Rx3 and Shh direct anisotropic growth and specification in the zebrafish tuberal/anterior hypothalamus

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Abstract

In the developing brain, growth and differentiation are intimately linked. Here we show that in the zebrafish embryo, the homeodomain transcription factor rx3 co-ordinates these processes to build the tuberal/anterior hypothalamus. Analysis of \textit{rx3} \textit{chk} mutant/\textit{rx3} morphant fish and EdU pulse-chase studies reveal that \textit{rx3} is required to select tuberal/anterior hypothalamic progenitors and to orchestrate their anisotropic growth. In the absence of \textit{rx3} function, progenitors accumulate in the 3\textsuperscript{rd} ventricular wall, die or are inappropriately-specified, the \textit{shh+} anterior recess does not form, and its resident \textit{pomc+}, \textit{ff1b+} and \textit{ otp+ TH+} cells fail to differentiate. Manipulation of Shh signalling shows that \textit{shh} co-ordinates progenitor cell selection and behaviour by acting as an on-off switch for \textit{rx3}. Together our studies show that \textit{shh} and \textit{rx3} govern formation of a distinct progenitor domain that elaborates pattern through its anisotropic growth and differentiation.
Introduction

The hypothalamus is an ancient part of the ventral forebrain. It centrally regulates homeostatic processes that are essential to survival and species propagation, including autonomic regulation of energy balance, growth, stress and reproduction. Such adaptive functions are dependent upon the integrated function of evolutionarily-conserved neurons (reviewed in Bedont et al., 2015; Biran et al., 2015, Burbridge et al., 2016; Lohr and Hammerschmidt, 2011; Machluf et al., 2011; Pearson and Placzek, 2013; Puelles et al., 2012) that in mouse, are located within defined nuclei, including the Arc and VMN of the tuberal hypothalamus, and the PVN of the anterior hypothalamus. In zebrafish, functionally-analogous neurons exist in the periventricular tuberal (pevTub) hypothalamus and the neurosecretory preoptic (NPO) area (Biran et al., 2015; Herget et al., 2014; see Materials and Methods and Discussion for terminology).

Many transcription factors and signalling ligands that govern differentiation of hypothalamic neurons from progenitor cells, likewise, have been largely conserved (reviewed in Bedont et al., 2015; Biran et al, 2015; Burbridge et al., 2016; Pearson and Placzek 2013). The mechanisms through which secreted signalling ligands and transcription factors define and build hypothalamic territories and cells remain enigmatic (see Bedont et al, 2015; Puelles et al., 2012; Pearson and Placzek 2013). Models based on the uniform growth and differentiation of patterned territories do not account for the complex spatial patterns, or the protracted period of neuronal differentiation in the hypothalamus and at present, little is known about how early patterning events are elaborated over time. In the hypothalamus, distinct neural progenitor domains that form around the 3rd (diencephalic) ventricle (3V) are not as well-characterised as are those in other regions of the CNS. Moreover, the 3rd ventricle is sculpted into the infundibular, optic, and other smaller and ill-defined recesses in mammals (Amat et al., 1992; O’Rahilly and Muller, 1990) and lateral (LR), posterior (PR) and anterior (AR) recesses in zebrafish (Wang et al., 2009; Wang et al., 2012). Three unexplored questions are when such hypothalamic recesses form, whether they are composed of distinct progenitor
cells and whether their appearance correlates with the emergence of particular neuronal subsets.

The \textit{Rx} paired-like homeodomain transcription factor and its fish orthologue, \textit{rx3} are expressed within retinal and hypothalamic progenitors (Bailey et al., 2004; Bielen and Houart 2012; Cavodeassi et al., 2014; Chuang et al., 1999; Furukawa et al., 1997; Lu et al., 2013; Mathers et al., 1997; Stigloher et al., 2006; Medina-Martinez et al., 2009; Muranishi et al., 2012; Pak et al., 2014; Zhang et., 2000) and play a central role in eye development. Disruption of \textit{Rx} leads to small or absent eyes in mouse (Bailey et al., 2004; Mathers et al., 1997; Medina-Martinez et al., 2009; Muranishi et al., 2012; Zhang et., 2000) and is associated with anophthalmia in humans (Voronina et al., 2004). In zebrafish, loss of function of \textit{rx3}, including mutation in the zebrafish \textit{rx3} gene (\textit{chk} mutant) disrupts eye morphogenesis (Kennedy et al., 2004; Loosli et al., 2003; Stigloher et al., 2006): retinal progenitors are specified, but remain trapped in the lateral wall of the diencephalon, failing to undergo appropriate migration (Rembold et al., 2006) and differentiation (Stigloher et al., 2006).

In addition to its well-documented role in eye formation, \textit{Rx/rx3} governs hypothalamic development. \textit{Rx}-null mice show variable penetrance, but all display abnormalities in the ventral hypothalamus (Mathers et al. 1997; Medina-Martinez et al. 2009; Zhang et al. 2000). Lineage-tracing studies demonstrate that \textit{Rx+} progenitors give rise to \textit{sf1+} VMN and \textit{pomc+} Arc tuberal neurons, while targeted ablation of \textit{Rx} in a subset of VMN progenitors leads to a fate switch from an \textit{sf1+} VMN identity to a \textit{dlx1+} DMN identity (Lu et al., 2013). These studies suggest that \textit{Rx} functions in progenitor cells to cell-autonomously select \textit{sf1+} VMN and \textit{pomc+} Arc identities. In zebrafish, \textit{chk} mutants and \textit{rx3} morphants similarly show reduced numbers of pevTub \textit{pomc+} neurons and, additionally decreased NPO \textit{avp+} (formerly \textit{vt}: arginine vasotocin) neurons (Dickmeis et al., 2007; Tessmar-Raible et al., 2007), although currently the underlying mechanism is unclear. These studies, together, raise the possibility that \textit{Rx/rx3} plays a widespread role in the differentiation of tuberal and anterior/NPO hypothalamic neurons.
In mice, expression of the secreted signalling ligand, Shh, overlaps with that of Rx (Shimogori et al 2010) and conditional ablation of Shh from the anterior-basal hypothalamus results in phenotypes that resemble the loss of Rx, including a reduction/loss of avp+ PVN and pomc+ Arc neurons (Shimogori et al., 2010; Szabo et al., 2009). As yet, however, the link between Shh and Rx/rx3 remains unclear and the mechanisms that operate downstream of Shh and Rx/rx3 to govern hypothalamic differentiation are unresolved.

Here we analyse rx3 and shh expression and function in the developing zebrafish hypothalamus. Analysis of chk mutant and rx3 morphant fish, together with EdU pulse-chase experiments, show that rx3 is required for a switch in progenitor domain identity, and for the survival and anisotropic growth of tuberal/anterior progenitors, including their progression to rx3-shh+ AR cells and pomc+, ff1b+ and otp+ TH+ tuberal/anterior fates. Timed delivery of cyclopamine or SAG reveals that shh signalling governs these processes via dual control of rx3 expression, inducing, then downregulating it. We demonstrate that rx3 downregulation, mediated by shh signalling, is an essential component of rx3 function: failure to downregulate rx3 leads to the failure of anisotropic growth, loss of the shh+rx3- AR and failure of tuberal/anterior cell differentiation. Together, our studies reveal a mechanism that elaborates early pattern around the hypothalamic ventricle by the selective growth of distinct progenitor cells.
Materials and Methods

Animals
Zebrafish were staged according to (Kimmel et al., 1995). Ckh\textsuperscript{w29} fish were kindly provided by Dr. Breandan Kennedy (University College Dublin, Ireland).

Nomenclature
We use the terms preoptic, anterior, tuberal and posterior to define the rostro-caudal domains of the hypothalamus. The region we define as anterior may overlap with the region that is conventionally termed the NPO (see Discussion).

In situ hybridisation
Single and double in situ hybridization methods were adapted from Thisse and Thisse 2008 and Lauter et al 2011 (details in supplementary methods), embryos post-fixed in 4% paraformaldehyde and visualised by Olympus Nomarski or confocal microscopy. For cryostat sectioning, embryos were re-fixed and equilibrated in 30% sucrose, and 15\mu m serial adjacent sections cut. n=10-40 embryos for wholemounts; n=4-6 embryos for sectioned data.

EdU analysis
Embryos were pulsed with 300mM EdU for 1hr on ice, chased for 1, 5 or 25h, then processed for cryostat sectioning and double EdU/in situ hybridisation analysis (details in supplementary methods) using the Click-iT EdU Alexa Fluor 488 Imaging Kit (Fisher Scientific).

Immunohistochemistry
Anti-phosP3 (Upstate) and anti-cleaved Caspase (Cell Signaling Technology) were used at 1:1000. Fixed embryos or sections were processed according to Liu et al., 2013 and mounted in VectaShield.

Length measurements
Length was determined through measurements of images, where in situ patterns could be detected relative to morphological landmarks (diencephalic-telencephalic junction (DTJ), optic commissure, lateral ventricle, posterior hypothalamus and adenohypophysis). For each experiment, length was normalized to the average length of age-matched sibling controls.
**Cell quantitation**
phosH3+ and EdU+ cell numbers were obtained through counts in serial adjacent sections through individual hypothalami using in situ patterns against morphological landmarks (above) to determine relative position. For chk mutants, section position was determined relative to unaffected posterior hypothalamus.

**Image acquisition**
DIC or fluorescence images were acquired using an Olympus BX60 with SPOT programme (Diagnostic Instruments Inc), Zeiss Confocal LSM510 Meta or Olympus Confocal. Data was processed with Adobe Photoshop CS3/Adobe Illustrator CS.

**Statistical Analysis**
Statistical analyses were performed using Prism 5. Each data value sampled were tested for Gaussian distribution prior to unpaired t-test by performing baseline subtraction of the two datasets and analyzed through D’ Agostino and Pearson omnibus normality test.

**Cyclopamine treatment**
Cyclopamine (in ethanol) was used at 50uM, optimized on basis of *ptch1* downregulation (20,50,100,120uM tested). Cyclopamine or ethanol were added to dechorionated embryos, kept in the dark.

**SAG treatment**
SAG (Millipore-EMD chemicals) in DMSO was used at 10µM, optimised on basis of *ptc1* upregulation (2,5,8,10uM tested). SAG or DMSO was added to dechorionated embryos in E3 medium, kept in dark.

**Morpholino**
0.25M *rx3* ATG (targets TSS) and 0.15mM *rx3* E212 (targets splice site) (GeneTools,LLC) (Tessmar-Raible et al., 2007) were injected into one-cell embryos and morphants selected on basis of absent eyes.
Abbreviations

| Abbreviation | Description                                      |
|--------------|--------------------------------------------------|
| AH           | anterior hypothalamus                            |
| AR           | anterior recess                                  |
| Arc          | arcuate nucleus                                  |
| cTub         | caudal tuberal hypothalamus                      |
| dA           | dorso-anterior                                   |
| DMN          | dorsomedial nucleus                              |
| DTJ          | diencephalic-telencephalic junction              |
| NPO          | neurosecretory preoptic (anterior) area          |
| oc           | optic commissure                                 |
| pevTub       | periventricular tuberal                          |
| PO           | preoptic hypothalamus                            |
| PH           | posterior hypothalamus                           |
| PR           | posterior recess                                 |
| PVN          | paraventricular nucleus                          |
| rTub         | rostral tuberal hypothalamus                     |
| SON          | supraoptic nucleus                               |
| TH           | tuberal hypothalamus                             |
| VMN          | ventromedial nucleus                             |
| vT           | ventral tuberal                                  |
| zli          | zone limitans intrathalamica                     |
| 3V           | 3\textsuperscript{rd} ventricle                  |
Results

**rx3 expression in 3rd ventricle cells**

Previous studies have described zebrafish *rx3* expression (Bielen and Houart 2012; Cavodeassi et al., 2014; Chuang et al., 1999; Kennedy et al., 2004; Loosli et al., 2003; Stigloher et al., 2006) but have not performed a detailed analysis in the 2-3 day embryo. Neurons in the hypothalamus, including *pomc*+ and *avp*+ neurons that are decreased/lost in the absence of *rx3* (Dickmeis et al., 2007; Tessmar-Raible et al., 2007) begin to differentiate over 2-3 days (Liu et al., 2003; Dickmeis et al., 2007; Tessmar-Raible et al., 2007) and we therefore focused on this period. At 55hpf, *rx3* is detected in three adjacent zones in the hypothalamus (Fig.1A-B”). In keeping with mouse nomenclature (Lu et al, 2013) we term these zones I, II, III, characterized by the thin strip of *rx3*(*weak*) cells in zone II. In zone I, *rx3* is expressed in neuroepithelial-like cells that form a T-shape, comprising the AR and LR of the 3rd ventricle (Fig.1B’-D’) but is excluded from the AR tips (Fig.1C’D’ arrowheads). In zone II, *rx3* labels cells that closely line the AR/LR, again excluded from the AR tips (Fig.1E,E’, arrowheads). In zone III, *rx3* marks neuroepithelial-like cells around the 3rd ventricle which in this region (between anterior and posterior recesses, see Fig.1A,B”) is small (Fig.1F,F’). At 30hpf, the entire 3rd ventricle is small and lined throughout by *rx3*+ neuroepithelial-like cells (Fig.1G-I). Thus the well-defined recesses of the 3rd ventricle, and characteristic *rx3*+ profiles, develop over 30-55hpf.

**Tuberal/anterior hypothalamus elongates from proliferating *rx3*+ progenitors**

To determine the position of *rx3*+ cells relative to other hypothalamic regions we compared expression to *emx2* and *fgf3*, which mark the posterior, ventro-tuberal and dorso-anterior hypothalamus (Herzog et al., 2004; Kapsimali et al., 2004; Liu et al., 2013; Mathieu et al., 2002) and to the adenohypophysis and DTJ, morphologically distinct landmarks. Over 30-55hpf, *rx3* expression is rostral and largely complementary to *emx2*, and sandwiched between ventro-tuberal and dorso-anterior *fgf3*+ cells (Fig.2A-H’, schematics O), and in zone III, overlies the adenohypophysis. This suggests that throughout 30-55hpf *rx3* demarcates cells at the boundary of posterior and tuberal/anterior hypothalamus.
Prior to 30hpf, $rx3$ is expressed in progenitor cells (Bielen and Houart 2012; Cavadeossi et al., 2013; Loosli et al., 2003; Rembold et al., 2006; Stigloher et al., 2006) and the 3rd ventricle is known to harbour cycling cells (Bosco et al., 2013; Lee et al., 2006; Wang et al., 2009, 2012; Wulliman et al., 1999). To directly address whether 30hpf $rx3^+$ cells proliferate, we pulsed fish with EdU, culled acutely, and analysed sections (Fig.2I,J). At 30hpf, 77% EdU+ cells are $rx3^+$ and the remainder immediately abut $rx3^+$ cells (Fig.2I,I'; n=110 cells, 4 embryos). Co-analysis of alternate sections with EdU and phosH3 shows that S-phase cells progress to M-phase (Fig.2J,J'). Analysis of control embryos with phosH3 and $rx3$ confirms that the majority of cycling cells at 30hpf are $rx3^+$ (68% phosH3+ cells co-express $rx3$; 32% phosH3+ cells abut $rx3^+$ cells; Figs.2K,N; n=76 cells, 4 embryos). Wholemount views of embryos double-labelled with $rx3$ and phosH3 suggest that by 55hpf, fewer cycling cells are $rx3^+$ (Fig.2L,L'). Sections confirm this, showing that at 55hpf, 35% cycling cells are $rx3^+$, 28% abut $rx3^+$ cells but 38% are now detected in the $rx3^-$ recess tips (Fig 2M,N; n=92 cells, 4 embryos).

Although expressed in proliferating cells, the length of $rx3$ in zones I and III does not change over 30-55hpf (Fig.2A,E) indicating its dynamic regulation. Proliferation correlates, though, with rostro-caudal growth of the tuberal/anterior hypothalamus (Figs.2A,E,O,P). Growth is greatest over 30-48hpf (Fig.1P), and is 2.5-fold greater than rostro-caudal growth of the posterior hypothalamus, or the dorsal diencephalon over this period (Fig.2Q). In summary, the tuberal/anterior hypothalamus shows anisotropic growth over 30-55hpf, driven from proliferating $rx3^+$ cells and their immediate neighbours.

**Development of $rx3$-$shh^+$ AR and tuberal/anterior immature neurons**

We next characterized the growing tuberal/anterior hypothalamus. At 30hpf, $shh$ is detected uniformly in the hypothalamus (Fig.3A,A'): double-FISH analysis reveals extensive co-expression with $rx3$ (Fig.3D,D', yellow arrowheads). $rx3^+shh^+$ cells are bounded rostrally and ventrally by $rx3^+shh^-$ cells (Fig.3D'red arrowheads) and caudally/dorsally by $shh^+$ cells (Fig3D' green arrowheads). In
the co-expressing region, \textit{rx3} is strongest dorso-caudally (Fig.3D’). Similar expression domains are detected at 55hpf (Fig.3B,E) but a \textit{shh+rx3}- domain now projects in the tuberal/anterior hypothalamus (Fig.3B,E,F,F’ white arrowheads). This domain appears to be composed of cells that have downregulated \textit{rx3}, resulting in the characteristic zone II, but is significantly (1.5-fold) longer at 55hpf compared to 30hpf (Fig.3D,E white arrows). Analysis of sections shows that \textit{shh+rx3}- cells line the AR tips (Fig.3F’ arrowheads; Supp.Fig.1A,A’,C red arrowheads). Together with our previous data, this suggests that AR tip cells are \textit{shh+/rx3-} progenitors that derive from adjacent \textit{rx3+shh+} progenitors.

In zebrafish, immature tuberal/anterior hypothalamic neurons can be characterized through expression of the transcription factor \textit{otpa} (Eaton and Glasgow, 2007; Lohr et al., 2009; Herget et al., 2014; Manoli and Driever, 2014), the nuclear receptor \textit{nr5a1/sf1} orthologue, \textit{ff1b} (Kuo et al, 2005) and the precursor polypeptide \textit{pomc} (Liu et al., 2003; Herzog et al., 2004; Dickmeis et al., 2007; Tessmar-Raible et al., 2007; Manoli and Driever, 2014). At 30hpf \textit{otpa} is detected in the posterior hypothalamus and at the DTJ (Fig.3G,G’) but by 55hpf, additional \textit{otpa+} cells are detected in the tuberal and anterior hypothalamus (Fig.3H,H’ white arrowheads; see Eaton and Glasgow, 2007) adjacent to the \textit{shh+} AR (Fig.3I). Ventral views show that these are periventricular, suggesting they are immature neurons (see Fig.4F; Herget et al., 2014). \textit{ff1b} expression is detected at 30hpf (Fig.3J,J’) and by 55hpf, is expressed broadly in the tuberal hypothalamus. Sections reveal that \textit{ff1b} is expressed in \textit{shh+} AR cells and adjacent periventricular cells (Fig.3K-K”). \textit{pomc+} cells cannot be detected in the 30hpf hypothalamus (Fig.3L,L’) but by 55hpf are detected in the tuberal hypothalamus (Fig.3M-M”) rostral to \textit{rx3+} progenitors (Fig.3N white arrowheads). Together, our data show that anterior elongation correlates with the development and growth of a \textit{shh+rx3}- AR and with the differentiation of \textit{otp+}, \textit{ff1b+} and \textit{pomc+} cells in the tuberal/anterior hypothalamus (schematized in Fig.3O,P).
rx3 is required for *shh*+ AR and neuronal differentiation

We next addressed the requirement for rx3 in development of the tuberal/anterior hypothalamus. Previous studies have shown that *pomc*+ and *avp*+ neurons are absent in embryos lacking rx3 (Dickmeis et al., 2007; Tessmar-Raible et al 2007) but a more extensive characterization of other progenitor/differentiating cells has not yet been performed.

Analysis of 55hpf *ckh* embryos shows that the *shh*+ AR fails to develop in *chk* mutants (Fig.4A-B", white asterisk; note that posterior *shh* expression in the floor plate and basal plate appears unaltered). *rx3* expression itself is markedly different in *ckh* mutant embryos compared to siblings: zones I-III cannot be clearly resolved, (Fig.4A',B',G-H"").

The failure in development of the *shh*+ AR correlates with a failure in differentiation. Mutant embryos lack *otp*+ cells in both the tuberal and anterior hypothalamus (Fig.4C-D" white arrowheads: note *otp*+ cells in the posterior hypothalamus and at the DTJ (green arrowheads) appear unaffected). Previous studies suggest that the anterior *otp*+ progenitors give rise to Group 2/3 *tyrosine hydroxylase* (*th*)+ dopaminergic neurons (Lohr et al., 2009), and in keeping with this, mutant embryos lack Group 2/3 *th*+ neurons (Fig.4E-F" white arrowheads: note Group 4-6 *th*+ neurons are not eliminated). *rx3* mutant embryos additionally lack *pomc*+ cells (Fig.4I-J" white arrowheads) and *ff1b*+ cells (Fig4K-L", white arrowheads) in the tuberal hypothalamus (note *pomc*+ cells in the adenohypophysis (green arrowheads) are still detected) Finally, *fezf1*, a HD gene that in mouse is regulated by *sf1* (Kurrasch et al 2007) and in fish, regulates *otpb* (Blechman et al., 2007), is markedly reduced (Fig.4M-N" white arrowheads); at the same time, ectopic expression is detected in the telencephalon. *rx3* morphant embryos closely phenocopy *ckh* mutants (SuppFigs.2,3; Fig.6F-I, Fig.7). Together, these analyses show that *rx3* is required for establishment of the *shh*+*rx3*+ AR and for the differentiation of tuberal/anterior cells (Fig.4O,P).
Rx3 represses dorsal and ventro-tuberal progenitors

We postulated that, as in mouse (Lu et al 2013), rx3 may switch the identity of other progenitor domains to select posterior tuberal/anterior progenitor fates, and that the absence of rx3 will lead to alterations in progenitor domains/increased alternate fates.

The transcription factor nkd2.1 (previously known as nkd2.1a (Manoli and Driever, 2014)), whose homologue in mouse is required for tuberal neuronal differentiation (Correa et al., 2015; Kimura et al., 1996; Yee et al., 2009), shows subtle difference in expression in ckh mutants at 25hpf: two sets of nkd2.1+ cells in the forming tuberal/anterior hypothalamus (Fig.5A,A‘ blue arrowheads) cannot be detected (Fig.5B,B‘). By 55hpf, this difference is pronounced: nkd2.1 is reduced in the anterior hypothalamus and not detected in the rostral tuberal hypothalamus (Fig.5C,D; position of tuberal/anterior hypothalamus confirmed through double-labelling with shh (Fig.5C‘,D‘)). nkd2.1 in the caudal tuberal, posterior hypothalamus and posterior tuberculum is seemingly unchanged.

Nkd2.1/2.4 and pax6 exert cross-repressive interactions in the hypothalamus (Manoli and Driever, 2014), prompting us to examine expression of pax6. In control embryos, pax6 is confined to the thalamus/dorsal hypothalamus and abuts the dorsal-most boundary of rx3 (Fig.5E,E‘). In the absence of rx3, pax6 is detected ectopically in the tuberal/anterior hypothalamus within and rostral to rx3+ cells (Fig.5F red arrowheads, Supp Fig.2). Thus the absence of rx3 leads to a ventral expansion of pax6+ progenitors.

Ectopic pax6+ domains do not extend throughout rx3 zone III (Fig.5F red dotted outline) raising the question of whether other progenitors are also affected by loss of rx3. The ets transcription factor, pea3, is expressed in the hypothalamus at 30hpf, and overlaps with rx3 zone III cells (Fig.3G,G‘). pea3 is downregulated at 55hpf in control embryos but expression persists in ckh mutants (Fig.3H,I). These results suggest that rx3 normally suppresses both dorsal pax6+ and ventro-tuberal pea3+ progenitors and predicts a widespread change in the
profile of other progenitor markers in *chk* mutants. In support of this idea, *asc1a* and *sox3* expression do not resolve in zones I, II and III in *ckh* mutant embryos, in contrast to their appearance in controls (Supp.Fig.4A-H white arrowheads).

In mouse, conditional ablation of Rx leads to a failure to select arcuate/VMN fates, and instead, additional *dlx*+ DMN cells form (Lu et al 2013). To determine whether the increase in *pea3*+ and *pax6*+ expression results in an increase in, respectively, ventro-tuberal and DMN-like cells, we examined the neurohypophyseal marker, *fgf3* and the DMN marker, *dlx1*. Both show slightly stronger expression in *ckh* mutants (Supp.Fig4I-L) and the ventro-tuberal hypothalamus appears longer in *chk* mutants (Supp.Fig4A,C) suggesting that in the absence of rx3, there is some expansion of ventral-tuberal and dorsal progenitors and their derivatives.

**rx3 is required for progenitor survival and anisotropic growth**

The increase in *fgf3* and *dlx1* in *ckh* mutants is, however, mild, suggesting that rx3 may play a role other than switching progenitor fates. In sectioning embryos we had noticed an unusually disorganized accumulation of *shh*+ cells (Fig.6A-C,G-I) suggesting that some ectopic progenitors may accumulate in the recess walls, rather than grow and progress to normal fates.

To examine this further, we compared proliferation and fate in control and *rx3*-null embryos. Loss of *rx3* led to a significant increase in phosH3+ cells in the 55hpf embryo (Fig.6D-F,J,K) that, in contrast to controls, were largely in or adjacent to *rx3*+ cells (Fig.6L). To better determine the fate of proliferating progenitors, we pulsed 30hpf fish with EdU, chased to 55hpf and, on serial adjacent sections, analysed whether EdU+ cells progressed to periventricular cells in the tuberal/anterior hypothalamus, were retained as *rx3*+ or *shh*+ progenitors, or assumed other fates. In *ckh* siblings, the majority (63%; n=156 cells, 6 embryos) EdU+ cells were laterally-oriented chains in the anterior (Fig.6M) or tuberal (Fig.6P) hypothalamus and were detected in/in the vicinity of *ff1b*+ and *pomc*+ cells (Fig.6M,O,P). A minority (27%) were *shh*+*rx3*- anterior (Fig.6N,O) or lateral (not shown) recess tip cells. No EdU+ *rx3*+ cells were
detected in zones I or III (Fig.6R and not shown). By contrast, in ckh mutant embryos, no EdU+ cells were detected in the region rostral to the adenohypophysis, ie the regions that would form part of the anterior/tuberal hypothalamus (Fig.6T-W). The majority (76%, n=165 cells, 6 embryos) EdU-labelling was detected in/adjacent to shh+ (Fig.6X) rx3+ (Fig.6Y) cells. EdU+ cells accumulated especially at the recess junctions and tips. No cleaved (c)Caspase+ cells were detected after the 25h chase period, but after a 5h chase, cCaspase+ cells, including EdU+cCaspase+ cells were detected in ckh mutants (Fig.6Z). No cCaspase was detected in siblings (Fig.6S).

These findings, together with our previous observations, suggest that rx3+ progenitors give rise to cells, including shh+ AR tip cells, that grow anisotropically and give rise to anterior/tuberal cells. Additionally, these finding show that in the absence of rx3 function, many progenitor cells accumulate in the recesses, where they either die, or fail to differentiate. Together, these observations point to a mechanism in which rx3 selects tuberal/anterior progenitors and governs their survival and growth (Fig.6 schematics).

**Shh is an ‘on-off’ switch for rx3**

Our findings demonstrate that rx3 is upstream of shh in the tuberal/anterior hypothalamus. However, given the critical role of shh in induction and early patterning of the hypothalamus (Bedont et al., 2014; Burbridge et al., 2016; Pearson and Placzek, 2013; Blaess et al., 2015) we wished to test whether at earlier stages of hypothalamic development, shh is upstream of rx3, a possibility suggested by the observation that at epiboly stages, shh is expressed on midline cells, close to the early zone of rx3 expression (Supp.Fig.5A,B).

*Ptch1*, a Shh-receptor and ligand-dependent antagonist, is weakly detected in the forming tuberal/anterior hypothalamus at 30hpf (Fig.7A), but not detected when embryos are exposed to cyclopamine over 10-28hpf (Fig.7G). Similar observations were made with *ptch2* (not shown). At the same time, cyclopamine treatment results in a marked downregulation of rx3 (Fig.7B,H) mimicking the phenotype of slow-muscle-omitted (smu) mutant zebrafish that lack essential
components of the Hh pathway (Supp.Fig.5C,D). Together these results suggest that shh induces rx3 in the early embryo.

By 55 hpf, strong ptch1 expression is detected in zones I and III (Fig.7C) and weaker expression in zone II (Fig.7C). ptch2 expression appears similar (not shown). To determine whether shh influences rx3 at this stage, we exposed embryos to cyclopamine over 28-55hpf. This resulted in an effective inhibition of shh signaling, as judged by ptc1 downregulation (Fig.7I) but led to a consistent increase in rx3 expression (Fig.7D,J). Increased rx3 expression was accompanied by changes that appeared to phenocopy loss of rx3, notably a significant decrease in tuberal/anterior territory (Fig.7D,J white lines and red arrows), a decrease in hypothalamic pomc+ cells (Fig.7E,K,M), the loss of ff1b expression (Fig.7F,L), a decrease in th+ Group2/3 neurons (Supp.Fig.5E,F; note Groups 4-6 in the posterior hypothalamus are unaffected) and a failure to resolve sox3 to zones I,II and III (not shown). These observations suggest that shh mediates rx3 downregulation in zone II, and that this is essential for differentiation of tuberal/anterior hypothalamic progenitors.

This idea predicts that provision of shh may be sufficient to rescue the phenotypic effects of rx3 morphant embryos, once the effects of the morpholino begin to disappear. To test this, we attempted a ‘late rescue’, in which rx3 morphant embryos were exposed to the small molecule shh agonist, SAG over 28-55hpf. SAG was effective in restoring a normal pattern of shh signaling in rx3 morphant embryos, as judged by expression of ptch1 (Fig.7N,T). Furthermore, both the normal pattern of ntx2.1 and the characteristic profile of rx3 in zones I, II and III were restored (Fig.7O,P,U,V). Both pomc+ and ff1b+ cells were restored in rx3 morphant embryos in response to SAG administration (Fig.7Q,Q',R,W,X). Finally, cellular homeostasis was restored: the enhanced numbers of phos-H3+ cells in rx3 morphants were reduced to normal, wild-type levels (Fig.7S,S',Y). This rescue is not seen in an early SAG-treated regime (10-28hpf; not shown), or in chk mutant embryos, treated with SAG over 28-55hpf (Supp.Fig.6) indicating that functional rx3 is required for the late rescue. Together these results suggest
that a Shh-rx3 ON and Shh-rx3 OFF feedback loop (Fig.7Z) is essential for the development of the tuberal/anterior hypothalamus.

Discussion

Here we show that *rx3* is required for morphogenesis of the tuberal/anterior hypothalamus and governs three aspects of cell behaviour: it re-specifies progenitor types to tuberal/anterior identities, promotes their survival and governs their anisotropic growth/migration. Shh co-ordinates tuberal/anterior progenitor selection and behaviour by acting as an on-off switch for *rx3*. Thus a shh-rx3 feed-forward/feedback loop generates tuberal/anterior progenitors that grow to expand the surface area of the 3rd ventricle and diversify the neuronal subtypes that differentiate around it.

**rx3 selects tuberal/anterior hypothalamic progenitors**

Our studies confirm that *rx3* is not required for induction or initial hypothalamic patterning (Kennedy et al., 2004), but show that it is essential to elaborate pattern. Our data suggest that *rx3* autonomously selects *nkx2.1*+ tuberal/anterior progenitors that grow anisotropically. In *ckh* mutant embryos, *pax6a* expands ventrally into *rx3*+ progenitors, a phenotype detected as early as 19hpf (Loosli et al., 2003). The ventral expansion of *pax6a* mimics the phenotype of *nkx2.1/nkx2.4a/nkx2.4b*-null embryos (Manoli and Driever, 2014) and suggests that *rx3* re-specifies progenitors that would otherwise assume a dorsal hypothalamic or prethalamic identity.

At the same time, *rx3* represses *pea3*. In wild type animals, *pea3* overlaps with ventral-most *rx3* at 30hpf, but is downregulated by 55hpf. In *ckh* mutant fish, *pea3* persists. Although we have not performed double FISH with *pea3* and *pax6* in *ckh* mutant fish, their expression patterns appear complementary. This suggests that *rx3* operates as a switch in at least two separate progenitor populations and provides a prosaic interpretation for the existence of two domains - the dorsal *rx3*+*shh*+ and ventral *rx3*+*shh*− domains.
Our studies reveal that rx3 promotes alternate fates in progenitor cells. Its loss leads to one of three outcomes: to undergo apoptosis or to be retained as a proliferating VZ cell (novel outcomes), or to initiate alternate adjacent differentiation programmes: after pulse-chase, some EdU+ cells are detected in periventricular regions in ckh mutants where it is likely that they contribute to nkx2.1+pea3+ progenitors and hence fgf-expressing neurohypophysis ventrally, and to dlx1+ cells dorsally. Dlx1+ cells are likely to be immature DMN-like neurons and notably, crf+ neurons persist in ckh mutants (Dickmeis et al., 2007). Together our studies suggest that rx3 selects tuberal/anterior neuronal progenitors and limits both ventro-tuberal neurohypophyseal and DMN-like progenitors.

In addition to promoting cell survival, rx3 regulates cellular homeostasis in the tuberal/anterior hypothalamus, orchestrating a balance of proliferation and differentiation. We surmise that the increased proliferation seen in the absence of rx3 reflects changes in Wnt or FGF signalling, both of which are upregulated in ckh mutants (Stigloher et al 2006; Yin et al., 2014; this study). fgf3, in particular, normally abuts neuroepithelial-like rx3+shh- cells both in zones I and III and is upregulated in rx3 mutants. Potentially, the driving force for proliferation resides in rx3+shh- cells in zones I and III that progress to rx3+shh+ cells in zone II.

Previous reports have shown that rx3 is required for retinal fate selection and that telencephalic fates are expanded in its absence (Bielen and Houart 2012; Cavadeossi et al., 2013). Our studies likewise show changes in the telencephalon/eye territory: fezf1 is upregulated in rx3 mutants, and both shh and nkx2.1 in the telencephalon/tuberal/anterior area are greatly reduced. Together, these studies suggest that rx3 selects fate in cells of distinct origins, anterior telencephalic and posterior diencephalic. Importantly, not all hypothalamic cells alter their identity in the absence of rx3: the posterior hypothalamus expresses nkx2.1, shh and otp as normal, the rostral-most hypothalamus expresses otp and nkx2.1 and the tuberal hypothalamus expresses nkx2.1, pea3 and fgf3, emphasising the fact that rx3 elaborates, rather than
initiates, hypothalamic pattern.

**Shh is an on-off switch for rx3**

Our study shows that shh is required for both the induction of *rx3* and the progression of *rx3*- to *rx3-shh* progenitors and demonstrates that both steps are required for tuberal/anterior hypothalamic neurogenesis. Downregulation of shh signalling over 10-30hpf leads to an almost complete loss of *rx3* expression. By contrast, downregulation over 30-55hpf leads to sustained *rx3* in zone II and a phenotype that is highly similar to that of *ckh* mutants: *sox3* is not downregulated, the *shh+rx3*- AR does not form, the tuberal/anterior hypothalamus is short and its resident neurons do not differentiate. Importantly, the shh agonist, SAG, can restore normal patterns of proliferating progenitors and neuronal differentiation in late *rx3* morphants. The most likely interpretation of these findings is that shh-mediated *rx3* upregulation is required to select tuberal/anterior progenitors but that shh-mediated *rx3* inhibition is required for these to realise their differentiation programme(s). Future studies are needed to establish whether the downregulation of *sox3*, *nkx2.1*, *ascl1* and *ptc1* that we observe in wild-type, but not *ckh* mutant fish, are similarly required for progression of tuberal/anterior progenitors. We predict that the downregulation of *ptc1*, in particular, supports shh active signaling from zone II cells and contributes to development of the *shh*+ AR. The intricate regulation of induction and cessation of Shh signaling in sets of neighbouring cells is emerging as a common theme within the CNS (Briscoe and Therond, 2013) and provides the opportunity to drive expansion of territories and build increasingly complex arrays of neurons.

In summary, our studies suggest that Shh plays a dual role in *rx3* regulation, inducing, then repressing it, and are consistent with a model in which shh deriving from AR cells, feeds back to *rx3*+ progenitors to promote their further differentiation.
Origins of hypothalamic neurons

Our studies show that the zebrafish tuberal hypothalamus includes regions analogous to the mouse Arc and VMN. Our EdU pulse-labelling studies suggest that shh+ AR cells and differentiating ff1b+ and pomc+ neurons derive from rx3+ cells. After a 25h chase, we detect strings of EdU+ cells, presumably of clonal origin, extending medio-laterally from the shh+ AR tips, to pomc+ and ff1b+ regions, favouring the idea that forming neurons derive from rx3+shh+ progenitors via rx3-shh+ progenitors. In mouse, Rax+ cells give rise to pomc and sf1+ neurons (Liu et al., 2013). Other mouse studies show that Shh+ hypothalamic cells give rise to tuberal neurons (Alvarez-Bolado et al., 2012), and that Shh- ablation in hypothalamic cells leads to the loss of pomc and sf1 (Shimogori et al., 2010) and a reduction in hypothalamic territory (Alvarez-Bolado et al., 2012; Zhao et al., 2012). These studies, together with observations that loss of Nkx2.1 results in loss of tuberal hypothalamic neurons (Correa et al., 2015; Kimura et al., 1996; Yee et al., 2009), disruptions to the infundibulum and a reduction in the size of the 3rd ventricle (Kimura et al., 1996) suggests a conserved differentiation route of pomc and ff1b/sf1 immature neurons and the tuberal hypothalamus from zebrafish to mouse.

In zone I, rx3 is expressed in the anterior hypothalamus, in a region that may be equivalent to the anterior-dorsal domain reported in mouse (Shimogori et al., 2010). Our work, together with previous studies, suggests that here, rx3 plays a role in a conserved differentiation pathway for Avp+ and Group2/3 th+ neurons. Avp+ and Group 2/3 th+ neurons localise within a discrete subregion of hypothalamic otp expression (Lohr et al., 2009; Herget et al., 2014; Herget and Ryu, 2015) and in fish, as in mouse, otp genes are required for AVP and TH neuronal differentiation (Acampora et al., 1999; Lohr et al., 2009; Fernandes et al., 2013). Avp+ neurons fail to differentiate in the absence of rx3 (Tessmar-Raible et al., 2007) and we now show a specific loss of an otpa+ subset and Group 2/3 th+ neurons. This suggests that rx3 governs a subset of otpa+ progenitors in the anterior hypothalamus that will give rise to avp+ and Group 2/3 th+ neurons. We have not yet asked whether this otp+ progenitor subset are dependent on shh. However, in mouse, conditional deletion of hypothalamic shh leads to a
reduction in \textit{otp} expression and \textit{Avp}+ neurons (Szabo et al., 2009) as well as a loss of \textit{Sim1} in the PVN (Shimogori et al., 2010), suggesting that the \textit{shh-rx3-shh} pathway that governs \textit{pomc} and \textit{ff1b} cell fate may likewise govern \textit{Avp}+ and Group 2/3 \textit{th}+ fates. A previous study has highlighted \textit{Sim1} and \textit{Otp} as core components of a conserved transcriptional network that specifies neuroendocrine as well as A11-related hypothalamic dopaminergic neurons (Lohr et al., 2009), suggesting that \textit{rx3} may be intimately linked to this pathway. Notably, since other NPO neurons, including oxytocin+ (previously known as isotocin) and somatostatin+ neurons are not affected by loss of \textit{rx3}, our data suggest that neurons that make up the NPO derive from discrete lineages. Our work adds to a growing body of evidence that directed cell migrations play a pivotal role in ventral forebrain/hypothalamic morphogenesis (Varga et al., 1999; Cavodeassi et al., 2014 and see Pearson and Placzek, 2013). We do not know the mechanisms that operate downstream of \textit{rx3} to govern appropriate migration, but Eph/ephrin signalling, expression of fgf and Netrin, all of which govern cell adhesion and migration of neural cells, are disrupted in \textit{ckh} mutant embryos (Cavadeossi et al., 2013; Yin et al., 2014; this study) and could contribute.

In conclusion, our study suggests a mechanism by which \textit{shh} elaborates pattern in the hypothalamus. Previous reports suggest that \textit{shh} patterns the early hypothalamus in many vertebrates, establishing early progenitor domains (reviewed in Pearson and Placzek, 2013; Blaess et al., 2012). Our study shows that in zebrafish, \textit{shh} elaborates early pattern by switching progenitor domain identity, and promoting the survival and anisotropic growth of the new progenitor cells. Recent studies in the developing spinal cord show that the coordination of growth and specification can elaborate pattern in an expanding tissue, if molecularly distinct neural progenitor domains undergo differential rates of differentiation (Kicheva et al., 2014), raising the possibility that \textit{shh} may govern differentiation rates in the tuberal/anterior hypothalamus. Studies in mice that reveal similarities in the phenotypes of mice in which Shh or Rax are conditionally ablated raise the possibility that features of the mechanism that we describe here may be conserved in other vertebrates.
Finally, the shh-rx3-shh loop that we describe provides a means to maintain a dynamic balance between proliferating and differentiating cells. Studies in mice show that at least a subset of Rax+ cells persist into adulthood as stem cells (Miranda-Angulo et al., 2014) that can direct hypothalamic neurogenesis even in postnatal life. The exquisite regulation of shh, fgf and wnt signalling, via rx3, is likely to hold the key to a better understanding of hypothalamic neurogenesis throughout the lifecourse and support a better understanding of complex human pathological conditions and dysfunctional behaviours that are underlain by tuberal/anterior hypothalamic cells and circuits.
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Figure 1: *rx3* expression around 3rd ventricle

(A) Schematic of 55hpf forebrain indicating subdivisions of hypothalamus relative to rostro-caudal axis and adenohypophysis (blue oval). Green and black show *shh* (Fig.3) and *rx3* expression. Dots depict rostro-caudal position of AR (blue) and PR (red) next to zone III (purple). (B-B”) Wholemount 55hpf embryo after *rx3* in situ hybridization. In (B) lines show planes of section in (C-F). In (B’,B”) side- and ventral-views are aligned (white lines) and show position of *rx3* relative to morphological landmarks. (C-F’) Representative serial sections through single embryo: bottom panels show high power views of boxed regions.
Red arrowheads point to $rx3^+$ cells; black arrowheads point to $rx3^-$ cells in AR tips. (G-I) Wholemount side-view of 30hpf embryo after $rx3$ in situ hybridization; lines in (G’) show positions/planes of sections in (H,I). Scale bars: 50μm
Figure 2: Anterior/tuberal hypothalamus elongates from rx3+ progenitors

(A-H’) Side views after single or double FISH at 30hpf (A-D) and 55hpf (E-H’: E’-H’ show high power views of boxed regions). Arrows in (A,C,E,G) show distances measured for growth comparisons. Coloured arrowheads in (E’,F’) indicate position of recesses. (I-K) MIP of representative sections through 30hpf embryos. (I,J) show serial adjacent sections. (I’,J’) show single channel views. Arrowheads show co-labelled (yellow) or single-labelled (green) cells. Small
white dotted lines indicate approximate positions of anterior and LR. (L, L’) Side views of 55hpf embryo: (L’) shows single channel view. (M, M’) Representative single plane views taken through zone II. (M’) shows single channel view. Yellow arrowheads show double-labelled cells; green arrowheads point to phosH3+ rx3-cells at recess tips. (N) Quantitative analyses of cycling cells at 30-55hpf. (O) Schematic, depicting rx3, fgf3 and emx2 expression, and change in length and axial orientation of hypothalamus. A ‘bending’ of the tuberal/anterior hypothalamus occurs over 30-55hpf, relative to the rostro-caudal axis. Red arrows indicate length of dorsal diencephalon or length of emx2+ PH; white arrow indicates length of ventral-most rx3+ zone III; blue arrow indicates distance from DTJ to rx3+ zone III. (P) Length from DTJ to rostral tip of rx3+ zone III (n=5 embryos each at 30, 40, 48, 55hpf). (Q) Tuberal/anterior hypothalamus grows approx. 2.5-fold more than dorsal diencephalon, emx2+ PH or ventral rx3+ zone III (n=10 each; p<0.0001). Scale bar: 50μm.
Figure 3: Differentiation in the 30-55hpf anterior/tuberal hypothalamus

Side-views (A,B,D,E,G-J,L-N), ventral-views (C,F), sagittal (K) or transverse (K”,M”) sections of 30hpf and 55hpf embryos. (A’,B’,D’E’G’,H’,J’-M’) show high power views of boxed regions. In (B’,E’,F) white arrowheads point to shh(weak+) AR cells, in (H’) to otp+ cells in the tuberal/anterior hypothalamus (H’), in (M’,N) to hypothalamic pomc+ cells. In (D’,E’) arrowheads point to rx3+shh+ cells (yellow), rx3+ cells (red) or shh+ cells (green). (O,P) Schematics, at 30hpf (O) or 55hpf (P). Scale bars: 50µm
Figure 4: *rx3* is required for *shh*+ AR and anterior/tuberal differentiation

Side or ventral views of 55 hpf *ckh* sibling or mutant embryos. Asterisk in (A',A") shows *shh*+ AR, absent in *ckh* mutant (asterisk, B'B'’). White arrowheads point to *otp*+ tuberal/anterior cells (C',C"), Group2/3 *th*+ anterior cells (E',E"), *pomc*+ cells (I',I"), *ff1b*+ cells (K,K") absent in *ckh* mutants (D',D";F',F";J',J;L,L";N,N") and to *fezf1*+ progenitors (M,M") reduced in *ckh* mutant (N,N"). (O,P) Schematics: boxed regions show areas shown in high power views. Scale bar: 50μm.
Figure 5: rx3 suppresses dorsal and ventro-tuberal progenitors

(A-I) Side views of control or rx3-absent embryos. (A’,B’,F’) show high power views of boxed regions in (A,B,F). Blue arrowheads and red arrows in (A-D) point to nkx2.1+ cells, absent in ckh mutants. Blue arrows in (C,D) point to nkx2.1+ ventral-tuberal domain. Red arrowheads in (F) point to ectopic pax6+ cells. Red dotted outline in (F,F’,I) indicates position of rx3+ cells, estimated relative to position of adenohypophysis and 3rd ventricle. Purple arrowhead in (G’) points to rx3+pea3+ cells. (H,I) show views of isolated neuroectoderm. (J,K) Schematics of ckh sibling (J) or mutant (K) 55hpf embryos. Scale bar: 50μm.
Figure 6: rx3 promotes progenitor survival and growth

Wholemount side views (A,G) or sections (B,C,H,I: planes and positions indicated by coloured asterisks) through 55hpf embryos in control or rx3 morphant embryos. (A’,G’) show high-power views of boxed regions in (A,G). Arrowheads in (I) show disorganized sshh+ cells around 3V. (D,J) Wholemount side views of phosH3 in 55hpf control or rx3 morphant embryo. (E,K) Representative sections after phosH3/rx3 co-labelling in 55hpf ckh sibling or mutant embryos. (E’,K’) show single channel views. (F,L) Quantitative analyses: (F) numbers of phosH3+ cells in ckh mutant or sibling embryos (n=6 each). Significantly more phosH3+ cells are detected in mutants compared to siblings (p<0.001). (L) proportion of phosH3+ cells that are rx3+ (black), adjacent to (hatched) or distant from (white) rx3+ cells in mutant vs sibling ckh embryos. (M-R,T-Y) Representative serial sections, from rostral-caudal (coloured asterisks denote approximate position of each section, see (A’,G’) of a 55hpf ckh sibling (M-R) or mutant (T-Y) embryos. White arrowheads point to EdU+ cells. (S,Z) Representative sections of a 35hpf ckh sibling or mutant embryos. 18+/−2 cCasp+ cells detected in chk mutants, n=8 embryos. Schematics summarise mutant vs sibling ckh embryos. Scale bar: 50μm.
Figure 7: Shh signaling functions as an rx3 ‘on-off’ switch

(A-C,F-L) Side or ventral views of 30hpf and 55hpf wild-type embryos, exposed to vehicle or cyclopamine over 10-28hpf or over 28-55hpf. Red arrows and white bars in D, J show distances measured for width and length of tuberal/anterior hypothalamus. (M) Quantitative analysis: significantly fewer pomc cells are detected after cyclopamine exposure (p<0.0001, n=30 embryos). (N-Y) Side or ventral views of 55hpf rx3 morphant embryos, exposed to DMSO vehicle (N-S) or SAG (T-Y) from 28hpf. Scale bar: 50 μm. (Z) Model for anterior/tuberal progenitor development.
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