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Pituitary stem cells produce paracrine WNT signals to control the expansion of their descendant progenitor cells

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Pituitary stem cells produce paracrine WNT signals to control the expansion of their descendant progenitor cells

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ABSTRACT

In response to physiological demand, the pituitary gland generates new hormone-secreting cells from committed progenitor cells throughout life. It remains unclear to what extent pituitary stem cells (PSCs), which uniquely express SOX2, contribute to pituitary growth and renewal. Moreover, neither the signals that drive proliferation nor their sources have been elucidated. We have used genetic approaches in the mouse, showing that the WNT pathway is essential for proliferation of all lineages in the gland. We reveal that SOX2+ stem cells are a key source of WNT ligands. By blocking secretion of WNTs from SOX2+ PSCs in vivo, we demonstrate that proliferation of neighbouring committed progenitor cells declines, demonstrating that progenitor multiplication depends on the paracrine WNT secretion from SOX2+ PSCs. Our results indicate that stem cells can hold additional roles in tissue expansion and homeostasis, acting as paracrine signalling centres to coordinate the proliferation of neighbouring cells.

KEY WORDS

SOX2, paracrine signal, WNT, pituitary gland, stem cell, feedforward regulation
INTRODUCTION

How stem cells interact with their surrounding tissue has been a topic of investigation since the concept of the stem cell niche was first proposed (Schofield, 1978). Secreted from supporting cells, factors such as WNTs, FGFs, SHH, EGF and cytokines, regulate the activity of stem cells (Nabhan et al., 2018; Palma et al., 2005; Tan and Barker, 2014). Furthermore, communication is known to take place in a bi-directional manner (Doupe et al., 2018; Tata and Rajagopal, 2016).

The anterior pituitary (AP) is a major primary endocrine organ that controls key physiological functions including growth, metabolism, reproduction and the stress responses and exhibits tremendous capability to remodel its constituent hormone populations throughout life, in response to physiological demand. It contains a population of Sox2 expressing stem cells that self-renew and give rise to lineage-committed progenitors and functional endocrine cells (Andoniadou et al., 2013; Rizzoti et al., 2013). During embryonic development, SOX2+ undifferentiated precursor cells of Rathke’s pouch, the pituitary anlage (Arnold et al., 2011; Castinetti et al., 2011; Fauquier et al., 2008; Pevny and Rao, 2003), generate all committed endocrine progenitor lineages, defined by the absence of SOX2 and expression of either POU1F1 (PIT1), TBX19 (TPIT) or NR5A1 (SF1) (Bilodeau et al., 2009; Davis et al., 2011). These committed progenitors are proliferative and give rise to the hormone-secreting cells. Demand for hormone secretion rises after birth, resulting in dramatic organ growth and expansion of all populations by the second postnatal week (Carbajo-Perez and Watanabe, 1990; Taniguchi et al., 2002). SOX2+ pituitary stem cells (PSCs) are most active during this period, but the bulk of proliferation and organ expansion during postnatal stages derives from SOX2+ committed progenitors. The activity of SOX2+ PSCs gradually decreases and during adulthood is minimally
activated even following physiological challenge (Andoniadou et al., 2013; Gaston-Massuet et al., 2011; Gremeaux et al., 2012; Zhu et al., 2015). By adulthood, progenitors carry out most of the homeostatic functions, yet SOX2+ PSCs persist throughout life in both mice and humans (Gonzalez-Meljem et al., 2017; Xekouki et al., 2018). The signals driving proliferation of committed progenitor cells are not known, and neither is it known if SOX2+ PSCs can influence this process beyond their minor contribution of new cells.

The self-renewal and proliferation of numerous stem cell populations relies upon WNT signals (Basham et al., 2019; Lim et al., 2013; Takase and Nusse, 2016; Wang et al., 2015; Yan et al., 2017). WNTs are necessary for the initial expansion of Rathke’s pouch as well as for PIT1 lineage specification (Osmundsen et al., 2017; Potok et al., 2008). In the postnatal pituitary, the expression of WNT pathway components is upregulated during periods of expansion and remodelling. Gene expression comparisons between neonatal and adult pituitaries or in GH-cell ablation experiments (Gremeaux et al., 2012; Willems et al., 2016), show that the WNT pathway is upregulated during growth and regeneration.

Our previous work revealed that during disease, the paradigm of supporting cells signalling to the stem cells may be reversed; mutant stem cells expressing a degradation-resistant β-catenin in the pituitary, promote cell non-autonomous development of tumours through their paracrine actions (Andoniadou et al., 2013; Gonzalez-Meljem et al., 2017). Similarly, degradation-resistant β-catenin expression in hair follicle stem cells led to cell non-autonomous WNT activation in neighbouring cells promoting new growth (Deschene et al., 2014). In the context of normal homeostasis, stem cells have been shown to influence daughter cell fate in the mammalian airway epithelium and the *Drosophila* gut via ‘forward regulation’
models, where the fate of a daughter cell is directed by a stem cell via juxtacrine Notch signalling (Ohlstein and Spradling, 2007; Pardo-Saganta et al., 2015). It remains unknown if paracrine stem cell action can also promote local proliferation in normal tissues. Here, we used genetic approaches to determine if paracrine stem cell action takes place in the anterior pituitary and to discern the function of WNTs in pituitary growth. Our results demonstrate that postnatal pituitary expansion, largely driven by committed progenitor cells, depends on WNT activation. Importantly, we show that SOX2+ PSCs are the key regulators of this process, acting through secretion of WNT ligands acting in a paracrine manner on neighbouring progenitors. Identification of this forward-regulatory model elucidates a previously unidentified function for stem cells during tissue expansion.

RESULTS

WNT-responsive cells in the pituitary include progenitors driving major postnatal expansion. To clarify which cells respond to WNT signals in the postnatal anterior pituitary, we first characterised the anterior pituitary cell types activating the WNT pathway at P14, a peak time for organ expansion and a time point when a subpopulation of SOX2+ stem cells are proliferative. The Axin2-CreERT2 mouse line (van Amerongen et al., 2012) has been shown to efficiently label cells with activated WNT signalling in the liver, lung, breast, skin, testes and endometrium among other tissues (Lim et al., 2013; Moiseenko et al., 2017; Syed et al., 2020; van Amerongen et al., 2012; Wang et al., 2015). Axin2 positive cells were labelled by GFP following tamoxifen induction in
Axin2\textsuperscript{CreERT2/+}, ROSA26\textsuperscript{mTmG/+} mice and pituitaries were analysed 2 days post-induction. We carried out double immunofluorescence staining using antibodies against uncommitted (SOX2), lineage committed (PIT1, TPIT, SF1), and hormone-expressing endocrine cells (GH, PRL, TSH, ACTH or FSH/LH) together with antibodies against GFP labelling the WNT-activated cells. We detected WNT-responsive cells among all the different cell types of the anterior pituitary including SOX2\textsuperscript{+} PSCs, the three committed populations and all hormone-secreting cells (Figure 1A, Figure 1 – figure supplement 1A).

To confirm if the three committed lineages as well as uncommitted SOX2\textsuperscript{+} PSCs all expand in response to WNT, we further lineage-traced Axin2-expressing cells for 14 days after tamoxifen administration at P14. Double labelling revealed an increase in all four populations between 2 and 14 days (Figure 1A, B). This increase reached significance for the PIT1 (13.7\% at 2 days to 30.3\% at 14 days, \(P=0.000004\)) and TPIT (3.78\% to 11.03\%, \(P=0.008\)) populations, but not SF1 (0.5\% to 4\%, n.s.). As this time course ends at P28 at the commencement of puberty, we repeated the analysis for SF1 cells to P42, which spans puberty and the expansion of gonadotrophs (Figure 1 – figure supplement 1B). This reveals a significant expansion in WNT-responsive SF1\textsuperscript{+} cells as a proportion of the total SF1\textsuperscript{+} population (\(P=0.0048, n=3\)). Lineage tracing of the PIT1-derivates (GH\textsuperscript{+} somatotrophs, PRL\textsuperscript{+} lactotrophs, TSH\textsuperscript{+} thyrotrophs) reveals that there is a preferential expansion of somatotrophs and thyrotophs (Figure 1 – figure supplement 1C). Only a minority of SOX2\textsuperscript{+} PSCs were WNT-responsive at 2 days (0.57\%) and this population expanded to 2\% at 14 days (n.s.), suggesting that these are self-renewing. GFP\textsuperscript{+} cells were traced for a period of 8 weeks post-induction, which revealed that WNT-responsive descendants continued to expand at the same rate as the rest of the pituitary (\(n=4\)-8 mice per time point at P16,
P21, P28, P42, P70) (Figure 1C, D). The time period between 2 and 7 days saw the greatest increase in GFP+ cells, during which, the labelled population nearly tripled in size (Figure 1D). The persistence of labelled cells was evident in longer-term traces using the ROSA26lacZ/+ reporter (Axin2CreERT2+/+;ROSA26lacZ/+) up to a year following induction at P14 (Figure 1E, n=4). Clonal analysis using the Confetti reporter, demonstrated that individual Axin2-expressing cells (Axin2CreERT2+/+;ROSA26Confetti+) gave a greater contribution after four weeks compared to lineage-tracing from Sox2-expressing cells (Sox2CreERT2+/+;ROSA26Confetti/+), in support of predominant expansion from WNT-responsive lineage-committed progenitors (Figure 1 – figure supplement 1D).

To establish if signalling mediated by β-catenin is necessary for organ expansion we carried out deletion of Ctnnb1 in the Axin2+ population from P14 during normal growth (Axin2CreERT2+/+;Ctnnb1lox(ex2-6)/lox(ex2-6) hereby Axin2CreERT2+/+;Ctnnb1LOF/LOF). Due to morbidity, likely due to Axin2 expression in other organs, we were limited to analysis up to 5 days post-induction. Deletion of Ctnnb1 resulted in a significant reduction in the number of dividing cells, marked by pH-H3 (40% reduction, Figure 1 – figure supplement 2A, P=0.0313, n=3), confirming that activation of the WNT pathway is necessary for expansion of the pituitary populations. This deletion did not result in significant differences in overall numbers among the three lineages, as determined by the numbers of PIT1+, SF1+ or ACTH+ cells among the targeted population (Figure 1 – figure supplement 2B, n=4 controls, 2 mutants). The number of SOX2+ stem cells and cells undergoing cell death also remained unaffected during the 5 day period (Figure 1 – figure supplement 2C and D). Taken together, these results confirm that postnatal AP expansion depends on
WNT-responsive progenitors across all lineages, in addition to SOX2+ PSCs (Figure 1F).

WNT/β-catenin signalling is required for long-term anterior pituitary expansion from SOX2+ pituitary stem cells.

We further explored the role of WNT pathway activation in postnatal SOX2+ stem cells. To permanently mark WNT-responsive cells and their descendants whilst simultaneously marking SOX2+ PSCs, we combined the tamoxifen-inducible Axin2\textsuperscript{CreERT2/+};ROSA26\textsuperscript{tdTomato/+} with the Sox2\textsuperscript{Egfp/+} strain, where cells expressing SOX2 are labelled by EGFP (Axin2\textsuperscript{CreERT2/+};Sox2\textsuperscript{Egfp/+};ROSA26\textsuperscript{tdTomato/+}). Following tamoxifen administration from P21, tdTomato- and EGFP-labelled cells were analysed by flow sorting after 72h, by which point all induced cells robustly express tdTomato (Figure 2A, Figure 2 – figure supplement 1). Double-labelled cells comprised 23.4% of the SOX2+ population (n=5 individual pituitaries) (Figure 2A, arrowheads), with the majority of tdTomato+ cells found outside of the SOX2+ compartment. It was previously shown that only around 2.5-5% of SOX2+ PSCs have clonogenic potential through in vitro assays (Andoniadou et al., 2012; Andoniadou et al., 2013; Perez Millan et al., 2016). To determine if WNT-responsive SOX2+ cells are stem cells capable of forming colonies, we isolated double positive tdTomato+;EGFP+ cells (i.e. Axin2+;Sox2+) as well as the single-expressing populations and plated these in equal numbers in stem cell-promoting media at clonal densities (Figure 2B). Double positive tdTomato+;EGFP+ cells showed a significant increase in the efficiency of colony formation compared to single-labelled EGFP+ cells (average 9% compared to 5%, n=5 pituitaries, \(P=0.0226\), Mann-Whitney U test (two-tailed)), demonstrating WNT-responsive SOX2+ PSCs have a greater clonogenic
potential under these in vitro conditions, confirming in vivo data in Figure 1B. As expected from previous work, none of the single-labelled tdTomato\textsuperscript{+} cells (i.e. SOX2 negative) were able to form colonies (Andoniadou et al., 2012).

To confirm that PSCs with active WNT signalling through β-catenin have a greater propensity to form colonies in vitro, we analysed postnatal pituitaries from TCF/Lef: H2B-EGFP mice, reporting the activation of response to WNT signals. This response is detected through expression of an EGFP-tagged variant of histone H2B, which is incorporated into chromatin and diluted in descendants with cell division (Ferrer-Vaquer et al., 2010). Therefore, cells responding to, or having recently responded to WNT, as well as their immediate descendants will be EGFP\textsuperscript{+}. At P21, EGFP\textsuperscript{+} cells were abundant in all three lobes and particularly in the marginal zone harbouring SOX2\textsuperscript{+} stem cells (Figure 2 – figure supplement 2A). Through double mRNA in situ hybridisation against Egfp and Sox2 in TCF/Lef: H2B-EGFP pituitaries, we confirmed that Sox2-expressing cells activate H2B-EGFP expression at this time point (Figure 2 – figure supplement 2B). Isolation by fluorescence-activated cell sorting and in vitro culture of the postnatal EGFP\textsuperscript{+} compartment revealed an enrichment of cells with clonogenic potential in the EGFP\textsuperscript{High} fraction compared to EGFP\textsuperscript{Low} or negative cells (Figure 2 – figure supplement 2C, n=5 pituitaries).

Together these results reveal that a proportion of postnatal SOX2\textsuperscript{+} stem cells respond to WNTs through downstream β-catenin/TCF/LEF signalling and that these cells have greater clonogenic capacity in vitro.

To further address the role of the canonical WNT response in the activity of SOX2\textsuperscript{+} PSCs in vivo, we expressed a loss-of-function allele of β-catenin specifically in Sox2-expressing cells (Sox2\textsuperscript{CreERT2/+};Ctnnb1\textsuperscript{lox(ex2-6)/lox(ex2-6)} hereby Sox2\textsuperscript{CreERT2/+};Ctnnb1\textsuperscript{LOF/LOF}) from P14. Twenty-two weeks following induction, at
P168, there was a substantial drop in the number of cycling cells in the pituitary of
Sox2<sup>CreERT2/+</sup>;Ctnnb1<sup>LOF/LOF</sup> mutants compared to Sox2<sup>+/+</sup>;Ctnnb1<sup>LOF/LOF</sup> controls
(Figure 2C, n=2 pituitaries per genotype). This was accompanied by anterior pituitary
hypoplasia following the loss of Ctnnb1 in SOX2<sup>+</sup> PSCs (Figure 2D). Therefore, in
this small sample size, the proliferative capacity of Ctnnb1-deficient SOX2<sup>+</sup> PSCs
and of their descendants was impaired long-term, leading to reduced growth. In vivo
genetic tracing of targeted cells over the 22-week period
(Sox2<sup>CreERT2/+</sup>;Ctnnb1<sup>LOF/+</sup>;ROSA26<sup>mTmG/+</sup> compared to
Sox2<sup>CreERT2/+</sup>;Ctnnb1<sup>LOF/LOF</sup>;ROSA26<sup>mTmG/+</sup> pituitaries) revealed that targeted
(Ctnnb1-deficient) SOX2<sup>+</sup> PSCs were capable of giving rise to the three committed
lineages PIT1, TPIT and SF1 (Figure 2 – figure supplement 2D), indicating that the
loss of Ctnnb1 does not prevent differentiation of SOX2<sup>+</sup> PSCs into the three
lineages. Downregulation of β-catenin was confirmed by immunofluorescence in
SOX2<sup>+</sup> (mGFP<sup>+</sup>) derivatives (Figure 2 – figure supplement 2E). Although limited by
a small sample size, we conclude that WNT/β-catenin signalling is likely required
cell-autonomously in SOX2<sup>+</sup> stem cells and their descendants (Figure 2E).

**SOX2<sup>+</sup> stem cells express WNT ligands.**

Having established that WNT activation is responsible for promoting proliferation in
the AP, we next focused on identifying the source of WNT ligands. Axin2 expressing
cells from Axin2<sup>CreERT2/+</sup>;ROSA26<sup>mTmG/+</sup> mice were labelled at P14 by tamoxifen
induction. Cells expressing Axin2 at the time of induction are labelled by GFP
expression in the membrane. Double immunofluorescence staining for GFP together
with SOX2 revealed that Axin2 expressing cells (mGFP<sup>+</sup>) are frequently located in
close proximity to SOX2<sup>+</sup> PSCs (Figure 3A). Two-dimensional quantification of the
two cell types revealed that over 50% of mGFP+ cells were in direct contact with SOX2+ nuclei (n=3 pituitaries, >500 SOX2+ cells per gland, Figure 3A). The analysis did not take into account the cellular processes of SOX2+ cells. These results led us to speculate that SOX2+ PSCs may be a source of key WNT ligands promoting proliferation of lineage-committed cells.

In order to determine if SOX2+ PSCs express WNT ligands, we carried out gene expression profiling of SOX2+ and SOX2- populations at P14, through bulk RNA-sequencing. Pure populations of Sox2-expressing cells excluding lineage-committed populations, were isolated from Sox2Egfp/+ male and female pituitaries at P14 based on EGFP expression as previously shown (Andoniadou et al., 2012) (Figure 3B, Figure 3 – figure supplement 1A). Analysis of global gene expression signatures using ‘Gene Set Enrichment Analysis’ (GSEA) (Subramanian et al., 2005) identified a significant enrichment of molecular signatures related to EMT, adherens and tight junctions in the EGFP+ fraction, characteristic of the SOX2+ population (Figure 3 – figure supplement 1B). The SOX2+ fraction also displayed enrichment for genes associated with several signalling pathways known to be active in these cells, including EGFR (Iwai-Liao et al., 2000), Hippo (Lodge et al., 2016; Lodge et al., 2019; Xekouki et al., 2019), MAPK (Haston et al., 2017), FGF (Higuchi et al., 2017), Ephrin (Yoshida et al., 2015; Yoshida et al., 2017) and p53 (Gonzalez-Meljem et al., 2017) (Figure 3 – figure supplement 1C, Supplementary File 1). Additionally, PI3K, TGFβ and BMP pathway genes were significantly enriched in the SOX2+ population (Figure 3 – figure supplement 1C, Supplementary File 1). Query of the WNT-associated genes did not suggest a global enrichment in WNT targets (e.g. enrichment of Myc and Jun, but not of Axin2 or Lef1) (Figure 3 – figure supplement 1D, Supplementary File 1). Instead, SOX2+ PSCs expressed a unique transcriptomic
fingerprint of key pathway genes including Lgr4, Znrf3, Rnf43 capable of regulating
WNT signal intensity in SOX2+ PSCs, as well as enriched expression of the receptors
Fzd1, Fzd3, Fzd4, Fzd6 and Fzd7 (Figure 3 – figure supplement 1D). The
predominant R-spondin gene expressed in the pituitary was Rspo4, specifically by the
EGFP-negative fraction (Figure 3 – figure supplement 1D). The gene profiling
revealed that Wls expression, encoding Gpr177/WLS, a necessary mediator of WNT
ligand secretion (Carpenter et al., 2010; Takeo et al., 2013; Wang et al., 2015), is
enriched in SOX2+ PSCs (Figure 3C). Analysis of Wnt gene expression confirmed
enriched expression of Wnt2, Wnt5a and Wnt9a in SOX2+ PSCs, and the expression
of multiple additional Wnt genes by both fractions at lower levels (SOX2+ fraction:
Wnt5b, Wnt6, Wnt16; SOX2- fraction: Wnt2, Wnt2b, Wnt3, Wnt4, Wnt5a, Wnt5b,
Wnt9a, Wnt10a, Wnt16) (Figure 3D). These results reveal that SOX2+ PSCs express
the essential components to regulate activation of the WNT pathway and express Wnt
genes as well as the necessary molecular machinery to secrete WNT ligands.

Paracrine signalling from SOX2+ stem cells promotes WNT activation.
We sought to conclusively determine if WNT secretion specifically from SOX2+
PSCs drives proliferation of surrounding cells in the postnatal pituitary gland. We
proceeded to delete Wls only in the Sox2-expressing population (Sox2CreERT2/+; Wls0/0)
from P14 by a series of tamoxifen injections. Due to morbidity, we limited analyses to
one week following induction. Pituitaries appeared mildly hypoplastic at P21 along
the medio-lateral axis (Figure 4 – figure supplement 1, n=4 controls and n=5
mutants). To determine if this is a result of reduced proliferation, we carried out
immunofluorescence using antibodies against Ki-67 and SOX2. This revealed
significantly fewer cycling cells in the SOX2- population of Sox2CreERT2/+; Wls0/0
mutant pituitaries compared to Sox2+/+;Wls0/0 controls (10.326% Ki-67 in control
(n=4) compared to 3.129% in mutant (n=5), P=0.0008, unpaired t-test) (Figure 4A).
Additionally, we observed a reduction of cycling cells within the SOX2+ population
(5.582% Ki-67 in control compared to 2.225% in induced Sox2CreERT2/+/Wls0/0 mutant
pituitaries, P=0.0121, unpaired t-test) (Figure 4A), resulting in a smaller SOX2+ cell
pool in mutants (23.425% SOX2+/total AP cells in Sox2+/+;Wls0/0 controls compared
to 19.166% SOX2+/total AP cells in induced Sox2CreERT2/+/Wls0/0 mutant pituitaries,
P=0.0238, Student’s t-test, n=5 mutants, 4 controls). To determine if reduced levels of
WNT activation accompanied this phenotype, we carried out double mRNA in situ
hybridisation using specific probes against Lef1 and Sox2. There was an overall
reduction in Lef1 expression in mutants compared to controls (n=4 per genotype), in
which we frequently observed robust expression of Lef1 transcripts in close proximity
to cells expressing Sox2 (arrows, Figure 4B). Together, our data support a paracrine
role for SOX2+ pituitary stem cells in driving the expansion of committed progeny
through the secretion of WNT ligands (Figure 4C).

DISCUSSION
Emerging disparities between the archetypal stem cell model, exhibited by the
haematopoietic system, and somatic stem cells of many organs, have led to the
concept that stem cell function can be executed by multiple cells not fitting a typical
stem cell paradigm (Clevers and Watt, 2018). In organs with persistent populations
possessing typical functional stem cell properties yet contributing minimally to
turnover and repair, the necessity for such classical stem cells is questioned. Here we
show that WNT signalling is required for postnatal pituitary growth by both SOX2+
PSCs as well as SOX2+ committed progenitors. We identify an additional discreet
function for SOX2+ PSCs, where these signal in a feedforward manner by secreting WNT ligands as cues to stimulate proliferation and promote tissue growth.

Consistent with previous reports, our data support that SOX2+ PSCs contribute, but do not carry out the majority of tissue expansion during the postnatal period (Zhu et al., 2015); instead, new cells primarily derive from more committed progenitors, which we show to be WNT-responsive. We demonstrate that this population of lineage-restricted WNT-responsive cells rapidly expands and contributes long-lasting clones from postnatal stages. It remains to be shown if cells among the SOX2 lineage-committed populations may also fall under the classical definition of a stem cell. Preventing secretion of WNT ligands from SOX2+ PSCs reveals that far from being dispensable, paracrine actions of the SOX2+ population that are inactive in their majority, are necessary for anterior lobe expansion from lineage-committed populations. In the adrenal gland, R-spondins are necessary for cortical expansion and zonation, where deletion of Rspo3, expressed by the capsule which contains adrenocortical stem cells, results in reduced proliferation of the underlying steroidogenic cells (Vidal et al., 2016). Corroborating a model where committed pituitary progenitors depend on the paracrine actions of SOX2+ PSCs, Zhu and colleagues observed that in pituitaries with reduced numbers of PSCs, proliferation among PIT1+ cells was significantly impaired (Zhu et al., 2015). It would be intriguing to see if there is a reduction in WNT signalling in this model, or following genetic ablation of adult SOX2+ PSCs (Roose et al., 2017).

We show that a sub-population of SOX2+ PSCs in the postnatal gland are also WNT-responsive and have greater \textit{in vitro} colony-forming potential under defined conditions. This colony-forming potential is normally a property of a minority of SOX2+ PSCs at any given age and reflects their \textit{in vivo} proliferative capacity.
Andoniadou et al., 2012; Rizzoti et al., 2013). A role for the WNT pathway in promoting SOX2\(^+\) cell activity is supported by studies showing that pathogenic overexpression of β-catenin promotes their colony-forming ability (Sarkar et al., 2016), and their *in vivo* expansion (Andoniadou et al., 2012). Additionally, elevated WNT pathway activation has been described for pituitary side-population cells, enriched for SOX2\(^+\) stem cells from young, compared to old pituitaries (Gremeaux et al., 2012). This is in line with our findings that the WNT pathway has an important function in promoting the activation of SOX2\(^+\) PSCs. It remains to be shown if this response relies on autocrine WNT-signalling as for other stem cells (Lim et al., 2013), however our results reveal reduced proliferation among SOX2\(^+\) PSCs and reduced SOX2\(^+\) cell numbers when WNT secretion from these cells is abolished, supportive of either autocrine signalling, or paracrine signalling between different subsets of the SOX2\(^+\) population.

The mechanism preventing the majority of SOX2\(^+\) PSCs from responding to WNT signals remains elusive but points to heterogeneity among the population. Such regulation could occur at the level of receptor signalling; we have shown by bulk transcriptomic profiling that SOX2\(^+\) PSCs express the receptors required to respond to the WNT pathway, but also express high levels of the frizzled inhibitor *Znrf3*, and the R-spondin receptor *Lgr4*. One conceivable scenario is that high levels of *Znrf3* inhibit frizzled receptors in the absence of R-spondin under normal physiological conditions, supressing a WNT response. In support of this, R-spondins have been shown to promote pituitary organoid formation (Cox et al., 2019). Whether the R-spondin/LGR/ZNRF3 module is active under physiological conditions needs to be determined. Furthermore, well-described factors expressed in PSCs are known to have inhibitory effects on β-catenin-mediated transcription, such as YAP/TAZ.
(Azzolin et al., 2014; Gregorieff et al., 2015) and SOX2 itself (Alatzoglou et al., 2011; Kelberman et al., 2008).

In summary, we demonstrate an alternative mechanism for stem cell contribution to homeostasis, whereby these can act as paracrine signalling hubs to promote local proliferation. Applicable to other organs, this missing link between SOX2+ PSCs and committed cell populations of the anterior pituitary, is key for basic physiological functions and renders stem cells integral to organ expansion.
MATERIALS AND METHODS

Mice

All procedures were performed under compliance of the Animals (Scientific Procedures) Act 1986, Home Office License (P5F0A1579). KCL Biological Services Unit staff undertook daily animal husbandry. Genotyping was performed on ear biopsies taken between P11 and P15 by standard PCR using the indicated primers. These experiments were not conducted at random and the experimenters were not blind while conducting the animal handling and assessment of tissue. Images are representative of the respective genotypes. For all studies, both male and female animals were used and results combined.

The Sox2<sup>CreERT2/+</sup> and Sox2<sup>Egfp/+</sup> strains were kept on a CD-1 background. Axin2<sup>CreERT2/+</sup> animals were kept on a mixed background of C57BL/6 backcrossed onto CD-1 for 5 generations and were viable and fertile, with offspring obtained at the expected Mendelian ratios. ROSA26<sup>mTmG/mTmG</sup>, ROSA26<sup>Confetti/Confetti</sup>, ROSA26<sup>tdTomato/tdTomato</sup>, Wls<sup>fl/fl</sup>, Ctnnb1<sup>fl(ex2-6)/fl(ex2-6)</sup> and TCF/LEF:H2B-EGFP mice were kept on a mixed background of 129/Sv backcrossed onto CD-1 for at least 3 generations. For lineage tracing studies, male Axin2<sup>CreERT2/+</sup> or Sox2<sup>CreERT2/+</sup> mice were bred with homozygous ROSA26<sup>mTmG/mTmG</sup> or ROSA26<sup>Confetti/Confetti</sup> dams to produce the appropriate allele combinations on the reporter background. Pups were induced at P14 or P15 with a single dose of tamoxifen (resuspended to 20mg/ml in Corn Oil with 10% ethanol) by intraperitoneal injection, at a concentration of 0.15mg per gram of body weight. Pituitaries were harvested at the indicated time points post induction and processed for further analysis as described below. Mice were harvested from different litters for each time point at random. For litters in which there was a...
surplus of experimental mice, multiple samples were harvested for each required time point.

For Wntless deletion studies, Sox2<sup>CreERT2/+;Wls<sup>fl/+;ROSA26<sup>tmG/mTmG males were bred with Wls<sup>fl/+;ROSA26<sup>tmG/mTmG dams, to produce Sox2<sup>CreERT2/+;Wls<sup>fl/+;ROSA26<sup>tmG/mTmG, Sox2<sup>CreERT2/+;Wls<sup>fl/+;ROSA26<sup>tmG/mTmG and Wls<sup>fl/+;ROSA26<sup>tmG/mTmG offspring. Pups of the indicated genotypes received intraperitoneal injections of 0.15mg of tamoxifen/gram body weight on 4 consecutive days, beginning at P14, and harvested 3 days after the final injection.

For the β-catenin loss-of-function experiments, either Sox2<sup>CreERT2/+;Ctnnb1<sup>fl(ex2-6)/+;ROSA26<sup>tmG/mTmG or Axin2<sup>CreERT2/+;Ctnnb1<sup>fl(ex2-6)/+;ROSA26<sup>tmG/mTmG males were crossed with Ctnnb1<sup>fl(ex2-6)/fl(ex2-6);ROSA26<sup>tmG/mTmG dams. Axin2<sup>CreERT2/+;Ctnnb1<sup>fl(ex2-6)/+;ROSA26<sup>tmG/mTmG pups were induced with a single dose of tamoxifen, at a concentration of 0.15mg per gram of body weight and kept alive for 7 days before harvesting. Sox2<sup>CreERT2/+;Ctnnb1<sup>fl(ex2-6)/+;ROSA26<sup>tmG/mTmG and Sox2<sup>CreERT2/+;Ctnnb1<sup>fl(ex2-6)/fl(ex2-6);ROSA26<sup>tmG/mTmG pups received two intraperitoneal injections of tamoxifen, at a concentration of 0.15mg/gram body weight, on two consecutive days and were kept alive for the indicated length of time before harvesting.

TCF/LEF:H2B-EGFP mice were culled and the pituitaries harvested at the indicated ages for the respective experiments. For fluorescence-activated cell sorting experiments, mice were harvested at 21 days of age. Axin2<sup>CreERT2/+;Sox2<sup>eGFP/+ males were crossed with ROSA26<sup>tdTomat/tdTomato dams to produce Axin2<sup>CreERT2/+;Sox2<sup>eGFP/+;ROSA26<sup>tdTomat/+ that were induced with single doses of tamoxifen at 21 and 22 days of age and harvested three days after the first injection for fluorescence-activated cell sorting experiments.
Flow cytometry analysis of lineage traced pituitaries

For the quantification of cells by flow cytometry, anterior lobes of
Axin2\textsuperscript{CreERT2/+};ROSA26\textsuperscript{mTmG/+} mice dissected at the indicated time points. The
posterior and intermediate lobes were dissected from the anterior lobes under a
dissection microscope. Untreated ROSA26\textsuperscript{mTmG/+} and wild type pituitaries from age-
matched litters were used as tdTomato only and negative controls, respectively.
Dissected pituitaries were incubated in Enzyme Mix (0.5% w/v collagenase type 2
(Lorne Laboratories), 0.1x Trypsin (Gibco), 50\textmu g/ml DNase I (Worthington) and
2.5\textmu g/ml Fungizone (Gibco) in Hank’s Balanced Salt Solution (HBSS)(Gibco)) in a
cell culture incubator for up to 3 hours. 850ml of HBSS were added to each
Eppendorf in order to quench the reaction. Pituitaries were dissociated by agitation,
pipetting up and down 100x at first with a 1ml pipette, followed by 100x with a 200\textmu l
pipette. Cells were transferred to a 15ml Falcon tube and resuspended in 9ml of HBSS
and spun down at 200g for 5 minutes. The supernatant was aspirated, leaving behind
the cell pellet that was resuspended in PBS and spun down at 1000rpm for 5 minutes
before being resuspended in a Live/Dead cell stain (Life Technologies, L34975)
prepared to manufacturer’s instructions, for 30 minutes in the dark. Cells were
washed in PBS as above. The pellet was resuspended in FIX & PERM Cell
Permeabilization Kit (Life Technologies, GAS003) prepared as per manufacturer’s
instructions for 10 minutes at room temperature. Cells were washed as above, and the
pellet was resuspended in 500\textmu l of FACS buffer (1% fetal calf serum (Sigma), 25mM
HEPES in PBS) and filtered through 70\textmu m filters (BD Falcon), into 5ml round
bottom polypropylene tubes (BD Falcon). 1 minute prior to analysis, 1\mu l of Hoechst
was added to the suspended cells and incubated. Samples were analysed on a BD
Fortessa, and gated according to negative and single fluorophore controls. Single cells were gated according to SSC-A and SSC-W. Dead cells were excluded according to DAPI (2ng/ml, incubated for 2 mins prior to sorting). GFP⁺, tdTomato⁺ and GFP⁺;tdTomato⁺ cells were gated according to negative controls in the PE-A and FITC-A channels.

**Fluorescence Activated Cell Sorting for sequencing or colony forming assays**

For fluorescence activated cell sorting, the anterior lobes from Sox2eGFP/+, TCF/LEF:H2B-GFP or Axin2CreERT2/++;Sox2eGFP/+;ROSA26tdTomato/+ and their respective controls were dissected and dissociated as above. After dissociation cells were spun down at 200g in HBSS and the pellet was resuspended in 500µl FACS buffer. Using an Aria III FACs machine (BD systems), samples were gated according to negative controls, and where applicable single fluorophore controls. Experimental samples were sorted according to their fluorescence, as indicated, into tubes containing either RNAlater (Qiagen) for RNA isolation or 1ml of Pit Complete Media for culture (Pit Complete: 20ng/ml bFGF and 50ng/ml of cholera toxin in ‘Pit Basic’ media (DMEM-F12 with 5% Fetal Calf Serum,100U/ml Penicillin and 100µg/ml Streptomycin). Cells were plated in 12-well plates at clonal density, approximately 500 cells/well. Colonies were incubated for 7 days total before being fixed in 10% neutral buffered formalin (NBF) (Sigma) for 10 minutes at room temperature, washed for five minutes, three times, mins with PBS and stained with crystal violet in order for the number of colonies to be quantified.

**RNA-sequencing**
Total RNA was isolated from each sample and following poly-A selection, cDNA libraries were generated using TruSeq (Clontech, 634925). Barcoded libraries were then pooled at equal molar concentrations and sequenced on an Illumina Hiseq 4000 instrument in a 75 base pair, paired-end sequencing mode, at the Wellcome Trust Centre for Human Genetics (Oxford, United Kingdom). Raw sequencing reads were quality checked for nucleotide calling accuracy and trimmed accordingly to remove potential sequencing primer contaminants. Following QC, forward and reverse reads were mapped to GRCm38/mm10 using Hisat2 (Kim et al., 2015). Using a mouse transcriptome specific GTF as a guide, FeatureCounts (Liao et al., 2014) was used to generate gene count tables for every sample. These were utilised within the framework of the Deseq2 (Love et al., 2014) and FPKM values (generated by FPKM count (Wang et al., 2012)) were processed using the Cufflinks (Trapnell et al., 2012) pipelines which identified statistically significant gene expression differences between the sample groups. Following identification of differentially expressed genes (at an FDR < 0.05) we focused on identifying differentially expressed pathways using a significance threshold of FDR < 0.05 unless otherwise specified. The gene lists used for Gene Set Enrichment Analysis (GSEA) were as found on the BROAD institute GSEA MSigDBv.7 ‘molecular signatures database’. The deposited dataset can be accessed through the following link:

https://dataview.ncbi.nlm.nih.gov/object/PRJNA421806?reviewer=kr90aklsdtikh3gkh3tdlpv30s

**Immunofluorescence and microscopy**

Freshly harvested pituitaries were washed in PBS for 10 minutes before being fixed in 10% NBF for 18 hours at room temperature. In short, embryos and whole pituitaries
were washed in PBS 3 times, before being dehydrated through a series of 1 hour washes in 25%, 50%, 70%, 80%, 90%, 95% and 100% ethanol. Tissues were washed in Neo-Clear (Sigma) at room temperature for 10 minutes, then in fresh preheated Neo-Clear at 60 °C for 10 minutes. Subsequently, a mixture of 50% Neo-Clear:50% paraffin wax at 60°C for 15 minutes followed by three changes of pure wax for a minimum of 1 hour washes at 60°C, before being orientated to be sectioned in the frontal plane. Embedded samples were sectioned at 5µm and mounted on to Super Frost+ slides.

For immunofluorescence, slides were deparaffinised in Neo-Clear for three times ten minutes, washed in 100% ethanol for three times five minutes, and rehydrated in a series of five minute ethanol washes up to distilled water (95%, 90%, 80%, 70%, 50%, 25%, H2O). Heat induced epitope retrieval was performed with 1x DeClear Buffer (citrate pH 6) in a Decloaking chamber NXGEN (Menarini Diagnostics) for 3 minutes at 110°C. Slides were left to cool to room temperature before proceeding to block for 1 hour at room temperature in Blocking Buffer (0.2% BSA, 0.15% glycine, 0.1% TritonX in PBS) with 10% serum (sheep or donkey, depending on secondary antibodies). Primary antibodies were diluted in blocking buffer with 1% of the appropriate serum and incubated overnight at 4°C. Slides were washed three times for 10 minutes with PBST. Slides were incubated with secondary antibodies diluted 1:400 in blocking buffer with 1% serum for one hour at room temperature. Slides were washed three times with PBST as above. Where biotinylated secondary antibodies were used, slides were incubated with streptavidin diluted 1:400 in blocking buffer with 1% serum for one hour at room temperature. Finally, slides were washed with PBST and mounted using Vectashield Antifade Mounting Medium (Vector Laboratories, H-1000).
The following antibodies, along with their dilutions and detection technique, were used: GFP (1:400, Alexa Fluor-488 or -647 secondary), SOX2 raised in goat (1:200, Alexa Fluor-488 secondary), SOX2 raised in rabbit (1:100, biotinylated secondary), SOX9 (1:500, biotinylated secondary), PIT1 (1:500, biotinylated secondary), SF1 (1:300, biotinylated secondary), TPIT (1:200, biotinylated secondary), Ki-67 (1:100, biotinylated secondary), pH-H3 (1:500, biotinylated secondary), GH (1:1000, biotinylated secondary), TSH (1:1000, biotinylated secondary), PRL (1:1000, biotinylated secondary), ACTH (1:400, Alexa Fluor-555 secondary), LH/FSH (1:300, biotinylated secondary), ZO-1 (1:300, Alexa Fluor-488), E-Cadherin (1:300, Alexa Fluor-488). Nuclei were visualized with Hoechst (1:1000). Images were taken on a TCS SPS Confocal (Leica Microsystem) with a 20x objective for analysis.

**mRNA In Situ Hybridisation**

All mRNA in situ hybridisations were performed using the RNAscope singleplex or duplex chromogenic kits (Advanced Cell Diagnostics) on formalin fixed paraffin embedded sections processed as described in the above section. The protocol followed the manufacturer’s instructions with slight modifications. ImmEdge Hydrophobic Barrier PAP Pen (Vector Laboratories, H-4000) was used to draw a barrier around section while air-drying following the first ethanol washes. Pretreatment followed the standard length of time for pituitaries (twelve minutes), while embryos were boiled for 10 minutes. For singleplex, the protocol proceeded to follow the instructions exactly. For duplex, Amplification 9 was extended to one hour and the dilution of the Green Detection reagent was increased to 1:30. For both protocols, sections were counterstained with Mayer’s Haematoxylin (Vector Laboratories, H-3404), left to dry at 60°C for 30 minutes before mounting with
VectaMount Permanent Mounting Medium (Vector Laboratories, H-5000). Slides were scanned using a Nanozoomer-XR Digital Slide Scanner (Hamamatsu) and processed using Nanozoomer Digital Pathology View (Hamamatsu).

**Quantification of cells**

Cell numbers were quantified in ImageJ using the cell counter plugin (Schindelin et al., 2012). At a minimum, three sections per pituitary were quantified, spaced no less than 100µM apart in the tissue.

**Statistics**

All statistical analyses were performed in GraphPad Prism. Data points in graphs represent the mean values of recordings from a single biological replicate unless otherwise stated.

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FIGURES

Figure 1. Axin2 expressing cells contribute to pituitary growth and expansion of all lineages

A. Immunofluorescence staining against GFP (green) with markers of PSCs or lineage commitment (magenta) in Axin2<sup>CreERT2</sup>/+; ROSA26<sup>mTmG</sup>/+ pituitaries harvested from mice induced at P14 and lineage traced for 2 days (top panel) and 14 days (bottom panel). Scale bar 10 µm.

B. Quantification of lineage expansion between 2 and 14 days following induction at P14. Graph shows that the proportion of lineage committed cells (either PIT1<sup>+</sup>, TPIT<sup>+</sup> or SF1<sup>+</sup>) and PSCs (SOX2<sup>+</sup>), i.e. that are transcription factor (TF)<sup>+</sup> cells that are GFP<sup>+</sup> increases between 2 days (black bars) and 14 days (grey bars) post induction. PIT1 <i>P</i>=0.000004, TPIT <i>P</i>=0.008 multiple <i>t</i>-tests. <i>n</i>=4 animals per time point.

C. Immunofluorescence staining against GFP (green) in pituitaries harvested from Axin2<sup>CreERT2</sup>/+; ROSA26<sup>mTmG</sup>/+ mice induced at P14 and lineage traced for 2 days, 2 weeks and 8 weeks. Bottom panel shows magnified fields of view of regions of interest indicated by white boxes in panels above. Scale bars 50 µm.

D. Top panel showing the quantification of the proportion of all cells in Axin2<sup>CreERT2</sup>/+; ROSA26<sup>mTmG</sup>/+ pituitaries that are GFP<sup>+</sup> at 2, 7, 14, 28 and 56 days.
post induction as analysed by flow cytometry. Day 2 to 7 \( P<0.0001 \) unpaired \( t \)-test. Data points show individual measurements from biological replicates, \( n=4-8 \) pituitaries per time point. (Bottom) Graph of the absolute number of GFP+ cells (green) and as a proportion of total cells (blue) at the time points indicated.

E. X-gal staining in \( Axin2^{CreERT2/+},ROSA26^{LacZ/+} \) pituitaries harvested from mice induced at P14 and lineage traced for 8 weeks (left) and 1 year (right). Scale bars 500\( \mu m \).

F. Model summarising the major contribution of WNT-responsive progenitors of all lineages to pituitary growth, in addition to that of SOX2+ PSCs.
Figure 2. Activation of WNT signalling in SOX2\(^+\) PSCs and their descendants is necessary for long-term growth

A. Schematic of the experimental timeline used in panels A and B. Endogenous expression of tdTomato (magenta, Axin2 targeted cells) and EGFP (green, Sox2 expressing cells) in Axin2\(^{CreERT2/+}\);Sox2\(^{Egfp/+}\);ROSA26\(^{tdTomato/+}\) pituitaries harvested at P24 sectioned in the frontal plane. Nuclei are counterstained with Hoechst in the merged panel. Scale bar 50\(\mu\)m.

B. A representative culture plate showing colonies derived from Tomato\(^+\), EGFP\(^+\) or Tomato\(^+\);EGFP\(^+\) cells that were isolated from Axin2\(^{CreERT2/+}\);Sox2\(^{Egfp/+}\);ROSA26\(^{tdTomato/+}\) pituitaries by FACS plated in stem cell promoting media at clonogenic densities and stained with crystal violet (left panel). The proportion of colony-forming cells in each subpopulation were quantified by counting the number of colonies per well (right panel). Each data point indicates individual wells, \(n=5\) separate pituitaries. \(P=0.0226\), Mann-Whitney \(U\) test (two-tailed). Scale bar 10mm.

C. Immunofluorescence staining against SOX2 (green) and Ki-67 (magenta) in Sox2\(^{+/+}\);Ctnnb1\(^{LOF/LOF}\) (control) and Sox2\(^{CreERT2/+}\);Ctnnb1\(^{LOF/LOF}\) (mutant) pituitaries from mice induced at P14 and analysed 22 weeks after induction (at P168) (bottom panel). Scale bar 50\(\mu\)m.

D. Dorsal view of whole mount pituitaries of Sox2\(^{+/+}\);Ctnnb1\(^{LOF/LOF}\) (control) and Sox2\(^{CreERT2/+}\);Ctnnb1\(^{LOF/LOF}\) (mutant), 22 weeks after induction (i.e. P168). Scale bars 1mm.

E. Model summarising the effect of Ctnnb1 deletion in SOX2\(^+\) PSCs.

PL, posterior lobe; IL, intermediate lobe; AL, anterior lobe.
Figure 3. SOX2+ PSCs are as a source of WNT ligands in the pituitary

A. Immunofluorescence staining against GFP (green) and SOX2 (magenta) in Axin2<sup>CreERT2</sup>/+; ROSA26<sup>mTmG</sup>/+ pituitaries 48 hours post induction. Graph representing a quantification of the proximity of individual GFP+ cells to the nearest SOX2+ cell as quantified by the number of nuclei separating them. Plotted data represents the proportion of GFP+ cells that fall in to each category of the total GFP+ cells, taken from n=3 separate pituitaries. Scale bars 50μm.

B. Experimental paradigm for RNA Seq analysis of Sox2 positive and negative cells.

C. Graphs representing the FPKM values of Wls and Porcupine in Sox2 positive and negative cells (black and grey bars, respectively). mRNA in situ hybridisation for Sox2 and for Wls on wild type sagittal pituitaries at P14, demonstrating strong Wls expression in the marginal zone epithelium. Scale bars 250μm.

D. Bar chart showing the FPKM values of Wnt genes in the Sox2+ and Sox2- fractions. Double mRNA in situ hybridisation against Wnt2, Wnt5a and Wnt9a (blue) together with Sox2 (red) validating expression in the Sox2+ population. Boxed regions through the marginal zone epithelium are magnified. Scale bars 100μm and 50μm in boxed inserts.
Figure 4. Paracrine secretion of WNTs from SOX2+ PSCs is necessary for expansion of committed cells

A. Immunofluorescence staining against SOX2 (green) and Ki-67 (magenta) in Sox2+/+;Wlsflo/flo (control) and Sox2CreERT2+/+;Wlsflo/flo (mutant) pituitaries induced from P14 and analysed after one week. Nuclei were counterstained with Hoechst. (i) and (ii) represent magnified fields of view of regions indicated by white boxes in top panels. Scale bar 50µm. Graph of quantification of cycling cells marked by Ki-67 among cells negative for SOX2. Values represent mean +/- SEM, P=0.0008, unpaired t-test. Graph of quantification of cycling cells marked by Ki-67 among SOX2-positive cells. Values represent mean +/- SEM, P=0.0121, unpaired t-test. Each data point shows the mean of one biological replicate, n=4 pituitaries from controls and 5 pituitaries from mutants.

B. Double mRNA in situ hybridisation using specific probes against Lef1 (blue) and Sox2 (red) in control and mutant pituitaries following tamoxifen induction from P14 and tracing for 7 days. Scale bars 250µm and 50µm in boxed regions.

C. Model summarising paracrine WNT secretion from SOX2+ PSCs to lineage-committed progenitors and the effects of abolishing WNT secretion from SOX2+ PSCs through the deletion of Wls.
SUPPLEMENTARY INFORMATION

Supplementary File 1. Gene lists of Gene Set Enrichment Analyses

Gene lists generated from Gene Set Enrichment Analyses of bulk RNA-sequencing data comparing Sox2⁺ and Sox2⁻ cells. Associated with Figure 3 – figure supplement 1.

SUPPLEMENTARY FIGURES

Figure 1 – figure supplement 1. Axin2 expressing cells contribute to pituitary growth and expansion of all lineages

A. Schematic of the combined experimental timeline used in panels A, B and C.
Immunofluorescence staining against GFP (green) and markers of hormone-secreting endocrine cells (GH (somatotrophs), ACTH (corticotrophs), PRL (lactotrophs), TSH (thyrotrophs), FSH/LH (gonadotrophs)) in Axin2CreERT2/⁺;Rosa26mTmG/⁺ pituitaries induced at P14 and lineage traced for 48 hours. Scale bar 10µm.

B. Graph of quantification of expansion of the WNT-responsive SF1⁺ population in Axin2CreERT2/⁺;ROSA26mTmG/⁺ pituitaries induced at P14 and lineage traced for 2 or 28 days. There is a significant increase of GFP⁺;SF1⁺ cells as a proportion of the total SF1⁺ cells at P28. P=0.0048, unpaired t-test (n=2 at 2 days, 3 at 28 days).

C. Immunofluorescence staining against GFP (green) and markers of hormone-secreting endocrine cells of the PIT1 lineage (GH (somatotrophs), PRL (lactotrophs), TSH (thyrotrophs)) in Axin2CreERT2/⁺;ROSA26mTmG/⁺ pituitaries induced at P14 and lineage traced for 14 days. Scale bars 50µm. Graph showing expansion of each of the Hormone⁺ cell types (Hormone⁺;GFP⁺) as a percentage of the total Hormone⁺ population between 2 and 14 days post-induction. There is
a significant increase in GH$^+$ somatotrophs ($P=0.000548$), and TSH$^+$ thyrotrophs ($P=0.0016$), whilst there is no significance (ns) between PRL$^+$ lactotroph populations between the two time points. multiple $t$-test ($n=3$ at 48 h, $n=4$ at 14 days post-induction).

D. Clonal analysis of individual cells targeted in $SOX2^{CreERT2/+};ROSA26^{Confetti/+}$ (left panel) and $Axin2^{CreERT2/+};ROSA26^{Confetti/+}$ pituitaries (right panel), induced at P14 and harvested after 4 weeks (P42). Arrows point to individual clones, numbered for the number of cells in the clone. Scale bar 100µm.

**Figure 1 – figure supplement 2. Axin2 expressing cells contribute to pituitary growth and expansion of all lineages**

A. Dorsal wholemount view of $Axin2^{CreERT2/+};Ctnnb1^{LOF/+};ROSA26^{mTmG/+}$ and $Axin2^{CreERT2/+};Ctnnb1^{LOF/LOF};ROSA26^{mTmG/+}$ pituitaries induced at P14 and lineage traced for 5 days. Scale bars 500µm. Immunofluorescence staining against GFP (green) and pH-H3 (magenta) in $Axin2^{CreERT2/+}$; $Ctnnb1^{LOF/+};ROSA26^{mTmG/+}$ and $Axin2^{CreERT2/+};Ctnnb1^{LOF/LOF};ROSA26^{mTmG/+}$ pituitaries. Scale bar 50µm. Quantification of the contribution of lineage traced cells in control and mutants. Each data point represents the mean from one individual. $P=0.0313$, unpaired $t$-test ($n=3$).

B. Immunofluorescence staining against GFP (green) and PIT1, SF1 and ACTH (magenta) in $Axin2^{CreERT2/+};Ctnnb1^{LOF/+};ROSA26^{mTmG/+}$ and $Axin2^{CreERT2/+};Ctnnb1^{LOF/LOF};ROSA26^{mTmG/+}$ pituitaries induced at P14 and lineage traced for 5 days. Quantification of the percentage of GFP$^+$ cells, double-positive for each of the lineage markers, showing no significant changes for each lineage
between controls and mutants (Unpaired t-test, PIT1 \( P=0.1729 \), SF1 \( P=0.9488 \), ACTH \( P=0.6186 \). \( n=4 \) controls, 2 mutants). Scale bars 50\( \mu \)m.

C. Immunofluorescence against GFP (green) and SOX2 (magenta) in Axin2\(^{CreERT2/+}\); Ctnnb1\(^{LOF/+}\);ROSA26\(^{mTmG/+}\) and Axin2\(^{CreERT2/+}\);Ctnnb1\(^{LOF/LOF}\);ROSA26\(^{mTmG/+}\) induced at P14 and lineage traced for 5 days (\( n=4 \) controls, 2 mutants). Scale bars 50\( \mu \)m.

D. Immunofluorescence against GFP (green) and Cleaved Caspase-3 (magenta) in Axin2\(^{CreERT2/+}\); Ctnnb1\(^{LOF/+}\);ROSA26\(^{mTmG/+}\) and Axin2\(^{CreERT2/+}\);Ctnnb1\(^{LOF/LOF}\);ROSA26\(^{mTmG/+}\) induced at P14 and lineage traced for 5 days (\( n=4 \) controls, 2 mutants). Scale bars 50\( \mu \)m.
A-E Step-wise gating strategy to isolate WNT-responsive, SOX2-EGFP\(^+\) cells by flow sorting.

A – B Single pituitary cells dissociated from \(Axin2^{CreERT2/+};ROSA26^{tdTomato/+};Sox2^{eGFP/+}\) mice were gated to exclude debris (A) and gated for single cells according to SSC-A and SSC-W (B).

C. Dead cells were excluded according to incorporation of DAPI.

D. Three populations of fluorescent cells were identified and sorted according to the following profiles: GFP\(^-\);tdTomato\(^+\), GFP\(^+\);tdTomato\(^+\) or GFP\(^+\);tdTomato\(^-\).

E. Quantification of the number of GFP\(^+\) cells out of all gated cells (left, \(n=5\) biological repeats), the proportion of all GFP\(^+\) cells that were found to be tdTomato\(^+\) (right, \(n=5\) biological repeats) and a representation of the gating used for quantification (bottom).

A. Confocal images of native GFP fluorescence in frontal sections from TCF/Lef:H2B-EGFP pituitaries at P21. Scale bar 50µm.

B. mRNA \textit{in situ} hybridisation in TCF/Lef:H2B-EGFP pituitaries at P21, detecting \(Egfp\) transcripts (red). Double mRNA \textit{in situ} hybridisation showing overlap between \(Sox2\) (red) and \(Egfp\) (blue) transcripts in pituitaries at P21. White arrowheads indicate double-positive staining. Scale bars 50µm.

C. Immunofluorescence staining against SOX2 (magenta) and GFP (green) in TCF/Lef:H2B-EGFP pituitaries harvested from P21 mice. White arrows indicate
double positive cells. Graph of quantification of the in vitro colony forming potential of GFP cells isolated from P21 TCF/Lef:H2B-EGFP pituitaries by flow sorting. Each data point represents single well replicates. Error bars show SEM, \( P<0.001 \) (One-way ANOVA, \( n=3 \) individual pituitaries). Scale bar 50\( \mu \)m.

Representative scatter plot showing gating used for fluorescence activated cell sorting and population percentages in each gate.

D. Immunofluorescence staining against PIT1, TPIT and SF1 (magenta) in Sox2\textsuperscript{CreERT2/+}; Ctnnb1\textsuperscript{LOF/+}; ROSA26\textsuperscript{mTmG/+} and Sox2\textsuperscript{CreERT2/+}; Ctnnb1\textsuperscript{LOF/LOF}; ROSA26\textsuperscript{mTmG/+} pituitaries 22 weeks post-induction at P14 (age P24). Arrows indicate double positive cells. Scale bar 50\( \mu \)m.

E. Immunofluorescence staining against \( \beta \)-catenin (magenta) and GFP (green) in Sox2\textsuperscript{CreERT2/+}; Ctnnb1\textsuperscript{LOF/+}; ROSA26\textsuperscript{mTmG/+} and Sox2\textsuperscript{CreERT2/+}; Ctnnb1\textsuperscript{LOF/LOF}; ROSA26\textsuperscript{mTmG/+} pituitaries 22 weeks post-induction. Arrowheads indicate double positive cells, arrows indicate GFP\textsuperscript{+} cells that have lost \( \beta \)-catenin expression in mutants. Scale bar 50\( \mu \)m.

PL, posterior lobe; IL, Intermediate lobe; AL, anterior lobe; Inf, infundibulum; RP, Rathke’s pouch; Sph, sphenoid bone.
Figure 3 – figure supplement 1. SOX2⁺ PSCs are as a source of WNT ligands in the pituitary

A. Native EGFP protein expression in frontal cryosection of a P14 Sox2<sup>Egfp/+</sup> pituitary. Schematic of the workflow used for bulk RNA-sequencing analysis of Sox2⁺ and Sox2⁻ cells. Genome browser views of reads aligning to the Sox2 and Pit1 loci in the positive and negative fractions indicating good separation of the EGFP⁺ population. Scale bar 50µm.

B. Sox2⁺ cells express a significant enrichment in markers associated with epithelial-to-mesenchymal transition (EMT), adherens and tight junctions, consistent with their epithelial nature. GSEA plots and immunofluorescence staining against E-Cadherin (adherens junction marker) and ZO1 (tight junction marker) in the marginal zone epithelium at P14. Scale bar 50µm. See Supplementary File 1 for full GSEA gene lists.

C. Sox2⁺ cells express a significant enrichment in several signalling pathways, shown with respective GSEA plots. See Supplementary File 1 for full GSEA gene lists.

D. Bar charts showing the FPKM values of components of the LGR/RNF43/ZNRF3/RESPONDIN module in the Sox2⁺ and Sox2⁻ fractions and the distribution of the Frizzled receptors. GSEA plot for components of the WNT pathway. Validation of sequencing: (i) mRNA <i>in situ</i> hybridisation with specific probes against Lgr4 (blue) and Sox2 (red) in P14 pituitaries showing co-expression. (ii) Double mRNA <i>in situ</i> hybridisation against Fzd4 (blue) and Sox2 (red) indicating co-expression in both the marginal zone epithelium and parenchymal Sox2⁺ cells. Boxed regions are magnified. Scale bars 250µm and 50µm in boxed inserts. (iii) mRNA <i>in situ</i> hybridisation against Rspo1, Rspo2, Rspo3 and Rspo4 in sagittal sections of wild type pituitaries at P14. Boxed regions
are magnified, only Rspo4 is detected. Scale bars 250µm and 100µm in boxed inserts.
Figure 4 – figure supplement 1. Paracrine secretion of WNTs from SOX2+ PSCs is necessary for expansion of committed cells

A. Schematic of time points for induction by tamoxifen induction and tissue harvesting of control Sox2+/+;Wls^fl/fl and mutant Sox2^CreERT2+/+;Wls^fl/fl pituitaries.

B. Whole mount, dorsal views of control Sox2+/+;Wls^fl/fl (top panel) and mutant Sox2^CreERT2+/+;Wls^fl/fl (bottom panel) pituitaries at P21, representative of n=4 controls and n=5 mutants. Scale bars 500μm.
| Key Resources Table |
|---------------------|
| **Reagent type (species) or resource** | **Designation** | **Source or reference** | **Identifiers** | **Additional information** |
| genetic reagent *(Mus musculus)* | \(Axin^{2\text{CreERT2/}}\) | Roel Nusse, Stanford University, The Jackson Laboratory | JAX:0188 67, RRID:IM SR_JAX:0 18867 | |
| genetic reagent *(Mus musculus)* | \(Sox^{2\text{CreERT2/}}\) | (Andoniadou et al. 2013) PMID: 24094324 DOI: 10.1016/j.stem.2013.07.004 | MGI:551 2893 | |
| genetic reagent *(Mus musculus)* | \(ROSA^{26\text{mTmG/mTmG}}\) | The Jackson Laboratory | JAX:0076 76, RRID:IM SR_JAX:0 07676 | |
| genetic reagent *(Mus musculus)* | \(ROSA^{26\text{Confetti/Confetti}}\) | The Jackson Laboratory | JAX:01749 2, RRID:IMS R_JAX:01 7492 | |
| genetic reagent *(Mus musculus)* | \(ROSA^{26\text{tdTomato/tdTomato}}\) | The Jackson Laboratory | JAX:0079 09, RRID:IM SR_JAX:0 07909 | |
| genetic reagent *(Mus musculus)* | \(Ctnnb^{1\text{fl(ex2-6)/fl(ex2-6)}}\) (\(Ctnnb^{\text{LOF/LOF}}\)) | The Jackson Laboratory | JAX:0041 52, RRID:IM SR_JAX:0 04152 | |
| genetic reagent *(Mus musculus)* | \(Wls^{0/0}\) | The Jackson Laboratory | JAX:0128 88, RRID:IM SR_JAX:0 12888 | |
| genetic reagent *(Mus musculus)* | \(Sox^{2\text{gfp/+}}\) | (Ellis et al., 2004) PMID: | MGI:3589 809 | |
| genetic reagent (Mus musculus) | TCF/Lef:H2B-GFP | The Jackson Laboratory | JAX:013752, RRID:IMSR_JAX:013752 |
|--------------------------------|-----------------|------------------------|----------------------------------|
| cell line (Mus musculus)      | Primary anterior pituitary cells | This paper | N/A | Freshly isolated from Mus musculus. |
| antibody                      | Anti-GFP, (Chicken Polyclonal) | Abcam | ab13970, RRID:AB_300798 | IF(1:400) |
| antibody                      | Anti-SOX2, (Goat Polyclonal) | Immune Systems Ltd | GT15098, RRID:AB_2195800 | IF(1:200) |
| antibody                      | Anti-SOX2, (Rabbit Monoclonal) | Abcam | ab92494, RRID:AB_10585428 | IF(1:100) |
| antibody                      | Anti-SOX9, (Rabbit Monoclonal) | Abcam | ab185230, RRID:AB_2715497 | IF(1:500) |
| antibody                      | Anti-POU1F1 (PIT1), (Rabbit Monoclonal) | Gifted by Dr S. J. Rhodes (IUPUI, USA) | 422_Rhodes, RRID:AB_2722652 | IF(1:500) |
| antibody                      | Anti-SF1 (NR5A1, clone N1665), (Mouse Monoclonal) | Thermo Fisher Scientific | 434200, RRID:AB_2532209 | IF(1:300) |
| antibody                      | Anti-TBX19 (TPIT), (Rabbit Polyclonal) | Gifted by Dr J. Drouin (Montreal Clinical Research Institute, Canada) | Ac1250 #71, RRID:AB_2728662 | IF(1:200) |
| Antibody     | Description                                      | Vendor                      | RRID                        | Dilution |
|--------------|--------------------------------------------------|-----------------------------|-----------------------------|----------|
| Anti-Ki67    | Rabbit Monoclonal                                | Abcam                       | ab15580, RRID:AB_443209    | IF(1:100) |
| Anti-pH-H3   | Rabbit Polyclonal                                | Abcam                       | ab5176, RRID:AB_304763     | IF(1:500) |
| Anti-GH      | Rabbit Polyclonal                                | National Hormone and Peptide Program (NHPP) | AFP-5641801 | IF(1:1000) |
| Anti-TSH     | Rabbit Polyclonal                                | National Hormone and Peptide Program (NHPP) | AFP-1274789 | IF(1:1000) |
| Anti-PRL     | Rabbit Polyclonal                                | National Hormone and Peptide Program (NHPP) | AFP-4251091 | IF(1:1000) |
| Anti-ACTH    | Mouse Monoclonal                                 | Fitzgerald                  | 10C-CR1096M1, RRID:AB_1282437 | IF(1:400) |
| Anti-LH      | Rabbit Polyclonal                                | National Hormone and Peptide Program (NHPP) | AFP-697071P | IF(1:300) |
| Anti-FSH     | Rabbit Polyclonal                                | National Hormone and Peptide Program (NHPP) | AFP-HFS6 | IF(1:300) |
| Anti-ZO-1    | Rat Monoclonal                                   | Santa Cruz                  | SC33725, RRID:AB_628459    | IF(1:300) |
| antibody                          | Antibody Type                        | Vendor          | Catalog #                  | Concentration |
|----------------------------------|--------------------------------------|-----------------|----------------------------|---------------|
| Antibody                        | Anti-E-CADHERIN, (Rabbit Monoclonal)  | Cell Signaling  | 3195S, RRID:AB _2291471    | IF(1:300)     |
| Anti-Rabbit 488, (Goat Polyclonal) | Life Technologies                     | Life Technologies | A11008, RRID:AB _143165    | IF(1:400)     |
| Anti-Rabbit 555, (Goat Polyclonal) | Life Technologies                     | Life Technologies | A21426, RRID:AB _1500929   | IF(1:400)     |
| Anti-Rabbit 633, (Goat Polyclonal) | Life Technologies                     | Life Technologies | A21050, RRID:AB _141431    | IF(1:400)     |
| Anti-Goat 488, (Donkey Polyclonal) | Abcam                                 | Abcam           | ab150133, RRID:AB _2832252 | IF(1:400)     |
| Anti-Chicken 488, (Goat Polyclonal) | Life Technologies                     | Life Technologies | A11039, RRID:AB _142924    | IF(1:400)     |
| Anti-Chicken 647, (Goat Polyclonal) | Life Technologies                     | Life Technologies | A21449, RRID:AB _1500594   | IF(1:400)     |
| Anti-Rat 555, (Goat Polyclonal)   | Life Technologies                     | Life Technologies | A21434, RRID:AB _141733    | IF(1:400)     |
| Anti-Mouse 555, (Goat Polyclonal)  | Life Technologies                     | Life Technologies | A21426, RRID:AB _1500929   | IF(1:400)     |
| Anti-Rabbit Biotinylated, (Donkey Polyclonal) | Abcam                                 | Abcam           | ab6801, RRID:AB _954900    | IF(1:400)     |
| antibody | Anti-Rabbit Biotinylated, (Goat Polyclonal) | Abcam | ab207995 | IF(1:400) |
|----------|------------------------------------------|-------|----------|-----------|
| antibody | Anti-Mouse Biotinylated, (Goat Biotinylated) | Abcam | ab6788, RRID:AB_954885 | IF(1:400) |
| sequence-based reagent | RNAscope probe \textit{M.musculus Axin2} | Advanced Cell Diagnostics | 400331 |
| sequence-based reagent | RNAscope probe \textit{M.musculus Lef1} | Advanced Cell Diagnostics | 441861 |
| sequence-based reagent | RNAscope probe \textit{M.musculus Wls} | Advanced Cell Diagnostics | 405011 |
| sequence-based reagent | RNAscope probe \textit{M.musculus Rspo1} | Advanced Cell Diagnostics | 401991 |
| sequence-based reagent | RNascope probe \textit{M.musculus Rspo2} | Advanced Cell Diagnostics | 402001 |
| sequence-based reagent | RNascope probe \textit{M.musculus Rspo3} | Advanced Cell Diagnostics | 402011 |
| sequence-based reagent | RNascope probe \textit{M.musculus Rspo4} | Advanced Cell Diagnostics | 402021 |
| sequence-based reagent | RNascope probe \textit{M.musculus Lgr4} | Advanced Cell Diagnostics | 318321 |
| sequence-based reagent | RNAscope probe | Advanced Cell Diagnostics |  |
|------------------------|----------------|---------------------------|------|
| sequence-based reagent | RNAscope probe M. musculus Wnt9a | Advanced Cell Diagnostics | 405081 |
| sequence-based reagent | RNAscope probe M. musculus Wnt2 | Advanced Cell Diagnostics | 313601 |
| sequence-based reagent | RNAscope probe M. musculus Wnt5a | Advanced Cell Diagnostics | 316791 |
| sequence-based reagent | RNAscope probe eGFP | Advanced Cell Diagnostics | 400281 |
| sequence-based reagent | RNAscope probe M. musculus Jun | Advanced Cell Diagnostics | 453561 |
| sequence-based reagent | RNAscope probe M. musculus Axin2 (Channel 2) | Advanced Cell Diagnostics | 400331-C2 |
| sequence-based reagent | RNAscope probe M. musculus Sox2 (Channel 2) | Advanced Cell Diagnostics | 401041-C2 |
| sequence-based reagent | RNAscope probe eGFP (Channel 2) | Advanced Cell Diagnostics | 400281-C2 |
| sequence-based reagent | RNAscope probe M. musculus Sox2 | Advanced Cell Diagnostics | 401041 |
| sequence-based reagent | RNAscope probe M. musculus Pou1f1 | Advanced Cell Diagnostics | 486441 |
| sequence-based reagent | RNAscope probe Duplex Positive Control *Ppib-C1, Polr2a-C2* | Advanced Cell Diagnostics | 321641 |
|------------------------|-------------------------------------------------------------|---------------------------|-------|
| sequence-based reagent | RNAscope probe Duplex Negative Control *DapB* (both channels) | Advanced Cell Diagnostics | 320751 |
| sequence-based reagent | RNAscope probe Singleplex Positive Control *Ppib* | Advanced Cell Diagnostics | 313911 |
| sequence-based reagent | RNAscope probe: Singleplex Negative Control *DapB* | Advanced Cell Diagnostics | 310043 |
| peptide, recombinant protein | Streptavidin 488 | Life Technologies | S11223 | IF(1:400) |
| peptide, recombinant protein | Streptavidin 555 | Life Technologies | S32355 | IF(1:400) |
| peptide, recombinant protein | Streptavidin 633 | Life Technologies | S21375 | IF(1:400) |
| commercial assay or kit | RNAscope 2.5 HD Assay -RED | Advanced Cell Diagnostics | 322350 |
| commercial assay or kit | RNAscope 2.5 HD Duplex Assay | Advanced Cell Diagnostics | 322430 |
| commercial assay or kit | LIVE/DEAD Fixable Near IR-Dead Cell | Life Technologies | L34975 |
| Category                          | Item Description                        | Manufacturer       | Code   |
|----------------------------------|-----------------------------------------|--------------------|--------|
| commercial assay or kit          | FIX and PERM Cell Permeabilization Kit  | Life Technologies | GAS003 |
| chemical compound, drug          | Tamoxifen                               | Sigma              | T5648  |
| chemical compound, drug          | Corn Oil                                | Sigma              | C8267  |
| chemical compound, drug          | Collagenase Type 2                      | Worthington        | 4178   |
| chemical compound, drug          | 10X Trypsin                             | Sigma              | 59418C |
| chemical compound, drug          | Deoxyribonuclease I                     | Worthington        | LS002172 |
| chemical compound, drug          | Fungizone                               | Gibco              | 15290  |
| chemical compound, drug          | Hank’s Balanced Salt Solution (HBSS)    | Gibco              | 14170  |
| chemical compound, drug          | Fetal Bovine Serum                      | Sigma              | F2442  |
| chemical compound, drug          | HEPES                                   | Thermo Fisher      | 15630  |
| chemical compound, drug | bFGF | R&D Systems | 233-FB-025 |
|------------------------|------|-------------|------------|
| chemical compound, drug | Cholera Toxin | Sigma | C8052 |
| chemical compound, drug | DMEM-F12 | Thermo Fisher | 31330 |
| chemical compound, drug | Penicillin/Streptomycin | Gibco | 15070-063 |
| chemical compound, drug | Neutral Buffered Formalin | Sigma | HT501128 |
| chemical compound, drug | Hoechst 33342 | Thermo Fisher | H3570 | 1:1000 |
| chemical compound, drug | Declere | Sigma | D3565 |
| chemical compound, drug | Neo-Clear | Sigma | 65351-M |
| software, algorithm | FlowJo | FlowJo, LLC | https://www.flowjo.com/ | RRID:SCR_008520 |
| software, algorithm | Prism 7 | GraphPad Software | https://www.graphpad.com/ |
| software, algorithm | Image Lab | Bio-Rad Laboratories | http://www.bio-rad.com/ |
|---------------------|-----------|---------------------|------------------------|
| software, algorithm | NDP View  | Hamamatsu Photonics | https://www.hamamatsu.com/ |
| software, algorithm | HISAT v2.0.3 | (Kim, Langmead, & Salzberg, 2015) | https://github.com/infphilo/hisat2
RRID:SC R_015530 |
| software, algorithm | DESeq2 v2.11.38 | (Love, Huber, & Anders, 2014) | https://github.com/Bioconductor-mirror/DESeq2
RRID:SC R_015687 |
| software, algorithm | featureCounts v1.4.6p5 | (Liao, Smyth, & Shi, 2014) | http://subread.sourceforge.net/
RRID:SC R_012919 |
| software, algorithm | The Galaxy Platform | (Afgan et al., 2016; Blankenberg et al., 2010; Goecks, Nekrutenko, & Taylor, 2010) | https://usegalaxu.org
RRID:SC R_006281 |
| software, algorithm | Gene Set Enrichment Analysis (GSEA) | (Subramanian et al, 2005) | software.broadinstitute.org/gsea/index.jsp
RRID:SC R_003199 |
| software, algorithm | Cufflinks | (Trapnell et al., 2012) | https://github.com/ccole-trapnell-lab/cufflinks | RRID:SCR_014597 |
|---------------------|-----------|-------------------------|--------------------------------------------------|------------------|
| other               | Deposited Data, RNA-Seq | BioProject (NCBI) | PRJNA42 1806 |
A Tamoxifen  In vitro assay  Axin2<sup>CreERT2/+;Sox2<sup>EGFP/+;ROSA26<sup>tdTomato/+  

B Tomato  TAM +72h  EGFP  Tomato/EGFP/Hoechst  

C Sox2<sup>+/+;Ctnnb1<sup>LOF/LOF  P14 +22 weeks  

D Sox2<sup>CreERT2/+;Ctnnb1<sup>LOF/LOF  P14 +22 weeks  

E Wild type  Ctnnb1 deletion in SOX2<sup>+  

- SOX2  Stem cell  Committed descendant  
  Normal growth  

- SOX2  Stem cell  Committed descendant  
  Hypoplasia  

B In vitro Assay of Clonogenic Potential  

% Colony Forming Units  

Tomato+  EGFP+  Tomato+;EGFP+  

*  

D Sox2<sup>+/+;Ctnnb1<sup>LOF/LOF  

α-SOX2/α-Ki-67  

D Sox2<sup>CreERT2/+;Ctnnb1<sup>LOF/LOF  

PL  AL  

D Sox2<sup>CreERT2/+;Ctnnb1<sup>LOF/LOF  

PL  AL  

A. Axin2CreERT2/+;ROSA26mTmG/+ 

α-GFP/α-SOX2

P14 + 2 days

Distance between GFP+ cells and nearest SOX2+ cell

% of GFP+ Cells

B. P14 Sox2Egfp/+ AP

M/F

dissociation/flow sorting

EGFP Pos

EGFP Neg

RNA-Seq

C. Expression of Wls and Porcn in the pituitary at P14

Expression of Wnt genes in the pituitary at P14

D. P14 Wnt2/Sox2 Wnt5a/Sox2 Wnt9a/Sox2
A

Tamoxifen → Analysis

P14  P17  P21

B

Sox2CreERT2+/+, Sox2+/+, WtSfl/fl, WtSfl/fl

P21