Development/Plasticity/Repair

# Gli3 Regulates Vomeronasal Neurogenesis, Olfactory Ensheathing Cell Formation, and GnRH-1 Neuronal Migration

©Ed Zandro M. Taroc,¹ Ankana S. Naik,¹ ©Jennifer M. Lin,¹ Nicolas B. Peterson,¹ David L. Keefe Jr.,² Elizabet Genis,¹ ©Gabriele Fuchs,¹ Ravikumar Balasubramanian,² and ©Paolo E. Forni¹

<sup>1</sup>Department of Biological Sciences; The RNA Institute, and the Center for Neuroscience Research; University at Albany, State University of New York, Albany, New York 12222, and <sup>2</sup>Harvard Reproductive Sciences Center and The Reproductive Endocrine Unit of the Department of Medicine, Massachusetts General Hospital, Boston, Massachusetts 02114

During mammalian development, gonadotropin-releasing-hormone-1 neurons (GnRH-1ns) migrate from the developing vomeronasal organ (VNO) into the brain asserting control of pubertal onset and fertility. Recent data suggest that correct development of the olfactory ensheathing cells (OEC) is imperative for normal GnRH-1 neuronal migration. However, the full ensemble of molecular pathways that regulate OEC development remains to be fully deciphered. Loss-of-function of the transcription factor Gli3 is known to disrupt olfactory development, however, if Gli3 plays a role in GnRH-1 neuronal development is unclear. By analyzing Gli3 extra-toe mutants (Gli3 Xt/Xt), we found that Gli3 loss-of-function compromises the onset of achaete-scute family bHLH transcription factor 1 (Ascl-1)<sup>+</sup> vomeronasal progenitors and the formation of OEC in the nasal mucosa. Surprisingly, GnRH-1 neurogenesis was intact in Gli3 Xt/Xt mice but they displayed significant defects in GnRH-1 neuronal migration. In contrast, Ascl-1 null mutants showed reduced neurogenesis for both vomeronasal and GnRH-1ns but less severe defects in OEC development. These observations suggest that Gli3 is critical for OEC development in the nasal mucosa and subsequent GnRH-1 neuronal migration. However, the nonoverlapping phenotypes between Ascl-1 and Gli3 mutants indicate that Ascl-1, while crucial for GnRH-1 neurogenesis, is not required for normal OEC development. Because Kallmann syndrome (KS) is characterized by abnormal GnRH-1ns migration, we examined whole-exome sequencing data from KS subjects. We identified and validated a *GLI3* loss-of-function variant in a KS individual. These findings provide new insights into GnRH-1 and OECs development and demonstrate that human *GLI3* mutations contribute to KS etiology.

Key words: Ascl-1; Gli3; GnRH-1; Kallmann syndrome; olfactory ensheathing cells; vomeronasal sensory neurons

### Significance Statement

The transcription factor Gli3 is necessary for correct development of the olfactory system. However, if Gli3 plays a role in controlling GnRH-1 neuronal development has not been addressed. We found that Gli3 loss-of-function compromises the onset of Ascl-1<sup>+</sup> vomeronasal progenitors, formation of olfactory ensheathing cells in the nasal mucosa, and impairs GnRH-1 neuronal migration to the brain. By analyzing Ascl-1<sup>null</sup> mutants we dissociated the neurogenic defects observed in Gli3 mutants from lack of olfactory ensheathing cells in the nasal mucosa, moreover, we discovered that Ascl-1 is necessary for GnRH-1 ontogeny. Analyzing human whole-exome sequencing data, we identified a *GLI3* loss-of-function variant in a KS individual. Our data suggest that *GLI3* is a candidate gene contributing to KS etiology.

### Introduction

The olfactory placode gives rise to multiple cell types, including migratory/pioneer neurons, terminal nerve cells, gonadotropin

releasing hormone-1 (GnRH-1) neurons (GnRH-1ns), olfactory, and vomeronasal sensory neurons (VSNs) (Wray et al., 1989; Miller et al., 2010; Casoni et al., 2016). GnRH-1ns play a central

Received Aug. 13, 2019; revised Oct. 18, 2019; accepted Nov. 17, 2019.

Author contributions: E.Z.M.T., A.S.N., J.M.L., G.F., R.B., and P.E.F. designed research; E.Z.M.T., A.S.N., J.M.L., N.B.P., D.L.K., E.G., G.F., R.B., and P.E.F. performed research; E.Z.M.T., A.S.N., J.M.L., N.B.P., D.L.K., E.G., G.F., R.B., and P.E.F. analyzed data; E.Z.M.T. and P.E.F. wrote the first draft of the paper; E.Z.M.T., A.S.N., J.M.L., D.L.K., G.F., R.B., and P.E.F. edited the paper; E.Z.M.T., G.F., R.B., and P.E.F. wrote the paper; P.E.F. contributed unpublished reagents/analytic tools.

This work was supported by the Eunice Kennedy Shriver National Institute of Child Health and Human Development of the National Institutes of Health (Grants 1R15HD09641101 and 1R01HD097331-01 to P.F.) and by the National Institute of Deafness and other Communication Disorders of the National Institutes of Health (Grant 1R01DC017149-01A1 to P.F.). R.B. is supported by the Eunice Kennedy Shriver National Institute of Child Health and Development (Grants K23 HD077043, R01 HD096324, and P50 HD HD028138). The content of this manuscript

role in controlling the reproductive axis of vertebrates. During embryonic development GnRH-1ns migrate along the terminal nerve (Schwanzel-Fukuda and Pfaff, 1989; Casoni et al., 2016; Taroc et al., 2017) from the vomeronasal organ into the hypothalamus. Once in the brain the GnRH-1ns control gonadotropin release from the pituitary gland. Defects in neurogenesis and/or migration of GnRH-1ns can lead to Kallmann syndrome (KS), a genetic disorder characterized by pubertal failure due to congenital hypogonadotropic hypogonadism (HH) and anosmia (impaired sense of smell), or to normosmic idiopathic hypogonadotropic hypogonadotropic hypogonadism (nIHH) (Pingault et al., 2013; Forni and Wray, 2015).

In the last three decades, various causal genes for KS have been identified and disruption of these genetic pathways lead to KS through aberrant development of olfactory placode derivatives and/or impaired migration of GnRH-1ns. Notably, recent studies show that the ability of the GnRH-1ns to migrate from the olfactory pit to the brain is also critically dependent on the correct development of a specialized neural crest derived glial cells termed "olfactory ensheathing cells" (Barraud et al., 2013; Pingault et al., 2013). In keeping with this notion, of the known KS genes, two genes, SOX10 and *ANOS1* have been recently identified as key factors controlling OECs development (Hu et al., 2019). Thus, understanding the genetic pathways governing OEC development could provide more mechanistic clues into the basis of aberrant GnRH neuronal migration and therefore, KS.

Gli3, together with Gli1 and Gli2, are key transcription factors involved in the sonic hedgehog (Shh) signaling. In the absence of Shh signaling, Gli3 and Gli2 act as transcriptional repressors. However, in the presence of Shh, Gli1,2 and Gli3 function as transcriptional activators (Sasaki et al., 1999; Niewiadomski et al., 2014). A spontaneous murine model of the *Gli3* gene, Gli3 pdn/pdn has previously implicated a potential role for Gli3 in GnRH neuronal migration. While the hypomorphic Gli3 pdn/pdn model shows a delayed, but not missing, GnRH-1 neuronal migration (Naruse et al., 1994), the more severe Gli3-null mutants Gli3 extra-toe (Gli3 xt/Xt) fail to develop a functional olfactory system (Keino et al., 1994; Balmer and LaMantia, 2004; Besse et al., 2011). However, GnRH1 neuronal development and migration in Gli3 xt/Xt has not been analyzed.

A candidate gene study in humans has revealed missense variants in the *GLI3* gene in HH (Quaynor et al., 2016) but no functional validation has been performed to ascertain the causality of these variants. Furthermore, *GLI3* mutations in humans have been reported in nonsyndromic forms of polydactyly and syndromic forms of polydactyly of which a subset of patients display neonatal hypogonadism (micropenis and undescended testes) (Johnston et al., 2010). Despite these observations, the precise role of Gli3 in GnRH neuronal biology remains to be fully elucidated.

By analyzing  $Gli3^{Xt/Xt}$  mutants across the key early developmental stages of embryonic olfactory and GnRH neuronal development, we found that Gli3 loss-of-function compromises the onset of Ascl-1 + vomeronasal progenitors and disrupts the for-

is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. We thank Dr. Hirotomo Saitsu (Hamamatsu University School of Medicine, Hamamatsu, Japan) for sharing Gli3 luciferase reporter plasmids; Dr. Susan Wray for sharing the anti GnRH-1 (SW) antibody; and Dr. Melinda Larsen, University at Albany, for sharing frozen samples of Gli1LacZ embryos.

 $Correspondence \ should \ be \ addressed \ to \ Paolo \ E. \ Forniat \ pforni@albany.edu.$ 

https://doi.org/10.1523/JNEUROSCI.1977-19.2019

Copyright © 2020 Taroc et al.

This is an open-access article distributed under the terms of the Creative Commons Attribution License Creative Commons Attribution 4.0 International, which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

mation of OECs in the nasal mucosa. Notably, in Gli3 Xt/Xt, although GnRH-1 neurogenesis was preserved as in controls, the GnRH-1ns were unable to migrate and populate the hypothalamus. To further dissect the precise roles of Gli3 and Ascl-1, we also analyzed *Ascl-1-null* mutants and found significant reduction for both vomeronasal and GnRH-1ns but less severe defects in OECs' development than in Gli3 Xt/Xt, suggesting the OEC development is critically dependent on *Gli3* rather than *Ascl-1*. By whole-exome sequencing of human KS patients, we identified and functionally validated the first *GLI3* loss-of-function variant in a KS individual. These findings provide new insights into GnRH-1 and OECs development and demonstrate that human *GLI3* mutations contribute to KS etiology.

#### **Materials and Methods**

Animals. Gli3 Xt (Schimmang et al., 1992) and Ascl-1 tm1.1(Cre/ERT2)Jeo/J mice (Kim et al., 2011) on C57BL/6J background were purchased from (Jackson Laboratories). Both colonies were maintained on C57BL/6J. The genotypes of Gli3 Xt mice were established by PCR analysis using the following primers: Gli3-C3F: GGCCCA AACATCTACCAACACA TAG, Gli3-C3R: GTTGGCTGCTGCATGAAGACTGAC; Gli3-XtJ580F: TACCCCAGCAGGAGACTCAGATTAG; Gli3-XtJ580F: AAACCCGT GGCTCAGGACAAG. Ascl-1 CreERT2 were genotyped following Ascl-1 tm1.1(Cre/ERT2)Jeo/J protocol available on jax.org website. Amplification products were analyzed by agarose gel electrophoresis. Animals were killed using CO<sub>2</sub>, followed by cervical dislocation. Mutant and wild-type mice of either sex were used, E15.0 frozen Gli1-LacZ embryos (Gli1 tm2Alj/J) (Bai et al., 2002) were kindly donated to us by Dr. Melinda Larsen. All mouse studies were approved by the University at Albany Institutional Animal Care and Use Committee.

*Tissue preparation.* Embryos were collected from time-mated dams where the emergence of the copulation plug was taken as E0.5. Collected embryos were immersion-fixed in 3.7% Formaldehyde/PBS at 4°C for 3–5 h. Postnatal animals were perfused with 3.7% Formaldehyde/PBS. All samples were then cryoprotected in 30% sucrose overnight, then frozen in O.C.T (Tissue-TeK) and stored at  $-80^{\circ}$ C. Samples were cryosectioned using CM3050S Leica cryostat and collected on Superfrost plus slides (VWR) at 12–16  $\mu$ m thickness for immunostainings and 18–25  $\mu$ m for *in situ* hybridizations (ISH).

Immunohistochemistry. Primary antibodies and dilutions used in this study were as follows: goat-α-Neuropilin-2 (1:3000; R&D Systems), goat- $\alpha$ -Neuropilin-1 (1:400; R&D Systems), mouse- $\alpha$ -ROBO2(1:50; Santa Cruz Biotechnology) chicken-α-peripherin (1:1500, Abcam), rabbit- $\alpha$ -peripherin (1:2000, Millipore), SW rabbit- $\alpha$ -GnRH-1 (1:6000, Susan Wray; NIH), goat- $\alpha$  olfactory marker protein (1:4000; WAKO), mouse- $\alpha$ -GAD67 (1:200; Santa Cruz Biotechnology), goat- $\alpha$ -AP-2 $\epsilon$ (1:1000; Santa Cruz Biotechnology), mouse- $\alpha$ -Meis2 (1:500; Santa Cruz Biotechnology), rabbit-α-phospho-Histone-H3 (1:400; Cell Signaling Technology), rat-α-phospho-Histone-H3 (1:500, Abcam), goat- $\alpha$ -Sox10 (1:100; Santa Cruz Biotechnology), goat- $\alpha$ -Collagen IV (1:800, Millipore), goat- $\alpha$ -Gli3 (1:200; R&D Systems), rabbit- $\alpha$ -Hes-1 (1:200; Cell Signaling Technology), and mouse- $\alpha$ -Ascl-1 (1:30; Santa Cruz Biotechnology). Antigen retrieval was performed in a citrate buffer before incubation with chicken- $\alpha$ -peripherin, rabbit- $\alpha$ -phospho-Histone-H3, rat- $\alpha$ -phospho-Histone-H3, goat- $\alpha$ -AP-2 $\epsilon$ , mouse- $\alpha$ -Meis2, goat- $\alpha$ -Gli3, rabbit- $\alpha$ -Hes-1, mouse- $\alpha$ -Ascl-1, goat- $\alpha$ -Sox10, mouse- $\alpha$ -ROBO2, and mouse- $\alpha$ -GAD67 antibodies. For immunoperoxidase staining procedures, slides were processed using standard protocols (Forni et al., 2013) and staining was visualized (Vectastain ABC Kit; Vector Labs) using diaminobenzidine (DAB) in a glucose solution containing glucose oxidase to generate hydrogen peroxide; sections were counterstained with methyl green. For immunofluorescence, species-appropriate secondary antibodies were conjugated with Alexa Fluor-488, Alexa Fluor-594, or Alexa Fluor-568 (Invitrogen and Jackson Laboratories) as specified in the legends. Sections were counterstained with 4',6'-diamidino-2phenylindole (1:3000; Sigma-Aldrich) and coverslips were mounted with Fluoro Gel (Electron Microscopy Services). Confocal microscopy pictures were taken on a Zeiss LSM 710 microscope. Epifluorescence pictures were taken on a Leica DM4000 B LED fluorescence microscope equipped with a Leica DFC310 FX camera. Images were further analyzed using FIJ/ImageJ software. Each staining was replicated on at least three different animals for each genotype.

In situ *hybridization*. Digoxigenin-labeled cRNA probes were prepared by *in vitro* transcription (DIG RNA labeling kit; Roche Diagnostics) from the following templates:

Semaphorins 3A (Kagoshima and Ito, 2001), Gli3 riboprobe RP\_080717\_03\_E04 (forward primer GCAGAATTATTCCGGTCAG TTC; reverse primer TCAGTCTTTGTGTTTGTGGTCC). In situ hybridization was performed as described previously (Lin et al., 2018) and visualized by immunostaining with an alkaline phosphatase conjugated anti-DIG (1:1000), and NBT/BCIP developer solution (Roche Diagnostics). Sections were then counter-immunostained with antibodies against both chicken- $\alpha$ -peripherin, and SW rabbit- $\alpha$ -GnRH-1, as described above for immunofluorescence.

*X-gal staining*. Sections were rehydrated in PBS and then incubated in a solution of 5 mm potassium ferrocyanide, 5 mm potassium ferricyanide, 2 mm MgCl<sub>2</sub>, 0.1% Tween, and 0.1% 5-bromo-4-chloro-3-indolyl-b-D-galactoside/dimethylformamide at 37°C O.N. After the enzymatic reaction, slides were counterstained with 1% Eosin-Y (Electron Microscopy Services).

Cell quantifications. Quantification of GnRH-1 neuronal distribution in mutant and control animals was performed in the nasal region (VNO, axonal tracks surrounding the olfactory pits), forebrain junction and brain (all the cells that accessed the olfactory bulb and were distributed within the forebrain). The number of cells was calculated for each animal as the average of the counted cells per series multiplied by the number of series cut per animal. Differences at each interval between genotypes were assessed by unpaired t test.

Number of vomeronasal apical and basal neurons was counted after AP-2 $\epsilon$  and Meis2 immunostaining, as cells/section on an average of 4 sections/embryo. Progenitor cells were counted after immunostaining for phospho-histone H3, Ascl-1, and Hes-1 and were quantified as described above for vomeronasal cells. Sox10  $^+$  cells were quantified in a 334,500  $\mu$ m<sup>2</sup> field area for an average of 4 VNO sections per animal, only Sox10  $^+$  cells proximal to the VNO were counted. Frequency of Sox10 cells on vomeronasal bundles was calculated by counting the number of Sox10 I.R. cells/ $\mu$ m fiber length. Means  $\pm$  SEs were calculated on at least three animals per genotype, t test was used to assess differences between groups.

Densitometric analysis of the median eminence. Brains of 5-month-old male Gli3  $^{\rm Xt}$  WT and heterozygous mice (n=3) were cryosectioned coronally at 20  $\mu$ m. Three series for each animal were stained for GnRH in DAB. Brightfield images of comparable sections for each series were taken for quantification. Images were processed with color deconvolution and the mean gray value was measured for a standardized ROI to calculate the average optical density for each animal.

Site-directed mutagenesis. The GLI-3 bs-2 plasmid carrying the human GLI3 (Kinzler et al., 1988) was purchased from Addgene. F387F\*fs mutation was generated in the vector pCDNA3 hGli3 using Q5 site directed mutagenesis kit (NEB) using primers Gli3 F387Ffs\* Fwd CCAACACA GAGGCCTATTCC and Gli3 F387Ffs\* Rev AAAGTTGGGGCAGGGT GG following the manufacturer's instruction. Mutation was validated by sequencing before use in experiments.

Transfection and luciferase reporter assay.  $1 \times 10^4$  HeLa cells (ATCC CCL2) were seeded into a 96-well plate 24 h before transfections. 100 ng of either reporter plasmid (8 × 3 GLI-BS Luc or 8 × 3 Mut GLI-BS Luc) and 100 ng of the effector plasmid (hGli3 WT or F387F\*fs) and 20 ng of pRL-SV40 were cotransfected using Lipofectamine 3000 (Thermo Fisher). pRL-SV40 coding for Renilla luciferase was used as a transfection control and for normalization. Cells were harvested 48 h posttransfection and Firefly and Renilla luciferase intensities were measured using Glomax 96 microplate luminometer (Promega) and Dual-Luciferase Reporter Assay System (Promega).

Human phenotyping and genotyping: WES analysis of KS individuals for GLI3 loss-of-function mutations. A total of 632 KS subjects at the Harvard Reproductive Endocrine Sciences Center were examined for GLI3 loss-

of-function mutations in this study. KS was defined by: (1) absent or incomplete puberty by age 18 years; (2) serum testosterone <100 ng/dL in men or estradiol <20 pg/ml in women in the face of low or normal levels of serum gonadotropins; (3) otherwise normal anterior pituitary function; (4) normal serum ferritin concentrations; (5) normal MRI of the hypothalamic-pituitary region (Balasubramanian and Crowley, 2011); and (6) impaired olfaction as determined by the University of Pennsylvania Smell Identification Test (UPSIT) (Lewkowitz-Shpuntoff et al., 2012; Doty, 2007). Clinical charts, biochemical testing (including overnight neuroendocrine testing for LH pulse profiles) and patient questionnaires of KS subjects were reviewed for phenotypic evaluation as described previously (Pitteloud et al., 2002). Whole-exome sequencing (WES) data in the KS cohort was performed on peripheral bloodderived DNA using either Nimblegen SeqCap target enrichment kit (Roche) or a custom Illumina capture kit (ICE) and the detailed variant-calling and annotation used have been described previously (Guo et al., 2018). WES data were then queried for GLI3 (RefSeq: NM\_000168.6) loss-of-function variants: defined as those variants resulting in protein alteration in the canonical GLI3 transcript as determined by Variant Effect Predictor (VEP) (127), resulting in nonsense, and essential splice site mutations; or (b) indels resulting in frame-shift mutations; AND (b) with ethnicity-specific minor allele frequency (MAF) <0.1% minor allele frequency (MAF) in gnomAD database (http://gnomad.broadinstitute.org) (Karczewski et al., 2019). To ascertain oligogenicity, KS proband(s) with GLI3 loss-of-function variants were also examined for rare sequence variants (RSVs) in other known KS/nIHH genes [ANOS1 (NM\_000216.4), CHD7 (NM\_017780.4), FEZF1 (NM\_001160264.2), FGF8 (NM\_006119.4), FGFR1 (NM\_023110.2), GNRH1 (NM\_000825.3), GNRHR (NM\_000406.2), HS6ST1 (NM\_ 004807.3), KISS1 (NM\_002256.4), KISS1R (NM\_032551.5), NR0B1 (NM\_000475.4), NSMF (NM\_001130969.1), PROK2 (NM\_001126128.2), PROKR2 (NM\_144773.3), SEMA3A (NM\_006080.3), SMCHD1 (NM\_ 015295.2), SOX2 (NM\_003106.4), TAC3 (NM\_013251.4), TACR3 (NM\_001059.3), and WDR11 (NM\_018117.12)]. RSVs in these other HH genes were defined as single nucleotide variants resulting in protein alteration in canonical transcript as determined by Variant Effect Predictor (VEP) (127) resulting in missense, nonsense, and essential splice site mutations; or (b) indels resulting in frame-shift mutations; AND (b) with ethnicity-specific minor allele frequency (MAF) < 1% in GnoMAD. Variants identified from above analysis were then confirmed by Sanger sequencing. This study was approved by the Partners Institutional Review Board at the MGH, and informed consent was obtained from all participants.

Statistical analyses. All statistical analyses were performed using GraphPad Prism7 software. Cell counts were done on serial sections immunostained for GnRH-1 at E13.5 (n = 3) and E15.0 (n = 4) and visualized under bright field (immunoperoxidase) or epi-fluorescence illumination (20×; Leica DM4000 B LED), according to their anatomical location: (1) nasal region (VNO, axonal tracks surrounding the olfactory pits, forebrain junction); (2) olfactory bulb/fibrocellular mass; and (3) brain (all the cells that accessed the olfactory bulb and were distributed within the forebrain). For each animal, counts were performed on three serial series. The average number of cells from these three series was then multiplied by the total number of series/animal to compute a value for each animal. These were then averaged  $\pm$  SE among animals of the same age and genotype. Means  $\pm$ SEs were calculated on at least three animals per genotype. The statistical difference between genotypes and groups were determined using an unpaired Student's t test. A gene-based burden testing for human GLI3 variants was performed between the KS and the gnomAD using a Fisher's-exact test. All data are represented as the mean  $\pm$  SEM from  $n \ge 3$  mice per genotype/age for each experiment. Values of p < 0.05 were considered to be statistically significant.

#### Results

### Gli3 is expressed by apical Hes-1 + progenitor cells in

Because GnRH-1ns originate in the region of the nasal pit that develops into the vomeronasal organ, we analyzed Gli3 expre-

ssion throughout GnRH-1ns and VSN development. GnRH-1ns arise between E10.5 and E11.5 and are immunodetectable before vomeronasal neuron formation (Fig. 1A,A1,M,P) (Forni et al., 2011a, 2013; Forni and Wray, 2015; Taroc et al., 2017). By E13.5–E15.5, we observed that the developing VNO was populated by Tfap2e (AP-2 $\epsilon$ ) + basal VSNs and Meis2 + apical VSNs (Fig. 1N,Q). However, most GnRH-1ns migrated out of the vomeronasal area by E13.5 (Fig. 1B,B1,C,C1).

We then performed in situ hybridization and immunohistochemistry (Data not shown) against Gli3 at embryonic stages E11.5, E13.5, and E15.5 (Fig. 1D-F). These experiments revealed that Gli3 mRNA and protein mostly localize in the apical regions of the developing vomeronasal epithelia. The proliferative apical cells in the developing olfactory pit (Cuschieri and Bannister, 1975a) localize Hes-1 + stem cell/progenitors that give rise, over time, to apical Ascl-1 neurogenic progenitors (Cau et al., 2000). By performing anti phospho-histone-H3 (PHH3) immunostaining, we only detected mitotic cells in the apical region of the VNO at E11.5. However, at E13.5 and E15.5, we detected mitotic cells along the margins of both the apical and basal regions (Cuschieri and Bannister, 1975b; Cau et al., 2000) (Fig. 1G-I). Immunostaining for Hes-1 and Ascl-1 in the developing VNO (Fig. 2A-D) showed a progressive enrichment of immunodetectable Ascl-1 + cells over time (Fig. 1J-L). We confirmed opposite expression patterns for Hes-1 and Ascl-1 at E15.5, as Hes-1 localized predominantly in the apical portions and Ascl-1 localized in the basal portions of the developing VNO (Fig. 2A-D). We confirmed Gli3 immunoreactivity in Hes-1 + cells and sparse immunoreactivity in the basal regions of the VNO (Fig.  $2G_1H$ ).

Following Hedgehog stimulation Gli3, as a transcriptional activator (Gli3a), induces the expression of Gli1 (which is the amplifying activator downstream Shh) (Lee et al., 1997). In the absence of Shh, Gli3 repressor (Gli3R) blocks Gli1 expression (Lee et al., 1997; Dai et al., 1999; Sasaki et al., 1999; Hu et al., 2006). Previous data have suggested that Gli3 acts as transcriptional repressor in the radial glia of the developing brain cortex (Hasenpusch-

Theil et al., 2018). To understand if, in the developing VNO, Gli3 acts as an activator or as a repressor we exploited a Gli1 knock-in LacZ mouse line (Bai et al., 2002) as a Hedgehog signaling reporter (Fig. 2*J*,*K*). X-Gal enzymatic reaction revealed Gli1 ex-

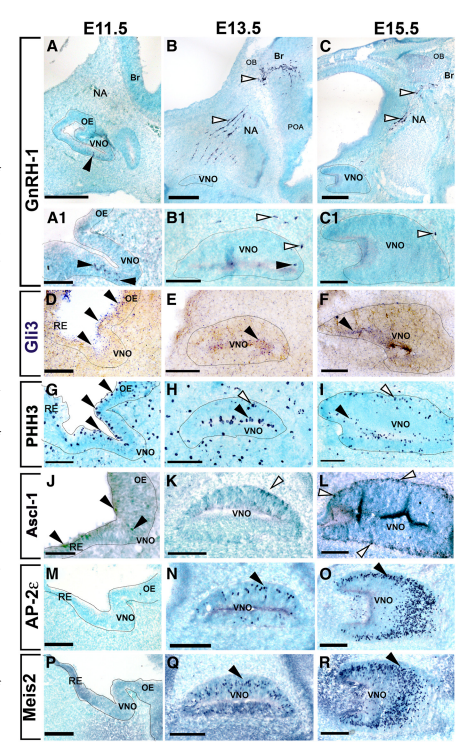
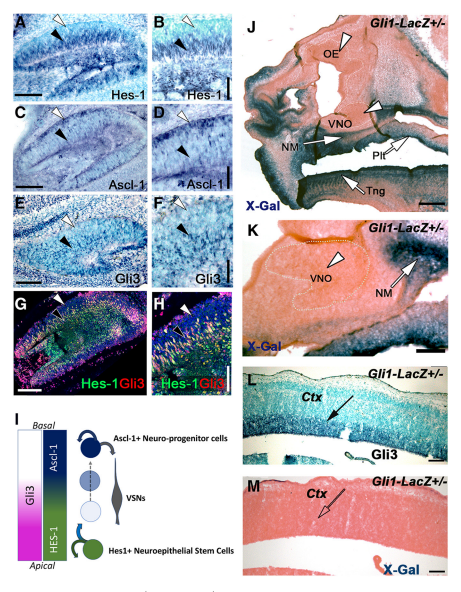


Figure 1. Gli3 is expressed in the developing vomeronasal area. GnRH-1ns can be detected at E11.5 (*A*, *A1*) in the ventral portion of the developing putative VNO and emerge from the VNO (white arrows) to the nasal area (NA) migrating toward the brain (Br) at E13.5 (*B*, *B1*) and E15.5 (*C*, *C1*). *D*–*F*, Gli3 mRNA expression is detected in cells located in the apical portion of the developing olfactory pit and VNO (black arrows) at all analyzed stages. *G*–*J*, PHH3 immunoreactivity highlights dividing cells in the apical (black arrow) and basal (white arrows) regions of the developing VNO.*J*–*L*, Immunostaining against Asd-1 highlights sparse Asd-1 + cells at E11.5 (*J*), neurogenic Asd-1 + cells were detected in the basal regions of the developing VNO at E13.5 and E15.5. Immunostaining against the basal transcription factor AP-2ε (*M*–*O*) and apical transcription factor Meis2 (*P*–*R*) reveal the lack of detectable vomeronasal neurons at E11.5, while a growing number of neurons were detected between E13.5 and E15.5. Scale bars in *A*–*C*,250 μm; *A*1, *D*, *G*, *J*, *M*, *P*, 50 μm; *B*1, *C*1, *E*, *F*, *H*, I, K, L, N, O, Q, R, 100 μm.

pression, which is indicative of active Hedgehog signaling (Fig. 2J, K), in the nasal mesenchyme (NM), cartilage and bones as well as in the oral palate and in the tongue (Sagai et al., 2017). However, even though Gli3 was detected both in the developing



**Figure 2.** Gli3 is expressed in Hes-1 +, but not in Ascl-1 +, progenitors. *A, B,* E15.5 immunostaining against Hes-1 expression in the cells located in the apical domains of the developing VNO (Black arrow). *C, D,* Immunostaining against Ascl-1 reveals expression of the neurogenic factor in cells in the basal territories of the developing VNO (White arrow), only background levels were found in the basal regions (Black arrow). *E, F,* Gli3 immunoreactivity was found in cells in the apical territories of the developing VNO (Black arrow). *G, H,* Double immunofluorescence against Gli3 and Hes-1 reveals strong Gli3 expression in the Hes-1 + cells (Black arrow) and lack of immunodetectable Gli3 in the most basal levels (white arrow). *I,* Proposed model with Gli3 expression in proliferative Hes-1 + cells and in daughter cells that exit the proliferative program and enter the Ascl-1 + controlled proneurogenic program in the basal regions of the developing VNO. *J,* X-Gal reaction on E15.5 sections of Gli1-LacZ Knock-in mouse shows active Shh signaling (Gli1 expression, blue, white arrows) in the palate (plt), nasal mesenchyme (NM), tongue (tng) but not in the olfactory epithelium (OE) norin VNO (white arrowheads). *K,* magnification of the VNO showing lack of Gli1 expression. *L, M,* Serial sections; (*L*) Immunostaining against Gli3 shows strong Gli3 expression in the ventricular zone of the developing brain cortex (arrow), (*M*) X-Gal reaction showing lack of Gli1 expression in the ventricular zone. Scale bars: *A, C, E, G, K, L, M,* 100 μm; *B, D, F, H, J,* 50 μm.

brain cortex (Fig. 2*L*) (Hasenpusch-Theil et al., 2018) and VNO (Fig. 2*E*–*H*), we did not detect Gli1 expression in these areas (Fig. 2*J*–*M*). Similar results were obtained using anti Gli1 immunostaining (data not shown). Lack of active Hedgehog signaling in the developing OE and VNO, suggests that, in these regions Gli3 acts as a transcriptional repressor.

# Gli3 loss-of-function affects VSN neurogenesis but retains VSN specification and differentiation

The expression pattern of Gli3 in the Hes1<sup>+</sup> progenitors in apical region of the VNO resembles its expression in the radial glia of

developing brain cortex (Hasenpusch-Theil et al., 2018). Gli3 immunoreactivity in the apical Hes1+ proliferative region of the VNO prompted us to analyze if Gli3 repressor activity contributes to initiating the differentiation of neurogenic progenitors (Wang et al., 2011; Hasenpusch-Theil et al., 2018). We speculated that Gli3 may play a similar role as in the cortex to stimulate the onset of neurogenic progenitors responsible for the genesis of GnRH-1ns, terminal nerve cells, and VSNs. VSNs differentiate into either apical or basal VSNs (Enomoto et al., 2011; Lin et al., 2018). To identify and quantify the cell bodies of the embryonic VSNs, we performed immunolabeling against the apical VSNs' transcription factors Meis2 and the basal VSNs' transcription factor Tfap2e (AP-2ε) on Gli3 Xt/Xt mutants and controls (Lin et al., 2018) (Fig. 3A, B). At E15, we detected both apical and basal VSNs in the developing VNOs of controls (Fig. 3A). However, we found a dramatic reduction ( $\sim$ 70%) in the number of VSNs in Gli3  $^{\text{Xt/Xt-null}}$  mutants. (Fig. 3*C*). Heterozygous Gli3 WT/Xt displayed an intermediate phenotype between controls and Gli3 Xt/Xt mutants. These data suggest that lacking functional Gli3 compromises VSNs neurogenesis, while permitting the differentiation of the two main types of VSNs (Fig. 3A–C).

Quantification of PHH3 + mitotic cells at E15 highlighted a significant decrease in the total number of mitotic progenitors in Gli3  $^{Xt/Xt}$  mutants (Fig. 3D–F). However, we found a significant reduction in the number of basal progenitors in Gli3 Xt/Xt compared with WT animals and found no significant difference in apical progenitor cells (Fig. 3F). Heterozygous Gli3 WT/Xt embryos displayed a nonsignificant intermediate phenotype compared with WT or Gli3 Xt/Xt. Because we found a significant reduction in proliferation in the basal portions of the VNO, we investigated whether Gli3 loss-of-function compromised the onset of Ascl-1 + basal neurogenic progenitors. Quantifying Ascl-1 + cells confirmed a dramatic (~90%) reduction of Ascl-1 + neurogenic progenitors in the

VNO (Fig. 3*G*–*I*). These data suggest that Gli3 loss-of-function in the VNO impairs the onset of neurogenic progenitors similar to what is reported for the developing brain cortex (Wang et al., 2011; Hasenpusch-Theil et al., 2018).

#### Gli3 loss-of-function disrupts GnRH-1 neuronal migration

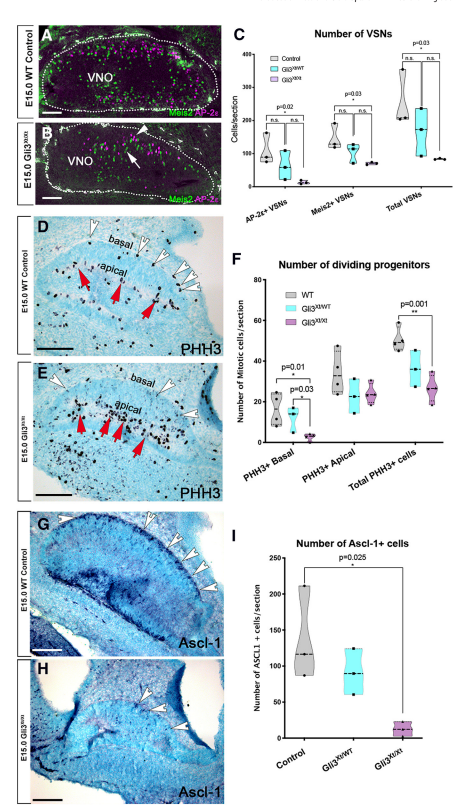
The observed defects in VSN development prompted us to investigate the impact of Gli3 loss-of-function on GnRH-1 neurogenesis and migration. In control conditions, GnRH-1ns migrate along the axons of the terminal nerve that invade the brain ventral to the olfactory bulb, cross the developing ventral telencephalon/

subpallium, then project to the preoptic area (Cariboni et al., 2011; Hanchate et al., 2012; Giacobini and Prevot, 2013; Taroc et al., 2017) (Fig. 4A, C, D, G). However, Gli3  $^{\text{Xt/Xt}}$  mutants show large clusters of cells in the nasal area (Fig. 4B, E), as most GnRH-1ns were unable to reach the brain. Some GnRH-1ns in these mutants migrated toward the developing brain but then formed cell clumps along the migratory track (Fig. 4F). Sparse GnRH-1ns did reach the putative forebrain junction without invading the brain (Fig. 4B, H).

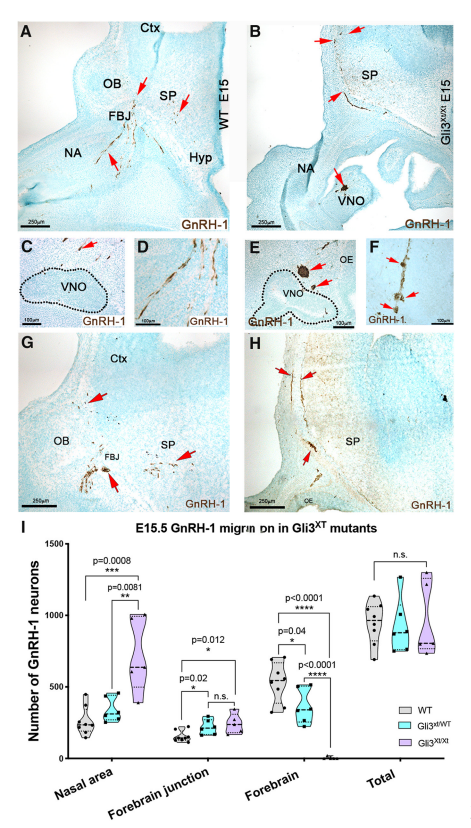
We then quantified the distribution of GnRH-1ns between the nasal area forebrain junction and brain on serial parasagittal sections after immunostaining against GnRH-1 (Fig. 41). These data revealed a lack of GnRH-1-immunoreactive cells in the brain of Gli3 mutants with a massive accumulation of GnRH-1 cells in the nasal area. We observed a small, but significant, reduction in the number of GnRH-1 + cells in the brain of heterozygous controls at E15 with a comparable increase in GnRH-1 cells in the nasal area. Notably, quantifying WT, Gli3 Xt/WT, and Gli3 Xt/Xt mutants at E15 revealed that the total number of GnRH-1ns was comparable across genotypes (Fig. 41). These data indicate a distinct lineage for VSNs and GnRH-1ns. We propose that Gli3 loss-offunction compromises the formation of Ascl-1 + vomeronasal progenitors (Fig. 3), but not GnRH-1 neurogenesis or differentiation.

## Gli3 loss-of-function impairs OECs development in the nasal mucosa

We investigated if the defective migration of GnRH-1ns in Gli3 Xt/Xt results from defective vomeronasal development and/or aberrant terminal nerve projections to the brain. So, we performed a series of immunolabeling experiments to discriminate between cell types. Immunolabeling with anti-peripherin provided a shared marker to highlight olfactory, vomeronasal, and terminal nerve neurons (Fig. 5A-D, M, N) (Taroc et al., 2017). To follow the projections of VSNs and the terminal nerve (see diagram in Fig. 5A-D), we performed immunolabeling against the guidance receptors, Neuropilin-2 (Nrp2), Roundabout-2 (Robo2), and Neuropilin-1 (Nrp1). Robo2 highlights projections of the basal VSNs (Fig. 5E,F,I,J), Nrp2 highlights apical VSNs (Fig. 5G-I) (Prince et al., 2009), while Nrp1 (Fig. 5K2,L2) (Taroc et al., 2017) labels projections of putative terminal nerve neurons (Fig. 5K1-L2). In control animals, we found TN fibers (Nrp1 +) invaded the



**Figure 3.** Gli3 loss-of-function impairs the formation of Ascl-1 <sup>+</sup> neuronal progenitor cells and VSN neurogenesis but not VSN terminal differentiation. **A**, **B**, Immunostaining against the transcription factors Meis2 and AP-2ε highlights maturing apical and basal vomeronasal sensory neurons in controls (**A**) and Gli3 <sup>Xt/Xt</sup> mutants (**B**). **C**, Quantifications of the average number of AP-2ε differentiated VSNs and Meis2 differentiated apical VSNs reveal a significant reduction in the number of differentiated VSNs in Gli3 <sup>Xt/Xt</sup> mutants. **D**, **E**, PHH3 immunostaining and quantification (**F**) of VNO from WT and Gli3 <sup>Xt/Xt</sup> reveals a reduction in the total number of dividing cells with a significant reduction of basal progenitors (white arrowheads), but not apical progenitors (red arrows). **G**, **H**, Immunostaining against Ascl-1 in controls and Gli3 <sup>Xt/Xt</sup> mutants reveals (**I**) a significant reduction in the number of Ascl-1 <sup>+</sup> cells in the basal domains of the developing VNO. Scale bars , 100 μm.



**Figure 4.** Gli3 <sup>Xt/Xt</sup> mutants display severely impaired GnRH-1 neuronal migration. Representative images of immunostaining against GnRH-1 on WT (*A***-G**) and Gli3 <sup>Xt/Xt</sup> (*B***-H**) at E15.5. *A***-G**, in WT GnRH-1ns migrate as a continuum from the vomeronasal organ (VNO) to the brain. GnRH-1ns from the nasal area (NA), cross the forebrain junction (FBJ), invade the brain ventral to the OB, migrate through the subpallium (SP) to eventually reach the hypothalamic area (Hyp). **B**, **C**, GnRH-1ns migrating in the nasal area and out of the VNO. **D**, Detail showing GnRH-1ns invading the subpallium (SP) ventral to the OB in the region corresponding to the putative accumbens (Ac), some GnRH-1ns migrate to the cortex (Ctx) and around the olfactory bulbs (OBs). E-H) In Gli3 <sup>Xt/Xt</sup>, a large portion of GnRH-1ns neurons form clusters of cells proximal to the VNO (*E*) while other migrate around the brain, occasional GnRH-1ns were found accessing the brain dorsal portion of the cortex (Ctx) (*H*). *F*, GnRH-1 immunostaining (brown) show less

brain, together with GnRH-1ns (Fig. 5A, C, E, G, I). However, Gli3  $^{Xt/Xt}$  mutants showed stalled olfactory and vomeronasal axons (positive for Robo2 and/or Nrp2, Fig. 5B, D, F, H, J) (Balmer and LaMantia, 2004) in front of the forebrain, which formed a fibro-cellular mass (FCM). However, we did observe that sparse putative terminal nerve axons and GnRH-1ns (Fig. 5B,L1,L2) did reach the putative forebrain junction and project around the dorsal portions of the brain. However, the majority of GnRH-1ns (Fig. 4E) and terminal nerve fibers (peripherin +; Nrp1 +, Fig. 5O1-O4; negative for Nrp2 and Robo2, Fig. 5N-N4) formed large tangles in the nasal area.

GnRH-1 neuronal development strictly depends on correct formation of OECs (Barraud et al., 2013; Pingault et al., 2013). Gli3 can act as a Sox10 modifier (Matera et al., 2008). Our data show that Gli3 loss-of-function leads to defective vomeronasal and terminal nerve trajectories, which phenocopies that described after OECs loss in Sox10-null mouse mutants (Barraud et al., 2013; Pingault et al., 2013). So, we tested the role of OECs in axonal misrouting and GnRH-1 migratory defects in Gli3 Xt/Xt mutants. So, we performed double staining against peripherin and Sox10 (Forni et al., 2011b; Pingault et al., 2013; Rich et al., 2018). In control animals, Sox10 + OECs were localized proximal to the basal lamina of the vomeronasal epithelium and correlated with the vomeronasal bundles emerging from the vomeronasal epithelium (Fig. 5P,R). In the OE and olfactory fibers, we observed a similar distribution with strong Sox10 immunoreactivity close to the basal regions of the OE and along the olfactory bundles projecting to the OBs (Fig. 5*P*). In Gli3 <sup>Xt/Xt</sup> mutants, we found a near complete lack of Sox10 immunoreactivity around the vomeronasal epithelium and no Sox10 + cells associated with vomeronasal axons or terminal nerve (Fig. 5Q,S). We also found no Sox10 immunoreactivity near the basal lamina of the olfactory epithelium (Fig. 5Q). However, in Gli3 Xt/Xt, we found Sox10 + en-

organized GnRH-1ns forming clumps while migrating toward the brain (cf.  $\emph{D}$ ).  $\emph{H}$ , Detail showing GnRH-1ns were unable to invade the brain migrating along the meninges.  $\emph{I}$ , Quantification of the distribution of GnRH-1ns in WT, Gli3  $^{\text{Xt/WT}}$  and Gli3  $^{\text{Xt/Xt}}$ . In Gli3  $^{\text{Xt/Xt}}$  the majority of the GnRH-1ns remain in the nasal area. Both Gli3  $^{\text{Xt/WT}}$  and Gli3  $^{\text{Xt/Xt}}$  have a significantly smaller number of GnRH-1ns in the brain compared with controls. Values are  $\pm$  SE. Dots indicate numbers of embryos/genotype, unpaired  $\emph{t}$  test, significant values  $\emph{p} < 0.05$ .

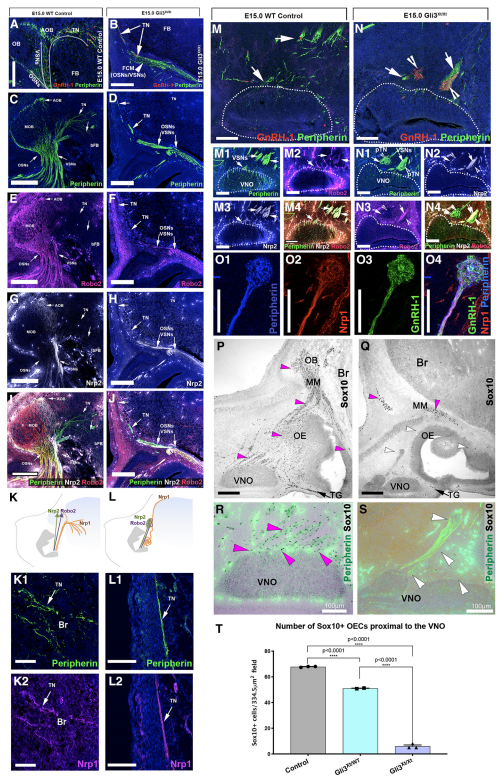
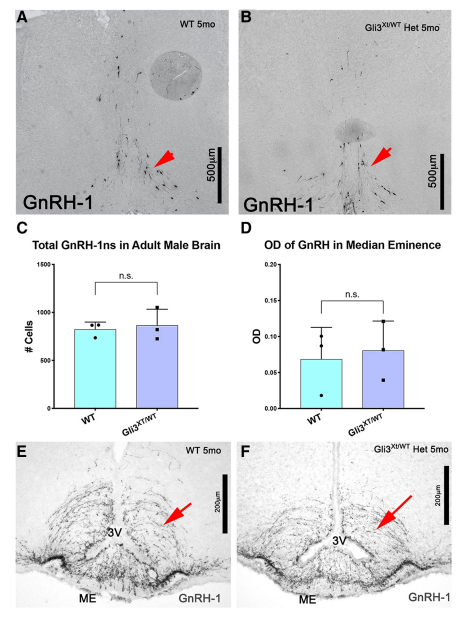


Figure 5. Misrouted vomeronasal and terminal nerve axons in Gli3 Xt/Xt are associated with lack of ensheathing cells in the nasal mucosa. **A, C,** Immunostainings against peripherin and GnRH-1. **B, D,** peripherin and GnRH-1 immunofluorescence, in Gli3 Xt/Xt mutants the OSNs and VSNs form a fibrocellular cellular mass (FCM). The GnRH-1ns (**D**) and the TN do not remain in the FCM but migrate around the FB. **C–I,** WT E15.5, immunostaining against peripherin, Robo2, Nrp2; the TN fibers are negative for Robo2 expression and negative for Nrp2 (see merge in **I**). **D–J,** Gli3 Xt/Xt E15.5 immunostaining against peripherin, Robo2, and Nrp2 and merge. **K, L,** Schematic of the phenotypes and fiber immunoreactivity in controls and Gli3 Xt/Xt mutants. **K1, K2,** Immunofluorescence showing Neuropilin-1 (Nrp1) immunoreactivity in TN fibers invading the brain in WT embryos. **L1, L2,** peripherin and Neruopilin1 + fibers of the putative TN project around the brain. **M1–M4,** In control animals the GnRH-1ns migrate along peripherin + bundles (**M1),** also immunoreactive for Robo2 (**M2),** Nrp2 (**M3),** Merge in **M4.** In Gli3 Xt/Xt, GnRH-1ns form large clumps of cells suggesting disorganized peripherin (**O1)** and Nrp1 (**O2)** and are associated with clusters of GnRH-1ns (**O3).** Merge in **O4.** P–**5,** GLI3 Xt/Xt lack of OECs associated with the vomeronasal nerve. peripherin F and Sox10-IHC are combined. **P,** In controls, Sox10-immunoreactive OECs (magenta arrowheads) were found around the VNO, proximal to the basal lamina of the OE, along the peripherin + vomeronasal and olfactory bundles and as part of the migratory mass (MM) in front of the brain. Sox10 + Schwann cells were found associated along nasal trigeminal projections (Figure legend continues.)



**Figure 6.** Gli3 XL/WT show no difference in GnRH-1ns numbers in the brain after birth. **A**, **B**, Immunostaining against GnRH-1 on postnatal WT and Gli3 XL/WT heterozygous animals shows comparable immunoreactivity in GnRH-1 cell bodies in the preoptic area (arrows). **C**, Quantification for GnRH-1-immunoreactive cell bodies in the preoptic area indicates no statistical differences between controls and Gli3 XL/WT heterozygous mutants. **D**, Quantification of area occupied by GnRH-1-immunoreactive fibers in the median eminence of WT controls and Gli3 XL/WT heterozygous mutants. **E**, **F**, Representative images showing GnRH-1 immunoreactivity of GnRH-1 axons in the median eminence (ME) of WT and Gli3 XL/WT heterozygous mutants.

sheathing cells proximal to the brain as part of the FCM (Fig. 5Q), consistent with previous reports (Balmer and LaMantia, 2004).

To determine whether the OECs in the nasal mesenchyme were missing or if they were just failing in expressing the maturation markers Sox10 (Rich et al., 2018), we performed immuno-

(Figure legend continued.) (white arrowheads). Notably Sox10  $^+$  cells were found in the migratory mass (MM). As in controls Sox10  $^+$  Schwann cells were found associated along nasal trigeminal projections (TG). **R**, **5**, Magnifications of the VNO. **R**, In control animals Sox10  $^+$  OECs were found proximal to the basal lamina of the VNO, along the vomeronasal bundles. **5**, In Gli3  $^{\rm Xt/Xt}$  Sox10  $^+$  OECs were rarely found in proximity of the basal lamina of the VNO and along the vomeronasal/terminal nerve bundles (white arrowheads). **7**, Quantification of Sox10  $^+$  OECs around the VNO, \*\*\*\*\*p < 0.0001. Scale bars, 100  $\mu$ m.

staining against BLBP which labels mature and immature OECs even in the absence of Sox10 (Pingault et al., 2013). This immunostaining indicated almost complete absence of BLBP + OECs in Gli3 Xt/Xt mice as observed with Sox10 (data not shown). These data suggest defective OECs specification in the nasal mucosa of Gli3 Xt/Xt.

As we detected dramatic defects in OECs formation in the developing olfactory system, we propose that the aberrant neuronal patterning and GnRH-1 migratory impairment observed in Gli3 Xt/Xt mutants is secondary to the lack of OECs in the nasal mucosa (Barraud et al., 2013; Pingault et al., 2013; Rich et al., 2018).

### Delayed, but not defective, migration in Gli3 Xt/WT heterozygous mutants

Partial defects in GnRH-1 neuronal migration in Gli3 Pdn/Pdn Gli3 hypomorphs (Naruse et al., 1994) suggest that defects in Gli3 levels can impair GnRH-1 neuronal migration. As we collected evidence indicating that Gli3 controls neurogenesis of VSNs, we further analyzed the development of the GnRH-1 system in Gli3 Xt/WT heterozygous embryos. Gli3 Xt/WT are viable and fertile, and have mild craniofacial defects and an extra toe (Veistinen et al., 2012). However, they do not display dramatic abnormalities in the olfactory system. We analyzed GnRH-1 neuronal migration in Gli3 Xt/WT heterozygous animals after development. Analysis of adult WT controls and Gli3 Xt/WT mutants (Fig. 6A-C) revealed no significant differences in the number and distribution of GnRH-1ns in the brain after development. Immunostaining against GnRH-1 and quantifying the average area of GnRH-1-immunoreactive axons also revealed comparable GnRH-1 innervation of the median eminence (Fig. 6D–F). These data suggest that Gli3 haploinsufficient mice show delayed, but not disrupted, GnRH-1 neuronal migration.

### Lack of TN and GnRH-1 invasion of the brain and altered Sema3A expression in Gli3 Xt/Xt mutants

In Gli3  $^{Xt/X\bar{t}}$ , the few TN and GnRH-1 neurons that reach the brain fail to invade it. Because brain development can be highly dysmorphic after Gli3 loss-of-function (Aoto et al., 2002; Balmer and LaMantia, 2004; Fotaki et al., 2006; Blaess et al., 2008; Veistinen et al., 2012; Hasenpusch-Theil et al., 2018), we analyzed how such aberrant development can indirectly compromise the ability of TN and GnRH-1 neurons that do reach the brain to invade it (Figs. 4*H*; 5*K*,*L*). In rodents, gammGABA-ergic interneurons form in the ventral telencephalon and subsequently migrate toward the cortex (Anderson et al., 1997; Lavdas et al., 1999; Wichterle et al., 1999, 2001; Markram et al., 2004). The transcription factor AP-2 $\varepsilon$  can identify the developing olfactory bulbs and

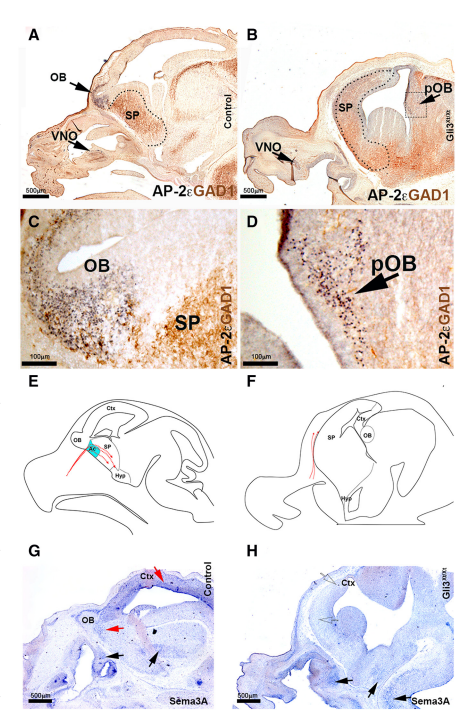
vomeronasal neurons (Feng et al., 2009; Enomoto et al., 2011; Lin et al., 2018) (Fig. 7A–D). We used antibodies against glutamate decarboxylase 1 (brain, 67 kDa) (GAD67/GAD-1) to label the ventral forebrain/subpallium. Consistent with previous reports (Yu et al., 2009; Besse et al., 2011), Gli3 Xt/Xt mutants exhibit a lack of rostral olfactory bulb protrusions but form ectopic olfactory bulb-like structures, positive for AP-2ɛ in the dorsal/ lateral portions of the developing telencephalon (Fig. 7B,D). At E15.5, GnRH-1ns invade the brain by crossing the putative accumbens (ventral to the OB) and then steering ventrally toward the hypothalamic area.

During embryonic development, Sema3A is expressed ventral to the OB in the putative accumbens/ "septostriatal transition area" (Fig. 7E,G) (Taroc et al., 2017). Lack of Sema3A is not compatible with normal invasion of the brain by the TN and GnRH-1ns (Cariboni et al., 2011). As the development of the brain appears highly dysmorphic in Gli3 mutants (See diagram in 7F), we analyzed the pattern of Sema3A expression compared with controls. Our analysis revealed that Gli3 Xt/Xt mutants showed comparable Sema3A expression to controls in nasal areas and the hindbrain. However, we found a near complete absence of Sema3A expression in the expanded subpallium and the dorsal portion of the forebrain (Fig. 7F,H). These data show aberrant Sema3A expression and patterning in the brain of Gli3 Xt/Xt mutants, which, alone, is a molecular condition sufficient to prevent migratory GnRH-1 neurons to invade the brain (Cariboni et al., 2011; Taroc et al., 2017). Analysis of Sema3A expression in heterozygous Gli3 Xt/WT animals did not highlight obvious changes in Sema3A expression pattern compared with controls (data not shown). These data, together with the delayed but correct migration of GnRH-1ns, suggest that Gli3 haploinsufficiency is not sufficient to disrupt the patterning of the guidance cue Sema3A.

### Ascl-1 loss-of-function compromises VNO and GnRH-1 neurogenesis but retains OECs formation in the nasal

The negative effects of Gli3 loss-offunction on VSNs' but not on GnRH-1ns' neurogenesis prompted us to test 1) if

Ascl-1 controls GnRH-1 neurogenesis (Kramer and Wray, 2000; McNay et al., 2006) and 2) if the lack of OECs in the nasal mucosa and impaired GnRH-1 migration, observed in Gli3 mutants, directly result from the compromised vomeronasal neurogenesis (Cau et al., 1997, 2002). For this purpose, we analyzed GnRH-1,



**Figure 7.** Gli3 XLIX mutants display an expansion of expression of forebrain ventral markers, form ectopic olfactory bulbs and lack of Sema3A expression in the forebrain. **A**, AP-2ε/GAD1 double immunostaining. Immunostaining against AP-2ε highlights mitral cells in the OB (magnified in **C**). GAD1 immunostaining highlights GABAergic neurons which are mostly located in the subpallium. **B**, In Gli3 XLIX, AP-2ε highlight extopic mitral cells in the posterior portion of the forebrain (boxed, see magnification in (**D**) showing the ectopic AP-2ε + OB in the posterior portion of the brain. GAD1 expression shows expansion of ventral forebrain maker. **E**, **F**, Illustration of GnRH-1ns migration in WT and Gli3-null animals. Area of Sema3A expression represented in light blue present in WT accumbens absent in the Gli3 KOs. **G**, **H**, In situ hybridization against Sema3A shows detectable Sema3A expression in the nasal area, basal brain (black arrows) and hindbrain of control and Gli3 mutants. No Sema3A expression was detected in Gli3 Xt mutants in the forebrain and cortex (compare red arrows in **I** with dotted arrows in **H**). Ac, Accumbens; Ctx, cortex; Hyp, hypothalamus; OB, olfactory bulb; pOB, putative olfactory bulb; SP, subpallium.

VSNs, and OECs development on Ascl-1 KOs and controls at E13.5.

In Ascl-1 KOs we found a ( $\sim$ 60%) reduction in VSNs compared with WT controls which is similar that observed in Gli3  $^{\text{Xt/Xt}}$  (Fig. 8 I, J, O). As for the reduced neurogenesis, in Ascl-1  $^{\text{null}}$  animals we

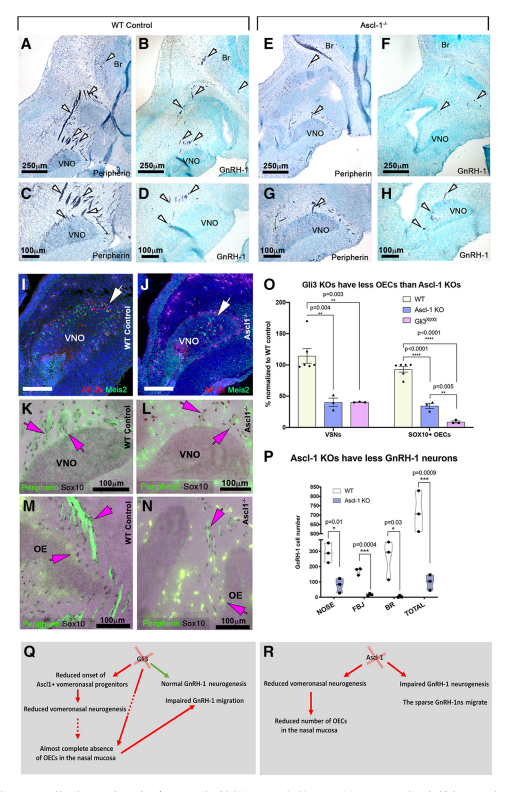


Figure 8. Ascl-1 KO have a comparable reduction in the number of vomeronasal and GnRH-1 neurons. *A*–*C*, Immunostaining against peripherin highlights neuronal projections (white arrows) emerging from the VNO of WT animals projecting toward the brain. *B*, *D*, In controls, immunostaining against GnRH-1 reveals migratory GnRH-1ns distributed between the VNO and the brain. *E*, *G*, In Ascl-1-null mutants, only sparse peripherin <sup>+</sup> projections emerge from the OE and VNO (white arrows). *F*–*H*, In Ascl-1 KOs, GnRH-1ns emerge from the VNO migrating in the nasal area (white arrows). *I*, *J*, AP-2ε and Meis2 immunostaining highlights the reduced number of differentiated VSNs in Ascl-1 KOs (white arrow). *K*–*N*, Reduced olfactory and vomeronasal neurons in Ascl-1 KOs does not prevent ensheathing cell formation in the nasal mucosa. IF anti peripherin and Sox10 IHC are combined. *K*, *M*, In control animals, many Sox10-immunoreactive OECs (magenta arrows) are found around the VNO, along the peripherin <sup>+</sup> vomeronasal bundles, (*M*) proximal to the developing OE and along the olfactory bundles. *L*, In Ascl-1 <sup>-/-</sup> mutants, Sox10-immunoreactive OECs were found associated with the sparse vomeronasal projections (magenta arrows) (*N*) proximal to the developing OE and along the olfactory bundles (magenta arrows). *O*, Quantification of GnRH-1ns in controls and Ascl-1 KOs shows dramatic reduction in the number of GnRH-1ns in Ascl-1 KOs. Forebrain junction (FBJ). Holm-Sidak's multiple-comparisons test. *P*, Quantification of VSNs, OECs, show similar defects in vomeronasal neurogenesis between Ascl-1 <sup>-/-</sup> and Gli3 <sup>Xt/Xt</sup> compared with WT controls however, more severe loss of OECs was found in Gli3 <sup>Xt/Xt</sup> compared with Ascl-1 <sup>null</sup> lines.

observed a comparable (-60%; SE  $\pm 1\%$ ; p=0.003) reduction in the number of OECS (Fig. 8K-N,O). However, though reduced in number, SOX10  $^+$  OECs were found to be distributed with similar frequency to the controls along the few vomeronasal projections (WT 0.080 cell/ $\mu$ m n=3; Ascl-1 KO = 0.082 cell/ $\mu$ m, n=3; p=0.89).

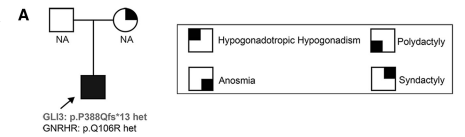
In Ascl-1 KOs, we found sparse GnRH-1-immunoreactive neurons emerging from the developing VNO and migrating toward the brain along peripherin fibers (Fig. 8*A*–*H*). However, cell quantifications revealed a dramatic (~86%) decrease in the total number of GnRH-1-immunoreactive neurons in Ascl-1 KOs (Fig. 8*P*). Analysis of Ascl-1 for revealed no differences compared with WT controls, suggesting no dose-dependent effect of Ascl-1 on vomeronasal and GnRH-1 neurogenesis (data not shown).

As we observed a severe reduction in the total number of GnRH-1ns in Ascl-1 null, but not in Gli3 Xt/Xt, mutants, our data suggest that the mechanisms controlling Ascl-1 expression in GnRH-1 neuronal progenitors are independent of Gli3 control. Further, these observations suggest that the near complete absence of Sox10 + OECs in the nasal mucosa of Gli3 Xt/Xt mutants is not a only the result of defective vomeronasal neurogenesis (Fig. 8O) (Miller et al., 2018; Rich et al., 2018), but likely secondary to neural crest defects (Matera et al., 2008). In support of this conjecture, we observed that Gli3 Xt/WT/Ascl-1 +/- double mutants were indistinguishable from Gli3 Xt/WT single mutants (data not shown).

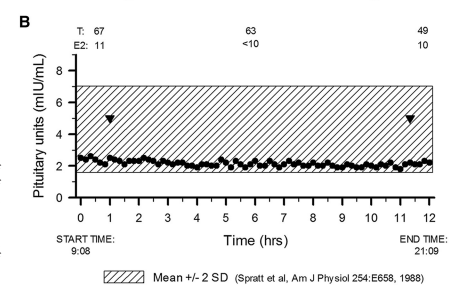
### A GLI3 loss-of-function mutation in a KS and polydactyl patient

Because KS features reproductive failure and various defects in GnRH-1 neuronal migration, we first examined the genetic constraint for *GLI3* gene in Genome Aggregation Database (gnomAD) v.2.1 (Karczewski et al., 2019). This population genetics data showed that *GLI3* human genetic variants were highly constrained for loss-of-function variants with a *pLI* of

1.0 and an observed/expected (*oe*) metric of 0.09 (upper bound *oe* confidence interval of 0.20). The suggested threshold for mendelian case analysis is an upper bound *oe* confidence interval of <0.35. These parameters suggest an extreme intolerance to loss-of-function variants reflecting the strength of selection acting on heterozygotes. So, we queried WES data from 632 KS probands for *GLI3* loss-of-function variants. We identified two qualifying *GLI3* loss-of-function variants from WES data. Sanger sequencing confirmed one novel heterozygous *GLI3* loss-of-function (frameshifting indel) variant (c.1161del; p.P388Qfs\*13; minor allele frequency in gnomAD: 0) in a KS male subject. This frame-



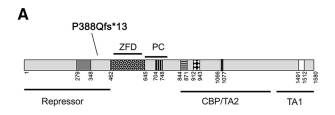
Testicular Volume (mL) = 8 (left)/10 (right)
FSH: 3.89 mIU/mL
Mean LH: 2.18 mIU/mL (%CV: 4.22)
SHBG: 8.3 nmol/L

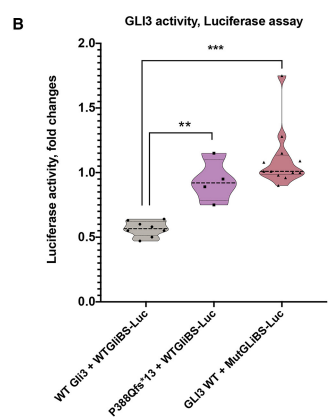


Additional Endocrine Test
TSH 1.38 (0.4-5.0) uU/mL
Prolactin 7.5 (0-15) ng/mL

**Figure 9.** *A*, Pedigree of Kallmann syndrome proband (shown with arrow) with reproductive and nonreproductive phenotypes. *B*, KS proband underwent neuroendocrine profiling (Q 10 min  $\times$  12 h) to chart GnRH-induced LH secretion showing low amplitude, low-frequency pulses (inverted triangles indicate LH pulses). Shaded region represents the normal reference range. NA, Not available; T, total testosterone; LH, luteinizing hormone; FSH, follicle stimulating hormone; SHBG, sex-hormone binding globulin' TSH, thyroid-stimulating hormone.

shift (fs) mutation leads to the formation of a premature TGA stop codon in position 1199–1201. Because parental DNA samples were not available, the precise mode of inheritance of the variant could not be evaluated. Additional KS/nIHH gene analysis showed that this KS proband also harbored a heterozygous GNRHR missense mutation (p.Q106R) (Fig. 9). Because gnomAD variants are not Sanger-confirmed, we performed a genebased burden test between the KS cohort and gnomAD for all qualifying GLI3 loss-of-function variants that showed enrichment in the KS cohort (p=0.002). We did not observe this enrichment when correcting for Sanger-confirmed variants only





**Figure 10.** Functional validation of GLI3 loss-of-function. **A,** Diagram illustrates *GLI3* protein, the mutation P388Qfs\*13 in the repressor domain of the protein. ZFD, Zinc finger domain; PC, protein cleavage; TA1, TA2, CBP-binding domain, transactivation domain 1 and 2 . **B,** Luciferase activity assay, the repressor activity of Gli3WT on the reporter (WTGLiBS-Luc) is lost in the truncated GLI3. No repression was found for WT Gli3 on reporter carrying nonfunctional Gli binding site MutGLiBS-Luc. \*\*p = 0.002, \*\*\*p < 0.0001, Holm—Sidak's multiple-comparisons to the comparison of the protein of

in the KS cohort but retaining all unconfirmed gnom AD variants ( p=0.06 ).

The patient harboring the GLI3 loss-of-function variant is a 38-year-old male, who presented to medical attention at age 18 years with anosmia and failure to enter puberty. He was undervirilized with a bilateral low testicular volume (Fig. 9). His biochemical investigations showed frank hypogonadotropic hypogonadism (serum testosterone: 67 ng/dL; LH: 2.18 mIU/ml and FSH: 3.89 mIU/ml) with neuroendocrine studies revealing low-amplitude low-frequency LH pulse profile consistent with hypogonadotropism (Fig. 9). The remaining pituitary hormone profile was normal and UPSIT showed complete anosmia. These results confirmed a clinical diagnosis of Kallmann syndrome. Consistent with the well established role of GLI3 in limb development, he reported in his history an extra toe on his right foot, for which he underwent surgery as a child, and bilateral webbed toes (second and third toes). Although the precise inheritance mode of the variant was not determined due to lack of parental samples, his mother also exhibited syndactyly affecting her feet, suggesting a possible autosomal-dominant mode of inheritance with variable penetrance for the reproductive phenotype (Fig. 9).

To test the biological effects of the identified GLI3 variant, we performed a luciferase assay (Fig. 10) to compare the activity of Wild-type (WT) GLI3 and mutated GLI3 using a reporter plasmid containing wild-type GLI binding site linked to firefly luciferase (WTGliBS-Luc). A reporter carrying a mutated/nonfunctional (MutGLiBS-Luc) Gli3 binding site linked to firefly luciferase (Sasaki et al., 1997; Saitsu et al., 2005) served as negative control. In contrast with a prior report (Krauss et al., 2008), luciferase assay on HeLa cells (ATCC CCL-2) revealed that Gli3 acts a transcriptional repressor. This analysis demonstrated the deleterious loss-of-function nature of our novel *GLI3* frameshift mutation (p.P388Qfs\*13), which affected the repressor ability of Gli3, consistent with a GLI3 loss-of-function as a putative mechanism for GLI3-related KS.

#### Discussion

Here, we found that Gli3 loss-of-function induces: (1) loss of Ascl-1 neuronal progenitors in the VNO and dramatically reduces the number of vomeronasal sensory neurons (VSNs), (2) normal neurogenesis for GnRH-1ns, 3() nearly complete absence of GnRH-1 neuronal migration, associated to aberrant formation of OECs in the nasal mucosa, and (4) loss of forebrain expression of Semaphorin 3A (Sema3A), a key axonal guidance molecule that controls GnRH-1 invasion of the brain (Cariboni et al., 2011; Hanchate et al., 2012; Young et al., 2012). To elucidate the underlying cellular substrates of the observed phenotypes, we analyzed Ascl-1<sup>-/-</sup> mutants. Even though we found reduced vomeronasal neurogenesis similar to Gli3 mutants, Ascl-1 -/mutants showed a severely reduced number of GnRH-1ns' and Sox10 + OECs (OECs) that formed along axonal bundles similar to controls (Fig. 8). Based on the observed phenotypes between Gli3 and Ascl-1 mutants, we conclude that Ascl-1 is crucial for GnRH-1 neurogenesis independent of Gli3 control. Ascl-1 mice showed that reduced neurogenesis in the VNO alone does not prevent the maturation of Sox10 + OECs in the nasal mucosa and the migration of GnRH-1ns in Gli3 mutants. These data suggest a direct role for Gli3 in controlling OECs development. These murine observations highlight a critical role for Gli3 distinctly in OEC development and GnRH neuronal migration, two features that underlie the pathogenesis of KS. In keeping with this notion, we also identified and functionally validated a novel GLI3 loss-of-function mutation in a KS patient who also exhibited polydactyly, a well recognized phenotypic feature of GLI3-related human diseases.

Analyzing Gli3 expression in the developing olfactory pit, we observed that, Gli3 is expressed by proliferative progenitors in the apical portions of the developing VNO positive for the transcription factor Hes-1 (Figs. 1,2). Hes1 inhibits neurogenesis by repressing the transcription factor Ascl-1 and maintaining neuroepithelial cells in undifferentiated/proliferative state (Ohtsuka et al., 1999; Cau et al., 2000; Ohtsuka et al., 2001). Neurogenic Ascl-1 progenitors were found to be distributed in the basal portions of the developing VNO (Fig. 2). In Gli3-null mutants we observed a dramatic reduction of VSNs secondary to a near complete loss (~91%) of Ascl-1 + neurogenic progenitors in the VNO. Surprisingly, the number of GnRH-1ns did not appear different from WT controls. These results indicate that loss of Gli3-mediated gene repression negatively regulates the onset of vomeronasal neurogenic progenitors, but not GnRH-1 neurogenesis.

The cyclin-dependent kinase gene Cdk6 was recently identified as a direct target of Gli3-mediated repression (Hasenpusch-Theil et al., 2018). Gli3 loss-of-function shortens the cell cycle and reduces neurogenesis (Pilaz et al., 2009; Hasenpusch-Theil et al., 2018). Because we detected Gli3 in Hes-1 proliferative cells, we hypothesize that Gli3 controls the onset of neurogenesis through a similar mechanism as in the cortex (Hasenpusch-Theil et al., 2018). Active Shh signaling induces Gli1 expression. By using a Gli1-Lac-Z mice, we observed lack of active Shh signaling in the VNO. This finding suggests, together with the absence of GnRH-1 phenotypes in Gli1/Gli2 KOs (Metz and Wray, 2010), that Gli3 acts as a transcriptional repressor in the developing VNO.

Gli3 heterozygous mice showed a delay in GnRH-1 migration without differences in cell number or distribution after birth and were fertile. These data indicate that heterozygous Gli3 loss-of-function mutations, alone, do not produce a GnRH-deficient state in mice, which is consistent with results from Gli3 hypomorphic mutant mice (Naruse et al., 1994).

OECs are a glial population that wraps bundles of olfactory and vomeronasal bundles (Rich et al., 2018). OECs promote normal GnRH-1 neuronal migration from the nasal area to the brain (Barraud et al., 2013; Geller et al., 2013; Pingault et al., 2013). OECs arise from the neural crest precursors (Barraud et al., 2010; Forni et al., 2011b; Miller et al., 2016, 2018; Rich et al., 2018). By analyzing GnRH-1ns migration in Gli3 Xt/Xt mutants, we found that most GnRH-1 cells were unable to migrate far from the VNO and formed large clumps of cells on tangled fibers of the putative TN. This phenotype matches that of Sox10-null mutants (Barraud et al., 2013; Pingault et al., 2013). Gli3 can act as a Sox10 modifier, which is consistent with the suggestion that Gli3 lossof-function can compromise OECs development (Matera et al., 2008). Here, we found virtually no Sox10 or BLBP + cells in the olfactory/vomeronasal bundles of Gli3 Xt/Xt mutants nor proximal to the basal lamina and lamina propria of the olfactory epithelia, suggestive of Gli3 loss impairing OECs development in the nasal mucosa. In Gli3 Xt/Xt mice, we found Sox10 + Schwann cells around the trigeminal nerve (Fig. 5) and around the fibers of the sensory Grueneberg ganglion (data not shown) (Liu et al., 2009; Matsuo et al., 2012; Moine et al., 2018). We also found Sox10 + OECs surrounding the FCM proximal to the OBs. These intriguing data suggest developmental differences between OECs of the nasal mucosa and olfactory bulb OECs (Richter et al., 2005; Ito et al., 2006; Mayeur et al., 2013).

In Gli3 Xt/Xt mutants, we found sporadic TN fibers and GnRH-1 cells could reach the brain but not invade it. These findings revealed a similar cellular behavior to that described after Sema3A loss-of-function. Analyses of Gli3 Xt/Xt mutants confirmed very dysmorphic forebrain structures with expanded regions positive for the early subpallial marker, GAD1, and ectopic OBs (Balmer and LaMantia, 2004; Fotaki et al., 2006; Blaess et al., 2008; Veistinen et al., 2012). *In situ* hybridization against Sema3A showed a near complete absence of Sema3A expression in the forebrain, a condition that alone is not permissive for normal GnRH-1 migration (Cariboni et al., 2011). We propose that altered Sema3A expression in Gli3 mutants directly reflects either the lack of functional Gli3 transcriptional control or overall aberrant brain development.

Analyzing Ascl-1 null mutants revealed a role for Ascl-1 in controlling GnRH-1 neuronal development, since we found an approximate 86% reduction in the number of GnRH-1ns in these mice.

We also found no vomeronasal defects together with normal GnRH-1 cell numbers and migration in Ascl-1  $^{+/-}$  heterozygous

mice. This finding indicates that Ascl-1 is not required in a dose-dependent manner for vomeronasal and GnRH-1 neurogenesis as described for other neurons (McNay et al., 2006). The lack of overlapping phenotype between Gli3 Xt/Xt (normal GnRH-1 but reduced VSN cell numbers) and Ascl-1 null mutants (reduced GnRH-1 and VSNs cell numbers) indicates that GnRH-1 neurogenic onset is not regulated in a Gli3-dependent fashion.

We observed that OECs defects in Ascl-1 null mutants appeared less severe than in Gli3 Xt/Xt and that OECs in the mucosa of Ascl-1 null mutants were positive for Sox10 (Fig. 80). Moreover, Gli3 Xt/WT/Ascl-1 +/- double mutants did not have exacerbated phenotypes compared with Gli3 Xt/WT suggesting that the effects seen in both Ascl-1 null and Gli3 Xt/Xt are independent of each other and not additive (data not shown). We conclude that the near complete absence of OECs in the nasal mucosa of Gli3 Xt/Xt mutants is not secondary to the decreased number of vomeronasal fibers (Miller et al., 2018).

Analysis of WES from KS/nIHH patients identified a novel frameshift (fs) GLI3 variant (p.P388Qfs\*13) that causes the formation of a premature TGA termination codon in position 1199-1201. Using site-directed mutagenesis and luciferase assays, we confirmed a loss of repressor ability of this mutation, which decreased the repressor ability of GLI3. This patient also displayed polydactyly and syndactyly, two hallmark clinical features of humans with GLI3-Greigs cephalopolysyndactyly syndrome (GCPS), which corresponds to the loss of GLI3 transcriptional repression during limb development (Johnston et al., 2010; Al-Qattan et al., 2017). This mutation affects the 5' part of the GLI3 gene upstream of the zinc finger domain which is well correlated with GCPS phenotype (Johnston et al., 2010; Al-Qattan et al., 2017). Reviewing previously reported GLI3 mutation subjects with GCPS or oral-facial-digital syndrome indicates that a small minority of these patients also display micropenis (Kroisel et al., 2001; Johnston et al., 2010), cryptorchidism (Kroisel et al., 2001; Johnston et al., 2010), or anosmia (Johnston et al., 2010). These phenotypes are consistent with an underlying KS. KS/nIHH may be under-recognized in these reports, as reproductive phenotypes cannot be assessed in prepubertal subjects harboring *GLI3* mutations. Our findings implicate heterozygous GLI3 loss-of-function mutations as a contributing genetic factor for KS/nIHH, which is further supported by the associated GLI3 missense variants in two subjects with HH (Quaynor et al., 2016).

We acknowledge that heterozygous *GLI3* mutations alone may be insufficient to cause KS/nIHH and may require additional genetic or nongenetic modifiers. Indeed, KS has a well established oligogenic basis (Sykiotis et al., 2010). Here, we found that the proband with *GLI3* loss-of-function mutation also harbored an additional rare *GNRHR* p.Q106R missense mutation. Although the *GNRHR* p.Q106R variant can be deleterious in cellular assays (Leaños-Miranda et al., 2005), *GNRHR* mutations typically cause autosomal recessive hypogonadotropic hypogonadism (Gianetti et al., 2012). So, the cooccurrence of *GLI3* and *GNRHR* variants in this proband suggests a potential oligogenic (*GLI3/GNRHR*) contribution to KS.

In summary, we revealed a crucial role for Gli3 in controlling vomeronasal sensory neurons neurogenesis, formation of OECs in the nasal mucosa, and GnRH-1 neuron migration. Our findings suggest a role for Gli3 in triggering the transition from a proliferative to neurogenic program in the developing VNO. Because we found a dramatic reduction in the number of Sox10 expressing OECs and aberrant Sema3A in the brain, we propose that Gli3 acts as a genetic modifier and contributes to the oligogenic nature of KS/nIHH. Future studies will delineate the gene

regulatory network controlled by the Gli3 transcriptional network in the developing nasal olfactory system and identify KS/nIHH candidate genes to confirm the role of this transcription factor as a modifier in the oligogenic nature of KS/nIHH.

#### References

- Al-Qattan MM, Shamseldin HE, Salih MA, Alkuraya FS (2017) GLI3-related polydactyly: a review. Clin Genet 92:457–466.
- Anderson SA, Eisenstat DD, Shi L, Rubenstein JL (1997) Interneuron migration from basal forebrain to neocortex: dependence on dlx genes. Science 278:474–476.
- Aoto K, Nishimura T, Eto K, Motoyama J (2002) Mouse GLI3 regulates Fgf8 expression and apoptosis in the developing neural tube, face, and limb bud. Dev Biol 251:320–332.
- Bai CB, Auerbach W, Lee JS, Stephen D, Joyner AL (2002) Gli2, but not Gli1, is required for initial shh signaling and ectopic activation of the shh pathway. Development 129:4753–4761.
- Balasubramanian R, Crowley WF Jr (2011) Isolated GnRH deficiency: a disease model serving as a unique prism into the systems biology of the GnRH neuronal network. Mol Cell Endocrinol 346:4–12.
- Balmer CW, LaMantia AS (2004) Loss of Gli3 and shh function disrupts olfactory axon trajectories. J Comp Neurol 472:292–307.
- Barraud P, Seferiadis AA, Tyson LD, Zwart MF, Szabo-Rogers HL, Ruhrberg C, Liu KJ, Baker CV (2010) Neural crest origin of olfactory ensheathing glia. Proc Natl Acad Sci U S A 107:21040–21045.
- Barraud P, St John JA, Stolt CC, Wegner M, Baker CV (2013) Olfactory ensheathing glia are required for embryonic olfactory axon targeting and the migration of gonadotropin-releasing hormone neurons. Biol Open 2:750–759.
- Besse L, Neti M, Anselme I, Gerhardt C, Rüther U, Laclef C, Schneider-Maunoury S (2011) Primary cilia control telencephalic patterning and morphogenesis via Gli3 proteolytic processing. Development 138:2079–2088.
- Blaess S, Stephen D, Joyner AL (2008) Gli3 coordinates three-dimensional patterning and growth of the tectum and cerebellum by integrating shh and Fgf8 signaling. Development 135:2093–2103.
- Cariboni A, Davidson K, Rakic S, Maggi R, Parnavelas JG, Ruhrberg C (2011) Defective gonadotropin-releasing hormone neuron migration in mice lacking SEMA3A signalling through NRP1 and NRP2: implications for the aetiology of hypogonadotropic hypogonadism. Hum Mol Genet 20:336–344.
- Casoni F, Malone SA, Belle M, Luzzati F, Collier F, Allet C, Hrabovszky E, Rasika S, Prevot V, Chédotal A, Giacobini P (2016) Development of the neurons controlling fertility in humans: new insights from 3D imaging and transparent fetal brains. Development 143:3969–3981.
- Cau E, Gradwohl G, Fode C, Guillemot F (1997) Mash1 activates a cascade of bHLH regulators in olfactory neuron progenitors. Development 124: 1611–1621.
- Cau E, Gradwohl G, Casarosa S, Kageyama R, Guillemot F (2000) Hes genes regulate sequential stages of neurogenesis in the olfactory epithelium. Development 127:2323–2332.
- Cau E, Casarosa S, Guillemot F (2002) Mash1 and Ngn1 control distinct steps of determination and differentiation in the olfactory sensory neuron lineage. Development 129:1871–1880.
- Cuschieri A, Bannister LH (1975a) The development of the olfactory mucosa in the mouse: electron microscopy. J Anat 119:471–498.
- Cuschieri A, Bannister LH (1975b) The development of the olfactory mucosa in the mouse: light microscopy. J Anat 119:277–286.
- Dai P, Akimaru H, Tanaka Y, Maekawa T, Nakafuku M, Ishii S (1999) Sonic hedgehog-induced activation of the Gli1 promoter is mediated by GLI3. J Biol Chem 274:8143–8152.
- Doty RL (2007) Office procedures for quantitative assessment of olfactory function. Am J Rhinol 21:460–473.
- Enomoto T, Ohmoto M, Iwata T, Uno A, Saitou M, Yamaguchi T, Kominami R, Matsumoto I, Hirota J (2011) Bcl11b/Ctip2 controls the differentiation of vomeronasal sensory neurons in mice. J Neurosci 31:10159–10173.
- Feng W, Simoes-de-Souza F, Finger TE, Restrepo D, Williams T (2009) Disorganized olfactory bulb lamination in mice deficient for transcription factor AP-2epsilon. Mol Cell Neurosci 42:161–171.
- Forni PE, Wray S (2015) GnRH, anosmia and hypogonadotropic hypogonadism-where are we? Front Neuroendocrinol 36:165–177.
- Forni PE, Fornaro M, Guénette S, Wray S (2011a) A role for FE65 in controlling GnRH-1 neurogenesis. J Neurosci 31:480–491.
- Forni PE, Taylor-Burds C, Melvin VS, Williams T, Wray S (2011b) Neural

- crest and ectodermal cells intermix in the nasal placode to give rise to GnRH-1 neurons, sensory neurons, and olfactory ensheathing cells. J Neurosci 31:6915–6927.
- Forni PE, Bharti K, Flannery EM, Shimogori T, Wray S (2013) The indirect role of fibroblast growth factor-8 in defining neurogenic niches of the olfactory/GnRH systems. J Neurosci 33:19620–19634.
- Fotaki V, Yu T, Zaki PA, Mason JO, Price DJ (2006) Abnormal positioning of diencephalic cell types in neocortical tissue in the dorsal telencephalon of mice lacking functional Gli3. J Neurosci 26:9282–9292.
- Geller S, Kolasa E, Tillet Y, Duittoz A, Vaudin P (2013) Olfactory ensheathing cells form the microenvironment of migrating GnRH-1 neurons during mouse development. Glia 61:550–566.
- Giacobini P, Prevot V (2013) Semaphorins in the development, homeostasis and disease of hormone systems. Semin Cell Dev Biol 24:190-198.
- Gianetti E, Hall JE, Au MG, Kaiser UB, Quinton R, Stewart JA, Metzger DL, Pitteloud N, Mericq V, Merino PM, Levitsky LL, Izatt L, Lang-Muritano M, Fujimoto VY, Dluhy RG, Chase ML, Crowley WF Jr, Plummer L, Seminara SB (2012) When genetic load does not correlate with phenotypic spectrum: lessons from the GnRH receptor (GNRHR). J Clin Endocrinol Metab 97:E1798–E1807.
- Guo MH, Plummer L, Chan YM, Hirschhorn JN, Lippincott MF (2018) Burden testing of rare variants identified through exome sequencing via publicly available control data. Am J Hum Genet 103:522–534.
- Hanchate NK et al. (2012) SEMA3A, a gene involved in axonal pathfinding, is mutated in patients with Kallmann syndrome. PLoS Genet 8:e1002896.
- Hasenpusch-Theil K, West S, Kelman A, Kozic Z, Horrocks S, McMahon AP, Price DJ, Mason JO, Theil T (2018) Gli3 controls the onset of cortical neurogenesis by regulating the radial glial cell cycle through Cdk6 expression. Development 145:dev163147.
- Hu MC, Mo R, Bhella S, Wilson CW, Chuang PT, Hui CC, Rosenblum ND (2006) GLI3-dependent transcriptional repression of Gli1, Gli2 and kidney patterning genes disrupts renal morphogenesis. Development 133: 569–578.
- Hu Y, Butts T, Poopalasundaram S, Graham A, Bouloux PM (2019) Extracellular matrix protein anosmin-1 modulates olfactory ensheathing cell maturation in chick olfactory bulb development. Eur J Neurosci 50: 3472–3486
- Ito D, Ibanez C, Ogawa H, Franklin RJ, Jeffery ND (2006) Comparison of cell populations derived from canine olfactory bulb and olfactory mucosal cultures. Am J Vet Res 67:1050–1056.
- Johnston JJ et al. (2010) Molecular analysis expands the spectrum of phenotypes associated with GLI3 mutations. Hum Mutat 31:1142–1154.
- Kagoshima M, Ito T (2001) Diverse gene expression and function of semaphorins in developing lung: positive and negative regulatory roles of semaphorins in lung branching morphogenesis. Genes Cells 6:559–571.
- Karczewski KJ et al. (2019) Variation across 141,456 human exomes and genomes reveals the spectrum of loss-of-function intolerance across human protein-coding genes. bioRxiv 531210. Available from: http://doi.org/10.1101/531210.
- Keino H, Masaki S, Kawarada Y, Naruse I (1994) Apoptotic degeneration in the arhinencephalic brain of the mouse mutant Pdn/Pdn. Brain Res Dev Brain Res 78:161–168.
- Kim EJ, Ables JL, Dickel LK, Eisch AJ, Johnson JE (2011) Ascl1 (Mash1) defines cells with long-term neurogenic potential in subgranular and subventricular zones in adult mouse brain. PLoS One 6:e18472.
- Kinzler KW, Ruppert JM, Bigner SH, Vogelstein B (1988) The GLI gene is a member of the kruppel family of zinc finger proteins. Nature 332:371–374.
- Kramer PR, Wray S (2000) Midline nasal tissue influences nestin expression in nasal-placode-derived luteinizing hormone-releasing hormone neurons during development. Dev Biol 227:343–357.
- Krauss S, Foerster J, Schneider R, Schweiger S (2008) Protein phosphatase 2A and rapamycin regulate the nuclear localization and activity of the transcription factor GLI3. Cancer Res 68:4658–4665.
- Kroisel PM, Petek E, Wagner K (2001) Phenotype of five patients with greig syndrome and microdeletion of 7p13. Am J Med Genet 102:243–249.
- Lavdas AA, Grigoriou M, Pachnis V, Parnavelas JG (1999) The medial ganglionic eminence gives rise to a population of early neurons in the developing cerebral cortex. J Neurosci 19:7881–7888.
- Leaños-Miranda A, Ulloa-Aguirre A, Janovick JA, Conn PM (2005) In vitro coexpression and pharmacological rescue of mutant gonadotropinreleasing hormone receptors causing hypogonadotropic hypogonadism

- in humans expressing compound heterozygous alleles. J Clin Endocrinol Metab 90:3001–3008.
- Lee J, Platt KA, Censullo P, Ruiz i Altaba A (1997) Gli1 is a target of sonic hedgehog that induces ventral neural tube development. Development 124:2537–2552.
- Lewkowitz-Shpuntoff HM, Hughes VA, Plummer L, Au MG, Doty RL, Seminara SB, Chan YM, Pitteloud N, Crowley WF Jr, Balasubramanian R (2012) Olfactory phenotypic spectrum in idiopathic hypogonadotropic hypogonadism: pathophysiological and genetic implications. J Clin Endocrinol Metab 97:E136–E144.
- Lin JM, Taroc EZM, Frias JA, Prasad A, Catizone AN, Sammons MA, Forni PE (2018) The transcription factor Tfap2e/AP-2epsilon plays a pivotal role in maintaining the identity of basal vomeronasal sensory neurons. Dev Biol 441:67–82.
- Liu CY, Fraser SE, Koos DS (2009) Grueneberg ganglion olfactory subsystem employs a cGMP signaling pathway. J Comp Neurol 516:36–48.
- Markram H, Toledo-Rodriguez M, Wang Y, Gupta A, Silberberg G, Wu C (2004) Interneurons of the neocortical inhibitory system. Nat Rev Neurosci 5:793–807.
- Matera I, Watkins-Chow DE, Loftus SK, Hou L, Incao A, Silver DL, Rivas C, Elliott EC, Baxter LL, Pavan WJ (2008) A sensitized mutagenesis screen identifies Gli3 as a modifier of Sox10 neurocristopathy. Hum Mol Genet 17:2118–2131.
- Matsuo T, Rossier DA, Kan C, Rodriguez I (2012) The wiring of Grueneberg ganglion axons is dependent on neuropilin 1. Development 139:2783–2791.
- Mayeur A, Duclos C, Honoré A, Gauberti M, Drouot L, do Rego JC, Bon-Mardion N, Jean L, Vérin E, Emery E, Lemarchant S, Vivien D, Boyer O, Marie JP, Guérout N (2013) Potential of olfactory ensheathing cells from different sources for spinal cord repair. PLoS One 8:e62860.
- McNay DE, Pelling M, Claxton S, Guillemot F, Ang SL (2006) Mash1 is required for generic and subtype differentiation of hypothalamic neuroendocrine cells. Mol Endocrinol 20:1623–1632.
- Metz H, Wray S (2010) Use of mutant mouse lines to investigate origin of gonadotropin-releasing hormone-1 neurons: lineage independent of the adenohypophysis. Endocrinology 151:766–773.
- Miller AM, Treloar HB, Greer CA (2010) Composition of the migratory mass during development of the olfactory nerve. J Comp Neurol 518:4825–4841.
- Miller SR, Perera SN, Benito C, Stott SR, Baker CV (2016) Evidence for a Notch1-mediated transition during olfactory ensheathing cell development. J Anat 229:369–383.
- Miller SR, Benito C, Mirsky R, Jessen KR, Baker CVH (2018) Neural crest Notch/Rbpj signaling regulates olfactory gliogenesis and neuronal migration. Genesis 56:e23215.
- Moine F, Brechbühl J, Nenniger Tosato M, Beaumann M, Broillet MC (2018) Alarm pheromone and kairomone detection via bitter taste receptors in the mouse Grueneberg ganglion. BMC Biol 16:12.
- Naruse I, Fukui Y, Keino H, Taniguchi M (1994) The arrest of luteinizing hormone-releasing hormone neuronal migration in the genetic arhinencephalic mouse embryo (Pdn/Pdn). Brain Res Dev Brain Res 81:178–184.
- Niewiadomski P, Kong JH, Ahrends R, Ma Y, Humke EW, Khan S, Teruel MN, Novitch BG, Rohatgi R (2014) Gli protein activity is controlled by multisite phosphorylation in vertebrate Hedgehog signaling. Cell Rep 6:168–181.
- Ohtsuka T, Ishibashi M, Gradwohl G, Nakanishi S, Guillemot F, Kageyama R (1999) Hes1 and Hes5 as notch effectors in mammalian neuronal differentiation. EMBO J 18:2196–2207.
- Ohtsuka T, Sakamoto M, Guillemot F, Kageyama R (2001) Roles of the basic helix-loop-helix genes Hes1 and Hes5 in expansion of neural stem cells of the developing brain. J Biol Chem 276:30467–30474.
- Pilaz LJ, Patti D, Marcy G, Ollier E, Pfister S, Douglas RJ, Betizeau M, Gautier E, Cortay V, Doerflinger N, Kennedy H, Dehay C (2009) Forced G1-phase reduction alters mode of division, neuron number, and laminar phenotype in the cerebral cortex. Proc Natl Acad Sci U S A 106: 21924–21929.
- Pingault V, Bodereau V, Baral V, Marcos S, Watanabe Y, Chaoui A, Fouveaut C, Leroy C, Vérier-Mine O, Francannet C, Dupin-Deguine D, Archambeaud F, Kurtz FJ, Young J, Bertherat J, Marlin S, Goossens M, Hardelin JP, Dodé C, Bondurand N (2013) Loss-of-function mutations in SOX10 cause Kallmann syndrome with deafness. Am J Hum Genet 92:707–724.
- Pitteloud N, Hayes FJ, Dwyer A, Boepple PA, Lee H, Crowley WF Jr (2002) Predictors of outcome of long-term GnRH therapy in men with idio-

- pathic hypogonadotropic hypogonadism. J Clin Endocrinol Metab 87:4128-4136.
- Prince JE, Cho JH, Dumontier E, Andrews W, Cutforth T, Tessier-Lavigne M, Parnavelas J, Cloutier JF (2009) Robo-2 controls the segregation of a portion of basal vomeronasal sensory neuron axons to the posterior region of the accessory olfactory bulb. J Neurosci 29:14211–14222.
- Quaynor SD, Bosley ME, Duckworth CG, Porter KR, Kim SH, Kim HG, Chorich LP, Sullivan ME, Choi JH, Cameron RS, Layman LC (2016) Targeted next generation sequencing approach identifies eighteen new candidate genes in normosmic hypogonadotropic hypogonadism and Kallmann syndrome. Mol Cell Endocrinol 437:86–96.
- Rich CA, Perera SN, Andratschke J, Stolt CC, Buehler DP, Southard-Smith EM, Wegner M, Britsch S, Baker CVH (2018) Olfactory ensheathing cells abutting the embryonic olfactory bulb express frzb, whose deletion disrupts olfactory axon targeting. Glia 66:2617–2631.
- Richter MW, Fletcher PA, Liu J, Tetzlaff W, Roskams AJ (2005) Lamina propria and olfactory bulb ensheathing cells exhibit differential integration and migration and promote differential axon sprouting in the lesioned spinal cord. J Neurosci 25:10700–10711.
- Sagai T, Amano T, Maeno A, Kiyonari H, Seo H, Cho SW, Shiroishi T (2017) SHH signaling directed by two oral epithelium-specific enhancers controls tooth and oral development. Sci Rep 7:13004.
- Saitsu H, Komada M, Suzuki M, Nakayama R, Motoyama J, Shiota K, Ishibashi M (2005) Expression of the mouse Fgf15 gene is directly initiated by sonic hedgehog signaling in the diencephalon and midbrain. Dev Dyn 232:282–292.
- Sasaki H, Hui C, Nakafuku M, Kondoh H (1997) A binding site for gli proteins is essential for HNF-3beta floor plate enhancer activity in transgenics and can respond to shh in vitro. Development 124:1313–1322.
- Sasaki H, Nishizaki Y, Hui C, Nakafuku M, Kondoh H (1999) Regulation of Gli2 and Gli3 activities by an amino-terminal repression domain: implication of Gli2 and Gli3 as primary mediators of shh signaling. Development 126:3915–3924.
- Schimmang T, Lemaistre M, Vortkamp A, Rüther U (1992) Expression of the zinc finger gene Gli3 is affected in the morphogenetic mouse mutant extra-toes (Xt). Development 116:799–804.
- Schwanzel-Fukuda M, Pfaff DW (1989) Origin of luteinizing hormonereleasing hormone neurons. Nature 338:161–164.
- Sykiotis GP, Plummer L, Hughes VA, Au M, Durrani S, Nayak-Young S, Dwyer AA, Quinton R, Hall JE, Gusella JF, Seminara SB, Crowley WF Jr, Pitteloud N (2010) Oligogenic basis of isolated gonadotropin-releasing hormone deficiency. Proc Natl Acad Sci U S A 107:15140–15144.
- Taroc EZM, Prasad A, Lin JM, Forni PE (2017) The terminal nerve plays a prominent role in GnRH-1 neuronal migration independent from proper olfactory and vomeronasal connections to the olfactory bulbs. Biol Open 6:1552–1568.
- Veistinen L, Takatalo M, Tanimoto Y, Kesper DA, Vortkamp A, Rice DP (2012) Loss-of-function of Gli3 in mice causes abnormal frontal bone morphology and premature synostosis of the interfrontal suture. Front Physiol 3:121.
- Wang H, Ge G, Uchida Y, Luu B, Ahn S (2011) Gli3 is required for maintenance and fate specification of cortical progenitors. J Neurosci 31:6440–6448.
- Wichterle H, Garcia-Verdugo JM, Herrera DG, Alvarez-Buylla A (1999) Young neurons from medial ganglionic eminence disperse in adult and embryonic brain. Nat Neurosci 2:461–466.
- Wichterle H, Turnbull DH, Nery S, Fishell G, Alvarez-Buylla A (2001) In utero fate mapping reveals distinct migratory pathways and fates of neurons born in the mammalian basal forebrain. Development 128: 3759–3771.
- Wray S, Grant P, Gainer H (1989) Evidence that cells expressing luteinizing hormone-releasing hormone mRNA in the mouse are derived from progenitor cells in the olfactory placode. Proc Natl Acad Sci U S A 86: 8132–8136.
- Young J, Metay C, Bouligand J, Tou B, Francou B, Maione L, Tosca L, Sarfati J, Brioude F, Esteva B, Briand-Suleau A, Brisset S, Goossens M, Tachdjian G, Guiochon-Mantel A (2012) SEMA3A deletion in a family with Kallmann syndrome validates the role of semaphorin 3A in human puberty and olfactory system development. Hum Reprod 27:1460–1465.
- Yu T, Fotaki V, Mason JO, Price DJ (2009) Analysis of early ventral telencephalic defects in mice lacking functional Gli3 protein. J Comp Neurol 512:613–627.