Imprinted **Dlk1** dosage as a size determinant of the mammalian pituitary gland

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**Abstract**

Co-regulated genes of the Imprinted Gene Network are involved in the control of growth and body size, and imprinted gene dysfunction underlies human paediatric disorders involving the endocrine system. Imprinted genes are highly expressed in the pituitary gland, among them, **Dlk1**, a paternally expressed gene whose membrane-bound and secreted protein products can regulate proliferation and differentiation of multiple stem cell populations. Dosage of circulating DLK1 has been previously implicated in the control of growth through unknown molecular mechanisms. Here we generate a series of mouse genetic models to modify levels of **Dlk1** expression in the pituitary gland and demonstrate that the dosage of DLK1 modulates the process of stem cell commitment with lifelong impact on pituitary gland size. We establish that stem cells are a critical source of DLK1, where embryonic disruption alters proliferation in the anterior pituitary, leading to long-lasting consequences on growth hormone secretion later in life.

**Introduction**

How organ size is determined and maintained throughout life is a key question in biology. This process is complex and depends on the balance of proliferation/differentiation and apoptosis during development, during which the eventual size is attained. Following this, stem cells must repopulate the differentiated cells to maintain a stereotypical number. In some cases organ size and cell composition can change dependent upon the life-cycle stage of the organism, or in response to the environment.
The mammalian anterior pituitary gland (AP) is a master endocrine organ that integrates hypothalamic and peripheral cues to drive the release of circulating hormones. Life events such as stress, puberty and pregnancy-lactation cause remodelling of the AP to allow appropriate levels of hormone production, and changes in cell number and gland size (Perez-Castro et al. 2012). Since the stem cell capacity of SRY-related HMG-box 2 expressing (SOX2+) tissue-resident pituitary stem cells (PSCs) was demonstrated (Andoniadou et al. 2013; Rizzoti, Akiyama, and Lovell-Badge 2013), many teams have integrated knowledge of developmental signalling pathways with information about their determination and maintenance (reviewed (Russell, Lodge, and Andoniadou 2018)). Importantly, the SOX2+ compartment acts not only as a source of cellular progenitors, but also in postnatal life directs the expansion of more committed cells by the production of paracrine signals, including WNT ligands (Russell et al. 2021). Due to these recent advances, the AP is an excellent system in which to study the fundamental process of organ size determination and homeostasis.

Imprinted genes are key determinants of body size in mammals, acting during early life to modulate growth and differentiation pathways (reviewed in (Tucci et al. 2019)). They represent ~100 transcripts in mammalian species that are epigenetically regulated such that only a single parental copy is expressed and the other silenced by a mechanism utilising DNA methylation. Imprinted gene functions converge on a small number of biological processes – neurobiology, placentation and growth (reviewed in (Tucci et al. 2019)). We and others have shown that maintaining imprinted gene expression dosage within a narrow range is key to attaining the balance of growth to organ maturation ((Charalambous et al. 2012; Plagge et al. 2004; Tsai et al. 2002)). We recently noted that imprinted gene expression is enriched in the developing and postnatal pituitary gland, and we proposed that mis-regulation of gene dosage may underlie many of the endocrine features of human imprinting disorders including Silver Russell and Prader Willi Syndromes (Scagliotti et al. 2021). Importantly, a previously identified co-regulated subset of these genes, known as the imprinted gene network (IGN, (Varrault et al. 2006)), are co-expressed and represent some of the most abundant transcripts in the developing pituitary gland (Scagliotti et al. 2021).

Delta-like homologue 1 (Dlk1), a paternally-expressed imprinted gene on mouse chromosome 12 (Schmidt et al. 2000; Takada et al. 2000), is a member of the IGN. The syntenic area on human chromosome 14 is the critical region for Temple Syndrome, and loss of DLK1 expression is thought to cause phenotypes associated with this disorder; pre-and postnatal growth restriction with increased truncal obesity and precocious puberty (Ioannides et al. 2014). Alternative splicing of Dlk1 results in several protein variants that have important functional differences. Full-length Dlk1 encodes a ~60kD protein that may be cleaved by the TACE protease ADAM17 at the extracellular cell surface to release...
soluble DLK1 into the circulation. Alternative splicing events skip the ADAM17 recognition site, resulting in a transmembrane form of DLK1 which cannot be cleaved (Smas, Chen, and Sul 1997). The signalling pathway by which DLK1 acts has yet to be elucidated. Dlk1 encodes a protein with high sequence similarity to the NOTCH ligand Delta, but it lacks the DSL domain critically required for NOTCH interaction (Smas and Sul 1993).

DLK1 is expressed in several progenitor cell populations where its expression is associated with the self-renewal to differentiation transition (reviewed in (Sul 2009)). For example, addition of soluble DLK1 to cultured preadipocytes results in failure of these cells to differentiate into adipocytes whereas deletion of this gene causes premature differentiation (Smas and Sul 1993). In the context of the postnatal neurogenic niche, the expression dosage of both membrane-bound and secreted DLK1 regulates the rate of self-renewal of the SOX2+ neural stem cells (Ferron et al. 2011).

In mice, Dlk1 expression is first detected from mid-gestation in a variety of mesodermal and neuroendocrine cell types, including the developing pituitary gland (da Rocha et al. 2007). After birth Dlk1 expression is rapidly reduced, serum levels fall, and expression becomes restricted to a limited number of cell types including some neurons, adrenal and AP cells (Sul 2009). In adult humans DLK1 is expressed in Growth Hormone (GH)-producing somatotrophs (Larsen et al. 1996). We and others have shown that Dlk1 knockout mice are small with reduced GH production ((Puertas-Avendano et al. 2011; Cheung et al. 2013; Cleaton et al. 2016)). However, Gh mRNA is not reduced following a conditional ablation of Dlk1 in mature somatotrophs (Appelbe et al. 2013). We previously demonstrated that serum GH levels are elevated in mice that overexpress Dlk1 from endogenous control elements at 6 months of age, and that their GH levels fall less markedly following high-fat diet feeding (Charalambous et al. 2014). In contrast, pregnant dams lacking circulating DLK1 in pregnancy have a blunted increase in pregnancy-associated GH production (Cleaton et al. 2016). These data led us to hypothesise that DLK1 dosage modulates the size of the GH reserve at critical life stages, and prompted us to investigate the molecular mechanism further.

Here we manipulate Dlk1 gene dosage in mice using both knock out and overexpression models. We show that Dlk1 dosage regulates AP size independently of whole body weight, suggesting an autocrine/paracrine role for the product of this gene. Loss of Dlk1 function leads to reduced AP volume by acting in a discrete developmental window to shift the balance of stem cell replication/commitment, by mediating the sensitivity of progenitor cells to the WNT pathway. Finally, increased Dlk1 expression dosage increases the SOX2+ stem cell compartment, influencing organ expansion throughout the life course and increases the rate of postnatal replication in committed cells. This indicates that DLK1 may
act postnatally to mediate paracrine signals between the stem and committed cell compartment of the AP. Overall we conclude that Dlk1 dosage determines pituitary size and postnatal stem cell homeostasis.

Results

Increasing the expression dosage of Dlk1 causes pituitary volume expansion without tumorigenesis.

We previously demonstrated that adult GH was elevated in a transgenic model where extra copies of the Dlk1 gene are introduced in the context of a 70kb bacterial artificial chromosome (BAC, TG\textsuperscript{Dlk1-70C}) (Charalambous et al. 2014). This transgene drives gene expression in a similar temporal and tissue-specific pattern to the endogenous gene, but imprinting is not maintained due to lack of inclusion of the imprinting control region for the chromosome 12 cluster ((da Rocha et al. 2009), Figure 1A). In the pituitary gland at 12 weeks of age, TG\textsuperscript{Dlk1-70C} hemizygous mice (hereafter WT-TG) had increased Dlk1 mRNA expression 2-3 fold compared to wild-type (WT, Figure 1B). This resulted in elevated protein expression of all of the major isoforms of DLK1 (Figure 1C). In the adult AP, DLK1 was expressed in a subpopulation of all hormone-producing cells except corticotrophs (Figure 1D). The transgenic model did not modify the proportion of cells in the population that were DLK1+ (WT 36.5±0.9%; WT-TG 36.3±0.8%), rather, the same fraction of cells produced more protein (Figure 1C). Immunohistochemical staining for DLK1 clearly indicated that a proportion of the protein is membrane-localised (Figure 1D).

To investigate the physiological basis of increased GH production we conducted a stereological analysis of the AP at 12 weeks in WT-TG mice and their WT littermates. Pituitary size was increased following Dlk1 overexpression (Figure 1E, F), but the proportions of hormone-producing cells were not affected (Figure 1D and Supplementary File 1). We saw no evidence of pituitary tumours in any animal examined up to 1 year of age. Overall, this resulted in a 38% expansion in the somatotroph population, potentially increasing the GH-secretory reserve in WT-TG animals (Figure 1G). Thus, increasing the expression dosage of Dlk1 causes pituitary volume expansion without a shift in hormone-producing cell fate. These data suggested that Dlk1 might be acting during pituitary development to modulate AP size.

Modulation of DLK1 gene dosage in the developing pituitary gland.

In order to understand how Dlk1 regulates pituitary size we generated embryos where the gene expression is increased (WT-TG), ablated (deletion of Dlk1 from the active paternal allele, PAT) or ablated but expressing the transgene (PAT-TG) from matched litters (Figure 2A). In the whole pituitary gland at E18.5 Dlk1 splice forms expressed from the TG\textsuperscript{Dlk1-70C} transgene appeared similar in type and abundance to those expressed from the endogenous locus (Figure 2B, C).
Dlk1 is expressed from early development of the AP, from at least E9.5 in Rathke’s pouch (RP) and the overlying ventral diencephalon (Figure 2D). As development proceeds, Dlk1 levels remain high both in progenitor cells of the cleft and lineage committing cells of the parenchyma (Figure 2D, WT and WT-TG). Dlk1 expression could not be detected in PAT embryos at any stage, confirming that genomic imprinting is maintained in this tissue. Surprisingly, when expression from the TGDlk1-70C transgene was examined in the absence of the endogenous allele, we could not detect Dlk1 in the early development of the AP or midbrain. Rather, Dlk1 expression was first detected in parenchymal cells at E15.5 of PAT-TG animals. In contrast, other tissues such as cartilage demonstrated robust expression of Dlk1 in a pattern similar to that of the endogenous gene (Figure 2D). DLK1 protein expression mirrored that of the mRNA at E18.5 (Figure 2E) and at postnatal day 21 (P21) a similar pattern was observed. These data indicate that endogenous Dlk1 is expressed from the onset of pituitary development, in both progenitor and lineage committing cells, and that some regulatory sequences necessary for the full repertoire of Dlk1 gene expression are not located within the 70kb genomic region delineated by the transgene.

Dlk1 exhibits complex temporal and spatial regulation in the AP which is only partially recapitulated by the TGDlk1-70C transgene.

Since DLK1 expression in the pituitary progenitor/stem population has not previously been described in detail, we examined its temporal co-expression with the stem cell marker SOX2. In late embryogenesis and postnatally SOX2+ PSCs occupy the periluminal marginal zone (MZ) of the pituitary gland. In WT and WT-TG E18.5 embryos, DLK1 is clearly co-expressed with SOX2 in this compartment (Figure 3A). However, by postnatal day 21 (P21), DLK1+ cells are largely absent from the MZ of animals with an intact Dlk1 gene (Figure 3B). As expected, deletion of Dlk1 from the paternal allele ablates all protein expression. However, expression from the TGDlk1-70C transgene was absent from the MZ at both stages.

We recently reanalysed publicly-available single-cell sequencing (scRNA-seq) data of the mouse pituitary gland generated by the Camper lab (Cheung et al. 2018), (Cheung and Camper 2020) in order to catalogue imprinted gene expression in the early postnatal (P4) and adult (P49) gland (Scagliotti, 2021). These data confirmed our immunohistochemical analyses (Figure 1D) of enriched expression of Dlk1 in somatotrophs, lactotrophs and thyrotrophs, as well as in a subset of Sox2-expressing stem cells (Figure 3C). The scRNA-seq data also appeared to confirm the reduction in Dlk1 gene expression in the stem cell compartment between early and late postnatal stages (17.6% at P4 to 14.8% at P49, Figure 3C). Similarly, the proportion of cells expressing DLK1 in the human postnatal PSC compartment decreases from youth to adulthood as shown by analyses on human snRNA-seq pituitary datasets (Zhang et al. 2022) (14.1% in infant, 5.7% in adult and 8.9% in aged, Figure 3D). As in the
mouse, DLK1 expression in the human pituitary is most abundant in somatotrophs, lactotrophs and thyrotrophs. These three cell types are derived from a common embryonic lineage progenitor expressing the transcription factor POU1F1, and the mature hormonal lineages continue to express this transcription factor to directly regulate hormonal genes (Camper et al. 1990; Li et al. 1990). We found a high level of co-expression between DLK1 and POU1F1 in the WT and WT-TG pituitary gland at E18.5, though DLK1+ cells were also clearly visible in the MZ and in POU1F1- parenchymal cells (Figure 3E). In contrast, in the PAT-TG pituitary, DLK1 expression was confined solely to POU1F1+ cells. These data indicate that the TG^{Dlk1-70C} transgene contains cis regulatory sequences necessary for Dlk1 expression in POU1F1+ cells but not those required for expression in stem cells and other hormonal lineages (Figure 3F). To determine whether Dlk1 might be a direct target of POU1F1 we explored a POU1F1 ChIP-seq dataset from GH-expressing cells derived from rats (Skowronska-Krawczyk et al. 2014). By mapping the rat POU1F1 binding sites onto the mouse genome we identified 3 potential POU1F1 binding sites within 120kb of the Dlk1 gene (Figure 3 – Figure supplement 1). These sites overlapped other genomic features indicative of regulatory activity including evolutionary conservation, CpG islands and previously mapped histone modifications associated with enhancer activity in the embryonic brain (Gorkin et al. 2020). One of these binding sites was localised within the 70kb included in the TG^{Dlk1-70C} transgene. We suggest that Dlk1 is likely to be a direct POU1F1 target in both the WT and TG^{Dlk1-70C} transgenic mice. 

Dlk1 dosage modulation controls AP volume in two distinct developmental periods and independently of body size. We next explored the consequences of Dlk1 dosage modulation on pituitary size by performing stereological volume measurements on a time course of embryonic and postnatal pituitaries. First addressing whole organism weight, we calculated total body weight of each genotype relative to WT littermates. PAT animals were growth restricted (~85% WT weight at E18.5) in late gestation, as previously reported (Cleaton et al. 2016). WT-TG mice were not significantly altered in mass, but the TG^{Dlk1-70C} transgene rescued some of the growth deficit on a Dlk1-deleted background; PAT-TG animals weighed ~92% WT mass at E18.5 (Figure 4A, and Supplementary File 1b). Postnatally PAT animals were further growth restricted, reaching a minimum proportion of WT weight at P14 (~55%), after which the animals commenced catch-up growth, reaching 82% WT weight by weaning at P21. Next we used stereological methods to measure pituitary volume and compared relative volumes by genotype. In contrast to body weight, the volume of the AP is already significantly reduced in size at E13.5 (61% WT), prior to embryonic growth restriction. As predicted by the lack of Dlk1 expression from the TG^{Dlk1-70C} transgene at this stage (Figure 2B), the WT vs WT-TG and PAT vs PAT-TG are
indistinguishable in volume (Figure 4B, C, Supplementary File 1c, d). Subsequently, the Dlk1-ablated AP maintains this deficit in volume (50-65% WT volume) into postnatal life, with some evidence of volume catch-up at weaning. The TG<sup>Dlk1-70C</sup> transgene is unable to consistently rescue the deficit in pituitary volume that is established during embryogenesis (PAT and PAT-TG pituitary volumes are broadly overlapping). In contrast, in WT-TG animals increased Dlk1 expression from the TG<sup>Dlk1-70C</sup> transgene did cause an increase in pituitary volume. This increase in volume was evident in the second postnatal week and became statistically significant at P21 (133% WT volume, consistent with 38% increased AP mass as adults, Figure 3C, 1F). Adjacent structures, the posterior pituitary and postnatal intermediate lobe volume did not differ between genotypes (Supplementary File 1c, d) indicating that Dlk1 acts selectively on pathways that regulate AP size. Moreover, AP volume expansion only develops when elevated Dlk1 expression in the parenchyma coexists with a Dlk1+ stem cell compartment.

Finally, we determined AP volume in a cohort of adult animals with both sexes represented (Figure 4D, and Supplementary File 1e). At 8 weeks of age, WT-TG male and female animals have reached ~140% WT volume, similar to that observed in the original 12-week old female cohort (Figure 1F). Consistent with the volume catchup observed at 21D, PAT and PAT-TG AP volume was no longer significantly different from WT, except for in PAT males. Importantly, PAT-TG AP volume did not expand beyond that of the PAT, further supporting the finding that Dlk1 is required in both stem and parenchymal compartments to drive increased volume. By stereological estimation of cell number in the adult anterior lobe, we found that WT-TG AP contained a larger total number of cells (Figure 4E), indicating that the expansion occurs by hyperplasia. PAT and PAT-TG anterior lobes contained fewer cells, suggesting a long-term impact of developmental size reduction.

Somatotroph and lactotroph volume expansion has previously been observed in mice following transgenic overproduction of growth hormone releasing hormone (Ghrh), (Stefaneanu et al. 1989). To test whether pituitary hyperplasia in WT-TG was secondary to alterations in hypothalamic neuropeptide production, we measured the expression of Ghrh and somatostatin (Sst) in the adult hypothalamus (Figure 4F). Consistent with the lack of TG<sup>Dlk1-70C</sup> -derived Dlk1 expression in the midbrain during embryogenesis (Figure 2D), Dlk1 mRNA levels were similar between WT and WT-TG mice, and undetectable in both PAT and PAT-TG mice. Expression of Ghrh and Sst did not differ between the genotypes (Figure 4F).

Dlk1 dosage modulation does not prevent hormone expression but modifies GH levels and pubertal physiology.

We surveyed the expression of pituitary hormones at E18.5, when all of the hormonal axes have been established. Loss or gain of Dlk1 dosage did not cause any gross change in hormonal cell localisation.
or expression at this stage (Figure 4 – Figure supplement 1A-Y). To investigate if the increased AP volume in the WT-TG animals has any physiological consequences we measured GH levels in female mice in the peri-pubertal period. In mice, GH levels rise around weaning in the third postnatal week to reach adult levels by the fifth postnatal week (Figure 4 – Figure supplement 1Z). GH levels were elevated in young WT-TG mice and though the trend was detected later it became obscured by considerable variability (Figure 4 – Figure supplement 1Z). We could not detect a reduction in GH levels in PAT mice at any stage (data not shown). Temple Syndrome and DLK1 mutations in humans are associated with precocious menarche (Ioannides et al. 2014; Dauber et al. 2017). Using day of vaginal opening (VO) as a proxy for onset of puberty we determined that, contrary to our expectations, increased expression of Dlk1 resulted in earlier pubertal onset. However, deletion of Dlk1 did not alter the timing of puberty, but females were relatively smaller when they entered this state (Figure 4 – Figure supplement 1AA).

Dlk1 regulates embryonic pituitary size by acting in a discrete developmental window to shift the balance of stem cell replication/commitment.

Dlk1 is expressed from the onset of pituitary morphogenesis in Rathke’s pouch and the overlying ventral diencephalon (Figure 2D). Despite this early expression, we could detect no difference in the expression of transcription factors and signalling molecules that have previously been shown to regulate morphogenesis and growth of the gland at this stage (Figure 5 – Figure supplement 1). From ~E13.5, as pituitary progenitor cells exit the cell cycle and differentiate they move ventrally away from the periluminal cleft (Davis, Mortensen, and Camper 2011). Proliferating cells proximal to the lumen of the cleft are SOX2+. DLK1 expression is found in the SOX2+ proliferating cells in the cleft as well as in the newly differentiating cells (Figure 3D). Loss of Dlk1 in PAT and PAT-TG animals caused a transient reduction in cell proliferation in the cleft at E13.5 (Figure 5A-D, Supplementary File 1f). There was a concomitant increase in the proportion of cells expressing the lineage marker POU1F1 (Figure 5E, G). Several molecular signalling pathways have been proposed to regulate the emergence of the POU1F1 lineage including NOTCH (Zhu et al. 2006) and WNT pathways (Potok et al. 2008), (Olson et al. 2006). While nuclear HES1, a marker of active NOTCH signalling, was abundantly expressed in the AP at this timepoint, we did not observe a difference in the proportion of HES1+ nuclei between Dlk1 genotypes (Figure 5F, H). However, we detected increased expression of Left1 and Axin2, downstream targets of the WNT/β-catenin pathway (Hovanes et al. 2001), (Jho et al. 2002) (Figure 5l). Taken together, we propose that loss of Dlk1 in PAT and PAT-TG animals reduces AP size by altering the balance of cells remaining in the replicating stem cell pool to those that commit to the Pou1f1 lineage (Figure 5J). This balance may be shifted by the action of Dlk1 on the WNT/β-catenin pathway (Figure 5l).
Elevated DLK1 dosage causes increased proliferation of the postnatal AP only when expression is retained in the stem cell compartment.

In postnatal life, overexpression of Dlk1 from the TGDlk1-70C transgene caused an increase in AP volume, but only when WT Dlk1 expression was retained (compare WT-TG with PAT-TG, Figure 3A). These data suggested that DLK1 produced by marginal zone stem cells and parenchymal cells, might act in concert to promote postnatal proliferation. We consistently observed that WT-TG mice have elevated parenchymal proliferation at P14, the time of maximal expansion of the committed cells prior to puberty (Taniguchi et al. 2001). This increase in proliferation was not observed in the PAT-TG mice (Figure 6A-C). Further, the SOX2+ stem cell compartment is modified in WT-TG and not PAT-TG mice; WT-TG animals at weaning have an increased proportion of SOX2+ cells (Figure 6D, F) and their overall compartmental volume (Figure 6E) in both the MZ and parenchymal clusters is increased. To determine if this translated to an increase in active adult stem cells we performed a stem cell colony-forming assay in adult mice (Andoniadou et al. 2012). In this assay the AP is dissociated and plated at low density in culture media that promotes only the survival of PSCs. The number of colonies after 7 days of culture is indicative of the proportion of active stem cells in the source gland (Figure 6G). In cultures from the AP of 12 week-old WT-TG mice we observed an increase in colony forming units (CFUs) compared to WT littermates, (Figure 5H, I, J) indicating that increasing Dlk1 expression dosage increases the lifelong stem cell reserve.

Discussion

We previously demonstrated that adult TGDlk1-70C (WT-TG) mice produce more pituitary Gh mRNA, have elevated fasting GH levels and concomitant changes to whole-body lipid oxidation pathways (Charalambous et al. 2014). The increase in the GH reserve can be explained by data presented in this study, since adult TGDlk1-70C mice have a hyperplastic pituitary gland associated with a ~40% increase in the size of the somatotroph population. We did not observe an increase in linear growth in the WT-TG mice (Figure 4A). This may be explained by the timing of AP volume expansion, which occurs in the third postnatal week (Figure 4B). In mice, the majority of GH action on linear growth occurs prior to P21 (Lupu et al. 2001). While we did observe increased circulating GH in WT-TG animals at P21 (Figure 4 – Figure supplement 1), this may have been too late to promote the pre-pubertal growth spurt. Moreover, growth signalling by GH requires regulated pulsatile secretion of the hormone (Huang, Huang, and Chen 2019), generated by the homotypic somatotroph cell network (Bonnefont et al. 2005). Future work could investigate how DLK1 dosage-mediated alterations in cell number influences the integrity of this network. In contrast to a role in growth, we have observed that Dlk1 dosage is elevated during life periods when enhanced peripheral lipid oxidation is beneficial for survival; during suckling
(Charalambous et al. 2012) and in the mother during pregnancy (Cleaton et al. 2016). These are periods of life when lipogenesis may be inhibited and tissue fatty acid (FA) oxidation promoted to spare scarce glucose for growth. We speculate that imprinting may be acting on DLK1 dosage to modulate the GH axis and shift the metabolic mode of the organism toward peripheral FA oxidation and away from lipid storage. Therefore, disruption of DLK1 dosage has important consequences for energy homeostasis and metabolic disease.

**Dlk1** imprinting is maintained in several embryonic cell populations that take part in the formation of the mature anterior pituitary gland. The endogenous gene is expressed exclusively from the paternally-inherited allele in the ventral diencephalon and in developing Rathke’s pouch at E10 (Figure 2D), then in the SOX2+ progenitor populations in mid-late gestation (Figure 3A). As development proceeds, the proportion of DLK1+SOX2+ cells decreases to approximately 10% of all stem cells in the mature gland (Figure 3B-D). Concomitantly, expression of *Dlk1* initiates in committed cells of the parenchyma from their appearance at ~E13.5 (Figure 2D), and is maintained in hormone producing cells, being particularly abundant in cells of the POU1F1 lineage, somatotrophs (GH+/DLK1+ = 93% of all GH+), lactotrophs (PRL+/DLK1+ = 37% of all PRL+), and thyrotrophs (TSH+/DLK1+ = 40% of all TSH+, Supplementary File 1a). Indeed, DLK1 shares considerable overlap with POU1F1 in late gestation and may be a direct target of this transcription factor, since three experimentally validated POU1F1 binding sites are located within 120kb of the *Dlk1* gene (Figure 3 – Figure Supplement 1), (Skowronska-Krawczyk et al. 2014)). Additional experiments, such as measurement of *Dlk1* levels in *Pou1f1*-deleted mice, or targeted deletion of the putative POU1F1 binding sites at the *Dlk1* locus would be required to definitively establish this finding. *Dlk1* may be part of an autoregulatory feedback loop since, at least *in vitro*, expression of *Dlk1* represses the action of POU1F1 on the *Gh* promoter (Ansell et al. 2007). The expression pattern of *Dlk1* in the progenitor compartment mirrors that of other imprinted genes in the ‘Imprinted Growth Network, IGN’, such as *Igf2, Cdkn1c* and *Grb10* (Scagliotti et al. 2021). These genes have been suggested to comprise the hub of a core pathway that regulates embryonic growth and is progressively deactivated in the perinatal period (Lui et al. 2008; Varrault et al. 2006). The retention of *Dlk1* expression in a small proportion of SOX2+ stem cells in adults suggests heterogeneity of this population, and that a small number of cells retain an embryonic progenitor cell-like phenotype. Well powered analysis of single-cell sequencing datasets of the PSC population over the life course will be required to validate this assertion.

We observed a profound reduction in the volume of the anterior pituitary gland (40-50%) of *Dlk1*-deficient animals at E13.5, before we observed any change in overall body weight (Figure 3B). *Dlk1* is expressed at very high levels in the developing AP compared to other embryonic tissues (compare for
example the AP expression with cartilage at E15.5 in Figure 2D). The relative volume of the Dlk1-
deficient AP is maintained at ~40-50% WT levels until the third postnatal week, when there appears to
be some catch-up in volume. Importantly and in contrast to the total body weight, addition of the TG
transgene does not consistently rescue pituitary volume in Dlk1-deficient mice (Figure 3B). These
data indicate that loss of Dlk1 in the SOX2+ progenitors in the early RP causes a persistent reduction to
AP volume. Consistently we observed a transient reduction in proliferation of the RP progenitor
population at E13.5. This was associated with an increase in the size of the nascent POU1F1+
population, cells which have recently left the dorsal proliferative zone and started along the lineage
commitment pathway (Figure 5E, G). Consistently, the expression of a terminal hormonal marker of the
POU1F1 lineage, GH, was observed earlier in Dlk1-deficient animals (Figure 5J). SOX2+ progenitor cell
cycle exit is regulated by multiple signalling pathways including the NOTCH and WNT/β-catenin
pathways. HES1 is a major downstream transcriptional target of NOTCH in the developing pituitary
gland (Goto et al. 2015). Despite robust nuclear expression of HES1 in the RP at E13.5, we saw no
difference in the proportion of HES1+ cells between genotypes, suggesting that NOTCH signalling
perturbation at this stage is not responsible for pituitary volume reduction in Dlk1-deleted embryos.

Whether DLK1 is a direct inhibitor of NOTCH signalling is still unresolved, since while some have
reported biochemical interactions between DLK1 and NOTCH1 (Baladron et al. 2005), others have
failed to co-immunoprecipitate the two proteins in a biologically-relevant context (Wang et al. 2010).

We observed that the expression downstream targets of WNT/β-catenin signalling, Lef1 and Axin2,
were increased in expression domain and intensity in PATs. Exit from the proliferating progenitor pool
for commitment to the POU1F1 lineage is regulated by an interplay between the transcription factor
Prophet of PIT1 (PROP1) and β-catenin binding to and activating the POU1F1 promoter (Olson et al.
2006). Increased activation of β-catenin targets in Dlk1-depleted embryos suggests that the normal
function of DLK1 may be to act to either reduce WNT production or reduce the sensitivity of cells to the
WNT pathway.

The expression of Dlk1 in both the stem cell and mature hormone-producing compartment of the
pituitary gland is perplexing, since it suggests an interaction between DLK1 expression in the progenitor
compartment and in the niche. A precedent for this lies in the postnatal neuronal subventricular zone,
where DLK1 expression is required in both the stem cells and in niche astrocytes to maintain adult stem
cell potency (Ferron et al. 2011). Here we observed that the postnatal WT-TG AP was expanded in
volume compared to WT littermates, driven by increased proliferation of the parenchymal cells in the
second postnatal week. A similar volume expansion was not observed when PAT-TG animals were
compared to PAT littermates (Figure 4C, 6B). These data suggest that excess Dlk1 can promote
pituitary hypertrophy, but only if DLK1 is expressed in the SOX2+ compartment when dosage is
increased in the parenchyma. Intriguingly, the increase in cell division mediated by excess DLK1 at P14
is in the parenchyma, and therefore DLK1 is not acting cell autonomously in stem cells to promote
proliferation, yet SOX2+DLK1+ cells are required for volume expansion. This suggests that the
SOX2+DLK1+ cells may produce a signal that promotes parenchymal proliferation, analogous to
previous data where we demonstrated that the postnatal stem cells promote committed cell proliferation
by a WNT-mediated pathway (Russell et al. 2021). We speculate that DLK1 production from stem cells
might comprise part of this paracrine signalling system that is required for the size regulation of the
mature pituitary gland. In concert, increased expression of DLK1 in the parenchyma might alter the
sensitivity of cells to proliferative signals.

Finally, we observed that the SOX2+ stem cell compartment was expanded in the WT-TG but not PAT-
TG postnatal and adult pituitary gland. This increase in stem cell number may be a result of 'sparking' of
stem cell proliferation earlier in development, since WT-TG pituitaries have significantly fewer dividing
cells in the marginal zone at P7. Regardless, the overall effect of increased DLK1 dosage is to increase
overall AP volume and increase the size of the stem cell population.

In conclusion, we have shown that DLK1 is an important determinant of anterior pituitary size, and acts
at multiple stages of development to modulate proliferative populations. The consequences of altered
Dlk1 dosage in the pituitary affect hormone production throughout life.

Methods
Mice. All animal procedures were carried out in accordance with the recommendations provided in the
Animals (Scientific procedures) Act 1986 of the UK Government. Mice were maintained on a 12-hour
light : dark cycle in a temperature and humidity-controlled room and re-housed in clean cages weekly.
All mice were fed ad libitum and given fresh tap water daily. Mice were weaned at postnatal day (P) 21,
or a few days later if particularly small. Thereafter, they were housed in single-sex groups (5 per cage
maximum) or occasionally singly housed. Embryos were generated through timed matings (either as a
pair or a trio). Noon on the day of the vaginal plug was considered as embryonic day (E)0.5.

All mice were maintained on a C57BL6/J background. The generation of the Dlk1-knockout (Dlk1^tm1Srbpa) and
the Dlk1-TG BAC transgenic (TgDlk1-70C) lines has been previously described (Raghunandan et al.
2008; da Rocha et al. 2009). For colony maintenance maternal heterozygotes (Dlk1^tm1Srbpa+/+, hereafter
called Dlk1+/- or MAT) and littermate wild-type (Dlk1^+/+, WT) animals were generated. TgDlk1-70C (WT-TG)
animals were maintained as heterozygotes by crossing WT females with MAT males. Dlk1^tm1Srbpa+ (hereafter called
Dlk1^+/+ or PAT) and littermate WT animals or embryos were generated by
crossing WT females with MAT males. Dlk1^tm1Srbpa+/+; TgDlk1-70C (PAT-TG) animals or embryos were
obtained by crossing WT-TG females with MAT males. From these crosses, WT, PAT, and WT-TG
littermates were also generated (shown in Fig2A).
Genotyping and Dlk1 isoform analysis. Genotyping was performed on ear and embryonic tail biopsies with DNA extracted using DNAReleasy (LS02, Anachem). PCR was performed using REDTaq® ReadyMix™ PCR Reaction Mix (R2523, Merck-SIGMA) using previously published primers (Charalambous et al. 2014).

Hypothalamus isolation. Animals were culled by terminal injection of Pentobarbital (0.1mL intraperitoneal EUTHATAL, Dopharma Research B. V. the Netherlands) followed by exanguination by cardiac puncture. The brain was removed and a 4mm coronal midbrain section isolated using a 1mm adult mouse brain matrix (Alto). A section containing the hypothalamus was isolated using the corpus callosum as the dorsal boundary, and the lateral ventricles as the lateral boundaries.

RNA extraction and cDNA synthesis. RNA from pituitary glands of E18.5 embryos and adults, and adult hypothalamus was extracted using TRizol™ LS (10296028, Invitrogen™) and treated with DNase I (M0303, New England Biolabs (NEB)), following the manufacturer’s instructions. Complementary DNA (cDNA) was obtained by Reverse Transcription (RT) using 200-300 ng of purified RNA as template and Moloney Murine Leukemia Virus (M-MuLV) Reverse Transcriptase (M0253, NEB). cDNA was synthesised using the standard first strand synthesis protocol with random hexamers (S1230, NEB).

Semi-quantitative PCR Dlk1 isoform usage was determined in embryonic pituitary cDNA as described (Charalambous et al. 2014), using tuba as a loading control.

Quantitative real time PCR. RT-qPCR was performed on cDNA isolated from adult hypothalamus using the Quantinova SYBR green kit (QIAGEN #208056) on a Quantstudio 6 thermocycler (Applied Biosystems). Data was analysed using the relative standard curve method as previously described (Scagliotti et al. 2021). Actb expression was used to normalise the expression of the target genes.

Details of the primers used in this study are included in Supplementary File 1i.

Histology. Fresh embryos and postnatal pituitaries were fixed with 4% w/v paraformaldehyde (PFA, P6148, Merck-SIGMA) in Phosphate-Buffered Saline (PBS, BR0014G, Thermo Scientific Oxoid) or Neutral Buffered Formalin (Merck-SIGMA, HT501128) overnight at 4°C and dehydrated through an increasing ethanol series the following day. Samples were stored at 4°C in 70% ethanol or dehydrated to 100% ethanol the day before the paraffin embedding. On the day of embedding, samples were incubated at room temperature (RT) with Histoclear II (National Diagnostics, HS202) (2 x 20 minutes for E9.5-E11.5 and postnatal pituitaries, 2 x 35 minutes for E13.5) or Xylene (VWR) (2 x 45 minutes for E15.5, 2 x 1 hour for E18.5). This was followed by 3 x 1-hour incubations at 65°C with Histosec® (1.15161.2504, VWR). 5μm histological sections were cut using a Thermo HM325 microtome, mounted on Menzel-Gläser Superfrost®Plus slides (Thermo Scientific, J1810AMNZ) and used for in situ hybridisation (ISH), RNAscope, immunohistochemistry (IHC) and immunofluorescence (IF).

mRNA in situ hybridisation. ISH was performed as previously described (Giri et al. 2017). Sections were hybridised overnight at 65°C with sense and antisense digoxigenin (DIG)-riboprobes against Dlk1 (da Rocha et al. 2009). Sections were washed and incubated overnight at 4°C with anti-Digoxigenin-AP antibody (45-1093274910 Merck-SIGMA, 1:1000). Staining was achieved by adding a solution of 4-Nitro blue tetrazolium chloride (NBT, 11383213001, Merck-SIGMA) and 5-Bromo-4-chloro-3-indolyl phosphate disodium salt (BCIP, 11383221001, Merck-SIGMA). Sections were mounted using DPX (6522, Merck-SIGMA). Sense controls for each probe were tested at E13.5 and showed no staining under identical conditions.

RNAscope mRNA in situ hybridisation. RNAscope experiments were performed as previously described (Russell et al. 2021). For the expression analysis of early developmental markers, mRNA
expression was assessed using the RNAScope singleplex chromogenic kits (Advanced Cell Diagnostics) on NBF fixed paraffin embedded sections processed as described in the previous section. The probes used for this experiment are listed in Supplementary File 1g. ImmEdge Hydrophobic Barrier PAP Pen (H-4000, Vector Laboratories) was used to draw a barrier around section while air-drying following the first ethanol washes. Sections were counterstained with Mayer’s Haematoxylin (Vector Laboratories, H-3404), left to dry at 60°C for 30 min before mounting with VectaMount Permanent Mounting Medium (Vector Laboratories, H-5000).

Immunohistochemistry and immunofluorescence. IHC on histological sections was performed as previously described (Giri et al. 2017). For IHC, unmasking was achieved by boiling the histological sections with 10 mM tri-sodium citrate buffer pH 6 for 20 minutes. Detection of the proteins was achieved by incubating the histological sections overnight at 4 °C with the primary antibodies described in Supplementary File 1h, and detected with secondary biotinylated goat α-rabbit, goat α-mouse or α -goat secondary (BA-1000, BA-9200 and BA9500, Vector Laboratories, 1:300), followed by 1-hour incubation at room temperature with Vectastain® Elite ABC-HRP kit (PK-6100, Vector Laboratories). Staining was achieved through colorimetric reaction using DAB Peroxidase Substrate Kit (SK-4100, Vector Laboratories). Slides were lightly counterstained with Mayer’s Haematoxylin (MHS16, Merck-SIGMA) and mounted using DPX. For double IF with hormonal markers an adult sections, unmasking was achieved as described above. For double IF on embryo sections, unmasking was performed with Tris-EDTA buffer pH 9 [10 mM Tris Base, 1 mM EDTA, 0.05% Tween 20]. Histological sections were incubated overnight at 4°C with the primary antibody, incubated sequentially with a goat biotinylated anti-rabbit (1:200) then with streptavidin-594 (SA5594, Vector Laboratories, 1:200) at RT. The second primary was added to the sections overnight at 4°C then detected with a goat α-rabbit DyLight® 488 (DI1488, Vector Laboratories, 1:200) and mounted using VECTASHIELD® Antifade Mounting Medium with DAPI (H-1200-10, Vector Laboratories). Isotype controls for each antibody showed no staining under identical conditions.

Stereological estimation of pituitary volumes and cell number. Pituitary volume was estimated using the Cavallieri method (Howard and Reed 1998). Briefly, the pituitary was exhaustively sectioned then sections were collected at regular, non-overlapping intervals throughout the gland from a random starting point and stained with H&E. Images of each H&E-stained section were acquired using a NanoZoomer HT (Hamamatsu) and processed using NDP.view2 software (Hamamatsu) to calculate the pituitary cross sectional area (CSA) at least 20 times per pituitary, which was then converted to volume. Samples were blinded prior to counting. Estimating total cell number: For each animal the maximal CSA was determined from the volumetric analysis above. At this level 3 independent sections were used to measure anterior pituitary CSA and count haematoxylin-stained nuclei (using the cell detection tool in QuPath v0.3.2 (Bankhead et al. 2017), standard Hematoxylin OD settings except with threshold intensity set to 0.05), giving a mean cells/CSA measurement per individual. This number was multiplied by the AP volume to give the total cell number.

Cell counting. Histological sections were immunostained as described above and imaged using the NanoZoomer HT (Hamamatsu). Images were exported using NDP.view2 and cell counted using the cell counter plugin in ImageJ (Schindelin et al. 2012). For IHC, cells that exhibited a clear brown staining were manually counted as positive. Samples were blinded prior to counting. Total number of cells was calculated by counting haematoxylin-counterstained cells. For IF, cells that exhibited a clear fluorescence signal were manually counted as positive. Total number of cells was calculated by counting DAPI-counterstained cells.
Imaging. Images of histological sections were acquired using a NanoZoomer HT (Hamamatsu). Fluorescence images and higher magnification bright-field images were acquired using a Zeiss Axioplan II microscope with a Luminera 3 digital camera and INFINITY ANALYSE software v6.5.6 (Luminera). Images were combined using Adobe Photoshop 23.4.1 release 2022 (Adobe).

Analyses of single cell and single nuclei RNA-seq data. Single-cell sequencing data from the mouse pituitary gland was downloaded from GEO using accession numbers GSE120410 (postnatal day 4, P4) and GSE142074 (postnatal day 49, P49) and analysed as previously described (Scagliotti et al. 2021). Human single-nuclei RNA sequencing pituitary data was obtained from GEO accession number GSE178454. Using Seurat (v4.1.0) (Satija et al. 2015), (Macosko et al. 2015), (Stuart et al. 2019), (Hao et al. 2021) in R, cells from male and female datasets of all ages were taken forward if they expressed between 1000–5500 genes and <10% mitochondrial transcripts, removing doublets and low-quality cells. All 6 filtered datasets were integrated using the SCTransform workflow (Hafemeister and Satija 2019). Clustering and visualisation for the integrated objects were carried out using the the default resolution and 1:15 principal components. Clusters were named according to known cell-type markers as previously reported (Cheung et al. 2018), (Zhang et al. 2022), (Scagliotti et al. 2021).

For both mouse and human data, Dlk1/DLK1 expression was plotted in the respective datasets using the “FeaturePlot” function in Seurat with a min.cutoff = 0 and split.by = ‘Age’. Percentage of stem cells expressing Dlk1/DLK1 was calculated under the ‘integrated’ assay of the Seurat object using the WhichCells function of grouped ‘Stem Cells’ expressing Dlk1/DLK1.

Mapping POU1F1 binding sites to the mouse Dlk1 region. POU1F1 binding sites identified using chromatin immunoprecipitation followed by sequencing (ChIP-seq) in growth hormone (GH)-expressing rat pituitary cell line were obtained from a published dataset (Skowronska-Krawczyk et al. 2014). POU1F1 binding sites overlapping the rat Dlk1 locus (chr6: 133828590-134274323, rn4 assembly) were selected and converted to the orthologues region on mouse chr12 (mm10 assembly) using University of California Santa Cruz Genomics Institute Genome Browser (UCSC). The reported POU1F1 binding sites displayed a rat/mouse sequence conservation higher than 98%. Data were visualised using Gviz R package (Hahne and Ivanek 2016).

Western blotting. Sample preparation, gel electrophoresis and western blotting were described previously (da Rocha et al. 2009), with Abcam anti-DLK1 (ab21682, which recognises the intracellular domain of the protein) at 1:500. Anti-alpha tubulin (Merck-SIGMA, clone B-5-1-2) was used as a loading control at 1:10000.

Pituitary SC cultures. Pituitary stem cells were isolated from anterior pituitaries collected from adult female mice at 13-16 weeks of age, essentially as described in Andoniadou, 2013 (Andoniadou et al. 2013). Briefly, mice were culled by CO2 asphyxiation, the skin overlying the skull cut and the skull bone opened with scissors. Using a dissecting microscope, the brain was carefully removed, as well as the posterior lobe of the pituitary gland. The remaining anterior pituitary was gently detached and transferred to a 1.5 ml microcentrifuge tube containing 200 µl of enzyme mix [0.5% w/v Collagenase type II (Worthington), 50 µg/ml DNase (Worthington), 2.5 µg/ml Fungizone (Gibco) and 0.1% v/v trypsin-EDTA solution 0.05% (Gibco) in Hank’s Balanced Salt Solution (HBSS, Gibco)]. Once transferred, the anterior pituitary was incubated at 37°C for up to 4 hours, under gentle agitation. Following the incubation, samples were mechanically dissociated into single cell suspensions by vigorously pipetting the solution up and down. 1 ml of HBSS was added to the solution to dilute the
enzyme mix and samples were centrifuged at 2500 rpm for 5 minutes. The cell pellet was re-suspended in Ultraculture Medium (Lonza), supplemented with 5% v/v fetal bovine serum (FBS, Invitrogen), 1% v/v penicillin/streptomycin (P/S, Merck-SIGMA), 20 ng/ml basic fibroblast growth factor (bFGF, R&D) and 50 ng/ml Cholera Toxin (Merck-SIGMA). Cells were counted using a haemocytometer, upon incubation with an equal volume of Trypan blue solution 0.4% (Merck-SIGMA) to assess cell viability. Viable cells were then plated at a density of 1000 cells/cm² and fresh media replaced every three days.

**Plasma GH determination.** ELISA kits were used for measurements Growth Hormone (Merck-SIGMA EZRMGH-45K) according to manufacturers’ instructions.

**Timing of puberty.** Day of puberty was determined as the time of first vaginal opening (VO) in postnatal female mice. Animals were checked daily in the morning from P18 to P35 and the first appearance of VO and body weight on that day recorded. For this experiment all mice were weaned at P21.

**Statistical analysis.** All statistical tests were performed using the GraphPad Prism Software version 9 for Windows, GraphPad Software, San Diego California USA, www.graphpad.com. Specific tests, significance values and number of samples analysed are indicated in the respective figure/table legends, and all error bars represent the standard deviation (SD). Data points in graphs represent individual animals as biological replicates. Outliers were removed using Grubb’s test within the Prism application.
Figure Legends

**Figure 1.** Increasing the expression dosage of **Dlk1** causes pituitary hyperplasia. (A) Schematic of part of the imprinted region on mouse chromosome 12 that contains the **Dlk1** and **Meg3** genes. **Dlk1** is expressed (blue box) from the paternally-inherited chromosome, **Meg3** from the maternally-inherited chromosome (red box). This expression pattern is established and maintained by an intergenic differentially methylated region (IG-DMR) which is paternally methylated (black oval) and maternally unmethylated (white oval). The TG<sup>Dlk1-70C</sup> transgene used in this study contains the whole **Dlk1** gene within ~70kb of surrounding DNA, but does not contain the IG-DMR and is not imprinted. (B) RT-qPCR for **Dlk1** in whole pituitary from 12-week old female hemizygous TG<sup>Dlk1-70C</sup> transgenic (WT-TG) mice and wild-type (WT) littermates. **Dlk1** expression is 2.7x higher in transgenic animals, **p < 0.01** compared by Mann-Whitney U test, n = 7 per genotype. Bar shows mean +/- SD. (C) Western blots of extracts from whole pituitary of 12wk female WT and WT-TG mice. Alpha tubulin (TUB) is used as a loading control. Full-length and membrane bound isoforms at 50-60kDa. (D) Fluorescence immunohistochemistry for DLK1 and pituitary hormones at 12wk. DLK1 in the adult AP is detected in all hormone-producing cells except corticotrophs; GH = somatotrophs, PRL = lactotrophs, TSH = thyrotrophs, FSH = gonadotrophs; ACTH = corticotrophs. Scale bar = 20um. Note some membrane localisation of DLK1. White arrows indicate co-expression. Proportion of hormone-labelled cells was quantified between mice of each genotype and shown as a pie chart on the right, UC = unclassified cell type. (E) Whole pituitary glands from adult WT-TG mice appear larger than those from WT littermates. Light field image of 12wk female glands, scale bar = 1mm. (F) Anterior pituitary volume, but not intermediate lobe or posterior pituitary volume is increased in WT-TG animals compared to WT littermates. 12-week old females, n = 12/13 animals per genotype, compared by 2-Way ANOVA with Sidak’s post-hoc multiple comparison test, *** p<0.001. (G) Overall cell proportion was not changed but absolute volume of hormone-producing cells was altered by **Dlk1** dosage (data shown in Supplementary File 1a). Groups are compared by 2-Way ANOVA and differ significantly according to genotype (p = 0.0009); genotypes are compared for each cell type Sidak’s multiple comparison test – WT and WT-TG animals have a significantly different proportion of GH-producing cells (**p < 0.001). F and G bars show mean values +/- SD.

**Figure 2.** **Dlk1** imprinting and expression in the developing pituitary gland from the endogenous locus and TG<sup>Dlk1-70C</sup>transgene. (A) Cross used to generate embryos and postnatal animals in the study. Males inheriting the deleted allele from the mother (maternal **Dlk1<sup>tm1Srbpa/+</sup>** heterozygotes or MATs) were crossed to females hemizygous for the TG<sup>Dlk1-70C</sup> transgene (WT-TG), generating 4 genotypes, WT, WT-TG, paternal **Dlk1**+/<sup>tm1Srbpa</sup> heterozygotes (PATs) and mice inheriting a deleted paternal allele and the transgene (PAT-TG). (B) Schematic showing the known splice variants of **Dlk1**, A-D. Splicing occurs internally in exon 5 of the **Dlk1** gene. **Dlk1**-A and B retain an extracellular cleavage domain (TACE), in **Dlk1**-C and D this region is spliced out. All versions contain a single pass transmembrane domain (TM). Red arrows indicate location of primers used in (C). (C) Semi-quantitative PCR on embryonic day (E)18.5 whole pituitary glands from the 4 genotypes shown in (A). Top – primers amplify the exon 4-5 region of **Dlk1** and can distinguish splice variants based on size. Bottom – alpha-tubulin (**Tuba1a**) was amplified as a loading control on each sample. (D) In-situ hybridisation for **Dlk1** in the developing pituitary gland from E9.5 to E18.5 in the 4 genotypes shown in (A). **Dlk1** expression is indicated by purple staining. Scale bars show 100um (E9.5 and E13.5, sagittal sections) and 200um (E15.5 and E18.5, frontal sections). (E) Immunohistochemistry (IHC) for **Dlk1** on frontal sections at E18.5 and postnatal day 21 (P21), counterstained with DAPI. Scale bars = 50um.
Figure 3. DLK1 expression in the pituitary gland is dynamically regulated in two distinct compartments, only one of which is recapitulated by the TGDlk1-70C transgene. (A) IF of DLK1 and SOX2 in the E18.5 pituitary. SOX2 expression is high in the epithelial cells lining the pituitary lumen, as expected. In WT and WT-TG mice there is co-expression of DLK1 and SOX2 (inset), as well as high levels of DLK1 expression in the SOX2-negative parenchyma. In the PAT-TG pituitary DLK1 expression is not detected in the SOX2-positive compartment. Scale bars = 25um. (B) IF of DLK1 and SOX2 in the P21 pituitary. The majority of DLK1 expression is outside the SOX2-positive compartment of all genotypes. Scale bars = 25um. (C) UMAP plot illustrating expression of Dlk1 in sc-RNAseq at P4 (Cheung and Camper 2020) and P49 mice (Cheung et al. 2018) in the postnatal mouse pituitary, indicating high expression in the POU1F1 lineage (S somatotrophs, T thyrotrophs, L lactotrophs), with additional expression in a subset of Sox2-positive stem cells (SC), outlined by red box. EC endothelial cells, IC immune cells, CT connective tissue, G gonadotrophs, SF1 steroidal factor 1 progenitors, C corticotrophs, M melanotrophs, PC proliferating cells, PP posterior pituitary. (D) UMAP plot illustrating expression of DLK1 in sn-RNAseq from infant, adult and aged human pituitary gland (Zhang et al. 2022), indicating high expression in the POU1F1 lineage (S somatotrophs, T thyrotrophs, L lactotrophs), with additional expression in a subset of SOX2-positive stem cells (SC). Additional cell labels as in (C). (E) IF of DLK1 and POU1F in the E18.5 pituitary. DLK1 and POU1F1 are highly co-expressed in all Dlk1-expressing genotypes (white arrows). In WT and WT-TG there is additional DLK1 expression in marginal zone cells (orange arrow) and parenchymal cells (yellow arrow) that are POU1F1-negative. This expression is absent from the PAT-TG. Scale bar = 100um, inset 25um. * indicates background autofluorescence from blood cells. (F) Summary of Dlk1 expression across embryonic and early postnatal development in the mouse, with contribution from the transgene.

Figure 4. DLK1 dosage during embryogenesis and early life modulates anterior pituitary volume independent of whole animal mass. (A) Time-course of proportional body weight by genotype from E13.5 to P21, derived from data in Supplementary File 1b, sexes are combined since body weight is not sexually dimorphic at these stages. Lines and points show mean proportional weight of embryo/pup for WT-TG, PAT and PAT-TG animals compared to WT littermates (dotted line at 1.0). The unmodified weight data was compared using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (*p<0.05, **p<0.001), and PAT with PAT-TG (## p<0.01). (B) Haemotoxylin and eosin-stained pituitary glands at E13.5 (sagittal view), E18.5, P7 and P21 (frontal view) across the 4 genotypes. Scale bars show 100um (E13.5), 200um (E18.5, P7) and 300um (P14, P21). (C) Time-course of proportional pituitary volume by genotype from E13.5 to P21, derived from data in Supplementary File 1c, d. Lines and points show mean volume of the anterior pituitary gland for WT-TG, PAT and PAT-TG animals as a proportion of WT littermates (dotted line at 1.0). Genotypes were compared in unmodified volume data using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (*p<0.05, **p<0.01, ***p<0.001), and PAT with PAT-TG (## p<0.01). (D) Anterior pituitary volumes of male (triangles, M) and female (circles, F) mice at postnatal week 8. Volumes were compared by sex and genotype with a 2-Way ANOVA and within genotypes using a Dunnett’s multiple comparison test. Volumes were significantly different by genotype (p< 0.0001), and p values of significant post-hoc comparisons are shown on the graph. (E) Estimated total cell numbers of APs from D. Sexes were combined and genotypes compared using One-Way ANOVA (p < 0.0001 overall). Each genotype was compared to WT using Dunnett’s multiple comparison test and p values shown on the graph. D, E Source data shown in Supplementary File 1e. (F) RT-qPCR quantitation of gene expression in the hypothalamus at 8 weeks in samples from D, normalised to actin. For each gene, expression levels were compared between genotypes as in E, post-hoc p values are shown on the graph. (D-F) Bars show the mean of the data +/- SD.
Figure 5. **Loss of Dlk1 expression in an embryonic time window shifts progenitor cells from proliferation to differentiation.** (A) Proportion of phospho-histone H3 (pHH3)-positive cells to total cells in the pituitary at E11.5. (B) Proportion of pHH3-positive cells in the morphological stem cell compartment (cleft/marginal zone (MZ)) at E13.5 and E18.5 across the 4 Dlk1 genotypes. (C) Proportion of pHH3-positive cells in the parenchymal region across the 4 Dlk1 genotypes. (B) and (C) for each time point genotypes were compared using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (*p<0.05, **p<0.01), and PAT with PAT-TG. (D) Representative image showing increased proliferation in the SOX2+ stem cell compartment of Dlk1-expressing (WT and WT-TG) embryos at E13.5 compared to non-expressing embryos (PAT and PAT-TG). (E) Representative image showing increased number of POU1F1 cells newly differentiating in the parenchyma of the Dlk1 non-expressing E13.5 pituitary (PAT and PAT-TG), compared with the Dlk1-expressing (WT and WT-TG) gland. (F) Representative image showing no difference in the number of cells expressing nuclear HES1 in Dlk1-expressing (WT and WT-TG) embryos at E13.5 compared to non-expressing embryos (PAT and PAT-TG). D-F scale bar = 50um. (G) Proportion of POU1F1-positive cells in the parenchymal region at E13.5 across the 4 Dlk1 genotypes. Genotypes were compared using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (**p<0.01). (H) Proportion of nuclear HES1-positive cells in the E13.5 pituitary of the 4 Dlk1 genotypes. Genotypes were compared using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (none significant). (I) Left: schematic showing how Dlk1 might interact with the WNT signalling pathway which promotes commitment towards the Pou1f1 lineage. Right: RNA scope in-situ hybridisation for Lef1 and Axin2 in E13.5 pituitary from WT and PAT littermates. Lef1 and Axin2 expression is indicated by pink staining, counterstained in blue (haematoxylin). Scale bar = 100um. (J) Representative IF for GH at e15.5 in WT and PAT pituitary (coronal section). A small number of GH-positive cells were detected in 6/6 PAT compared to 0/4 WT animals. Scale bar = 100um or 50um (inset). (K) Schematic summarising the impact of loss of Dlk1 expression in the embryonic pituitary gland.

Figure 6. **Increased proliferation and stem cell number occurs when Dlk1 dosage is elevated in both stem and parenchymal cells.** (A) Proportion of phospho-histone H3 (p-HH3)-positive cells in the marginal zone (MZ) of the anterior pituitary at postnatal days 7, 14 and 21. (B) Proportion of phospho-histone H3 (p-HH3)-positive cells in the parenchymal zone (MZ) of the anterior pituitary at postnatal days 7, 14 and 21. (A) and (B) for each time point genotypes were compared using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (**p<0.01). (C) Representative images of immunohistochemistry for p-HH3 at P14 (brown staining, blue counterstain with haematoxylin), showing increased proportion of positive staining in the parenchyma of the WT-TG anterior pituitary. Scale bars = 250um. (D) Proportion of SOX2-positive cells in the anterior pituitary at postnatal day 21. Genotypes were compared using a One-Way ANOVA with Bonferroni’s posthoc test comparing each genotype with WT (*p<0.05). (E) Volumes of P21 anterior pituitaries subdivided into categories dependent on location and SOX2 staining. Genotypes were compared using a 2-Way ANOVA and differ significantly according to genotype (p < 0.0002), category (p <0.0001) and the interaction between them (p<0.0001). Genotypes are compared in each category to WT using Dunnett’s multiple comparison test. WT-TG animals have a significantly increased cell volume in all categories, whereas PAT and PAT-TG animals differ only in parenchymal cell volume (*p < 0.05). Bars show mean +/-SD, n = 4-8 animals per genotype. (F) Representative images of immunohistochemistry for SOX2 at P21 (brown staining, blue counterstain with haematoxylin), showing increased proportion of positive staining in the WT-TG anterior pituitary. Scale bars = 250um. (G) Methodology for determining
the number of tissue-resident stem cells in the AP. Whole pituitary is dissected and the posterior lobe
removed. The tissue is enzymatically dissociated and counted. A fixed number of cells are seeded at
low density into culture media that promotes stem cell growth. After 7 days colonies are counted. Each
colony represents a stem cell from the original organ. (H) Image of pituitary stem cell cultures stained
after growth for 7 days, derived from WT (top) and WT-TG (bottom) adult animals. Black dots in the
plate indicate colonies derived from a single colony forming unit (CFUs). (I) Number of CFUs from
pituitary stem cells following 7 days of culture on 4 separate occasions from a total of n = 6 WT and n =
9 WT-TG, 13-16wk adult animals. (J) Area of CFUs from the stem cell cultures in (I). Genotypes were
compared using a Student’s t-test, *p<0.05. (K) Schematic summarising the action of Dlk1 in the
different compartments of the AP.

Supplementary Figure Legends

**Figure 3 – Figure supplement 1.** Dlk1 may be a direct transcriptional POU1F1 target. Schematic
showing relative genomic positions of sequences in the TGDlk1-70C transgene and experimentally-verified
binding sites for POU1F1. 3 predicted POU1F1 binding sites are within 120kb of the Dlk1 promoter,
each within highly conserved regions of the genome and overlapping genomic sites enriched for
enhancer-specific histone marks in embryonic forebrain. Only one POU1F1 binding site is predicted to
be present in the transgene, mapping ~10kb upstream of the Dlk1 promoter.

**Figure 4 - Figure supplement 1.** Dlk1 dosage modulates hormone levels and early life endocrine
physiology. (A-Y) Representative images of IHC showing hormone expression at e18.5 of ACTH (A-D,
A'-D'), GH (E-H, E'-H'), LH (I-L, I'-L'), POU1F1 (M-P, M'-P'), PRL (Q-T, Q'-T'), TSH (U-Y, U'-Y') with 2-3
animals per genotype examined. Marker expression is stained in brown with haemotoxylin
counterstaining in blue. Scale bar = 200um, boxed = 100um. (Z) Circulating GH measured in plasma
from WT-TG mice and their WT littermates at postnatal day 21 (P21, n = 9/10), P28 (n = 7/9) and P35
(n = 10/11). Genotypes had significantly different levels of GH (Two-Way ANOVA p = 0.036 by
genotype) with Bonferroni post-hoc testing at each time point **p<0.01. (AA) Day of detection of vaginal
opening (VO) in female mice (left) and body weight on that day (right) in WT (n=8-10), WT-TG (n = 8)
and PAT (n = 6) mice. WT-TG entered puberty by this measure earlier than WT littermates. PAT mice
weighed significantly less than WT when VO was detected. Genotypes were compared using a One-
way ANOVA with Dunn’s multiple comparison test, *p<0.05, **p<0.01. Bars show mean and SD.

**Figure 5 – Figure supplement 1.** Altered dosage of Dlk1 does not affect expression of early
morphological signalling genes in Rathke’s pouch. Representative images of mRNA in situ
hybridisation in E9.5 wild type (WT) and Dlk1-paternal heterozygote (PAT) embryos using specific
probes against Bmp4 (A, A'; F, F'), Shh (B, B'; G, G'), Fgf8 (C, C'; H, H'), Fgf10 (D, D'; I, I'), all red with
blue haemotoxylin counterstaining, and Dlk1 (E, E'; J, J') purple, no counterstaining. At least 3 animals
per genotype were examined. Scale bar = 200um for main images and 50um for boxed images.

Supplementary File 1 Legends.

**Supplement File 1a.** Proportions of AP cells labelled with hormonal markers in WT and WT-TG female
animals at 12 weeks of age.
**Supplementary File 1b.** Total body mass of animals from matched litters sacrificed from E11.5 to P21. Individuals in each age group were compared by One-Way ANOVA with post-hoc pairwise testing WT vs WT-TG, PAT, PAT-TG and PAT vs PAT-TG, corrected for multiple comparisons using Bonferroni's adjustment.

**Supplementary File 1c.** Pituitary volumes acquired by stereological estimation in the embryo. Data from males and females is combined.

**Supplementary File 1d.** Pituitary volumes acquired by stereological estimation in of the intact postnatal gland. Individuals in each age group were compared by One-Way ANOVA with post-hoc pairwise testing WT vs WT-TG, PAT, PAT-TG and PAT vs PAT-TG, corrected for multiple comparisons using Bonferroni's adjustment. Data from males and females is combined.

**Supplementary File 1e.** Pituitary volumes acquired by stereological estimation in of the intact adult gland. All animals were compared by Two-Way ANOVA with post-hoc pairwise testing WT vs WT-TG, PAT, PAT-TG and PAT vs PAT-TG, using Dunnett's multiple comparison test.

**Supplementary File 1f.** Proportion of proliferating cells (IHC positive for p-HH3) in the embryonic and postnatal pituitary gland. Data from males and females is combined.

**Supplementary File 1g.** RNAscope probes used in this study

**Supplementary File 1h.** Primary antibodies used in the study.

**Supplementary File 1i.** RT-PCR and RT-qPCR primers used in this study

**Legends for Source data**

**Source data for Figure 1**

**Figure 1-source data 1.** Zipped Excel file containing raw data used to generate graphs in Figure 1.

**Figure 1B.** RT-qPCR data for Dlk1 expression from whole pituitary gland at 12 weeks, females. Normalised to mean WT = 1, with B-actin as housekeeping gene.

**Figure 1D&G.** Raw and proportional volumes of adult pituitary cell types in WT and WT-TG.

**Figure 1F.** Pituitary volumes in WT and WT-TG adult pituitary glands.

**Figure 1-source data 2.** Zipped file containing source data for Figure 1C – gels with cropped bands highlighted and original gel images.

**Source data for Figure 2**

**Figure 2-source data 1.** Zipped file containing the source data for Figure 2D -gels with cropped areas highlighted and original gel images.

**Source data for Figure 4**

**Figure 4-source data 1.** Zipped file containing raw data used to generate graphs in Figure 4.

**Figure4A_TableS2_raw weights.** Weight data from all animals in the developmental study.

**Figure4A_proportions.** Body weight as a proportion of time-matched WT weight.

**Figure4C_pituitary_volumes.** Raw pituitary volumes generated by stereological measurement for animals in the developmental study.
Figure 4D_E_volume_number. Raw pituitary volumes and cell number for adult animals generated by stereological measurement.

Figure 4F_RTqPCR. RT-qPCR data for Dlk1 expression from hypothalamus at 8 weeks, females and males. Normalised to B-actin as a housekeeping gene.

Source data for Figure 5 and 6

Figure 5&6-source data 1. Zipped file containing raw data used to generate graphs in Figure 5 and Figure 6.

Fig 5ABC&FigAB_pHH3. Cell counts of pH3+ cells in developmental time course.

Fig 5G_POU1F1. Cell counts of POU1F1+ cells at e13.5.

FigSH_HES1. Cell counts of HES1+ cells at e13.5.

Fig6DE_SOX2. Cell counts for %SOX2+ cells and SOX2+ compartmental volume at postnatal day 21.

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Competing interests

We have no competing interests to declare.
References


A) Schematic representation of different genotypes:

- **Dlk1<sup>−/−</sup>** (MAT)
- **Dlk1<sup>−/−;</sup>Tg<sub>Dlk1-70C</sub>** (WT-TG)
- **Dlk1<sup>+/+</sup>** (WT)
- **Dlk1<sup>+/−</sup>** (PAT)
- **Dlk1<sup>+/−;</sup>Tg<sub>Dlk1-70C</sub>** (PAT-TG)

B) Exon structures with splicing variants:

- Exon 4
- Exon 5: TACE TM

- **Dlk1-A**
- **Dlk1-B**
- **Dlk1-C**
- **Dlk1-D**

C) Expression analysis at E18.5:

- **WT**
- **WT-TG**
- **PAT**
- **PAT-TG**

D) H&E staining at different time points:

- **E9.5-E10**
- **E13.5**
- **E15.5**
- **E18.5**

E) Immunostaining for DLK1 and DAPI at E18.5:

- **WT**
- **WT-TG**
- **PAT**
- **PAT-TG**

**Legend:**

- **Dlk1**
- **WT**
- **WT-TG**
- **PAT**
- **PAT-TG**
- **C-**
- **Exon4**
- **Exon5**
- **TACE TM**
- **DLK1 DAPI**
- **Tuba1a**
Genomic location

70C-BAC transgene

Dlk1 gene

POU1F1 ChIP

CpG islands

ENCODE regulatory activity embryonic brain

ENCODE key

- Distal enhancer active
- Enhancer poised
- Heterochromatin polycomb associated
- Transcription permissive

Supplementary Figure 1
Plasma GH

Postnatal days

Day of VO detected

Body weight at VO detected

**Z**

Plasma GH

0
5
10
15

Postnatal days

WT

WT-TG

PAT

PAT-TG

Body weight (g)

AA

Day of VO

Body weight at VO

WT

WT-TG

PAT

WT

WT-TG

PAT
A) Proliferation MZ

B) Proliferation parenchyma

C) P14 p-HH3

D) P21 SOX2+ cells

E) P21 pituitary volume

F) P21 SOX2

G) Dissociate AP

H) Plate at low density in stem cell medium

I) Colony forming units

J) Colony size (arb)

K) Stem cells

Committed cells

DLK1-high

DLK1+ve

DLK1-ve