In vivo clonal analysis reveals spatiotemporal regulation of thalamic nucleogenesis

Short title: In vivo clonal analysis of thalamic nucleogenesis

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ABSTRACT

The thalamus, a crucial regulator of cortical functions, is composed of many nuclei arranged in a spatially complex pattern. Thalamic neurogenesis occurs over a short period during embryonic development. These features have hampered the effort to understand how regionalization, cell divisions and fate specification are coordinated and produce a wide array of nuclei that exhibit distinct patterns of gene expression and functions. Here, we performed an *in vivo* clonal analysis to track the divisions of individual progenitor cells and spatial allocation of their progeny in the developing thalamus. Quantitative analysis of clone compositions revealed evidence for sequential generation of distinct sets of thalamic nuclei that are associated with the location of the founder cell. Furthermore, we identified intermediate progenitor cells that produced two to four neurons populating more than one thalamic nuclei, indicating a late specification of nuclear fate. Our study reveals an organizational principle that governs the spatial and temporal progression of cell divisions and fate specification, and provides a framework for studying cellular heterogeneity and connectivity in the thalamus.

INTRODUCTION

A large portion of the vertebrate brain is organized so that groups of neurons that share similar properties are arranged in aggregates Developmental mechanisms underlying the formation of such nuclear brain structures are much less understood compared with the mechanisms that regulate the formation of laminar structures including the cerebral cortex and retina. The thalamus contains dozens of nuclei that play crucial roles in controlling cortical functions (Jones, 2007; Sherman, 2016). Most thalamic nuclei are populated by excitatory neurons that project axons to the cerebral cortex. Some of these nuclei receive substantial subcortical input and convey sensory and motor information to specific areas in the neocortex, whereas other nuclei receive input mostly from the cortex and serve as a hub for cortico-cortical communications. Thalamic nuclei can be distinguished by morphology and gene expression as well as areal and laminar specificity of their axonal projections to the cortex. Genetic lineage tracing at the population level revealed that neurons that populate cortex-projecting nuclei are derived from a progenitor domain marked by the proneural basic helix-loop-helix (bHLH) transcription factors NEUROG1 and NEUROG2 (Fig.1A, pTH-C domain; (Vue et al., 2007)). In addition to these "excitatory nuclei", the thalamus also contains "inhibitory nuclei" that are exclusively composed of GABAergic neurons, which include the intergeniculate leaflet (IGL) and ventral lateral geniculate (vLG) nucleus. Notably, recent studies identified a small progenitor domain (pTH-R) that is located rostral to the pTH-C domain during neurogenesis and expresses the proneural bHLH factors ASCL1 and TAL1. Population-level lineage tracing studies using these markers found that pTH-R contributes to IGL and vLG nuclei (Fig.1A; (Jeong et al., 2011; Vue et al., 2007)). However, due to the complex spatial arrangement of all thalamic nuclei and the narrow time window during which neurons for these nuclei are produced (Angevine, 1970), little is known about how individual progenitor cells undergo divisions and fate specification and eventually produce neurons and glia that populate thalamic nuclei.

As demonstrated in invertebrate brains (Kohwi and Doe, 2013), determination of precise cell lineage at the single-cell level is a key to understanding the molecular and cellular mechanisms that underlie the formation of complex brain structures like the mammalian thalamus. In this study, we performed a clonal analysis of neural progenitor cells of the thalamus using MADM (Mosaic Analysis of Dual Markers)-based genetic fate mapping. Quantitative analysis of clone size and composition obtained with two inducible Cre drivers allowed us to determine the division patterns of individual neuroepithelial and radial glial progenitor cells and their differential contributions to distinct thalamic nuclei. Furthermore, we used a third Cre driver to label more differentiated, basally dividing progenitor cells and captured the fate of their progeny generated in the last few cell divisions. These results reveal how patterning, cell division and cell fate specification are coupled in a spatially and temporally specific manner to generate distinct thalamic nuclei during development.

RESULTS

Birthdating of thalamic neurons using a thymidine analog

To investigate the temporal progression of thalamic nucleogenesis, we first performed EdU (ethynyldeoxyuridine) birthdating by labeling progenitor cells during the S-phase of the cell cycle at embryonic day 10.5 (E10.5), E11.5, E12.5, E13.5 and E14.5. We analyzed the distribution of birthdated, EdU-positive cells at postnatal day 0 (P0) (Fig.1B). The lateral-to-medial neurogenic gradient was evident at all rostrocaudal and dorsoventral levels, consistent with previous studies (Altman and Bayer, 1988; Angevine, 1970). With the exception of the earliest-generated (dorsal lateral geniculate; dLG, lateral posterior; LP, lateral dorsal; LD) and the latest-generated (paraventricular; PV) nuclei, most thalamic nuclei became postmitotic between E11.5 and E13.5. Some nuclei (e.g., dLG, LP, LD) were generated over at least 48 hours (E10.5 to E12.5), whereas others (e.g., centromedian; CM, reuniens/rhomboid; Re/Rh, mediodorsal; MD) were generated within one day around E13.5. Thus, thalamic neurogenesis occurs over a short span of embryonic development and likely involves a complex pattern of division and fate specification at the level of individual progenitor cells.

Establishment of clonal lineage tracing strategy using MADM

In order to determine the temporal and spatial dynamics of neural progenitor cells in the thalamus, we performed genetic fate mapping at the clonal level by using the mosaic analysis with double markers (MADM) system (Fig.S1A; Hippenmeyer et al., 2010; Zong et al., 2005). When the frequency of Cre-mediated recombination is

kept low by titrating the tamoxifen dosage, the sparse G/R clones generated by "G2-X" type segregation would allow us to determine the division pattern and fate of the two daughter cells that are derived from the originally recombined progenitor cell.

In this study, three inducible Cre driver lines, *Gli1*^{CreERT2}, *Olig3*^{CreERT2} and *Neurog1*^{CreERT2} were combined with *MADM-11* mutants. Both *Gli1* and *Olig3* are expressed in neuroepithelial and radial glial cells in the thalamus (Vue et al., 2009). *Gli1* is a direct target gene of Sonic Hedgehog signaling pathway and is expressed highly in the rostral part of the thalamus as well as in other regions in the forebrain including the prethalamus and hypothalamus. OLIG3 is expressed in the entire thalamic ventricular zone and in the zona limitans intrathalamica (ZLI), whereas its expression is minimal in other forebrain regions (Vue et al., 2007; see Fig.1A). In the thalamus, NEUROG1 is largely excluded from Notch-ICD (intracellular domain of NOTCH)-expressing neuroepithelial and radial glial cells, and is expressed in basally dividing progenitor cells in the pTH-C domain (Wang et al., 2011).

To validate the recombination patterns of each CreER driver, we performed a population analysis of reporter mice. The *Rosa-H2B-GFP* reporter line crossed with the *Gli1*^{CreERT2} driver, and the *Ai6 ZSGreen* reporter crossed with *Olig3*^{CreERT2} or *Neurog1*^{CreERT2} driver showed labeling of progenitor cells within the thalamic primordium and production of labeled progeny in the thalamus a day after the tamoxifen administration (E10.5 for *Gli1*, E11.5 for *Olig3* and E12.5 for *Neurog1*) and 18.5 (Fig.2A).

We then crossed each CreER driver with $MADM-11^{GT/GT}$ mice, and the CreER/+: $MADM-11^{GT/GT}$ mice were crossed with $MADM-11^{TG/TG}$ mice. A day after

the tamoxifen administration, MADM brains for all three CreER drivers had G/R and Y clones in thalamic primordium (Fig.2B), and the apical and basal processes of the labeled cells in Gli1^{CreERT2} MADM brains were positive for Nestin, a neuroepithelial and radial glial cell marker (Fig.S2A). Thus, using these three CreER drivers would allow us to fully trace the lineage of spatially and temporally divergent progenitor cell populations in the thalamus at different stages of differentiation. By adjusting the dosage of tamoxifen for each CreER driver, we achieved a low frequency of labeling a pair of progeny generated by a single recombined progenitor cell in the thalamus (Fig.2C). The percentage of hemispheres that had one G/R or Y clone one day after the Cre activation was less than 25% (~20% for Gli1CreER, ~12% for Olig3CreER and ~14% for Neurog1CreER), and almost no hemispheres had two clones (~2% for Gli1^{CreER}, ~3% Olig3^{CreER} and ~5% for Neurog1^{CreER}), which validated our methodology for clonal analysis of thalamic progenitor cells. For the *Olig3*^{CreER} driver, we analyzed 186 hemispheres (27 litters) of CreER+ E18.5 mice and 112 hemispheres (23 litters) of CreER+ P21 mice. Out of these brains, we found 72 and 22 G/R clones, respectively. For the Gli1^{CreER} driver, we analyzed 194 hemispheres (37 litters) of CreER+ E18.5 mice and 44 hemispheres (14 litters) of CreER+ P21 mice. Out of these brains, we found 27 and 9 G/R clones, respectively. In order to avoid analyzing potentially mixed clones that arise from two or more recombined cells, we abandoned two litters of Olig3^{CreER} E18.5 brains in which every hemisphere had fluorescently labeled cells in the thalamus.

Analysis of clonal division patterns of thalamic progenitor cells

A major advantage of using the MADM system for lineage tracing is that it allows for analysis of the division pattern of the recombined single progenitor cell based on the numbers of green cells and red cells in each G/R clone. In early embryonic neocortex, neuroepithelial cells undergo proliferative symmetric divisions to increase the pool size of progenitor cells. As neurogenesis starts, progenitor cells divide asymmetrically and produce a self-renewed radial glial cell and a more differentiated cell, either a postmitotic neuron or an intermediate progenitor cell (IPC). IPCs in embryonic neocortex express the T-box transcription factor TBR2 and divide symmetrically in basal locations away from the surface of the lateral ventricle (Kriegstein and Alvarez-Buylla, 2009; Florio and Huttner, 2014; Mihalas et al., 2016). Although TBR2 is not expressed in embryonic thalamus, basally dividing cells do exist that express the proneural basic helix-loop-helix transcription factors NEUROG1 and NEUROG2 (Wang et al., 2011). Since neocortical IPCs also express NEUROG2 (Miyata et al., 2001), we hypothesized that NEUROGENIN-expressing, basally dividing cells in the thalamus are IPCs. In E18.5 Neurog1^{CreERT2} MADM brains, 78% of G/R clones had 4 or fewer cells (Fig.3A) and all labeled cells had a neuronal morphology. Thus, a majority of *Neurog1*-expressing thalamic progenitor cells are intermediate neuronal progenitor cells with a limited proliferative capacity.

In contrast to the small size of *Neurog1* MADM clones, most clones in *Gli1* and *Olig3* MADM brains had more than 10 cells (Fig.3B). Large G/R clones with more than 4 cells in both green and red progeny are likely to be originated from a recombined progenitor cell that produced two radial progenitors, not *Neurog1*-expressing progenitor cells. We defined these clones as symmetric proliferative

(Fig.3C). G/R clones in which either green or red hemi-clone contained 4 or fewer cells were defined as asymmetric self-renewing/neurogenic, which likely produced one radial progenitor and an IPC, or a one radial progenitor and a postmitotic neuron (Fig.3C, "Asym Neuro"). G/R clones with 4 or fewer cells in both hemi-clones were defined as symmetric neurogenic (Fig.3C, "Sym Neuro").

Both *Gli1* and *Olig3* clones exhibited a combination of the three division patterns described in Fig.3C. When recombination was induced in *Gli1*^{CreER} mice at E9.5, ~57% of the clones were of the symmetric proliferative type, indicating that a majority of recombined progenitor cells at E9.5 were neuroepithelial cells undergoing symmetric expansions. In *Olig3*^{CreER} mice, this fraction was ~45% at E10.5 and ~28% at E11.5. In addition, at E11.5, ~61% of the clones were of the symmetric neurogenic type (Fig.3D). The average clone size was smaller for a later Cre activation (22.5 for E10.5 and 9.5 for E11.5). These results indicate a transition of cell division mode in the thalamus from symmetric proliferative to asymmetric to symmetric neurogenic, similar to the neocortex (Gao et al., 2014).

Spatial organization of clonal units reveals principles of thalamic nucleogenesis

To investigate the spatial organization of clonal units during thalamic nucleogenesis, we performed three-dimensional (3D) reconstruction of serial brain sections containing all labeled cells (Fig.4A, Fig.S3A-D) and assigned each labeled cell to a specific thalamic nucleus using a published atlas (Paxinos, 2006), data from our previous study (Nakagawa and O'Leary, 2001) and a newly generated custom

atlas (Fig.S1B). Clones that included cells in the thalamus did not contribute to the prethalamic structures including the reticular nucleus or zona incerta, confirming the separate embryonic origins of the thalamus and the prethalamus. Asymmetric G/R clones generated 11.9 (± 0.87) cells (Fig.4B) that populated 3.3 (± 0.26) nuclei (Fig.4C). This demonstrates that each thalamic radial glial cell produces neurons that populate multiple nuclei after asymmetric divisions. The average number of nuclei populated by progeny of asymmetrically dividing progenitor cells was significantly smaller than for symmetric proliferative ones (Fig.4C), which suggests that the transition from symmetric proliferative to asymmetric division restricts the nuclear fate potential. Within asymmetric clones, "minority" hemi-clones (i.e. G or R hemiclones containing 4 or fewer cells) exhibited significantly more fate restriction than "majority" hemi-clones based on the average number of populated nuclei (Fig.4C). However, some minority hemi-clones still populated more than one nuclei, implying that the fate of thalamic nuclei may not be fully restricted even in terminally dividing cells. The three principal sensory nuclei (ventral posterior; VP, dLG, ventral part of medial geniculate; MGv) are among the earliest-born in the thalamus (Fig.1B). A large proportion of majority hemi-clones was composed of both principal sensory nuclei and other, later-born nuclei, whereas a larger proportion of minority hemiclones was composed exclusively of neurons in principal sensory nuclei (Fig.4D). This suggests that thalamic radial glial cells sequentially generate neurons that populate different nuclei as they undergo asymmetric divisions. This pattern is reminiscent of the sequential generation of deep- and upper-layer neurons from individual radial glial cells in the embryonic neocortex (Gao et al., 2014).

Individual thalamic clones exhibit distinct patterns of nuclear contributions

We next tested if individual radial glial clones contribute to specific subsets of thalamic nuclei. To analyze our results computationally, we generated a data matrix in which rows and columns represented individual G/R clones and thalamic nuclei, respectively. The numeric value in each cell in the matrix was computed (see Materials and Methods) so that it indicated the frequency of progeny in a given clone that was located in each thalamic nucleus. This configuration is analogous to the matrix used for single-cell RNA-seq analysis, in which rows and columns represent individual cells and genes, respectively. We then applied principal component analysis or unsupervised hierarchical clustering analysis for classification of thalamic MADM clones.

Principal component analysis of asymmetric G/R clones identified five clusters that show distinct patterns of nuclear contribution (Fig.5A). Three of the five clusters had major contributions to principal sensory nuclei (green, yellow and red clusters as shown in Fig. 5A). Of these, clones in the green cluster included cells in principal sensory nuclei and medial ventral field (see Methods for its definition). The yellow cluster contained clones that contributed to principal sensory nuclei and other nuclei including ventromedial (VM) and reuniens/rhomboid (Re/Rh). The medial ventral field, VM and Re/Rh are all located medial to principal sensory nuclei, and neurons in these nuclei are born later than those in principal sensory nuclei (Fig.1A). Importantly, in seven out of eight green clones, the minority hemi-clones contributed exclusively to principal sensory nuclei. This result demonstrates that asymmetrically

dividing thalamic radial progenitor cells first generate early-born, principal sensory nuclei and then late-born, medially located nuclei. It also demonstrates a strong lineage relationship among the three principal sensory nuclei, which share many properties including laminar and areal patterns of axon projections to primary sensory areas (Clasca et al., 2012; Jones, 2007), gene expression (Frangeul et al., 2016; Nagalski et al., 2016) and patterns of propagating spontaneous activity during embryogenesis (Moreno-Juan et al., 2017).

The other two clusters included clones that have few cells in principal sensory nuclei (Fig.5A). All the clones in the grey cluster contributed to the ventrolateral (VL) nucleus. The remaining nuclei represented in this cluster are diverse and include centromedian/intermediodorsal (CM/IMD), mediodorsal (MD) and Re/Rh. Many of these nuclei are located more dorsally and caudally within the thalamus compared with principal sensory nuclei, demonstrating that nuclei that are close to each other in physical location are likely to share the same lineage. This result provides a single-cell based evidence for our previous population-based study, in which a gradient of gene expression within the pTH-C domain corresponded to preferential generation of progeny along the rostro-ventral to caudo-dorsal axis of the thalamus (Vue et al., 2007).

The blue cluster contained clones that were relatively more diverse. Here, two clones contained cells in intergeniculate leaflet (IGL) and either MGv or VP. The IGL nucleus receives axonal projections from non-image forming retinal ganglion cells and are known to be derived from the pTH-R domain at the population level (Delogu et al., 2012; Jager et al., 2016; Jeong et al., 2011; Vue et al., 2007). The surprising

co-existence of pTH-C derived cells (in principal sensory nuclei) and pTH-R derived cells (in IGL) in the same clones suggests that during early stages, some progenitor cells might still be uncommitted to either pTH-C or pTH-R fate.

The existence of clone clusters suggests that some thalamic nuclei are more likely to share the cell lineage than other nuclei. In order to assess the "lineage distance" between nuclei, we transposed the data matrix and clustered thalamic nuclei instead of clones. Because a majority of the clones that contributed to lateborn nuclei were of the symmetric proliferative type, we included both asymmetric and symmetric G/R clones in this analysis in order to cluster the entire set of thalamic nuclei. We found that certain groups of nuclei exhibited higher propensity to originate from common progenitor cells at E18.5 (Fig.5B). Examples of such clusters included vLG and IGL (pTH-R derived), VM, VL, centromedian/intermediodorsal (CM/IMD) and Re/Rh (pTH-C derived and located medially to principal sensory nuclei), LP, MD, LD, Po and parafascicular (PF) (pTH-C derived and located dorsally), and VP, dLG, MGv and medial ventral field. The presence of lineage-based clusters of nuclei shown here suggests that the distribution of progeny generated from thalamic progenitors appears to follow distinct nuclei-specific patterns.

Position of radial progenitor cells dictates domain-specific distribution of clones

To determine if the clustering of clones based on cell lineage reflects the distinct positions of the starting progenitor cells within the embryonic thalamus, we

took advantage of the finding that some clones, especially the ones labeled at E9.5 in the *Gli1*^{CreER} line, retained a cell at the ventricular surface with a long radial process spanning the entire thalamic wall.

In order to use the locations of these cells as a proxy for the position of the founder progenitor cell at early embryonic stages, we defined four domains of these radial glial cells within E18.5 thalamus (Fig.6B). "Dorsal-rostral (DR)" corresponded to the most rostral domain in relatively dorsal sections, whereas the "dorsal-middle (DM)" and "dorsal-caudal (DC)" domains were located progressively more caudally. The "ventral (V)" corresponded to the domain in more ventral sections near the basal plate. A hierarchical clustering analysis revealed that radial glial cell-containing clones in the DR and DM domains heavily contributed to principal sensory nuclei (Fig.6A); many of these clones contained cells in both principal sensory nuclei and more medially located non-sensory nuclei including VM, Re/Rh and the medial ventral field. Some DR clones contained cells in both "excitatory nuclei" composed of cortex-projecting neurons and "inhibitory nuclei" containing GABAergic neurons. Thus, DR and DM clones appear to correspond to the green, yellow and red clones in Fig.5A. DC and V clones had a heavy contribution to dorsally located, non-sensory nuclei and showed little contribution to principal sensory nuclei except MGv. These results suggest that thalamic nuclei that are related in ontogeny originate from similar locations within the embryonic thalamus and support the model wherein individual thalamic radial glial cells form a radial column of progeny that spans a defined subset of thalamic nuclei.

Analysis of glia-containing clones reveals rare segregation of neurons and glia across nuclei

Out of the 37 G/R P21 clones obtained from *Gli1*^{CreER} and *Olig3*^{CreER} drivers, 12 clones were composed of both neurons and astrocytes. Clones containing astrocytes were distributed in multiple clusters, suggesting that the ontogeny of thalamic glial cells appears to be largely heterogeneous and that gliogenesis is not restricted to specific thalamic progenitor cell populations (astrocyte-containing clones in asymmetric G/R clones are indicated by an asterisk in Fig.5A).

Clonal lineage tracing of basal progenitor cells shows late specification of thalamic nuclear fate

Analysis of asymmetric clones of *Gli1^{CreER}* and *Olig3^{CreER}* MADM brains revealed minority hemi-clones that included cells in more than one thalamic nuclei (Fig.4C). This suggested that progenitor cells undergoing their last few rounds of divisions are still not yet specified to the fate of a single nucleus. As described above (Fig.4A), a large majority of *Neurog1^{CreER}* MADM clones contained 4 or fewer cells with a neuronal morphology, indicating that *Neurog1*-expressing thalamic progenitor cells are IPCs. Taking further advantage of the MADM system, we analyzed the division patterns and fates of these cells.

At the population level, timed activation of Cre recombinase at E11.5 resulted in expression of the ZSGreen reporter in the entire thalamus except the medial-most region including PV, CM and Re/Rh (Fig.2A, 7A). These medial nuclei start to be labeled by ZSGreen with the Cre activation at E12.5 (Fig.7A). This is one day before

they become robustly labeled in EdU birthdating (Fig.1B). Similarly, VL starts to be labeled by ZSGreen at E11.5 and by EdU at E12.5, indicating that at least for the above thalamic nuclei, progenitor cells express *Neurog1* before they undergo the final division.

We next performed the clonal lineage tracing with MADM. Activation of the Cre recombinase at E11.5, E12.5, E13.5 or E14.5 followed by analysis at E18.5 revealed that a vast majority of MADM clones contained 4 or fewer cells (43 out of 55 G/R clones; Fig.3A,7B). Surprisingly, many of these clones (13 out of 36 G/R clones with 2-4 cells) contained neurons that populated more than one thalamic nuclei (Fig.7C). These multi-nuclei clones were abundant with E11.5 activation, but there were some with the activation at E13.5 (Fig.7D). Four of the multi-nuclei clones were composed of two or all three principal sensory nuclei, while others included late-born nuclei like PV and Re/Rh. By taking advantages of the two-color labeling of the progeny in the MADM system, we also captured 58 terminal divisions for which we could determine whether the last division generated cells in the same nucleus or two different nuclei. Of these 58 final divisions, 13 generated cells in two different nuclei, of which 4 were in dLG and VP and one was in MGv and VP (Fig.7E), consistent with the ontogenic proximity of these nuclei revealed in the analysis of Gli1^{CreER} and Olig3^{CreER} MADM samples (Fig.5B). For individual nuclei, we found that 4 out of 8 terminal divisions that produced a neuron in dLG also produced a neuron in another nucleus (all in VP). Out of 13 terminal divisions that produced a neuron in VP, 7 produced a neuron in another nucleus (4 in dLG, 1 in MGv, 1 in LP, 1 in CL/PC). Thus, dLG and VP nuclei are particularly close to each other in cell lineage and their fates may not diverge at least until the final progenitor cell division. In contrast, only 2 out of 9 terminal divisions that produced a neuron in MGv also produced a neuron in another nucleus (1 each in VP and LP), suggesting that the MGv lineage is largely specified by the time of the final division of *Neurog1*-expressing IPCs.

Taken together, analysis of *Neurog1*^{CreER} clones demonstrated that the thalamus contains intermediate neuronal progenitor cells and that many of these cells are not yet specified for the fate of a specific nucleus. Strikingly, even the last division of such progenitor cells can generate cells in two distinct nuclei, indicating late mechanisms of nuclear fate specification in developing thalamus.

DISCUSSION

In this study, we tracked the lineage of thalamic progenitor cells at the clonal level by using the MADM system with three different inducible Cre drivers. The dual-color labeling of progeny derived from individual progenitors allowed us to examine the temporal progression of progenitor cell differentiation based on division patterns. Our results revealed several cardinal features of thalamic nucleogenesis; 1) contribution to multiple nuclei by individual radial glial cells, 2) temporal progression of nucleogenesis within the lineage, 3) relationship between the position of radial glial cells and the contribution to specific sets of nuclei, and 4) presence of neuronal IPCs and late specification of nuclear fate. We propose that these features are fundamental to the organizing principles of complex brain structures like the thalamus.

Individual radial glial cells in the thalamus produce many neurons that populate multiple nuclei

Thalamic neurogenesis occurs over a short period of time during embryonic development. In mice, neurons in the earliest-born thalamic nuclei, including dLG and MGv, are first generated at around E10.5, while later-born nuclei are born by E13.5 with an overall "outside-in" pattern (Angevine, 1970; Fig.1B). Due to this short time span of thalamic neurogenesis and the spatially complex arrangement of dozens of nuclei, it was previously unknown whether each neural progenitor cell produces a limited number of progeny of a single nuclear identity or sequentially generates a large number of neurons that populate multiple nuclei with different

birthdates. With the MADM approach, we defined clones that originated from an asymmetrically dividing radial glial cell, and found that each asymmetric clone contains an average of 11.9 cells that populated 3.3 thalamic nuclei. This result provides a general principle of thalamic nucleogenesis in which a single radial glial cell produces many neurons that populate multiple nuclei. By distinguishing the differentially colored hemi-clones derived either from the self-renewed radial glia or the differentiated daughter cell, we found that many radial glial cells labeled at E9.5, E10.5 or E11.5 first produced neurons that populate early-born, principal sensory nuclei including dLG, VP and MGv. Then, in many clones, the same lineage produced neurons that occupy more medially located, later-born nuclei such as Re/Rh and VM (Fig.1B). A recent work by Gao et al. (2014) analyzed neurogenesis in the neocortex using the MADM approach and found that each asymmetrically dividing radial glia sequentially produces an average of 8.4 cells that populated multiple layers in an inside-out fashion. This indicates that despite the difference between nuclear versus laminar organizations of these two structures, the thalamus and neocortex might share a basic strategy in cell division and differentiation.

Regional heterogeneity of thalamic progenitor cells in nuclear contributions and neurogenesis

In order to elucidate the heterogeneity among individual cell lineages, we performed principal component analysis of asymmetric MADM clones obtained with $Gli1^{CreER}$ and $Olig3^{CreER}$ drivers. This revealed several clusters of clones each of which contributed to a shared set of thalamic nuclei (Fig.5A). Out of the five clusters

identified, three showed significant contributions to principal sensory nuclei, dLG, VP and MGv. One of these clusters (yellow) was composed of principal sensory nuclei and later-born nuclei that lie medial to principal sensory nuclei, indicating that some radial glia are capable of sequentially generating cells that contribute to different sets of nuclei located along the radial axis. The green cluster included few nuclei other than the principal sensory nuclei. Thus, some radial glial lineages may be depleted once the early-born sensory nuclei are generated. The third such cluster (red) had a strong contribution to MGv, the principal auditory nucleus. These clones contributed to a different set of late-born nuclei including PF and Po from the clones of green and yellow clusters. The remaining two clusters of clones had more contributions to dorso-caudally located nuclei than principal sensory nuclei. Thus, although the principal sensory nuclei generally shared a progenitor cell lineage, the actual pattern of nuclear contribution differed between clones. Clustering of nuclei based on contributions by both symmetric and asymmetric clones revealed additional nuclear groups (Fig.5B), including IGL/vLG, the two nuclei populated by GABAergic neurons, and LP/MD/Po/LD/PF, nuclei located more caudo-dorsally than principal sensory nuclei.

Recently, Shi et al. (2017) reported MADM-based lineage tracing in the postnatal mouse thalamus using the *NestinCreER* driver. They used an unsupervised hierarchical clustering method to cluster both MADM clones and thalamic nuclei based on the distribution of the cells of each clone in different nuclei. The patterns of clone clusters are generally similar to ours (Fig.5A, B). They found the three principal sensory nuclei and the motor nuclei VA/VL are ontogenetically

segregated from three other clusters (MD/Po/LP/CM etc., VM/Re, and AM/AV/PV/LD, etc.). Sherman and colleagues (Sherman et al, 2016 and Sherman and Guillery, 2013) have proposed that the three principal sensory and VA/VL nuclei in the thalamus receive strong subcortical input and relay the information to primary sensory and motor areas ("first-order" nuclei). Therefore, Shi et al. concluded that the first-order and higher-order nuclei, which receive most driver input from the cortex, are ontogenetically segregated. Our study found clusters of nuclei (Fig.5B) that are similar to those detected by Shi et al.. However, there were some differences in the results as well as interpretation. First, our analysis did not find a strong clustering of VL with principal sensory nuclei. Instead, VL more closely segregated with CM, Re and VM (Fig.5A, B). indicating that the ontogenetic relationship of thalamic nuclei does not necessarily follow the distinction between the first-order versus higher-order nuclei. In addition, another first-order nucleus, anterior nuclear complex, particularly the anterodorsal (AD) nucleus, which receives subcortical driver input from the mammillary body (Sherman and Guillery, 2013, Sherman 2016, Petrof and Sherman, 2009), was ontogenetically distant from the principal sensory nuclei both in our study and in Shi et al., (2017). Thus, we propose that the ontogenetic relationship between thalamic nuclei are defined primarily by their rostrocaudal and dorsoventral positions instead of functions. Patterns of gene expression (Frangeul et al., 2016; Nagalski et al., 2016; Nakagawa and O'Leary, 2001; Suzuki-Hirano, 2011), laminar specificity of efferent projections to the cortex (Clasca et al., 2012) and sources and types of afferent projections (Sherman, 2016; Sherman and Guillery, 2013) are likely depend on both cell lineage and postmitotic, and potentially extrinsic, mechanisms.

In addition to simply clustering clones and nuclei, we attempted to correlate the locations of original progenitor cells with their nuclear contributions by analyzing clones that retained radial glial cells at E18.5 (Fig.6). This analysis revealed that the location of the original radial glia predicts the set of nuclei it populates. For example, rostrally located clones (shown as "DR" and "DM" clones in Fig.6B) strongly contributed to principal sensory nuclei, whereas more caudally located, DC clones contributed to caudo-dorsal nuclei including Po, PF, MD and CM/IMD. Interestingly, some of the most rostrally located (DR) clones included both principal sensory nuclei and IGL/vLG, the two nuclei that are mostly populated by inhibitory neurons. This was unexpected because previous population-based lineage studies found that IGL and vLG nuclei originate from a distinct progenitor domain, pTH-R, not pTH-C that contribute principal sensory nuclei (Delogu et al., 2012; Jeong et al., 2011; Vue et al., 2007; Vue et al., 2009). Although we did not determine if the neurons in pTH-Cderived principal sensory nuclei that shared the lineage with those in IGL/vLG are glutamatergic or GABAergic, there are almost no GABAergic neurons in VP and MGv in mice and those in dLG are derived from the midbrain and appear only postnatally (Jager et al., 2016). In addition, we observed this type of clone not only at E18.5, but also at P21, making it unlikely that all the neurons in the pTH-C domains in clones of this cluster at E18.5 were still on route to their final destinations in pTH-R derived nuclei. Thus, it is likely that some clones do contribute to both glutamatergic pTH-C nuclei and GABAergic pTH-R nuclei. One possible mechanism for this is that some early progenitors are still uncommitted for either the pTH-C or pTH-R fate. This is consistent with the fact that the pTH-R domain emerges only after neurogenesis starts in the thalamus (Vue et al., 2009).

The location of the originating radial glia not only had a correlation with the set of nuclei it populates, but also the temporal patterns of cell division. Most of the clones that contained caudo-dorsally located, late-born nuclei such as mediodorsal (MD), centromedian (CM) or paraventricular (PV) nuclei were found to be of the symmetric proliferative type (26 out of 32 clones containing cells in these nuclei were of symmetric type), demonstrating that upon early Cre activation, progenitor cells that produce neurons of these nuclei are still expanding the pool size by symmetric proliferative divisions. Analysis of clones containing a residual radial glia (Fig.6) indicated that cells of MD, CM and PV nuclei are derived from dorso-caudally located progenitor cells. In contrast, progenitor cells in the rostral part contributed to principal sensory nuclei, many of which underwent an asymmetric division immediately after the recombination at E9.5 to E11.5 (28 out of 77 clones containing cells in these nuclei were of asymmetric type). Our results indicate that the temporal pattern of transition from symmetric to asymmetric divisions differs among progenitor domains within the thalamus, where the transition occurs earlier in rostro-ventrally located progenitor cells than in the caudo-dorsal ones and produce a cohort of earlier-born nuclei.

Late specification of nuclear fates in the thalamus

Our previous study showed that similar to the neocortex, the thalamus has a large population of progenitor cells dividing basally away from the surface of the third

ventricle (Wang et al., 2011). However, the division patterns and the fate of these cells had remained unknown. By using the Neurog1^{CreER} driver for clonal lineage analysis of these basal progenitor cells in the thalamus, we found that most of the labeled clones contained four or fewer cells of neuronal morphology. This demonstrates that Neurog1-expressing basal progenitor cells in the thalamus are neuron-generating IPCs. The existence of IPCs fits the model of sequential neurogenesis in the thalamus, in which each radial glial cell divides limited number of times during a short neurogenic period and generates IPCs, and that these IPCs divide once or twice to expand the neuronal output. In the MADM study by Gao et al., (2014), the average size of asymmetric clones in the embryonic neocortex was 8.4 when they were defined as those with 3 or fewer G or R cells. Using the same criteria, the number for the embryonic thalamus in the current study was 11.4, which indicates that the asymmetrically dividing radial glia in the thalamus produce more progeny within a shorter period of neurogenesis than the neocortical counterpart. Interestingly, many of the Neurog1^{CreER} MADM clones populated 2-3 different thalamic nuclei, demonstrating that each round of radial glia division provides a pool of neurons that share similar birthdates and sometimes populate multiple nuclei. This is an efficient strategy to produce a large variety of neuronal types with a limited number of radial glial cells in a short period of time (summary schematics in Fig.S4).

We further took advantage of the dual-color labeling of the progeny of IPCs and determined whether the terminal division of these cells produced neurons in the same or different thalamic nuclei. Although 78% of the final divisions resulted in neurons in the same nucleus, the remaining 22% produced neurons in two different

nuclei. The most frequent pair of these nuclei was VP and dLG, the two principal sensory nuclei that share patterns of gene expression (Frangeul et al., 2016; Gezelius et al., 2016; Nagalski et al., 2016) and functional interactions (Moreno-Juan et al., 2017). Thus, the last division of thalamic IPCs may not be strictly symmetric, producing two neurons that have different nuclear fates. Alternatively, the division is symmetric but postmitotic mechanisms result in diversification of the fate of two daughter neurons. Further studies will be needed to distinguish these possibilities.

Materials and Methods

Animals

Gli1^{CreERT2} (Ahn and Joyner, 2005), Olig3^{CreERT2} (Storm et al., 2009) and Neurog1^{CreERT2} (Koundakjian et al., 2007; Bluske et al., 2012) mutants were found to provide an optimal labeling of thalamic progenitor cells at a clonal level when combined with the MADM-11 system (Hippenmeyer et al., 2010). We bred Gli1^{CreERT2/+}; MADM-11^{GT/GT} mice, Olig3^{CreERT2/+}; MADM-11^{GT/GT} mice or Neurog1^{CreERT2/+}; MADM-11^{GT/GT} mice with MADM-11^{TG/TG} mice. A single dose of tamoxifen (132mg/kg body weight for Gli1^{CreERT2}, 24mg/kg or 34mg/kg for and *Neurog1^{CreERT2}*) was administered to pregnant females (intraperitoneal injection for Gli1CreER^{T2} and oral gavage for Olig3^{CreERT2} and Neurog1^{CreERT2}) at various time points (embryonic day 9.5 (E9.5) for Gli1^{CreERT2}, E10.5 or E11.5 for Olig3^{CreERT2} and E11.5 to E14.5 for Neurog1^{CreERT2}). Embryos or postnatal pups were fixed and frozen. Neurog1^{CreERT2} mice were also bred with Ai6 ZSGreen reporter mice for a population-based lineage tracing with tamoxifen administration at E11.5, E12.5, E13.5 or E14.5 (60mg/kg body weight) and the embryos were perfusion fixed at E18.5. For a birth-dating study, ethynyldeoxyuridine (EdU; Invitrogen or Carbosynth) was injected intraperitoneally (50mg/kg body weight) into pregnant CD1 female mice at E9.5, 10.5, E11.5, E12.5, E13.5 or E14.5 and pups were perfusion fixed at on the day of birth (P0).

Immunostaining, EdU staining, Confocal Imaging, and 3D reconstruction

Serial coronal brain sections (40µm in thickness) through the entire forebrain were

cut by a cryostat or a sliding microtome, and were immunostained with the following antibodies: GFP (1:200 or 500; goat; Rockland), GFP (1:500; chicken; Aves Labs), RFP (1:200 or 1,000; rabbit; Rockland; to detect tdTomato), anti-Nestin (1:500; chicken; Aves Labs), anti-goat/chicken Cy2, anti-rabbit Cy3 and anti-goat Cy5 (1:200 or 500; donkey; Jackson ImmunoResearch). EdU was visualized on cryosections in the detection solution (5uM Sulfo-Cy3 azide (Lumiprobe), 0.1M Tris pH7.5, 4mM copper sulfate, 100mM sodium ascorbate) for 30min after permeabilization in 0.5% Triton-X100 for 30min. Consecutive sections covering individual clones were imaged using Zeiss LSM 710 confocal microscope (Carl Zeiss). For 3D reconstruction, optical stacks from the entire diencephalon were serially aligned along the rostrocaudal axis using Reconstruct 1.1.0 (J.C. Fiala, NIH), followed by import into Imaris (Bitplane) for further analysis.

Axial nomenclature and identification of thalamic nuclei

We adopted the axial nomenclature used in the prosomeric model of forebrain organization ((Puelles et al., 2013; Vue et al., 2007); Fig 1A). In this system, the rostral border of the thalamus is the zona limitans intrathalamica (ZLI), and in the frontal view of cross sections, the bottom part (closer to the ZLI) is ventral-rostral and the top part (closer to the pretectum) is dorsal-caudal. Axial nomenclature of E18.5 sections (e.g., Fig.6B) also follows this rule in order to retain consistency. For identification of thalamic nuclei within tissue sections, we referred to ("Atlas of the Developing Mouse Brain at E17.5, P0 and P6" (Academic Press, 2006) and Nakagawa and O'Leary, (2001). Furthermore, we performed in situ hybridization

(Vue et al., 2007) on E18.5 and P21 sections of wild type brains using the same orientation and thickness as used for the MADM brains, and generated a custom thalamic atlas (Fig.S1B). The mRNA probes for *Gbx2*, *Calb2*, *RORα*, *Foxp2*, *Igsf21*, Slc6a4 and Zyx were used to label thalamic nuclei. At E18.5, Gbx2 was expressed in PV, MD, CL, CM, Re/Rh, Po, LP, VM, MGv and the "medial ventral field" (shown by a white asterisk), which is not labeled in the available atlas (Paxinos, 2006) or book (Jones, 2007). Because this region is marked by ZSGreen in Olig3^{CreERT2}: Ai6 reporter mice at E18.5 (Fig.S2B), we consider it to be thalamus-derived and included it in the analysis; Calb2 was expressed in PV, LD, CL, CM, Re/Rh, Po, LP and medial ventral field; $ROR\alpha$ was expressed in AD, MD, dLG, VP, MGv; Foxp2 was expressed in PV, LD, CM, Re/Rh, MD, Po, LP, PF; Isaf21 was expressed in AD, AV, AM, PV, LD, MD, Po, LP, VP, dLG, MGv, PF; Slc6a4 was expressed in VP, dLG, MGv; Zyx was expressed in AV, PV, LD, MD, VL, VM, CM, Re/Rh, Po, PF. At P21, expression patterns remained essentially unchanged from E18.5 for all markers except $ROR\alpha$, which was expressed in an additional set of nuclei including PT, AD, AV, AM, LD, VL, VM, Po, LP and MGd.

Computational Quantitative Analyses

Each clone was first assigned a vector of values representing its cellular contribution to each nucleus (nucleus X, number of labeled cells in nucleus X). Analogous to analysis of large datasets of gene expression in single-cell RNA sequencing in which each cell has a corresponding vector of values based on relative expression levels for each gene (gene X, expression level of gene X), we performed logarithmic

transformation, followed by principal component analysis and unsupervised hierarchical clustering analysis. For principal component analysis, the clonal distribution data were rescaled such that the average number of cells in each nucleus was centered to zero with principal components as normalized eigenvectors of the covariance matrix of the cellular distributions. The clones were then ordered according to their contribution to the variance in the dataset. For unsupervised hierarchical clustering analysis, we calculated Euclidean distance among clones for plotting dendrograms and heatmaps. Since there were no negative values in the dataset, the Canberra distance was further analyzed for hierarchical clustering with Ward's linkage, which yielded similar results.

Statistical Analyses

Computational analysis of MADM clones was performed using Excel and R softwares. GraphPad Prism ver.5.0 was used for other statistical tests were In Fig.4C, statistical significance was assessed by two-tailed unpaired Student's t-tests. *P < 0.05; **P < 0.01; ***P < 0.001.

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Figure Legends

Figure 1. Outside-in temporal specification of thalamic nucleogenesis

(A) A schematic of the thalamus (TH) in the caudal diencephalon of the mouse embryo. In the sagittal schematic (left two panels), the embryo is facing left. The thalamus is bordered by the pretectum caudally, the habenula dorsally, the basal plate ventrally, and by the zona limitans intrathalamica (ZLI) rostrally. The three dashed lines represent the planes of sections shown in the frontal schematic (right two panels). TH = thalamus, Tel = telencephalon, PTH = prethalamus, Hyp = hypothalamus, ZLI = zonas limitans intrathalamica, HB = habenula, PT = pretectum, M = midbrain, Cx = cortex, RP = roof plate, LV = lateral ventricle, 3V = third ventricle.

(B) Frontal sections of the forebrain showing EdU birthdating in the embryonic mouse thalamus at E18.5 with EdU injections at E10.5, E11.5, E12.5, E13.5 or E14.5. Three representative levels (dorsal, middle and ventral) are shown for each stage of EdU injection. In all three levels, there is a general trend in which neurons of laterally located nuclei are born before neurons of more medially located nuclei, demonstrating the outside-in pattern of neurogenesis. Scale bar: 500μm. Schematic on the right shows a graphical representation of EdU birthdating data.

Figure 2. Clonal lineage tracing of thalamic progenitors using MADM

(A) Sample images of frontal sections from *Gli1^{CreERT2};H2B-GFP*, *Olig3^{CreERT2};ZSGreen* and *Neurog1^{CreERT2};ZSGreen* mouse brains counter-stained with DAPI to show labeling of progenitor cells or their progeny. The left three panels show labeling one day after tamoxifen administration (E10.5 for *Gli1^{CreERT2}*, E11.5 for

Olig3^{CreERT2} and E12.5 for Neurog1^{CreERT2}). Scale bar: 500 µm. **(B)** Bar graphs showing regional distribution of labeled clones in the forebrain of *Gli1^{CreERT2}* and Olig3^{CreERT2} mouse lines crossed with *MADM11* reporter mice one day after tamoxifen administration. Telen, telencephalon; Tha, thalamus; PTh, prethalamus; Hypo, hypothalamus. **(C)** Bar graphs showing frequencies of thalamic clones (green/red or yellow) per hemisphere labeled one day after tamoxifen administration for the *Gli1^{CreERT2};MADM11*, *Olig3^{CreERT2};MADM11* and *Neurog1^{CreERT2};MADM11* mice. More than 70% of hemispheres had no recombined clones. The curves represent the best non-linear fit.

Figure 3. Temporal change of cell divisions during thalamic nucleogenesis

(A) Bar graph showing a breakdown of the cell number per clone for all Neurog1^{CreERT2} labeled clones. (B) Top: Bar graphs showing breakdown of the cell number per clone for all Gli1^{CreERT2} and Olig3^{CreERT2} labeled clones. Bottom: Bar graphs showing breakdown of the cell number per clone for all Gli1^{CreERT2} and Olig3^{CreERT2} labeled green/red hemi-clones. (C) A schematic depicting the three defined types of clonal division patterns: symmetric proliferative, asymmetric neurogenic and other. (D) Bar graph showing the % of division pattern within clones labelled using Gli1 and Olig3 driver lines with tamoxifen induction at E9.5 for Gli1 and E10.5/E11.5 for Olig3.

Figure 4. Cell division patterns reveal the fate specification of early-born neurons in thalamus

(A) Brains were reconstructed from serial sections cumulatively containing all labelled cells within a single clone using IMARIS. Shown is a sample representative 2D projection (left panel) and 3D reconstruction (right panel) of thalamic clone derived from *Gli1*^{CreERT2} MADM E18.5 brain. Violescent stereostructure sketches the contours of the hemisphere, the green filament shows the radial process of RGC, and the green and red spots represent labelled GFP+ and tdTomato+ cells, respectively. Scale bar: 300μm. (B) Bar graph showing a summary of the average cell number within asymmetric clones. (C) Left: bar graph showing the average number of nuclei populated in symmetric and asymmetric G/R clones. Symmetric proliferative clones contributed to more thalamic nuclei than asymmetric neurogenic clones. Right: bar graph showing the average number of nuclei populated in majority ("major") and minority ("minor") hemi-clones within asymmetric clones. (D) Bar graph showing the percentage of clones (left panel) and number (right panel) of cells populating principal sensory nuclei (PSN) in majority ("major") and minority ("minor") hemi-clones within asymmetric clones.

Figure 5. Spatial specification of thalamic nucleogenesis

(A) Principal component analysis of asymmetric clones and summary of nuclei populated in each cluster. Asterisks label clones that contain both neurons and astrocytes. (B) Unsupervised hierarchical clustering analysis of all types of G/R clones. The heatmap shows the distance between different thalamic nuclei.

Figure 6. RGC position dictates domain-specific distribution of clones

(A) Clustering analysis of clones based on the position of radial glial cells (RGC). (B) The four micrographs show frontal sections of E18.5 forebrain, in which colored solid bars indicate the four zones, DR (dorsal rostral), DM (dorsal middle), DC (dorsal caudal) and V (ventral), of GC positions. Scale bar: 500µm. Bottom: four faying surface plots illustrate representative shapes of the respective clones.

Figure 7. Clonal lineage tracing of basal progenitor cells shows late nuclear specification of thalamic cells

(A) Left: sample images of frontal sections from E18.5 *Neurog1*^{CreERT2}; *ZSGreen* mouse brains to show labeling of progeny of *Neurog1*-expressing progenitor cells when tamoxifen is administered at E11.5, E12.5 or E13.5. Sections at three different dorsoventral levels are shown for each stage of tamoxifen administration. (B) Sample image of a *Neurog1*^{CreERT2} clone at E18.5. The colonization of two nuclei within a single clone is shown. Scale bar = 300μm. (C) Bar graph showing the number of nuclei populated for *Neurog1*^{CreERT2} labeled clones with 2 to 4 cells. (D) Bar graph showing the percentage of clones populating single or multiple nuclei for *Neurog1*-labeled clones with 2-4 cells. The left bar indicates clones that are activated at E11.5 or E12.5, and the right bar indicates clones activated at E13.5 or E14.5. (E) Pie chart showing the percentage of clones populating single or multiple nuclei for clones undergoing terminal divisions. The breakdown of nuclei populated for clones colonizing multiple nuclei is shown.

Figure S1

(A) In the MADM system, inducible Cre mice are used to drive interchromosomal recombination of the gene locus that contains partial coding sequences of EGFP and tdTomato (MADM-11GT and MADM-11TG). The recombination causes the formation of fluorescent EGFP or tdTomato proteins. When the recombination occurs in a progenitor cell during G2 phase of the cell cycle, the two daughter cells express different combinations of fluorescent proteins depending on the mode of segregation of sister chromatids. Upon "G2-X" type segregation, one daughter cell expresses EGFP (green or "G" cell) and the other expresses tdTomato (red or "R" cell). If these daughter cells undergo further divisions, respective fluorescent proteins continue to be expressed in the progeny, generating a "G/R clone" that contains a mixture of green and red cells. The other, "G2-Z" type segregation produces one daughter cell that expresses both EGFP and tdTomato (yellow or "Y" cell) and the other daughter cell that expresses neither of the proteins. Thus, the G2-Z segregation generates a "Y clone", which allows us to trace only half of the progeny of the recombined progenitor cell. (B) A custom atlas generated by in situ hybridization to define individual thalamic nuclei at E18.5 (upper panel) and P21 (lower panel). Expression of seven representative markers is shown. This is a consecutive set of 40µm-thick frontal sections. The left column is most dorsal and the right column is the most ventral (see Fig.1A for axial orientation within the thalamus). Scale bar: 1mm.

Figure S2

(A) Immunostaining of an E10.5 section of Gli1^{CreERT2} labeled clones with anti-Nestin

antibody. Tamoxifen was activated at E9.5. (C) An image of frontal section from

Olig3^{CreERT2};ZSGreen brain to show labeling of medial ventral field (asterisk) by

ZSGreen. Tamoxifen was administered at E9.5.

Figure S3

(A-B) Sample serial confocal images from thalamic clones showing the labeling of

multiple thalamic nuclei within individual clones. (more explanations) Scale bar: 200

μm. (C) 2D projection and 3D reconstruction of a thalamic clone without a retained

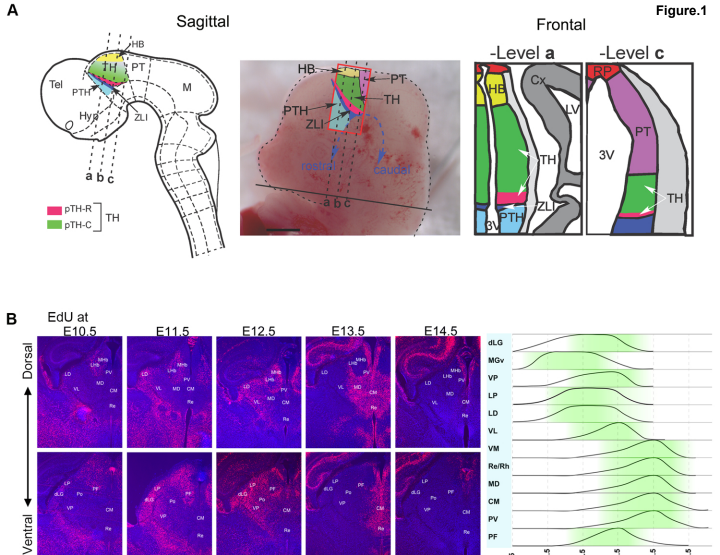
RGC is shown.

Figure S4

A schematic summary of the current study showing the main principles underlying

spatiotemporal regulation of thalamic progenitor specification at (A) E10.5 and (B)

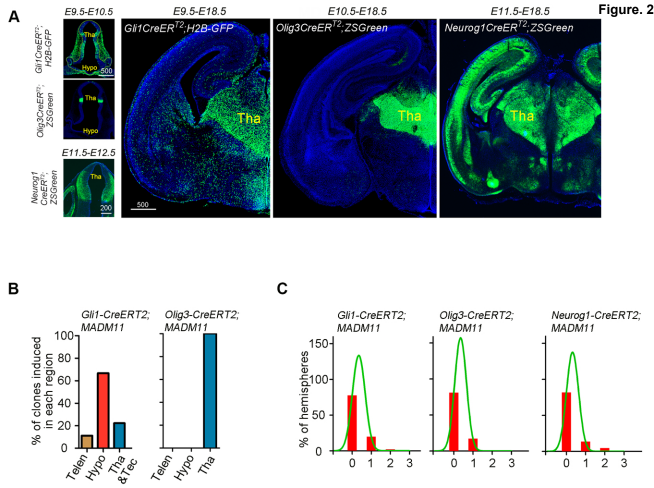
E18.5.

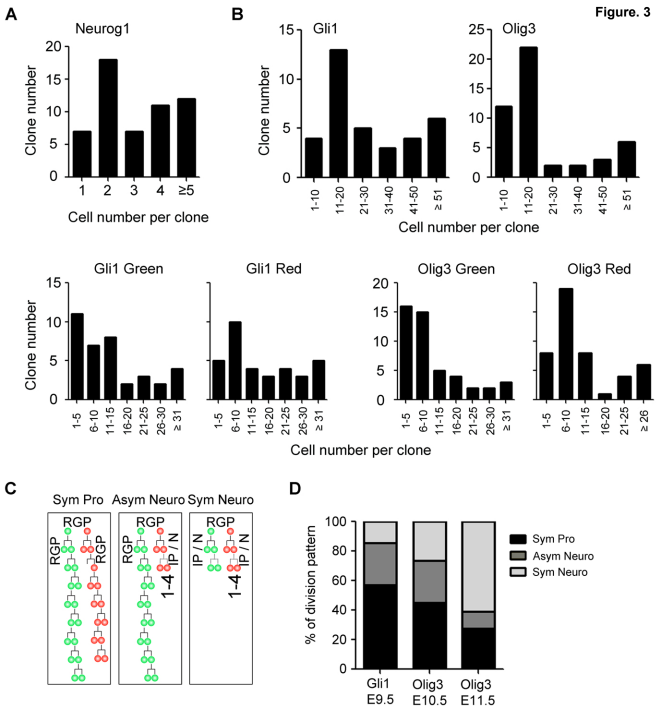


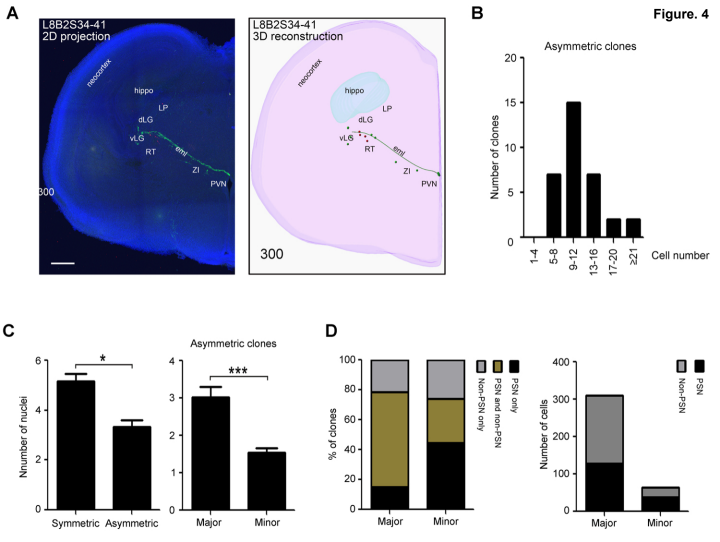
E12.5

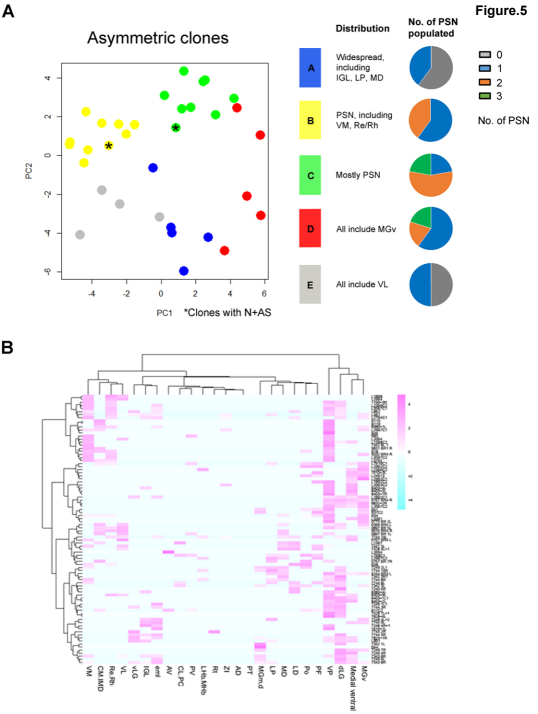
E11.5

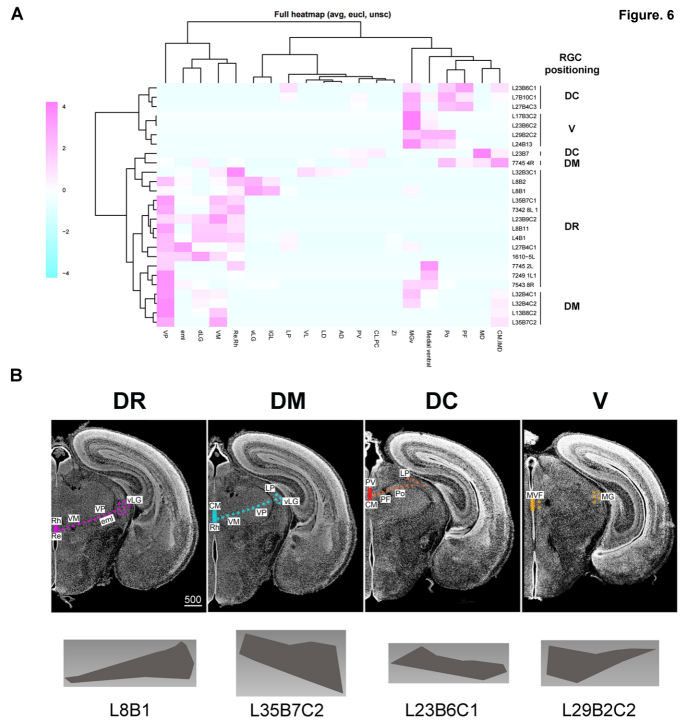
E13.5

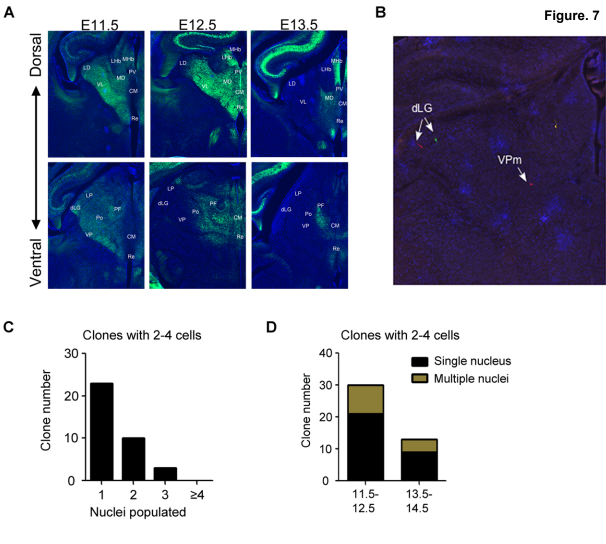


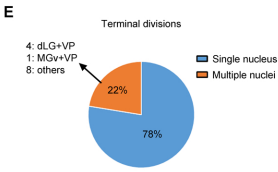


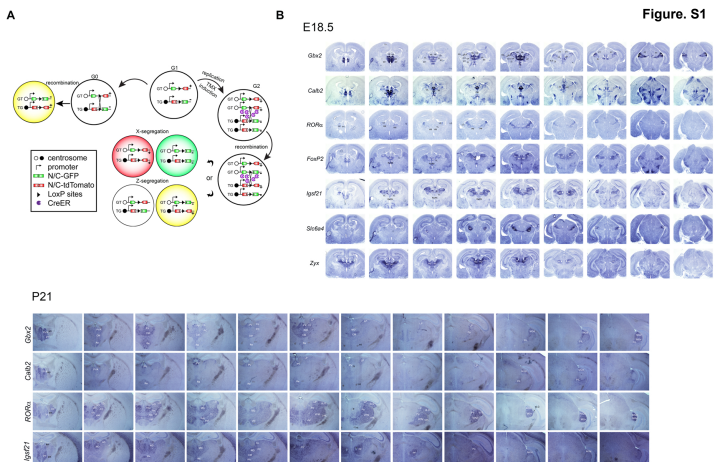












Zyx

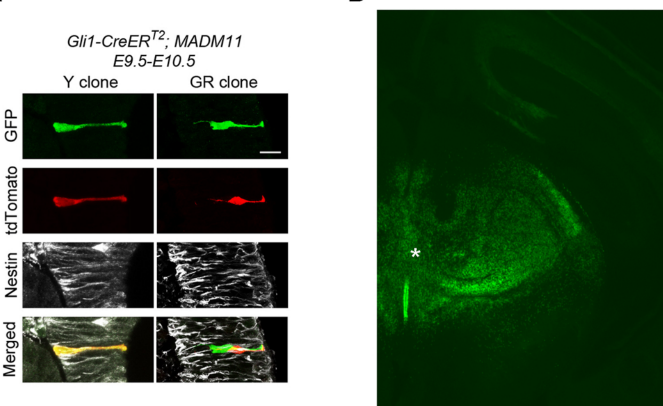
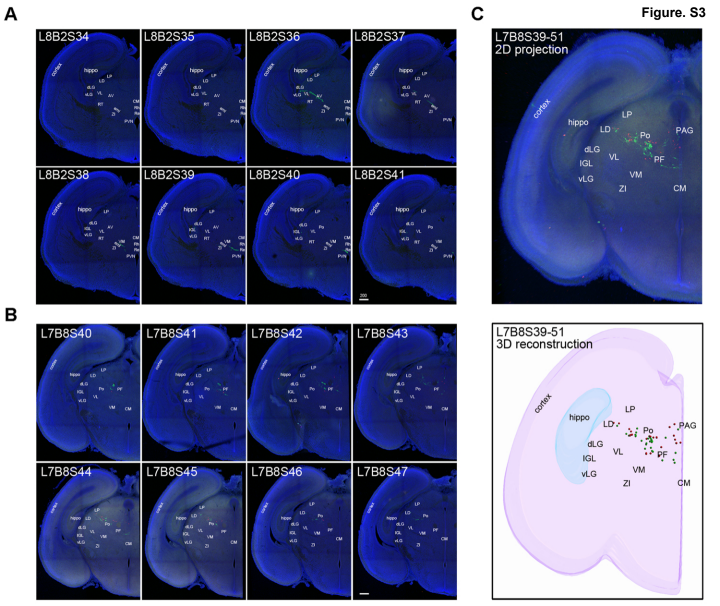
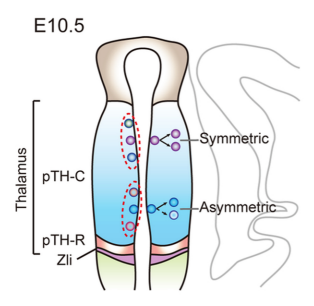


Figure. S2







В

E18.5