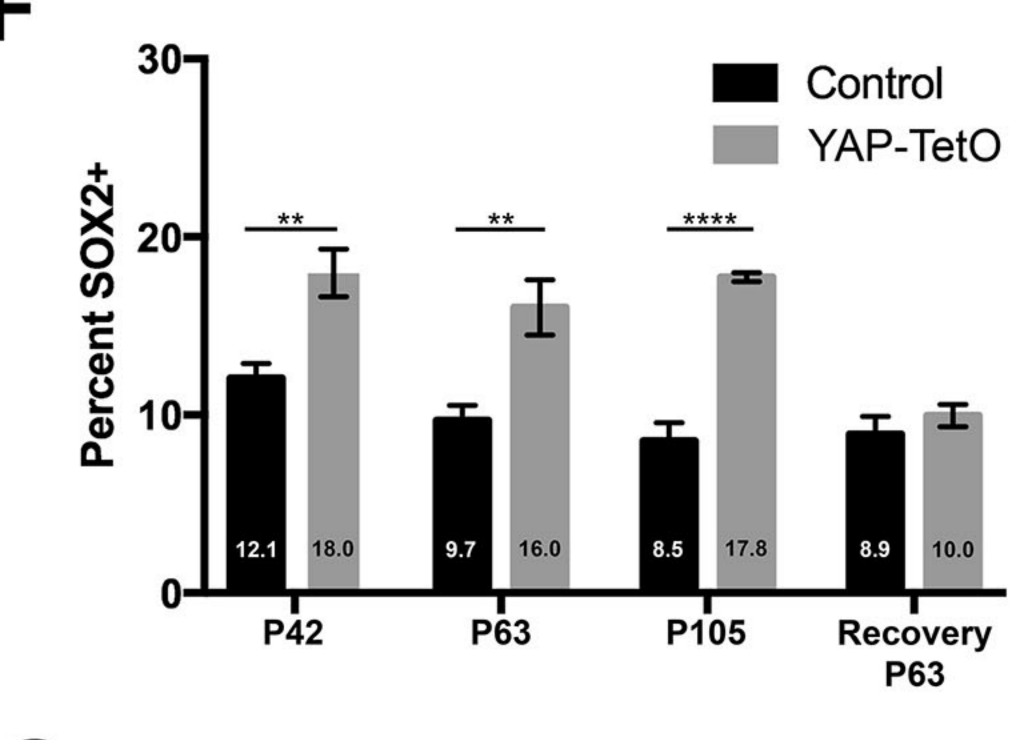
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(ii')

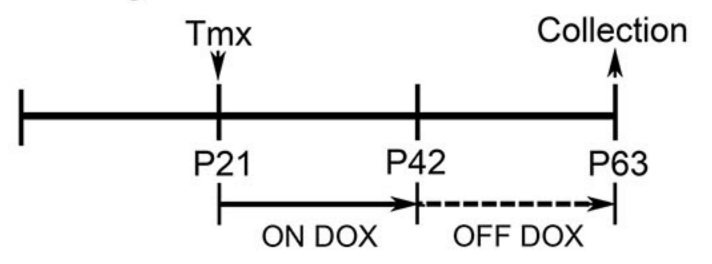
(ii)

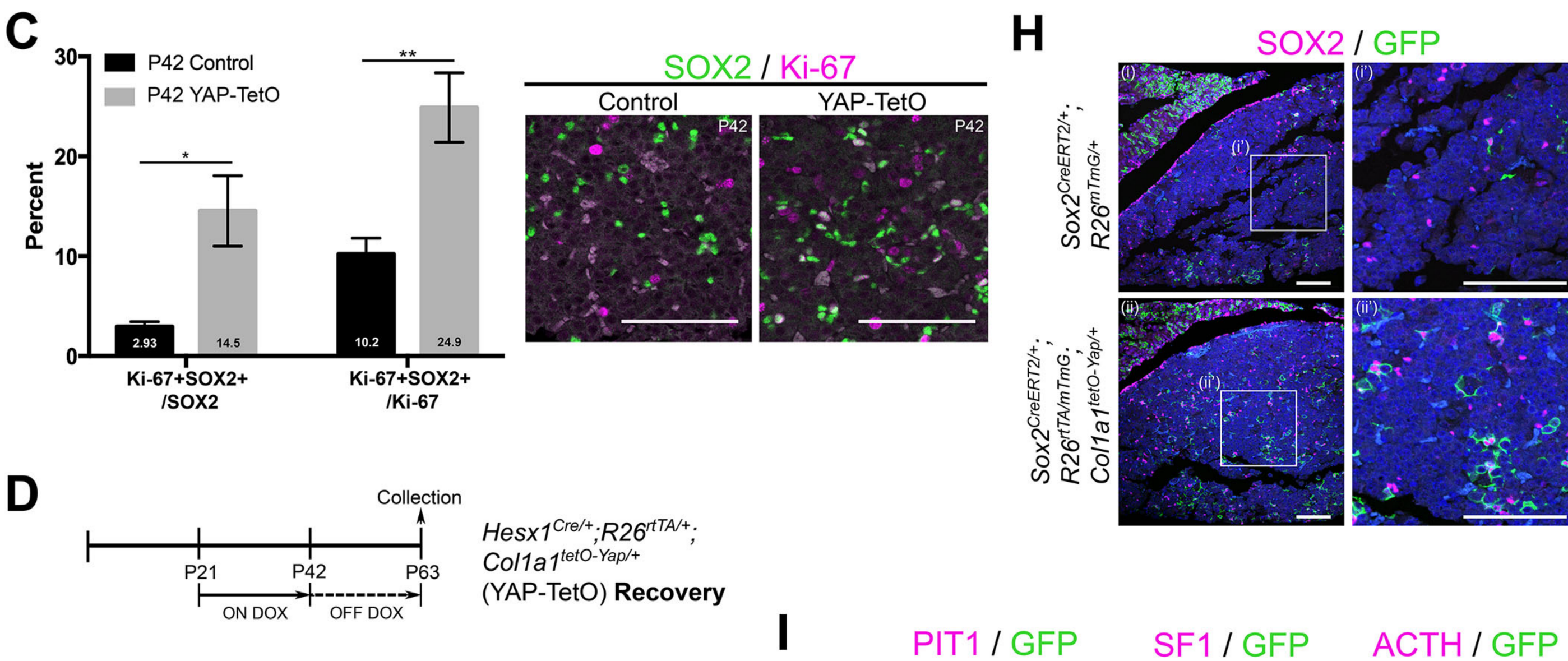






Sox2<sup>CreERT2/+</sup>;R26<sup>rtTA/mTmG</sup>;Col1a1<sup>tetO-Yap/+</sup> Recovery





1tetO-Yar

B

-

Col

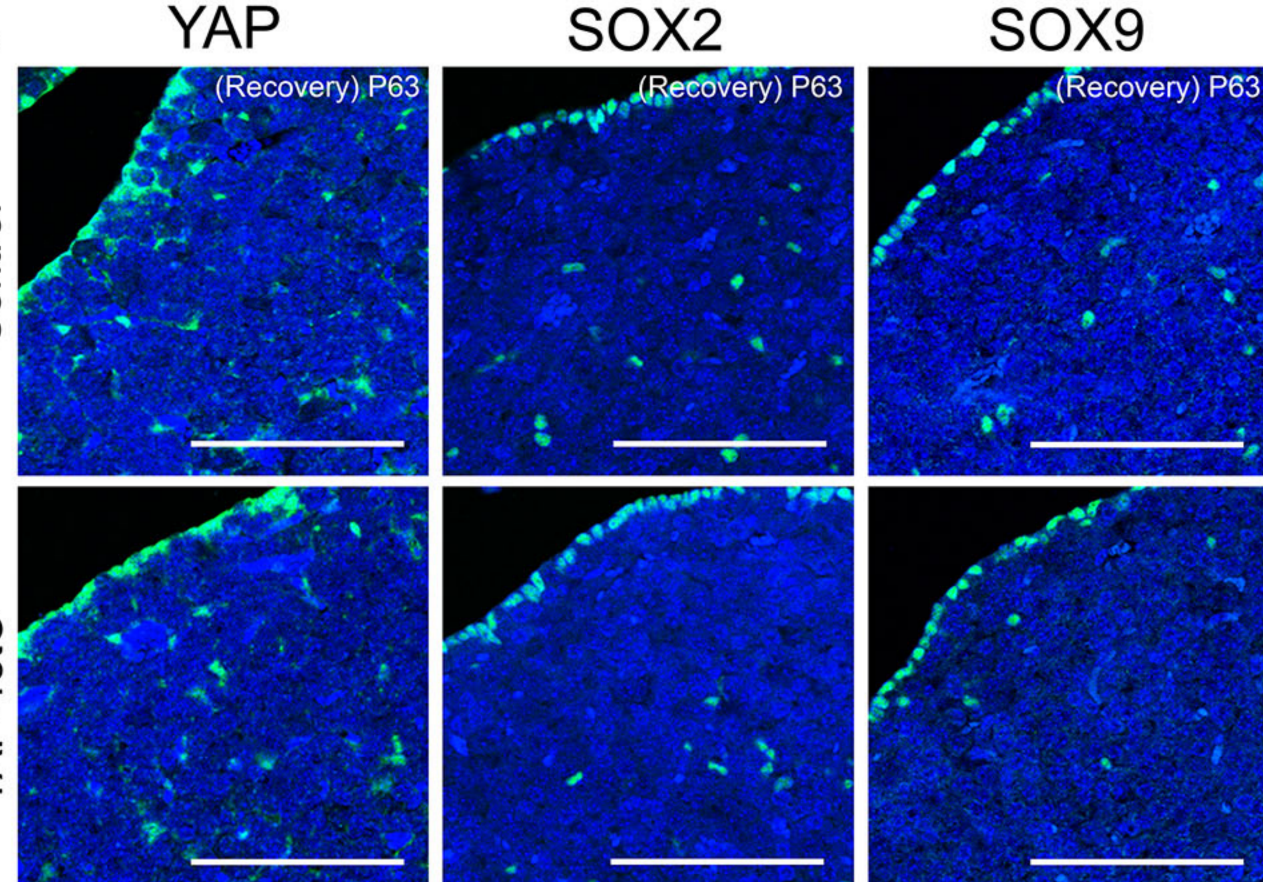
R26<sup>rtTA/mTmG</sup>

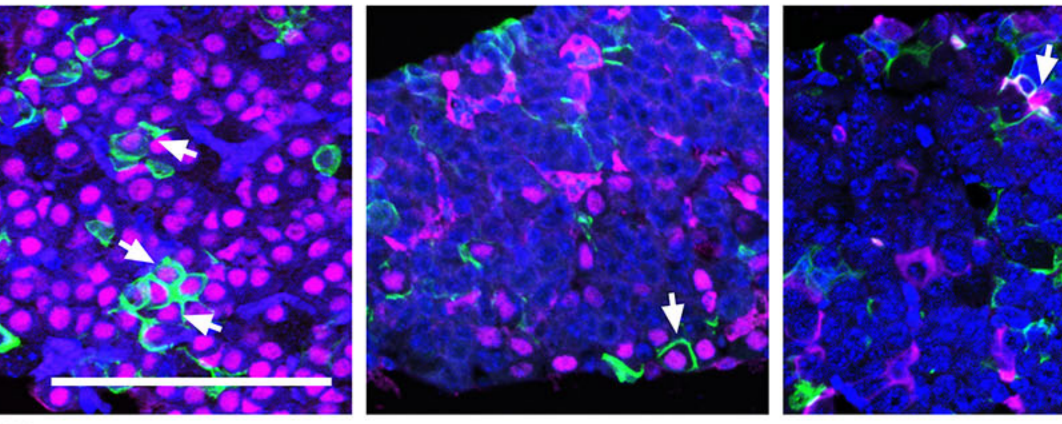
Sox2<sup>CreERT2/</sup>

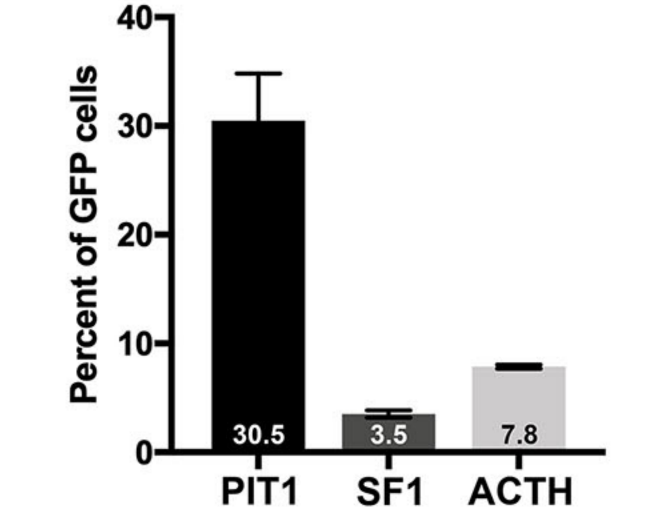
# YAP-TetO

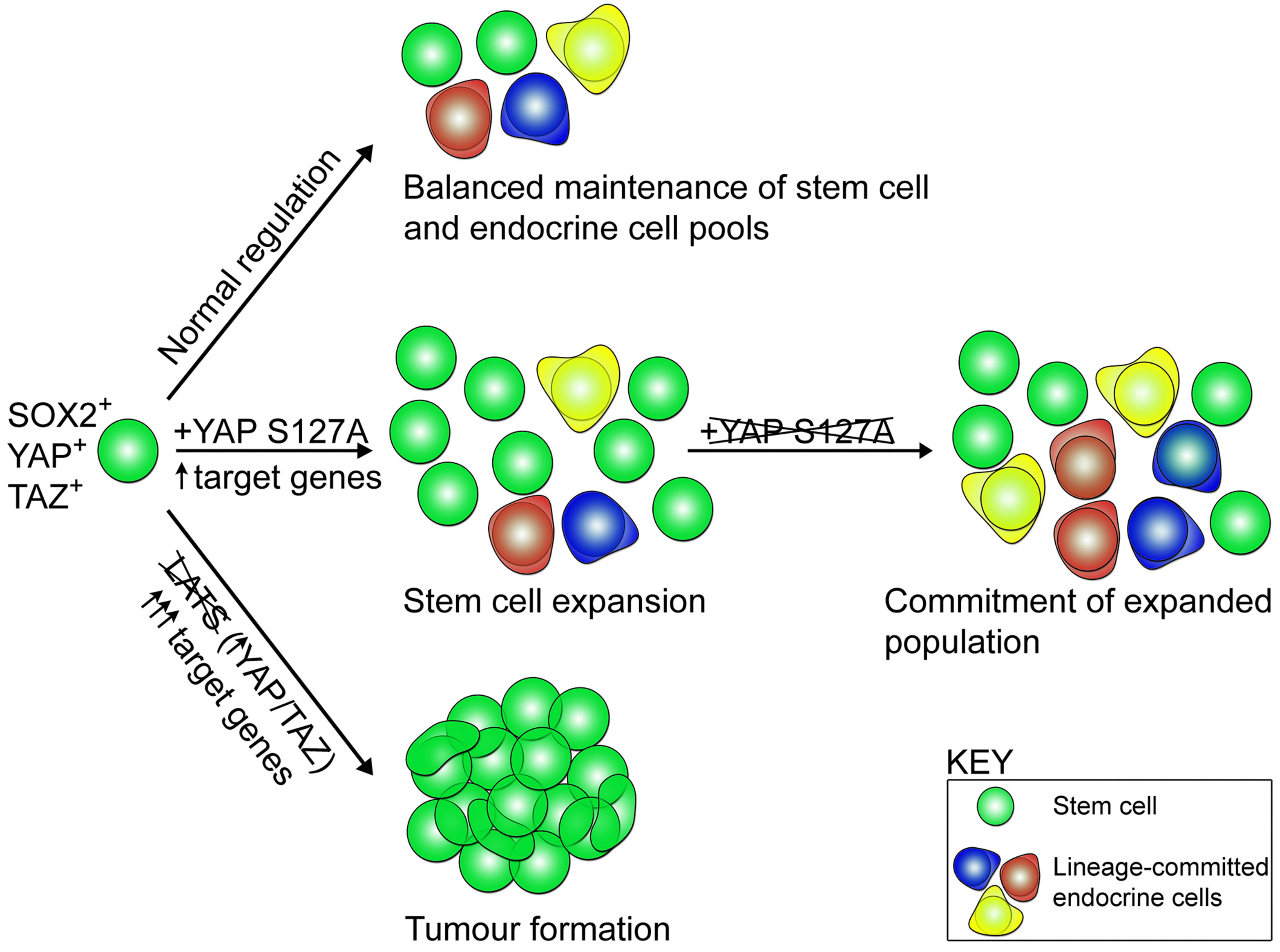


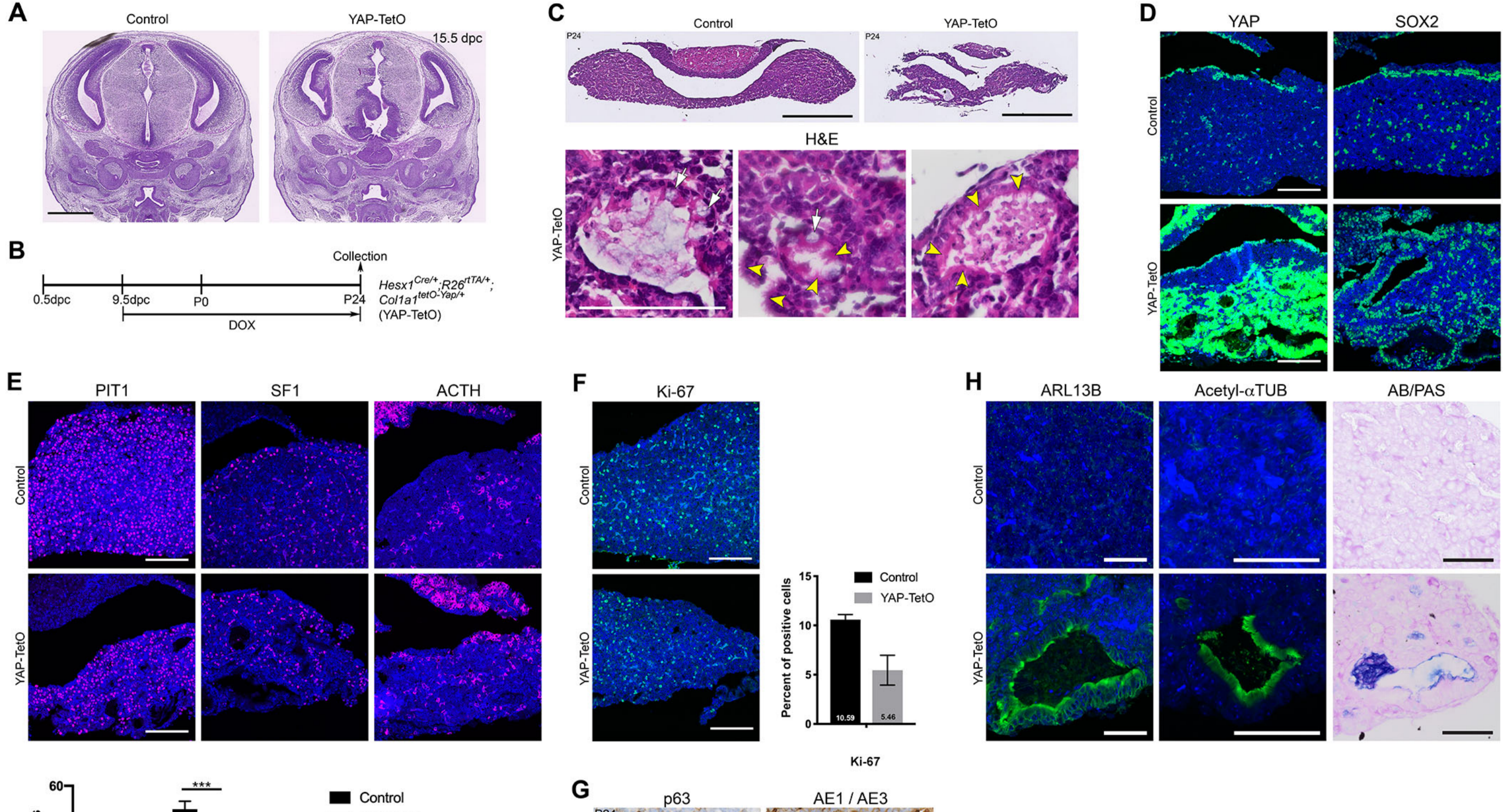
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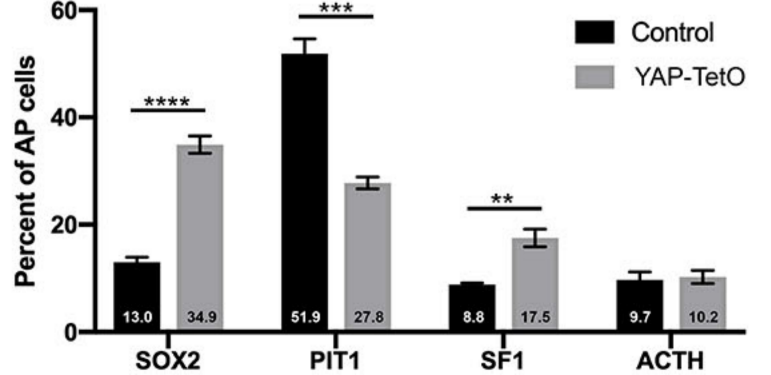




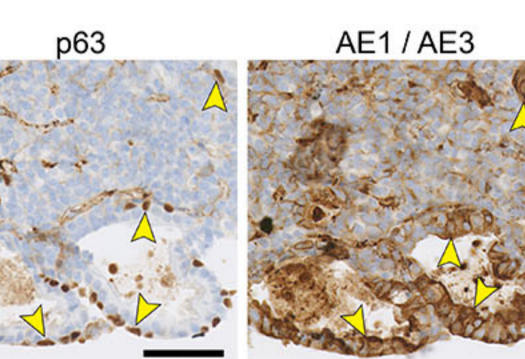


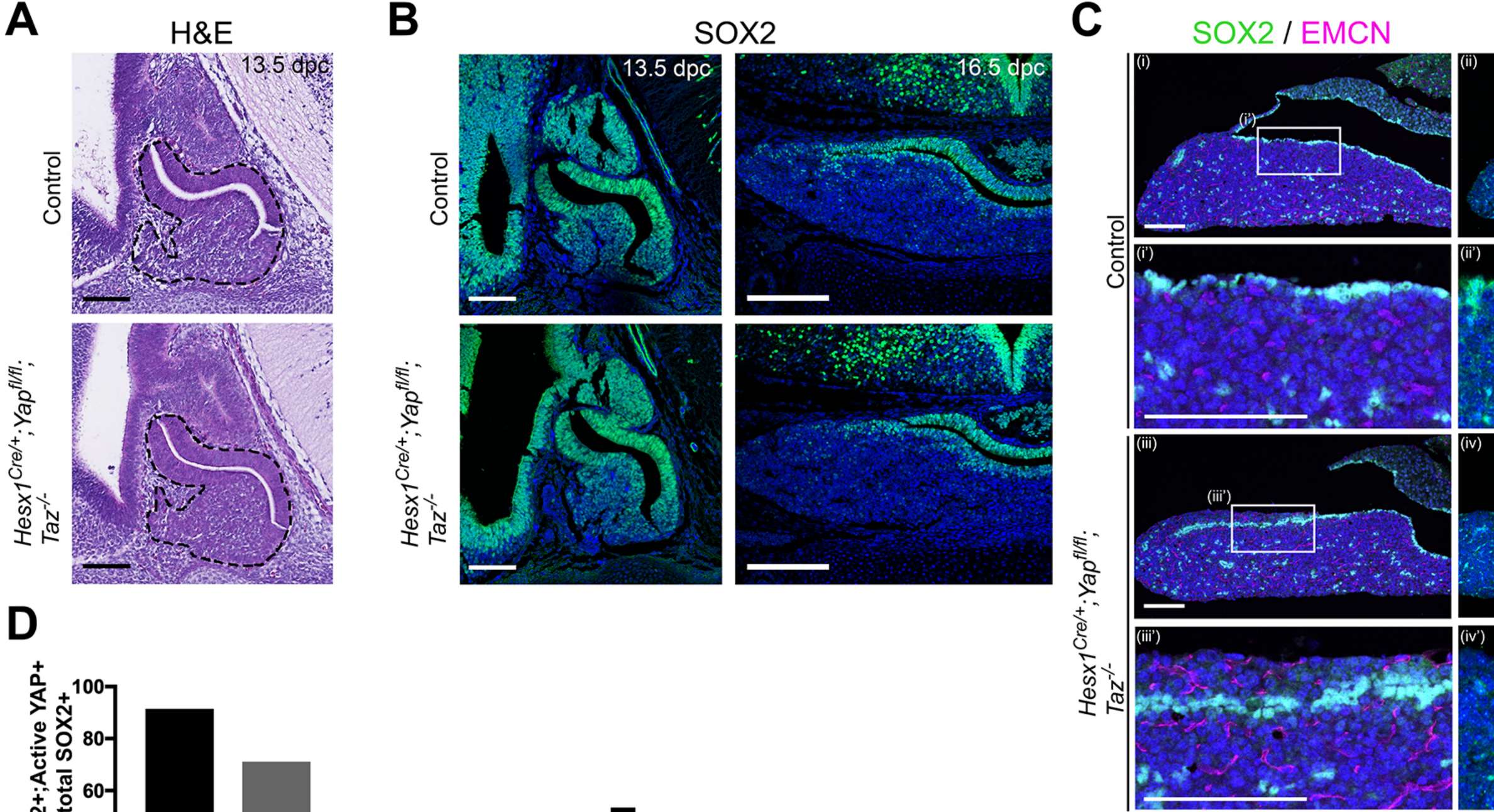


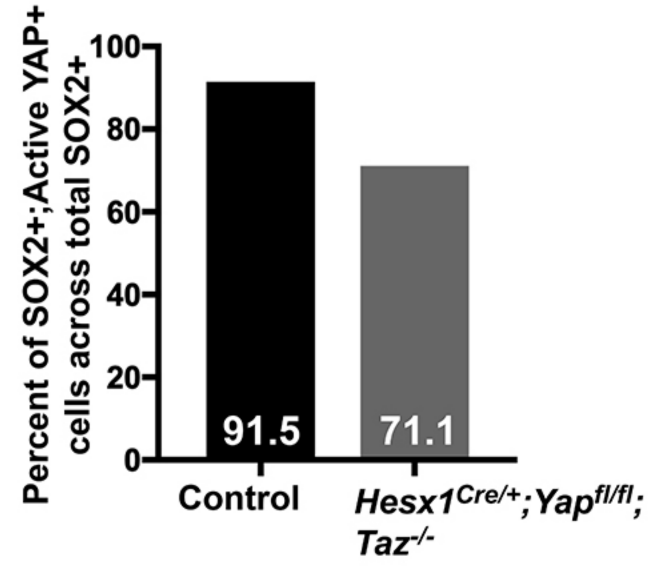


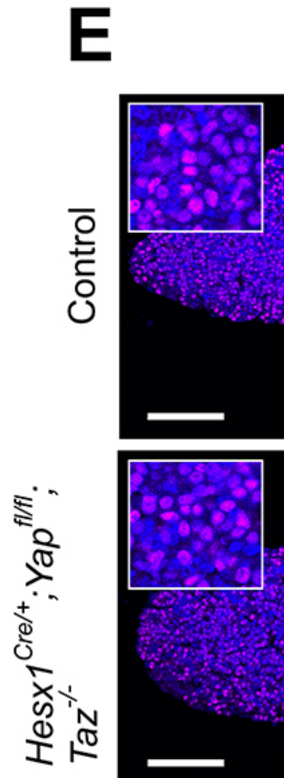


P24 (AP-TetO



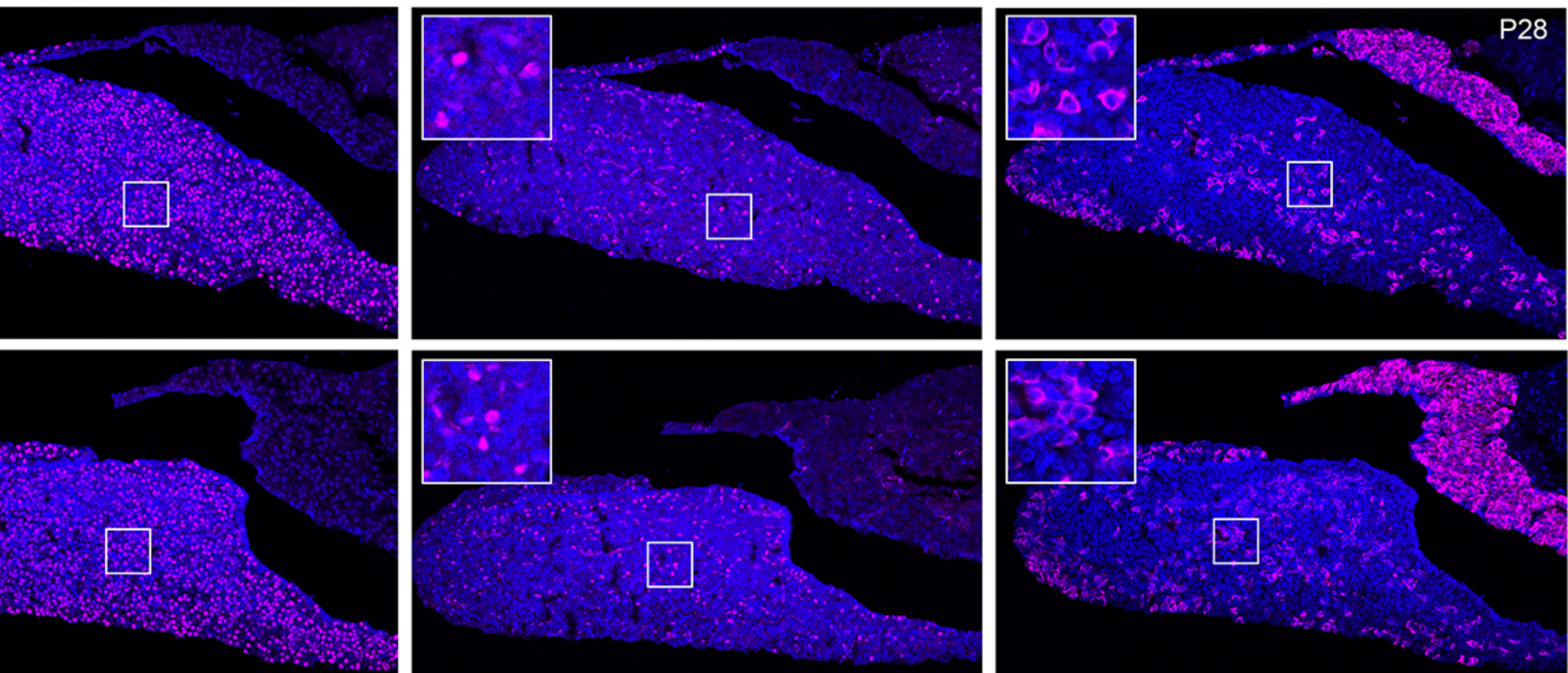




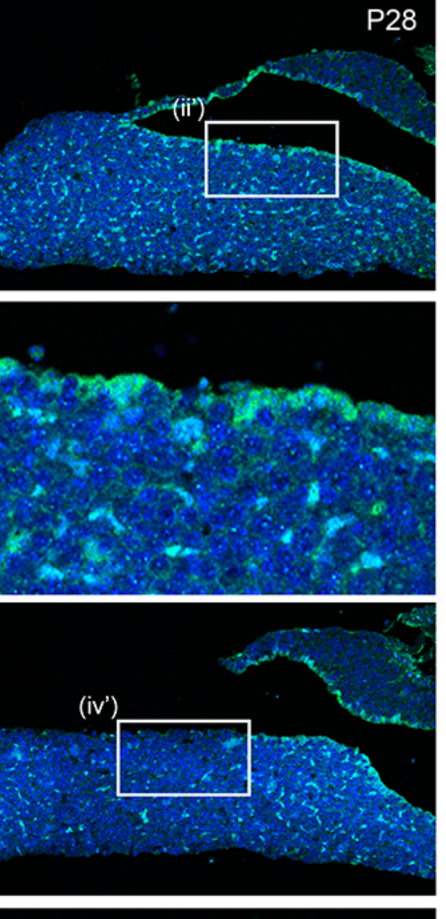


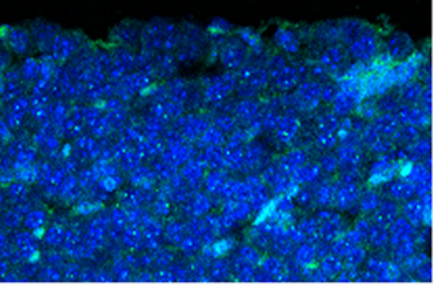
PIT1

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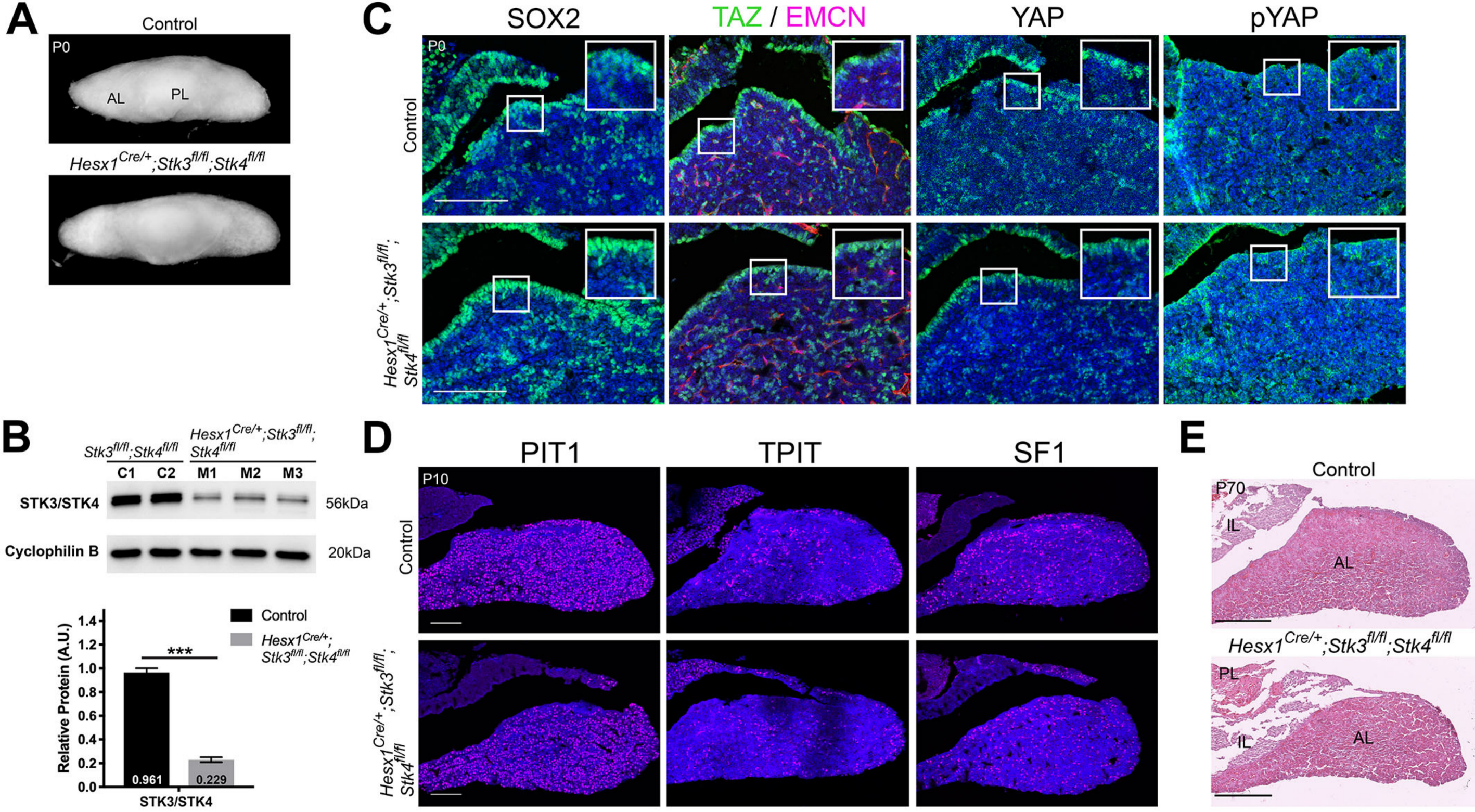


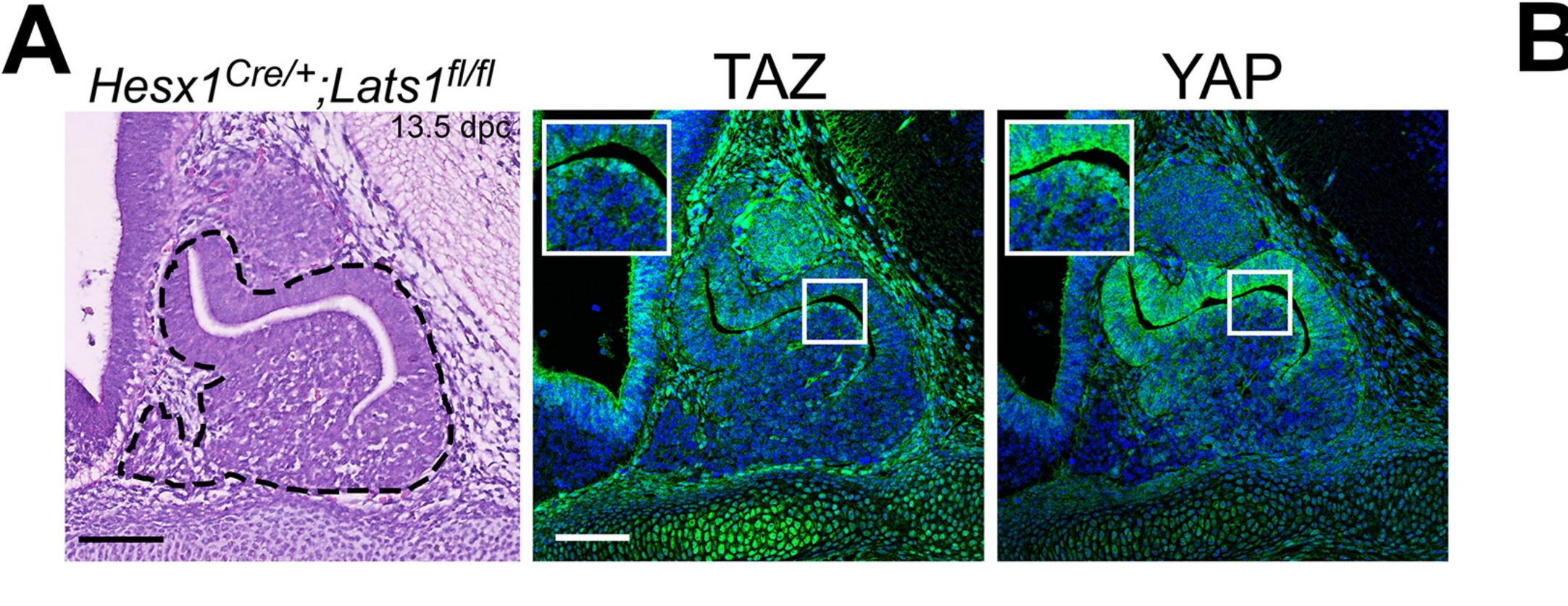


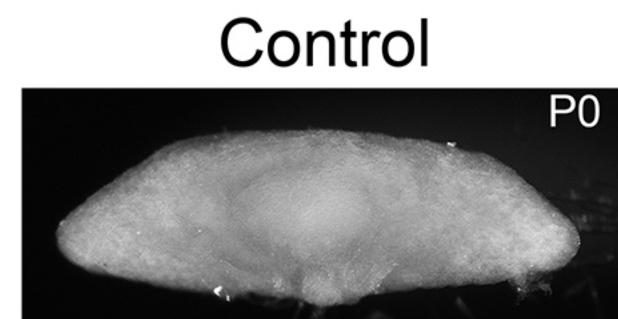


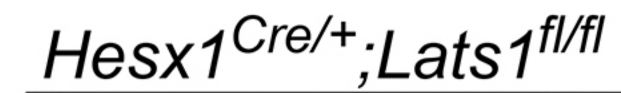


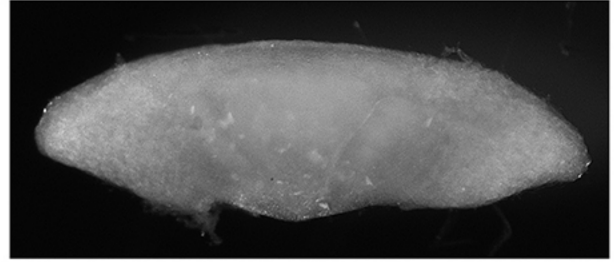


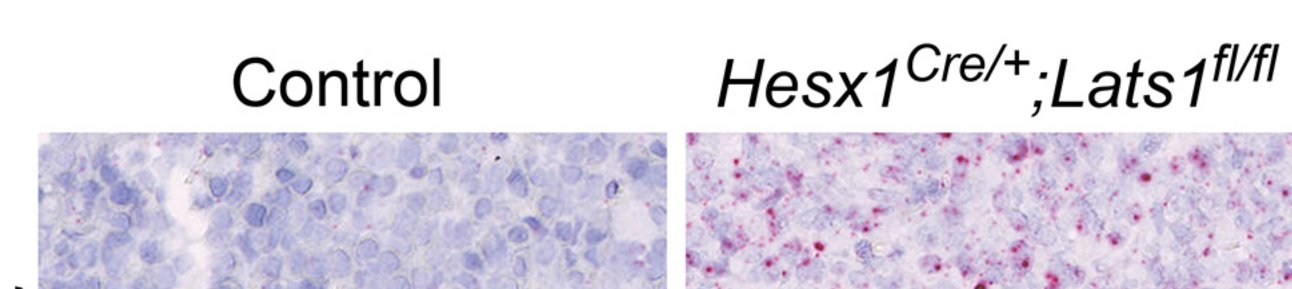


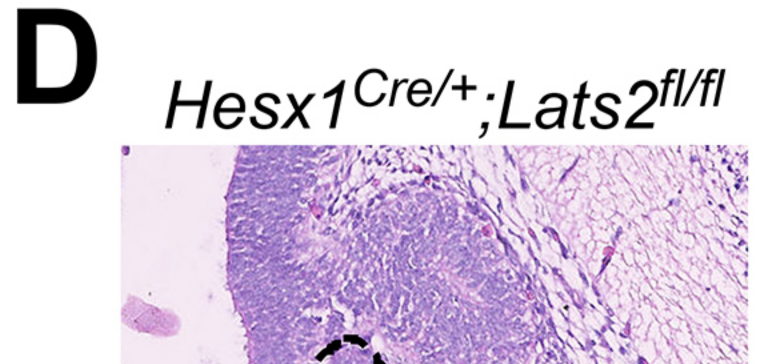


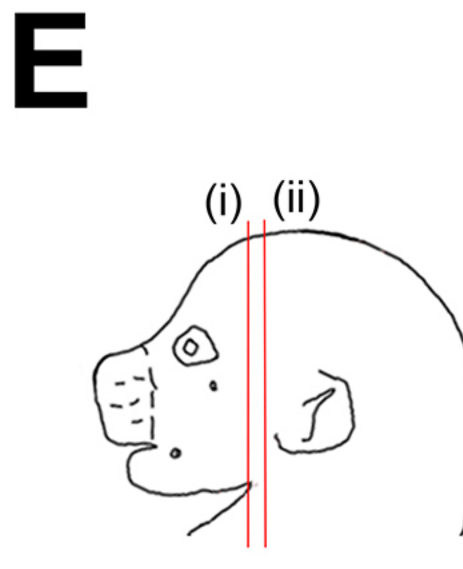




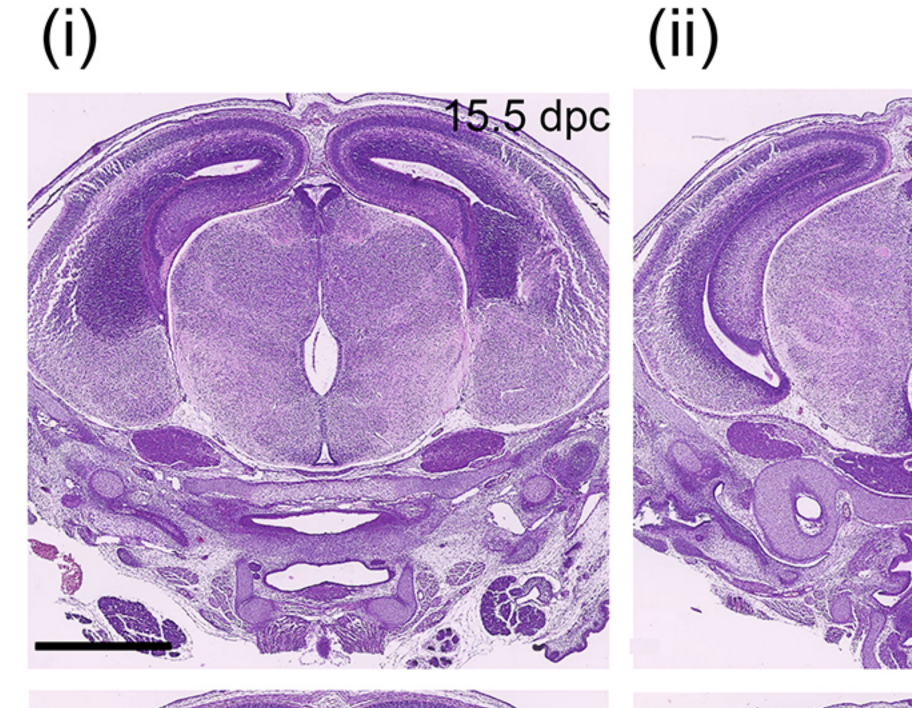


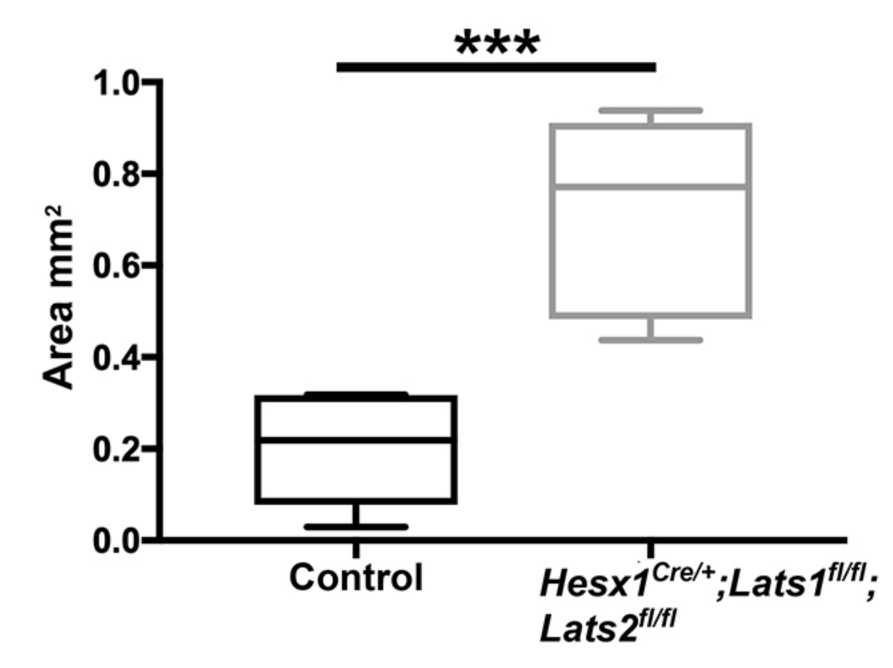


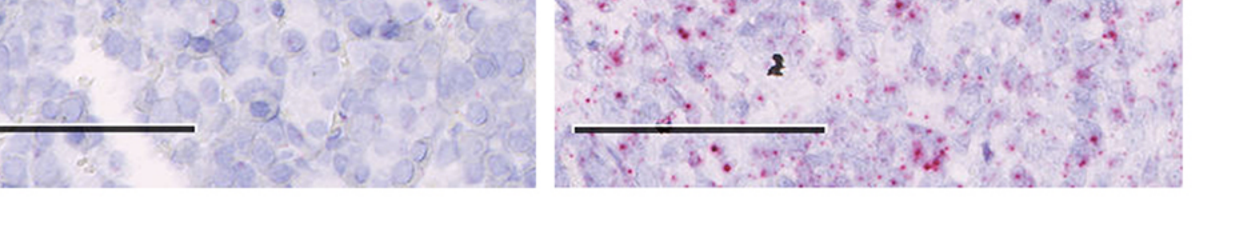


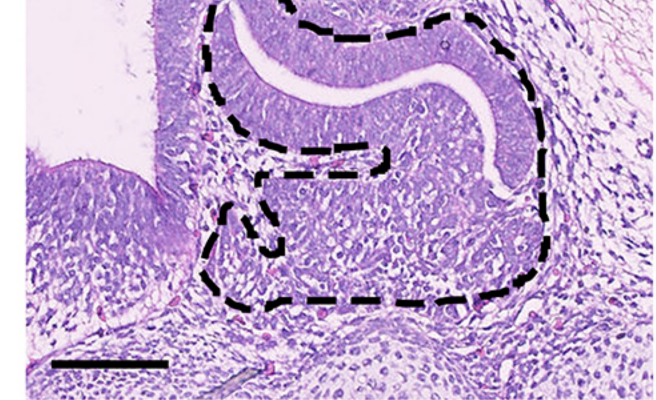


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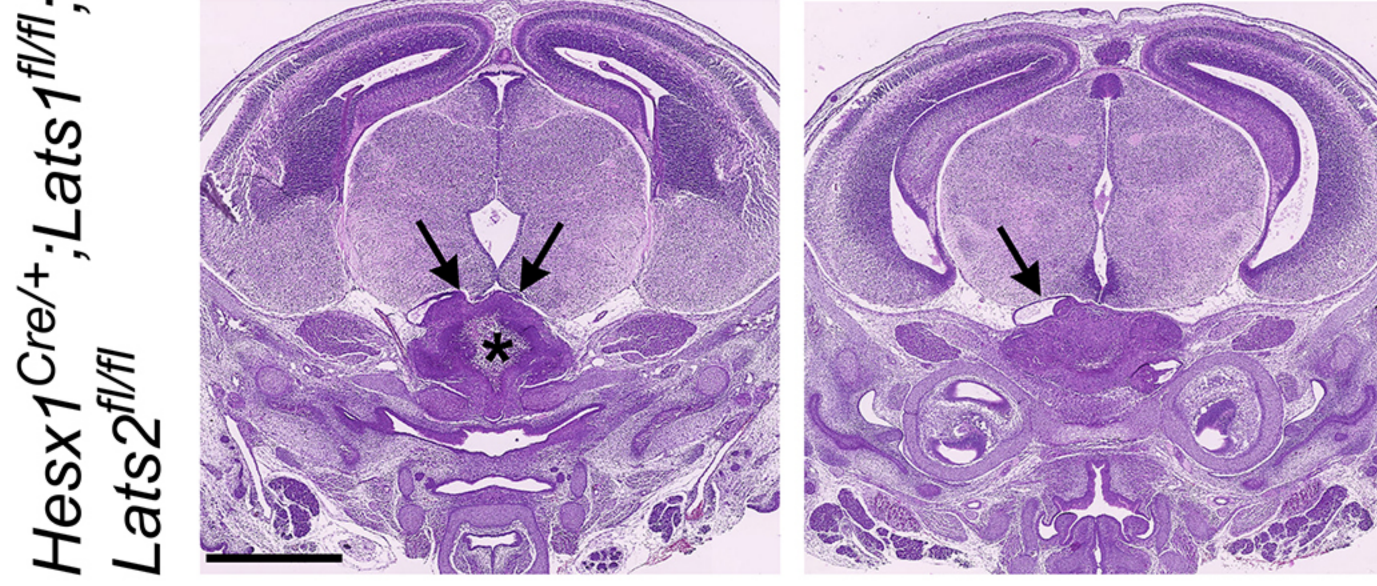


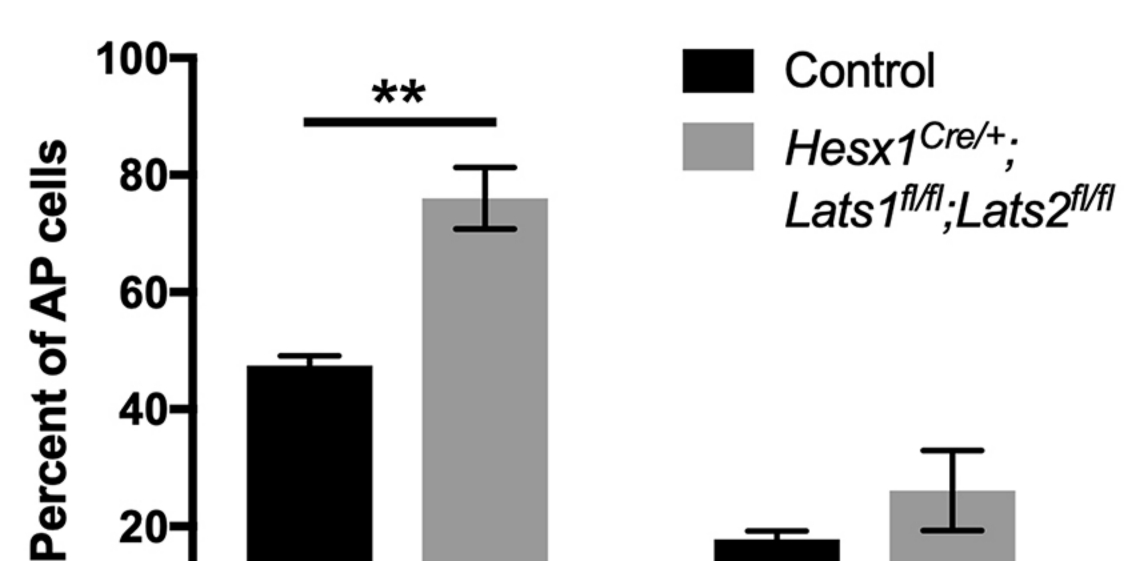


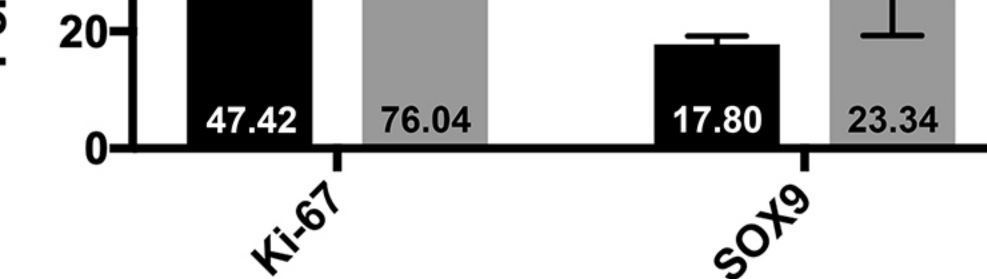


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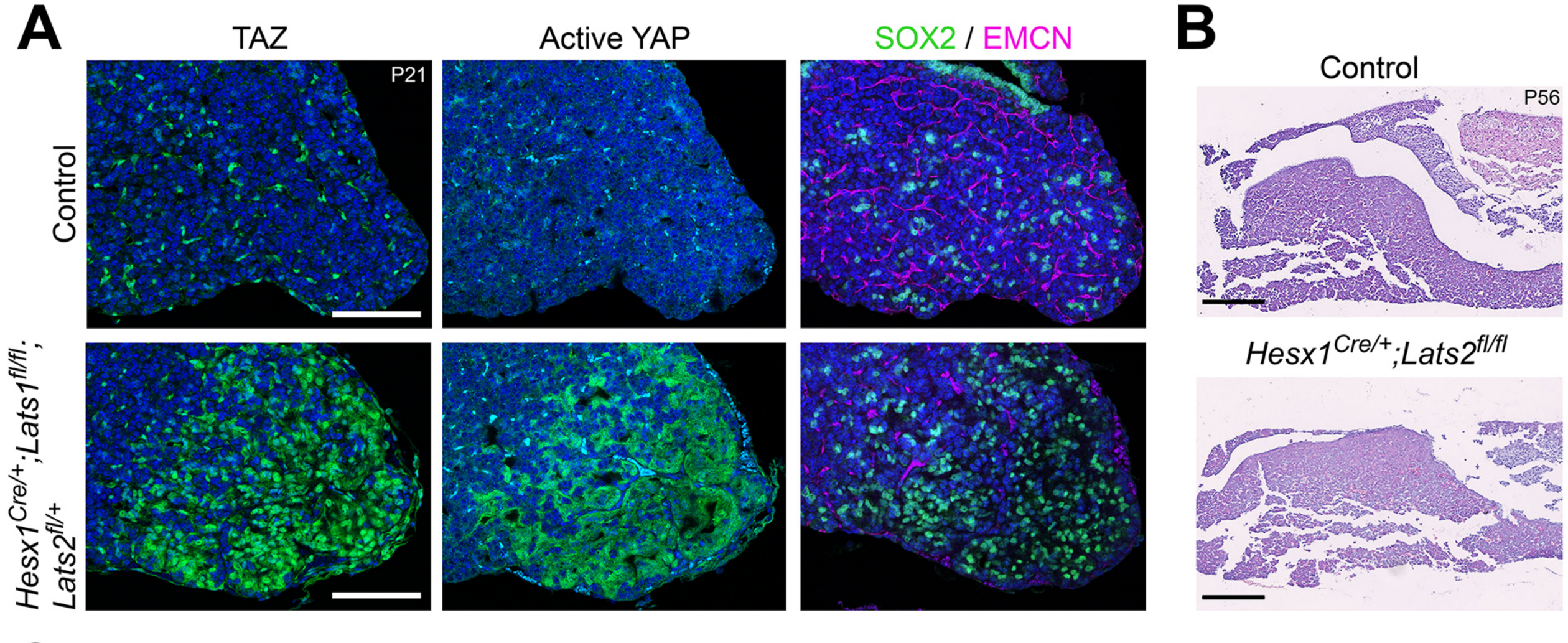
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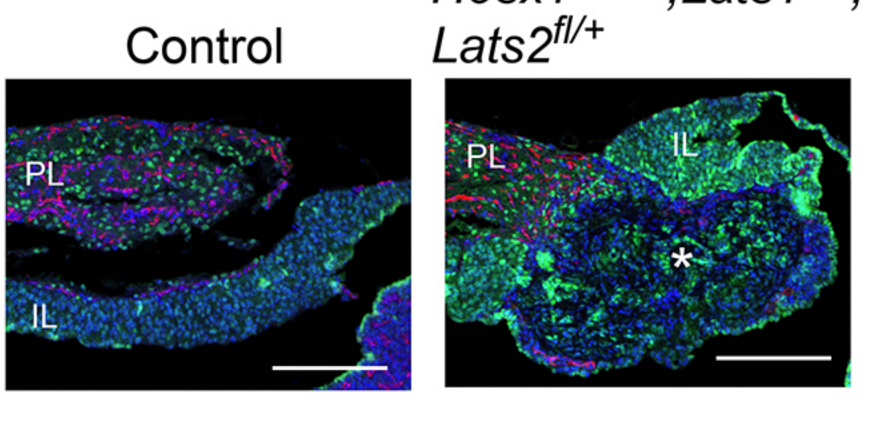




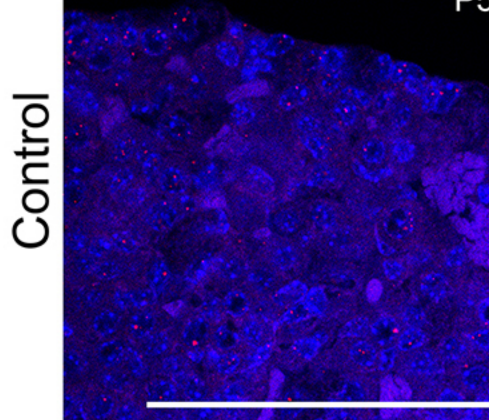
sote



- С SOX2 / EMCN Hesx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup>;



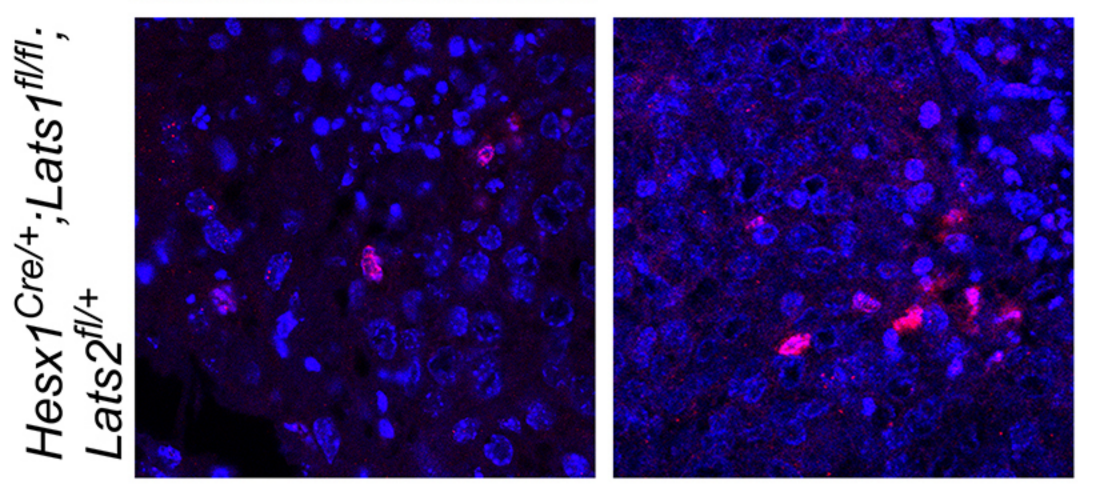
## gamma H2A.X P56



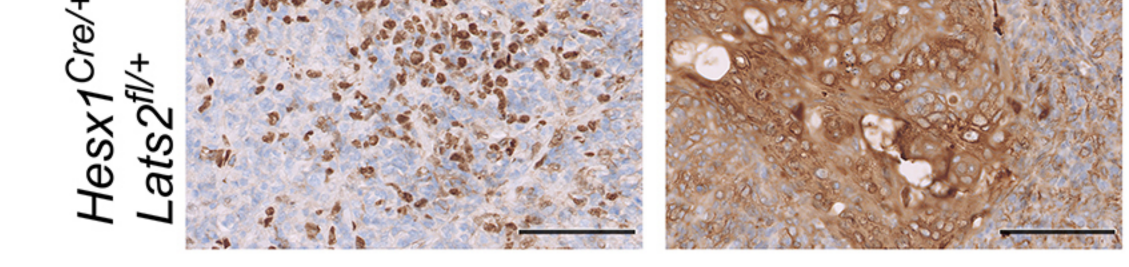
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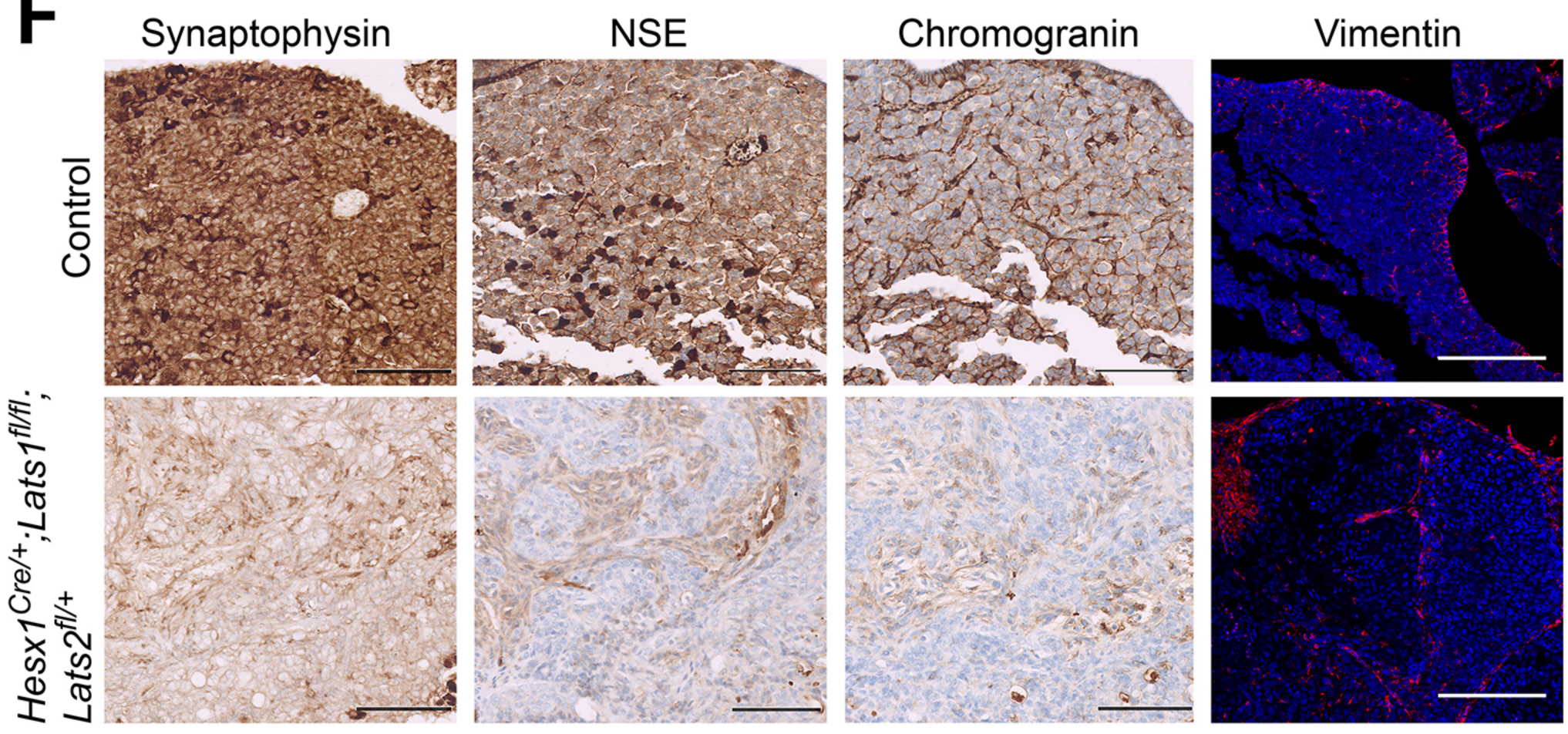
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esx1<sup>Cre/+</sup>;Lats1<sup>fl/fl</sup>;

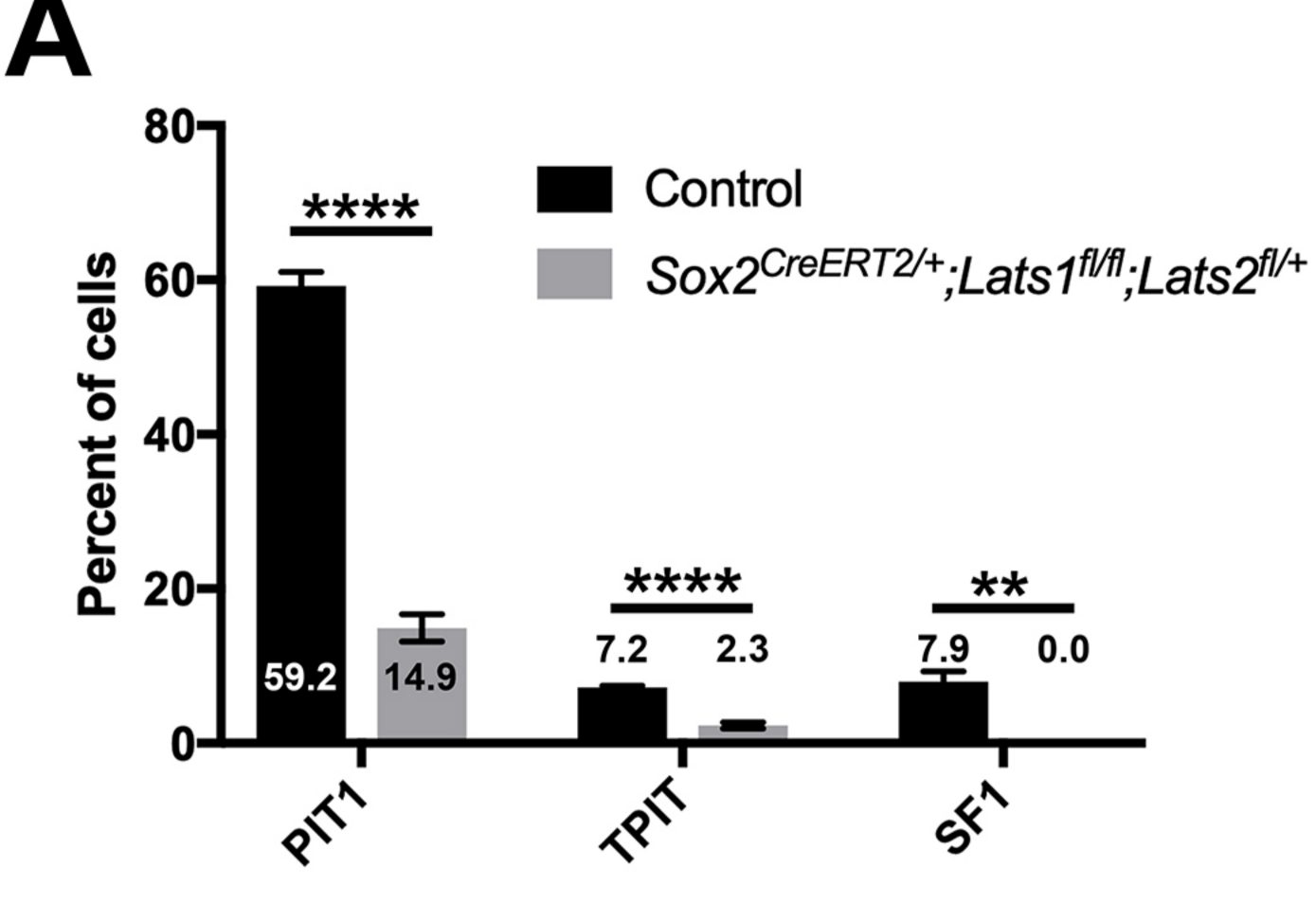


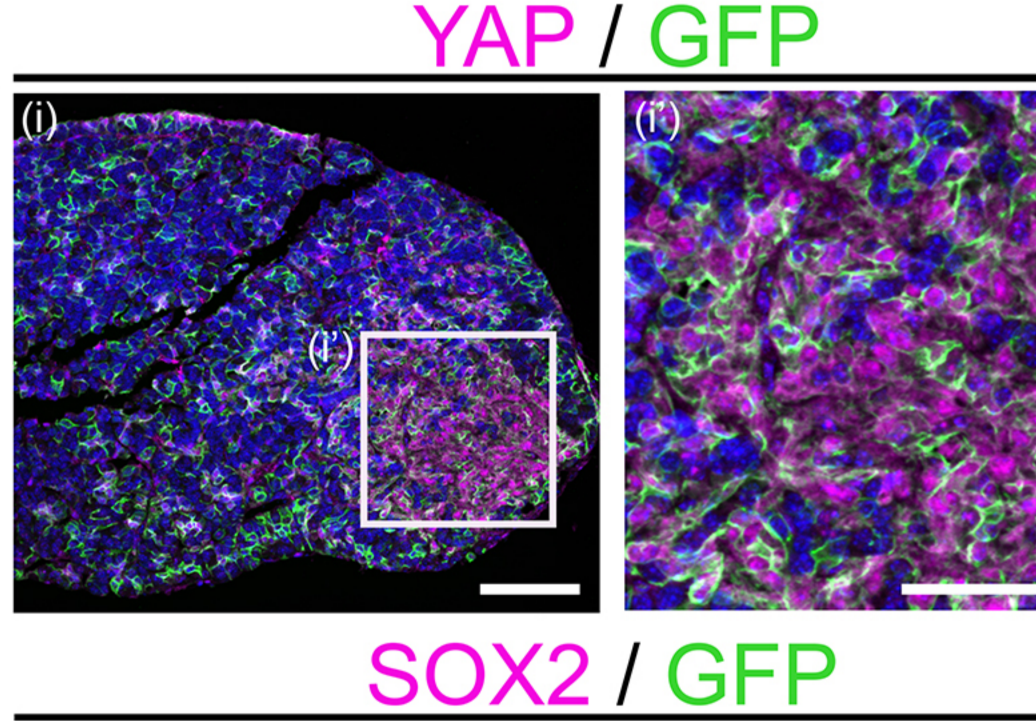
## Ε AE1/AE3 p63 9 Contr ;Lats1<sup>fl/fl</sup>

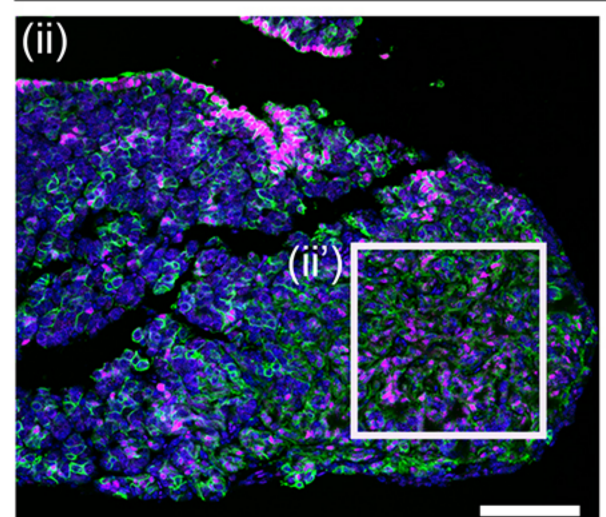


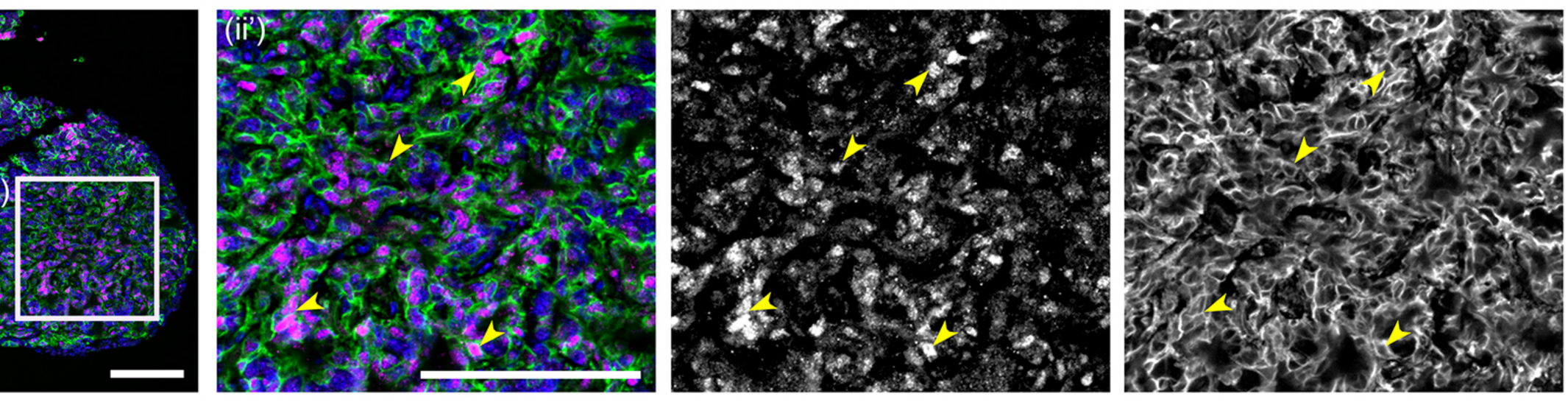








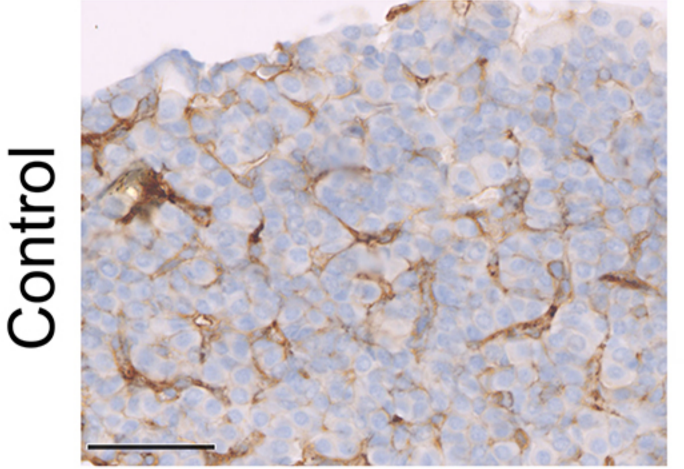




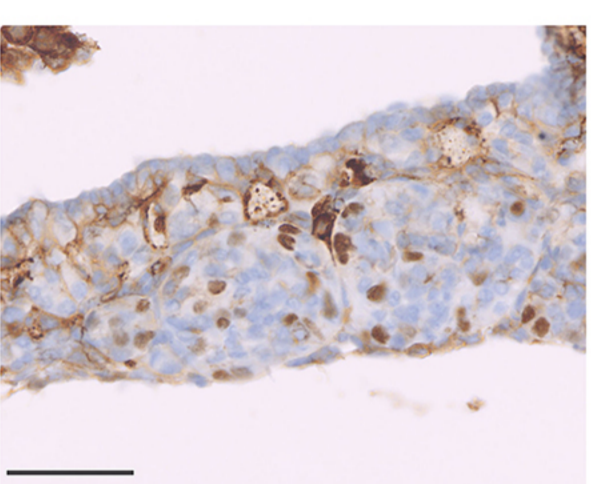


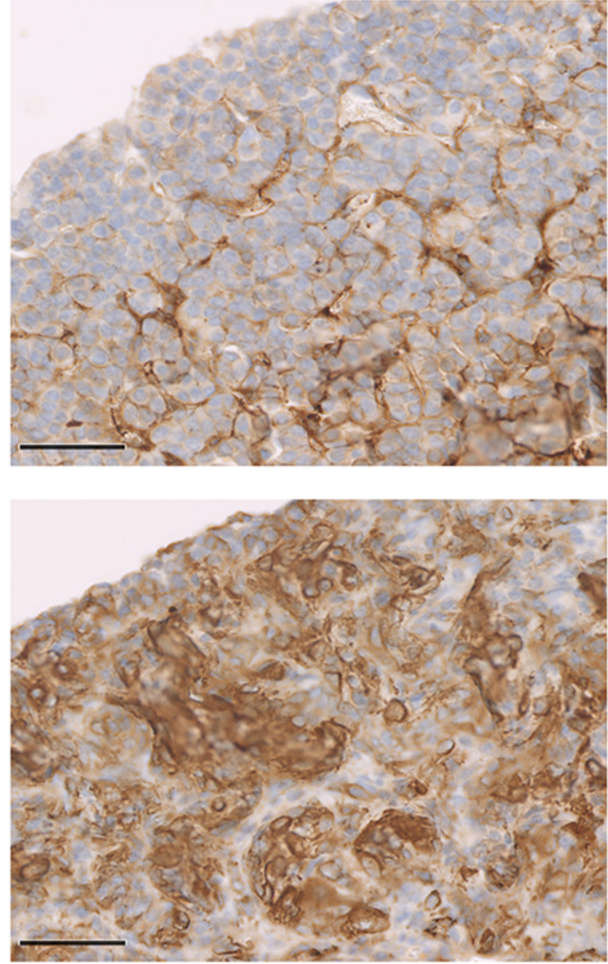
## p63



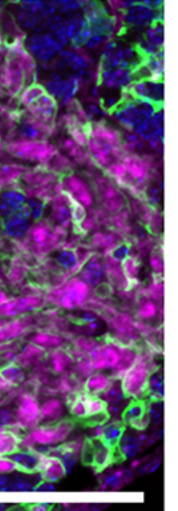


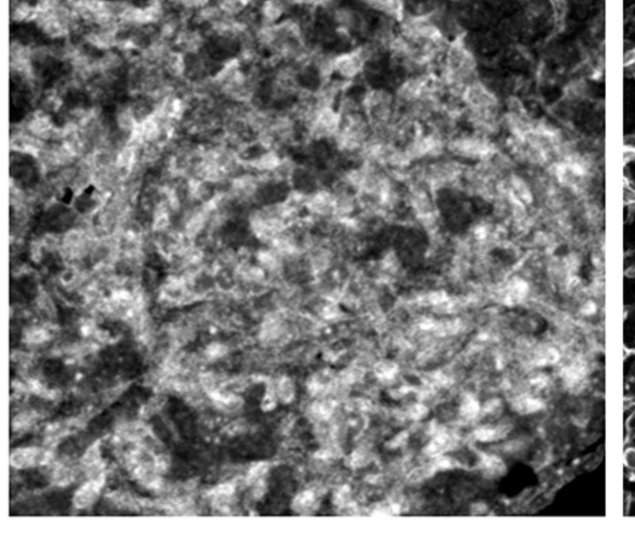






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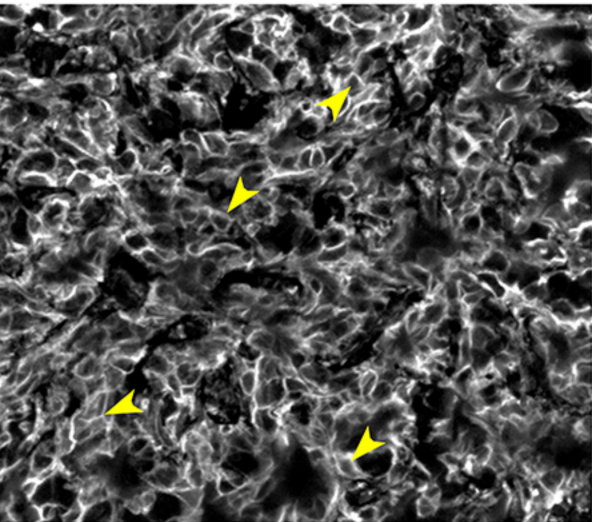




YAP

SOX2





## AE1/AE3

