RESEARCH ARTICLE

Defining developmental diversification of diencephalon neurons through single cell gene expression profiling

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ABSTRACT

The embryonic diencephalon forms integration centers and relay stations in the forebrain. Anecdotal expression studies suggest that the diencephalon contains multiple developmental compartments and subdivisions. Here, we utilized single cell RNA sequencing to profile transcriptomes of dissociated cells from the diencephalon of E12.5 mouse embryos. We identified the divergence of different progenitors, intermediate progenitors, and emerging neurons. By mapping the identified cell groups to their spatial origins, we characterized the molecular features of cell types and cell states arising from various diencephalic domains. Furthermore, we reconstructed the developmental trajectory of distinct cell lineages, and thereby identified the genetic cascades and gene regulatory networks underlying the progression of the cell cycle, neurogenesis and cellular diversification. The analysis provides new insights into the molecular mechanisms underlying the amplification of intermediate progenitor cells in the thalamus. The single cell-resolved trajectories not only confirm a close relationship between the rostral thalamus and prethalamus, but also uncover an unexpected close relationship between the caudal thalamus, epithalamus and rostral pretectum. Our data provide a useful resource for systematic studies of cell heterogeneity and differentiation kinetics within the diencephalon.

KEY WORDS: Mouse, Transcription regulation, Transcriptome, Cell fate specification, Thalamus

INTRODUCTION

Residing in the posterior part of the forebrain, the diencephalon produces distinct classes of neurons, which assemble into multiple nuclei. These functional units are important for linking the anterior forebrain with the rest of the nervous system. For example, the thalamus and epithalamus, which arise from the center of the diencephalon, serve important functions such as learning, motor control, regulation of the sleep-wake cycle, and modulation of emotion and motivation (Hikosaka, 2010; Sherman, 2007). Therefore, understanding the development of the diencephalon is necessary to improve our comprehension of this integral component of forebrain circuitry.

Based on gene expression patterns and morphological landmarks, the embryonic diencephalon is divided into at least three transverse segments called prosomeres (Puelles and Rubenstein, 2003; Rubenstein et al., 1994). During embryogenesis, proliferating

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progenitor cells in the alar plate of the respective prosomere (p) give rise to the pretectum (p1), the thalamus and epithalamus (p2), and the prethalamus (p3). Additional intra-prosomere subdivisions have been proposed (Ferran et al., 2007; Kiecker and Lumsden, 2005). For instance, the zona limitans intrathalamica (ZLI), which is wedged in between p2 and p3, forms a compartment that is delineated by rostral and caudal cell-lineage restriction boundaries (Zeltser et al., 2001). Importantly, the ZLI acts as an organizer by producing Shh, a secreted protein, to regulate regionalization and patterning of the entire diencephalon (Kiecker and Lumsden, 2004). Shh acts in concert with other signaling molecules emanating from the roof plate, including those in the Fgf, Bmp and Wnt families, to regulate the development of the diencephalon (Chatterjee and Li, 2012; Scholpp and Lumsden, 2010). Despite some progress, our understanding of the pattern formation and neuronal differentiation of the diencephalon remains incomplete. For instance, although much is known about the molecules that define populations of postmitotic neurons arising from different areas of the diencephalon, considerably less is clear about the molecular features that delineate their respective progenitor populations. Prior studies have revealed highly variable progression in cell proliferation and neurogenesis, the so-called morphogenetic gradients, across the developing diencephalon (Jones, 2007). Because of this developmental asynchrony, traditional expression profiling, which measures averaged gene expression in bulk tissue, lacks the resolution necessary to gain full insight into the transcription regulation that underlies the formation of different diencephalic lineages. However, single cell gene expression analysis, which provides such resolution, has only been performed in diencephalon-derived neurons from postnatal mice (Kalish et al., 2018; Phillips et al., 2018 preprint; Rosenberg et al., 2018; Zeisel et al., 2018).

In this study, we sought to systematically define cellular heterogeneity and transcriptional landscapes during cell specification and differentiation in the embryonic diencephalon. We performed single cell RNA sequencing (scRNAseq) of dissociated cells from the mouse diencephalon at embryonic day (E) 12.5. Our unbiased scRNAseq allowed characterization of the molecular features across different cell types and cell states that arise from various diencephalic domains. Furthermore, we inferred developmental trajectories of diverse cell lineages, and thereby identified the genetic cascades and gene regulatory networks underlying the progression of the cell cycle, neurogenesis, and cellular diversification. To the best of our knowledge, this is the first reported scRNAseq analysis of mouse embryonic diencephalon. Our data provide a valuable resource for systematic studies of cell heterogeneity and differentiation kinetics within the diencephalon.

RESULTS AND DISCUSSION

Classification of cell groups of the mouse diencephalon

We used a mouse line, in which bicistronic *creER-ires-EGFP* transcripts are expressed from the *Gbx2* locus so that thalamic



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neurons specifically produce both creER and enhanced green fluorescence protein (EGFP) (Chen et al., 2009). Utilizing EGFP as a guide, we dissected the thalamus and surrounding tissues from mouse embryos at E12.5. Using the Chromium Drop-Seq platform (10x Genomics), we profiled the transcriptome for over 7500 single cells. After applying quality filters, we obtained a dataset with 7365 cells and 14,387 genes for subsequent analysis. Using the Seurat algorithm (Butler et al., 2018; Satija et al., 2015), we partitioned the 7365 cells into 18 clusters, which were visualized with t-distributed stochastic neighbor embedding (t-SNE; Fig. 1A). Differential gene expression analysis identified genes that were significantly enriched in each cell cluster (Fig. 1B, Table S1). We used a set of cell cyclerelated genes (Tirosh et al., 2016) to calculate cell-cycle scores and thereby to assign cell-cycle status (G2/M, S or postmitotic) to each cell (Fig. 1C). In t-SNE projections, the distribution of cells with various mitotic statuses showed a trend reflecting the progression from proliferating progenitors to postmitotic cells (Fig. 1C). Inspection of the average gene counts revealed a trend of decreasing transcript levels from progenitors to postmitotic cells (Fig. 1D), indicating that the dividing progenitors have higher gene counts than their progeny. We classified cluster 11 as low-quality cells, as they had much lower gene counts than the others, and

contained few cluster-specific genes (Fig. 1B,D). Besides brain cells, we recovered non-neural cell types, such as endothelial cells (cluster 17) and microglia (cluster 18). Hierarchical analysis classified the 18 cell clusters into five groups: postmitotic neurons, neuron precursors or intermediate progenitor cells (IPCs), neural progenitors, non-neural cells and low-quality cells (Fig. 1E). We identified *Id3* and *Ptn* as the markers that were common for progenitors; *Gadd45 g* and *Hes6* for newly committed progenitors, or neuron precursors; *Stmn2* and *Gap43* for postmitotic neurons (Fig. 1F, Table S1). Therefore, our scRNAseq data illustrate the heterogeneity of cells within the mouse diencephalon at E12.5.

Characterization of the molecular features of postmitotic neurons

Next, we related postmitotic cell groups to their endogenous positions by inspecting RNA *in situ* hybridization data in the Allen Developing Mouse Brain Atlas (Thompson et al., 2014) or by examining expression ourselves. By comparing the expression of at least two markers in t-SNE projections and *in situ* hybridization on serial sagittal sections of mouse embryonic brains, we assigned cell cluster 3 to caudal thalamus, cluster 1 to rostral thalamus, cluster 2 to prethalamus, cluster 4 to ZLI, cluster 7 to epithalamus, and cluster 5 to

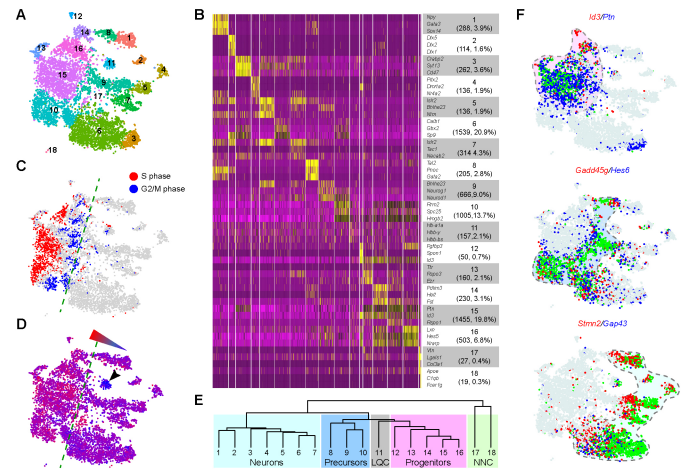


Fig. 1. Identification of major cell groups in E12.5 mouse diencephalon by scRNAseq. (A) Visualization of 18 classes of cells using t-SNE. Each dot represents a single cell; similar cells are grouped and shown in color. (B) Heatmap showing expression of marker genes across cell groups. The number and percentage of cells are shown in brackets under the cluster number. Relative expression from 2 to –2 are shown in yellow and purple, respectively. (C,D) t-SNE plots showing the inferred cell cycle phase (C) and average gene counts (D). The dashed lines delineate between proliferating cells (to the left) and postmitotic cells (right). In D, the arrowhead indicates the low-quality cells, and the triangle illustrates the gradient of gene counts. (E) Dendrogram showing the relationship between cell groups recovered by scRNAseq. LQC, low-quality cells; NNC, non-neural cells. (F) Expression of the genes marking cell clusters corresponding to neural progenitors, neuron precursors and postmitotic neurons, respectively (top to bottom).

pretectum (Fig. 2A-F). Clusters 6, 8 and 9 apparently represented intermediate cell states in transition to more differentiated cells of clusters 3, 1 and 5/7, respectively (see below). As the Allen Developing Mouse Brain Atlas has limited information for E12.5 mouse brains, we curated a list of markers for different diencephalic cell types at E12.5 based on published studies (Chatterjee et al., 2012; Delogu et al., 2012; Mallika et al., 2015; Suzuki-Hirano et al., 2011; Virolainen et al., 2012). Examination of these known markers confirmed our annotation of postmitotic cell clusters (Fig. 2G-I). The ZLI acts as the primary organizing center in the caudal forebrain by producing Shh as a morphogen (Kiecker and Lumsden, 2004). Unexpectedly, we failed to detect any enrichment of Shh transcripts in cluster 4 cells, but we found Shh expression in the presumed floor plate cells (cluster 12, see below). This indicates that, although scRNAseq at the current sequencing depth is sufficient to distinguish different cell groups, it may be not sensitive enough to provide a comprehensive transcriptional profile of a given cell group.

The pretectum is commonly subdivided into rostral and caudal parts, or tripartite subdivisions, in avian embryos (Ferran et al.,

2007; Rendahl, 1924). Although the mammalian pretectum has not been well characterized, evidence suggests a similar organization between mice and chicken (Ferran et al., 2008). C1ql2 and Bhlhe23, which represent cluster 5, are expressed in the rostral part of the pretectum, immediately posterior to the fasciculus retroflexus, or habenular peduncular tract (Fig. 2F). On coronal sections, the expression domain of Bhlhe23 resides above that of Pou4f1 (Fig. 2E,F, insets), which marks the epithalamus-derived lateral habenular nucleus (Chatterjee et al., 2014; Quina et al., 2009). The caudal part of the pretectum expresses Sox14, Gata3 and Gad2, which are absent from the rostral part of the pretectum (Fig. 2B,C). Therefore, our analysis suggests that the mouse pretectum can be subdivided into rostral and caudal domains with distinct molecular features. Notably, most genes that are specific for cluster 1 are present in both the rostral thalamus and the caudal pretectum (Fig. 2B.C). It has been shown that these two domains express the same set of developmental regulators (Ascl1, Helt, Gata2/3, Tal1/2 and Sox14), and produce GABAergic neurons (Delogu et al., 2012; Virolainen et al., 2012). However, the molecular underpinning of

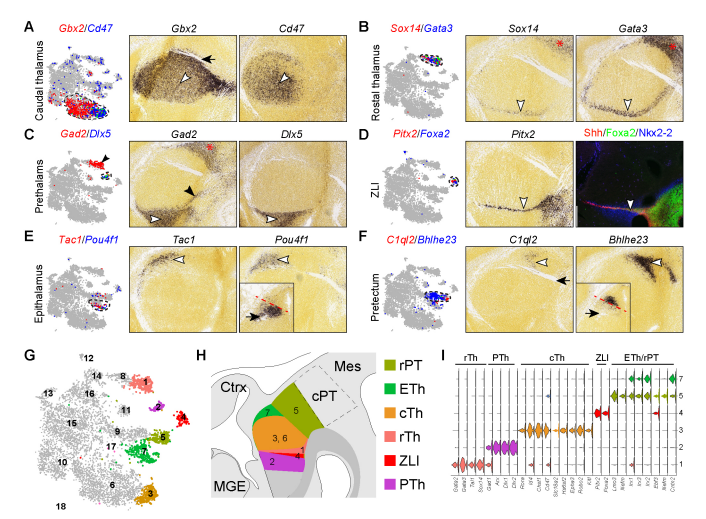


Fig. 2. Relating postmitotic cell groups to their spatial positions in the diencephalon. (A-F) Expression of cell-specific markers with t-SNE plotting (left) and on sagittal sections of E13.5 mouse diencephalon (right; from the Allen Developing Mouse Brain Atlas). White arrowheads indicate the marker expression domain; black arrow denotes the fasciculus retroflexus; black arrowheads in C show that Gad2 is expressed in the rostral thalamus, as well as the prethalamus; asterisks indicate the caudal pretectum, which exhibits a similar gene expression profile as rostral thalamus. Insets in E and F show the expression of *Pou4f1* and *Bhlhe23* on coronal sections of E13.5 diencephalon; the red dashed line separates their expression domains. (G) t-SNE projections of diencephalic neurons. (H) Schematic summary of the postmitotic neurons identified by scRNAseq in their endogenous positions in the diencephalon. (I) Violin plot showing expression of genes that are known to mark different diencephalic neurons at E12.5. cPT, caudal pretectum; cTh, caudal thalamus; ZLI, zone limitans intrathalamica.

GABAergic differentiation may be different, as the deletion of *Ascl1* or *Gata2* caused distinct phenotypes in the rostral thalamus and posterior pretectum (Virolainen et al., 2012). As our microdissection likely included tissues from the rostral thalamus due to its close proximity to the thalamus, we provisionally assign cluster 1 to the rostral thalamus. However, we are uncertain whether this cluster may contain cells from the caudal pretectum. Future scRNAseq of the entire diencephalon will be necessary to discern the molecular features between the rostral thalamus and caudal pretectum at E12.5.

Determination of the molecular characteristics of diencephalic progenitor cells

Compared with postmitotic neurons, diencephalic progenitors display less distinctive expression profiles among different groups (Fig. 1B,E). We had to rely on combinations of multiple genes to annotate the progenitor cell groups. For example, Neurog1 and Dll1 are both expressed in presumptive IPCs, or basal progenitors, in the caudal thalamus (cluster 10) and neuron precursors in the epithalamus and rostral pretectum (cluster 9; Fig. 3A); Dll1 is also expressed in neuron precursors within the rostral thalamus (cluster 8; Fig. 3A). As described previously (Wang et al., 2011), Neurog2 and Olig3 are expressed in both apical progenitors next to the ventricular surface (cluster 15) and IPCs (cluster 10) in the subventricular zone of the p2 domain (Fig. 3B). In addition, Olig3 is expressed in the ventricular zone of the rostral thalamus (cluster 14), whereas *Neurog2* is expressed in newborn neurons in the epithalamus and rostral pretectum (clusters 9 and 16; Fig. 3B). Gbx2 and Lhx2 are co-expressed in newborn thalamic neurons (cluster 6; Fig. 3C). In addition, *Lhx2* is expressed in IPCs of the caudal thalamus (cluster 10) and newborn neurons from the epithalamus and rostral pretectum (cluster 9; Fig. 3C). Pax3 and Cdon are co-expressed at the dorsal midline (cluster 13), but the expression domain of *Pax3* extends ventrally, including progenitors (cluster 16) and newborn neurons (cluster 9) in the rostral pretectum (Fig. 3D). Cluster 12 cells express Shh and Foxa2, which are specific for the ZLI and basal plate (Fig. 2D, Fig. S1A, Table S1). In addition, cluster 12

cells express *Ednrb* and *Ntn1*, which are only present in the basal plate (Fig. S1B,C), indicating that cluster 12 represents the basal plate. The annotation of progenitor cell groups is summarized in Fig. 3E,F. A summary of the annotation and number of all 18 cell groups is provided in Fig. S2. It is worth noting that the cell number and percentage may not be an accurate measurement of different populations of diencephalic cells at E12.5 because we mostly focused on the thalamus when we collected tissues for the current scRNAseq (see Materials and Methods for details).

Expression data of many feature genes are not available in Allen Developing Mouse Brain Atlas. Even fewer exist for coronal brain sections, which are important for assessing expression in progenitor domains. Future studies with improved cell clustering algorithms and additional gene expression data should help to refine the division of different diencephalic progenitor domains.

Identification of spatial gene expression patterns in the roof plate and p2

It is generally believed that the epithalamus and the thalamus arise from the dorsal and ventral parts of p2 (Chen et al., 2009; Mallika et al., 2015). However, cell clustering could not reliably subdivide the presumed p2 progenitor cells (cluster 15). This suggests that the epithalamic and thalamic progenitors share similar molecular features, or that these two groups of progenitors contain gradual expression changes of relatively few genes. To test the latter hypothesis, we explored the 'spatial structure' in our scRNAseq data with different analytical methods.

Fgf8 is expressed in two narrow longitudinal stripes flanking the roof plate overlaying p1-2, whereas *Wnt1* is expressed at the roof plate covering the p1 and mesencephalon (Martinez-Ferre and Martinez, 2009; Martinez-Ferre et al., 2013). We found that the presumed roof-plate cells (cluster 13) were adjacent to the p2 cells (cluster 15) in t-SNE projections (Fig. 3E). *Fgf8* and *Wnt1* were detected in two small discrete subgroups within the roof-plate cluster (Fig. S1D,E). Moreover, *Wnt1*⁺ cells were embedded with *Pax3*⁺ cells, indicating a p1 origin of the *Wnt1*⁺ cells (Fig. S1D). Finally, we detected a decreasing gradient of *Rspo3* from the *Fgf8*⁺ cells to the rest of the p2

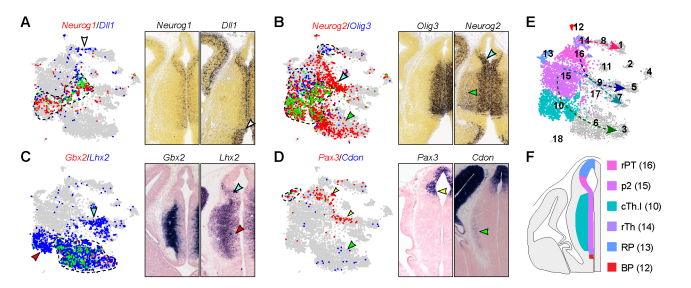


Fig. 3. Relating progenitor cell groups to their original positions in the diencephalon. (A-D) Expression of cell-specific markers on t-SNE plots (left) and on coronal sections of mouse diencephalon (right) at E13.5 (A,B; from Allen Developing Mouse Brain Atlas) and E12.5 (C,D). On t-SNE plots, the cells that express a given marker are shown in red or blue as indicated; those that express both are shown in green and outlined by dashed lines. Cell groups expressing a single marker are indicated by an arrowhead with matching color on both t-SNE plots and sections. (E) t-SNE plots of diencephalic progenitor cell groups. The dashed arrows show developmental trajectories. (F) Schematic of scRNAseq-identified progenitor groups within the diencephalon. BP, basal plate; cTh, caudal thalamus; p2, prosomere 2; RP, roof plate; rPT, rostral pretectum; rTh, rostral thalamus.

progenitor cells in t-SNE projections, recapitulating the endogenous expression pattern (Fig. S1E,F). Collectively, these observations suggest that some spatial information is preserved among cells in t-SNE projections. Therefore, we applied trendsceek, a statistical method that tests points (cells in low-dimensional projections) in a pairwise manner to identify genes that exhibit distance-dependent expression in single cell gene expression datasets (Edsgärd et al., 2018). Trendsceek identified 92 genes with significant spatial trends (P<0.01, Benjamini-Hochberg adjusted), including at least six different patterns exemplified by Ttr, Fgf8, Bmp4, Pax3, Irx3 and Olig3, respectively (Fig. 4A-F). By inspecting the endogenous expression in the Allen Developing Mouse Brain Atlas, we found that trendsceek correctly predicted the spatial expression of several genes (Fig. 4K,L). Ttr, which encodes a transporter protein that carries thryroxin and retinol, is normally expressed in precursor cells of the choroid plexus (Herbert et al., 1986). Therefore, we classified the Ttr^+ cell group as choroid plexus epithelium.

We uncovered 36 genes that displayed opposing gradients in p2 progenitors along the presumptive dorsoventral axis, similar to the expression patterns of *Irx3* or *Olig3* (Fig. S3A,B). Some of these genes have been implicated in p2 regionalization. For instance, *Irx3* and *Pax6*, which exhibit a dorsal^{high}-vental^{low} gradient (Fig. 4E, Fig. S3A,B), are known to establish differential competence for Shh signaling in the patterning of the p2 progenitor zone (Robertshaw et al., 2013). *Tcf7l2*, which shows a ventral^{high}-dorsal^{low} gradient (Fig. S3B), is important for thalamus development as its deletion results in a thalamic-to-epithalamic fate conversion (Lee et al.,

2017b). Collectively, our data suggest that the prospective epithalamus and thalamus arise from two p2 progenitor domains with graded, rather than discrete, gene expression differences.

It has been shown that Fgf8 signaling induces Sprv1 but represses Fgfr3 (Liu et al., 2003). We observed opposing gradients of Spry1 (dorsal^{high}) and Fgfr3 (ventral^{high}) (Fig. 4G). This indicates the presence of graded Fgf8 activity in the p2 progenitor domain, in agreement with the known function of *Fgf8* in patterning of p2 (Martinez et al., 1999). A prior report has demonstrated a complex pattern of Wnt/β-catenin signaling in the p2 domain, with lower activity in the ventral and rostral regions (Bluske et al., 2009). However, the pattern of Wnt/β-catenin activity does not correlate with the expression pattern of known Wnt ligands. How the spatial pattern of Wnt/ β -catenin activity is established in the p2 progenitor domain remains largely unknown. Trendsceek predicted significant spatial trends of Rspo1, 2 and 3, which code for secreted proteins of the R-spondin family. Indeed, we confirmed a caudalhigh-rostrallow gradient of Rspo1, restricted expression of Rspo2 near the roof plate, and a dorsal^{high}-ventral^{low} gradient Rspo3 (Fig. 4H-J). As R-spondin proteins are known to potentiate Wnt/β-catenin signaling (de Lau et al., 2014), the spatial expression of Rspo1-3 may account for the differential activity of Wnt/β-catenin found in the p2 domain. Collectively, our findings suggest that the morphogen gradients of Fgf8 and Wnt/β-catenin signaling may orchestrate graded gene expression in the p2 progenitor zone, leading to the formation of the epithalamus and thalamus from the dorsal and ventral parts of p2, respectively.

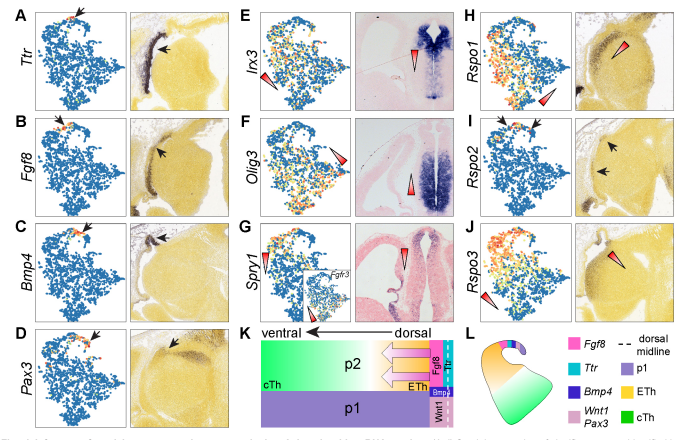


Fig.. 4. Inference of spatial gene expression patterns in the p2 domain with scRNAseq data. (A-J) Spatial expression of significant genes identified by trendsceek on scRNAseq data projected with t-SNE (left) and on sagittal brain sections (right). Expression levels are scaled from 0 (blue) to 1 (red); arrows indicate the matching expression domain on the section and cells on the t-SNE; triangles denote the expression gradient. (K,L) Schematics of the spatial expression patterns in the diencephalon (K) and t-SNE data (L). The arrow indicates the morphogen gradients of Fgf8 and Wnt/β-catenin signaling. Images in A-D and H-J are from Allen Developing Mouse Brain Atlas.

Progenitor cells in the caudal thalamus display a unique pattern of cell cycling and neurogenesis

Prior studies have demonstrated the existence of a unique and significant population of IPCs in the caudal thalamus, forming a defined subventricular zone as found in the developing neocortex and basal ganglia (Smart, 1972; Wang et al., 2011). As expected, we recovered abundant presumptive IPCs (cluster 10) from the caudal thalamus (Fig. 3E,F). This cluster contained cells in S and G2/M phases, despite the robust expression of the proneural genes *Neurog1* and *Neurog2* (Figs 1C, 3A,B), indicating that these neurogenic progenitors are highly proliferative. It has been shown that IPCs in the thalamus depend on *Neurog1/2* and *Pax6* (Wang et al., 2011). However, as these genes are expressed in other areas that lack IPCs, other unknown factors must be involved in the amplification of IPCs in the caudal thalamus.

To investigate the mechanism underlying IPC development, we digitally isolated cells that presumably originated from the caudal thalamus (clusters 3, 6, 10 and 15) and ordered them according to pseudotime using the Monocle 2 algorithm (Qiu et al., 2017). For comparison, we performed the same analysis with the rostral pretectal cells (clusters 5, 7, 9 and 16). Monocle inferred a trajectory that was composed of thalamic cells from clusters in the $16\rightarrow10\rightarrow6\rightarrow3$ order, and a rostral pretectal trajectory that was composed of cells from clusters in the $16\rightarrow9\rightarrow7/5$ order (Fig. S4A,B). Inspection of the pseudotime and expression of known markers, *Id3* (apical progenitors), *Neurog1* (IPCs), *Gbx2* (newborn neuron

precursors) and *Hs6st2* (postmitotic neurons), showed that Monocle correctly ordered cells according to the developmental trajectory of the thalamic lineage (Fig. 5A-C). Interestingly, the apical progenitor $(Id3^+)$ and IPC (Neurog1⁺) groups each contained one group of S-phase cells, and two separate groups of M-phase cells in t-SNE projections (Fig. 5D), suggesting that both apical and IPCs in the caudal thalamus undergo actively symmetric division. Ordering cells according to pseudotime revealed sequential changes of gene expression and temporal coupling between cell cycle and neurogenesis in the caudal thalamus and rostral pretectum (Fig. 5E,F). Remarkably, the expression of *Birc5* and *Atad2*, which mark the M- and S-phase, respectively (Tirosh et al., 2016), displayed at least two out-of-phase waves (Fig. 5E). Furthermore, the last wave of the cell cycle genes overlapped with the peak of *Neurog1* and *Neurog2* expression (Fig. 5E). In sharp contrast to the caudal thalamus, the expression of the cell-cycle genes and apical progenitor markers sharply decreased before the induction of *Neurog1* and *Neurog2* in the rostral pretectum (Fig. 5F). We detected co-expression of Insm1 and Neurog 1/2 in both the caudal thalamus and rostral pretectum (Fig. 5E,F), indicating that *Insm1* is not specific for IPCs as suggested previously (Wang et al., 2011). To study candidate molecules that regulate the development of neural progenitors in the caudal thalamus, we performed differential gene expression analyses. We found that cluster 16 cells were enriched for genes that are involved in stem cell maintenance, Wnt signaling and Hedgehog signaling, whereas IPCs were enriched for genes in the FoxO

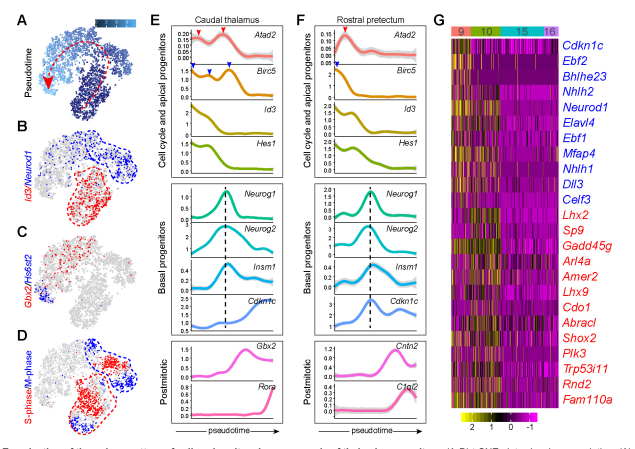


Fig. 5. Examination of the unique pattern of cell cycle exit and neurogenesis of thalamic progenitors. (A-D) t-SNE plots showing pseudotime (A), gene expression (B,C) and inferred cell cycle states (D) of cells originating from the caudal thalamus. The dashed arrow indicates the increase of pseudotime predicated by Monocle. (E,F) Temporal expression profiles of the genes that represent cell cycle and apical progenitors, IPCs (basal progenitors), and postmitotic neurons, respectively, in the caudal thalamus (E) and rostral pretectum (F). The arrowheads indicate the peak expression of genes representing the S (red) and M (blue) phases; the vertical dashed lines show the pseudotime position that corresponds to the peak of *Neurog1/2* expression. (G) Heatmap showing the genes that are highly expressed in clusters 9 and 10. Cluster number is indicated at the top; clusters 15 and 16 are shown for comparison.

signaling pathway, which plays an important role in cell cycle progression (Table S2). Pairwise comparisons among clusters 9, 10, 15 and 16 identified genes that were specific to cluster 9 and 10 (Fig. 5G). Among cluster 10-specific genes, Sp9 (trans-acting transcription factor 9), Gadd45g (growth arrest and DNA-damageinducible 45 gamma) and Plk3 (polo-like kinase 3) have been implicated in cell cycle regulation. Sp9 is expressed and plays an essential role in IPC proliferation within the basal ganglia (Zhang et al., 2016). Furthermore, Rnd2, a transcriptional target of Neurog2 (Heng et al., 2008), and Fam110a are enriched in the subventricular zone of the cortex (Fietz et al., 2012). Remarkably, *Cdkn1c*, which codes for a cyclin dependent kinase inhibitor (p57^{kip2}) important for neuronal cell cycle exit (Dyer and Cepko, 2001; Gui et al., 2007), was highly expressed in cluster 9 (Fig. 5G). By contrast, the induction of Cdkn1c was delayed in the caudal thalamus compared with that in the rostral pretectum (Fig. 5E,F).

We next compared high-order gene transcription regulation in the caudal thalamus and rostral pretectum through weighted gene co-expression analysis (WGCNA) (Langfelder and Horvath, 2008; Zhang and Horvath, 2005). Because scRNAseq recovers only a small fraction of the transcripts in a cell, the technical noise, commonly called dropout, can obscure gene-gene relationships (Stegle et al., 2015). In fact, we found that *Gbx2* and *GFP*, which are transcribed from the same *Gbx2* locus in *Gbx2^{creER/+}* embryos (Chen et al., 2009), showed only moderate correlation (r=0.49; Fig. S5A). To address the dropout issue, we tested three different expression recovery methods, MAGIC (Markov affinity-based graph imputation of cells) (van Dijk et al., 2018), SAVER (single cell analysis via expression recovery) (Huang et al., 2018), and ALRA (adaptive-thresholded low-rank approximation) (Linderman et al., 2018 preprint). All three methods improved the linear coefficient between Gbx2 and GFP (r=1, MAGIC; 0.8, SAVER; 0.94, ALRA; Fig. S5). However, MAGIC imputation distorted the relationship between Gbx2 and Sox14, which were present in different cell types (Fig. S5). Following the standard procedure as described in our previous publication (Heng et al., 2017), we used the ALRA-imputed expression matrix as inputs for WGCNA analysis and identified co-expressed gene modules, which were labeled by colors (Fig. 6A,B). The purple module of the caudal thalamus network, and the light-cyan module of the rostral pretectum network were enriched for Neurog1, Neurog2 and other factors that promote neurogenesis (Table S3). Furthermore, the distribution of module eigengene, the first principal component, of these two modules was closely correlated with clusters 10 and 9, respectively (Fig. S6A,B), and the eigengene changes paralleled the expression of *Neurog1* in cells arranged according to pseudotime (Fig. 6C,D, Fig. S6C,D). By examining the number of genes that fell into modules of the caudal thalamus versus modules of the rostral pretectum, we detected a strong agreement in assignments of most modules between the networks (Fig. 6E). However, the thalamus purple module and pretectal light-cyan module, as well as a few related modules, were poorly preserved (Fig. 6E, Fig. S6C,D). To quantify the overall significance of the network preservation statistics, we computed the composite Z_{summary} scores (Z_{summary}>10 for preserved; Z_{summary}<2 for non-preserved modules) (Langfelder et al., 2011). The thalamus purple module and pretectum light-cyan module were among the least preserved modules in their counterpart networks (Fig. 6F,G). Collectively, our data show that the gene

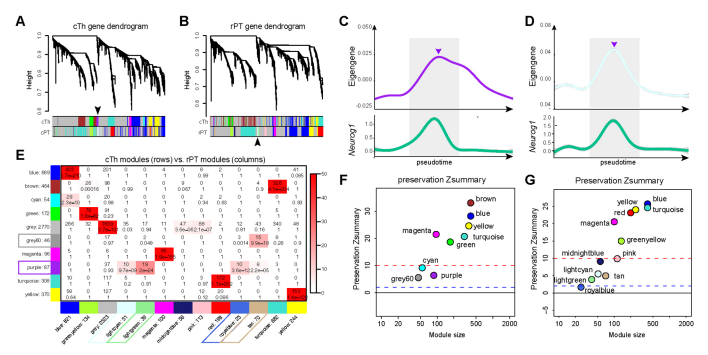


Fig. 6. Comparison of gene co-expression networks between caudal thalamic and rostral pretectal cells. (A,B) Hierarchical clustering trees of genes based on the co-expression network in the caudal thalamus (cTh; A) and the rostral pretectum (rPT; B). The colored rows below the dendrograms indicate module membership in each network. The arrowheads indicate that the purple module of the cTh network is absent from the rPT network, whereas the light-cyan module of the rPT network is absent from the cTh network. (C,D) Expression of thalamic purple module (C) and pretectal light-cyan module (D) eigengene values over pseudotime. The lower panel shows *Neurog1* expression, as shown in Fig. 5E,F. The gray area defines *Neurog1*-expressing cells. The arrowheads indicate the peak of purple and light-cyan modules at the center of the gray area. (E) A contingency table of cTh (rows) and rPT (column). Each row and column is labeled by the corresponding module color and the total number of genes in the module. In the table, numbers indicate gene numbers in the intersecting row-column module. The table is color-coded by $-\log(p)$ of the Fisher exact test *P*-value, according to the color key on the right. (F,G) Preservation statistics of cTh modules in rPT network (F) or rPT modules in cTh network (G). The red and blue dashed lines indicate Z_{summary} cutoff at 10 and 2 (Z_{summary}>10 for preserved; Z_{summary}<2 for non-preserved modules).

network associated with the onset of neurogenesis is markedly divergent between the caudal thalamus and the rostral pretectum.

In summary, by using pseudotime analysis and gene co-expression network analysis, we demonstrate the unique pattern of cell cycle progression and neurogenesis in the caudal thalamus. Importantly, we have identified a set of genes that may contribute to the distinctive coupling of neurogenesis and cell proliferation in the caudal thalamus. We suggest that the delayed induction of Cdkn1c is involved in the significant expansion of IPCs in the caudal thalamus, which may be responsible for the generation of the vast number and diversity of thalamic neurons in mammals.

Inference of developmental trajectories of lineage specification

To reconstruct developmental trajectories of various diencephalic cell types, we applied the URD algorithm (Farrell et al., 2018) to the entire dataset minus the low-quality cells and non-neural cells. Building on an extension of the diffusion map framework (Haghverdi et al., 2015, 2016), URD uses discrete random walks and graph searches to order cells in pseudotime and reconstruct complex developmental trajectories (Farrell et al., 2018). We assigned apical progenitors (mostly clusters 15 and 16) as the root, and postmitotic neurons of clusters 1-5 and 7 as tips, resulting in a multi-branching trajectory (Fig. 7A). The hierarchical tree was visualized with a force-directed layout based on cells' visitation frequency by the random walks from each tip (Fig. 7B). Inspection of the expression of known markers showed progressive changes along consecutive segments of the respective trajectory, as expected

(Fig. 7C, Fig. S7), suggesting that we had successfully reconstructed the developmental trajectory of the diencephalon.

The reconstructed tree closely reflects the development of different populations of diencephalic neurons. For example, the URD-inferred tree predicts a close relationship between the rostral thalamus and prethalamus (Fig. 7A,B). This is in agreement with the notion that these two regions may share a common pool of progenitors but give rise to different subtypes of GABAergic neurons (Kataoka and Shimogori, 2008). Indeed, GABAergic precursors that arise from the rostral thalamus or pretectum retain latent plasticity for subtype identity upon cell cycle exit; removal of various transcription factors, such as Gata2, Helt and Ascl1, Tal2, and Dlx1/2 results in cell fate switching between rostral thalamus and prethalamus (Delogu et al., 2012; Lee et al., 2017a; Sellers et al., 2014; Song et al., 2015; Virolainen et al., 2012). Furthermore, the predicted close relationship between the epithalamus and the thalamus is also in line with the fact that they both arise from the p2 domain (Puelles and Rubenstein, 2003; Rubenstein et al., 1994). In addition, we and others have reported that deletion of Gbx2 or *Tlc7l2* causes cell-fate conversion from thalamic neurons to those of an epithalamic fate (Lee et al., 2017b; Mallika et al., 2015).

The predicted close relationship between the rostral pretectum and the thalamus or the epithalamus is somewhat unexpected as they belong to p1 and p2, respectively. However, evidence from genetic studies in mice and fish has recently suggested a latent cell-fate plasticity between thalamus- and pretectum-derived neurons. For example, deletion of lhx2/9 or increased Wnt signaling causes re-specification of the thalamic to a pretectal fate in the zebrafish embryo (Peukert et al., 2011). Conversely, ectopic expression of *Shh*

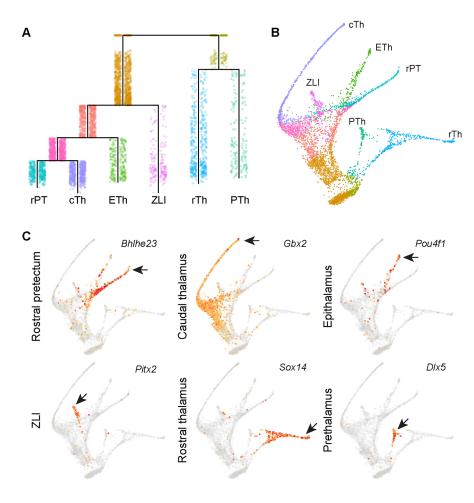


Fig. 7. Reconstruction of developmental

trajectories of the diencephalon. (A) URD-inferred hierarchical tree of diencephalic lineages. (B) Forcedirected layout of the trajectory tree colored by segments as shown in A. (C) Expression of lineage markers overlaid on the trajectory tree. The arrows indicate robust expression in each specific lineage. cTh, caudal thalamus; ETh, epithalamus; PTh, prethalamus; rPT, rostral pretectum; rTh, rostral thalamus; ZLI, zona limitans intrathalamica. can convert the pretectum into a thalamic fate (Kiecker and Lumsden, 2004; Vieira and Martinez, 2006; Vieira et al., 2005). Finally, although it has been demonstrated that the thalamic neurons are converted to an epithalamic fate in mice lacking *Gbx2* or *Tcf7l2* (Lee et al., 2017b; Mallika et al., 2015), our new evidence suggests that the rostral pretectum-specific markers were also ectopically increased in the thalamus in the absence of *Gbx2* and *Tcf7l2* (see below). Collectively, these observations support the view that the thalamus and pretectum, as least the rostral part of the pretectum, are closely related.

Therefore, the single cell-resolved trajectories not only confirm a close relationship between the rostral thalamus and prethalamus, but also uncover an unexpected close relationship between the caudal thalamus, epithalamus and rostral pretectum. It is worth noting that the deduced lineage relationship is based on gene expression. The actual ancestor-progeny relationship will need to be determined in the future by emerging technology combining lineage tracing and scRNAseq (Kester and van Oudenaarden, 2018).

Determination of gene expression landscapes along developmental trajectories

To determine gene expression dynamics during the specification of diencephalic cell lineages, we searched for genes for which expression levels significantly changed along a given URD-recovered trajectory. We identified a total of 647 such so-called 'cascade genes' in each trajectory leading to the formation of the epithalamus (n=313), caudal thalamus (n=361), prethalamus (n=100), rostral pretectum (n=335), rostral thalamus (n=94) and ZLI (n=157; Table S4). For the cascade genes in each lineage, we applied impulse response fitting to determine their temporal dynamics, leading to identification of markers specific for each lineage (Fig. 8A-F, Table S2). Assessing the expression of 20 thalamus-specific markers predicted by URD showed that they correctly highlighted the caudal thalamus branch in the trajectory tree (Fig. S8). Notably, 13 of them (65%) have been shown to be specific for thalamic neurons according to previous studies (Mallika et al., 2015; Suzuki-Hirano et al., 2011), indicating the accuracy of lineage-specific markers recovered by URD.

It has been demonstrated that Gbx2 is essential for the development of the thalamus (Chatterjee et al., 2012; Chen et al., 2009; Mallika et al., 2015; Miyashita-Lin et al., 1999). Through microarray transcriptome profiling, we previously identified a set of genes that are mis-regulated in the mouse thalamus at E12.5 as a consequence of deletion of Gbx2 (Mallika et al., 2015). Remarkably, the downregulated genes showed significant enrichment for URDpredicted markers of the caudal thalamus (hypergeometric tests, Bonferroni-corrected $P=1.37\times10^{-23}$), whereas the upregulated genes were significantly enriched for those of the epithalamus $(P=2.41\times10^{-05};$ Fig. 8A,B). This is in agreement with the conclusion that the loss of Gbx2 results in a thalamus-toepithalamic cell fate change (Mallika et al., 2015). Unexpectedly, we found that the upregulated genes in the Gbx2-deficient thalamus were also enriched for rostral pretectum markers ($P=4.51\times10^{-05}$; Fig. 8C). Therefore, this analysis extends our previous findings by showing that the loss of *Gbx2* causes re-specification of the thalamic fate to a cell fate of rostral pretectum, as well epithalamus. We observed that many genes, such as Irx2, Islr2, Lmo3 and Ebf3, which were reported to be expressed in thalamus lacking Tcf7l2 (Lee et al., 2017b), were indeed markers for the rostral pretectum (Fig. 8C). This suggests that loss of Tcf7l2 may result in a partial cell fate switch from thalamus to rostral pretectum. Collectively, these observations further support the idea that the formation of the epithalamus, caudal

thalamus and rostral pretectum are closely related in terms of transcriptional regulation.

To gain insight into the molecular underpinnings of cell fate decisions, we generated impulse models for transcriptional regulators that exhibited the most dynamic changes in each diencephalic lineage (Fig. S9). We identified 14 such transcription factors associated with development of the rostral thalamus (Fig. 8G). Remarkably, the predicted temporal expression profile of these genes recapitulated their known expression behavior in vivo (Delogu et al., 2012; Sellers et al., 2014; Virolainen et al., 2012). It has been demonstrated that many of these proteins play an essential role in GABAergic neuron differentiation in the rostral thalamus or in other brain regions (Achim et al., 2012; Delogu et al., 2012; Lee et al., 2017a; Sellers et al., 2014; Song et al., 2015; Virolainen et al., 2012). Our analyses also revealed dynamic expression of several transcription factors, such as Otx1, Emx2, Six3, Uncx, Nr4a2 and *Irx1*, without described roles in the rostral thalamus. These genes would be candidates for future genetic studies.

For validation, we focused on the expression of known molecules at the branch point separating the prospective rostral thalamus and prethalamus (Fig. 8H). As expected based on previous studies (Lee et al., 2017a; Sellers et al., 2014; Song et al., 2015; Virolainen et al., 2012), most cells fell along the bifurcation from progenitors into rostral thalamic and prethalamic fates with sequential gene expression changes corresponding to the progression of lineage development (Fig. 8H). Notably, a small fraction of cells that expressed *Dlx2* were found at the early stage of the rostral thalamic lineage, indicating a latent fate plasticity of these cells. Therefore, repression of *Dlx2*, possibly by *Gata2*, is important for the commitment of the rostral thalamic fate, as shown previously (Sellers et al., 2014; Virolainen et al., 2012).

Collectively, our analyses have successfully determined the gene regulation dynamics associated with the development of different diencephalic lineages. Our data provide a valuable resource for interrogation of cell fate specification and differentiation in the developing diencephalon.

Examination of transcriptional networks in single cell-resolved trajectories

Next, we investigated how the URD-inferred developmental trajectories are characterized by gene regulatory networks regulated by transcription factors. We applied the single cell regulatory network inference and clustering (SCENIC) (Aibar et al., 2017) pipeline to uncover the gene regulatory network that underlies the cell-state transition in different diencephalon lineages. We first identified sets of genes that were co-expressed with transcription factors using the GENIE3 algorithm (Huynh-Thu et al., 2010). We then applied RcisTarget (Aibar et al., 2017) to select the co-expression modules, called regulons, that exhibited significant enrichment for binding motifs of the corresponding transcription factors. This resulted in 230 regulons in our dataset. The regulon activity in each cell was calculated with the AUCell computational method (Aibar et al., 2017). By fitting regulon activity changes with an impulse model, we determined their temporal dynamics. As expected, sequential changes of regulon activity reflected the expression dynamics of the corresponding transcription regulators remarkably well (Figs 8G, 9A,B). The congruent changes of gene expression and regulatory state provide additional evidence for the important role of these transcription factors in development of the rostral thalamus. We calculated the binary (on or off) activity of regulons and selected those that were enriched for each lineage (Fig. 9C, Fig. S9A-F). Therefore, by combining URD and gene regulatory network

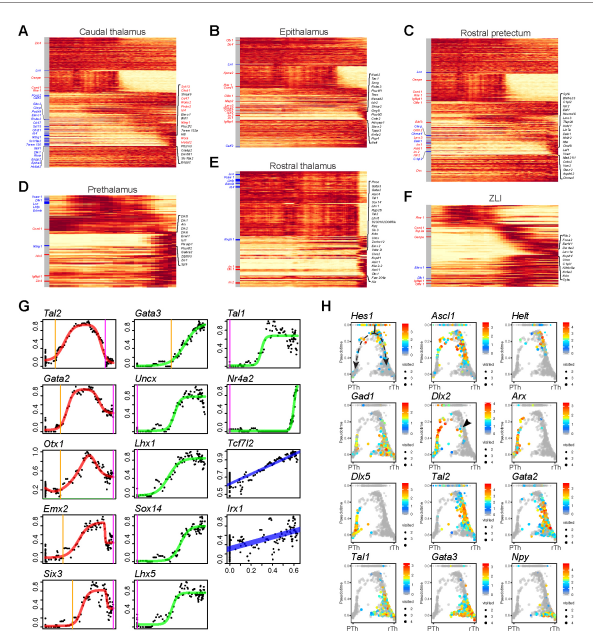


Fig. 8. Gene expression cascades involved in the development of different diencephalic lineages. (A-F) Heatmaps showing gene expression dynamics in different diencephalic lineages. Pseudotime increases from left to right. Lineage-specific genes are shown to the right. To the left are the genes that are significantly altered in the thalamus by *Gbx2* deletion (red, upregulated; blue, downregulated; gray ,unchanged). (G) Temporal dynamics of 14 transcription factors that exhibit the most significant changes in the specification of rostral thalamic neurons. Each dot represents the moving-window average of expression levels; the line shows impulse response fitting: blue for genes that change at a constant rate, green for a single sigmoid impulse, and red for a convex impulse; vertical lines indicate the onset (orange) and offset (magenta) of gene expression. (H) Expression of key developmental regulators overlaid on the branch point forming the rostral thalamus (right) and the prethalamus (left), respectively. The dashed arrows indicate the lineage bifurcation and trajectory; the arrowhead shows the presence of *Dlx2* expression in rostral thalamic cells near the branch point.

analyses, we uncover the transcriptional cascades that accompany the development of progenitors into differentiated cells, and highlight both previously characterized and newly identified trajectory-enriched genes.

In conclusion, our current study has characterized the molecular and cellular heterogeneity of the developing mouse diencephalon. We have identified the key developmental regulators and their associated regulatory networks that define specific cell lineages. The resulting compendium of gene expression data provide opportunities to use genetic studies to interrogate the mechanisms that control cell fate decisions and cell differentiation in the developing diencephalon. In addition, our approach of combining multiple state-of-the art statistical tools provides a framework with which we can classify cell types and cell states, study spatial pattern of gene expression, and reconstruct the specification trajectories of highly complex developmental systems with scRNAseq data.

MATERIALS AND METHODS

Generation and maintenance of mouse lines

All animal procedures described herein were approved by the Animal Care Committee at the University of Connecticut Health Center. The generation of *Gbx2^{creER/+}* (*Gbx2^{tm1.1(cre/ERT2)Jyhl/J*, The Jackson Laboratory #022135) mice were described previously (Chen et al., 2009). PCR conditions and primer}

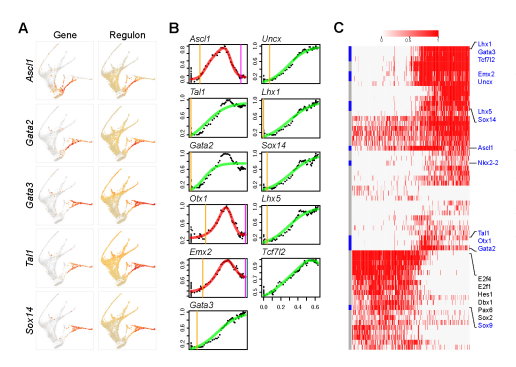


Fig. 9. Gene regulatory networks underlying caudal thalamic

development. (A) Expression of genes (left) and regulons (right) overlaid on the trajectory tree. (B) Temporal changes of regulon activity in association with development of the rostral thalamus. Each dot represents expression level as a moving-window average in pseudotime. The line shows impulse response fitting: green for a single sigmoid impulse, and red for a convex impulse; vertical lines indicate the onset (orange) and offset (magenta) of gene expression. (C) Heatmap showing the binary activity of regulons in the rostral thalamic cells ordered according to pseudotime (increasing from left to right). The URD-identified cascade genes are highlighted in blue.

information are provided at www2.jax.org/protocolsdb/f?p=116:5:0::NO:5: P5_MASTER_PROTOCOL_ID,P5_JRS_CODE:28952,022135. The mouse line was maintained on a CD1 genetic background (Charles River Lab, Wilmington, MA, USA). Noon of the day when a vaginal plug was detected was designated as E0.5 in staging of embryos.

Generation of single cell suspensions

Brains of Gbx2^{creER/+} embryos were dissected in ice-cold PBS. The neural tube was opened after cutting along the dorsal midline. Under a fluorescent dissecting microscope, the thalamus, which was labeled by EGFP expressed from the Gbx2 locus, was dissected together with surrounding tissues. Tissues from four animals were pooled, cut into pieces smaller than 1 mm across, and transferred to ice-cold MACS Tissue Storage Solution (Miltenyi Biotec). For cell dissociation, the storage solution was exchanged with RPMI 1640 Medium (Thermo Fisher Scientific); tissue was pelleted and digested with 500 µl of pre-warmed Accumax (Innovative Cell Technologies) in a 1.5-ml tube at 37°C for 5-10 min. At the end of digestion, the tissue pieces were dissociated by gentle trituration with a wide-bore pipet tip. The cell suspension was strained using a 100-µm cell strainer (Corning), and transferred to 1.5 ml of ice-cold re-suspension buffer (Lebovitz L15 medium with 2% fetal bovine serum, 25 mM HEPES, 2 mM EDTA). The cell clumps that were unable to pass through the filter were placed into a new 500 µl pre-warmed Accumax solution, and the digestion and filtering process was repeated twice. Following dissociation, cells were stained with Trypan Blue, and counted and visualized with Countess II Automatic Cell Counter (Thermo Fisher Scientific). The single cell suspension had over 77% viability.

Single cell RNA-sequencing library construction and sequencing

Libraries were prepared using the Chromium Single Cell Kit v1 (PN-120233, 10x Genomics) and sequenced on an Illumina NextSeq 500. The raw reads were processed to molecule counts using the Cell Ranger pipeline (version 1.3.1, 10x Genomics) with default setting (Zheng et al., 2017).

Cell clustering and classification

The raw unique molecular identifier (UMI) counts from Cell Ranger were processed with the Seurat R package (version 1.4.0.9) (Satija et al., 2015). Genes that were detected in fewer than three cells were removed. Cells in which over 5% of the UMIs were mapped to mitochondrial genes and those that contained over 2500 genes were discarded. Library-size normalization was performed on the UMI-collapsed gene expression for each barcode by scaling the total number of transcripts per cell to 10,000. The data were then

log2 transformed. In total, 7365 cells and 14,387 genes (an average of 1147 detected genes/cell) were used in cell clustering. On average, we obtained 68,275 sequencing reads per cells, which represented median 1758 unique genes per cells. We followed the standard Seurat workflow as described by the Satiji lab (https://satijalab.org/seurat/) to perform cell clustering and marker gene detection. For direct partitioning of roof-plate cells, we used the new interactive plotting function of Seurat (version >2.0).

For cell-cycle score assignment, Seurat's 'RunPCA' function was used to perform principal component analysis with a list of cell cycle-dependent genes (Tirosh et al., 2016). The resulting first principal component, which correlated with the activity of known S- and G2/M-phase genes, was referred to as a 'cell-cycle score'. Cells with a cell-cycle score below -3 or above 3 were assigned as S or M phase, respectively.

Examination of spatial structures of scRNAseq data

We created a new Seurat object by extracting clusters 13 and 15 from the full dataset and performed normalization, scaling and t-SNE analyses. As inputs to trendsceek, the positions of the 4261 cells in the t-SNE were used as locations of the points, and the normalized expression matrix was used as marks of the points. We used 10,000 permutations of the marks to obtain the null distribution representing marks being conditionally independent of the spatial location of the cells. We selected significant genes with Benjamini–Hochberg-adjusted $P \le 0.05$ for at least one of the four statistic tests (Edsgärd et al., 2018).

Inference of developmental trajectories

We used Monocle 2 (Qiu et al., 2017) to perform pseudotime analysis. For the caudal thalamus, we generated a new Seurat object by extracting clusters 3, 6, 10 and 15. For the rostral pretectum, we created a new Seurat object by extracting clusters 5, 7, 9 and 16. The resulting Seurat objects were exported to Monocle. We reconstructed the developmental trajectories in these two lineages by following the standard procedure described in the Trapnell lab (http://cole-trapnell-lab.github.io/monocle-release/tutorials/). Temporal gene expression profiles were plotted using the ggplot2 package (Wickham, 2016).

To perform URD analysis, we followed the procedure described previously (Farrell et al., 2018). We assigned presumptive apical progenitors (mostly clusters 15 and 16) as the root, and postmitotic neurons (clusters 1-5 and 7) as tips, resulting in a branching trajectory tree. The URD 'aucprTestAlongTree' function was used to identify the 'cascade genes' that were progressively increased along each trajectory. To identify

lineage-specific genes, we used all the cascade genes for each lineage and selected those that were highly enriched in the given lineage compared with the rest of the diencephalon. We used the area under the precision-recall curve (AUC) for those genes that were greater than the area expected for a random feature by a factor of 3. To examine the enrichment of lineage markers in Gbx2 target genes, we performed hypergeometric tests using the R package piano (Väremo et al., 2013).

For the genes identified as members of a gene expression cascade, we used the URD 'geneCascadeProcess' function to smooth their expression using a moving window average (18 cells per window, moving five cells at a time) along the trajectory. The value was then scaled to the maximum window average for each gene in the trajectory. The smoothed expression of each gene was fitted with an impulse model to determine its temporal expression dynamics (Farrell et al., 2018).

WGCNA

To impute dropout values of the scRNAseq expression matrix, we tested three different expression recovery methods by following the recommended procedures: MAGIC (https://github.com/KrishnaswamyLab/ MAGIC#python), SAVER (https://mohuangx.github.io/SAVER/articles/ saver-tutorial.html) and ALRA (https://github.com/KlugerLab/ALRA). Although all methods restored the linear coefficient between Gbx2 and GFP expression, only ALRA maintained the expected relationship between Gbx2 and Sox14 (Fig. S5). SAVER moderately improved the expression matrix. Therefore, we used the ALRA-imputed matrix to produce gene co-expression networks by following the WGCNA procedures described in our previous study (Heng et al., 2017). We used softpower=12 to generate a 'signed' WGCNA network for both datasets. Modules were defined as branches of a hierarchical cluster tree using the top-down dynamic tree cut method (Langfelder et al., 2008). For each module, the expression patterns were summarized by the module eigengene, defined as the singular vector of the standardized expression patterns. Pairs of modules with high module eigengene correlations (r>0.85) were merged. The module membership for each gene with respect to each module was then defined as the Pearson correlation between the expression level of the gene and the module eigengene (Oldham et al., 2008). We used the 'modulePreservation' function of the WGCNA package (Langfelder et al., 2011) to calculate the preservation of gene modules across networks with 200 permutations.

Inference of single cell gene regulatory network

We used the SCENIC pipeline (Aibar et al., 2017) to determine the gene regulatory network in each diencephalic lineage. We first applied GENIE3 (Huynh-Thu et al., 2010) to the gene count matrix to construct a coexpression network. As input to GENIE3, we filtered genes with at least three counts in 10% cells, and cells with at least 90.15 UMI, resulting a final matrix of 8320 genes and 7162 cells. We used the 'RcisTarget' algorithm to select potential direct-binding targets (regions) based on DNA-motif analysis, and then the 'AUCell' algorithm to calculate the network activity in each individual cell, resulting in an activity matrix in which the columns represent cells, and the rows are regions (Aibar et al., 2017). Using a default threshold, we converted the regulon activity to on/off binary activity matrix, which was then used to produce heatmaps. We examined the temporal changes of regular activity by applying the URD's 'geneCascadeProcess' function to the continuous AUC matrix.

In situ hybridization and immunohistochemistry

Standard protocols were used for immunofluorescence, and *in situ* hybridization, as described previously (Chen et al., 2009). Detailed protocols are available on the Li Laboratory website (https://health.uconn.edu/li-lab/ protocols/). The following antibodies were used: goat anti-Foxa2 (Santa Cruz Biotechnology, sc-6554); mouse anti-Nkx2-2 (Developmental Studies Hybridoma Bank, 74.5A5); rabbit anti-Shh (Santa Cruz Biotechnology, sc-9024); Alexa Fluor secondary antibodies (Invitrogen). To generate riboprobe templates, cDNA of E13.5 mouse brain was used to PCR amplify the 3' end of the target gene, ranging between 500-700 bp with the T7 RNA polymerase recognition site incorporated into the product. Standard *in vitro* transcription methods using T7 polymerase (Promega) and digoxigenin-UTP RNA labeling mix (Roche) were used to produce antisense riboprobes.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: J.Y.H.L.; Methodology: J.Y.H.L.; Software: J.Y.H.L.; Formal analysis: Q.G., J.Y.H.L.; Data curation: Q.G.; Writing - original draft: J.Y.H.L.; Writing - review & editing: Q.G., J.Y.H.L.; Visualization: J.Y.H.L.; Supervision: J.Y.H.L.; Project administration: J.Y.H.L.; Funding acquisition: J.Y.H.L.

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Data availability

The sequencing files and raw gene count matrix have been deposited in NCBI's Gene Expression Omnibus and are accessible through accession number GSE122012. All the computer codes and necessary data associated with the manuscript are provided in the supplementary information, and are also available at https://github.com/JLiLab/scRNAseq_Diencephalon.

Supplementary information

Supplementary information available online at http://dev.biologists.org/lookup/doi/10.1242/dev.174284.supplemental

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