Hypothalamic Rax\(^+\) tanycytes contribute to tissue repair and tumorigenesis upon oncogene activation in mice

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Hypothalamic tanycytes in median eminence (ME) are emerging as a crucial cell population that regulates endocrine output, energy balance and the diffusion of blood-born molecules. Tanycytes have recently been considered as potential somatic stem cells in the adult mammalian brain, but their regenerative and tumorigenic capacities are largely unknown. Here we found that Rax\(^+\) tanycytes in ME of mice are largely quiescent but quickly enter the cell cycle upon neural injury for self-renewal and regeneration. Mechanistically, Igf1r signaling in tanycytes is required for tissue repair under injury conditions. Furthermore, Braf oncogenic activation is sufficient to transform Rax\(^+\) tanycytes into actively dividing tumor cells that eventually develop into a papillary craniopharyngioma-like tumor. Together, these findings uncover the regenerative and tumorigenic potential of tanycytes. Our study offers insights into the properties of tanycytes, which may help to manipulate tanycyte biology for regulating hypothalamic function and investigate the pathogenesis of clinically relevant tumors.

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Hypothalamic tanycytes share many common features with ependymal cells but display a unique morphology and distinct functional features. They are polarized cells, with cell bodies lining the third ventricle, elongated processes extending into the parenchyma, and endfeet contacting the pial surface of the brain. In the median eminence (ME), a structural link between the hypothalamus and pituitary gland, tanycytes contribute to the regulation of multiple hypothalamic functions including the diffusion of blood-borne molecules, neuroendocrine output, energy balance, and reproductive ageing. Tanycytes in ME have been implicated in the maintenance of hypothalamus-mediated body homeostasis, but have also been recently identified as a possible key component of stem cell niche due to their prominent expression of neural progenitor markers and their potency of proliferation and differentiation. While somatic stem cells, in general, are characterized by their self-renewal, regenerative, and tumorigenic potential, it remains largely unknown how ME tanycytes maintain themselves and to what extent the disturbance of such homeostasis contributes to diseases such as cancer.

Single-cell RNAseq technology has recently been applied to dissect the self-renewal and regenerative capacity of ependymal cells. In particular, the combination of cell lineage tracing with single-cell profiling shows that ependymal cells do not function as latent neural stem cells (NSCs) to self-renew and repair the damaged neural tissue. As specialized ependymal cells, the role of tanycytes in tissue repair is still unknown. Interestingly, the ultrastructural analysis identifies tanycytes as uniciliated ependymal cells, implicating a potentially different function from typical ependymal cells.

Another important characteristic of somatic stem cells is their tumor-initiating potential, but the tumorigenic capacity of tanycytes has never been reported. Craniohypophyseal tumors are a benign but aggressive intracranial tumor that occurs in between the hypothalamus and pituitary gland. Clinically, craniohypophyseal tumors are subdivided into adamantinomatous and papillary subtypes, which differ in genetic mutations and onset of age.

Papillary craniohypophyseal tumors are restricted to adult-onset with a much higher mortality rate and frequently driven by somatic Brf1v600E mutation. Although the clinical endoscopic studies imply that papillary craniohypophyseal tumors may originate from the hypothalamus–pituitary axis including ME, the cell of origin for adult-onset craniohypophyseal tumors remains unknown. We wondered whether ME tanycytes could develop into craniohypophyseal-like tumors upon oncogene activation in the adult brain.

Retina and anterior neural fold homeobox transcription factor (Rax) is selectively expressed in hypothalamic tanycytes, especially those at the ventral part of the third ventricle. Here, we aim to deconstruct the regenerative and tumorigenic potential of Rax+ tanycytes and speculate that whether tanycytes in ME serve as cells-of-origin for craniohypophyseal. We thereby manipulated tanycyte biology and performed lineage tracing using Rax-CreERT2 knock-in mice, and revealed that Rax+ tanycytes responded to neural injury for regeneration and contributed to tumorigenesis upon Braf oncogene activation.

**Results**

Transcriptomic analysis of tanycytes in ME and their niche cells. Adult ME has been reported to represent a robust proliferative structure, encompassing a 15-fold higher density of dividing cells compared to other hypothalamic regions. Recently, a molecular census of arcuate nucleus and ME cells focused on the heterogeneity of neurons and their roles in regulating energy homeostasis. However, the mitotic activity of tanycytes in the niche remains contentious. To profile the transcriptome of tanycytes in ME (containing β2 subtype) and its neighboring hypothalamic area of interest (Fig. 1a), we microdissected this particular region from adult mice and acutely dissociated the cells for single-cell RNAseq (Supplementary Fig. 1; “Methods”). Our unsupervised analysis of 990 qualified cells clearly discriminated ten distinct clusters (Fig. 1b). Using expression patterns of cell-type-specific marker genes, we assigned a single identity to each cluster: tanycytes (Rax+), vascular and leptotemporalneural cells (VMLCs, Lam+), erythrocytes (Hbα+, Hbβ+), neurons (Syp+), oligodendrocyte precursor cells (OPCs, Cspg4+), pars tuberalis cells (PTCs, Slc4a7A1+, Tshb+), natural killer cells (NK cells, Krg+), macrophages/microglia (Aif1+), and oligodendrocytes (Mog+) (Supplementary Fig. 2a and Supplementary Data 1). The putative cell stem markers Sox2, Slc1a3, Vimentin (Vim), and Alodc were enriched but not exclusively expressed in tanycytes (Supplementary Fig. 2d). Moreover, tanycytes shared a set of stemness-associated genes with NSCs and ependymal cells (Supplementary Fig. 2f and Supplementary Data 2).

**Adult Rax+ tanycytes are largely quiescent.** Actively dividing cells in ME have been reported to control systemic energy balance, but whether ME tanycytes are mitotically active remain controversial. To determine their mitotic activity, we first confirmed Rax as a molecular marker for ME tanycytes and revealed Scn7a, Col25a1, and Mia as specific markers (Fig. 1c), confirming the recent census of cell types in the hypothalamus.

The molecular signatures were validated by single-molecule fluorescent in situ hybridization (smFISH) and Rax-driving lineage tracing, showing the spatial distribution of tanycytes in both ependymal (EZ) and subependymal zone (SEZ) (Fig. 1d–f and Supplementary Fig. 2g). Next, we mapped the expression of cell cycle-related genes (e.g., Mki67 and Mcm2) using single-cell transcriptomic analysis and unexpectedly found that the mitotic genes were predominantly enriched in OPCs but rarely expressed in ME tanycytes (Supplementary Fig. 3a, b).

To reveal the identity of dividing cells, we labeled mitotically active cells by intraperitoneal ethynyldeoxyuridine (EdU) administration in adult mice and costained EdU-positive cells with cell-specific markers such as Rax (tanycytes), Sox2 (NSCs), NG2 (OPCs), Olig2 (OPCs and oligodendrocytes), NeuN (neurons), CC1 (oligodendrocytes), S100β (astrocytes), or Iba1 (microglia) in the hypothalamus. Our results showed that Rax+ tanycytes did not undergo active cell division (Fig. 1d). Further analyses uncovered that up to 60% of labeled dividing cells expressed Sox2, in accordance with the ratio of Olig2+ cells, and approximately 40% of mitotic cells were positive for NG2 (Fig. 1g, h and Supplementary Fig. 3c, d). Given the striking coexpression of NG2 (encoded by Cspg4) or Olig2 with Sox2 revealed by both immunostaining and single-cell transcriptomic analysis (Supplementary Fig. 3e–h), we conclude that Sox2+Olig2+ OPCs rather than Sox2+Olig2– tanycytes predominate the mitotically active cells in ME.

Furthermore, we traced the cell lineage of tanycytes using the Rax-CreERT2–Rosa26-Stop-tdTomato (also known as Ai14) mouse line (Supplementary Fig. 4a). We tested the specificity of instant genetic labeling at 1 day post tamoxifen induction (dpi) by smFISH and found that 99% of tdTomato+ cells in ME were positive for tanycyte marker Snc7a (Fig. 2a), despite the sparse leaky labeling of cells with neuronal morphology in the hypothalamic nuclei (Supplementary Fig. 4b). The cell fate of Rax+ tanycytes was then analyzed at 3, 7, 14, and 30 days following a single dose of tamoxifen injection. We first analyzed the spatial distribution of induced cells and found that the starter cells included ~80% of ventricular cells along EZ and 20% of parenchymal cells in SEZ (Fig. 2b). Notably, the proportion of cells distributed in EZ and SEZ did not change during 30 days of lineage...
tracing (Fig. 2b, c). Second, we assessed the identity of traced cells with multiple cell-specific markers and found that Rax+ tanyocytes did not generate neurons, oligodendrocytes and astrocytes under physiological conditions (Fig. 2d, e). Lastly, we combined lineage tracing with EdU labeling assay to examine whether tanyocytes divide and self-renew. Our data showed that cell division of traced tanyocytes was not detectable, or very rare (Fig. 2f, g). Collectively, these data support our hypothesis that Rax+ tanyocytes are relatively quiescent under physiological conditions.

Rax+ tanyocytes respond to neural injury. It has been reported that adult ependymal cells are postmitotic and remarkably stable without any proliferative potential, even after stroke-induced
Tanyocytes in ME are slow-cycling cells. a Representative image of fluorescent immunoreactivity in adult coronal brain sections from GFAP-CreER<sup>T2</sup>:Ai14 mice. White box indicates the magnified view of the median eminence (ME) on the right. The mice were induced with tamoxifen at postnatal day 60 (P60) and sacrificed at 7 days post induction (dpi) to guide microdissection of ME. Scale bars: 1000 µm (left) and 50 µm (right). b Spectral t-distributed stochastic neighbor embedding (tSNE) representation of the single-cell transcriptomic data with clusters colored and annotated according to known cell-type markers. PTCs pars tuberalis cells, VLMCs vascular and leptomeningeal cells, OPCs oligodendrocyte precursor cells, NKC natural killer cells, MGs microglia, ODs oligodendrocytes. c tSNE scatter plots covering neural cell clusters show the specific expression of Rax, Col25a1, Scn7a and Mib in tanyocytes. d Representative images of smFISH probing for Rax mRNA display no colocalization of Edu with Rax. Numbers refer to subregions shown in the magnified images (right). Yellow arrowheads indicate Rax<sup>+</sup> tanyocytes and dashed white circles signify Edu<sup>+</sup> mitotic cells. EZ ependymal zone, SEZ subependymal zone. Scale bar: 40 µm (left) and 10 µm (right). e, f Shown are Sox2 and Col25a1 gene expression in ME detected by smFISH and three-dimensional (3D) simulation of Scn7a- and Col25a1-positive signals. Scale bars: 50 µm. g Representative images stained for Edu with multiple cell markers including Sox2, NG2, Olig2, NeuN, C11, S100β, and Iba1. Three daily Edu administrations were applied to label dividing cells. Scale bars: 10 µm. h Quantification of the percentage of different cell types among mitotic cells. Values represent mean ± SEM (n = 7, 7, 3, 3, 4, 4, and 4 mice from bottom to top bars).

As specialized ependymal cells, whether tanyocytes are involved in the repair of local injury is not clear. To investigate the regenerative capacity of tanyocytes, we induced a mechanical injury by penetrating an acupuncture needle into ME (Fig. 3a). In contrast to the quiescence of tanyocytes in sham control mice, mechanical injury caused a robust activation of Scn7a<sup>+</sup> tanyocytes in both EZ and SEZ (Fig. 3b). Cell-type analysis of dividing cells revealed that the number of cycling tanyocytes (Sox2<sup>+</sup>Olig2<sup>−</sup> or Scn7a<sup>+</sup>), OPCs (Sox2<sup>+</sup>Olig2<sup>+</sup> or PDGFRα<sup>+</sup>) and astrocytes (S100β<sup>+</sup>) were strikingly increased after injury (Fig. 3c, d and Supplementary Fig. 5a). The results of fate mapping using Rax-CreER<sup>T2</sup>:Ai14 mice further showed that neural injury induced a predominant self-renewal of tanyocytes and drove a small number of tanyocytes to differentiate into OPCs or astrocytes (Fig. 3e and Supplementary Fig. 5b–d), implicating the multipotent differentiation of activated tanyocytes into glial cells. Notably, given that our heat-based antigen retrieval approach deteriorated the tdTomato signal to a certain extent, whether ME tanyocytes could robustly differentiate into OPCs or astrocytes upon injury requires further confirmation.

We then induced targeted neural injury via genetic cell ablation of tanyocytes in Rax-CreER<sup>T2</sup>:Rosa26-Stop-diphtheria toxin receptor (DTR)−2A-GFP (iDTR) mice (Supplementary Fig. 5e–g), whereby the tamoxifen and diphtheria toxin dosage was optimized to avoid a complete ablation of tanyocytes. Subsequently, we evaluated the mitotic activity of cells in the damaged tissue and found that targeted neural injury also induced a significant increase in the number of dividing cells (Fig. 3f and Supplementary Fig. 5h). The further cell-type assessment revealed that tanyocytes self-renewed following induced cell ablation, indicating that residual tanyocytes are capable of regenerating to compensate for their cell loss (Fig. 3f–h). Quantitative analysis confirmed that neural injury increased the number of cycling oligodendrocyte lineage cells and astrocytes (Fig. 3i). We also tracked the fate of Rax<sup>+</sup> tanyocytes after genetically-induced injury and found that a very small number of tanyocytes could transform into astrocytes or OPCs (Supplementary Fig. 5i). Nevertheless, neurogenesis in ME was very limited even after neural injury (Fig. 3i). These results demonstrate that Rax<sup>+</sup> tanyocytes expand themselves with environmental insult for tissue damage repair.

Igf1r signaling is required for tanyocyte preservation and injury-induced tissue repair. Given that tanyocytes possess regenerative capacity, it is important to investigate the regulatory mechanism of their preservation and tanyocyte-mediated tissue repair. To determine the molecular basis for maintaining tanyocyte population, we assessed the expression of receptor tyrosine kinases (RTKs) in our single-cell transcriptomic dataset and found that insulin-like growth factor 1 receptor (Igf1r) and neurotrophic receptor tyrosine kinase 2 (Ntrk2) were enriched in tanyocytes among the 51 RTKs (Supplementary Fig. 6a). Further analysis showed that the expression of Igf1r in tanyocytes was relatively specific, compared to epithelial growth factor receptors (EGFRs), fibroblast growth factor receptors (FGFRs), platelet-derived growth factor receptors (PDGFRs), and neurotrophic receptors (Fig. 4a, b).

To investigate whether signaling through Igf1r is critical for preserving the number of tanyocytes, we used Rax-CreER<sup>T2</sup>:Igf1r<sup>−/−</sup> mice to delete Igf1r in adult tanyocytes by applying tamoxifen at P60 and examined the cell population of tanyocytes at 2 months post genetic deletion. Our data showed that the population of tanyocytes (Sox2<sup>+</sup>Olig2<sup>−</sup> or Scn7a<sup>+</sup>) in SEZ was remarkably declined by loss of Igf1r, but there was no change in OPC number (Fig. 4c, d and Supplementary Fig. 6b, c). Further analysis showed that the deficiency in Igf1r signaling increased the number of apoptotic cells in ME (Supplementary Fig. 6d, e) and pharmacological inhibition of Igf1r signaling in cultured neural progenitors subtly compromised the expression level of stemness-related protein Sox2 (Supplementary Fig. 6f–h). Collectively, these findings suggest that Igf1r signaling is critical for maintaining the tanyocyte population.

We further induced a mechanical injury in Igf1r-deficient mice to observe the role of Igf1r signaling in neural tissue repair (Fig. 4e). The results revealed that genetic ablation of Igf1r in tanyocytes significantly reduced the number of cycling tanyocytes and OPCs in damaged ME (Fig. 4f, g), indicating that inhibition of Igf1r signaling impairs tissue regeneration by tanyocytes.

Adult Rax<sup>+</sup> tanyocytes can contribute to tumor formation. Beyond the regenerative ability, somatic stem cells have frequently been found to generate tumors upon targeted oncogenic mutations. We, therefore, determined to investigate the tumorigenic potential of Rax<sup>+</sup> tanyocytes. Considering the specific anatomic location of craniopharyngioma in between the hypothalamus and pituitary<sup>25,35</sup>, we wonder whether Rax<sup>+</sup> tanyocytes in ME contribute to its formation. A vast majority of adult-onset craniopharyngiomas have been reported to harbor the oncogenic Braf<sup>V600E</sup> point mutation, which is rarely found in meningioma and ependymoma<sup>25,26,36–38</sup>. We introduced an inducible mutant Braf into tanyocytes using Rax-CreER<sup>T2</sup>:Braf<sup>V600E</sup> mice and observed the fate of tanyocytes at 1 week post tamoxifen induction (Fig. 5a). Strikingly, the constitutive activation of Braf quickly incited the mitotic activity of tanyocytes in both EZ and SEZ (Fig. 5b, c). Further cell-type mapping analysis revealed that Braf activation enhanced the expansion of tanyocytes per se but did not promote their differentiation into neurons and astrocytes in the short-term (Fig. 5d–i).

Furthermore, we examined the pathology of ME at 2 months post induction (mpi) of Braf<sup>V600E</sup> expression and found the formation of tumor bearing Braf mutation at the floor of the third

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ventricle (Fig. 6a and Supplementary Fig. 7a, b). The pathological features of induced tumors in ME resembled papillary craniopharyngioma. The width and height of ME were significantly enlarged in the tanycyte-specific mutant mice (Fig. 6b). To validate whether the neoplasm originates from tanycytes, we generated Rax-CreERT2::BrafV600E::Ai14 mice and induced oncogene activation during adulthood. As expected, these animals developed identifiable, well-circumscribed tumors which were clearly positive for tdTomato and Vimentin (Fig. 6c and Supplementary Fig. 7c), suggesting the origin of papillary craniopharyngioma.
craniopharyngioma-like neoplasm from Rax<sup>+</sup> tanycytes. We next assessed the expression pattern of cell-specific markers in the center and marginal zone of tumors generated in the Rax-CreER<sup>T2</sup>•Braf<sup>V600E</sup> model. Importantly, we observed that the tumor cells in the center zone displayed a prominent enrichment of stem cell markers Sox2 and Sox9, but did not express differentiated cell markers Olig2, CC1, S100β, and PDGFRα as observed in oligodendrocytoma and/or astrocytoma (Fig. 6d, e and Supplementary Fig. 7d–f). The cell division detected by EdU incorporation and Ki67 staining occurred in both the central and peripheral zone of the tumor (Fig. 6d, e and Supplementary Fig. 7e). Our quantitative analysis showed that the center zone of tumor tissue mainly consisted of tanycyte-like cells but lacked multiple neural lineage cells, while glia-like cells contributed to the cell expansion in the tumor marginal zone (Fig. 6f). A substantial enhancement of OPC division and differentiation was also observed in the neighbor region of tumor tissues (Fig. 6g), implying the potential activation of OPCs in proximity to tumor tissue.

To characterize the malignancy and invasiveness of papillary craniopharyngioma-like tumor, we further performed xenograft transplantation, histological analysis of multiple organs...
Susceptible to tumor metastasis and in vitro culture of tumor cells. Our data showed that tumor cells derived from Rax-CreER^T2::Braf^{V600E} mice did not cause neoplasms in naked mice or proliferate massively in vitro, and neither did the neoplasms invade into other organs at 6 mpi (Supplementary Fig. 8).

We then dissected the mouse tumor tissues, constructed cDNA libraries, and performed bulk RNAseq. Principal component analysis revealed that the mRNA expression profile of tumor tissues was distinguishable from control ME tissues (Fig. 7a). After applying a standard filtering approach that compared tumor with normal tissues, we identified 291 differentially expressed genes (Fig. 7b–d). The tumor tissues displayed an enriched expression of transcription factors (e.g., Dlx2, Gbx2, Foxb1, and Nkx2.2) critical for brain development, tanyocyte markers (e.g., Ptprz1, Crym, and Gja1), and differentiation markers (e.g., Dcx, Olig1, and Csgp5) (Fig. 7d). The gene ontology analysis of tumor-enriched genes demonstrated that the transcriptional programs involved in cell division, forebrain, diencephalon, and midbrain development were upregulated in the papillary craniopharyngioma model (Fig. 7e and Supplementary Fig. 9a). Surprisingly, genes engaged in immune response were downregulated in tumor tissues (Fig. 7f and Supplementary Fig. 9b). Together, our results suggest that tanyocytes may serve as a cell-of-origin for papillary craniopharyngioma in the adult brain and use the transcriptional programs activated during brain development to evolve into tumor cells.

Discussion

Manipulation of tanyocyte biology could provide a valuable tool for regulating hypothalamic function and modeling disease. Understanding the property, plasticity, and potential of tanyocytes is fundamentally important to maintain their stemness, enhance their function and restrain their tumorigenesis. Here we show that Rax^+ tanyocytes in ME robustly transit from a quiescent to an active state for tissue regeneration when subjected to mechanical injury. Loss of Igf1r signaling in tanyocytes leads to a substantial decrease of tanyocyte population and impairs tissue repair after injury. Importantly, Rax^+ tanyocytes are susceptible to Braf oncogene activation and display tumorigenic potential (Fig. 8).

A combination of bioinformatics analysis, lineage tracing, and pulse-chase assay demonstrate that Rax^+ tanyocytes are largely quiescent, supporting previous studies showing that β-tanyocytes residing in ME cannot proliferate but NG2 glia are actively dividing cells there^{12,31}. Nevertheless, adult tanyocytes are not permanently dormant like ependymal cells in lateral ventricles^{19}. While αSMA^+ ependymal cells fail to activate under growth factor infusion and striatal injury condition^{19}, targeted neural stem cells not only maintain their own stemness and promise their own maintenance and regenerative capacity. These findings provide a potential druggable target to preserve tanyocyte population during aging and promote tissue regeneration after traumatic brain injury.

Somatic stem cells not only maintain their own stemness and tissue homeostasis after injury but also possess the potential of oncogenic transformation^{15,21,22}. Previous reports suggest that Braf^{V600E} expression in neural progenitors is not sufficient for tumorigenesis and needs to cooperate with Ink4a or Cdkn2a locus deficiency in oligodendrocyte or astrocyte precursor cells to induce glioma formation^{40-45}. Unexpectedly, a remarkable result reported here is the discovery that an introduction of mutant Braf^{V600E} into tanyocytes is sufficient to drive tumor formation at the floor of the third ventricle, demonstrating that tanyocytes have tumorigenic potential and are susceptible to Braf mutation. Craniopharyngioma is a heterogeneous brain tumor of uncertain origin and its tumor biology is poorly understood^{35}. Recent studies show that pituitary stem/progenitor cells carrying Cnntb1 but not Braf mutation contribute to the development of pituitary tumors and/or adamanatinomatous craniopharyngioma via
Interestingly, our results reveal that tanycyte-derived tumors in ME mimic papillary craniopharyngioma (frequently carrying BrafV600E mutation) with respect to their anatomic location, genetic mutation, and pathological features, suggesting that hypothalamic Rax\(^+\) tanycytes could serve as a cell-of-origin for papillary craniopharyngioma. In contrast to the salient OPC features in glioma\(^{37,48}\), the tumor cells in our disease model do not abundantly express differentiated cell markers such as Olig2, S100\(\beta\), and CC1. Notably, craniopharyngioma was previously assumed to derive from squamous...
Fig. 5 Introduction of BrafV600E stimulates mitotic activation of Rax+ tanycytes. a Experimental scheme describing the induction of somatic genetic mutation in tanycytes using Rax-CreERT2::BrafV600E mice and detection of cell division in ME. b Sample confocal images showing the mitotic cells in the ME of control and Braf-mutant mice. Scale bar: 50 µm. c Quantification of EdU+ cell number in EZ and SEZ of control and mutant mice. Boxes represent IQR, whiskers extend to ±1.5 IQR, and significance was analyzed by unpaired two-tailed Student’s t test (n = 12 and 14 sections from three mice for control and mutant groups). ***P < 0.001. d Representative confocal images stained for EdU with Sox2, Olig2, CC1, and S100β in Rax-CreERT2::BrafV600E mice receiving tamoxifen injection. Scale bar: 20 µm. e Quantification of the number of mitotic cells expressing different cell markers (n = 20 sections from at least three mice from bottom to top boxes). *P < 0.05; ***P < 0.001; N.S. not significant. f Sample images showing mutant BrafV600E activates mitotic division of Sox2+ Olig2− tanycytes in ME. White arrowheads signify Sox2+ Olig2+ OPCs while yellow arrowheads indicate tanycytes. Scale bar: 10 µm. Source data are provided as a Source Data file. The precise P values are summarized in Supplementary Data 3.
epithelial cells by clinical researchers, but we did not find a robust transcriptomic similarity between tanycytes and esophageal squamous epithelia, or between tanycyte-derived tumor and esophageal squamous cell carcinoma (Supplementary Fig. 10). The data from our transcriptomic profiling further demonstrate that tanycyte-derived tumors display enrichment of genes involved in the brain but not pituitary development. Due to the lack of transcriptomic data of human papillary craniopharyngioma in previous studies, further investigation is required to uncover the similarity and difference between mouse and human tumor tissues.

Given that previous studies have subdivided tanycytes into α1, α2, β1, and β2 subtypes along the third ventricle, here we have to emphasize that our study focuses on Rax+ tanycytes,
Fig. 6 Rax<sup>+</sup> tanycytes may serve as a cell-of-origin of papillary craniopharyngioma. a Hematoxylin and eosin (HE) staining of ME in control and Rax-CreERT2::BrafV600E mutant mice at 2 months post tamoxifen injection. Scale bar: 50 µm. b Quantification of the width and height of ME in control and mutant mice. Boxes represent IQR, whiskers extend to ±1.5 IQR, and significance was analyzed by unpaired two-tailed Student’s t test (n = 24 and 7 sections from four mice for control and mutant groups). **P < 0.01. c Representative confocal images showing the coating of tdTomato, Vimentin, and EdU in Rax-CreERT2::BrafV600E::Ai14 mice at 2 months post tamoxifen injection. Scale bar: 100 µm. d Representative confocal images of tumor tissues stained for Olig2, Sox2, S100, and water ad libitum. The animal facility was maintained at a temperature of 21 °C and 50–60% humidity. All mice in the study were backcrossed to the C57BL/6N background for at least six generations. To perform lineage tracing, targeted neural progenitors were dissected from ten female adult C57BL/6N mice and pooled together. Single-cell RNA sequencing (scRNA-seq) was performed according to the manufacturer’s instructions. The library preparation and sequencing were done by Beijing Genomics (BGII). The reads were analyzed with R package. The first step is to normalize the data by a global-scaling normalization method that normalizes, multiplies a scale factor (10,000), and log-transforms the feature expression. Next, features were selected to exhibit high cell-to-cell variation in the aME dataset by Seurat “FindVariableGenes” command with the following parameters: mean function equal to ExpMean, dispersion function equal to LogVMR, x.low.cutoff equal to 0.02, x.high.cutoff equal to 8, and y.cutoff equal to 0.3. The number of scRNA expression data in Supplementary Fig. 3. Second, we amplified the target DNA fragment and verified the activation of Braf using XbaI restriction enzyme (NEB, R0145) which cut the recognition site introduced by the source paper reporting the generation of BrafV600E mouse line21. Amplified the target DNA fragment and verified the activation of Braf using XbaI restriction enzyme (NEB, R0145) which cut the recognition site introduced by the source paper reporting the generation of BrafV600E mouse line21. Amplified the target DNA fragment and verified the activation of Braf using XbaI restriction enzyme (NEB, R0145) which cut the recognition site introduced by the source paper reporting the generation of BrafV600E mouse line21.
3 days in adult Rax-CreERT2::iDTR mice that had received a single tamoxifen (66 mg/ml) injection 3 days before. The procedure has been described in our previous study52.

Tissue section preparation, EdU staining, and immunohistochemistry. The experimental mice were anesthetized by intraperitoneal injection of 4% chloral hydrate and then transcardially perfused with saline followed by 4% paraformaldehyde (PFA) in phosphate-buffered saline (PBS). Mouse brains were dissected, post-fixed for 4–6 h in 4% PFA at 4 °C, and subsequently cryo-protected in 20% sucrose in PBS for 12 h followed by 30% sucrose for 24 h. Tissue blocks were prepared by embedding in Tissue-Tek O.C.T. Compound (Sakura 4583). The brain sections (20 μm in thickness) were prepared using a cryostat microtome (Leica, CM3050S), dried for 30 min at room temperature in the dark and stored in −20 °C freezer. For immunostaining, the tissue sections were washed with 1×TBS (pH 7.4, containing 3 mM KCl, 25 mM Trisma base, and 137 mM NaCl) and pre-blocked with 1×TBS++ (TBS containing 5% donkey serum and 0.3% Triton X-100) for 1 h at room temperature, followed by incubation with primary antibodies diluted in TBS++ over night at 4° C. The primary antibodies used in this study included Sox2 (Goat; R&D Systems; #KOY0317071; 1:500), NG2 (Rabbit; Millipore; #3018740; 1:200), Olig2 (Goat; R&D Systems; #UPA0617051; 1:200), NeuN (Mouse; Abcam; #ab104224; 1:1000), HuC/D (Mouse; Molecular Probes; #A21271; 1:1000), CCI (Mouse; Calbiochem; #OP98; 1:200), S100 (Rabbit; Abcam; #ab686; 1:400), Ibα1 (Rabbit; Wako; #019-19741; 1:1000), GFAP (Goat; Rockland; #33302; 1:1000), RFP (Rabbit; Rockland; #35055; 1:1000), GFAP (Rabbit; DAKO; #Z0334; 1:1000), IgG1, Cspg5 (Goat; Thermo; 1:1000), Iba1 (Rabbit; Zymed; #019-19741; 1:1000), Sox9 (Rabbit; Abcam; #ab185966; 1:250); PDGFβ (Goat; R&D Systems; #AF464; 1:200), Vimentin (Rabbit; DAKO; #Z0334; 1:500), and others.

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Fig. 7 Bulk RNAseq analysis of tumor tissues. a–c Principal component (PC) analysis (a), MA plot (b), and volcano plot (c) showing the differential gene expression in ME tissues of control and Rax-CreERT2::BrafV600E mutant mice at 2 months post tamoxifen injection. d A heatmap showing the differential gene expression of control and papillary craniopharyngioma-like tumor tissues. A set of upregulated and downregulated transcription factors, tanycyte markers, and differentiation markers in the mouse papillary craniopharyngioma are highlighted in the table. e, f Gene ontology network analysis of upregulated (e) and downregulated (f) genes in the tumor tissues. The major clusters with different gene ontology are labeled with different colors.

3 days in adult Rax-CreERT2::iDTR mice that had received a single tamoxifen (66 mg/ml) injection 3 days before. The procedure has been described in our previous study52.
To combine smFISH with the pulse-chase assay, we further treated the tissue sections with 0.5% Triton-X-100 and EdU detection solution. After staining, the slides were mounted and imaged for further analysis.

To detect the mRNA expression level of Miu, we used the hybridization chain reaction (HCR) approach. We designed the probe sequences (Supplementary Table 2) using the coding and 3’UTR regions of Miu and synthesized the probes in Sangon Biotech, China. The brain sections were permeabilized in 70% ethanol for 16 h at 4 °C, followed by 0.5% Triton X-100 in 1×PBS at 37 °C for 1 h, and treated with 10 µg/mL Protease K to improve mRNA accessibility. After two washes with 1×PBS at room temperature, sections were pre-hybridized in probe hybridization buffer (30% formamide, 5×SSC, 9 mM citric acid, 0.1% tween 20, 50 µg/mL heparin, 1×Denhardt’s solution, and 10% dextran sulfate) for 1 h at 37 °C and then incubated in probe hybridization buffer containing HCR probes (10 µM for each) at 37 °C for 3 h. After mRNA hybridization, the washing and amplification steps were performed as previously described.

**TUNEL assay.** For TUNEL assay, cryopreserved sections were fixed in 4% PFA for 20 min at room temperature and incubated with sodium citrate solution containing 0.3% Triton X-100 for 30 min. Subsequently, TUNEL reaction mixture (Roche, 11684795910) was added and incubated with brain sections for 1 h at 37 °C in the dark. After staining, sections were coverslipped with a mounting medium and observed under a microscope.

**Western blotting analysis.** The neural progenitor cells were cultured at 37 °C with 5% CO2 in NeuroCult Basal Medium (STEMCELL, #05700) supplemented with NeuroCult Proliferation Supplement (STEMCELL, #05701) EGF (PeproTech, AF-115-09) and bFGF (PeproTech, AF-450-33). The cells were treated with 1 µM OSI-906 (MedChemExpress, HY-10191) to block Igf1r signaling for 48 h. Subsequently, the cell lysates were spun down at 20,000×g for 20 min and the supernatants were denatured at 95 °C for 10 min. The equivalent denatured samples were subjected to SDS-PAGE, transferred to PVDF membranes and subjected to antigen retrieval. Subsequently, the tissue sections were dehydrated with 20–30% DEPC-treated sucrose for 24 h. Subsequently, the tissues were rapidly frozen on dry ice, embedded in O.C.T. compound, cryosectioned at a thickness of 20 µm, and mounted onto SuperFrost Plus microscope slides. The probes targeting against Rax (19046 A), Scn7a (18199B), and Col25a1 (19032A) were designed and validated by Advanced Cell Diagnostics. RNAscope v2 Assay (Advanced Cell Diagnostics, #320511) was used for all smFISH experiments according to the manufacturer’s protocol. Briefly, the brain sections were dried at 55 °C for 2 h, rinsed with 1×PBS, treated with 3% hydrogen peroxide in methanol, and subjected to antigen retrieval. Subsequently, the tissue sections were dehydrated with 100% ethanol and incubated with mRNA probes for 2 h at 40 °C. The specific signals were then amplified with multiplexed amplification buffer and detected with TSA Plus fluorophore (Perkin Elmer, #NEL753001KT) for 30 min. To combine smFISH with the pulse-chase assay, we further treated the tissue sections with 0.5% Triton-X-100 and EdU detection solution. After staining, the slides were mounted and imaged for further analysis.

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and probed with the antibodies against Sox2 (goat; R&D; AF2018; 1:2000), GFRα1 (rabbit; CST; 30272; 1:1000), phosphorylated GFRα1 (rabbit; CST; 30243; 1:1000) or α-Tubulin (mouse; CST; 3875; 1:5000). Blots were visualized with SuperSignal West Pico chemiluminescence (Thermo, 3457) and imaged with a Bio-Rad ChemiDoc XRS system. The intensity of stained bands was quantified using Image J software.

In vitro culture of tumor cells and xenograft transplantation. The tumor tissues were microdissected from Rax-CreERT2;BrafV600E mice at 2 months post tamoxifen induction and normal and tumor tissues, we microdissected the ME region from three control and locally, we only had a limited number of brain sections (number of cells expressing various cell markers. Given that tumor cells expanded quantification.

Mechanical injury. Adult C57BL/6N mice, Rax-CreERT2;Ai14 mice, and Rax-CreERT2;TgIgf1rf/b mice were anesthetized with 4% chloral hydrate and subjected to mechanical injury with an acupuncture needle penetrating the midline of brains 2.3 mm posterior to bregma and 7 mm below dura. At 24 h after mechanical injury, the mice were intraperitoneally injected with EdU (Ark Pharm, AK163060-1g) for 3 consecutive days (50 mg/kg body weight), followed by tissue collection to analyze the mitotic activity of cells in ME.

Histology analysis. The mouse brains, liver, spleen, lung, and kidney embedded in O.C.T. compound were sectioned at a thickness of 10–20 µm with cryostat microtome (Leica, CM3050S). The tissue sections were washed with 1×PBS buffer, sequentially stained with hematoxylin and eosin (H&E, Beyotime, #C0105), and then dehydrated with gradient ethanol. Subsequently, we clarified the stained tissue sections with xylene and mounted the slides with DPX mounting medium (Sigma, #06522). The images were taken with a Leica SP8 microscope and Nikon ECL IPSE Ci-L microscope.

Cell quantification. To quantify the number of cells expressing cell-type-specific markers, we serially sectioned the ME and analyzed the positive cells in one out of five brain sections spanning the whole ME (from ~1.58 mm to ~2.3 mm posterior to bregma). Quantification of cells labeled by EdU and expressing different cell-type markers (Sox2, Olig2, NG2, CC1, S100β, and NeuN) in ME was conducted using Carl Zeiss AIM 2.3 software, Leica LAS X software or Image J 1.52p (NIH software). Generally speaking, at least three animals and five brain sections from each animal were used for quantification. To investigate the spatial distribution of cells, we respectively deblinded in this study. Rstudio, GraphPad Prism (v8), and Microsoft Excel 2016 were used to test covariates and normality, calculate statistical significance, and prepare quantitative graphs. Group data are presented as bar plots showing mean ± standard error of the mean (SEM) or box plots wherein boxes represent interquartile range (IQR), whiskers extend to ±1.5 IQR, dots represent outliers and bold black lines indicate median values. Statistical analysis was calculated using unpaired two-tailed Student’s t test or one-way ANOVA with Sidak’s multiple comparison test. If not otherwise noted, statistical significance was indicated as follows: *P < 0.05; **P < 0.01; ***P < 0.001; N.S. not significant. The precise P values and sample size were provided as an additional Supplementary Data S3. All results shown in the study are representative of at least two independent experiments with similar results.

Reporting summary. Further information on experimental design is available in the Nature Research Reporting Summary linked to this paper.

Data availability. The single-cell and bulk RNAseq data in this study have been deposited in the Gene Expression Omnibus (GEO) with accession number GSE132943. All other relevant data that support the findings of this study are available within the article and its Supplementary Information files or from the corresponding authors upon reasonable request. A reporting summary for this article is available as a Supplementary Information file. Source data are provided with this paper.

Code availability. The computational code used in this work is available upon request.

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45. Author contributions
The authors declare no competing interests.

46. Correspondence
Q.F.W., W.M., and S.L. designed all experiments. W.M., S.L., and J.X. performed the experiments and statistical analyses. Q.F.W., H.W., Z.C., and X.G. performed bioinformatics analysis of single-cell and bulk transcriptomic datasets. F.L. and L.Q. contributed to the generation of the cDNA library for sequencing from tanyctye-derived tumor tissues. G.H. and C.L. provided technical advice. Q.F.W and S.L. wrote the paper.

47. Competing interests
The authors declare no competing interests.

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